Se2017

Program and Abstract Book
Conference venue

The Berzelius and Scheele laboratories, with lecture halls **Samuelsson**, **Vesalius** and **Retzius**, as well as **Poster and Exhibition room**.
Welcome!

The Se2017 Conference celebrates 200 Years of Selenium Research and is held at Karolinska Institutet in Stockholm, Sweden, on August 13-17, 2017. The conference covers all major fields of current selenium research, with a special focus on biology, medicine, biomedicine and the environment. The event is officially composed of two serial symposia; for the first time held in parallel at the same site and having overlapping plenary sessions, poster sessions and social events. The two symposia are The 11th International Symposium on Selenium in Biology and Medicine and The 5th International Conference on Selenium in the Environment and Human Health.

Please consult the conference website for further information and last-minute changes: www.Se2017.se

Welcome to Se2017!

Elias Arnér, MD PhD
Chair, Se2017
Acknowledgements

Co-organizer

Gold sponsors
Silver sponsors

Additional contributions

Cancerfonden

Vetenskapsrådet

Stockholms stad

Si. Swedish Institute

nutrients

The Packer/Wentz Endowment

The Berzelius Society
Organizing committees

Chair:
Elias Arnér, Karolinska Institutet, Sweden

Executive Committee:
Elias Arnér, Karolinska Institutet, Sweden
Gary Bañuelos, USDA, USA
Arne Holmgren, Karolinska Institutet, Sweden
Zhi-Qing Lin, Southern Illinois University, Illinois, USA

Local Organizing Committee, Karolinska Institutet, Sweden:
Elias Arnér
Arne Holmgren
Aristi Fernandes
Jan Trofast

Co-Chair:
Xuebin Yin, University of Science and Technology of China, China

International Scientific Advisory Committee:
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John Hesketh, University of Newcastle, United Kingdom
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Michael Rother, Technical University of Dresden, Germany
Lutz Schömburg, Charité-Universitätsmedizin Berlin, Germany
Ulrich Schweizer, University of Bonn, Germany
Dieter Söll, Yale, USA
Roger Sunde, University of Madison, USA
Anatoly Skalny, University of Orenburg, Russia
Instructions for speakers and chairs

The allotted time for all speakers is including questions. All speakers are kindly asked to plan accordingly and Chairs are kindly asked to be stringent in keeping the time frames in their sessions. For presentations and short biographies of invited speakers and Chairs, see http://se2017.se/confirmed-speakers-and-chairs/.

Oral presentations – format and technical restrictions
The presentation should be prepared as a PowerPoint presentation in a 16:9 format (due to the format of the screen).

We will only use PC computers (no Mac!), Windows and PowerPoint software, so please provide file(s) compatible with this setup. We recommend that you also bring a PDF file of your presentation. Please do not bring your own computer!

If any videos will be included in your presentation, please hand these in as separate files when you hand in your presentation. Do not include them in the PowerPoint presentation.

We kindly ask you to hand in your presentation no later than the break before your session and to be present in the lecture hall at a minimum of 10 minutes before your lecture. If your lecture is in the first session of the day, please provide your presentation no later than 30 minutes before the session begins.

All presentations must be given in English.

Instructions for posters

Posters should be displayed in portrait (=standing) orientation. The poster board area is 110 cm wide and 200 cm high. The material on the poster board is felt. Pins will be provided. Each poster has been given a number and should be fixed on the board marked with the same number.

Poster mounting and dismounting
All posters shall be on display throughout the whole meeting. You will be able to put up your poster during the following hours:
Sunday 13 August at 15:30 to 17:00.
Monday 14 August from 08:00.
Please note that your poster must be mounted before 10.30 on Monday 14 August.

We kindly ask you to take down your poster on Wednesday, August 16 after the last coffee break, i.e. 16.30. If you wish, you may leave your poster to the person in charge of the poster exhibition on Wednesday and retrieve it on Thursday in Aula Medica.

Language
The poster should be prepared in the English language.
Practical information

Conference office:
Academic Conferences
Universities in cooperation: Karolinska Institutet, Swedish University of Agricultural Sciences and Uppsala University
Phone: +46 18 67 10 03
E-mail: info@se2017.se for questions.

Official language
The language of the conference is English.

Conference venue
The Se2017 conference will be held at Karolinska Institutet Campus Solna, in Aula Medica, in two lecture halls of the Berzelius Laboratory, Gustaf Retzius and Andreas Vesalius, and in one lecture hall of the Scheele laboratory, Samuelsson.

Addresses

<table>
<thead>
<tr>
<th>Aula Medica</th>
<th>Berzelius Laboratory</th>
<th>Scheele Laboratory</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nobels väg 6</td>
<td>Berzelius väg 3</td>
<td>Tomtebodavägen 6</td>
</tr>
<tr>
<td>171 65 Solna</td>
<td>171 65 Solna</td>
<td>171 65 Solna</td>
</tr>
</tbody>
</table>

Badges
The participant name badge will be provided at the registration desk. All participants are requested to wear the badge throughout the conference. Only badge holders will be admitted to the sessions. On the badge you will also see what social events you have registered for.

Meals
Lunches and refreshments are included in the registration fee.

Time zone
Sweden is in the Central European Time zone.

Business hours & shopping
Shops are typically open between 10.00 and 18.00 hrs on weekdays and from 10.00 to 15.00 on Saturdays. Shops in the city center have extended opening hours, some even on Sundays between 12.00 and 16.00 hrs.

Transportation between Arlanda Airport and Stockholm
A high-speed train called Arlanda Express runs non-stop between Stockholm and Arlanda in 20 minutes. If you are travelling from Stockholm Central Station to Arlanda Airport, you can buy your ticket at the train station’s information desks or using the Arlanda Express self-service machines. Price: Adult one way 280 SEK, round trip 540 SEK (Prices as per June 2017).

Smoking policy
Sweden has a non-smoking policy, i.e. smoking is prohibited in public buildings, public transport, taxis, buses and trains.

Tourist information
Stockholm Visitor Center
Kulturhuset, Sergels Torg 3
103 27 Stockholm
E-mail: touristinfo@stockholm.se
Tel: 08-508 28 508

Public transportation
Stockholm has a well-developed public transport system. For more information please visit www.sl.se/en.
Transportation to the venue

Metro
From the metro station T-Centralen, located below Stockholm Central Station, you can take train 17, 18, and 19 to St. Eriksplan. Change to bus no. 3 (towards Karolinska sjukhuset), 73 (towards Karolinska Institutet), 77 (towards Karolinska sjukhuset), or take a 10-minute walk to the venue.

Bus
Bus lines 3 (towards Karolinska sjukhuset), 67 (towards Frösundavik), 73 (towards Karolinska Institutet) and 77 (towards Karolinska sjukhuset) run between central Stockholm and the east side of Karolinska Institutet, close to Aula Medica. The bus stop closest to the venue is called Karolinska Institutet Östra.

Bus line 69 runs between central Stockholm and the west side of Karolinska Institutet, close to the Berzelius and Scheele Laboratories. You can get on bus 69 by the Central Station or at Norra Bantorget and you will get off at bus stop Karolinska Institutet Västra.

Taxi
The organising committee recommend the following taxi companies:
Taxi Stockholm (+46 8-15 00 00)
Taxi Kurir (+46 771-860000)
Sverigetaxi (+46 20-20 20 20)

WiFi at the venue
Username: KI-Guest
Password: Stockholm17

Force majeure
The organisers are not liable for any claims for damages and/or losses if the entire conference has to be cancelled due to a force majeure incident.

Disclaimer
The organisers are not liable for damages and/or losses of any kind which may be incurred by the conference delegates or by any other individuals accompanying them, both during the official activities as well as going to/from the conference. Delegates are responsible for their own safety and belongings.
## Se2017 – 200 Years of Selenium Research

### Program at a Glance

#### Sunday 13 August 2017
- 15:30 – 17:00: Registration
- 17:00 – 20:00: Inaugurating Plenary Session
- 20:00 – ca. 22: Get-together

#### Monday 14 August 2017

<table>
<thead>
<tr>
<th>Time</th>
<th>Samuelsson</th>
<th>Vesalius</th>
<th>Retzius</th>
</tr>
</thead>
<tbody>
<tr>
<td>08:00 – 09:00</td>
<td></td>
<td>Time to mount posters in the poster and exhibition room</td>
<td></td>
</tr>
<tr>
<td>09:00 – 10:30</td>
<td>1.1. Inorganic selenium chemistry</td>
<td>2.1. Metabolism of selenium in living cells</td>
<td>3.1. Selenium supplementation for animal and livestock health</td>
</tr>
<tr>
<td>10:30 – 11:00</td>
<td></td>
<td>Coffee break</td>
<td></td>
</tr>
<tr>
<td>11:00 – 12:30</td>
<td>1.3. Local geological selenium sources and global cycling</td>
<td>2.2. Molecular mechanisms of selenium toxicity</td>
<td>3.2. Epidemiology of selenium related health and disease</td>
</tr>
<tr>
<td>12:30 – 14:00</td>
<td>1.4. Relationships of selenium between soils, water, and vegetation (Session I)</td>
<td>Poster Session with lunch and exhibition</td>
<td></td>
</tr>
<tr>
<td>14:00 – 16:00</td>
<td>1.4. Relationships of selenium between soils, water, and vegetation (Session II)</td>
<td>2.4. Selenoprotein synthesis pathways (Session I)</td>
<td>3.3. Nutritional selenium intervention studies in human</td>
</tr>
<tr>
<td>16:00 – 16:30</td>
<td></td>
<td>Coffee break</td>
<td></td>
</tr>
<tr>
<td>16:30 – 19:00</td>
<td>1.5. Excessive selenium accumulation from natural or anthropogenic sources and remediation technologies</td>
<td>2.4. Selenoprotein synthesis pathways (Session II)</td>
<td>3.4. Selenium based medical therapeutics (Sessions II and III)</td>
</tr>
<tr>
<td>19:00 – ca. 21</td>
<td></td>
<td>Poster session with dinner and exhibition</td>
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</tr>
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</table>
## Tuesday 15 August 2017

<table>
<thead>
<tr>
<th>Time</th>
<th>Samuelsson</th>
<th>Vesalius</th>
<th>Retzius</th>
</tr>
</thead>
<tbody>
<tr>
<td>09:00 - 10:30</td>
<td>1.6. Strategies to improve selenium accumulation and biofortification (Session I)</td>
<td>2.6. Selenoprotein function (Session I)</td>
<td>3.5. Selenium metabolism or selenoprotein function in health and disease (Session I)</td>
</tr>
<tr>
<td>10:30 – 11:00</td>
<td></td>
<td>Coffee break</td>
<td></td>
</tr>
<tr>
<td>11:00 – 12:30</td>
<td>1.6. Strategies to improve selenium accumulation and biofortification (Session II)</td>
<td>2.6. Selenoprotein function (Session II)</td>
<td>3.5. Selenium metabolism or selenoprotein function in health and disease (Session II)</td>
</tr>
<tr>
<td>12:30 – 14:00</td>
<td></td>
<td>Post Session with lunch and exhibition</td>
<td></td>
</tr>
<tr>
<td>14:00 – 16:00</td>
<td></td>
<td>2.7. The systems biology of selenium and selenoproteins</td>
<td>3.5. Selenium metabolism or selenoprotein function in health and disease (Session III)</td>
</tr>
<tr>
<td>16:30 – ca 00.00</td>
<td></td>
<td>Excursion to Mariefred (pre-booked participants only)</td>
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</table>

Coaches leave from outside the Berzelius and Scheele laboratories at 16.30.
### Se2017 – 200 Years of Selenium Research

#### Program at a Glance

**Wednesday 16 August 2017**

<table>
<thead>
<tr>
<th>Time</th>
<th>Vesalius</th>
<th>Retzius</th>
</tr>
</thead>
<tbody>
<tr>
<td>09:00 - 11:00</td>
<td>3.5. Selenium metabolism or selenoprotein function in health and disease (Session IV)</td>
<td></td>
</tr>
<tr>
<td>11:00 – 11:30</td>
<td>Coffee break</td>
<td></td>
</tr>
</tbody>
</table>
| 11:30 – 13:30 | 2.8. Selenium based biotechnological applications  
2.9. Additional and emerging topics of selenium in molecular life sciences  
3.6. Clinical genetics of selenium or selenoprotein-encoding genes | 3.7. Additional and emerging topics of selenium in health or disease |
| 13:30 – 15:00 | Lunch in poster and exhibition room                                      |                                                                         |
| 15:00 – 18:00 | Planning for *The 12th International Symposium on Selenium in Biology and Medicine* and Information on *The 6th International Conference on Selenium in the Environment and Human Health*  
Introducing *International Society for Selenium Research* (ISSR)  
Natural Biofortification Program (NBP) satellite meeting | Elsevier Workshop  
*How to Write a Great Research Paper, and Get it Accepted by a Good Journal* |
| 18:30       | Coaches leave from outside the Berzelius and Scheele laboratories for transport to the Stockholm City Hall |                                                                         |
| 19:00 – ca 20:30 | **Reception in the Stockholm City Hall**                               |                                                                         |

**Thursday 17 August 2017**

<table>
<thead>
<tr>
<th>Time</th>
<th>Aula Medica</th>
</tr>
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<tbody>
<tr>
<td>8:30 – 10:30</td>
<td>Plenary Session I</td>
</tr>
<tr>
<td>10:30 – 11:00</td>
<td>Coffee break</td>
</tr>
<tr>
<td>11:00 – 12:30</td>
<td>Plenary Session II</td>
</tr>
<tr>
<td>12:30 – 13:00</td>
<td>Closing Session and Awards</td>
</tr>
</tbody>
</table>
## Sunday 13 August 2017

**Aula Medica**

<table>
<thead>
<tr>
<th>Time</th>
<th>Event</th>
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<tbody>
<tr>
<td>15.30 – 17.00</td>
<td>Registration</td>
</tr>
<tr>
<td>17:00 – 20:00</td>
<td><strong>Inaugurating Plenary Session, Aula Medica</strong></td>
</tr>
<tr>
<td></td>
<td>Chair: Elias Arnér</td>
</tr>
<tr>
<td>17:00 – 17:10</td>
<td>Welcome address</td>
</tr>
<tr>
<td></td>
<td>Elias Arnér and Gary Bañuelos</td>
</tr>
<tr>
<td>17:10 – 17:30</td>
<td>Berzelius and his discovery of Selenium</td>
</tr>
<tr>
<td></td>
<td>Jan Trofast</td>
</tr>
<tr>
<td>17:30 – 18:00</td>
<td>O1 - The global cycle of selenium</td>
</tr>
<tr>
<td></td>
<td>Lenny Winkel</td>
</tr>
<tr>
<td>18:00 – 18:30</td>
<td>O2 - Selenium metabolism in plants</td>
</tr>
<tr>
<td></td>
<td>Philip John White</td>
</tr>
<tr>
<td>18:30 – 19:00</td>
<td><strong>Thressa C. Stadtman Memorial Lecture</strong></td>
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<tr>
<td></td>
<td>O3 - Selenium utilization in diverse animals</td>
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<td></td>
<td>Vadim N. Gladyshev</td>
</tr>
<tr>
<td>19:00 – 19:30</td>
<td>O4 - Phenotypes and molecular pathogenesis of disorders of human selenoprotein synthesis</td>
</tr>
<tr>
<td></td>
<td>Krishna Chatterjee</td>
</tr>
<tr>
<td>19:30 – 20:00</td>
<td>Performance – “Probing the mind of Berzelius”</td>
</tr>
<tr>
<td></td>
<td>1+1=3 (Stephen Whitmarsh and Jean-Louis Huhta)</td>
</tr>
<tr>
<td>20.00 – ca. 22</td>
<td>Get-together</td>
</tr>
</tbody>
</table>
### Monday 14 August 2017

#### 08:00 – 09:00

*Time to mount posters in the poster and exhibition room*

<table>
<thead>
<tr>
<th>Samuelsson</th>
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</tr>
</thead>
<tbody>
<tr>
<td><strong>09:00 - 09:20</strong></td>
<td><strong>1.1. Inorganic selenium chemistry,</strong> <em>Chair: Ingrid Pickering</em></td>
<td><strong>2.1. Metabolism of selenium in living cells,</strong> <em>Chair: Aristi Fernandes</em></td>
</tr>
<tr>
<td><strong>09:20 - 10:30</strong></td>
<td><strong>O5 - Inorganic selenium species separation and preconcentration with a new nanosilica-ionic liquid hybrid,</strong> <em>Mauricio Llaver</em></td>
<td><strong>O9 - Selenium requirements and upper limits in mammals and avians from enzyme and molecular biomarkers,</strong> <em>Roger A Sunde</em></td>
</tr>
<tr>
<td><strong>09:00 - 09:20</strong></td>
<td><strong>1.2. Imaging of selenium and analytical methodologies,</strong> <em>Chair: Ingrid Pickering</em></td>
<td><strong>O15 - New opportunities for selenium speciation from advanced X-ray spectroscopy,</strong> <em>Graham George</em></td>
</tr>
<tr>
<td><strong>09:20 - 10:30</strong></td>
<td><strong>O6 - A novel operon involved in selenite reduction in Geobacter sulfurreducens,</strong> <em>Hisaaki Mihara</em></td>
<td><strong>O9 - Selenium requirements and upper limits in mammals and avians from enzyme and molecular biomarkers,</strong> <em>Roger A Sunde</em></td>
</tr>
<tr>
<td><strong>09:00 - 09:50</strong></td>
<td><strong>O16 - Development of fluorescent ROS probe based on redox reaction of selenol group,</strong> <em>Noriyuki Suzuki</em></td>
<td><strong>O17 - Advances in large scale speciation of selenium by plasma and molecular mass spectrometry,</strong> <em>Joanna Szpunar</em></td>
</tr>
<tr>
<td><strong>09:50 - 10:30</strong></td>
<td><strong>O7 - Regulation of selenium metabolism in Archaea,</strong> <em>Michael Rother</em></td>
<td><strong>O10 - Effects of feeding cows Se-yeast or Se-enriched alfalfa hay on baby calves Se status and IgG titers,</strong> <em>Jean Hall</em></td>
</tr>
<tr>
<td><strong>10:10 - 10:30</strong></td>
<td><strong>O8 - Anaerobic reduction of Se(IV) by bacterial strain isolated from Spanish bentonites: multidisciplinary approach characterization,</strong> <em>Mohamed Larbi Merroun</em></td>
<td><strong>O11 - Thirty years of selenium fertilization: A Finnish solution to ensure an adequate selenium intake,</strong> <em>Merja Eurola</em></td>
</tr>
<tr>
<td><strong>10:30 - 11:00</strong></td>
<td><strong>O17 - Advances in large scale speciation of selenium by plasma and molecular mass spectrometry,</strong> <em>Joanna Szpunar</em></td>
<td><strong>O12 - Selenium bioaccessibility in biofortified crops and food supplements: a factor to take into account?</strong> <em>Gijis Du Laing</em></td>
</tr>
<tr>
<td><strong>10:00 - 10:15</strong></td>
<td><strong>O18 - Selenium requirements of the turkey based on enzymatic biomarkers and next-generation sequencing,</strong> <em>Rachel M Taylor</em></td>
<td><strong>O13 - Selenium requirements of the turkey based on enzymatic biomarkers and next-generation sequencing,</strong> <em>Rachel M Taylor</em></td>
</tr>
<tr>
<td><strong>10:15 - 10:30</strong></td>
<td><strong>O14 - Selenium supplementation of feed for Atlantic salmon,</strong> <em>Heidi Amlund</em></td>
<td><strong>O14 - Selenium supplementation of feed for Atlantic salmon,</strong> <em>Heidi Amlund</em></td>
</tr>
</tbody>
</table>
## Monday 14 August 2017

### Samuelsson

<table>
<thead>
<tr>
<th>Time</th>
<th>Title</th>
<th>Speaker</th>
</tr>
</thead>
<tbody>
<tr>
<td>11:00</td>
<td>1.3. Local geological selenium sources and global cycling</td>
<td>Chair: Lenny Winkel</td>
</tr>
<tr>
<td>11:30</td>
<td>O18 - Selenium speciation in rainwater from high altitude locations</td>
<td>Elke Suess</td>
</tr>
<tr>
<td>11:15</td>
<td>O19 - Effect of conservation agriculture on solubility and fractionation of soil selenium</td>
<td>Ivy Sichinga - Ligowe</td>
</tr>
<tr>
<td>11:30</td>
<td>1.4. Relationships of selenium between soils, water, and vegetation (Session I)</td>
<td>Chair: Zhi-Qing Lin</td>
</tr>
<tr>
<td>11:45</td>
<td>O27 - Key role of selenium (0) nanoparticles in soil and as therapeutic agent</td>
<td>Laurent Charlet</td>
</tr>
<tr>
<td>12:00</td>
<td>O28 - Adsorption of selenate and selenide in cultivated and native soils of the Brazilian Cerrado</td>
<td>Guilherme Lopes</td>
</tr>
<tr>
<td>12:15</td>
<td>O29 - Selenium and barium in brazil nuts: unravelling the spatial distribution</td>
<td>Luiz Roberto Guimarães Guilherme</td>
</tr>
</tbody>
</table>

### Vesalius

<table>
<thead>
<tr>
<th>Time</th>
<th>Title</th>
<th>Speaker</th>
</tr>
</thead>
<tbody>
<tr>
<td>11:00</td>
<td>2.2. Molecular mechanisms of selenium toxicity</td>
<td>Chair: Hugh H Harris</td>
</tr>
<tr>
<td>12:00</td>
<td>O20 - Selenocysteine in mammalian thioredoxin reductase and application of eselen as a therapeutic</td>
<td>Arne Holmgren</td>
</tr>
<tr>
<td>11:30</td>
<td>O21 - Characterization of novel selenium compounds as a therapeutic approach for improved cancer treatment</td>
<td>Aristi Fernandes</td>
</tr>
<tr>
<td>11:50</td>
<td>O22 - Impairment of protein homeostasis accounts for selenomethionine toxicity in Saccharomyces cerevisiae</td>
<td>Myriam Lazard</td>
</tr>
<tr>
<td>12:10</td>
<td>2.3. Molecular consequences of selenium deficiency</td>
<td>Chair: Hugh H Harris</td>
</tr>
<tr>
<td>12:30</td>
<td>O30 - Role of the selenium in articular cartilage metabolism, growth, and maturation</td>
<td>Caroline Bissardon</td>
</tr>
</tbody>
</table>

### Retzius

<table>
<thead>
<tr>
<th>Time</th>
<th>Title</th>
<th>Speaker</th>
</tr>
</thead>
<tbody>
<tr>
<td>11:00</td>
<td>3.2. Epidemiology of selenium related health and disease</td>
<td>Chair: Roger Sunde</td>
</tr>
<tr>
<td>11:30</td>
<td>O23 - Selenium status of Russian population: interactions with demography</td>
<td>Anatoly Skalny</td>
</tr>
<tr>
<td>11:30</td>
<td>O24 - Selenium status of school children for Kaschin–Beck disease endemic areas in Tibet, China</td>
<td>Linsheng Yang</td>
</tr>
<tr>
<td>12:30</td>
<td>O25 - Impact of early-life selenium status on children’s cognitive abilities</td>
<td>Helena Skröder</td>
</tr>
</tbody>
</table>

### Poster Session with lunch and exhibition

12.30 – 14.00
### Monday 14 August 2017

#### Vertebrai

<table>
<thead>
<tr>
<th>Time</th>
<th>Session</th>
<th>Chair/Presenter</th>
</tr>
</thead>
<tbody>
<tr>
<td>14:00</td>
<td>1.4. Relationships of selenium</td>
<td>Milton Ferreira Moraes</td>
</tr>
<tr>
<td>14:00</td>
<td>O31 - Functional analysis of plant hyperacumulators</td>
<td>Marinus Pilon</td>
</tr>
<tr>
<td>14:00</td>
<td>O32 - Influence of sulfate on selenium</td>
<td>Michele Schiavon</td>
</tr>
<tr>
<td>14:00</td>
<td>O33 - Selenolanthionine is the main selenium species</td>
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### 14:00 – 15:00

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<td>Milton Ferreira Moraes</td>
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<td>O31 - Functional analysis of plant hyperacumulators</td>
<td>Marinus Pilon</td>
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<td>O32 - Influence of sulfate on selenium</td>
<td>Michele Schiavon</td>
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### 15:00 – 16:00

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### 16.00 – 16.30

#### Coffee break
# Monday 14 August 2017

## Samuelsson

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<tbody>
<tr>
<td>16:30</td>
<td>1.4. Relationships of selenium between soils, water, and vegetation (Session III)</td>
<td>Chair: Luiz Roberto Guimarães Guilherme</td>
<td>Zhengyu BAO</td>
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<tr>
<td>16:30</td>
<td>O50 - Selenium distribution and its correlation to geochemical factors in East China intertidal zone</td>
<td>Zhongyu BAO</td>
<td>Jing Wang</td>
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<tr>
<td>16:30</td>
<td>O51 - Distribution and translocation of selenium from soil to highland barley in Kashin-Beck disease area</td>
<td>Jing Wang</td>
<td>Abdolbaset Karim</td>
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<tr>
<td>16:30</td>
<td>O52 - The status of selenium in Iraqi Kurdistan and feasibility of Se bio fortification using 77-Se</td>
<td>Abdolbaset Karim</td>
<td>Saeed Ahmad</td>
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<tr>
<td>17:50</td>
<td>17:10</td>
<td>O53 - Geochemistry of selenium in Gilgit-Baltistan (North East Pakistan)</td>
<td>Saeed Ahmad</td>
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<td>2.4. Selenoprotein synthesis pathways (Session II)</td>
<td>Chair: Michael Rother</td>
<td>Norman Metanis</td>
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<tr>
<td>16:30</td>
<td>O54 - Selenocysteine chemistry and total chemical synthesis applied for accessing human selenoproteins</td>
<td>Norman Metanis</td>
<td>Qing Cheng</td>
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<td>16:30</td>
<td>O55 - Redefining UAG for selenocysteine widens the scope of recombinant selenoprotein production in E.coli</td>
<td>Qing Cheng</td>
<td>Ryuta Tobe</td>
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<td>16:30</td>
<td>O56 - Delivery of selenide to selenophosphate synthetase for selenoprotein biosynthesis in bacteria</td>
<td>Ryuta Tobe</td>
<td>Mark Pinkerton</td>
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<td>O57 - Reconstitution of processive selenoprotein P synthesis in the wheat germ lysate system</td>
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<td>Chair: Björn Åkesson</td>
<td>Ralph Muecke</td>
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<td>O58 - Selenium in radiation oncology – 10 years of experiences in Germany</td>
<td>Ralph Muecke</td>
<td>Oliver Micke</td>
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<td>O60 - Selective targeting of leukemia stem cells by selenium</td>
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<td>O61 - Methylselenol suppressed the metastatic potential of B16F10 melanoma by reducing integrin expression</td>
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<td>O62 - Selenium enhances auranofin-mediated Nrf2 induction in lung epithelial cells</td>
<td>Trent Tipple</td>
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<td>O63 - Preclinical chemopreventive efficacy of a novel hybrid p-XSC-aspirin compound in a lung cancer model</td>
<td>Daniel Plano</td>
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<td>O64 - Selenofolate is a promising novel agent in targeted chemotherapy</td>
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### 17:50 – 18:00

**Short break**
### Monday 14 August 2017

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<td>O65 - Environmental remediation of selenium contamination in water and soil&lt;br&gt;&lt;br&gt;Zhi-Qing Lin</td>
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<td>O66 - Integrated passive biological selenium treatment system: Results of a 1-year pilot study&lt;br&gt;&lt;br&gt;James Bays</td>
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<td>O68 - Role of antioxidant defense system and mitochondrial activity in selenium toxicity tolerance in wheat&lt;br&gt;&lt;br&gt;Sucheta Sharma</td>
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*Poster session with dinner and exhibition*
### Tuesday 15 August 2017

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<td>O76 - Agronomic strategies affect the efficacy and quality of selenium biofortification</td>
<td>Graham Lyons</td>
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<td>O77 - Improving selenium supply in food systems</td>
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<td>O78 - Agronomic Biofortification with Selenium in Intercropping Systems would address Low Selenium Intake</td>
<td>Allan Chilimba</td>
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<td>O79 - To bio or not to bio? Strategies for agronomic biofortification with Se in tropical agroecosystems</td>
<td>Luiz Roberto Guimarães Guilherme</td>
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<td>O80 - Soil and foliar application of selenium in upland rice aiming agronomic biofortification</td>
<td>Andre Rodrigues dos Reis</td>
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<td>O81 - A model for Glutathione peroxidase 4-catalyzed reduction of lipid hydroperoxides in membranes</td>
<td>Marcus Conrad</td>
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<td>O83 - The key ferroptosis regulatorGPX4: cellular mechanisms and in vivo relevance</td>
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<td>O84 - GPx4 depleted cell death involves different cell death pathway from ferroptosis</td>
<td>Hirota KImai</td>
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<td>O82 - Selenium versus sulfur: reversibility of chemical reactions and resistance to permanent oxidation in proteins and nucleic acid</td>
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<td>09:00</td>
<td>O88 - Health Benefit Values (HBV) reliably indicate effects of seafood consumption</td>
<td>Anna Kipp</td>
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<tr>
<td>09:30</td>
<td>O85 - Selenoproteins expressed in intestinal stem cells and cancer stem cells</td>
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<tr>
<td>10:00</td>
<td>O89 - Molecular and Metabolic Mechanisms for Steatosis and Obesity Induced by Overexpression of Glutathione Peroxidase-1 in Mice</td>
<td>Xingen Lei</td>
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<td>10:10</td>
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<tr>
<td>10:10</td>
<td>O86 - Roles of the thioredoxin and glutathione systems in reduction of inorganic- and Cys-polysulfide spec</td>
<td>Péter Nagy</td>
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<td>10:30</td>
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</table>

**10.30 – 11.00** Coffee break
## Tuesday 15 August 2017

<table>
<thead>
<tr>
<th>Time</th>
<th>Session</th>
<th>Speaker</th>
</tr>
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<tbody>
<tr>
<td>11:00</td>
<td>1.6. Strategies to improve selenium accumulation and biofortification</td>
<td>Samuelsson</td>
</tr>
<tr>
<td>11:00</td>
<td>2.6. Selenoprotein function (Session II)</td>
<td>Vesalius</td>
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<tr>
<td>11:00</td>
<td>3.5. Selenium metabolism or selenoprotein function in health and disease (Session II)</td>
<td>Retzius</td>
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<tr>
<td>11:00</td>
<td>Chair: Andre Rodrigues dos Reis</td>
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<tr>
<td>11:00</td>
<td>Chair: Ulrich Schweizer</td>
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<td>11:00</td>
<td>Chair: Anna Kipp</td>
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<tr>
<td>11:00</td>
<td>Chair: Chang Liao</td>
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<tr>
<td>11:00</td>
<td>Chair: Girish Kirimanjeswara</td>
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<tr>
<td>11:00</td>
<td>Chair: Tejo Prakash Nagaraja</td>
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<tr>
<td>11:00</td>
<td>11:20 O90 - Selenium use efficiency by wheat cultivars</td>
<td>Milton Ferreira Moraes</td>
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<tr>
<td>11:20</td>
<td>11:20 O91 - Effects of selenium biofortification on mineral nutrients in grains of wheat and oat</td>
<td>Tao Li</td>
</tr>
<tr>
<td>11:20</td>
<td>11:40 O92 - Characterization on rhizosphere bacteria from a selenium-hyperaccumulator Cardamine hupingshanesis</td>
<td>Linxi Yuan</td>
</tr>
<tr>
<td>11:20</td>
<td>11:40 O93 - Assessment and biofortification of wheat grain selenium in staple food production regions of China</td>
<td>Sen Wang</td>
</tr>
<tr>
<td>11:20</td>
<td>11:40 O94 - Selenium supplemented kale and kohlrabi sprouts as possible ingredients for potent functional food.</td>
<td>Pawel Zagrodzki</td>
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<tr>
<td>11:20</td>
<td>11:20 O95 - Selenoproteins in hypothalamic energy homeostasis</td>
<td>Matthew Pitts</td>
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<td>11:20</td>
<td>11:20 O96 - Role of hypothalamic selenoprotein M in leptin signaling and calcium regulation</td>
<td>Ting Gong</td>
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<td>11:20</td>
<td>11:40 O97 - Exploring Selenoprotein N structure and function</td>
<td>Alain Lescure</td>
</tr>
<tr>
<td>11:20</td>
<td>11:00 O98 - Disruption of cancer cell redox homeostasis promoted by S-nitrosylation of thioredoxin reductase</td>
<td>Moran Benhar</td>
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<tr>
<td>11:20</td>
<td>11:00 O99 - Influence of small intestinal thioredoxin and thioredoxin reductase on intestinal permeability</td>
<td>Liangwei Zhong</td>
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<tr>
<td>11:20</td>
<td>11:00 O100 - The intricate role of selenoproteins in stress erythropoiesis</td>
<td>Chang Liao</td>
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<td>11:20</td>
<td>11:00 O101 - Selenoproteins restrict the replication of Francisella tularaensis in macrophages.</td>
<td>Girish Kirimanjeswara</td>
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<td>11:20</td>
<td>11:00 O102 - Se-dietary matrices can upregulate the anti-inflammatory responses in RAW macrophages</td>
<td>Tejo Prakash Nagaraja</td>
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<tr>
<td>12:30</td>
<td>12:00 O103 - The intricate role of selenoproteins in stress erythropoiesis</td>
<td>Chang Liao</td>
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<tr>
<td>12:30</td>
<td>12:00 O104 - Selenoproteins restrict the replication of Francisella tularaensis in macrophages.</td>
<td>Girish Kirimanjeswara</td>
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<tr>
<td>12:30</td>
<td>12:00 O105 - Se-dietary matrices can upregulate the anti-inflammatory responses in RAW macrophages</td>
<td>Tejo Prakash Nagaraja</td>
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**Poster Session with lunch and exhibition**
## Tuesday 15 August 2017

### Vesalius

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<thead>
<tr>
<th>Time</th>
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<tbody>
<tr>
<td>14:00</td>
<td><strong>2.7. The systems biology of selenium and selenoproteins</strong></td>
<td>Chair: Vadim Gladyshev</td>
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<tr>
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<td>O103 - Selenium in human and vertebrate evolution</td>
<td>Sergi Castellano</td>
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<td>O104 - Evolution of selenoproteins across the tree of life</td>
<td>Marco Mariotti</td>
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<td>O105 - The human selenomicobiome</td>
<td>Didac Santesmasses</td>
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<td>O106 - Selenoprotein extinction in Drosophila occurred concomitantly to genome catastrophes</td>
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<td>O107 - Metabolomics of selenium</td>
<td>Sun Hee Yim</td>
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<td>O108 - Selenium-encoded chemical proteomics</td>
<td>Chu Wang</td>
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<td>O109 - Characterization of Atlantic salmon (Salmo salar) selenoproteins using bioinformatics and hyphenated</td>
<td>Veronika Sele</td>
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<tr>
<td>15:50</td>
<td><strong>3.5. Selenium metabolism or selenoprotein function in health and disease (Session III)</strong></td>
<td>Chair: Lutz Schomburg</td>
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<tr>
<td>14:00</td>
<td>O110 - The cellular location of selenoproteins in human prostatic tissue and their role in prostate cancer.</td>
<td>Alan Diamond</td>
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<tr>
<td>14:20</td>
<td>O111 - Dietary selenium deprivation oppositely impacts longevity and healthspan in telomere dysfunctional mice</td>
<td>Wen-Hsing Cheng</td>
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<tr>
<td>14:40</td>
<td>O112 - Selenium and cataract</td>
<td>Kaixun Huang</td>
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<tr>
<td>15:00</td>
<td>O113 - Role of selenoprotein P in Alzheimer's disease</td>
<td>Xiubo Du</td>
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<td>15:20</td>
<td>O114 - Role of selenoprotein P in function of pancreatic β cell: Improving effects of neutralizing antibody</td>
<td>Yoshiro Saito</td>
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<tr>
<td>15:40</td>
<td>O115 - Tissue-specific pools of selenoprotein P differentially modify colitis-associated carcinogenesis</td>
<td>Sarah Short</td>
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<tr>
<td>15:50</td>
<td>O116 - Selenoprotein P in cord serum: wide disparity between ELISA and HPLC</td>
<td>Margaret Rayman</td>
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### Excursion to Mariefred (pre-booked participants only)

Coaches leave from outside the Berzelius and Scheele laboratories at 16.30
Wednesday 16 August 2017

<table>
<thead>
<tr>
<th>Time</th>
<th>Session</th>
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</table>
| 09:00 - 11:00| 3.5. Selenium metabolism or selenoprotein function in health and disease (Session IV)  
Chair: Ed Schmidt |
| 09:00 - 09:20| O117 - Expression and activity of enzymes of selenium metabolism in the selenoprotein P knockout mouse  
Marla J. Berry |
| 09:20 - 09:40| O118 - The Role of Selenoprotein K in Progression and Metastasis of Melanoma  
Peter Hoffmann |
| 09:40 - 10:00| O119 - Targeting thioredoxin reductase 1 as a basis for anticancer therapy  
Elias Arnér |
| 10:00 - 10:15| O120 - Redox regulation of protein kinase C by selenium and selenoprotein thioredoxin reductase influences the cancer-preventive efficacy of selenium  
Rayudu Gopalakrishna |
| 10:15 - 10:30| O121 - Thioredoxin reductase 2 inhibition by tamoxifen-like metalloconfins drives Jurkat cells to apoptosis  
Valeria Scalcon |
| 10:30 - 10:45| O122 - Dietary selenium and the 15 kDa selenoprotein influence initiation/promotion of colon carcinogenesis  
Petra Tsuji |
| 10:45 - 11:00| O123 - Selenoprotein T is a novel neuroprotective antioxidant enzyme in Parkinson’s disease  
Youssef Anouar |

11.00 – 11.30 Coffee break
Wednesday 16 August 2017

### Vesalius

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</table>
| 11:30 - 11:50 | 2.8. Selenium based biotechnological applications  
Chair: Elias Arnér                                                                                      |                        |
| 11:30 - 11:50 | O124 - Selenium atom-specific functionalization of nucleic acids for structure and function studies  
Zheng Huang                                                                                               |                        |
| 11:50 - 13:15 | 2.9. Additional and emerging topics of selenium in molecular life sciences  
Chair: Elias Arnér                                                                                      |                        |
| 11:50 - 12:20 | O126 - Selenocysteine and the genetic code  
Dieter Söll                                                                                               |                        |
| 12:20 - 12:40 | O127 - Cysteine polysulfidation governed by cysteinyll-tRNA synthetase (CARSs)  
Takaaki Akaike                                                                                           |                        |
| 12:40 - 13:00 | O128 - Direct observation of methylmercury and auranofin binding to selenocysteine in thioredoxin reductase  
Ingrid Pickering  
Hugh Harris                                                                                              |                        |
| 13:00 - 13:15 | O129 - X-ray fluorescence imaging and X-ray absorption spectroscopy combined yield insight on mammalian selenium biochemistry  
Hugh Harris                                                                                              |                        |
| 13:15 - 13:30 | 3.6. Clinical genetics of selenium or selenoprotein-encoding genes  
Chair: Elias Arnér                                                                                      |                        |
| 13:15 - 13:30 | O125 - Interrelationships among SELENOF genotype, cellular localization, serum selenium and race in prostate cancer  
Dede Ekoue                                                                                               |                        |

### Retzius

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</table>
| 11:30 - 13:30 | 3.7. Additional and emerging topics of selenium in health or disease  
Chair: Regina Brigelius-Flohé                                                                                     |                        |
| 11:30 - 11:50 | O130 - Se-speciation investigations at neural barrier (NB)  
Bernhard Michalke                                                                                               |                        |
| 11:50 - 12:10 | O131 - Selenium-binding protein 1 in serum may signify a heart at risk  
Eike Kuehn                                                                                                   |                        |
| 12:10 - 12:30 | O132 - Of dogs and men: Review of translational impact of dog studies on selenium and prostate cancer risk  
David Waters                                                                                               |                        |
| 12:30 - 12:45 | O133 - High selenium induces endothelial dysfunction via endoplasmic reticulum stress  
Matthesiiso Zachariah                                                                                          |                        |
| 12:45 - 13:00 | O134 - Selenium and sex: competition between brain and testes for selenium results in male-specific consequences in mice and men  
Marla Berry                                                                                                 |                        |
| 13:00 - 13:15 | O135 - Ceramide analog S14 causes a coordinate downregulation of selenoproteins in a murine psoriasis model  
Jack L. Arbiser                                                                                             |                        |
| 13:15 - 13:30 | O136 - Maternal nutrition and transcript abundance of selenium related genes in fetal bovine hepatic tissues at d 50 of gestation  
J. S. Caton                                                                                                 |                        |

13.30 – 15.00 Lunch in poster and exhibition room
Wednesday 16 August 2017

Vesalius

15:00 - 16:30  Planning for The 12th International Symposium on Selenium in Biology and Medicine and Information on The 6th International Conference on Selenium in the Environment and Human Health

Chairs: Elias Arnér and Gary Bañuelos

- Decision of venue and date for The 12th International Symposium on Selenium in Biology and Medicine.
- Presentation by Dongli Liang on The 6th International Conference on Selenium in the Environment and Human Health, to be held in Yangling/Xi'an, China, 2019.

Introducing International Society for Selenium Research (ISSR)

16:30 - 18:00  Natural Biofortification Program (NBP) satellite meeting

Chairs: Gary Bañuelos and Zhi-Qing Lin

16:30 - 16:50  O137 - A ten-year study and practice of functional agriculture in China

Xuebin Yin

16:50 - 17:00  Introduction of NBP and open proposal call to participate

Linxi Yuan

17:00 - 17:10  Does selenium intake relate to human longevity: A case study in Shitai, Anhui, China

Zedong Long

17:10 - 17:20  Selenium products produced from natural Se-rich areas

Seran Group (Gold Sponsor)

17:20 - 17:30  Leading functional agriculture practices in China

Setek Co. Ltd. (Gold Sponsor)

17:30 - 18:00  Concluding words

Zhi-Qing Lin

Retzius

15.00 - ca. 18:00  Elsevier Workshop

O144 – How to write a great research paper, and get it accepted by a good journal

Anthony Newman

16:30 - 18:30  Time to dismount posters

18:30  Coaches leave at 18:30 from outside the Berzelius and Scheele laboratories for transport to the Stockholm City Hall

19:00 – ca 20:30  Reception in the Stockholm City Hall, by invitation of the City and the Council of Stockholm.
# Thursday 17 August 2017

## Aula Medica

<table>
<thead>
<tr>
<th>Time</th>
<th>Session</th>
<th>Speaker(s)</th>
</tr>
</thead>
</table>
| 08:30 - 10:30 | **Plenary Session I, Aula Medica**<br>
Chair: Arne Holmgren | O87 - Nutritional aspects of selenium in human beings<br>
*Raymond F Burk* | 09:00 - 09:30 | O139 - Selenium from ocean fish provides protection against mercury toxicity<br>
*Nicholas Ralston* | 09:30 - 10:00 | O138 - From atom to field: how and why plants hyperaccumulate selenium, and how this affects ecosystems<br>
*Elizabeth Pilon-Smits* | 10:00 - 10:30 | O140 - Selenium at the redox interface of the genome and exposome<br>
*Dean Jones* |
| 10.30 – 11.00 | **Coffee break** | |
| 11:00 - 12:30 | **Plenary Session II, Aula Medica**<br>
Chair: Raymond Burk | O141 - Enzymology and biological functions of glutathione peroxidases<br>
*Regina Brigelius-Flohé* | 11:00 - 11:30 | O142 - Selenium versus Sulfur in GSH Peroxidases<br>
*Fulvio Ursini* | 11:30 - 12:00 | O143 - Nrf2 improves leptin and insulin resistance provoked by selenocysteine-trna knockout in hypothalamus<br>
*Masayuki Yamamoto* |
| 12:30 - 13:00 | **Closing Session and Awards, Aula Medica**<br>
Chair: Elias Arnér and Gary Bañuelos | |
Monday 14 August (Afternoon posters)

Presenting authors of posters listed below are kindly asked to be present at the respective poster during the first 30 min of the session.

12:30 - 14:00  Poster Session with lunch and exhibition, Poster and Exhibition Room

P1 - In vitro Generation of Superoxide by Selenofolate in MDA-MB-468 Breast Cancer Cells
Soni Khandelwal

P2 - A preliminary study of selenium species in natural selenium-enriched garlic
Shaozhan Chen

P3 - Catalytic redox activity of selenium compounds generating superoxide assessed by chemiluminescence
Soni Khandelwal

P4 - Quantification of selenoprotein P in human serum using isotope dilution analysis
Christian Deitrich

P5 - Relevancy of using Se speciation for geographical authentication purpose: example of red wines
Veronique Vacchina

P6 - Selenium Bicentennial: Two Hundred Years of Selenium Discovery
Mikael Bjornstedt

P7 - Simultaneous determination of selenium and sulfur species in biological samples
Nina Kroepfl

P8 - The fractions and distribution of soil selenium in Heilongjiang province and its' impacting factors
Fengqin Chi

P9 - Zoning pollution-free and selenium-rich land resources with geochemistry
Tao Yu

P10 - Selenium, Nitrogen and Carbon remobilization study following litter decomposition during 10 months
Maryse Castrec-Rouelle

P11 - The Precipitation Impact on the Selenium Speciation in Surface Soil
Dacheng Wang

P12 - Effect of pH, iron plaque and phosphorus on selenium uptake by rice seedling
Ju Min

P13 - Analysis of Se (IV) and Se(VI) absorption kinetics of different genotypes of Nicotiana tabacum L.
Shuxin Tu

P14 - Agronomic biofortification of carrots with selenium in oxidic soil
Ediu Carlos Silva Junior

P15 - Effects of different selenium application on selenium accumulation in Lentinula edode
Dongli Liang

P16 - Apples: a suitable target for selenium biofortification?
Diemo Daum

P17 - Optimising fertiliser formulations for selenium biofortification of wheat grain
Chandnee Ramkissoon

P18 - Effects of foliar spraying selenite or selenate at different stage on the selenium uptake and distri
Xinwei Liu

P19 - Se and its antagonists Hg, As, Cd in hair of Taiwanese and Russian residents
Margarita Skalnaya

P20 - Selenium and mercury interactions in the apex predators from the Gulf of Trieste (northern Adriatic)
Jadran Faganelli

P21 - Environmental Selenium Influences Fish Methylmercury Bioaccumulation and Risks
Nicholas Ralston

P22 - Study of geochemical characteristics and influencing factors of soil selenium in the typical soil
Zhongfang Yang

P23 - Interaction of selenium and cadmium in soil-corn system in natural selenium and cadmium rich area
Zezhou Zhang
Monday 14 August (Evening posters)

Presenting authors of posters listed below are kindly asked to be present at the respective poster during the first 30 min of the session.

19:00 - 21:00  Poster Session with dinner and exhibition, Poster and Exhibition Room

P24 - Impact of Initial Se Status and Gpx1 Genotype on Selenoenzyme and Transcript Expression in Mice
Roger A Sunde

P25 - Insight into the microbial respiration of selenium. A metagenomics study
Simon Mills

P26 - Selenium subcellular distribution, speciation and antioxidant response in rice booting stages
Shuxin Tu

P27 - Explore the subcellular distribution and speciation of Se in pakchoi (Brassica chinensis L.)
Dongli Liang

P28 - Comparison in bioavailability of nine bioselenocompounds
Kazuaki Takahashi

P29 - Characterization of rhodanese-like protein from Geobacter sulfurreducens
Olajumoke Kadiri

P30 - A novel Geobacteraceae-specific outer membrane protein required for selenite and tellurite reduction
Mst. Ishrat Jahan

P31 - Effect of inorganic and organic selenium compounds on tumor infiltrating lymphocytes
Deepika Nair

P32 - Small Selenium Species: Toxicity, Bioavailability and Modes of Action in Caenorhabditis elegans
Isabelle Rohn

P33 - Diabetic fish on selenium supplements
Lynn Weber

P34 - Biochemical characterization of novel methylseleno derivatives
Prajakta Khalkar

P35 - Metabolomics Profiling of the Liver of the Selenocysteine Lyase Knockout Mouse
Lucia Seale

P36 - Cardiac muscle necroptosis induced by selenium deficiency implicates miR-200a activating
Yang Tianshu

P37 - Selenophosphate synthetase 1 is an essential protein with roles in regulation of redox homoeostasis
Byeong Jae Lee

P38 - Influences of TRIT1 catalysed tRNA-modification i6A37 on translation efficiency of selenoproteins
Simon Bohleber

P39 - Genetic screening for unkown factors in selenium metabolism in Archaea
Vivien Quitzke

P40 - A role of bacterial thioredoxin in selenide delivery to selenophosphate synthetase in vitro
Atsuki Shimizu

P41 - Selenoprotein S is involved in degradation of C99 through ERAD.
Jun Ki Jung

P42 - Computational identification of the selenocysteine tRNA (tRNA-Sec)
Didac Santemasasses

P43 - The use of dimedone to study redox states of selenoproteins
Robert J. Hondal

P44 - Site specific acetylation of thioredoxin reductase 1
David Wright

P45 - Thioredoxin reductase 1 and NADPH directly protect protein tyrosine phosphatase 1B from inactivation
Markus Dagnell

P46 - Details in the catalytic mechanism of mammalian selenoprotein thioredoxin reductase 1 revealed using point mutations and juglone reducing activities
Jianqiang Xu
P47 - Probing the role of TRP14 (TXNDC17) in cellular signaling pathways
   Belén Espinosa Fernández

P48 - Apatone inhibited GPx activity and triggered AIF-mediated cell death pathway in cancer cells
   Xiaoyuan Ren

P49 - The significance of selenoprotein P expression in pancreatic beta-cell line MIN6
   Shohei Nakao

P50 - Interaction between selenoprotein W and 14-3-3 is regulated by oxidative stress.
   Kwan Young Ko

P51 - Immortalized human prostate cell line RWPE-1 as a model to study selenoprotein regulation and function in normal prostatic tissue
   Lenny Hong

P52 - Glutathione depletion in HEK293T addresses cytosolic GPx4 to the membrane
   Ana-Marija Vuckovic

P53 - Synthesis and efficacy of conjugated selenium ADC monoclonal antibodies in vitro
   Soni Khandelwal

P54 - Antifungal Activity of Selenium Nanoparticles Synthesized by B. subtilis Against P. syringae
   Miao Li

P55 - Exosome-mediated methylmercury detoxification accelerated by selenium compound, selenoneine in aquatic organism
   Shintaro Imamura

P56 - Studying selenoproteome regulation using selenium stable isotope labeling
   Laurent Chavatte
Tuesday 15 August (Afternoon posters)

Presenting authors of posters listed below are kindly asked to be present at the respective poster during the first 30 min of the session.

12:30 - 14:00  Poster Session with lunch and exhibition, Poster and Exhibition Room

P57 - Progress in selenium nutrition in China during the last thirty years
Weiming Shi

P58 - Protective effect of organic selenium-enriched extract from cardamine violifolia on carbon tetrachloride-induced hepatic damage in mice
Xin Cong

P59 - Comparison of Trace Elements and Oxidant Status in Dairy Cows at Different Physiological Stages
Zongping Liu

P60 - Relative bioavailability of selenium sources for beef cattle using glutathione peroxidase activity in liver
Marcus Zanetti

P61 - Organic Se has better protective effects than inorganic Se against AFB1/OTA-induced immunotoxicity
Fang Gan

P62 - Dietary selenium forms influence selenogenome expression in broiler chickens
Mickael Briens

P63 - Eisenia fetida - a novel organic selenium-biofortified soil animal
Shizhong Yue

P64 - Dietary selenium supplies in Malawi
Edward Joy

P65 - Selenium status in healthy elderly from the Northwest of Algeria
Nouria Dennouni

P66 - Association between Selenium Levels and Antioxidant Capacity in An Elderly Chinese Population
Liqin Su

P67 - Selenium and Alzheimer’s disease: facts and effects
Barbara R Cardoso

P68 - Selenomethionine improves synaptic deficit in an Alzheimer’s disease mouse model
Zhonghao Zhang

P69 - Effect of selenium supplementation in Hungarian patients with autoimmune thyroiditis and endocrine orbitopathy
Jeannette Molnar

P70 - Selenium supplementation as a defense method against posttraumatic stress disorder development in combats
Vladimirs V. Voicehovskis

P71 - Selenium as a mercury antidote
Xin Fu

P72 - Gender specific differences in urinary level of 8-oxodG in selenium supplemented subjects
Ewa Jablonska

P73 - Cystine-glutamate antiporter expression as a potential target for cancer therapy using redox active selenium compounds
Arun Kumar Selvam

P74 - Finding What’s Important In Selenium Drug Development
G. Michael Wall

P75 - Seleno heterocyclic compounds as antitumoral and radical scavenging agents
Daniel Plano

P76 - Selenocyanate and diselenide amides: A new class of potent antichagasic agents
Carmen Sanmartin

P77 - Cytotoxicity, oxidative stress and antioxidant enzyme activity in pancreatic cancer cells treated with organic selenium compounds
Jeremy Braude

P78 - Altered Chemoresistance in an Ex Vivo Pancreatic Cancer Model Compared to In Vitro Cell Culture
Rim Jawad
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Abstracts for Oral Presentations
O1 - The global cycle of selenium
Inaugurating plenary session

Keywords: global biogeochemical cycle, atmosphere, soils

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Introduction: The distribution and chemical speciation of selenium in the environment is of key importance in both environmental health issues and selenium status of crops and thus dietary intakes. However, there is a lack in understanding of the biogeochemical selenium cycle and especially the role of the atmosphere in this cycle. The atmosphere is a global transient reservoir of selenium that can function as a source of selenium to terrestrial environments, and thus food chains, via wet and dry deposition.

Method: This talk will give new insights into the atmospheric sources, sinks and fluxes of selenium and how these are linked. Biogenic sources of atmospheric selenium will be discussed as well as atmospheric pathways and deposition onto the Earth’s surface.

Result: Climate influences selenium distributions in soils, both indirectly (by affecting retention and mobility of Se in soils) and directly (atmospheric sources), and therefore, we further investigated broad-scale relationships between climatic factors, soil properties, and soil selenium concentrations. In this talk it will be shown how climate and soil parameters can be used to predict selenium concentrations in areas where these have not been systematically analysed. Predictions of current soil selenium concentrations and future changes in these concentrations due to climatic changes indicate that soil Se distributions are dynamic.

Discussion: As changes in spatial selenium distributions may ultimately affect nutritional quality of food crops and thus human and animal health, such broad-scale predictions will help in the prevention of future health hazards related to unsafe levels of Se in soils.
O2 - Selenium Metabolism in Plants
Inaugurating plenary session

Keywords: genetics, hyperaccumulator, metabolism, nutrition, toxicity

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1
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**Introduction:** Selenium (Se) is not an essential element for flowering plants (angiosperms), although there is evidence that Se can benefit their growth and survival under some circumstances. However, excess Se accumulation is toxic to most angiosperms, presumably because the indiscriminate incorporation of selenocysteine and selenomethionine into proteins impairs their function.

**Method:** There are differences between angiosperm species in their ability to tolerate Se in their tissues, which are related to their Se metabolism. These differences have shaped the ecology of seleniferous soils.

**Result:** This lecture will first describe the proteins involved in Se uptake, translocation and metabolism in angiosperms, their location within the plant and their regulation. It will then identify differences in metabolic pathways between angiosperm species that might account for variation in their ability to tolerate large tissue Se concentrations and the genetics that underpins these differences. It will emphasise that angiosperm species that hyperaccumulate Se generally exhibit constitutive expression of genes encoding Se transporters and enzymes involved in primary Se assimilation, the biosynthesis of non-toxic Se metabolites, and Se volatilisation.

**Discussion:** Finally, strategies for exploiting differences in the abilities of plants to accumulate Se in their tissues for either the phytoremediation of Se-contaminated soils or Se-biofortification of edible crops for the nutrition of animals and humans will be discussed.

**Selected references**
O3 - Selenium Utilization in Diverse Animals
Inaugurating plenary session
Thressa C. Stadtman Memorial Lecture

Keywords: selenium utilization, evolution

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Introduction: The Gladyshev lab works in the areas of selenium and redox biology as applied to aging and cancer. Selenium is a trace element that exhibits both beneficial and toxic effects in human health. Importance of this micronutrient in the diet is primarily due to the fact that selenium is used in selenoproteins in the form of selenocysteine.

Method: In this presentation, discussion will be focused on functions and evolution of selenium and selenoproteins. Comparative and functional genomics methods allow assessing its use at the levels of proteins, cells, organs and entire organisms.

Result: Selenoproteins with known functions are oxidoreductases, and the tight link between selenium and redox biology offers an opportunity to better understand protein function and use this information to examine questions central to the thiol-based redox control of cellular processes.

Discussion: We use high throughput approaches, including genome sequencing, transcriptomics, metabolomics and ionomics, to better understand the systemic role of selenium in diverse animals and other organisms.
O4 - Phenotypes and molecular pathogenesis of disorders of human selenoprotein synthesis

Inaugurating plenary session

Keywords: SECISBP2; TRU-TCA1-1; SEPSECS

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Introduction: Mutations in three human genes (SECISBP2, TRU-TCA1-1, SEPSECS) within the selenocysteine (Sec) biosynthesis and incorporation pathway, resulting in impaired synthesis of selenoproteins, have been described.

Method: We review the genetic architecture, molecular pathogenesis and clinical phenotypes of these different disorders.

Result: Nine families, harbouring compound heterozygous or homozygous SECISBP2 defects, have been described to date. Many mutations generate aminoterminally truncated mutant proteins with reduced Sec incorporation function, mediating global selenoprotein deficiency. Growth retardation in childhood, abnormal thyroid function tests due to lack of Sec-containing thyroid deiodinase enzymes and low plasma selenium levels reflecting deficiencies of SEPP1 and GPx3, are universal features of cases. Muscular dystrophy and male infertility reflect loss of tissue-specific selenoproteins. Cutaneous photosensitivity, increased fat mass with preserved insulin sensitivity and hearing loss may be manifestations of elevated cellular ROS, secondary to lack of antioxidant selenoenzymes. Degeneration and dilatation of the thoracic aorta is a newly-recognised adult phenotype.

A single child with a homozygous TRU-TCA1-1 mutation, altering expression and base modification of tRNA[Ser]Sec, has been described. Compared to SECISBP2 defect cases, synthesis of some selenoproteins (TXRNDs, GPx4) is preserved, with some clinical features (e.g. growth retardation, abnormal thyroid hormone and selenium levels) being similar.

Ten families with compound heterozygous or homozygous SEPSECS defects, associated with progressive cerebellocerebral atrophy, mental retardation, spasticity and seizures, have been recorded.

Discussion: SECISBP2 and TRU-TCA1-1 defects mediate a multisystem disorder with a thyroid signature, due to lack of tissue-specific selenoproteins, antioxidant selenoenzymes and thyroid deiodinase enzymes. SEPSECS defects mediate a predominantly neurological phenotype.

Selected references

O5 - Inorganic selenium species separation and preconcentration with a new nanosilica-ionic liquid hybrid

1. Selenium chemistry and geochemistry
1.1. Inorganic selenium chemistry

Keywords: Preconcentration, Nanomaterial, Ionic Liquid, ETAAS

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Introduction: The functionalization of porous silica nanoparticles with the ionic liquid (IL) 3-methyl-1-dodecylimidazolium bromide and its application as a sorption material for the development of a novel separation and preconcentration method based on dispersive micro solid phase extraction (D-µ-SPE) technique for Se speciation is presented in this work.

Method: The hybrid material was obtained by direct contact of 1 mg of powdered nanosilica with an IL aqueous solution. The process was characterized by FT-IR and UV spectroscopies, as well as electronic microscopies. The D-µ-SPE method started with the selective formation of a complex between Se(IV) and APDC¹ at pH 4, followed by the extraction from 5 ml of aqueous sample with the IL-functionalized nanosilica under vortex stirring. After a centrifugation step, the material was dried and eluted with 100 µL of an ethyl acetate-Triton X-114 solution prior to Se determination by ETAAS. An identical procedure was applied for total Se determination with a pre-reduction step and Se(VI) was calculated by difference between total Se and Se(IV).

Result: Several factors affecting the functionalization, extraction and elution steps were evaluated. Thus, a retention efficiency of 100% and an enhancement factor of 45 were obtained, yielding LODs in the ppt range. The proposed method was successfully applied for the determination inorganic Se species in drinking and natural waters.

Discussion: Considering the low mass of sorbent material and the negligible amount of organic solvent used, the proposed method can be considered not only as an outstanding analytical tool, but also as a “green”² alternative to less environment-friendly methods.³,⁴

Selected references
O6 - A novel operon involved in selenite reduction in Geobacter sulfurreducens

2. Selenium in the molecular life sciences
2.1 Metabolism of selenium in living cells
Keywords: selenite, porin, heme, rhodanese

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Introduction: Geobacter sulfurreducens is an obligate anaerobic Gram-negative bacterium, which grows by respiratory reduction of various metal compounds. The genome of this bacterium contains 111 ORFs coding for c-type cytochromes and 10 ORFs for selenoproteins. In the previous study, we identified a novel multiheme-containing selenoprotein (MHSEP), which carries five hemes and one selenocysteine residue per subunit in G. sulfurreducens. MHSEP is encoded by an ORF located within an operon-like gene cluster containing 10 genes (gsu2940-gsu2930), which codes for a rhodanese-like protein (GSU2940), a porin-like protein (GSU2939), c-type and b-type cytochromes (GSU2935-GSU2930). However, little is known about the function of those encoded proteins.

Method: Gene-disrupted strains were constructed for mhsep, gsu2939, gsu2940, and gsu2935-gsu2930, and their phenotypes were examined. Each of MHSEP, porin-like protein, and rhodanese-like protein was recombinantly produced, purified to homogeneity, and characterized.

Result: Phenotype analysis of the gene-disrupted mutant strains showed that mhsep, gsu2939, and gsu2935-gsu2930 are important for the reduction of selenite and tellurite by G. sulfurreducens. We found that purified MHSEP exhibited selenite-reducing activity in vitro, suggesting that it is a novel-type of selenite reductase. We also found that GSU2939 is a Geobacteraceae-specific novel porin that plays a role in the reduction of selenite and tellurite probably as a porin-cytochrome protein complex. In addition, GSU2940 showed a rhodanese activity, though it has only a slight sequence identity with already characterized rhodaneses.

Discussion: Our results suggest that MHSEP, GSU2940, and GSU2935-GSU2930 may form a novel porin-cytochrome protein complex to function in the electron transfer during dissimilatory selenite/tellurite reduction.
O7 - Regulation of selenium metabolism in Archaea

2. Selenium in the molecular life sciences
2.1 Metabolism of selenium in living cells

Keywords: Archaea, Methanococcus, transcription, regulation

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Introduction: The only known Archaea that synthesize selenocysteine (sec)-containing proteins (selenoproteins) are members of the Methanococcales and Methanopyrales. The model archaean *Methanococcus maripaludis* employs several selenoproteins in its primary energy metabolism, methanogenesis. Upon selenium deprivation, or when the pathway for selenoprotein synthesis is disrupted, they are replaced by cysteine-containing isoforms, thus allowing for selenium-independent growth. Selenium-responsive transcriptional regulation of the encoding genes appears to be the underlying principle, but the mechanism and the factors involved are largely unknown.

Method: Here, we will present our results of characterizing HrsM, a transcription factor involved in selenium-dependent gene regulation of *M. maripaludis* JJ.

Discussion: A possible scenario of how selenium is sensed in the environment, of how this information is transduced in the cell, thus ultimately effecting antagonistic transcriptional regulation of the selenoprotein genes and their cysteine-encoding isogenes will be discussed.
O8 - Anaerobic reduction of Se(IV) by bacterial strain isolated from Spanish bentonites: multidisciplinary approach characterization

2. Selenium in the molecular life sciences
2.1 Metabolism of selenium in living cells
Keywords: Se(IV), Se nanoparticles, bacterial reduction, microscopy, spectroscopy

Mohamed Larbi Merroun

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Introduction: The long-term disposal of radioactive wastes in a deep geological repository (DGR) is the international accepted solution for the treatment of these residues. Selenium is a long-lived radionuclide contained in high-level radioactive wastes. It exist in four oxidation states i.e. +VI, +IV, 0, and –II in nature. The speciation of Se depends on different factors including oxidation state, Eh, pH, microbial processes, etc. Elucidation of interactions mechanisms of Se with bacteria isolated from different barriers of DGR (e.g. bentonites) is crucial to understand the role of microbial processes in the safety case of this disposal system.

Method: The present study aimed to investigate the reduction of Se(IV) by Stenotrophomonas bentonitica BII-R7 isolated from Spanish bentonites under anaerobic relevant conditions for deep repositories. Therefore, we used a multidisciplinary approach combining microscopy (STEM/HAADF, HRTEM, etc.), spectroscopy (UV-Vis, Infrared, XRD, etc.) microbiology, biochemistry, etc.

Result: Using acetate as electron donor, XRD analysis have clearly demonstrated that the cells of the studied strain are able to reduce Se(IV) to Se(0) forming Se nanoparticles (SeNPs). Using a combination of STEM-HAADF and SEM techniques allowed for the first time to identify two different sized Se NPs (30 and 100 nm), located at the cell surface, the extracellular space and intracellularly. In addition, flow cytometry was used to evaluate the toxicity of Se(IV) on cell viability and metabolic activity.

Discussion: The results obtained indicated the role of bacteria isolated from bentonites in affecting the speciation of selenium under deep geological repository relevant conditions.
**O9 - Selenium Requirements and Upper Limits in Mammals and Avians From Enzyme and Molecular Biomarkers**

3. Selenium in animal and human health and disease
3.1 Selenium supplementation for animal and livestock health

Keywords: biomarkers, glutathione peroxidase, requirements, transcripts, upper limits

**Roger A Sunde** 1

1 Univ of Wisconsin

**Introduction:** To gain insights into nutrient biomarkers and dietary requirements, Se biomarker levels and requirements were compared in weanling rats and mice, 2-day-old lambs, and day-old turkeys and chickens in response to multiple graded levels of dietary Se provided as Na₂SeO₃.

**Method:** When fed truly Se-deficient diets (<0.007 µg Se/g), liver Gpx1 (glutathione peroxidase) activity falls in all species to <4% of Se-adequate levels, plasma Gpx3 activity falls to <3% in all species except mice, and liver Gpx4 activity falls to <10% in avians but only to ~50% of Se-adequate levels in rodents. Robust biomarkers should be specific for the nutrient, fall dramatically in deficiency, and reach well-defined plateaus.

**Result:** Se-response curves for these biomarkers reach well-defined plateaus with increasing Se supplementation in all species, collectively indicating minimum Se requirements of 0.06-0.10 µg Se/g for rats, mice, and lambs, but 0.10-0.13 µg Se/g for chicks and 0.23-0.33 µg Se/g for turkeys. In contrast, increasing dietary Se does not result in well-defined plateaus for erythrocyte Gpx1 activity and liver Se in most species. Se-response curves for Gpx1 mRNA for rodents and avians have well-defined plateaus and similar breakpoints. Gpx4 mRNA, in contrast, is not significantly regulated by dietary Se in rodents, but Gpx4 mRNA in avians decreases in Se deficiency to ~35% of Se-adequate plateau levels.

**Discussion:** Notably, no selenoprotein activities nor transcripts are robust biomarkers for supernutritional Se status; small clusters of non-selenoprotein transcripts, identified by microarray and RNA Seq, are identifying potential high-Se biomarkers and pathways associated with response to high Se.

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O10 - Effects of feeding cows Se-yeast or Se-enriched alfalfa hay on baby calves Se status and IgG titers

3. Selenium in animal and human health and disease
3.1 Selenium supplementation for animal and livestock health
Keywords: cows, calves, IgG, Se-yeast, agronomic biofortification

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Introduction: Selenium (Se) is an essential trace mineral important for immune function and overall health of cattle. Two methods of Se-delivery to pregnant cows are organic Se-yeast supplementation and agronomic Se biofortification, whereby the Se content of hay is increased through the use of Se-containing fertilizer amendments (Hall et al., 2013a and 2013b). Our objective was to evaluate the effect of these two Se-delivery methods in cows on passive transfer of IgG to calves.

Method: Se-Yeast Supplementation: During the last 8-wk before calving, dairy cows were fed either 0 (n=17) or 105 mg Se-yeast once weekly (n=20), in addition to Na-selenite at 0.3 mg Se/kg DM in their ration (Hall et al., 2014). The Se-yeast dosage was calculated to provide 15 mg of Se/d (5× the maximal FDA-permitted level). After birth, calves were fed pooled colostrum from control or supranutritional Se-yeast supplemented cows. Concentrations of whole-blood (WB)-Se and serum-IgG were measured at birth, 48-h, and 14-d of age.

Agronomic Biofortification: During the last 8-wk before calving, beef cows were fed alfalfa hay fertilized with 0 (calculated Se intake: 8.3 mg Se/head/d; n=15), 45.0 (27.6 mg Se/head/d; n=15), or 89.9g Se/ha (57.5 mg Se/head/d; n=15). Concentrations of colostrum-Se and IgG1 were measured at birth, and concentrations of calf WB-Se and serum-IgG1 were measured at birth and 12, 24, 36, and 48-h of age.

Result: Both Se supplementation strategies for cows during the dry period were effective for maximizing WB-Se and serum-IgG concentrations in calves.

Discussion: The more economical alternative is Se-agronomic biofortification.

Selected references


O11 - Thirty years of selenium fertilization: A Finnish solution to ensure an adequate selenium intake

3. Selenium in animal and human health and disease
3.1 Selenium supplementation for animal and livestock health

Keywords: fertilization, food, feed, serum

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Introduction: The average selenium (Se) intake in Finland was one of the lowest in the world in 1970’s due to the low Se content of Finnish agricultural products (Koivistoinen 1980). This was considered to pose a risk to human and animal health. Soils in Finland are not exceptionally poor in Se, but the soil and climatic conditions depress Se uptake by plants.

Method: To improve Se uptake, fertilizers have been supplemented with sodium selenate since 1985.

Result: The Se supplemented fertilization has increased Se concentration in wheat flour to about 0.14 mg kg⁻¹ dry matter, 14 times higher than prior to Se fertilization, whereas levels in beef and pork have increased 3 and 1.5 times. The average daily Se intake has been about 0.08 mg/day in recent years which meet well the recommendations and is considered adequate and safe level. Excessive Se intake from any type of diet is not probable. In spite of Se fertilization, the amount of soluble Se in soil has not increased. In Finnish conditions Se is quickly transformed in less soluble forms and annual Se fertilization is necessary to maintain the sufficient Se levels in crops. The average serum Se content has been at the level of 1.4 µmol l⁻¹ during the 2000s which is among the highest in Europe.

Discussion: A systematic follow-up study was established in 1984. It enables an adequate revision of Se levels in fertilizers when needed. Se fertilization has been an effective, safe and economical way to improve the Se situation in Finland.

Selected references
O12 - Selenium bioaccessibility in biofortified crops and food supplements: a factor to take into account?

3. Selenium in animal and human health and disease
3.1 Selenium supplementation for animal and livestock health
Keywords: digestion, bioavailability, bioaccessibility, gastrointestinal tract, intestine

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Introduction: The amount of Se released in the gastrointestinal tract may affect its bioavailability. Therefore, we assessed Se bioaccessibility in different Se-enriched food supplements and crop samples, including an extensive set of Se-rich rice and leek samples.

Method: The human gastrointestinal digestive process was simulated in vitro by conducting sequential batch extractions mimicking stomach, small intestine and colon conditions. Bioaccessibility in each extraction step was calculated as the ratio (%) between Se released into the liquid phase and total Se added through the food crop or supplement sample in the first step (stomach). The food crop samples originated from field experiments and samplings in Brazil, China and Belgium.

Result: The bioaccessibility of Se was highest in the small intestine. It is shown to be highly variable (10-60% in stomach and 20-90% in small intestine), depending on the properties of the food supplements and growth conditions of the crops. A lower Se bioaccessibility in the colon is observed (Sun et al., 2017). Some food supplements show a very low Se bioaccessibility throughout the entire intestine (Lavu et al., 2017).

Discussion: Lower bioaccessibilities may result from formation of elemental Se particles at higher supplementation doses during production of the foodstuffs. Part of the Se is transferred with remaining food matrix from small intestine to colon. The lower Se bioaccessibility in the colon may be attributed to its accumulation by gut microorganisms. To what extent Se is essential to protect the colonic environment and whether Se deficiency plays a role in colonic diseases should still be further investigated.

Selected references
O13 - Selenium Requirements of the Turkey Based on Enzymatic Biomarkers and Next-generation Sequencing

3. Selenium in animal and human health and disease
3.1 Selenium supplementation for animal and livestock health
Keywords: glutathione peroxidase, next-generation sequencing, requirement, transcript, turkey

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Introduction: The current National Research Council selenium (Se) requirement for the turkey is 0.2 μg Se/g diet.

Method: To determine dietary Se requirements using selenoenzyme activities, and to assess the potential of transcript biomarkers from next-generation sequencing (NGS) of the liver transcriptome, we fed day-old male poults a Se-deficient torula diet (<0.007 μg Se/g) supplemented with graded levels of Se (0-1 μg/g), measured selenoenzyme activities in liver, kidney and gizzard, and conducted NGS.

Result: Se deficiency decreased activity of the glutathione peroxidases (GPX1, GPX4) to <16% of Se-adequate values in liver, kidney and gizzard. Se supplementation resulted in well-defined GPX plateaus, and minimum Se requirements based on liver and kidney GPX1 are 0.30–0.33 μg Se/g diet. Se deficiency significantly altered 16 of 14,288 NGS transcripts expressed in the liver, including downregulating the selenoproteins DIO1, GPX1, GPX4, SEPP1, SELU (FAM213A) and TXNRD1. Se-response curves for regulated selenoprotein transcripts decreased in Se deficiency to 20-51% of the Se-adequate plateau values. For selenoprotein transcripts, minimum Se requirements as determined by qPCR were 0.05–0.09 μg Se/g for liver, 0.13–0.19 μg Se/g diet for kidney and 0.06–0.15 μg Se/g for gizzard. In NGS, 5 μg Se/g altered only 67 transcripts (<0.5% transcriptome), unlike the rat where 4% of the transcriptome was significantly affected by 5 μg Se/g diet.

Discussion: The minimum Se requirement of the turkey poult is at least 0.3 μg Se/g diet. Minimal changes in the liver NGS transcriptome when turkeys are fed 5 μg Se/g diet indicate that the turkey is resistant to Se toxicity.

Selected references
O14 - Selenium supplementation of feed for Atlantic salmon

3. Selenium in animal and human health and disease
3.1 Selenium supplementation for animal and livestock health
Keywords: feed legislation, fish, speciation, toxicity, transfer

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Introduction: Feed formulations for farmed marine fish are changing. Plant ingredients are increasingly replacing the traditional marine feed ingredients fishmeal and fish oil. This leads to lower levels of selenium in the feeds, and it may be necessary to add selenium to meet the dietary requirement of farmed fish. The supplementation of selenium is regulated through the European feed legislation setting maximum levels for total selenium in feed and, in some cases, maximum levels for the supplementation. The legislation is based on risk assessments, which often lack studies of selenium supplementation to feeds for fish.

Method: The present work investigated the transfer of dietary selenium to fish and possible effects in Atlantic salmon (Salmo salar). Salmon were fed experimental diets supplemented with inorganic selenium (as sodium selenite) or organic selenium (as selenised yeast) at low (1-2 mg Se kg\(^{-1}\) feed) and high (15 mg Se kg\(^{-1}\) feed) levels for 12 weeks. Muscle samples were analysed for total selenium and selenium species, while possible toxic effects were studied in liver samples.

Result: At the high supplementation level, both organic and inorganic selenium accumulated in muscle, and oxidative stress was observed in the fish. Differences between inorganic and organic selenium were observed at the low inclusion level, where selenium accumulated in fish fed organic selenium, but not in fish fed inorganic selenium. Interestingly, the accumulation of organic selenium did not lead to oxidative stress.

Discussion: An overview of the study will be given, and the implications for feed and food safety will be discussed.
Abstracts

O15 - New opportunities for selenium speciation from advanced X-ray spectroscopy
1. Selenium chemistry and geochemistry
1.2 Imaging of selenium and analytical methodologies
Keywords: X-ray absorption spectroscopy, high energy resolution fluorescence detection, speciation

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Introduction: Synchrotron X-ray absorption spectroscopy (XAS) can provide details of the local physical and electronic structure of heavier elements in a wide variety of samples. One unique advantage of XAS is that it can be used to probe chemistry in situ in a range of systems with little or no pre-treatment. As such, Se K-edge XAS has been used to develop an understanding of diverse systems including metalloenzyme active sites, intact biological tissues or organisms, and environmental samples. Despite its success, applications of the method can be limited by spectroscopic lifetime broadening, giving poor energy resolution, and by lack of access to low concentrations.

Method: In high energy resolution fluorescence detected XAS (HERFD-XAS), the spectroscopic lifetime broadening is overcome by measuring the emitted fluorescence (Se K-alpha fluorescence for the Se K-edge) at very high resolution. Measurements were conducted on beamline 6-2 of the Stanford Synchrotron Radiation Lightsource.

Result: Selenium HERFD-XAS shows exquisite spectral resolution compared with selenium XAS of the same species, enabling much greater distinction of chemical forms than is possible with XAS. In addition, the high brightness of the beamline coupled with the superior rejection of background lead to a very high signal to background ratio for HERFD-XAS, with adequate spectra from Se concentrations as low as 100 nM.

Discussion: Improvements in both spectral resolution and sensitivity indicate that HERFD-XAS shows superior selenium speciation potential, while maintaining attributes such as flexibility of sample state that make XAS such a valuable speciation tool.
O16 - Development of Fluorescent ROS Probe Based on Redox Reaction of Selenol group

1. Selenium chemistry and geochemistry
1.2 Imaging of selenium and analytical methodologies
Keywords: reactive oxygen species, fluorescent probe

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Introduction: Reactive oxygen species (ROS) are continuously generated in biological systems, and cause damage to lipids, proteins and DNA. On the other hand, ROS also act as signaling molecules in the maintenance of physiological functions. In order to investigate the behavior of ROS in vivo, we used selenol group as a sensor of ROS in our fluorescent probe. Selenium can exist in various oxidation states in biological systems. By using selenium as a ROS-sensing motif in a fluorescent probe, it enables us to observe reversible fluorescence that relevant to biological ROS signals.

Method: We have designed and synthesized “SeMF”, based on the intramolecular spiro-cyclization of selenol group. Because of high acidity of the selenol group, SeMF is expected to form a spiro ring in reductive condition due to the high nucleophilicity of selenide, and the fluorescence of SeMF is completely quenched under the condition. When SeMF is oxidized by ROS, the spiro ring form is converted to the quinonoid form that shows the strong fluorescence.

Result: SeMF was specifically oxidized to fluorescent form by hypochlorite, while the fluorescence was quickly reduced in the presence of glutathione to regenerate the reduced form. This redox-induced reversible fluorescence response of SeMF allowed us to detect the intracellular hypochlorite of HepG2 cells.

Discussion: This fluorescent probe is expected to be useful for monitoring the dynamics of ROS production continuously in vivo.
O17 - Advances in large scale speciation of selenium by plasma and molecular mass spectrometry

1. Selenium chemistry and geochemistry
1.2 Imaging of selenium and analytical methodologies
Keywords: speciation, mass spectrometry, electrospray MS, ICP MS, coupled techniques

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Introduction: Distinguishing between different selenium chemical forms present in biological systems presents a particular challenge for analytical chemists because - besides having several possible oxidation states in its inorganic form - selenium can be incorporated into tens of known (and probably many more not discovered yet) low molecular weight (below 1000 Da) selenometabolites, many SeCys- and SeMet-containing proteins, selenosugars, selenolipids and, the last but not least, present as selenium nanoparticles. Each of these forms has a distinct biological function and as such should be individually detected and determined in the presence of a rich biological matrix. A particular importance has been attributed to the selenium speciation in food sources and its metabolic pathways leading to synthesis of physiologically important selenoproteins and low molecular weight selenometabolites.

Method: Developments in selenium speciation analysis by coupled techniques combining separation methods with parallel elemental and molecular mass spectrometry have resulted in identification and quantification of a growing number of selenium species opening a way to fully comprehensive species-selective characterization of biological samples.

Discussion: The presentation will address the remaining problems in the field and propose the solutions for (1) the quantitative assessment of selenium incorporation into proteins with a focus on the preservation of the stability of SeCys residues by carefully optimized derivatization, (2) the structural elucidation of selenoproteins isoforms, (3) the speciation of high molecular selenium species in organic media and (4) quantification of nanoparticulate selenium fraction in biological samples. The role of developments in both mass spectrometry instrumentation and custom designed sample preparation methods will be highlighted.
O18 - Selenium speciation in rainwater from high altitude locations

1. Selenium chemistry and geochemistry
1.3 Local geological selenium sources and global cycling

Keywords: selenium, precipitation, speciation, rainwater, HPLC-ICP-MS

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Introduction: Atmospheric transport and wet deposition play a major role in supplying selenium (Se) and other essential (micro)nutrients to terrestrial environments, including agricultural soils. To investigate the temporal variation in total Se concentrations and speciation in rainfall, we collected and analyzed weekly precipitation over 14 and 5 months at two high-altitude locations in Europe, i.e., Jungfraujoch (JFJ, Alps, Switzerland) and Pic du Midi (PDM, Pyrenees, France), respectively. Both sites are often in the free troposphere, which enables investigation of long range atmospheric transport.

Method: We analyzed total concentrations and speciation of Se as well as of sulfur (S), iodine (I) and bromine (Br) in precipitation using (HPLC)-ICP-QQQ, dissolved organic carbon (DOC) concentrations (Shimadzu TOC-L CSH) and isotopic carbon signatures of DOC ($\delta^{13}$C, IRMS). 

Result: Selenium showed generally higher concentrations at PDM (0.6 nM) compared to JFJ (0.3 nM). At both locations, mainly inorganic and anionic species of Se, S, I, and Br were found, without clear seasonal differences. In the productive period (April-September) we found significant positive correlations between Se and biogenic non-sea-salt sulfate (for both sites) and for Se with $\delta^{13}$C-values (only for JFJ).

Discussion: The results indicate that Se in rainfall may have major contributions from marine biogenic sources at both locations. Interestingly, at the more continental location (JFJ), Se, S, I, and Br concentrations were highest when $\delta^{13}$C-signatures were less negative, indicating that marine biogenic sources bring in highest Se (and S, I, and Br) concentrations to this region. Currently, analysis of air mass sources is underway to test this hypothesis.
O19 - Effect of conservation agriculture on solubility and fractionation of soil selenium

1. Selenium chemistry and geochemistry
1.3 Local geological selenium sources and global cycling

Keywords: Minimum tillage, Mineralization

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Introduction: The benefits of Conservation Agriculture (CA) to soil fertility have been widely reported (Wu et al., 2005; Chivenge et al., 2006; Ngwira et al., 2012). However, the effects of CA on the nutritional quality of maize still present a significant knowledge gap. This study is part of an ongoing investigation into the trace element dynamics within a CA trial in Malawi.

Method: Soil and plant samples from a 9 year long-term CA trial in Malawi were analyzed for selenium and other parameters. Soil selenium data (‘total’ (HF digest), ‘exchangeable’ (phosphate-extractable) and ‘soluble’ (in 0.01 M KNO₃) from the trial plots are presented here.

Result: The total selenium concentration in topsoil (0 - 20 cm) was greater in tilled treatments, as compared to the CA minimum-tilled plots whereas, by contrast, soluble and phosphate-exchangeable Se were greater in minimum tilled plots.

Discussion: Minimum tillage is thought to increase mineralization and solubility of some nutrients in soil (Wolf and Snyder, 2003). This may include selenium, which is known to cycle between organic and inorganic forms. Thus both the adsorbed phosphate-exchangeable Se pool and the soluble soil Se were significantly greater in plots receiving minimum tillage across all crop treatments. Selenium solubility may be enhanced under conservation agriculture cropping systems. Current evidence suggests that dietary Se is at sub-optimum levels in Malawi (Hurst et al., 2012) and thus CA systems may present benefits beyond improved soil management practices.

Selected references


**O20 - Selenocysteine in mammalian thioredoxin reductase and application of ebselen as a therapeutic**

**2. Selenium in the molecular life sciences**

**2.2 Molecular mechanisms of selenium toxicity**

Keywords: thioredoxin reductase, ebselen, bacteria, silver

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**Introduction:** A thioredoxin system consisting of thioredoxin (Trx), NADPH and thioredoxin reductase (TrxR) is present in all living cells with many functions (1) as a disulfide reductase for reductive enzymes like peroxiredoxins or ribonucleotide reductase (RNR). Most cells also have a glutaredoxin system comprising glutathione (GSH), NADPH, and glutathione reductase plus glutaredoxin (Grx) acting as an efficient electron donor for RNR in DNA synthesis. Mammalian TrxRs are three large dimeric isoenzymes with an essential selenocysteine (Sec) residue in a C-terminal active site. In contrast TrxR from yeast, plants and bacteria have a small TrxR with cysteine residues in the active site and a different structure and mechanism. Auranofin a gold-containing FDA-approved drug targeting the essential Sec residue in TrxR1 in the cytosol and TrxR2 in mitochondria causes cancer cell death. Ebselen is a selenazol drug with GSH peroxidase and antioxidant activity, a superfast oxidant of reduced Trx and a substrate of mammalian TrxR (2) but in contrast a competitive inhibitor of bacterial TrxR (3).

**Method:** Here I will discuss therapeutic ebselen use.

**Result:** Ebselen has been used as a new antibiotic principle and can effectively kill bacteria like Staphylococcus aureus and MRSA or Helicobacter pylori lacking GSH and Grx (3). Recent unpublished data from our laboratory show strong synergistic effects with silver, which enables targeting multi-resistant Gram-negative bacteria like E.coli (4). Silver was a strong inhibitor of both thioredoxin and thioredoxin reductase.

**Discussion:** In combination with ebselen, silver became selectively toxic to Gram-negative bacteria as shown by mouse peritonitis model experiments (4).

**Selected references**

O21 - Characterization of novel selenium compounds as a therapeutic approach for improved cancer treatment

2. Selenium in the molecular life sciences
2.2 Molecular mechanisms of selenium toxicity

Keywords: cytotoxicity, programmed cell death, anticancer agents

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Introduction: It is well established that tumor cells in general are particularly vulnerable to oxidative stress. Increased ROS production to preferentially target cancer cells has hence emerged as an attractive strategy for new cancer therapeutics (1). Based on this knowledge, redox active selenium (Se) compounds, acting as prooxidants, have gained substantial attention with promising chemotherapeutic potential in both solid and hematological malignancies (2). Novel compounds are however warranted to increase the efficacy and specificity and to minimize the systemic effects. The better pharmacological and toxicological profiles recently disclosed by some organic Se compounds has fostered our research towards the development of this type of Se-based drugs.

Method: Following the synthesis of novel organic Se compounds, we have biochemically characterized and cell biologically evaluated these compounds based on their, cytotoxicity profile, cell death pathways, efficacy and specificity index in in vitro and ex vivo models.

Result: Several of the novel organic Se compounds synthesized have shown superior effects compared to naturally occurring Se metabolites or compared to analogues lacking the selenium moiety.

Discussion: The novel Se compounds examined has provided valuable insights into their molecular mode of action that will help in the development of new efficient Se containing chemotherapeutic agents.

Selected references


O22 - Impairment of protein homeostasis accounts for selenomethionine toxicity in Saccharomyces cerevisiae

2. Selenium in the molecular life sciences
2.2 Molecular mechanisms of selenium toxicity

Keywords: Selenomethionine, selenocysteine, toxicity, protein aggregation, yeast

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Introduction: Selenium supplementation in human diet has been proposed to exert beneficial effects against several diseases, including cancer (1). As a result, selenomethionine (SeMet) has become a popular dietary supplement. However, excessive intake is toxic. The origin of this toxicity is not yet understood, although it may help understanding the basis of SeMet beneficial properties. The high level of conservation of its genes and metabolic pathways with those of higher organisms makes yeast an attractive model organism to study the molecular basis of selenium toxicity.

Method: To identify biological pathways involved in SeMet toxicity, we performed a genome-wide screen in a yeast deletion collection. Fluorescence microscopy using a GFP-tagged Hsp104 marker was used to monitor protein aggregation in vivo. Selenium incorporation into proteins was analyzed by mass spectrometry.

Result: Genes involved in protein degradation and synthesis were enriched in the sensitive and resistant datasets, suggesting that SeMet causes a proteotoxic stress. Accumulation of protein aggregates was observed upon SeMet exposure. Reduction of translation rates was accompanied by a reduction of protein aggregation and SeMet toxicity. Protein aggregation and toxicity were suppressed in a Δcys3 mutant unable to synthesize selenocysteine. Lastly, SeCys incorporation in a reporter polypeptide was observed in cells exposed to SeMet (2,3).

Discussion: These results suggest that SeMet toxicity is caused by protein aggregation resulting from the metabolization of SeMet to selenocysteine followed by translational incorporation in the place of cysteine. The hypothesis that selenocysteine misincorporation disrupts protein homeostasis is reinforced by studies in higher eukaryotes (4).

Selected references
Abstracts

O23 - Selenium status of Russian population: interactions with demography

3. Selenium in animal and human health and disease
3.2 Epidemiology of selenium related health and disease
Keywords: hair, population, demography, mortality

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Introduction: It is interesting to study how Se status has an impact on population health and demography.

Method: The objective was to assess Se status of the Russian population using ICP-MS analysis of hair Se content from more than 60,000 adults and 15,000 children.

Result: The median (25-75 percentile) hair Se content (µg/g) in adult men and women was 0.439 (0.393-0.484) and 0.343 (0.305-0.383), whereas in boys and girls the respective values were 0.397 (0.340-0.443) and 0.388 (0.327-0.445). The lowest hair Se values in adults were detected in the Orenburg region (South Urals), whereas the highest in the Kabardino-Balkar Republic (North Caucasus). In children, the lowest Se levels were detected in Kaliningrad region, whereas the highest in Chechen Republic (North Caucasus) and Republic of Mordovia (Volga region). The IUPAC calculated reference values for hair Se levels in adult Russian population were 0.093-0.482 µg/g. Hair Se levels significantly correlated to demographic indices. In particular, despite the absence of significant association with birth rate (p=0.602) and total morbidity (p=0.312), hair Se inversely correlated with mortality (p=0.022) and was positively associated with increased life span (p<0.001) in adults. In children, a significant inverse relationship between hair Se and total morbidity (p=0.002) was observed.

Discussion: It is notable that the observed associations were more tight after adjustment for population levels of toxic trace elements (especially, Hg), being indicative of the outcome of Hg-Se antagonism. The present findings demonstrate that modulation of Se status of the population is expected to be beneficial to improve demography and population health.

Selected references


O24 - Selenium status of School Children for Kaschin–Beck Disease Endemic Areas in Tibet, China

3. Selenium in animal and human health and disease
3.2 Epidemiology of selenium related health and disease
Keywords: Children, Kaschin–Beck disease, Tibet

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Introduction: Selenium (Se) deficiency is an important environmental factor for the etiology of Kaschin–Beck disease (KBD). Although KBD is presently controlled in most regions of China, it is still active in Tibet. The study aims to explore selenium status of Se in school children in KBD areas in Tibet, China.

Method: Hair samples of 221 school children were collected in KBD areas of Lhasa, Shigatse and Chamdo during 2013-2015. The Se contents in the hair samples were measured by inductive coupled plasma mass spectrometry.

Result: The mean Se level in children’s hair was 0.252 ± 0.082 µg/g, which was 0.77 times higher than that of reported in 2003 (0.142 µg/g) in KBD areas in Tibet. The average hair Se level in boys (0.270 ± 0.088 µg/g) was significantly higher than that in girls (0.237 ± 0.075 µg/g) in the studied areas (t = 2.971,  P < 0.01). The detectable X-ray of children aged 7-12 in KBD areas in Tibet was declined from 37.86% in 2000 to less than 3% since 2010.

Discussion: The rising trend of Se level in children was in good agreement with the decline of KBD, which likely as a result of high Se content staple food substitution with the enforcement of Free Education Policy and Nutrition Improvement Plan in Tibet. However, 22.17% of students were still deemed to be in Se deficiency (hair Se <0.20 µg/g), which indicated that Se status of school children was also partly affected by low Se environment in KBD areas in Tibet.

Selected references
O25 - Impact of early-life selenium status on children’s cognitive abilities

3. Selenium in animal and human health and disease
3.2 Epidemiology of selenium related health and disease
Keywords: child, cognition, neurodevelopment, pregnancy, hair selenium

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Introduction: We recently discovered that mothers’ selenium status in pregnancy was positively associated with their children’s cognitive function at 1.5 years. The present aim was to assess whether this association persisted in later childhood, and if the children’s selenium status was of importance.

Method: This cohort study, including 1405 children, was nested in a population-based, randomized food and micronutrient supplementation trial in pregnancy in rural Bangladesh. Selenium in maternal blood (erythrocyte fraction, Ery-Se) at gestational week 14, and in children’s hair at 10 years, was measured using inductively coupled plasma mass spectrometry. Children’s cognitive function at 5 and 10 years was assessed using the Wechsler Pre-school and Primary Scale of Intelligence (3rd edition) and the Wechsler Intelligence Scale for Children (4th edition), respectively.

Result: Multivariable-adjusted linear regression analyses revealed a persistent association between prenatal selenium status and cognitive function at 5 and 10 years. A 0.5 µg/g Hb increase in maternal Ery-Se (median: 0.45 µg/g Hb) was associated with an increase in full developmental score corresponding to 0.2 and 0.3 SD at 5 and 10 years, respectively. Children’s concurrent selenium status was also positively associated with cognitive function at 10 years, possibly through interactions with thyroid hormones. Moreover, there were indications of inverse associations at the highest hair selenium concentrations (>0.7 mg/kg), although very few children (~2%) had such high concentrations. Further adjustments for other nutritional factors (maternal BMI, erythrocyte zinc, urinary iodine, hemoglobin) did not affect the associations.

Discussion: Adequate selenium status during early life appears to be beneficial for children’s cognitive function.

1. Selenium chemistry and geochemistry
1.4 Relationships of selenium between soils, water, and vegetation
Keywords: Dietary deficiency, fertilizers, food systems, geochemistry, geography, soil

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Introduction: Our research has shown that selenium (Se) deficiency is widespread in Malawi. We have been supported by many individuals and groups, including the International Selenium Society, to whom we have presented updates at all meetings since 2009. This paper summarizes our past and current research.

Method: (1) Surveying soils and food crops (Chilimba et al. 2011; Joy et al. 2015a); (2) studying human Se intakes and status (Hurst et al. 2013; Siyame et al. 2013; Gibson et al. 2015); (3) analyzing food systems (Joy et al. 2014); (4) linking food and household survey data (~12,000 households; Joy et al. 2015b); (5) studying soil geochemistry and crop responses to Se fertilizers (Chilimba et al., 2012a,b; Ligowe et al., unpubl.); (6) assessing population blood/urine biomarkers of Se status (n=~3000, Phiri et al., unpubl.); (7) evaluating the potential of minor crops to address Se deficiency (Kumssa et al., 2017).

Result: Human Se deficiency in Malawi is widespread, and influenced strongly by geography, including long-range variation due to soil-type and proximity to Lake Malawi, and short-range variation due to age, gender, urbanization and wealth.

Discussion: Research is ongoing to provide evidence to support potential policy responses to address this potential health burden, and to determine the wider regional scale of dietary Se deficiency in sub Saharan Africa.

Selected references
O27 - Key role of Selenium (0) nanoparticles in soil and as therapeutic agent

1. Selenium chemistry and geochemistry
1.4 Relationships of selenium between soils, water, and vegetation
Keywords: Nanoparticle reactivity, soil, cancer cells, XANES

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Introduction: Metallic selenium nanoparticles (SeNP) form in hydromorphic soils. via Se reduction and retention within soil aggregates, while SeNP dissolution may lead to food fortification via a Se slow delivery systems to plants. Conversely, selenium SeNP have been shown to be a promising therapy for specific ovarian and prostate cancers and a mechanistic understanding of Se methylation within cancer cell may enhance the development of cancer therapy.

Method: Mineral specific investigations were conducted with selenite ion reacted in anoxic conditions with ferrous iron in presence of soil minerals. The coupling between physical and biogeochemical processes in soil aggregates is investigated by a flow-through reactor system and a 3D reactive transport model. Soil and cancer cells are investigated by synchrotron X-ray absorption near edge spectromicroscopy (XANES) linear combination fitting (LCF).

Result: Selenite ions react with aqueous Fe2+ ion (a major ion in hydromorphic soils) in presence of various soil minerals (clays1, calcite2), or with Fe2+ rich minerals (siderite magnetite, pyrite)3,4 to form elemental SeNP and/or FeSe NP species. Within soil aggregates, a radial redox gradient is observed. The reactive transport model details the interplay between intra-aggregate Se diffusion and sorption/reduction.

Discussion: In soil, Se reduction increased towards the core of aggregates, and in clay rich soils slow Se delivery system to plants cis investigated. Conversely Se-NPs dissolution in cancer cells were shown5,6,7 to significantly inhibit the cell growth, to impact the regulation of the cell cycle e.g. with tubuline depolymerization8, to stimulate apoptosis and to inhibit tumor cell migration and invasion in vitro.

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O28 - Adsorption of selenate and selenite in cultivated and native soils of the Brazilian Cerrado

1. Selenium chemistry and geochemistry
1.4 Relationships of selenium between soils, water, and vegetation
Keywords: selenium sorption, oxidic soils, soil fertility management

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Introduction: The availability of selenium (Se) for plants - and as a result for humans/animals - is influenced by several factors, such as the chemical species of Se (e.g., selenate or selenite) and soil characteristics, as well as the contents of other ions, especially anions, soil texture, pH, organic matter content, among others (Yasin et al., 2015).

Method: This study evaluated the adsorption of selenate, Se(VI) and selenite, Se(IV) in cultivated and uncultivated soils of the Brazilian Cerrado biome. Adsorption tests were performed adding both Se species as sodium salts (selenate or selenite) at 500 µg L$^{-1}$, using 15 mmol L$^{-1}$ NaCl as the electrolyte solution (pH=5.5).

Result: The results showed that selenite was much more adsorbed on the studied soils compared with selenate. Moreover, uncultivated soils adsorbed higher amounts of both Se forms, which can be attributed to the presence of competing anions (e.g., phosphate and sulphate) added through soil fertility management in cultivated soils. Still, adsorption of selenate seems to be more affected by these competitive anions, when compared with selenite.

Discussion: The higher adsorption of Se(IV) compared with Se(VI) indicates that selenate is the key Se species to be applied in Se-poor soils to increase Se levels in plants (biofortification) and, as a results, in animals and humans (Boldrin et al., 2012; Ramos et al., 2011). Moreover, adding selenate in fertilizers containing other anions to compete with Se for sorption sites should be considered a good strategy for studies concerning agronomic biofortification with Se. (Sponsored by FAPEMIG, CNPq, and CAPES).

Selected references
O29 - SELENIUM AND BARIUM IN BRAZIL NUTS: UNRAVELLING THE SPATIAL DISTRIBUTION

1. Selenium chemistry and geochemistry
1.2 Imaging of selenium and analytical methodologies
Keywords: Amazon region, elemental distribution, potential toxicity, XRF

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Introduction: Brazil nuts (Bertholletia excelsa) from Amazônia are the richest food in selenium and accumulate significant concentrations of barium as well (Parekh et al., 2008). Selenium is a trace element essential for humans but Ba is not, and can be toxic (Suwa et al., 2008; Rayman, 2012). This study aimed to evaluate total concentration and map the elemental distribution of Se and Ba in Brazil nuts.

Method: Brazil nut samples were collected in the Amazon region (Acre, Amazonas, Roraima, and Amapá) between 2014 and 2015 and analysed in the Department of Soil Science (Lavras - MG). For total selenium and barium concentration, acid digestion was performed using 6 mL of HNO₃ + HClO₄ (2:1) (50-200°C over 2 h). Selenium was analysed via GF-AAS and barium via ICP-OES. Selenium and barium mapping was performed in 2D using μ-XRF (Tsui et al., 2015).

Result: Total Se and Ba concentration (mg kg⁻¹) were on average respectively: Acre (3.09; 2731.90), Roraima (14.93; 772.41), Amapá (59.35; 421.49) and Amazonas (69.09; 48.42). Highest Ba and Se concentrations (in Acre and Amazonas respectively) require bioavailability research to evaluate potential toxicity.

Discussion: Se concentration varied inversely with Ba concentration. Brazil nuts with highest Se content (Amapá and Amazonas) possessed higher concentrations in the extremities - the stem apex of the seed - while in other samples Se was more homogeneously distributed. Ba was concentrated around the edge of the nuts (Figure 1).

Selected references


O30 - Role of the selenium in articular cartilage metabolism, growth, and maturation

2. Selenium in the molecular life sciences
2.3 Molecular consequences of selenium deficiency

Keywords: Articular Cartilage, Osteoarthritis, Synchrotron Fourier Transform Infrared Microscopy, Synchrotron

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Introduction: In China, strong correlations exist between Se-deficient soil locations and KBD-distribution (KBD=musculoskeletal disease) in the population. This trace-element remains an essential component of antioxidant/anti-inflammatory-related proteins which seems to be indirectly involved in normal cartilage growth and homeostasis. In the USA, a clinical study showed strong evidence that Se-deficiency influences cartilage metabolism, inducing a favorable environment for knee osteoarthritis progression. The present work aims at better understand the Se-impact on articular tissue organization/function during cartilage maturation using an in-vitro model.

Method: In-vitro articular cartilage culture based on an accelerated maturation model was used. To induce maturation in immature cartilage explants, they were cultured in serum-free medium supplemented with growth factors during 21 days in classic or Se-depleted medium.

Result: Synchrotron X-ray fluorescence microscopy shows a fundamental recurrent pattern of Se-distribution with punctual localization at chondrocyte-matrix interfaces that may highlight a particular role of Se in chondrocyte function. From Fourier-Transform-Infrared-microscopy absorption maps, we observed that explants placed in Se-depleted medium display abnormal molecular distributions, potentially related to metabolic changes/disruption. In Se-absence, abnormal surface chondrocyte organizations, similar to patterns observed in early stage of osteoarthritis, and a stiffness reduction in cartilage surface (nanomechanical-tests/Atomic-force-microscopy) were observed.

Discussion: Se-deficiency induces morphological matrix changes during the fast maturation-like processes which could be related to cartilage degenerative-like morphology, and could potentially be associated with degenerative changes that occur in KBD-patients during childhood. This work highlights the Se-importance in articular cartilage maturation. Se deserves to be studied for future enhancement of regenerative/preventive treatments for specific musculoskeletal diseases with a metabolic component.

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O31 - Functional Analysis of Plant Hyperaccumulator Genes SpSultr1;2 and SpATPS2 via Microbial Expression

1. Selenium chemistry and geochemistry
1.4 Relationships of selenium between soils, water, and vegetation
Keywords: hyperaccumulation, selenate transporter, selenium assimilation

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Introduction: RNA sequencing revealed hyperaccumulator Stanleya pinnata to highly overexpress putative selenate transporter Sultr1;2 and ATP sulfurylase 2 (ATPS2), relative to nonaccumulator Stanleya elata. We address here whether S. pinnata and S. elata SULTR1;2 and ATPS2 also differ in kinetic properties.

Method: Sultr1;2 and ATPS2 from S. pinnata and S. elata were PCR-amplified from root cDNA libraries and sequenced. The Sultr1;2 genes were expressed in a Saccharomyces cerevisiae YSD mutant lacking sulfate transporters, using plasmid pYES2 with C-terminal MycHis tags. The Atps2 genes (without targeting sequence) were expressed in E. coli with N-terminal MycHis tags. Yeast expressing the plant Sultr1;2 genes were tested for sulfate and selenate uptake capacity, kinetics and competition. SpATPS and SeATPS are purified and ATPS kinetics characterized for selenate, sulfate, or both.

Result: SpSULTR1;2 and SeSULTR1;2 differ in 13 aminoacids (1.9%): in MSD3, 6, 10 (1 AA each), in C-terminal regulatory domain STAS (5 AA), in N terminus (2), and between MSDs (3). The Sultr1;2 genes restored YSD sulfate uptake capacity and enhanced selenate sensitivity and Se accumulation. Kinetic properties for selenate uptake did not differ between the plant proteins, nor did inhibition by sulfate. SpATPS2 and SeATPS2 differed mainly in the N and C terminus, including a stop codon in the SpATPS plastid transit sequence.

Discussion: The Se hyperaccumulation capacity of S. pinnata is caused by high expression of SpSultr1;2 and not by differences in Vmax or Km for selenate, or altered selenate:sulfate specificity. S. pinnata ATPS2 appears purely cytosolic, likely affecting plant S/Se metabolism.

Selected references
O32 - Influence of Sulfate on Selenium Uptake in Hyperaccumulator Stanleya pinnata and Non-Accumulators

1. Selenium chemistry and geochemistry
1.4 Relationships of selenium between soils, water, and vegetation

Keywords: hyperaccumulation, sulfur, uptake, gene expression

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Introduction: Some plant species called hyperaccumulators contain selenium (Se) up to 0.1- 1.5% of their dry biomass. These plants display high Se to sulfur (S) ratios in tissues. Our hypothesis is that they may possess specific mechanisms for Se uptake.

Method: In our study we used Stanleya pinnata (Brassicaceae) as the model hyperaccumulator, and S. elata and Brassica juncea as non-hyperaccumulator species. We studied the capacity of plants to absorb Se either over a long period or in the short-term using different selenate and sulfate concentration. Distinct sulfate pre-treatments were tested for their effect on Se uptake, and gene expression of sulfate transporters was assayed under varying ambient Se/S ratios.

Result: S. pinnata exhibited higher capacity to absorb selenate compared to non-hyperaccumulators in both long and short-term assays, particularly under high S ambient concentration. In the long-term period, high sulfate reduced Se uptake in all species, but the effect was more pronounced in non-hyperaccumulators. The inhibitory effect of high sulfate on Se uptake was also observed in the short-term. In this case, Se uptake was totally repressed in S. elata by high sulfate, while it was moderately decreased in S pinnata and B. juncea. S pinnata plants had maximum values of root Se/S ratio and constitutive up-regulation of several sulfate transporter genes.

Discussion: These findings suggest that selenate uptake in hyperaccumulators is less sensitive to the inhibition exerted by sulfate. Hyperaccumulation may be due to high expression of sulfate transporters and/or higher specificity of these transporters for selenate over sulfate.

Selected references
O33 - Selenolanthionine is the main selenium species of the hyperaccumulator plant Cardamine violifolia

1. Selenium chemistry and geochemistry
1.4 Relationships of selenium between soils, water, and vegetation
Keywords: speciation, HPLC-ICP-MS, HPLC-ESI-QTOF-MS, synthesis, water soluble

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Introduction: Selenium hyperaccumulation in plants involve different selenium species to be stored in a chemical form serving for the elimination of excess selenium from plant metabolism to protect plant homeostasis. Organic forms, volatile species and inorganic forms are all concerned in this process.

Method: The study aimed at the identification of the main selenium species of the selenium hyperaccumulator plant Cardamine violifolia. This plant is from the family of Brassicaceae and it is common in the highly seleniferous region of Enshi, China. A sample of this plant, containing 3.6 g Se / kg d.w. was prepared with several methods including ultrasonic assisted water extraction, proteolytic digestion, sodium sulphite and carbon disulphide extraction. The extracted selenium species were identified and quantified (when the relevant standard was available) with orthogonal chromatographic purification followed by either LC-ESI-QTOF-MS or LC-ICP-MS set-ups.

Result: The Cardamine violifolia sample did not contain in considerable amount any of the organic selenium species that are often formed in hyperaccumulator plants and the inorganic selenium content (mostly in the form of elemental selenium) accounted only for less than 20% of total Se content. The most abundant selenium compound, accounting for more than 50% of total Se showed the accurate mass, elemental composition and chromatographic behaviour of selenolanthionine, a selenium species that has never been unambiguously identified before from any selenium containing sample. The identification process was completed with chemical synthesis as well.

Discussion: Finding selenolanthionine as the main selenium species in a hyperaccumulating plant unveils a new way of metabolism in the selenium hyperaccumulation process.
O34 - Two selenium hyperaccumulators and their influence on their plant communities

1. Selenium chemistry and geochemistry
1.4 Relationships of selenium between soils, water, and vegetation
Keywords: hyperaccumulator, plants, soil selenium

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Introduction: Selenium hyperaccumulators are capable of concentrating and tolerating tissue Se levels as high as 15,000 mg/kg dry weight (DW), while most plants show toxicity from tissue concentrations in the range of 20-100 mg/kg DW. Through root and shoot turnover, hyperaccumulators change distribution, form and concentration of soil Se. How does this affect their plant communities?

Method: We used transect methods to survey the plant communities of the hyperaccumulators Stanleya pinnata (Brassicaceae) and Astragalus bisulcatus (Fabaceae), as well as communities near, but without, hyperaccumulators. Both sites were also surveyed for soil and plant Se concentrations.

Result: In areas with Se hyperaccumulators, canopy cover was lower and % bare ground higher, yet plant species richness was higher. These hyperaccumulator effects depend on spatial scale and are much less evident at larger scale. Some plant species show positive or negative co-occurrence with hyperaccumulators. Several species that positively co-occurred showed extreme Se tolerance and contained more Se when growing next to hyperaccumulators (sometimes >1000 mg/kg DW).

Discussion: Most plants are sensitive to tissue Se concentrations from 20 to 100 mg/kg (DW). Perennial hyperaccumulators S. pinnata and A. bisulcatus appear to have a local effect on plant communities, as they over time concentrate, transform and redistribute Se through root and shoot turnover. Their influence appears most pronounced for nearby plant communities, and may exert selection pressure, Se-tolerant populations becoming more abundant and sensitive less abundant. They may also select for alleles involved in Se tolerance and thus the seed pool in the greater community.

Selected references
O35 - The effect of different Selenium fertilizers on garlic

1. Selenium chemistry and geochemistry
1.4 Relationships of selenium between soils, water, and vegetation

Keywords: Se-enriched slate powder, Cultivation, Cadmium

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Introduction: Selenium (Se) is an essential element for human and crops are believed to be the main sources of Se. Thus, understanding the effect of different Se fertilizers on vegetables is the premise of Se agro-enhancement. In this study, the effect of different Se fertilizers on garlic and the relationship between Se and Cd were studied.

Method: Garlic was grown in soils with different dose of activated Se-enriched slate powder as fertilizer in 2015 and with sodium selenite in 2014. Total contents of cadmium (Cd) and Se in the garlic were determined by ICP-MS and HG-AFS, respectively.

Result: 1. Both the activated Se-rich slate and sodium selenite were effective. The highest Se concentration of the garlic with the Se-fertilizer was about 4-times higher than the blank group (Se0). When the dose (Se) is less than 70 mg/m² (Se70), Se-enriched slate powder was more efficient, and when the dose (Se) is larger, sodium selenite was more valid (Fig.1 a).
2. Se and Cd concentration of garlic treated by activated Se-enriched slate powder showed a synergistic effect when the dose (Se) is less than 35mg/m²(Se35) and antagonistic effect when the dose (Se) is larger than 35mg/m²(Se35) (Fig.1). When treated by sodium selenite, this trend still presented. But Cd concentration of garlic treated with Se-enriched slate powder was always lower than that of garlic treated with sodium selenite (Fig.1 b).

Discussion: Se-enriched mineral powder has the ability to detoxify the garlic grown under Cd stress, which may be used to remediation of Cd poisoning soil.

Selected references
O36 - Biofortification of maize and beans with selenium in central Kenya highlands

1. Selenium chemistry and geochemistry
1.4 Relationships of selenium between soils, water, and vegetation

Keywords: Selenium deficiency, biofortification

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Introduction: Dietary Se supply for Kenya is estimated between 23-35 µg capita-1 d-1 and risk of dietary Se deficiency between 91-100 % (1). Narrow food choices exacerbates the suboptimal Se intake further increasing the deficiency risk (2). Selenium interacts with zinc (Zn), and iodine (I) which are common mineral deficiencies in Sub-Saharan Africa (3). Selenium status assessment in Kenya identified four regions with low soil Se concentration; 0.227, 0.233, 0.249, 0.323, mg Se kg-1. Staple foods from these regions recorded lowest Se concentration; maize at 0.035, and beans 0.040 mg Se kg-1.

Method: Biofortification trials were conducted to determine crops response: Soil application dosages for Se, Zn, and I were 0, 5, 10, 20 g ha-1; 0, 25, 50 kg ha-1; and 0, 5, 10 g ha-1 respectively. A factorial design was used for foliar application: 4*Se (0, 5, 10, 20 g ha-1), 3*Zn (0, 2, 4 kg ha-1), 3*I (0, 5, 10 g ha-1), 2*replicates.

Result: Se concentration in maize and beans gradually increased with increased Se fertilizer dosage. Crops response was best for foliar application. Beans responded significantly better compared to maize. Application of DAP did not improve Se uptake by crops. A combined fertilizer application for Se, Zn, and I increased Se uptake in maize and beans for both soil and foliar applications

Discussion: The above results will determine the necessary Se application dosage for a human intervention trial in 2017 and whether Se deficiency can be addressed alongside other mineral deficiencies through agronomic biofortification

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O37 - Selenophosphate synthetase 1 regulates cellular redox state and cell defense system
2. Selenium in the molecular life sciences
2.4 Selenoprotein synthesis pathways
Keywords: selenophosphate synthetase, reactive oxygen species, cell growth, cancer

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Introduction: Selenophosphate synthetase (SPS) was initially detected in bacteria and was shown to synthesize selenophosphate, the active selenium donor. However, higher eukaryotes including mammals and Drosophila have two SPS paralogues, which are designated SPS1 and SPS2. Between these two isoforms, only SPS2 has the catalytic activity of selenophosphate synthesis. Interestingly, SPS1 has been known to play an essential role in the cell, although it does not contain selenophosphate synthesis activity.

Method: SPS1 deficiency was induced by targeting the removal of SPS1 mRNA or its gene either by RNAi technology or by gene knockout technology. Cell growth and morphology were observed. Reactive oxygen species (ROS) levels were measured. Differentially expressed genes were identified from microarray analysis and pathways that could be regulated by SPS1 were predicted and validated by employing various molecular and biochemical analyses.

Result: Induction of SPS1 deficiency in Drosophila, mouse and human cell resulted in common phenotypic changes: the inhibition of cell growth/proliferation and accumulation of reactive oxygen species (ROS), specifically and the growth/proliferation inhibition was caused by ROS. Therefore, it seems that the primary function of SPS1 is to regulated intracellular ROS levels. SPS1 deficiency also activated genes participating in cell defense. However, the mechanisms to control ROS scavenging cell defense system are different from cell to cell.

Discussion: Our observations suggest that SPS1 regulates intracellular redox state and the ROS levels affect cell viability and/or cell defense system.

Selected references
O38 - Translation regulation of human selenoproteins

2. Selenium in the molecular life sciences
2.4 Selenoprotein synthesis pathways
Keywords: Selenoprotein, glutathione peroxidase, oxidative stress, senescence

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Introduction: Selenium is incorporated as a rare aminoacid, selenocysteine (Sec), into an essential family of antioxidant enzymes, the selenoproteins. Selenocysteine is co-translationally inserted using a UGA codon, normally used to stop protein synthesis. Several cis- and trans-acting components of the selenocysteine insertion machinery have yet been identified, including the stem-loop-stem-loop structure in the 3'UTR of selenoprotein mRNAs referred to as SECIS element.

Method: Our goal is to evidence whether the SECIS element modulates the efficiency of UGA-selenocysteine recoding in response to various stimuli. This elongation step is indeed supposed to be limiting for selenoprotein expression. We used a battery of luciferase based reporter constructs, containing an in frame UGA codon, to evaluate how selenocysteine insertion is regulated in response to selenium variation, oxidative stress, aging and viral infection.

Result: The regulation of the selenoproteome has been characterized in response to oxidative stress, cellular senescence and selenium bioavailability. We confirmed that most of the regulation of the selenoproteome occurs at the level of translation, and more precisely at the UGA-selenocysteine stage.

Discussion: Our work support a translational control of selenoprotein expression at the level of selenocysteine insertion. We confirm that the SECIS element, together with its interacting partner, plays a key role in the translational control of the selenoproteome.

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O39 - Structural and mechanistic insights into the mechanism of decoding of the Sec UGA codon in humans
2. Selenium in the molecular life sciences
2.4 Selenoprotein synthesis pathways
Keywords: selenocysteine, selenoprotein, tRNAsec, eEFSec, UGA

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Introduction: Selenocysteine (Sec) is encoded by an in-frame UGA codon across kingdoms. Instead of operating as the translational stop codon, the Sec UGA codon mediates the co-translational incorporation of Sec into nascent selenoproteins. This process requires a specialized translation elongation factor, eEFSec in eukaryotes and SelB in prokaryotes. The mechanism by which eEFSec facilitates decoding of Sec UGA and selenoprotein synthesis is not well understood.

Method: In order to advance our understanding of mechanisms governing co-translational incorporation of Sec, we characterized the major functional states of human eEFSec using X-ray crystallography, small-angle X-ray scattering, molecular modeling, and in vitro binding and functional assays.

Result: Our structural results revealed that four domains of human eEFSec fold into a chalice-like structure (Fig. 1A). Domains 1-3 resemble EF-Tu and the C-terminal domain 4 adopts the OB-fold structure. The binding assays showed that eEFSec has similar binding affinities towards GDP, GTP, and GTP analogs. Further, the activity assays established that the ability of eEFSec to promote decoding is sensitive to mutations in the GTPase site and the Sec-binding pocket. Surprisingly, unlike in eEF1A and EF-Tu, the guanine nucleotide exchange did not induce a canonical conformational change in domain 1 of eEFSec, but instead it caused a swing of domain 4 (Fig. 1B).

Discussion: We propose that eEFSec employs a non-canonical mechanism for the release of Sec-tRNA Sec during decoding on the ribosome. Given its dependence on the dynamics of domain 4, we suggest that the eEFSec-based mechanism may be similar to that of the universal initiation protein factor IF2/eIF5B.

Selected references
O40 - Identification of determinants regulating processive Sec incorporation in Selenoprotein P (SELENOP)

2. Selenium in the molecular life sciences
2.4 Selenoprotein synthesis pathways
Keywords: SECIS, Processivity, Efficiency, SELENOP

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Introduction: Selenoprotein P (SELENOP), is an interesting selenoprotein because it contains multiple selenocysteine (Sec) residues and two SECIS elements in its 3′-untranslated region (UTR). Significant progress has been made toward deciphering the mechanism of single Sec incorporation, but processive Sec incorporation still poses a conundrum. Interestingly, the SELENOP 3′UTR is highly conserved, especially SECIS-1. Prior work showed that SECIS-1 but not SECIS-2 is capable of processive Sec incorporation. We hypothesize that the conserved SECIS-1 may either contain sequence determinants that specifically dictate processivity or recruit ribonucleoprotein (RNP) complexes to modulate processive Sec incorporation in cells.

Method: We used biochemical and molecular biology approaches such as deletion mutagenesis, chimeras, qRTPCR, affinity chromatography and CRISPR gene editing in cultured cells.

Result: In this study, we compared SELENOP SECIS-1 and 2 to a non-SELENOP SECIS from Gpx4 for their ability to perform processive Sec incorporation. We found that both GPX4 and SELENOP SECIS-1 are capable of processive Sec incorporation in vitro while SECIS-2 terminated at second UGA. However in cells, GPX4 SECIS is defective for processivity despite sufficient RNA abundance, strongly suggesting a role for processivity regulation by cellular factors. We are currently investigating RNP complexes recruited by SECIS-1 and identifying sequence determinants within SECIS-1 that dictate processivity.

Discussion: SELENOP serves as the major source of selenium into the brain and solely for testes. Understanding the determinants of processive Sec incorporation will allow us to modulate its expression. This study provides new insights into basic questions regarding processive Sec incorporation in SELENOP and its mechanistic aspects.

Selected references

O41 - Codon-specific roles for cis-acting elements during translation of selenoprotein P

2. Selenium in the molecular life sciences
2.4 Selenoprotein synthesis pathways

Keywords: Recoding, Ribosome profiling, RNA structure, SECIS

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Introduction: Gene-specific expansion of the genetic code allows for UGA codons to specify the amino acid selenocysteine (Sec). A striking example of UGA recoding occurs during translation of the mRNA encoding for the selenium transport protein, selenoprotein P, which in vertebrates may contain up to 22 UGA codons.

Method: Here we present data from phylogenetic analyses, reporter assays, and ribosome profiling to support distinct codon-specific roles for multiple cis-acting elements located within the coding sequence and 3’ UTR of the mRNA for selenoprotein P.

Result: The data supports a model originally presented by Stoycheva, Tujebajeva, Harney and Berry (2006) wherein the 3’ UTR proximal selenocysteine insertion sequence (SECIS1) supports Sec incorporation at all UGA codons downstream of the first UGA, whereas the distal SECIS element (SECIS2) effects Sec incorporation at the first UGA. Further, we describe increased probability of nucleotide pairing across the first and last ~20% of the coding sequence, identify a cis-acting element near the start codon that overlaps the signal peptide sequence and impacts translation initiation, and a secondary structure downstream of the first UGA codon that impacts recoding of that codon. Additional, discreet structures of unknown function are found in proximity of the region containing multiple UGA codons.

Discussion: Selenoprotein P has evolved multiple cis-acting elements in both the coding sequence and 3’ UTR that coordinately act to facilitate and regulate multiple occurrences of UGA recoding. Selenoprotein P provides a unique example of how cis-acting elements may alter ribosome decoding during translation of different regions within the same mRNA.

Selected references

Cis-acting elements affecting Selenop translation. A) Red arrows indicate a negative effect of translation initiation, green arrows indicate stimulators of recoding and the location of action along the mRNA. B) Schematic of Selenop mRNA cds (rectangle) and UTRs (line), UGA codon positions (red vertical lines), signal peptide (sp), and two Secis elements. The probability of internal RNA pairing is enriched near the 5’ and 3’ ends of the CDS.
O42 - Coding region determinants regulate Selenoprotein P (SELENOP) expression in cells

2. Selenium in the molecular life sciences
2.4 Selenoprotein synthesis pathways

Keywords: SELENOP, processivity, RNA stability, ribonucleoprotein complexes

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Introduction: Selenoproteins incorporate the 21st amino acid, selenocysteine (Sec) in response to a stop codon. Under selenium depleting conditions, when selenoprotein translation is dramatically inhibited, most selenoprotein transcripts are predicted targets for mRNA surveillance or decay pathways but many are resistant. Selenoprotein P (SELENOP) is one of 6 selenoprotein transcripts that display resistance to mRNA decay during low selenium. This study focusses on identifying determinants and mechanism that SELENOP employs to resist cellular RNA decay.

Method: We used several biochemical and molecular biology approaches such as deletion mutagenesis, chimeras and CRISPR gene editing in cultured cells.

Result: We focussed on the SELENOP coding region since 1) its highly conserved among vertebrates and 2) SECIS-deleted mutants do not effect RNA levels. In-frame coding region deletions within SELENOP N-terminal sequences reduced RNA abundance in cells by 10 to 35-fold even with adequate selenium. Similarly, fusing SELENOP native N-terminus sequence with an artificial Sec-containing C-terminus also increased RNA abundance. We are currently investigating the role of coding region ribonucleoprotein (RNP) complexes including SBP2 and key RNA regulation factors in SELENOP expression in selenium inadequate conditions.

Discussion: Upon selenium starvation, differential selenoprotein regulation occurs in a tissue specific manner. This allows production of essential selenoproteins critical for cell viability. Our study provides new mechanistic insight into how selenoproteins maintain hierarchy during limiting selenium.

Selected references


O43 - Effect of long-term selenium supplementation on mortality: results from a multiple-dose, RCT

3. Selenium in animal and human health and disease
3.3 Nutritional selenium intervention studies in human

Keywords: randomised controlled trial, Denmark, mortality, cancer mortality

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Introduction: Selenium is an essential trace element that is incorporated into selenoproteins with a wide range of health effects¹. Selenoprotein concentration plateaus at plasma selenium around 125 µg/L²; whether levels higher than this are beneficial, or otherwise, is unknown.

Method: In a double-blind, randomized, controlled trial of selenium supplementation and mortality in Denmark, a country of moderately-low selenium status, 491 men and women aged 60-74 years were treated with 100, 200, or 300 µg selenium/d as selenium-enriched yeast or placebo-yeast. From 1998-1999, active treatment was provided for 5 years and post-treatment follow-up for 10 additional years.

Result: During 6,871 person-years of follow-up, 158 deaths occurred (31 during active treatment, 127 after treatment cessation). The hazard ratio (95% confidence interval) for all-cause mortality comparing 300 µg selenium/d to placebo was 1.62 (0.66 to 3.96) after 5 years of treatment and 1.59 (1.02 to 2.46) over the entire follow-up period, rising to 2.20 (1.16 to 4.17) in those with baseline plasma selenium ≥ 82 µg/L. Selenium doses of 100 and 200 µg/d non-significantly decreased mortality during the intervention period but their effects vanished after treatment cessation. The effects on cancer and cardiovascular mortality were similar but less precise.

Discussion: A dose of 300 µg/d of selenium (as high-selenium yeast) taken for five years in a country with moderately-low selenium status increased all-cause mortality 10 years later. Lower doses showed a non-significant reduction in mortality which dissipated after treatment discontinuation. Total selenium intake (diet plus supplements) over 300 µg/d should be avoided.

Selected references

O44 - Selenium supplementation blocked the crosstalk among pathogenic pathways in Alzheimer's disease mice

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease

Keywords: Alzheimer's disease (AD); oxidative stress; protein phosphatase of type 2A (PP2A)

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Introduction: Oxidative stress can induce the imbalance of protein tyrosine kinase/phosphatase [1], leading to phosphorylation of protein phosphatase of type 2A (PP2A) on tyrosine 307 and thus inactivation of the enzyme in Alzheimer’s disease (AD). Selenium is well known for its antioxidative property in vivo [2], but its effect on AD has not been investigated systematically.

Method: Different forms of selenium, including sodium selenate, selenomethionine, ebselen and Se-methylselenocysteine were supplemented to the triple transgenic mouse model of AD (3×Tg-AD) in our laboratory. Primary neurons isolated from the hippocampus of AD mouse were cultured with different forms of selenium. The pathological hallmarks of AD were detected by immunohistochemistry, immunofluorescence and Western blot.

Result: Both inorganic and organic forms of selenium could ameliorate cognitive deficits of AD mice by attenuating tauopathic pathway, amyloid cascade and synapse loss [3,4]. Selenomethionine was further found to activate autophagy-based pathway [5]. Increased phosphorylation of PP2A on tyrosine 307 and decreased enzyme activity in AD brain were inhibited after selenium treatment, together with inhibition of GSK-3b activity and dephosphorylation of amyloid precursor protein on threonine 668 which respectively result in tau hyperphosphorylation and b-amloid generation. LB-100, an inhibitor of PP2A, could eliminate the effect of sodium selenate on AD in the cultured primary neurons of AD mice.

Discussion: These findings revealed that supplementation of selenium could enhance PP2A activity in vivo through antioxidation, leading to activation of Wnt/β-catenin signaling and blockage of the crosstalk among Aβ generation, tau hyperphosphorylation and neuronal apoptosis. This work was financially supported by NSFC (No. 31470804).

Selected references
O45 - Selenium, coenzyme Q10 and cardiovascular health – results from a 4-year-intervention in elderly

3. Selenium in animal and human health and disease
3.3 Nutritional selenium intervention studies in human

Keywords: Coenzyme Q10, deficiency, supplementation, health effects

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Introduction: Selenium is essential for the cell. However, in Europe selenium soil content is low. Coenzyme Q10 is an important anti-oxidant, and the two are interrelated to get optimal cellular function.

Method: 443 healthy elderly participants were randomized to receive supplementation with selenium and coenzyme Q10 or placebo during 4 years. Blood samples, echocardiography, and quality-of-life registrations were obtained. Underlying mechanisms were traced by use of laboratory biomarkers.

Result: A follow-up period of 5.2 years showed reduced cardiovascular mortality, better cardiac function and improved physical and emotional quality-of-life. Specific biomarkers (sP-selectin, hs-CRP) showed less inflammatory activity, less oxidative stress (copeptin, MR-proADM), increased levels of IGF-1, and less fibrosis. MicroRNA analysis demonstrated significant difference in expression, presumed to constitute an underlying mechanism for other observations.

Discussion: We propose a model illustrating relations between observed laboratory effects and clinical endpoints (Figure). It is seen that a suboptimal selenium status increases cardiovascular risk, which was attenuated by the supplementation. The intervention influenced the expression of microRNA, the level of inflammatory activity, oxidative stress, IGF-1 levels and fibrosis activity. Apparently, the intervention improved cardiomyocyte function reflected by better cardiac function and less NT-proBNP biomarker concentration. We also found a reduced cardiovascular mortality after 10 years although the intervention was limited to 4 years. The proposed model illustrates the complex relationship between some of the effects resulting in the positive clinical endpoints.

Selected references
O46 - Antibacterial redox selenium coatings, covalent small molecules and antibody drug conjugates (ADCs)

3. Selenium in animal and human health and disease
3.4 Selenium based medical therapeutics

Keywords: Selenium Toxicity, Selenium Coatings, Selenium Antibodies

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Introduction: Selenium (Se) has a 100+ year history as an anti-cancer element both before and after its nutritional requirement was discovered by Schwarz (1957). In the 1960’s debate ensued over the experimental reports of selenium’s anti-cancer attributes in animals and humans. The famous 1996 Clark study of human cancer with Se-yeast and failure of the SELECT Trial (Lu, 2016) have seemingly dashed hope of dietary selenium supplementation ever having an anti-cancer future. With understanding selenium redox chemistry, selenides (Spallholz, 2015) and isoselenocyanates (Crampsie, 2012) can be covalently “targeted” and delivered to cells offering a renewed assessment of selenium’s role as a therapeutic treatment, especially for cancer.

Method: Selenium toxicity research really begins in the 1930’s among horses and sheep on the Great Plains of the USA by Moxon (1937). The first chemical equation of WHY selenium is toxic, superoxide generation, was by Seko (1989) for selenite (Figure1). Understanding this chemistry by organic selenium compounds permits them to be incorporated into polymers, covalently attached to “targeting” small molecules, and recently clinical antibodies, Herceptin® and Avastin®.

Result: Attachment of redox selenium to proteins is accomplished with a Se-modified Bolton-Hunter reagent, as developed in the 1970’s for ¹²⁵I-radioimmunoassays (Lane, 2011). This chemistry permits antibodies to deliver redox selenium, generate superoxide inducing oxidative stress and cell death. Using redox probes, Se-antibody treated cancer cells are observed to generate intracellular superoxide and H₂O₂.

Discussion: Redox selenium and its applied “targeted” therapeutic applications require a renewed assessment of its anti-cancer and medical applications against viruses and antibiotic-resistant bacterial infections.

Selected references
O47 - Selenium in the treatment of cancer

3. Selenium in animal and human health and disease
3.4 Selenium based medical therapeutics

Keywords: Redox-active selenium compounds, ex vivo, Pharmacokinetics, Chemotherapy, Clinical trials

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Introduction: The prognosis for visceral malignancies is poor, especially for carcinoma of the lung, pancreas and liver due to resistance to chemotherapy. Over several decades, a plethora of preclinical investigations have demonstrated high degree of tumor-specific cytotoxic effects of selenium compounds, especially to therapy-resistant tumor cells [1]. Herein, we have undertaken translational approaches to establish the efficacy of selected selenium-based candidate chemotherapeutics into clinical practice.

Method: The efficiency of the potent candidate drugs are evaluated both in vitro and in vivo. In vitro studies span the spectrum of cell lines, isolated primary tumor cells and ex vivo tumor tissues in cultures. In parallel, we perform clinical trials with selected selenium compounds on cancer patients.

Result: Serum samples from the SECAR study on cancer patients [2] have been extensively analysed for selenium compounds. Selenite was detected in serum and selenosugar was the major organic metabolite. Pharmacokinetic analyses indicated that plasma selenite concentrations increased in a linear dose-dependent manner. An unprecedented increase in plasma SEPP level was recorded. In another study, we documented favourable anti-cancer effects of an organic selenium compound in liver cancer models. A novel target specific drug release principle might serve as a promising therapeutic intervention in liver cancer and a phase I clinical trial is underway.

Discussion: Implementation of selenium-based candidate chemotherapeutics in clinical practice underscores the absolute requirement of both safety and efficacy evaluations. To this end, our founding clinical studies can greatly accelerate the transition of selenium compounds from experimental drugs to clinically relevant chemotherapeutics.

Selected references
O48 - Selective targeting of redox dysregulation of cancer cells by redox-active selenium compounds

**3. Selenium in animal and human health and disease**

**3.4 Selenium-based medical therapeutics**

**Keywords:** Redox balance, redox-active selenium compounds, thiols, animal model, leukemia

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**Introduction:** Altered redox homeostasis is an acquired physiological adaptation of proliferating malignant cells. An increased cellular proliferation necessitates high energy supplies which culminate into elevated reactive oxygen species (ROS) production. Malignant cells counterbalance the cytotoxic effects of elevated ROS levels by augmenting activities of ROS-detoxifying pathways. However, such adaptive mechanisms compromise the cellular tolerance to further increase in intracellular ROS levels. Herein, we investigated whether further manipulations of cellular redox balance could inflict selective cytotoxic effects on malignant cells by two redox-active selenium compounds implicated in ROS generation.

**Method:** Sixteen different cancer cell lines and isolated primary normal cells were used for screening purposes. We employed cell viability assays, gene and protein expression analyses, knock-down and overexpression of target genes to investigate the underlying mechanisms of cytotoxicity.

**Result:** The selected selenium compounds were cytotoxic to all the tested cancer cell lines, however, with varying efficacy. Small molecules-induced specific modulations of cellular redox homeostasis resulted in dramatic potentiation (up to 70-fold) of the cytotoxic effects of these selenium compounds, independent of the origins or oncogene status of these cells. No such potentiation of cytotoxic effects was observed in normal cells under identical test conditions. Gene and protein expression analyses indicated involvement of key redox-regulatory enzymes in the observed cytotoxic effects on cancer cells. Animal studies are underway to investigate the *in vivo* efficacy of these compounds in experimental cancer models.

**Discussion:** Specific modulations of key redox-regulatory pathways of cancer cells can greatly potentiate the cytotoxicity of redox-active selenium compounds with promising cancer chemotherapeutics applications.
O49 - Selenium: A potentially powerful tool to design potent anticancer molecules-Discovery of Se-Aspirins

3. Selenium in animal and human health and disease
3.4 Selenium based medical therapeutics
Keywords: Se-Aspirin, cancer, apoptosis, NF-κB, ROS

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Introduction: We have shown that rational incorporation of Se into small molecules can substantially enhance their anti-cancer efficacy1-5. Recently, through extensive SAR studies based on Se incorporation into NSAIDs, we identified a Se-aspirin hybrid molecule, which was >300 times more potent than aspirin in inhibiting viability of various cancer cells5.

Method: Further SAR studies and investigations directed towards efficacy, toxicity, and drug-likeness has now led to the identification of a novel Se-Aspirin molecule AS-10 as the most efficacious compound.

Result: AS-10 (47 mg/kg, i.p) inhibited subcutaneous colon tumor growth by ~70% and was lethal to a variety of cancer cells including pancreatic cancer (PC) for which no effective therapy currently exists. AS-10 inhibited PC cell growth (IC50 2.5-5 µM), induced G1/G2 cell cycle arrest which was associated with increase of cell cycle inhibitory proteins p21 and p27, and induced apoptosis as evidenced by caspase 3/7 activity, PARP cleavage and Annexin V staining. AS-10 inhibited NF-κB DNA binding activity as well as NF-κB translocation to the nuclei upon stimulation by TNFα. Notably, AS-10 potentiated cytotoxic activity of gemcitabine in PC cells. Furthermore, in LNCaP prostate cancer cells, AS-10 decreased protein level of AR and its best known target PSA, and led to increased caspase-mediated apoptosis and expression of p53-DNA damage response proteins such as p21 and p-H2A.X. AS-10 induced ROS in cancer cells as likely primary biochemical event.

Discussion: AS-10 may represent a promising intervention and therapeutic agent for pancreatic, prostate, and colon cancers and other cancer types through multiple targets.

Selected references
O50 - Selenium Distribution and its Correlation to Geochemical Factors in East China Intertidal Zone

1. Selenium chemistry and geochemistry
1.4 Relationships of selenium between soils, water, and vegetation

Keywords: Selenium distribution, Intertidal zone, East China

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Introduction: The east China intertidal zone (ITZ) has a total area of 11056.48 km², about half of the ITZ in China. In this paper, we are focused on the distribution of selenium (Se) in the east China ITZ and its correlation with other geochemical factors.

Method: 1906 soil/mud samples had been taken by China Geological Survey from the east China ITZ, which includes, from north to south, 1199 samples from Jiangsu Province, 102 from Shanghai municipal city, 252 from Zhejiang Province and 353 from Fujian Province, and 54 geochemical indexes had been measured.

Result: 1. Se concentration in the east China ITZ was quite low, with a range of 0.01-0.50 mg/kg and a mean value of 0.1-0.16 mg/kg, Jiangsu Province being the lowest, with a range of 0.01-0.20 mg/kg and a mean value of 0.06 mg/kg, reflecting that East China terrestrial landmass is severely selenium deficient.
2. Se concentrations in ITZ soils are found to be positively correlated to the content of organic carbon (OrgC), aluminum and iron, but negatively correlated to pH values (Fig. 1).

Discussion: Absorption by organic materials, clay minerals and iron oxides seems to be the mechanism of Se accumulation in the ITZ soils and the low contents of OrgC, aluminum and iron and high pH values in East China ITZ soils may another reason to explain its low concentration of Se.

Fig. 1 Scatter plots of Se with other geochemical indexes in East China ITZ to show their correlation
O51 - Distribution and translocation of selenium from soil to highland barley in Kashin-Beck disease area

1. Selenium chemistry and geochemistry
1.4 Relationships of selenium between soils, water, and vegetation

Keywords: Highland barley; Translocation; Kashin-Beck disease

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Introduction: Kashin-Beck disease (KBD), which is still active in the Tibetan Plateau, is considered to be relevant to insufficient selenium intake. Highland barley as the most popular staple food in Tibetan Plateau KBD areas is one of the dominant selenium sources for local people.

Method: In this study, samples of intact highland barley plant and corresponding topsoil were collected from both non-KBD and KBD endemic areas of Songpan County. Total selenium content in plant and soil samples and fractions of soil selenium were determined and analysed.

Result: The results showed that selenium levels in different fractions of highland barley were lower in KBD areas than non-KBD areas (grain $P = 0.090$; straw $P = 0.131$; root $P = 0.001$), while no significant difference was observed in their corresponding soil ($P = 0.993$). The comparison of selenium transfer factors indicated that the restricting step for selenium translocation was from soil to root. Water-soluble, exchangeable and fulvic acid-bound selenium fractions in soil were the key fractions dominating in this transfer process. Stronger positive correlation was observed between tetravalent selenium speciation and plant selenium, suggesting the closer affinity to selenite for highland barley. Additionally, selenium transfer from soil to root significantly increased as the soil pH increased ($P = 0.007$) and the soil organic matter content decreased ($P = 0.019$).

Discussion: The information obtained may have considerable significance for understanding the patterns of selenium transfer from soil to grain in low-selenium areas, providing a theoretical and experimental basis for proposing effective agricultural measures to increase crops selenium in KBD endemic areas.

Selected references
O52 - The status of selenium in Iraqi Kurdistan and feasibility of Se bio fortification using 77-Se

1. Selenium chemistry and geochemistry
1.4 Relationships of selenium between soils, water, and vegetation

Keywords: Soil, Plant, biofortification, Iraqi Kurdistan

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Introduction: Selenium (Se) is an essential trace element for human and animals. Since selenium is absorbed initially by plants and consumed by animals. Consequently, soil can be accounted as preliminary driver of human selenium status. No data exists on the Se status of the population of Iraq. To the aim of this work was to investigate Se content of soils, plants and waters in Iraqi Kurdistan and establish if Se biofortification is desirable.

Method: A total of 97 soil, 300 plant and 20 water samples were collected. Soil characteristics, total Se, and Se fractionation were measured. Plant uptake experiments were also conducted using a 10 g ha^{-1} 77-Se spike. Total Se and Se speciation was determined by HPLC-ICP-MS.

Result: Total soil Se ranged from 179-617 µg kg^{-1} (average 309 µg kg^{-1}) less that the global average (Figure 1a). Less than 1% of the total soil Se was in the soluble fraction, 2% was absorbed and 45% was organic Se (TMAH extractable). Plant Se ranged from 4-678 µg kg^{-1} with greater concentrations in vegetables grown in higher pH soils (Figure 1b). Vegetable species had different 77-Se uptakes (Figure 1c). Post-harvest 60% of the inorganic 77-Se spike had been converted to organic Se. Soil to plant transfer factors ranged from 7-50 fold greater than for natural soil Se. Total recovery of 77-Se in the biomass was 4-24 %.

Discussion: Soils in Iraqi Kurdistan are typically below global average Se concentrations and in regions where soil pH is < 8 Se biofortification may be advisable.
O53 - Geochemistry of selenium in Gilgit-Baltistan (North East Pakistan)

1. Selenium chemistry and geochemistry
1.4 Relationships of selenium between soils, water, and vegetation

Keywords: soil, plants

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Introduction: Selenium (Se) is an important micronutrient for human and animal health. It is essential for normal functioning of the thyroid gland and thyroid hormone synthesis.

Method: Samples of soil and wheat grain were collected from Gilgit-Baltistan (N. Pakistan). Samples were digested in nitric acid and extracted in 10% TMAH prior to analysis for Se concentrations by ICP-MS.

Result: Soils were alkaline, with pH values c. 8.0, and low organic carbon contents (0.12% - 0.59%). Average total soil Se concentration was 290 µg kg⁻¹, 25% less than the global average (400 µg kg⁻¹).

Discussion: Only 5% of the soil Se was extractable by TMAH suggesting that the majority of the Se is not associated with soil organic matter, or as surface-adsorbed selenite or selenate, but exists in a poorly available form. No relationship was observed between soil organic carbon and total soil selenium concentrations but soil pH was negatively correlated with TMAH-extractable Se. Selenium concentration in wheat grain was exceptionally low (1-95 µg kg⁻¹). Considering that wheat consumption in the region (~350 g) provides 75% of calorie intake, it is likely that the population does not receive sufficient Se from their diet in the absence of supplementation.

Further investigation of locally grown foods, including vegetables, fruit, meat and dairy products, is required to fully audit dietary Se sources. We also plan to analyse human biomarkers for Se. From this data we will assess whether sufficient dietary Se is available or whether supplementation through Se biofortification of the staple crop (wheat) is required.

Selected references

O54 - Selenocysteine Chemistry and Total Chemical Synthesis Applied for Accessing Human Selenoproteins

2. Selenium in the molecular life sciences
2.4 Selenoprotein synthesis pathways
Keywords: Chemical protein synthesis, selenocysteine, selenopeptide, selenoprotein, native chemical ligation

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Introduction: Once considered a toxic element, selenium is now known as an essential element for life. For example, in humans it is incorporated in proteins known as selenoproteins containing the 21st encoded amino acid selenocysteine (Sec). The differences between selenium and sulfur in their redox potentials, pKas, and nucleophilicities and electrophilicities give selenium interesting chemistry.

Method: We use chemical protein synthesis as our main technique to access selenoproteins. This technology is based on two main methods, solid-phase peptide synthesis (SPPS) and chemical ligation reactions; most famous is native chemical ligation (NCL).

Result: Sec can be used as a tool for chemical protein synthesis, and allowing for site selective modifications. It can be also incorporated into protein sequences to enhance oxidative protein folding.

Discussion: This lecture will discuss our recent studies on chemical protein synthesis using Sec and selective deselenization reactions, which convert Sec into Ala or Ser. These advances in chemical protein synthesis bring us closer to accessing naturally occurring selenoproteins, especially human selenoproteins that still await functional characterization.

Selected references
O55 - Redefining UAG for selenocysteine widens the scope of recombinant selenoprotein production in E.coli

2. Selenium in the molecular life sciences
2.4 Selenoprotein synthesis pathways

Keywords: Recombinant selenoprotein, E coli, Genetic code, Thioredoxin reductase, Glutathione peroxidase

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Introduction: Selenoproteins, containing one or several Sec residues, are found in all three domains of life (1). It is important to characterize selenoprotein functions in order to understand their roles in life, but scarce availability of selenoproteins has remained a major bottleneck for such research. Here we illustrate a surprising flexibility of selenoprotein synthesis, which opens up a powerful method for selenoprotein research using recombinant selenoproteins (2).

Method: In nature, Sec is co-translationally inserted at predefined UGA opal codons by specific translation machineries (3). Here we found that Sec can be equally efficiently incorporated at predefined UAG amber codons, thereby competing with RF1 rather than RF2. By utilizing this system within a RF1-depleted E. coli strain C321.AA (4) we could produce a number of selenoproteins with unsurpassed purity and yield.

Result: We have here produced several form of mammalian TrxR isoenzymes with much higher specific activity compared to previous methods (5). We believe this new routine could be widely employed for production of a wide variety of selenoproteins carrying penultimate or ultimate Sec residues, as in TrxRs. Interestingly, we also found that a SECIS element was no longer absolutely required in our system. This also allowed production of selenoproteins with internal Sec, such as human GPx1, and we could confirm a previously proposed catalytic tetrad in the enzyme using directed mutagenesis (6).

Discussion: We conclude that our new methodology (2) should constitute a powerful resource enabling many forthcoming studies of recombinant selenoproteins, considering its versatility and comparable ease of use.

Selected references

O56 - Delivery of selenide to selenophosphate synthetase for selenoprotein biosynthesis in bacteria

2. Selenium in the molecular life sciences
2.4 Selenoprotein synthesis pathways

Keywords: selenophosphate, thioredoxin, bacteria, selenoprotein biosynthesis, selenide

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Introduction: Selenophosphate synthesized by selenophosphate synthetase (SelD) from selenide (HSe⁻) and ATP is essential for selenoprotein biosynthesis. In mammals, glutathione (GSH) and thioredoxin reductase (TrxR) have been proposed to serve as a HSe⁻ donor for selenophosphate synthesis. However, little is known about the HSe⁻ donor to SelD in bacteria. Here, we investigated the mechanism of HSe⁻ generation and HSe⁻ supply to SelD in the selenoprotein biosynthesis in bacteria.

Method: To identify the genes/proteins that affect selenoprotein biosynthesis in E. coli, 57 candidate genes potentially participating in selenite reduction, such as those encoding nitrate reductase, glutaredoxin, and GSH-related enzymes, were selected. The mutant strains defective in each of the 57 candidate genes were tested for the ability to synthesize selenoprotein by measuring formate dehydrogenase (FDH) activity.

Result: A significant decrease of FDH activity was seen in a thioredoxin (TrxA)-deficient mutant, whereas the deficiency of genes related to GSH and other reductases did not affect the FDH activity. To clarify the function of TrxA in selenite reduction, the purified recombinant TrxA of Pseudomonas sp. F2a was analyzed. We found that TrxA exhibited selenite-reducing activity coupled with a TrxR reaction and that HSe⁻ generated by TrxA was efficiently supplied to SelD for selenophosphate synthesis.

Discussion: Our data provide the first evidence for a pivotal role of TrxA in selenoprotein biosynthesis in bacteria.
O57 - Reconstitution of processive selenoprotein P synthesis in the wheat germ lysate system

2. Selenium in the molecular life sciences
2.4 Selenoprotein synthesis pathways
Keywords: Selenoprotein P, SECIS Binding Protein 2, wheat germ lysate, in vitro, translation

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Introduction: A UGA stop codon is recoded to accommodate the incorporation of the 21st amino acid selenocysteine (Sec). For UGA to be recoded a specialized set of cis and trans factors are required and consist of: an mRNA with an in frame UGA codon, a selenocysteine insertion sequence (SECIS) in the 3’ untranslated region, a SECIS binding protein 2 (SBP2), a specific translation elongation factor (eEFSec), and a selenocysteine tRNA. The N-terminus of SBP2 is believed to have no direct role in Sec incorporation because the C-terminus of SBP2 is sufficient for the incorporation of Sec into selenoproteins that have one Sec codon. Having recently developed an in vitro translation system in wheat germ requiring the addition of all Sec-incorporation factors, we found that translation of SELENOP, which contains 10 Sec codons, is defective in that only early termination products are made. This contrasts with mammalian systems where full length protein is predominantly found. This observation, combined with the fact that neither SELENOP nor the N-terminus of SBP2 are found in invertebrates, points to a specialized function for the N-terminus of SBP2 related to SELENOP synthesis.

Method: To validate the relationship between the N-terminus of SBP2 and SELENOP, C-terminal and full length SBP2 were compared for in vitro translation of SELENOP.

Result: We found that the production of full length SELENOP is dependent on the presence of the SBP2 N-terminus.

Discussion: Our results suggest that the N-terminus of SBP2 was appended during evolution to allow for processive incorporation of multiple Sec residues into SELENOP.

Selected references
O58 - Selenium in Radiation Oncology – 10 years of Experiences in Germany

3. Selenium in animal and human health and disease
3.4 Selenium based medical therapeutics

Keywords: Selenium Supplementation, Radiation Oncology, Tumor Patients, Lymphedema, Radioprotection

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Introduction: Se supplementation in radiation oncology is a controversial issue. We summarize the major results obtained by the German Working Group Trace Elements and Electrolytes in Oncology (AKTE).

Method: We analyzed blood and tissue Se-levels of tumor patients (n=170) before and after Se-supplementation.

Result: The majority of tumor patients (carcinomas of the uterus, head and neck, lung, rectal or prostate cancer) showed a relative Se deficiency in whole blood and serum. In prostate cancer, tissue Se levels were lower in the compartment surrounding the carcinoma in comparison to patients with benign prostatic hyperplasia. Se supplementation successfully corrected Se-deficiency and decreased radiotherapy-induced diarrhea in a randomized study of radiotherapy patients with carcinomas of the uterus. Survival data imply that Se supplementation did not interfere with radiation success. Limited effects of supplemental Se in the prevention of ageusia (loss of taste) and dysphagia due to radiotherapy were noted in a second randomized trial in head and neck cancer. In comparison, positive effects of Se supplementation were detected on radiation-associated secondary lymphedema in patients with limb edemas as well as in the head and neck region, including endolaryngeal edema. No adverse effects of supplemental Se were noted.

Discussion: Se supplementation yielded promising results concerning radioprotection and edema prevention in tumor patients and should be considered as a meaningful adjuvant treatment option in subjects with a low Se status.

Selected references

O59 - Selenium in the treatment of radiation-associated secondary lymphedema - An Update

Keywords: lymphedema, radiation, oncology

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Introduction: Lymphedema can be observed after surgery and/or radiotherapy or as a result of tumor compression. The antiedematous effect of selenium is well-known, but not very well defined. The aim of this explorative study was to further evaluate the impact of selenium in the treatment of lymphedema after radiotherapy.

Method: Between June 1996 and December 2016, 32 patients with edema of the arm and 46 patients with edema of the head-and-neck region were treated with selenium for therapy-related lymphedema. Of these 46 patients, a total of 30 had interstitial endolaryngeal edema. All patients received sodium selenite (500 µg per day) over 4 to 6 weeks.

Result: Self-assessment using a visual analog scale showed a reduction of 4.3 points when comparing pre- and posttreatment values. Of 30 patients with endolaryngeal edema, 20 underwent no tracheostomy, 7 underwent a temporary tracheostomy, and only 3 underwent a permanent tracheostomy. Overall, 28 of 32 patients with arm edema showed a circumference reduction of the edematous limb and improvement in the Skin-Fold Index by 23.6 points. An improvement of one stage or more was shown by the Földi or the Miller score (n=28) in 22 (Földi score) and in 24 (Miller score) patients. Side effects of treatment were not observed.

Discussion: Treatment with sodium selenite is well tolerated, cost-effective and easy to deliver. Additionally, our results suggest that sodium selenite has a positive effect on secondary lymphedema after radiotherapy.

Selected references


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O60 - Selective targeting of leukemia stem cells by selenium

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Introduction: Supplementation with non-toxic doses of selenium, as selenite, inhibits the progression of chronic myelogenous leukemia (CML) in mice via the selective elimination of leukemia stem cells (LSCs). Use of non-steroidal anti-inflammatory drugs, which blocked the effect of selenite, have suggested a key role for the endogenous cyclopentenone prostaglandins (CyPGs), delta-12 prostaglandin J₂ (Δ₁₂-PGJ₂) and 15-deoxy-delta₁₂,₁₄-prostaglandin J₂ (15d-PGJ₂). Here we show that these CyPGs, produced by mice maintained on selenium supplemented diets (0.4 ppm Se), inhibit leukemic progression CML through their ability to activate the nuclear hormone receptor, peroxisome proliferator activated receptor gamma (PPARγ).

Method: C57BL/6 mice on diets with adequate Se or supplemented with Se were transplanted with LSCs expressing the fusion oncoprotein, BCR-ABL. Changes in WBC counts, flow cytometric evaluation of LSCs, splenomegaly, LSC-colony forming units, gene expression profiling, western immunoblotting, and LC-MS/MS analyses of serum and LSCs were performed.

Result: GW9662, a potent PPARγ-antagonist, blocked the anti-leukemic effect of selenium supplementation by significantly reducing CyPGs. Selenium supplementation decreased 15-prostaglandin dehydrogenase (15-Pgdh), which oxidizes and inactivates CyPGs. In contrast, pioglitazone, a PPARγ agonist, mimicked selenium supplementation and decreased 15-Pgdh activity leading to increased CyPG levels resulting in the inhibition of CML progression. Selenium-dependent activation of PPARγ mediated by endogenous CyPGs decreased Stat5a expression leading to the downregulation of Cited2, a master regulator of LSC quiescence.

Discussion: Our studies provide a novel method to eliminate LSCs in-vivo that are otherwise not targeted by existing therapies. Selenium-dependent inhibition of quiescence of LSCs via the activation of PPARγ was key to this process.

Selected references
NIH R01-DK077152 (KSP) and R01-DK080040 (RFP) and supported by USDA National Institute of Food and Agriculture Hatch Project #4736 (R.F.P.) and 4605 (K.S.P)
O61 - Methylselenol suppressed the metastatic potential of B16F10 melanoma by reducing integrin expression

3. Selenium in animal and human health and disease
3.7 Additional and emerging topics of selenium in health or disease
Keywords: methylselenol, metastasis, B16F10 melanoma, integrin expression

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Introduction: Our previous studies have shown that selenite inhibits the invasion of HT1080 fibrosarcoma cells by reduced expression of matrix metalloproteinase (MMP)-2 and -9 (1). Methylselenol induces apoptosis of B16F10 melanoma cells (2) and suppresses the invasive ability of the cells by reducing integrin expression (3).

Method: Methylselenol can be produced by selenomethionine plus methioninase (SeMet-METase) in a cell culture system and by Se-methylselenocysteine (Se-SMC) in an animal model. The expression of integrins was analyzed by flow cytometry. The enzymatic activities of MMP-2 and -9 were assayed by gelatin zymography. Four-week-old female C57BL/6J mice were fed a Se-MSC-supplemented diet or water. After four weeks, B16F10 melanoma cells were injected intravenously via the tail vein. On day 20 after the tumor cell injection, the pulmonary metastasis were macroscopically observed, and survival was checked 40 days later.

Result: Treatment with SeMet-METase induced a substantial decrease in the expression of integrins, and fibronectin receptors in B16F10 melanoma cells. Moreover, this compound suppressed gelatinase activity and invasive ability in the cell culture system. Mice given a mash diet or water supplemented with Se-MSC almost completely diminished the pulmonary metastasis of the melanoma cells and enhanced survival compared with the control group. Collectively, our results indicates that methylselenol contributes to blocking the metastasis of B16F10 melanoma cells by the suppression of integrin expression.

Discussion: The results suggest that Se-MSC can be used for cancer treatment as nutritional adjuvant. More studies are required to use Se-MSC as nutritional adjuvant for determination of an adequate amount without toxicity.

Selected references
O62 - Selenium Enhances Auranofin-mediated Nrf2 Induction in Lung Epithelial Cells

3. Selenium in animal and human health and disease
3.7 Additional and emerging topics of selenium in health or disease

Keywords: Nrf2, thioredoxin reductase, acute lung injury, bronchopulmonary dysplasia

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Introduction: Bronchopulmonary dysplasia (BPD) is common in preterm infants and acute lung injury (ALI) is associated with significant mortality in critically ill patients. Thioredoxin reductase-1 (TrxR1) inhibition by auranofin (AFN) activates Nrf2-dependent responses in murine transformed club cells (mtCC), decreases lung damage, and improves survival in murine models of BPD and ALI. TrxR1 activity is selenium (Se) dependent and Se deficiency is common in preterm infants and critically ill patients. We tested the hypothesis that Se supplementation would enhance Nrf2 induction by AFN.

Method: MtCCs, supplemented with 0, 25, or 100nM Se, were treated with 0.5 mM AFN or vehicle for 1h. TrxR1 activity was assessed and nuclear Nrf2 protein amounts determined. Data (mean±SEM) were analyzed by ANOVA.

Result: We detected a concentration-dependent effect of Se supplementation on TrxR1 activity in control-treated mtCCs (R²=0.97, p<0.0001). AFN inhibited TrxR1 activity in control and Se-treated groups (p<0.0001 vs vehicle). Nuclear Nrf2 protein was increased in all AFN-treated groups compared to respective vehicle-treated controls (0 nM: 2.7±0.1 vs 1.0±0.2; 25 nM: 5.6±0.6 vs 1.5±0.4; 100 nM: 4.6±0.4 vs 0.6±0.1; all p<0.05). The magnitude of AFN-induced increases in nuclear Nrf2 was greatest in Se-supplemented mtCCs (25 nM: 13.9±2.3 vs 5.8±0.2; and, 100 nM: 13.5±1.5 vs 5.8±0.2, p=0.02).

Discussion: Our findings support the hypothesis that Se supplementation enhances Nrf2 activation by TrxR1 inhibition. We speculate that Se status may modulate the efficacy of TrxR1 inhibitors as therapeutic agents to prevent or treat BPD and/or ALI. Optimization of Se status could enhance the therapeutic efficacy of TrxR inhibition.

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Effects: Treatment (p<0.0001); Na₂SeO₃ (p=0.0002)
Interaction: p=0.0032
* vs control, same [Se]
# vs AFN + 0 nM Se and AFN + 10 nM Se
O63 - Preclinical chemopreventive efficacy of a novel hybrid p-XSC-aspirin compound in a lung cancer model

Keywords: Chemoprevention, lung cancer, aspirin, hybrid compounds

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Introduction: 1,4-Phenylenebis(methylene)selenocyanate (p-XSC) has been shown to inhibit tobacco carcinogen NNK induced lung cancer development in several animal models[1-2]. We hypothesized that p-XS-Asp would cleave in vivo to release the active p-XSeH, not releasing undesired HCN but the aspirin (Figure 1), thus making the compound less toxic and more potent than p-XSC or aspirin alone.

Method: Chemopreventive efficacy of dietary p-XS-Asp was assessed using a NNK-induced A/J mouse model. Se content in plasma and tissues were carried out by atomic absorption spectroscopy. DNA adducts in lung tissues were performed by MS/MS.

Result: Orally fed mice by gavage showed higher bioavailability for p-XS-Asp compared with p-XSC along with greater inhibition of DNA adducts in lung and liver. MTD study reflected lower toxicity for the hybrid derivative (MTD values between 74.5 and 111.7 versus 39.8 mg/Kg). p-XS-Asp groups at doses of 15 ppm and 7.5 ppm Se showed a significantly marked decrease in the percentage of lung cancer incidence in vivo with only 50% and 87% of tumor incidence, as compared to p-XSC (79% and 100%), respectively. NNK-control showed a 100% tumor incidence at both the doses. Likewise, the multiplicity for p-XS-Asp was 0.87 and 1.93 tumors/mouse as compared with NNK-control (11.53) and p-XSC (1.66 and 4.10 tumors/mouse, respectively) at the two doses tested. Notably, blood and tissue analyses showed no systemic toxicity for the p-XS-Asp fed group.

Discussion: In conclusion, p-XS-Asp is less toxic while being more effective chemopreventive compound than p-XSC and is a promising candidate to future clinical evaluation.

Selected references

Figure 1. Mechanistic formation of active bis-selenol (p-XSeH) from (A) p-XSC: associated with the release of HCN, (B) p-XS-Asp: releasing aspirin as side product.
O64 - Selenofolate is a Promising Novel Agent in Targeted Chemotherapy

3. Selenium in animal and human health and disease
3.4 Selenium based medical therapeutics
Keywords: Selenofolate, Folate Receptor Alpha (FOLR1), Ovarian Cancer

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Introduction: Selenium compounds at high concentration have shown growth-modulating properties with high specificity for tumor cells. Thus, they have been reported to strongly improve the efficiency of chemotherapeutics. Folate receptor alpha (FOLR1) is a membrane-bound transport protein with high affinity for folate. FOLR1 expression levels are highly upregulated in cancer cells of epithelial origin and positively correlate with tumor progression. To date, the appealing concept of therapeutic FOLR1 pathway intervention is explored extensively, as FOLR1 expression is largely absent in normal tissue. Hence, selective targeting of FOLR1 presents a unique and promising option in chemotherapy.

Method: Here we used a synthetic selenofolate to investigate its effect on a FOLR1 high-expressing ovarian adenocarcinoma cell line, IGROV-1 cells, compared to a FOLR1 negative malignant melanoma cell line, A-375 cells. We tested the half maximal inhibitory concentration (IC50) of selenofolate by a luminescence-based viability assay.

Result: Surprisingly, selenofolate was more cytotoxic to the FOLR1 negative A-375 cells than to the FOLR1 high-expressing IGROV-1 cells, suggesting a FOLR1-independent pathway for cellular selenofolate uptake. Furthermore, siRNA-mediated knockdown of FOLR1 expression did not impact the growth of IGROV-1 cells despite alterations in specific mitogenic gene expression.

Discussion: Our results indicate that the role of FOLR1 in malignant cells is still poorly understood and that the transport of folates might not be its foremost function. Currently, we investigate the cytotoxicity of selenofolate in combination with other drugs interfering with the folate metabolism, such as methotrexate. Our work will uncover the potential use of selenofolate as a supporting agent in chemotherapy.
O65 - Environmental Remediation of Selenium Contamination in Water and Soil

1. Selenium chemistry and geochemistry
1.5 Excessive selenium accumulation from natural or anthropogenic sources and remediation technologies
Keywords: Treatment wetlands, Phytoremediation, Accumulation, Volatilization, Plant and microbial interaction

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Introduction: Selenium (Se) is essential to humans and animals, but becomes toxic to wildlife at high concentrations in the environment. Previous studies demonstrated that plants and plant-associated microbes can be applied to remove excessive Se from contaminated waters and soils. For example, the vegetated constructed wetlands were capable of significantly reducing Se from the inflow waste water. Biogenic volatilization of Se represents an environmentally sound remediation strategy to remove Se from the contaminated environment.

Method: The selected Se remediation technologies include constructed treatment wetlands, different soil-plant systems, and the algal Se reduction pond. This presentation focused on the plant-soil microbial interactions and their effects on Se bioaccumulation and chemical transformation. Different environmental conditions and chemical characteristics of Se pollutants (including nanoscale elemental Se) have been evaluated for the processes of Se immobilization, bioaccumulation and volatilization.

Result: The constructed treatment wetland was able to remove about 70% of the total Se mass in selenate-contaminated water. About 55% of the Se was immobilized in sediment, 5% of the Se accumulated in plants, and 10% of the Se volatilized to the atmosphere. The chemical transformation of selenate to elemental Se is the mechanistic process for the Se removal in treatment wetlands or algal retention ponds. Recent studies showed that microbial-derived nanoscale elemental Se could become bioavailable and further volatilized in soil-plant systems.

Discussion: The transport and fate of Se removal was discussed, and using Se-laden plant materials from phytoremediation for producing Se-biofortified edible mushrooms has been demonstrated.

Selected references
O66 - Integrated Passive Biological Selenium Treatment System: Results of a 1-year Pilot Study

Keywords: passive treatment, biological reduction, pilot study

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Introduction: Passive biological systems comprised of anaerobic organic media reactors create conditions supportive of dissimilatory reduction, adsorption and volatilization of Se at a lower cost compared to active biological treatment systems (Bays et al, 2013; Bays et al. 2014). Integrated passive biological treatment systems (BTS) combine anaerobic treatment with aerobic polishing to meet multiple regulatory water quality criteria. This paper summarizes a one-year BTS study in Centennial CO using membrane concentrate from the Joint Water Purification Plant (JWPP) for the Cottonwood Water & Sanitation District (CWSD).

Method: CH2M piloted a BTS at the JWPP comprised of two parallel trains. Water in each train flowed passively through passive vertical downflow anaerobic biochemical reactors (BCRs) comprised of wood chips, sawdust, straw, horse manure, and limestone chips for primary removal of Se, followed by a sequence of organic and inorganic media cells configured for removal of metals, phosphorus, and nitrogen. The BTS was operated and monitored under the direction of CH2M in three phases from January through December 2016. Samples were collected weekly and laboratory analyses performed using US EPA methods.

Result: Inflow Se averaged 68, 47 and 37 µg/L and outflow Se averaged 4, 3.4, and 3.3 µg/L for Phases 1-3, e.g. Train 1 (Figure 1). Substrates sequestered Se as predominantly adsorbed selenite, elemental and organic Se. Phosphorus and other pollutants met discharge criteria.

Discussion: The BTS pilot effectively met the 30-day average discharge criterion of 4.6 µg Se/L, as well as criteria for phosphorus and other parameters, demonstrating the benefit of passive anaerobic and aerobic treatment.

Selected references
O67 - Industrial selenium pollution: challenges to treat Flue Gas Desulfurization effluents

1. Selenium chemistry and geochemistry
1.5 Excessive selenium accumulation from natural or anthropogenic sources and remediation technologies
Keywords: FGD, Industrial pollution, Ion exchange, Desulfurization

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Introduction: Selenium (Se) pollution has been linked to numerous episodes of severe aquatic ecosystem deterioration [1]. Flue Gas Desulfurization (FGD) effluents produced by coal-fired power facilities are arguably the most complex and difficult to treat [2]. In this paper, a real FGD effluent was characterized and the Se removal potential was investigated for the first time by chemical precipitation/adsorption using a commercial ion exchange resin.

Method: The effluent was characterized for Se and metal content, major anions and other parameters [3]. A commercial ion exchange resin (gel type II hybrid strong base anion exchange resin impregnated with iron oxide) was employed as the adsorbent. Sulfate was precipitated (desulfurization) using BaCl₂. The adsorption experiments were performed in the batch mode.

Result: In the studied FGD, sulfate, a known competitor of Se, was found to be present almost 3 orders of magnitude in excess of Se (1.1 g L⁻¹ vs 1.2 mg L⁻¹). The resin performed poorly (~3% Se removal) against the raw FGD effluent. Desulfurization followed by ion exchange treatment led to a significant increase in Se removal (~80%). However, complete desulfurization using equimolar BaCl₂ could not be achieved due to the presence of bicarbonate that acts as a sulfate competitor for Ba. In addition to Se and SO₄²⁻, several toxic metals were efficiently removed (Cd: 91%; Cr: 100%; Zn: 99%) by the combined treatment.

Discussion: Due to the complexity of FGD effluents, this study underscores the necessity of a multistage treatment process as a high-performance Se treatment platform.

Selected references
O68 - Role of antioxidant defense system and mitochondrial activity in selenium toxicity tolerance in wheat

1. Selenium chemistry and geochemistry
1.5 Excessive selenium accumulation from natural or anthropogenic sources and remediation technologies

Keywords: Oxidative stress, Wheat, Selenate, Selenite

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Introduction: Selenium (Se) induced oxidative stress as well as synthesis of non-specific selenoproteins has been attributed to its toxicity in plants. Crops grown on Se-rich soil accumulate Se to the levels, considered highly toxic to plants and for animal and human consumption. This study reveals the effect of selenium on its uptake, leaf physiology, antioxidant defense system, isoenzymic patterns and mitochondrial activity in wheat.

Method: Selenium treatment-induced changes in growth, pigments, mitochondrial activity, antioxidant enzymes, antioxidants and other related metabolites were studied by standard procedures reported in the literature at tillering and ear-initiation stages.

Result: Selenate-treated wheat plants accumulated higher Se concentration in leaves than control and selenite treatment. Selenium-treated plants exhibited lower Se tolerance index, reduced growth, impaired photosynthesis and mitochondrial activities, high levels of hydrogen peroxide and lower reactive oxygen species scavenging activities leading to a range of antioxidant responses that varied depending on the form and doses of Se used. There was presence of two extra catalase, one superoxide dismutase, two peroxidase and one glutathione reductase isozyme in different Se-treatments.

Discussion: Wheat plants could adapt to applied selenite concentrations by developing antioxidant defense system but selenate-treated plants exhibited toxicity tolerance up to 2 mg kg⁻¹ and died at high concentrations due to damage to tissue development and function. The higher Se uptake in selenate as compared to selenite treatment was due to its mobile nature. Significant increase in hydrogen peroxide and antioxidant enzymes activities in leaves was due to the presence of Se-induced oxidative stress in wheat plants.

Selected references


O69 - Insufficient documentation for clinical efficacy of selenium supplementation in chronic autoimmune thyroiditis

3. Selenium in animal and human health and disease
3.4 Selenium based medical therapeutics
Keywords: selenium supplementation, chronic autoimmune thyroiditis, thyroid, meta-analysis

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Introduction: Selenium supplementation effectively reduces thyroid autoantibody concentrations in chronic autoimmune thyroiditis (AIT), but its effects on clinically relevant outcomes are unclear. The purpose of this systematic review and meta-analysis was to investigate clinically relevant effects of selenium supplementation in patients with AIT.

Method: Controlled trials in adults (≥ 18 years) with AIT, comparing selenium with or without levothyroxine substitution (LT4), versus placebo and/or LT4, were eligible for inclusion. Identified outcomes were serum thyrotropin (TSH) levels in LT4-untreated patients, thyroid ultrasound (US) and health-related quality of life (HRQL). Eleven publications, covering nine controlled trials, were included in the systematic review. Random effects model meta-analyses were performed in weighted mean difference (WMD) for TSH, US and HRQL. Quality of evidence was assessed per outcome, using GRADE.

Result: Meta-analyses showed no change in TSH, or improvements in HRQL or thyroid echogenicity (US), between LT4-untreated patients assigned to selenium supplementation or placebo. Three trials found some improvement in wellbeing in patients receiving LT4, but could not be synthesized in a meta-analysis. The quality of evidence ranged from very low to low for TSH as well as US outcomes, and low to moderate for HRQL, and was generally downgraded due to small sample sizes.

Discussion: We found no effect of selenium supplementation on TSH, HRQL or thyroid US, in LT4-untreated individuals, and sporadic evaluation of clinically relevant outcomes in LT4-treated patients. Future well-powered RCTs, evaluating e.g. disease progression or HRQL, are warranted before determining the relevance of selenium supplementation in AIT.
O70 - Dihydroxy-1-selenolane protects cells from radiation-induced mitotic death: role of GPx
3. Selenium in animal and human health and disease
3.4 Selenium based medical therapeutics

Keywords: Organoselenium, radioprotector, glutathione peroxidase, DNA repair

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Introduction: In recent times, researchers are exploring synthetic organoselenium compounds as novel therapeutic agents.

Method: Dihydroxy-1-selenolane (DHS) an organoselenium compound [1] was evaluated for radio-protective effect in cellular models employing cellular, biochemical and molecular assays.

Result: DHS treatment at 25µM for 16 hours significantly protected CHO cells of epithelial origin from radiation (4-11 Gy)-induced cell death and this effect was associated with increase in glutathione peroxidase (GPx) both at activity and mRNA level. The addition of mercaptosuccinic acid, a pharmacological inhibitor of GPx abrogated the radio-protective activity of DHS (fig. 1). On contrary increasing the GPx level in CHO cells through treatment with DHS-C₆, a lipophilic conjugate known to increase the cellular uptake of DHS and there by GPx level, showed significantly higher protection (fig. 1). Surprisingly both DHS and DHS-C₆ although increased the GPx level in spleen lymphocytes (radiosensitive cells) did not protect them from the radiation-induced apoptosis as monitored by propidium iodide (PI) and DNA fragmentation assays. Since CHO cells undergo radiation-induced cell death through mitotic catastrophe mediated through chromosomal aberration and G2/M arrest, it was anticipated that radio-protective effect of DHS might be related to its effect on DNA damage/repair and cell cycle arrest. Supporting this assumption, our results revealed that treatment with DHS and/or DHS-C₆ in CHO cells led to faster repair of DNA following radiation exposure and subsequently inhibited the G2/M arrest. The GPx inhibitor, mercaptosuccinic acid reversed these effects.

Discussion: Taken together it is confirmed that DHS-induced GPx level protects cells from mitotic death but not from apoptosis.

Selected references

Fig. 1. (A) Effect of a GPx inhibitor, mercaptosuccinic acid (MSA) on the radio-protective effect of DHS and DHS-C₆ in CHO cells calculated in terms of survival fraction by clonogenic assay. (B) Representative images showing colonies of CHO cells under different treatment conditions. The cells treated with DHS (25 µM) and DHS-C₆ (25 µM) for 16h were incubated with MSA (10 mM) for 2h and then exposed to irradiation at 11 Gy. Following this, cells were cultured for 7 days and stained with crystal violet (0.5%). Results are presented as means ± SEM, n = 3. * p < 0.05 as compared to radiation control. # p < 0.05 as compared to radiation + DHS or DHS-C₆ treated group.

CN - Control, IR – Radiation control, MSA – Mercaptosuccinic acid.
O71 - Inhibition of metalloenzymes, i.e., angiotensin-converting enzyme and tyrosinase, by selenol-metal ion interaction of selenoneine

3. Selenium in animal and human health and disease
3.4 Selenium based medical therapeutics
Keywords: selenoneine, ACE, tyrosinase

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Introduction: Selenoneine is a novel selenium containing compound discovered in the blood of bluefin tuna. Since selenoneine has strong antioxidant activity and detoxification ability of methyl mercury, it is expected to be used for functional foods as antioxidant and selenium supplements. This study characterized its biological activities against hypertension and skin melanism.

Method: Inhibition of angiotensin converting enzyme (ACE) and tyrosinase was assayed in the presence of selenoneine. Melanin generation was measured in mice melanoma cells and human melanoma cells.

Result: Selenoneine inhibited activities of ACE and tyrosinase. The Km values of ACE were 1.396 ± 0.103 mM with selenoneine and 0.724 ± 0.051 mM without selenoneine. Lineweaver-Burk plots for inhibition of ACE activities by selenoneine indicated a competitive inhibition. Melanin generation was also inhibited by selenoneine in enzyme activity assay and bioassay with human cultured keratinocytes. On the other hand, selenite and ergothioneine (thiol-analog of selenoneine) did not inhibit these enzyme activities and melanin generation.

Discussion: ACE and tyrosinase are metalloenzymes. ACE has a zinc ion and tyrosinase has two copper ions in their active centers. Since selenoneine inhibited metalloenzymes, selenol group in selenoneine may interact with metal ions situated active sites. Therefore, selenoneine has a novel biological functions in metalloenzyme inhibition, relating to prevention of hyper-tension by ACE inhibition and reduction of melanin production by tyrosinase inhibition.
O72 - Selenium reduces macrophage and B-cell responses in vitro and suppresses germinal center B-cell responses in vivo: A potential for therapy in lupus

3. Selenium in animal and human health and disease
3.4 Selenium based medical therapeutics

Keywords: Systemic Lupus Erythematosus, B-Cell, Macrophages, Germinal center

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Introduction: A systems biology approach applied to determine selenium-responsive markers in circulation revealed alterations in proteins critical for regulating systemic lupus erythematosus (SLE), in selenium-supplemented healthy men¹. For the current study we investigated the impact of selenium on B-cells and macrophages for its potential as a therapeutic supplement for SLE patients.

Method: Bone marrow cells from B6.129F2/J mice were differentiated into macrophages in-vitro with or without selenium. Alternately, bone marrow cells were differentiated into macrophages and subsequently treated with selenium. Macrophage maturation was assessed by flow cytometry in both conditions. Additionally, B-cells from B6.129F2/J mice were treated with selenium, and simultaneously activated with LPS and anti-CD40 and B-cell survival/activation was measured by flow cytometry. Selenium compounds used for this study included: Se-methylselenocysteine, selenomethionine, sodium selenite, and methylseleninic acid (MSeA) (dose range 0.125 - 100 µM). SLE-prone mice, B6.Sle1b, were supplemented with MSeA (3 ppm, orally, 5 days a week) for 3 months. Splenic cellularity, germinal center (GC) B-cells and serum autoantibody titers were investigated.

Result: In vitro, selenium reduced macrophage and B-cell survival at higher doses. Se-methylselenocysteine and MSeA inhibited macrophage differentiation/maturation and B-cell activation. MSeA-treated B6.Sle1b mice showed significantly decreased splenic weights and cellularity (Fig. A-B) including reduced GC B-cells (Fig. C) and macrophages (Fig. D). Consistently there were reduced autoantibodies in MSeA-treated mice compared to untreated mice (Fig. E-F).

Discussion: Selenium reduces macrophage and B-cell responses in vitro and suppresses GC B-cell responses in vivo. These observations suggest a promising therapeutic potential for selenium supplementation in SLE patients.

Selected references
O73 - The significance of selenoproteins for human health revealed by inborn errors of metabolism

2. Selenium in the molecular life sciences
2.5 Selenoprotein genetics

Keywords: TXNRD; neurodegeneration; PCH2; familial glucocorticoid deficiency; Sedaghatian

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Introduction: The significance of the 25 genes encoding selenoproteins for human health is increasingly recognized through the identification of patients with inborn errors in selenoprotein biosynthetic factors or in individual selenoproteins.

Mutations in selenoprotein N lead a spectrum of disorders nowadays called SEPN1(selenon)-related myopathy.

Mutations in glutathione peroxidase 4 cause respiratory failure and bone defects (Sedaghatian syndrome). Patients carrying mutations in thioredoxin reductase 2 (TXNRD2) suffer from familial glucocorticoid deficiency.

Method:

Result: Recently, we have identified patients with hypomorphic mutations in TXNRD1 who suffer from familial epilepsy. As with mutations in TXNRD2, modifier genes may modulate the phenotype.

There are also inborn errors of selenoprotein biosynthesis that affect many selenoproteins. For example, pathogenic mutations in selenocysteine synthase cause neurodevelopmental and neurodegenerative disorders. Mostly endocrine phenotypes result from mutations in SECIS-binding protein 2 and tRNA[Ser]Sec, which will be covered in another lecture. Mutations in the latter two genes involve impaired metabolism and action of thyroid hormones leading to delayed bone growth and maturation. In addition, mutations in SECISBP2 sometimes affect nervous system development, muscle, inner ear, skin, and immune system function underlining the significance of selenoproteins for many organ systems.

Discussion: Mouse models have been generated for several of these mutations and will be compared to human phenotypes where possible.

Selected references

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2. Homozygous mutation in TXNRD1 is associated with genetic generalized epilepsy.
   Free Radical Biology and Medicine 2017 106:270–277
O74 - Mouse models lacking the selenoprotein thioredoxin reductase-1

2. Selenium in the molecular life sciences
2.5 Selenoprotein genetics
Keywords: thioredoxin, glutathione, ribonucleotide reductase, NADPH

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Introduction: Oxidation of energetic nutrients sustains high intracellular NADPH/NADP⁺ ratios. NADPH-dependent reduction of thioredoxin-1 (Trx1)-disulfide and glutathione-disulfide by the selenoprotein thioredoxin reductase-1 (TrxR1) and glutathione reductase (GR), respectively, fuels ribonucleotide reductase (RNR) and cytosolic antioxidant systems.

Method: Here we investigated mouse models with liver-specific co-disruptions of the genes encoding TrxR1, Trx1, and GR.

Result: Mouse livers lacking both TrxR1 and GR sustained reduced glutathione (GSH) pools using an NADPH-independent methionine-consuming system. Liver-specific co-disruption of Trx1, TrxR1, and GR, paradoxically, causes a 100-fold increase in hepatocyte proliferation, indicating that, in the absence of Trx1, methionine-fueled GSH/glutaredoxin supports mammalian S phase RNR.

Discussion: In the triple-null livers, like in many cancers, RNR places a critical yet relatively low-volume demand on cytosolic reducing power, thereby favoring high hepatocyte turnover over sustained hepatocyte integrity.

Selected references
O75 - Mutated selenocysteine synthase creates a Sedaghatian-type spondylometaphyseal dysplasia mouse model

2. Selenium in the molecular life sciences
2.5 Selenoprotein genetics

Keywords: SepSecS, pontocerebellar hypoplasia, PLP

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Introduction: Mutations in SEPSECS (O-Phosphoseryl-tRNA:selenocysteinyl-tRNA synthase) have been found in different human populations. These mutations are the cause of a severe early-onset neurodegenerative disease, termed now pontocerebellar hypoplasia type 2D (PCH2D).

SEPSECS is a PLP (pyridoxal phosphate) dependent-enzyme which catalyzes the final step of selenocysteine (Sec) biosynthesis using selephosphate as selenium donor. Sec is mostly found in the active site of selenoproteins, which are essential for brain function.

PCH2D patients suffer from severe spasticity, profound mental retardation and progressive microcephaly. One of the SEPSECS mutations found in Sephardic Jewish patients of Iraqi ancestry consists on a substitution of a highly conserved tyrosine at position 334 with cysteine (Y334C). This mutation was suggested to affect the binding of PLP cofactor, required for the enzyme catalysis.

Method: We generated mice carrying the Y334C mutation, using a targeting vector created by recombineering.

Result: Unlike human PCH2D patients, homozygous mice (Sepsecs²³₃₄₄/Y334C) died the day after birth, did not show a milk spot, developed cyanosis, and showed abnormal electrocardiograms. Analysis of tissue expression revealed that selenoproteins, including Gpx4, were mainly reduced in brain, especially in neurons, but not in liver. In primary culture, PLP supplementation did not rescue neuronal selenoprotein levels.

Discussion: The Sepsecs²³₃₄₄/Y334C mouse phenotype resembles Sedaghatian-type spondylometaphyseal dysplasia (SSMD), a neonatal lethal disease caused by truncating mutations in GPX4. This disease is characterized by cardio-respiratory insufficiency and neurological anomalies.
O76 - Agronomic strategies affect the efficacy and quality of selenium biofortification

1. Selenium chemistry and geochemistry
1.6 Strategies to improve selenium accumulation and biofortification

Keywords: biofortification, selenoamino acids, vegetables

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Introduction: The Se content of food is highly dependent on agronomic strategies, soil Se bioavailability, and the ability of plants to take up and accumulate Se as selenoamino acids. The dietary intake of Se is strongly influenced by the consumption of food products containing Se. To mitigate a low intake of Se, biofortification strategies such as applying foliar Se, amending soils with Se fertilizers or cropping in Se rich soil, have all been used to increase Se accumulation in crops.

Method: In three multi-year field studies, the following biofortification strategies have been evaluated: 1) foliar application of selenate, selenite, or selenomethionine; 2) soil application of selenate, selenite, or selenomethionine; and 3) growing crops in soils naturally rich in Se.

Result: Results indicated that with foliar application of Se, total Se content in Allium cepa ranged from 0.03 (control) to a high of 0.93 mg/kg DM with selenate treatment and greatest percentages (%) of selenoamino acids were γ-glu-MeSeCys (67), MeSeCys (15) and SeMet (13). With soil applications of Se, analyses for Se in onion are in progress. In soil naturally rich in Se, total Se in five different Brassica vegetables ranged from 4.8 to a high of 17.2 mg/kg DM in Brassica oleracea var. capitata f. rubra where the greatest percentages (%) of selenoamino acids were SeMet (22), MeSeCys (16), and γ-glu-MeSeCys (13).

Discussion: Selenium biofortification occurred with all three sources of Se, however, Se accumulation was greatest but more variable when Brassica vegetables were grown in soils naturally rich in Se.

Selected references

O77 - Improving selenium supply in food systems

1. Selenium chemistry and geochemistry
1.6 Strategies to improve selenium accumulation and biofortification

Keywords: Biofortification, food system, selenium deficiency, Moringa

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Introduction: Selenium intakes are suboptimal in many countries. Strategies to increase Se supply include process fortification, biofortification, increasing dietary diversity and individual supplementation.

Under the HarvestPlus food system program, agronomic biofortification trials of Se and other micronutrients were conducted in China, Colombia and Australia, and orange sweet potatoes were researched and promoted in Melanesia.

Method: Potential improvements to food system Se delivery are presented, derived from studies by the author and colleagues from 2001-2015.

Result: Recovery of Se in edible crop parts was less than 15% of soil-applied Se. Foliar Se is usually more efficient, especially when applied with urea. Selenium-biofortified flour has been marketed in Australia since 2005.

In Tibet, where Se deficiency is widespread and where vast grasslands are unfertilised while cropland is often overfertilised, the most efficient method to increase Se may be to add with iodine in salt. This would exploit Se and iodine thyroid synergy, reach a high proportion of people and livestock and minimise wastage.

Genotype x environment studies of nutritious leafy vegetables in Pacific countries found that Moringa oleifera (drumstick tree) had Se leaf levels 12-fold higher than other plants on the same soil, including low-Se soils. In sub/tropical regions with low-Se soils, consumption of drumstick leaves/leaf powder could increase Se status along with sulphur, vitamins and protein at minimal cost.

Discussion: Improving the efficiency of biofortification, exploiting synergies with other micronutrients and employing natural Se biofortifiers could improve Se supply in food systems.
O78 - Agronomic Biofortification with Selenium in Intercropping Systems would address Low Selenium Intake

1. Selenium chemistry and geochemistry

1.6 Strategies to improve selenium accumulation and biofortification

Keywords: Bio-fortification, inter-cropping systems, selenium intake

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Introduction: Selenium (Se) is an essential human micronutrient with critical roles in immune functioning and antioxidant defense (Fairweather-Tait et al. 2011). There is widespread dietary Se deficiency in Sub-Saharan Africa (SSA) (Chilimba et al. 2011; Joy et al. 2014). Sole crops of maize were used in bio-fortification (Chilimba et al. 2012) but smallholder farmers intercrop maize with legumes. Therefore, agronomic bio-fortification with Se in intercrops was conducted to determine efficiencies of Se uptake in intercrops or sole crops.

Method: The experiment was conducted at two sites consisted monocrop maize, monocrop groundnuts, moncroc soybean, intercrop maize/groundnuts and intercrop maize/soybean with two rates of Se, 0 and 10 g ha⁻¹ using sodium selenate and laid out in a randomized complete block design with three replicates. Grain samples were analyzed for selenium (78Se) by ICP-MS (X-SeriesII, Thermo Fisher Scientific Inc., Waltham, MA, USA).

Result: Se application increased Se concentration and soybean gave the highest followed by groundnuts and maize (Figure 1). Se application in intercrops significantly increased Se recovery to 7.2 % compared to monocrops (Figure 2) and Selenium intake also increased (Table 1).

Discussion: The results indicate that Se recovery in monocropped maize is inefficient and applied Se is leached as selenate, adsorbed as selenite or immobilized into organic forms. Legumes are efficient in uptake of the applied Se compared to maize hence higher Se recovery. The Se intake range between 66 and 162 µg cap⁻¹ d⁻¹ which is optimum (Fairweather-Tait et al., 2011).

Selected references


O79 - To bio or not to bio? Strategies for agronomic biofortification with Se in tropical agroecosystems

1. Selenium chemistry and geochemistry
1.6 Strategies to improve selenium accumulation and biofortification
Keywords: Se-fertilizers, soil and foliar application, Se sorption, Oxisols

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Introduction: The addition of selenium (Se) in agroecosystems is important to increase Se levels in food crops, especially in Se-poor regions, e.g., Brazil. The potential of different plant species and agronomic strategies for Se biofortification have been addressed in several studies worldwide. Soil application of Se-containing fertilizers seems to be an effective strategy in soils from temperate regions, yet such approach may result in poor efficiency of the Se source in oxidic soils, where anionic forms of Se are strongly adsorbed. Under these circumstances, the application of Se together with other selected elements/nutrients may be considered in order to increase the efficiency of plants for taking up soil-Se.

Method: This work will discuss different strategies that can be used for increasing Se content in crops of great social/economic relevance in Brazil, through adoption of different agronomic biofortification practices. For that, we are setting a Research Network (BioSeBrazil) of greenhouse/field experiments with food and feed crops in different soils to test different methods (soil and/or foliar) and sources (inorganic and/or organic Se) for Se biofortification.

Result: Our efforts for addressing the best soil application approach focus on the use of Se together with P-fertilizers (at planting time), and/or through split nitrogen (N) fertilizer applications, using different N sources, as this play an important role in a nutrient management strategy as well as Se acquisition by plants.

Discussion: The rationale for adding Se with P and/or N fertilizers will be discussed, with a focus on the factors affecting Se availability in tropical (oxidic) soils (Figure 1).
O80 - Soil and foliar application of selenium in upland rice aiming agronomic biofortification

1. Selenium chemistry and geochemistry
1.6 Strategies to improve selenium accumulation and biofortification

Keywords: rice, agronomic biofortification, foliar application, soil application

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Introduction: Agronomic biofortification is a good strategy to increase selenium in plants. This study aimed to evaluate the levels of Se and application forms on rice growth, grain yield, and Se accumulation in seeds.

Method: The experiment was performed under field condition located at Selviria city, Mato do Grosso do Sul State, Brazil. The experimental design was a completely randomized $2 \times 6$ factorial scheme as follows: 2 forms of Se application (soil and foliar) and 6 levels of Se (0; 10; 20; 40; 80; e 160 g ha$^{-1}$). The rice seeds were harvested and prepared for chemical analysis. The Se analysis was performed as described in Chilimba et al. (2011).

Result: The total Se concentration ranged from 0.01 to 0.54 mg kg$^{-1}$ in the leaves and 0.03 to 0.36 mg kg$^{-1}$ in the seeds when applied at the level of 20 g ha$^{-1}$ via soil application. On the other hand, the Se level 20 g ha$^{-1}$ applied through foliar application the Se concentration in seeds ranged from 0.01 to 1.65 mg kg$^{-1}$.

Discussion: In this study, the calculated daily intake of Se from the biofortified rice grains ranged from 2.05 to 24.7 µg day$^{-1}$, which represent an increase from 3.72% to 44.9% of the daily requirement of Se per day when applied 20 g ha$^{-1}$ through soil application. Despite that the recommendation for daily Se intake for adults is 55 µg day$^{-1}$, the present study show relevant information on agronomic biofortification to increase Se content in edible parts, which can benefits human health.

Selected references
O81 - A model for Glutathione peroxidase 4-catalyzed reduction of lipid hydroperoxides in membranes

2. Selenium in the molecular life sciences
2.6 Selenoprotein function

Keywords: PHGPx, SPR analysis, Molecular Dynamics, Phospholipid hydroperoxide.

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Introduction: Reduction of PLOOH by GPx4/GSH is vital. Although details of the reaction have been unraveled, what is still unknown is how the enzyme interacts with membrane and how this impacts on catalytic cycle.

Method: SPR analysis to evaluate the affinity of GPx4 for different bilayers; in silico analysis of the docking and dynamics (MD) of the interaction during the catalytic cycle.

Result: SPR analysis indicates cardiolipin as the phospholipid with the maximal affinity to GPx4. This complies with the notion that GPx4 contains a cationic surface adjacent to the catalytic center. Consistently, KCl increases the $k_D$ of the GPx4 - CL complex. MD simulation shows that a hydroperoxide group posed in the core of the membrane moves up and is stabilized at the lipid-water interface. Docking analysis, validated by MD simulation, indicates two seemingly oscillating interaction patterns of the GPx4 cationic surface with the anionic head of CL. These interactions precisely address water-exposed 13-OOH or 9-OOH groups of tetra-linoleoyl CL toward catalytic redox center. However, in this pose the catalytic site would be inaccessible to the reducing substrate but the redox chemistry inside the complex and the anionic nature of GSH contribute to a displacement. This permits the last reaction of the cycle, when the enzyme is regenerated and GSSG released.

Discussion: GPx4 binds to the membrane by electrostatic interactions precisely addressing the redox center toward the phospholipid hydroperoxide, and GSH affects the binding. From this set of data, a model describing how the peroxidase jumps on the membrane seeking for hydroperoxide, is produced.
O82 - Selenium versus sulfur: reversibility of chemical reactions and resistance to permanent oxidation in proteins and nucleic acid

2. Selenium in the molecular life sciences

2.6 Selenoprotein function

Keywords: selenocysteine, cysteine, selenouridine, thiouridine, reversibility, resistance to oxidation

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Introduction: The central question in the field of selenium biochemistry is: “What chemical advantage does selenium confer to a protein, tRNA, or other biomolecule relative to the more abundant element sulfur?”

Method: One possible answer to this question is that chemical reactions with selenium are more readily reversible than those of sulfur. This is especially true for the oxidation of selenium relative to sulfur as Se-oxides are more rapidly reduced compared to S-oxides.

Result: Here, the evidence that selenium confers resistance to irreversible oxidation in selenium-containing enzymes is reviewed.

Discussion: New evidence will also be presented that resistance to permanent oxidation is extended to selenouridine-containing tRNAs.
O83 - The key ferroptosis regulator GPX4: cellular mechanisms and in vivo relevance
2. Selenium in the molecular life sciences
2.6 Selenoprotein function
Keywords: ACSL4, cell death, ferroptosis, PHGPx

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Introduction: The selenoenzyme glutathione peroxidase 4 (GPX4) has emerged as the key regulator of ferroptosis [1,2], a recently recognized form of regulated necrotic cell death [3]. Ferroptosis is clearly distinct from other paradigms of regulated cell death, such as apoptosis and necroptosis, and is characterized by an iron-dependent lipid peroxidation. Initially being recognized as a cell death modality highly relevant for a subset of cancer entities, ferroptosis has been implicated in a variety of degenerative disease contexts including tissue ischemia/reperfusion injury and neurodegeneration.

Method: CRISPR/Cas9-mediated genome-wide screening efforts and expression analysis of ferroptosis-resistant cell lines have recently identified acyl-CoA synthetase long-chain family member 4 (ACSL4) as a novel player in the ferroptotic process downstream of GPX4 deletion/inhibition [4].

Result: Using an interdisciplinary approach combining redox global phospholipidomics, reverse genetics, bioinformatics and systems biology, 15-hydroperoxy-arachidonoyl- and 15-hydroperoxy-adrenoyl residues esterified in phosphatidylethanolamines were identified as proximate signals in the ferroptotic death program [5]. Genetic ACSL4 disruption and pharmacological inhibition of ACSL4 prevented the accumulation of these death signals in cells and conferred an unprecedented resistance to ferroptosis induction triggered by GPX4 inactivation.

Discussion: Despite its outstanding importance for ferroptosis, little is known about the cellular and in vivo mechanisms that regulate GPX4 stability and potential implications in various disease contexts, which shall be discussed here.

Selected references
O84 - GPx4 depleted cell death involves different cell death pathway from ferroptosis

2. Selenium in the molecular life sciences
2.6 Selenoprotein function

Keywords: GPx4, ferroptosis, lipid peroxidation, vitamin E, DFO

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Introduction: GPx4 is an intracellular antioxidant selenoprotein that directly reduces peroxidized phospholipids produced in cell membranes. We previously showed that disruption of GPx4 gene in embryo, testis and retina of mice induces cell death in normal tissues. Administration of vitamin E in testis GPx4 KO mice suppressed the cell death of spermatogenic cells, indicating that suppression of lipid peroxidation by GPx4 or vitamin E is required for the fate of the normal cells. On the other hand, anti-mutated Ras cancer drug erastin and RSL3 induce the iron dependent novel cell death (Ferroptosis). Since erastin and RSL3 indirectly or directly could suppress the activity of GPx4, GPx4 is a regulator of ferroptosis by suppression of lipid peroxidation. However, we found that tamoxifen inducible GPx4 gene disruption enhanced lipid peroxidation until 26hr, resulting in cell death at 48~72hr in our established MEF cells, on the other hand, erastin or RSL3 could induce cell death until 24hr. The differences of cell death mechanism between ferroptosis and GPx4 depleted cell death remained to be clarified.

Method: We analyzed this differences of cell death mechanism by several strategies using our established tamoxifen inducible GPx4 gene disruption MEF cells without 15-LOX expression.

Result: We found that DFO, an iron chelator, could suppress lipid peroxidation and ferroptosis but not GPx4 depleted cell death. Several our identified Lipo genes that could suppress GPx4 depleted cell death, could not inhibit ferroptosis.

Discussion: These results indicated that GPx4 gene disruption induced cell death involves different cell death pathway from ferroptosis by anti cancer drug.

Selected references

Lipid peroxidation-dependent cell death regulated by GPx4 and ferroptosis
O85 - Selenoproteins expressed in intestinal stem cells and cancer stem cells

Keywords: GPX2, SELENOH, cancer, stem cells, intestine

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Introduction: Both healthy and cancer stem cells are highly susceptible to changes in the cellular redox status. Because of that, they up-regulate antioxidant enzymes to protect them from damage and to modulate their signaling pathways. We aimed to identify selenoproteins relevant for stem cell maintenance and to study their role during intestinal differentiation or tumor development, respectively.

Method: We compared selenoprotein expression profiles of intestinal epithelial cells of the crypt base to those of differentiated epithelial cells. The identified selenoproteins were studied by means of a shRNA-mediated knockdown in cancer cell lines, by xenograft models, or by the characterization of knockout mice.

Result: Both glutathione peroxidase 2 (GPX2) and selenoprotein H (SELENOH) were highly expressed in undifferentiated intestinal epithelial cells. In colorectal cancer cells, a stable knockdown of either GPX2 or SELENOH resulted in less differentiation and stem cell maintenance. Also in the healthy intestine of GPX2 knockout mice and in 3D cultures with loss of GPX2 expression, differentiation into the different lineages was found to be disturbed. However, while GPX2 knockdown cells almost completely lost their competence to form colonies or tumor xenografts, SELENOH knockdown cells grew better under both circumstances. These effects appear to be modulated by different redox-sensitive pathways being affected by loss of either of these selenoproteins.

Discussion: Selenoprotein expression and thus selenium supply both are involved in the maintenance of stem cell physiology. However, the specific role of individual selenoproteins further needs to be clarified in the future to more efficiently counteract processes such as tumorigenesis.
O86 - Roles of the thioredoxin and glutathione systems in reduction of inorganic- and Cys-polysulfide species

2. Selenium in the molecular life sciences
3.5 Selenium metabolism or selenoprotein function in health and disease

Keywords: Thioredoxin reductase, thioredoxin, hydrogen sulfide, polysulfide protein persulfide

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Introduction: The pivotal roles of Reactive Sulfur Species, in particular inorganic- and Cys-polysulfides, as mediators of thiol-based redox signaling and protecting agents against oxidative stress is an emerging field of study. Production of these sulfur species is highly regulated in vivo and their biological actions in most cases is related to redox reactions on regulatory or functional protein Cys residues.¹,²

Method: We will discuss the underlying molecular mechanisms of the reductions of inorganic- and Cys-polysulfides species in a biological context.

Result: We found that the thioredoxin and glutathione systems are key players in the biological functions of endogenous polysulfide species via orchestrating their reductions to H₂S and the corresponding cysteine thiol derivatives.³ We developed novel methods to detect polysulfide species in cells and tissue samples. With the aid of these new methods we collected substantial evidence for the roles of the different components of the thioredoxin and glutathione systems in i) maintaining sulfane sulfur homeostasis, ii) regulation of redox signaling and iii) Cys protecting functions of polysulfides under oxidative stress.

Discussion: Our results shed light on novel pathways in the protecting and signaling functions of the NADPH driven thioredoxin and glutathione reducing machineries. Polysulfide reduction by Thioredoxin reductase, and/or by its concerted actions with other components of the thioredoxin system is critically Se-dependent. A number of proposed models will be shown for different biological scenarios.

Selected references
1) Nagy P. Methods in enzymology 554, 3-29
2) Ono K. et al. Free Radical Biology and Medicine 77, 82-94
3) Dóka É. et al. Science Advances 2 (1), e1500968
O87 - Nutritional aspects of selenium in human beings
Plenary Session I

Keywords: cellular and whole-body selenoprotein hierarchies, selenoprotein P, selenium nutritional status

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Introduction: Se deficiency causes a variety of diseases in animals but is known only as a cause of Keshan Disease in humans. Thus, Se status in humans might affect other diseases as well.

Method: Because most work on Se homeostasis has been carried out in animals, animal research will be covered and, then, those findings will be discussed in humans.

Result: Regulation of Se metabolism is controlled by its availability. When supply is more than is needed for selenoproteins, excess Se is excreted. There are several cellular mechanisms that regulate Se use for synthesis of individual selenoproteins. Some selenoproteins are favored over others for Se incorporation, creating a cellular hierarchy. Selenoprotein P (SELENO-P) is higher in the liver cellular hierarchy than Gpx1, which contains 25% of mouse Se. The liver secretes SELENO-P into the plasma to supply other tissues with Se. SELENO-P binds to the endocytic receptor apoER2. ApoER2 varies among tissues, creating an organ hierarchy for Se (in SELENO-P) uptake.

Discussion: Together, the two hierarchies create a whole-body selenoprotein hierarchy. This is how liver Se can maintain selenoproteins in other parts of the body. Human plasma contains most of its Se in two selenoproteins: Gpx3, which originates in the kidney and SELENO-P, which originates in the liver. Plasma also contains unregulated selenomethionine in all its methionine-containing proteins. Thus, plasma Se does not accurately mirror the regulated Se pool. Plasma Gpx3 mirrors kidney Se and SELENO-P mirrors liver Se. Plasma SELENO-P appears to be the most accurate indicator of whole-body Se.
O88 - Health Benefit Values (HBV) Reliably Indicate Effects of Seafood Consumption

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease

Keywords: Seafood, mercury, neurodevelopment

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Introduction: Methylmercury (MeHg) affinities for selenium (Se) are $\sim10^6$ greater than its affinity for sulfur. Thus, MeHg inhibits the Se-dependent enzymes required for healthy brain development and functions. This appears to explain why laboratory and epidemiological studies find adverse effects of MeHg are associated with exposures to amounts in substantial molar excess of Se. Future risk assessments need to apply the Health Benefit Value (HBV) that considers MeHg in relation to Se, and thus reliably predicts benefits or risks associated with maternal seafood intakes.

Method: We conducted a study of Hg and Se contents of over 14,000 ocean and freshwater fish samples collected from all over North America. The updated equation for calculating the HBV (Ralston et al., 2016) reflects the positive yield of dietary Se biologically available to consumers or the negative effects of MeHg in molar excess of Se.

Result: Calculation of the HBVs of seafoods consumed in New Zealand (typical varieties of ocean fish as well as great white sharks) and the Faroe Islands (pilot whale meats and cod) indicated these populations were exposed to substantial amounts of MeHg in excess of Se (negative HBV seafoods).

Discussion: Because it considers the beneficial effects associated with improved dietary intakes of Se, the HBV equation correctly predicts the adverse effects of pilot whale or shark meats as well as beneficial outcomes now known to be associated with consumption of typical varieties of ocean fish. Meanwhile, all other epidemiological studies involved seafoods with positive HBV's, consistent with their uniform findings of benefits.

Selected references

O89 - Molecular and Metabolic Mechanisms for Steatosis and Obesity Induced by Overexpression of Glutathione Peroxidase-1 in Mice

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease
Keywords: Calorie, ER stress, Glutathione peroxidase, Metabolic syndrome

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Introduction: Mice overexpressing Se-dependent glutathione peroxidase-1 (OE) developed fatty liver and obesity. This study was to reveal molecular dysregulation of tissue lipid metabolism and body energy expenditure shift leading to the disorder in these mice.

Method: Male OE and the wild-type (WT) mice were fed an Se-adequate Torula yeast-sucrose diet and were tested for energy metabolism at ages of 3 wk and 2, 6, 9, and 12 mo. Plasma, liver, and adipose tissue were collected at those ages for histology and assays of lipid profiles and functional expression of 20 key factors involved in lipid metabolism and ER stress.

Result: Obesity started to appear at 4 mo of age, whereas steatosis developed between 6 and 12 mo of age in the OE mice as shown by liver histology and total triglyceride accumulation. The OE mice displayed elevated levels of O₂ consumption and CO₂ production, but decreased activity after 6 mo of age than the WT mice. Compared with the WT, the OE mice showed upregulation (P < 0.05, 2-3 fold) of FASN, ChREBP and SREBP1c, but down-regulation (P < 0.05) of PPARβ, CPT1α, and ATGL in the liver and adipose tissue. The OE mice also had elevated expression (P < 0.01) of ER stress-related biomarkers (IRE1 and XBP1) and selenoproteins (K and S) in both tissues over the WT.

Discussion: Development of obesity and steatosis in the OE mice were associated with a stimulation of ER stress-related selenoproteins and enhanced triglyceride synthesis, along with decreased fatty acid oxidation, lipolysis, and activity.
O90 - Selenium use efficiency by wheat cultivars

1. Selenium chemistry and geochemistry
1.6 Strategies to improve selenium accumulation and biofortification

Keywords: Agronomic biofortification, micronutrients, food quality, plant nutrition

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Introduction: Agronomic biofortification is considered one of the best ways to raise the levels of nutrients and vitamins in foods, because of its practicality, speed and low cost.

Method: An experiment was conducted with 8 wheat cultivars, previously selected in field for contrasts in the accumulation of selenium (Se) in the grains. A completely randomized experimental design was used, with application of 0.5 mg dm⁻³ of Se and without Se application. Each experimental unit was represented by a 3 kg⁻¹ pot of soil, with three replicates.

Result: There was a positive effect of applying Se on the concentration and accumulation of Se in grains of all cultivars, but without significant difference in the concentration. Cultivars BRS Guamirim and Mentana presented the highest average Se concentration (2.8 mg kg⁻¹), followed by Abalone and BRS Parrudo (2.6 mg kg⁻¹), Londrina (2.5 mg kg⁻¹), CD 150 and Embrapa 21 (2.1 mg kg⁻¹) and BRS 210 (2.0 mg kg⁻¹). In the treatments without Se application, it was not possible to detect Se in the grains.

Discussion: Grain yield did not vary among the cultivars, regardless of Se application. There was small variation in the concentrations of macronutrients and micronutrients. Higher concentrations of Se and micronutrients were found in the diagnostic leaves than in the grains. Se addition in the soil was efficient to enrich the Se in the grains, proving to be an effective and practical way to biofortify wheat cultivars. There was no genetic variability in the accumulation of Se in the wheat cultivars.
O91 - Effects of selenium biofortification on mineral nutrients in grains of wheat and oat

1. Selenium chemistry and geochemistry
1.6 Strategies to improve selenium accumulation and biofortification

Keywords: Wheat, Oat, Selenium biofortification, Mineral nutrition

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Introduction: Wheat is one of the three major staple crops in the world, and oat is a nutrient dense crop. Se-biofortification of wheat and oat is conducive to relieve selenium (Se) deficiency in human diet

Method: Four levels of Se dosage, 0 mg·kg⁻¹ (CK), 100 mg·kg⁻¹ (Se10), 200 mg·kg⁻¹ (Se20) and 300 mg·kg⁻¹ (Se30), were foliar sprayed uniformly onto 114 wheat varieties and 22 oat varieties for the first time at booting stage and a second time at sprouting stage. The concentrations of Se, Ca, Mg, Cu, Fe, Mn, Zn and S in the grains were measured using Inductively Couple Plasma (ICP) method

Result: Se concentrations in grains of both crops were significantly increased (p<0.01) after Se application (Fig.1). The average Se contents in wheat grains of CK, Se10, Se20, Se30 were 1.54, 5.70, 10.01, and 13.10 mg·kg⁻¹, respectively, and 1.85, 4.59, 8.95, 14.66 mg·kg⁻¹ in oat, respectively. Se application decreased overall concentrations of the other seven nutrients in wheat. Concentrations of Cu, Fe, Zn, Mg were also decreased after Se spray in oat grains, however, Ca and S contents were unaffected and Mn content was increased

Discussion: Similar total Se were accumulated in wheat and oat grains but the effects of Se application on Ca, S and Mn concentration differed between the two crops. However, oat may be superior to wheat because oat grains can be consumed by humans, and leaf and stem can be used for animal silage, which is good for the welfare of animals and finally good for human health.

![Fig.1 Selenium concentration in grain of wheat and oat after foliar spray of selenate](image)

Fig.1 Selenium concentration in grain of wheat and oat after foliar spray of selenate

* *, ** *, and *** indicate the differences were significant between wheat and oat at p<0.05, p<0.01 and p<0.001, respectively.
O92 - Characterization on rhizosphere bacteria from a selenium-hyperaccumulator Cardamine hupingshanesis

1. Selenium chemistry and geochemistry
1.6 Strategies to improve selenium accumulation and biofortification
Keywords: Enshi, China; Cardamine hupingshanesis; Rhyzosphere

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Introduction: In the Se-mine drainage area from Enshi, China, a novel Se hyperaccumulating plant, Cardamine hupingshanesis, was discovered in 2013, and the plant could accumulate Se up to 4414 mg/kg in the root dominated by selenocystine (SeCys2) (Yuan et al., 2013). Previous studies by Pilon-Smists and her team revealed that there were 24 Se-tolerant fungus strains in the rhizosphere of Stanleya Pinnata, a typical Se-hyperaccumulator, and those strains could survive in a medium with Se up to 10 mg/L (Wangeline et al., 2011)

Method: High-performance 16S rRNA analysis was performed on rhizosphere bacteria from C. hupingshanesis to demonstrate their roles on Se-hyperaccumulation by C. hupingshanesis.

Result: Compared with the Se non-accumulating plants in study site, there have much higher contents of microorganisms in the rhizosphere of C. hupingshanesis, and it was predominant by alpha-proteobacteria class (15-22 %), beta-proteobacteria class (10-16 %), actinobacteria class (10-18 %), acidobacteria class (8-15 %), delta-proteobacteria class (5-16 %). Moreover, some special microorganism with nitrospira class (2-5 %), gemmatimonadetes class (2-4 %), verrucomicrobiae class (2-4 %), planctomycetacia class (1-2 %) and others (opitutae class, sphingobacteria class, bacilli class, clostridia class) (3-4 %).

Discussion: The diversity of rhizosphere microorganisms from C. hupingshanesis was plenty and some of them could be Se-tolerant to relate with Se transformation from soil to plant roots
O93 - Assessment and biofortification of wheat grain selenium in staple food production regions of China

1. Selenium chemistry and geochemistry
1.6 Strategies to improve selenium accumulation and biofortification
Keys: China, wheat, soil, foliar application

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Introduction: Selenium as an essential nutrient for human, its deficiency has become a worldwide concern and especially in developing countries like China. It is crucial to investigate the wheat grain selenium level and its relevant soil factors and potential biofortification measures.

Method: 655 wheat grain samples were collected from different wheat production regions in China during 2008-2009, 2009-2010, and 2010-2011, meanwhile the grain yield and selenium concentration were measured. And in the 2010-2011 season, another field experiment was conducted in 30 representative agriculture research stations, to investigate the effects of foliar applied water and 0.017% sodium selenite solution at jointing stage, on wheat grain yield and Se concentration at maturity.

Result: Overall, the average wheat grain Se concentration was 67.5 and 64.2 µg kg⁻¹ for spring and winter wheat, and 63%, 19% and 8% was evaluated as deficient, low, and enriched Se respectively. The wheat grains of northern and western regions were higher than that of southern and eastern regions. Foliar Se application of had no effect on wheat grain yield, but significantly increased grain Se concentration, on average from 31.0 to 647.8 µg kg⁻¹ by 116 g Se ha⁻¹. Grain Se was positively correlated with soil available Se and shoot Se concentration before jointing.

Discussion: In different wheat production areas, both soil available Se and shoot Se accumulation before jointing are important for grain Se at maturity. Therefore, soil application of Se at sowing and foliar Se application at middle or late jointing stage, are both effective measures for wheat grain Se biofortification.

<table>
<thead>
<tr>
<th>Region</th>
<th>Location</th>
<th>Cultivar</th>
<th>Grain yield (kg ha⁻¹)</th>
<th>Grain Se (µg kg⁻¹)</th>
<th>Control</th>
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Different letters in the same rows indicate significant difference between Control and Se treatment at P < 0.05. Short dash line “-” means data missing.
O94 - Selenium supplemented kale and kohlrabi sprouts as possible ingredients for potent functional food.

1. Selenium chemistry and geochemistry
1.6 Strategies to improve selenium accumulation and biofortification

Keywords: sprouts supplemented with selenium, antioxidant activity, phenolic compounds

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Introduction: The objective of this project was to examine biochemical properties of kale and kohlrabi sprouts supplemented with different doses of selenium.

Method: The kale and kohlrabi sprouts were grown in EQMM-Easy-Green-Microfarm for 5-7 days, and watered with tap water or selenium supplemented water (10 mg/L, 15 mg/L, 30 mg/L). We determined in sprouts: selenium concentration (AFS-230 spectrometer); phenolic acids (Dionex HPLC), antioxidant activity (DPPH, FRAP) and total phenolic compounds (TP) (Synergy-2 reader).

Result: Selenium was in the range 11-16 mg/g. Highest concentrations of sinapic acid were in kale sprouts supplemented with 15 or 30 mg/L Se (120.0±1.8 and 175.7±1.4 mg/100 g d.w., respectively). Chlorogenic acid occurred solely in kale sprouts and the highest concentration (10.3±0.1 mg/100 g d.w.) was for samples supplemented with 30 mg/L Se. Izo-chlorogenic acid was quantitatively determined only in samples supplemented with selenium, being one order of magnitude higher in kale than in kohlrabi sprouts (2349.8±448.1 vs. 383.1±116.9 mg/100 g d.w.), showing dose dependent positive relationship with selenium. The differences in TP followed the pattern of izo-chlorogenic acid. FRAP and DPPH showed slight increasing tendency following increase of selenium concentration, irrespectively of sprout species.

Discussion: We measured the antioxidant potential of selenium-supplemented kale and kohlrabi sprouts and identified some candidates (among phenolic acids) for most biologically active compounds they contained, and determined the influence of selenium on their synthesis. Next stage of the study will include the assessment of the impact of the examined sprouts on the viability, proliferation and metabolic activity of human normal and cancer cells.

Selected references
O95 - Selenoproteins in hypothalamic energy homeostasis

2. Selenium in the molecular life sciences
2.6 Selenoprotein function
Keywords: SELENOM, hypothalamus, leptin, energy metabolism

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\textsuperscript{2} University of Hawaii

\textbf{Introduction:} Leptin is a hormone secreted from adipose tissue that is a key regulator of energy metabolism. Upon binding to its cognate receptor in the hypothalamus, leptin activates cell signaling pathways that serve to decrease appetite and stimulate energy expenditure. One of the cardinal features of obesity is leptin resistance, in which increasing leptin levels fail to reduce feeding behavior and stimulate energy metabolism. The exact mechanism underlying hypothalamic leptin resistance is not entirely clear, but ER stress, inflammation, and impaired leptin transport across the blood-brain barrier have been implicated as contributing factors. Given the fact that selenoproteins are known to mitigate ER stress and inflammation, hypothalamic selenoproteins may protect against leptin resistance. Moreover, knockout studies in mice have shown that several selenoproteins (DIO2, GPX1, SELENOM, SELENOP, SELENOT) significantly impact energy homeostasis.

\textbf{Method:} This lecture will discuss experimental evidence supporting the beneficial influence of hypothalamic selenoproteins on global energy metabolism.

\textbf{Result:} Recent studies have shown that constitutive deletion of SELENOM and hypothalamic-specific knockout of selenocysteine-tRNA (Trsp) both lead to the development of obesity and hypothalamic leptin resistance in mice. Moreover, preliminary findings indicate that SELENOM promotes leptin signaling and Ca\textsuperscript{2+} homeostasis in an Agrp-expressing hypothalamic cell line (mHypoE-44).

\textbf{Discussion:} These findings strongly suggest that selenoproteins promote leptin signaling in the hypothalamus.
O96 - Role of hypothalamic selenoprotein M in leptin signaling and calcium regulation

2. Selenium in the molecular life sciences
2.6 Selenoprotein function

Keywords: selenoprotein M, leptin signaling, endoplasmic reticulum stress, calcium regulation, Hypothalamus

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Introduction: The prevalence of obesity is growing rapidly and has become an indisputable problem worldwide. One hallmark of obesity is hypothalamic leptin resistance, of which ER stress is a major contributing factor. Selenoprotein M (SELENOM) is an ER-resident thiol-disulfide oxidoreductase that is highly expressed in the brain. Our group has reported that deletion of SELENOM in mice results in elevated body weight, adiposity, and serum leptin levels, symptoms that are indicative of leptin resistance [1]. Moreover, SELENOM is highly expressed in the arcuate and paraventricular nuclei of the hypothalamus, two key regions involved in neural control of energy metabolism. In this study, we investigated the function of hypothalamic SELENOM using an immortalized hypothalamic cell line (mHypoE-44) that expresses agouti-related peptide, neuropeptide Y, and the leptin receptor.

Method: Responses to leptin stimulation and ER stress were assessed by Western blotting in mHypoE-44 cells where SELENOM expression was manipulated. Further studies utilized Ca^{2+} imaging to assess changes in cytosolic Ca^{2+} flux in response to both leptin and ER stress.

Result: Our studies show that SELENOM knockdown significantly attenuates leptin signaling in mHypoE-44 neurons, while overexpression promotes leptin-stimulated increase in pSTAT3, suggesting that SELENOM promotes leptin signaling in hypothalamus. Ca^{2+} imaging studies showed that SELENOM-depletion augmented cytosolic Ca^{2+} elevations in response to thapsigargin-induced ER stress, but mitigated leptin-induced reductions of cytosolic Ca^{2+} levels.

Discussion: These results provide clear evidence that SELENOM has an important role in leptin signaling, ER stress, and Ca^{2+} regulation in hypothalamic neurons. This study may provide future therapeutic applications for the obese populations.

Selected references
O97 - Exploring Selenoprotein N structure and function

2. Selenium in the molecular life sciences
2.6 Selenoprotein function

Keywords: Selenoprotein N, endoplasmic reticulum, membrane protein topology, structure function analysis

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Introduction: The Selenoprotein N (SelenoN) is a membrane glycoprotein localized in the endoplasmic reticulum (ER) and it is ubiquitously expressed in all tissues. However, mutations in the SELENON gene, which encodes SelenoN, leads to a selective set of muscular diseases (SELENON-related myopathies), which suggests that, despite its widespread expression, SelenoN is particularly important for molecular processes relevant to muscle development and maintenance. At the molecular level, the presence of a thioredoxin reductase like motif in the core of the protein indicates a redox function, but its catalytic activity, including its substrate(s) and co-factor(s), remains unknown.

Method: We are conducting structural and molecular studies to address the SelenoN activity and to analyze this activity in the context of cellular metabolic or signaling pathways, such as ER oxidative stress control or calcium homeostasis.

Result: We characterized the topology of SelenoN within the ER membrane, and showed the importance of its post-translational modifications. In parallel, we have developed eukaryotic expression systems for the production and purification of recombinant protein to be used in crystallographic studies and in vitro enzymatic assays. At the cellular level, we have screened for conditions controlling the transcription of the SELENON gene, and using SelenoN depleted models, we showed that the unfolded protein response, one important activity within the ER, is not affected by SelenoN loss of function.

Discussion: A better understanding of the precise SelenoN enzymatic activity is fundamental to deciphering the pathogenesis of SELENON-related myopathies.
**O98 - Disruption of cancer cell redox homeostasis promoted by S-nitrosylation of thioredoxin reductase**

2. Selenium in the molecular life sciences
2.6 Selenoprotein function

Keywords: Nitrosylation; Thioredoxin reductase; Redox regulation; Cell death

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**Introduction:** Nitric oxide (NO)-induced signaling and cytotoxic responses are mediated in part by S-nitrosylation of protein cysteine residues, to form S-nitrosothiols (SNOs). Protein denitrosylation is primarily carried out by glutathione (GSH), thioredoxin (Trx), and their associated redox systems. SNO reduction by GSH and Trx is increasingly implicated in the regulation of NO/SNO-mediated cellular signaling and in protection from SNO-related cellular stress.

**Method:** We used biochemical and cellular approaches to study the effects of NO/SNO donors on recombinant and endogenous Trx reductase (TrxR).

**Result:** NO/SNO donating agents rapidly and effectively inhibited the activity of recombinant rat TrxR1. In particular, the NADPH-reduced TrxR1 was partially and reversibly inhibited upon exposure to low micromolar concentrations of S-nitrosocysteine and markedly and continuously inhibited at higher doses. Further analyses indicated that its active site selenocysteine residue renders TrxR1 highly susceptible to nitrosylation-mediated inhibition, and revealed both thiol and selenol modifications at the two redox active centers of the enzyme. Studies in HeLa cancer cells further confirmed the susceptibility of TrxR to inhibitory nitrosylation. Notably, depletion of cellular glutathione with L-buthionine-sulfoximine (BSO) synergized with nitrosating agents in promoting rapid and sustained nitrosylation and inactivation of TrxR. These events were accompanied by significant oxidation of Trx1, activation of multiple cellular stress pathways, and induction of cell death.

**Discussion:** Our findings uncover novel aspects of the interplay between NO and the Trx and GSH redox systems. The observations suggest the utility of BSO/NO combination therapy for inhibition of tumor growth through the disruption of cancer cell NO-redox homeostasis.
O99 - Influence of small intestinal thioredoxin and thioredoxin reductase on intestinal permeability

2. Selenium in the molecular life sciences
2.6 Selenoprotein function
Keywords: small intestine, thioredoxin, thioredoxin reductase

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Introduction: Obesity is associated with increased intestinal permeability, but the underlying mechanism remains largely unclear. Here we show that the development of high-fat diet-induced obesity in mice was accompanied by a decrease in small intestinal thioredoxin (Trx) and its reductase (TrxR).

Method: Although Trx and TrxR are components of Trx system that participates in many cellular processes, the relationship between Trx system and intestinal permeability remains unknown. Thus, we made two model intestinal epithelial cells with increased or decreased activity of Trx system via selenium supplementation or Trx-knocking down.

Result: Comparison of these model cells showed distinct patterns. Na⁺-K⁺ ATPase α-subunit is a crucial determinant in regulating intestinal permeability. In the cells with lower activity of Trx system, Na⁺-K⁺ ATPase α-subunit expression was lower, but its phosphorylation level was higher, consisting with a decrease in Na⁺-K⁺ ATPase activity. In the same cells, the increased activity of protein kinase Cα and decreased activity of protein phosphatase-2A were in favor of Na⁺-K⁺ ATPase phosphorylation. In addition, the cells with decreased activity of Trx system had higher level of reactive oxygen species, abnormal shape, lower level of Zonula Occludens-1 and increased transepithelial transport.

Discussion: The observed differences reveal a previous unrecognized role of Trx/TrxR in regulating intestinal permeability.
O100 - The intricate role of selenoproteins in stress erythropoiesis
3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease
Keywords: Erythropoiesis, proerythroblasts, anemia, erythroblastic islands, selenoproteins

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Introduction: Se, in the form of selenoproteins, functions as a cellular redox gatekeeper to modulate a wide array of physiological processes. Stress erythropoiesis is a systemic response to hypoxia stress that involves the rapid generation of new erythrocytes. Our data show that Se and selenoproteins impacts stress erythropoiesis at several levels.

Method: Acute anemia model was induced by phenylhydrazine administration. Microarray was applied for transcriptomic analysis. Flow cytometry was used for phenotypic study.

Result: Se deficiency severely impairs stress erythropoiesis and compromises the response to acute anemia in mice. Se deficient animals fail to expand stress erythroid progenitors in the spleen and exhibit a defect in the differentiation of proerythroblasts to basophilic erythroblasts during terminal maturation. Molecular analysis showed that Se deficiency alters the progression of terminal erythroid differentiation leading to augmented cleavage of transcription factor GATA1 and defects in enucleation. Transcriptomic analysis revealed significant differences in proerythroblasts as a function of Se status. In particular, the expression of selenoprotein W (SELENOW) was decreased in Se-deficient cells suggesting that it may play a role in the terminal differentiation of stress erythroid progenitors. Se deficiency also decreased development of erythroblastic islands, which form the erythropoietic niche. To further demonstrate a role for selenoproteins in stress erythropoiesis, mice mutant for tRNA^{Ser}Sec (Trsp) exhibited impaired stress erythropoiesis and delayed recovery from anemia.

Discussion: Our data suggest a fundamental role for selenoproteins in supporting effective stress erythropoiesis and that adequacy of Se is required throughout the recovery process in the erythropoietic niche.

Selected references
NIH R01-DK077152 (KSP) and R01-DK080040 (RFP) and supported by USDA National Institute of Food and Agriculture Hatch Project #4736 (R.F.P.) and 4605 (K.S.P)
O101 - Selenoproteins restrict the replication of Francisella tularensis in macrophages.

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease

Keywords: bacteria, infection, selenoprotein, autophagy

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Introduction: The micronutrient selenium (Se) is known to regulate immune functions via selenoproteins. However, the mechanisms by which selenoproteins regulate immune functions during an acute infection are not clear. Therefore, we investigated the role of macrophage (Mac) selenoproteins during an acute bacterial infection.

Method: Francisella tularensis (Ft.), the causative agent of tularemia, is a gram-negative intracellular bacterium. Since Ft. infects and replicates primarily in Macs, we measured the bacterial growth in Macs derived from Trsp⁻⁻ mice that are unable to synthesize selenoproteins. The disease progression was monitored in wild-type (WT) and Trsp⁻⁻ mice to determine the influence selenoproteins in an acute infection. Since Ft. growth is closely associated with autophagy, we investigated the kinetics of autophagy in infected cells by western blot.

Result: Macs maintained on deficient level of Se had a ~50 fold higher bacterial numbers than the Macs maintained on adequate or supplemented levels of Se. Similarly, Trsp⁻⁻ Macs had a significantly higher growth of Ft. compared to WT Macs. Moreover, Trsp⁻⁻ mice were more susceptible to Ft. infection and harbored significantly higher levels of bacteria in their livers and spleens as compared to WT mice. Our data also indicate that selenoproteins inhibit autophagy in Macs, which may limit the availability of nutrients to the bacteria.

Discussion: These data indicate that Mac selenoproteins are essential for restricting bacterial growth and promoting host survival. Our studies demonstrate that dietary Se affect the outcome of an infection by influencing host immune response, and reveal potential novel targets for antibacterial therapies.
O102 - Se-dietary matrices can upregulate the anti-inflammatory responses in RAW macrophages

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Introduction: Dietary Se supplementation downregulates expression of proinflammatory genes. However, the extent of Se bioavailability from dietary matrices such as Se-rich wheat, and its impact on inflammation is not known.

Method: We tested the ability of Se-rich wheat extracts to impact the pathways of inflammation and pro-resolution in RAW264.7 macrophages stimulated with bacterial endotoxin lipopolysaccharide (LPS). Cells were supplemented with Se in the form of sodium selenite (SS), seleniferous wheat extract (SeW) and seleniferous wheat extract treated with rMETase (SeW+rMET) in three increasing concentrations (500nM>100nM>50nM). The expression of GPx-1, COX-2, HPGDS, mPGES-1, iNOS were examined in qPCR or western blot analyses.

Result: Cells supplemented with 500 nM SS and SeW+rMET showed 9.0 and 8.0 fold increase in GPx-1 expression, respectively, as compared to SeW treated cells (1.1 fold) at 4h of LPS incubation. rMETase pretreatment increased the bio-availability of Se from wheat matrices and showed effects equivalent to SS supplementation. 500nM SeW+rMET downregulated the LPS-induced expression of Cox-2, mPges-1 and iNos up to 4.0, 12.0 and 10.0 fold compared to Se deficient cells, respectively, at both mRNA and protein levels. SeW+rMET treatment significantly upregulated (4.4 fold) the expression of HPGDS, which produces anti-inflammatory and pro-resolving eicosanoid 15-deoxy-prostaglandin J2.

Discussion: Our studies show that treatment of seleniferous wheat extracts with rMetase increases the bioavailability of Se to downregulate pathways of inflammation, while increasing the pro-resolution responses to effectively mitigate inflammation. These results highlight the potential application of selenium supplementation through dietary matrices to alleviate diseases where inflammation contributes to the underlying pathology.

Selected references

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O103 - Selenium in human and vertebrate evolution
2. Selenium in the molecular life sciences
2.7 The systems biology of selenium and selenoproteins
Keywords: selenium deficiency, adequacy, toxicity, adaptation

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Introduction: Selenium, a sparse element on earth, is an essential micronutrient in the diet of humans and other vertebrates and its intake depends on its content in the soils and waters across the world. Selenium is required due to its function in selenoproteins, which contain the amino acid selenocysteine (the 21st amino acid in the genetic code) as one of their constituent residues. Selenocysteine is analogous to the amino acid cysteine, which uses the abundant element sulfur in place of selenium. Despite the irregular and often scant distribution of selenium in the world, the distinct biochemical properties of selenium have made the substitution of selenocysteine for cysteine rare in vertebrate proteins. Still, vertebrate species have lived in environments with different amounts of selenium and may have adapted in different ways to it.

Method: Using comparative and population genomics approaches, we investigate whether selenoprotein genes and genes involved in the metabolism of selenium have genetic signatures compatible with adaptive evolution.

Result: Fishes have genetic signatures compatible with adaptation to the variable but generally abundant selenium in waters throughout the world, whereas humans have genetic signatures compatible with adaptation to the varying but generally scarce selenium in lands across the world. In both cases, the adaptive signatures are shared among genes that use (selenoproteins) or regulate selenium, suggesting that it is the overall metabolism and homeostasis of selenium that adapts to its worldwide variation.

Discussion: Dietary selenium has thus distinctly shaped the evolution of vertebrates.
O104 - Evolution of selenoproteins across the tree of life

2. Selenium in the molecular life sciences
2.7 The systems biology of selenium and selenoproteins
Keywords: Evolution, genomics, SECIS, SPS, Archaea

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Introduction: Rapid technological advances have brought an unprecedented wealth of sequences from organisms across the whole tree of life. Through the lens of computational genomics, these resources can provide insights about biological processes at evolutionary timescales. When applied to the biology of selenium, these studies can outline the evolution of selenoprotein genes, elucidate the origin of the selenocysteine pathway, and delineate new roles for its components.

Method: In the last few years we developed several tools for the computational identification of selenoproteins, SECIS elements and Sec machinery genes [1-3]. We applied these methods for evolutionary studies at diverse scales, from a relatively recent history [4] to the origin of Life itself [5,6].

Result: Through the study of SPS proteins (selenophosphate synthetases), we produced the first portrait of selenium utilization traits encompassing all sequenced organisms at once [5]. We uncovered a peculiar process of gene duplication/subfunctionalization explaining the multiple appearances of SPS1 proteins in metazoans. We also discovered selenoproteins in Lokiarchaeota, an archaeal lineage related to the first eukaryotes [6], which shed light on the origin of the eukaryotic Sec encoding system. Despite having typical archaeal selenoproteins, Lokiarchaeota possess SECIS elements strikingly similar to those of eukaryotes, but exhibit no obvious SBP2 partner. We traced back this proto-eukaryotic structure to the archaeal SECIS of one specific selenoprotein, which shows its core key motif in methanogenic archaea.

Discussion: Our studies provide broad insights on selenium utilization and selenoprotein function, and frame them in the context of evolutionary processes at large.

Selected references
O105 - The human selenomic microbiome
2. Selenium in the molecular life sciences
2.7 The systems biology of selenium and selenoproteins
Keywords: human microbiome, selenium utilization, selenoprotein prediction, bSECIS

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Introduction: The human microbiota comprises bacteria, archaea, viruses, and microbial eukaryotes that play fundamental roles in human health and disease. Selenium is an essential trace element for humans and for organisms inhabiting our body. For the first time, we characterized the composition and distribution of selenoproteins and other selenium utilization forms in the human microbiome.

Method: We analyzed whole metagenome assemblies from the Human Microbiome Project (HMP)[1]. The HMP provides a catalog of the microbial communities across five body areas from a large cohort of healthy adult individuals. By applying computational methods developed in our group, we identified and quantified selenoproteins and other genetic markers of selenium utilization.

Result: We found that selenoprotein genes are commonly present in the bacterial microbiota throughout the entire human body. Selenoproteins are remarkably abundant in the oral cavity, and are depleted in stool samples, where the usage of selenium as cofactor is surprisingly more spread than selenocysteine. By applying our new method bSeblastian, we also identified novel bacterial selenoproteins.

Discussion: The effect of selenium in human health cannot be fully understood without taking into account the human microbiome. Our results provide a profile of selenium utilization by the microbiota inhabiting our body. Selenium availability might be important for the regulation, not only of the selenoproteins encoded in our genome, but also for those encoded in our microbiome.

Selected references
O106 - Selenoprotein extinction in Drosophila occurred concomitantly to genome catastrophes

2. Selenium in the molecular life sciences
2.7 The systems biology of selenium and selenoproteins

Keywords: Drosophila, selenoprotein loss, genome sequencing, comparative genomics, evolution

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Introduction: Drosophila willistoni was the first animal discovered to lack selenoproteins [1]. The three selenoproteins of D. melanogaster were either lost or converted to cysteine homologs in D. willistoni, as the selenoprotein synthesis machinery degenerated. Other selenoproteinless animals have been identified [2,3,4,5], but little is known about the evolutionary path leading to the complete loss of selenocysteine.

Method: We sequenced eight Drosophila genomes from the saltans group, the sister lineage of D. willistoni, and performed RNAseq in ten Drosophila species. Using a multispecies design, we further conducted various functional experiments at diverse levels, ranging from biochemical to behavioural assays.

Result: From the newly sequenced genomes, we identified three additional selenoprotein extinction events in Drosophila. Although independent, these events happened in a single lineage: saltans/willistoni. Interestingly, various other genomic features set saltans/willistoni apart from the rest of Drosophila (most remarkably their lower GC content). Based on comparative transcriptomics and other observations, we hypothesized that saltans/willistoni underwent an ancestral event of adaptation to hypoxia, setting in motion a “genome catastrophe” that triggered, among other effects, the eventual loss of selenoproteins. We show various lines of evidence supporting our hypothesis, including assays in which saltans/willistoni species exhibit phenotypes similar to a D. melanogaster strain artificially adapted to hypoxic conditions [6].

Discussion: By high-throughput sequencing and comparative analyses, we delineated the evolutionary events leading to selenocysteine losses in Drosophila, and we propose a comprehensive explanation involving major adaptive changes occurred at the root of the saltans/willistoni lineage.

Selected references
O107 - Metabolomics of selenium

2. Selenium in the molecular life sciences
2.9 Additional and emerging topics of selenium in molecular life science
Keywords: selenoproteins, metabolite profiling, lifespan, miRNA

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Introduction: Selenium is an essential trace element in mammals due to its presence in 25 selenoproteins in the form of selenocysteine residue. Both selenium deficiency, which leads to compromised selenoprotein functions, and high levels of selenium, resulting in selenium toxicity, have been associated with oxidative stress and adverse health effects, although molecular mechanisms are not fully understood.

Method: Here, we subjected mice to diets spanning selenium deficiency and toxicity and examined global changes in metabolism using metabolite profiling and miRNA assays. Long-term effects of dietary selenium were also investigated.

Result: Most significant changes in metabolite patterns were associated with selenium deficiency, which altered redox homeostasis, reduced the levels of free amino acids, and affected miRNA expression. Dietary selenium more strongly affected liver metabolism than brain metabolism.

Discussion: These changes, along with the activation of metabolic pathways associated with xenobiotic metabolism and redox homeostasis, were consistent with a pro-longevity state induced by selenium deficiency. Indeed, both male and female mice subjected to selenium deficiency for their entire adult life, while dramatically reducing selenoprotein expression, did not shorten lifespan. Thus, selenium deficiency, while associated with adverse health effects, is compensated for by pro-longevity mechanisms supporting normal lifespan of mice.
O108 - Selenium-encoded chemical proteomics

2. Selenium in the molecular life sciences
2.7 The systems biology of selenium and selenoproteins
Keywords: chemical proteomics, selenoprotein, activity-based probes, selenocysteine

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Introduction: Selenium is one of the indispensable trace elements for human health and its dominant form in human body is selenocysteine, which serves as critical active-site residues in selenoproteins in regulating redox balance. Selenium excess and deficiency are both implicated with severe diseases. However, it remains challenging to profile selenoproteins by traditional shotgun proteomics tools due to its low abundance and versatile activity states.

Method: We have developed a computational program to detect the characteristic isotope envelope of selenium-containing peptides from complex proteomic data and use these information to guide the proteomic analysis in a targeted mode. With a selenium-encoded enrichment tag, this method can be seamlessly combined with current chemical proteomic technologies using activity-based or metabolic analog probes to profiling sites of post-translational modifications.

Result: We showed that the targeted analysis can dramatically improve the sensitivity of detecting the sites of post-translational modification or probe adduction, as well as to improve identification of natural selenoproteins in cellular and tissue proteomes. The method could also be applied to aid target analysis of selenium-containing metabolites.

Discussion: Our selenium-encoded chemical proteomic strategy will be a novel tool and great resource for in-depth proteomic analysis of selenoproteins and other selenium-containing biomolecules, which will aid functional studies of selenoproteins and their implications in human health.
O109 - Characterization of Atlantic salmon (Salmo salar) selenoproteins using bioinformatics and hyphenated techniques

Introduction: Selenium is an essential element, important for the function of several selenoproteins. The number of selenoproteins may vary between different animal species. For example, compared to vertebrates, teleost fish such as Atlantic salmon (Salmo salar) feature a high number of selenoproteins. The biological functions of many of these selenoproteins are yet to be elucidated. With the recently published Atlantic salmon genome (Lien et al. 2016) it has become possible to predict selenoproteins using bioinformatics tools. However, a pure in silico description of the Atlantic salmon selenoproteome is challenging. The selenoproteins’ key amino acid, selenocysteine, is coded for by messenger RNA (mRNA) comprising the exact same nucleic acid triplet (TGA) that signals the termination of translation into proteins. This may lead to selenoproteins being missannotated or completely missed. Therefore, the validation of the theoretical prediction through analytical data is essential.

Method: Recently, hyphenated analytical techniques, which employ laser ablation coupled to inductively coupled plasma mass spectrometry (LA-ICP-MS) and high performance liquid chromatography (HPLC) coupled to both ICP-MS and high-resolution tandem mass spectrometry (HR-MS) to analyse selenoproteins in human cells (Bianga et al. 2014). In the present study, a similar methodological approach for the analysis of the salmon liver selenoproteome was implemented.

Result: By combining the theoretical genome derived bioinformatics data with the results from the analytical techniques, we expect to obtain a phenotypically anchored description of the salmon selenoproteome.

Discussion: These findings will ultimately contribute to the advancement of our understanding of selenium biology in marine organisms.

Selected references
O110 - The cellular location of selenoproteins in human prostatic tissue and their role in prostate cancer.

Keywords: prostate, cancer, SELENOF, GPX1, SBP1

Introduction: Two selenocysteine-containing proteins, SELENOF and GPX1, and a non-selenocysteine-containing protein that binds selenium, SBP1, have been implicated in cancer etiology. Given human epidemiological data indicating an inverse association between selenium intake and prostate cancer risk, these proteins were imaged in human cell lines and prostate tissue obtained from men who underwent radical prostatectomy.

Method: Specificity of anti-SELENOF, -GPX1 and -SBP1 antibodies was verified by western blot analysis as well as by peptide blocking. Proteins were visualized by either confocal microscopy of selected cell lines or immunohistochemistry of human prostate tissue microarrays obtained from the Cooperative Prostate Cancer Tissue Resource.

Result: SBP1 was localized to the nucleus and cytoplasm in prostatic tissues, with the nuclear/cytoplasm ratio being inversely associated with tumor grade, while lower levels of SBP1 were associated with recurrence (OR 0.38, 95% CI 0.20-0.72). SELENOF was localized to the endoplasmic reticulum in breast and prostate cancer cell lines, but principally localized to the outer cell membrane in benign prostate tissue, as well as in immortalized or primary human prostate cultured cells. There was dramatically lower levels in benign tissue vs. tumor ($p=1x10^{-7}$). GPX1 was primarily localized to the cytoplasm and mitochondria in all cells examined, although a small number of cells exhibited nuclear localization that occasionally segregated asymmetrically in dividing cells.

Discussion: These data indicate that cellular localization and abundance should be considered when trying to understand the role these proteins play in prostate cancer.

Selected references
O111 - Dietary Selenium Deprivation Oppositely Impacts Longevity and Healthspan in Telomere Dysfunctional Mice

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease
Keywords: Aging, Genome stability, metabolism, hormesis, selenoprotein

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Introduction: Selenium (Se), a trace metalloid essential for life, plays nutritional and physiological roles mainly through selenoproteins. While telomere attrition provokes DNA damage response and replicative senescence, such an aging process is restricted in mice due to their comparatively long telomeres.

Method: To circumvent this limitation and employ a mouse model with humanized telomeres, weanling third generation telomerase RNA component knockout mice carrying short telomeres were fed a Se-deficient, Torula-yeast basal diet or the diet supplemented with 0.15 ppm Se as sodium selenate throughout their life.

Result: Dietary Se deprivation delayed wound healing and accelerated a wide range of age-related degenerations including osteoporosis, grey hair, alopecia, cataract, and hyperglycemia. Surprisingly, long-term dietary Se deprivation paradoxically promoted longevity in both sexes. Plasma microRNA profiling revealed a circulating signature of Se deprivation and aging, and subsequent ontological analyses predicted dominant changes in metabolism. Consistent with this, dietary Se deprivation accelerated age-dependent declines in glucose tolerance, glucose-stimulated insulin production and insulin sensitivity, as well as DNA damage and senescence responses in the pancreas. Selenotranscriptomic and metagenomic analyses identified key selenoprotein mRNAs and gut bacteria in the response to dietary Se deprivation in males and females at old age.

Discussion: These results suggest a novel model of aging with conceptual advances, whereby Se at low or nutritional levels of intake may be considered a hormetic chemical and decouple healthspan and longevity.

Selected references
O112 - Selenium and Cataract

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease

Keywords: SelR, Sep15, cataract, endoplasmic reticulum stress, oxidative stress

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Introduction: Much of selenium’s beneficial influence on health is attributed to its presence in 25 selenoproteins. Previous reports showed that Sep15 knockout mice developed a prominent nuclear cataract; selenium supplementation could slow the development of naphthalene cataract. However, the mechanism remains unclear. Cataract is the leading cause of blindness worldwide—it accounts for approximately 42% of all blindness. More than 17 million people are blind because of cataract. Approximately 25% of the population over 65 and about 50% over 80 have serious loss of vision because of cataract. Meanwhile, Cataract is one of major complications of both type 1 and 2 diabetes, a severe metabolic disorder characterized by hyperglycemia. Although the precise mechanism of cataract formation is not well understood, lens epithelial cell apoptosis might be a common cellular basis for initiation of noncongenital cataract formation in humans and animals and oxidative stress is believed to play a major role in cataract formation.

Result: This presentation will review the research progress of selenium and cataract, especially, the roles of SelR and Sep15 in cytoprotection against oxidative stress and endoplasmic reticulum stress-induced lens epithelial cell damage and apoptosis in human lens epithelial cells and rats, the effects of SelR or Sep15 gene knockdown on the FGF/MAPK and FGF/PI3K signaling pathways, and selenium supplementation on redox and endoplasmic reticulum homeostasis.

Discussion: Our results imply that these selenoproteins play a role in protecting lens epithelial cells against differentiation dysfunction and selenium supplementation might maintain redox and endoplasmic reticulum homeostasis via increased expression of selenoproteins.

Selected references
O113 - Role of selenoprotein P in Alzheimer's disease

3. Selenium in animal and human health and disease
3.7 Additional and emerging topics of selenium in health or disease

Keywords: Selenoprotein P (SeP), Alzheimer's disease (AD), metal, Aβ, tau

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Introduction: Dyshomeostasis of metal ions clearly occurs in AD brains. Previously, we reported SeP inhibited metal-induced aggregation and neurotoxicity of both Aβ and tau by its excellent metal binding properties in vitro.1-3 In this study, we evaluated the potential of SeP in the treatment of cognitive dysfunction of triple transgenic AD mice.

Method: Brain-targeted SelP-H gene was delivered via recombinant adeno-associated virus serotype 9 (rAAV9). Cognitive ability and activity of the AD mice were examined by Morris Water Maze test and Open Field test. Concentration and distribution of metal ions in the brains were determined by ICP-MS and SR-XRF. Protein levels of SeP, Aβ, tau, p-tau, GSK3β, PP2A, APP, BACE1 and synaptic proteins were measured by western blot.

Result: Overexpression of SeP-H: 1) significantly improved the cognitive ability and activity of AD-mice; 2) decreased Aβ deposition in the brains but did not alter expression levels of APP and Bace-1; 3) inhibited tau hyperphosphorylation and NFT formation by decreasing GSK3β but promoting PP2A expression; 4) mitigated the decrease of synaptic proteins; 5) changed the concentration and distribution of metal ions in the mouse brains.

Discussion: Our study suggested that SeP ameliorates Aβ and tau pathology by regulating metal homeostasis in the brains and finally improved cognitive decline of AD model mouse.

Selected references
O114 - Role of selenoprotein P in function of pancreatic β cell: Improving effects of neutralizing antibody

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease

Keywords: Selenoprotein P, diabetes, hepatokine, neutralizing antibody, insulin secretion

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Introduction: Selenoprotein P (SeP), a selenium (Se)-supply protein, is synthesized mainly by the liver and functions as a Se-supply protein to maintain Se levels in peripheral tissue. SeP is identified as a hepatokine, causing insulin resistance in type 2 diabetes. Thus, the suppression of Se-supply activity of excess SeP might improve insulin resistance and glucose metabolism.

Method: We established neutralizing antibodies (Abs) with activity against Se-supply of SeP by using purified human SeP protein and cultured cells. Human SeP protein was administrated to induce glucose intolerance and insulin resistance in treated mice.

Result: Anti-human SeP monoclonal Ab AE2 was identified as having strong neutralizing activity. Administration of AE2 to mice significantly improved glucose intolerance and insulin resistance that were induced by human SeP administration. Furthermore, excess SeP administration significantly decreased in pancreas insulin levels and high glucose-induced insulin secretion, which were improved by AE2 administration. The impaired effects of excess SeP on insulin secretion and protective effects of AE2 were also observed in isolated islets and a MIN6 cell model of pancreatic β cells. Neutralizing Ab of mouse SeP improved glucose intolerance and insulin secretion in a mouse model of diabetes, such as KKAy mice and mice fed a high-fat, high-sucrose diet.

Discussion: The present study demonstrated that administration of SeP-neutralizing Ab can improve glucose metabolism, insulin resistance, and insulin secretion in vivo and established a molecular basis for the development of therapeutic agents that target SeP for treatment of type 2 diabetes.

Selected references

*Fig. Improving effects of SeP-neutralizing antibody for insulin resistance and insulin secretion*
SeP expression was induced by high glucose and high lipids, and excess SeP increased insulin resistance in skeletal muscle and decreased insulin secretion in the pancreas. Administration of SeP-neutralizing Ab can improve glucose metabolism, insulin resistance, and insulin secretion in vivo.
O115 - Tissue-specific pools of Selenoprotein P differentially modify colitis-associated carcinogenesis

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease

Keywords: Selenoprotein P, colitis-associated carcinoma, intestine, cancer

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Introduction: Patients with inflammatory bowel disease demonstrate nutritional Se deficiencies and are at increased risk for colon cancer due to heightened inflammation and oxidative stress. Previously, we determined that global loss of the selenium-containing protein, Selenoprotein P (SELENOP), exacerbates experimental colitis-associated cancer (CAC). However, because SELENOP loss altered both immune and epithelial cellular function, we sought to delineate tissue-specific contributions of SELENOP to intestinal inflammatory carcinogenesis.

Method: Conditional Selenop floxed mice were crossed with Albumin-Cre, Villin-CreER, or LysM-Cre lines to delete Selenop in the liver, intestinal epithelium, or myeloid lineage, respectively. Resulting cohorts were placed on an azoxymethane/dextran sodium sulfate (AOM/DSS) experimental CAC protocol.

Result: Although liver-specific Selenop deletion decreased levels of colonic Se, it failed to modify AOM/DSS-mediated tumorigenesis. Conversely, complete loss in the intestinal epithelium increased tumor incidence, number, size, degree of dysplasia, and DNA damage, while no effect was observed in heterozygous mice. However, much like the global Selenop model, hemizygosity in the myeloid population increased tumor number and degree of dysplasia, while complete loss resulted in smaller, less aggressive tumors than both WT and heterozygous cohorts.

Discussion: Despite the vast majority of total body SELENOP being produced by the liver, these results implicate locally-expressed SELENOP as the primary mediator of experimental CAC. As epithelial-specific loss increased DNA damage and apoptosis, this pool of SELENOP likely attenuates tumor initiation through local antioxidant mechanisms. However, myeloid-specific loss closely recapitulated the results observed with the global SELENOP model, suggesting that these SELENOP pools are the largest contributor to inflammatory-carcinogenesis.

Selected references
O116 - Selenoprotein P in cord serum: wide disparity between ELISA and HPLC

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease
Keywords: SELENOP, delivery, cord blood.

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Introduction: Selenoprotein P (SELENOP) is the largest fraction of selenium (Se) in blood1. The maternal-neonatal transfer of SELENOP at the time of birth has not been thoroughly elucidated2. Our aim was to determine total Se and SELENOP in healthy women and their newborns at delivery.

Method: A cross-sectional study included 83 healthy mother-baby couples. Total Se and SELENOP were measured in maternal serum and umbilical-cord serum by an in-series three-dimensional size exclusion/affinity high-performance/anion exchange liquid chromatography (3D/SE-AF-AEC-HPLC). SELENOP concentrations were also measured by ELISA.

Result: Total Se was significantly higher in maternal serum compared to cord serum (68.9±15.2 and 56.1±14.6 µg/L respectively; p<0.01). SELENOP concentration determined by HPLC significantly correlated between mothers and newborns, although they also show significant differences (42.5±9.5 vs 28.1±7.7 µg/L in maternal serum and cord serum respectively, p<0.01). SELENOP concentration determined by ELISA was significantly different in maternal and cord serum (38.27±12.34 µg/L and 1.93±0.98 µg/L respectively, p<0.01). Although SELENOP concentration in maternal serum measured by HPLC and ELISA were not significantly different, there was a significant difference between these two methods of determination in cord serum (28.06 ± 7.69 µg/L by HPLC and 1.93 ± 0.98 µg/L by ELISA, p<0.01). There was no correlation between SELENOP concentrations measured by HPLC and ELISA in either maternal or cord serum3.

Discussion: Total Se and SELENOP concentrations are higher in mother than newborns. SELENOP concentration determined by ELISA in cord serum does not reflect the real Se status in newborns. Circulating SELENOP in newborns needs further investigation.

Selected references
O117 - Expression and activity of enzymes of selenium metabolism in the selenoprotein P knockout mouse

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease

Keywords: Sepp1, Scly, CTH, CBS, TrxR1

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Introduction: Selenoprotein P (Sepp1) is produced primarily in the liver and transports selenium (Se) to other tissues. Male mice lacking Sepp1 (Sepp1 KO) have decreased total hepatic selenium. We hypothesized that absence of Sepp1 activates the transsulfuration pathway whereby selenocompounds are decomposed by either selenocysteine lyase (Scly), cystathionine γ-lyase (CTH), cystathionine β-synthase (CBS), or thioredoxin reductase 1 (TrxR1) to provide selenide for selenoprotein production. To further understand the role of Sepp1 in hepatic Se metabolism, we evaluated the expression and activity of several enzymes involved in selenium metabolism in the Sepp1 KO mouse model.

Method: Juvenile male Sepp1 KO and C57/J wild type (WT) mice on an adequate Se diet were probed for Scly, CTH, CBS, and TrxR1 expression in the testes, lung, kidney, and liver. Scly activity was measured in the same tissues.

Result: Sepp1 KO mice exhibit a decrease in liver mRNA expression of Scly, however, there was no significant difference in Scly protein expression in the testes, lung, kidney, and liver. Expression of CTH and CBS in the liver does not differ significantly in Sepp1 KO versus WT mice, while there was a ~40% decrease in TrxR1 protein expression.

Discussion: The transsulfuration pathway is not activated in the liver of the Sepp1 KO mice through Scly, CTH, and CBS, while TrxR1 upregulation suggests selenite is more efficiently utilized.
O118 - The Role of Selenoprotein K in Progression and Metastasis of Melanoma

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease

Keywords: cancer, melanoma, migration, metastasis, calcium

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Introduction: Selenoprotein K (SELENOK) has been implicated in calcium flux in immune cells during activation and migration, and we hypothesized that it also plays a critical role in the progression of primary and secondary melanoma tumors.

Method: We developed an in vitro human melanoma cell model as well as a mouse model of spontaneous melanoma. For the first studies, we used CRISPR/Cas9 to generate a SELENOK-null clone in the NCI-60 validated human melanoma cell line, SK-Mel28. For in vivo studies, we utilized a spontaneous melanoma transgenic (Tg) mouse model involving the overexpression of the glutamate receptor 1 (Grm1) in melanocytes. We generated littermate controlled experiments consisting of Grm1/SELENOK+/+, Grm1/SELENOK+/-, and Grm1/SELENOK−/− mice that could be compared for the development of primary and secondary tumors.

Result: The SELENOK-null SK-Mel28 cells were compared to wt control SK-Mel28 cells for proliferation, migration, and calcium flux. All three functions were impaired in the SELENOK-null cells compared to controls. Results showed for both males and females the Grm1/SELENOK−/− mice had significantly lower levels of tumors compared to Grm1/SELENOK+/+ and Grm1/SELENOK+/- littermate controls, which both were similar to each other. Also, the Grm1/SELENOK−/− mice had significantly lower levels of metastasis to draining lymph nodes compared to Grm1/SELENOK+/+ and Grm1/SELENOK+/- littermate controls, which both were similar to each other.

Discussion: These data suggest that SELENOK is crucial for melanoma progression and this selenoprotein may serve as a therapeutic target for treating melanoma in humans.
O119 - Targeting thioredoxin reductase 1 as a basis for anticancer therapy

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease
Keywords: thioredoxin reductase, TrxR1, TXNRD1, cancer, SecTRAPs

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Introduction: Many studies have implicated that the thioredoxin system may be important in cancer progression, but the molecular mechanisms underpinning this relation are highly intricate and complex. Because thioredoxin reductases are easily targeted by electrophilic anticancer agents it is possible that the inhibition of these enzymes can often be a correlative finding rather than a mechanistic basis for anticancer efficacy.

Method: We have studied the targeting of cytosolic thioredoxin reductase (TrxR1, TXNRD1) in pure form, cellular context and in mouse models with the overriding aim of asking whether the enzyme may be a bona fide anticancer target or not. This presentation will present some of our results from these studies.

Result: Using newly developed small molecule inhibitors of TrxR1 with hitherto unsurpassed selectivity in TrxR1 over TrxR2 inhibition, our results suggest that TrxR1 targeting, and the conversion of the enzyme to SecTRAPs (Selenium compromised thioredoxin reductase-derived apoptotic proteins) can indeed serve as a mechanistic basis for anticancer therapy.

Discussion: Targeting of TrxR1 with small molecules can provide anticancer efficacy without overt toxic side effects, at least in mouse models. Challenges for the further development of this principle of therapy include complete assessment of toxicology profiles, combinatory effects with classical chemotherapeutic agents, additional SAR analyses and development towards trials in human subjects.

Selected references


O120 - Redox regulation of protein kinase C by selenium and selenoprotein thioredoxin reductase influences the cancer-preventive efficacy of selenium

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease
Keywords: thioredoxin reductase, protein kinase C, prostate cancer, cancer prevention

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Introduction: Although experimental studies demonstrated cancer prevention by supplemental dietary selenium, clinical trials did not support this. Previously, we have shown that methylselenol, active metabolite of selenium, reacts with membrane lipid hydroperoxides and converts to methylseleninic acid (MSA). This locally generated MSA selectively oxidizes critical cysteine sulfhydryls and inactivates PKC. Due to a catalytic role of selenium in the peroxidatic redox cycle, a low concentration of selenium inactivates membrane-bound PKC that is compartmentally separated from cytosolic glutathione. Thioredoxin (TRX), which binds with a high affinity to PKC, along with selenoprotein thioredoxin reductase (TR), reverses the inactivation of PKC.

Method: We used various human prostate cancer cell lines, with a common lineage, exhibiting different degrees of malignancy to determine whether there is a correlation in the distribution of PKC isoenzymes and the TRX-TR system and sensitivity of tumor cells to selenium.

Result: We found that advanced malignant cells overexpressing TR1, TR2, and TRX may escape the cancer-preventive actions of selenium by reversing selenium-induced inactivation of PKC. Low concentrations of selenium were required to inactivate antiapoptotic PKC isoenzymes that have a higher number of vicinal thiol residues and to induce apoptosis. However, higher concentrations of selenium were required to inactivate less sensitive proapoptotic enzymes that have limited number of critical thiol residues. This makes tumor cells, paradoxically, resistant to selenium at higher concentrations.

Discussion: These studies suggest that in some cases, certain selenoproteins may counteract selenometabolites to protect tumor cells. These studies also explain the intriguing U-shaped curve seen with dietary selenium intake and cancer prevention.

Possible mechanism for biphasic effects of selenium in inducing apoptosis in tumor cells

Selected references
O121 - Thioredoxin reductase 2 inhibition by tamoxifen-like metallocifens drives Jurkat cells to apoptosis

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease
Keywords: Thioredoxin reductase, ROS, cancer cells, mitochondria, selenoenzyme

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Introduction: The thioredoxin system is a chief regulator of cell redox homeostasis and is formed by NADPH, thioredoxin and the selenoenzyme thioredoxin reductase (TrxR) which is often overexpressed in cancer cells especially in tumors resistant to classic chemotherapeutics. Therefore, the research for potential specific cytosolic and mitochondrial TrxRs inhibitors is a challenging task to contrast chemoresistance.

Method: The mechanism of action of three tamoxifen-like metallocifens (TLMs) of Fe, Ru, Os was investigated. Firstly, the activity of the compounds was assessed on isolated enzymes. Afterward, TLMs effect on a lymphoblastoid cell line (Jurkat cells) was estimated. In particular, their impact on TrxRs, on cellular redox state and on mitochondrial physiology was explored.

Result: TLMs act as pro-drugs showing a strong inhibitory activity on isolated TrxRs, after enzymatic oxidation with H₂O₂/HRP. The inhibition occurs through the interaction with the selenocysteine in the active site of TrxRs. In Jurkat cells, TLMs selectively inhibit mitochondrial TrxR hindering thioredoxin reduction. The consequent ROS rise and the decrease of mitochondrial membrane potential (MMP), drive cytochrome c release and caspase 3 activation. Moreover, metal quantitation demonstrated that TLMs accumulate mainly in the mitochondrial fraction and to a lower extent in the cytosolic compartment.

Discussion: TLMs are endowed with two functional groups: the amino chain, positively-charged at neutral pH which favors their mitochondrial accumulation, and the metallocene unit, characterized by redox properties that promotes reactivity with TrxR. Indeed, according to their subcellular localization, TLMs determined a preferential inhibition of mitochondrial TrxR and showed impairment of mitochondrial functions inducing apoptosis in Jurkat cells.

Selected references
O122 - Dietary selenium and the 15 kDa selenoprotein influence initiation/promotion of colon carcinogenesis

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease

Keywords: colon cancer, intestinal microbiome, 15 kDa selenoprotein

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Introduction: The 15kDa selenoprotein (SELENOF) plays roles in cancer prevention and promotion. Previously, SELENOF knockout (KO) mice were shown to be resistant to formation of chemically-induced pre-neoplastic lesions [1].

Method: In our current inflammatory tumorigenesis study, KO mice and littermate controls were maintained on diets with deficient (0.02 ppm) or adequate (0.1 ppm) dietary selenium (Se), and injected with azoxymethane (AOM) or saline, and treated with dextran sulfate sodium (DSS) or water.

Result: Formation of pre-neoplastic lesions was dramatically reduced in KO mice compared to controls. Additionally, fewer KO mice developed tumors when Se-deficient, but formed a similar number of tumors as controls when Se-replete. Thus, KO mice may be protected against cancer initiation, but not against tumor formation, and dietary Se may modify the outcome. Further analyses suggest that CYP2E1, the hepatic enzyme responsible for activating AOM to its mutagenic form, has lower expression in untreated KO mice than controls under Se-deficient conditions, whereas no differences were detected in AOM/DSS-treated mice. Because intestinal microbiota are suspected to affect colon tumorigenesis, bacterial DNA was isolated from mouse feces. Subsequent amplification of the 16S rRNA gene and sequencing with Illumina MiSeq revealed an increased abundance of Bacteriodetes in KO mice fed Se-deficient diets, and a substantial increase in Verrucomicrobia, especially the mucin-degrading species Akkermansia muciniphila, in Se-adequate KO mice.

Discussion: Levels of both SELENOF and dietary selenium have substantial effects on colon tumorigenesis and inflammation, which may be modulated by the intestinal microbiome. Their contribution to the regulation of colorectal cancer will be further examined.

Selected references
O123 - Selenoprotein T is a novel neuroprotective antioxidant enzyme in Parkinson’s disease

Keywords: Selenoprotein, neuroprotection, neurodegeneration, dopamine, transgenic mice

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Introduction: Oxidative stress is central to the pathogenesis of Parkinson’s disease (PD), but the mechanisms involved in the control of this stress in dopaminergic cells are not fully understood. There is increasing evidence that selenoproteins play a central role in the control of redox homeostasis and cell defense, but the precise contribution of members of this family of proteins during the course of neurodegenerative diseases is still elusive.

Method: We used the SH-SY5Y dopaminergic cell model and brain-specific selenoprotein T (SelT)-deficient mice to demonstrate the importance of this protein in neuroprotection.

Result: In the SH-SY5Y cell model of dopaminergic neurons, both silencing and overexpression of SelT affected oxidative stress and cell survival. Treatment with PD-inducing neurotoxins such as MPTP or rotenone triggered SelT expression in the nigrostriatal pathway of wild-type mice, but provoked rapid and severe parkinsonian-like motor defects in conditional brain SelT-deficient mice. This motor impairment was associated with marked oxidative stress and neurodegeneration, and decreased tyrosine hydroxylase activity and dopamine levels in the nigrostriatal system. Finally, in PD patients, we report that SelT is tremendously increased in the caudate putamen tissue.

Discussion: These results reveal the activity of a novel selenoprotein enzyme that protects dopaminergic neurons against oxidative stress and prevents early and severe movement impairment in animal models of PD. Our findings indicate that selenoproteins such as SelT play a crucial role in the protection of dopaminergic neurons against oxidative stress and cell death, providing insight into the molecular underpinnings of this stress in PD.

Selected references
O124 - Selenium Atom-specific Functionalization of Nucleic Acids for Structure and Function Studies

2. Selenium in the molecular life sciences
2.8 Selenium based biotechnological applications

Keywords: selenium derivatized nucleic acid, modified DNA & RNA, crystal structural biology, molecular sensor

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Introduction: There are total five essential elements (H, C, N, O and P) in nucleic acids. Single Se-atom replacement (or atom-specific mutagenesis) of nucleic acids means the substitution of O with Se atom. Atom-specifically functionalized nucleic acids by introducing the sixth element (such as Se) can offer nucleic acids with many unique and novel properties (such as facilitated crystallization and phase determination) without significant perturbation of 3D structures of nucleic acids and their protein complexes.

Method: Nucleic acids possess not only the ability to store genetic information and participate in transcription and translation, but also the capacity to adopt well-defined 3D structures, which can be readily adjusted to meet various functional needs (such as catalysis and therapeutics). Although the importance of numerous nucleic acids in catalysis, gene expression, protein binding and therapeutics has been acknowledged by the entire scientific society, current understanding of nucleic acid-protein functions and structures is still limited, especially high-resolution structures.

Result: Thus, this novel atom-specific mutagenesis provides important tools to investigate nucleic acid structure/folding, recognition and catalysis, to study nucleic acids and their protein interactions, to improve biochemical and biophysical properties of nucleic acids, to facilitate gene silencing and RNA & DNA nanotechnology, and to explore potential nucleic acid therapeutics and diagnostics.

Discussion: Our presentation will focus on the most recent selenium-atom functionalization of nucleic acids and their potential applications in 3D structure-and-function studies and anticancer therapeutics in molecular medicine.

Selected references
O125 - Interrelationships among SELENOF genotype, cellular localization, serum selenium and race in prostate cancer

3. Selenium in animal and human health and disease
3.6 Clinical genetics of selenium or selenoprotein-encoding genes

Keywords: prostate, cancer, SELENOF, genetics, race

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Introduction: Polymorphisms in the endoplasmic reticulum (ER) resident selenoprotein SELENOF are associated with prostate cancer mortality and are 5-fold more prevalent in African Americans, who are also at greater risk of dying of the disease. A role of SELENOF in prostate cancer was investigated in human prostate tissues and cell lines.

Method: Plasma selenium levels were quantified by atomic absorption spectroscopy. DNA from 129 men who had undergone prostatectomy were genotyped for SELENOF and the interactions between SELENOF genotype, serum selenium, and race were assessed. Levels and cellular location of SELENOF were analyzed in cultured cells by western blotting and in human tissue microarrays by immunohistochemistry scored by computer-assisted image analysis.

Result: SELENOF predominated in the outer cell membranes (OCM) of cells in benign glands and in primary benign human epithelial cells, but not in human cancer-derived cell lines and tumors tissue. Protein levels were significantly lower, and generally absent in the OCM in cancer compared to benign prostatic tissue (p=1x10^{-7}). SELENOF levels were lower in tumor cells from African Americans (n=33) compared to Caucasians (n=295) (p<0.02). The SELENOF allele that was more frequent in African Americans (n=33) compared to Caucasians (n=295) (p<0.02). The SELENOF allele that was more frequent in African Americans was associated with both higher prostate specific antigen levels (p<0.05) and Gleason score. African American men in this cohort had significantly lower mean serum selenium levels (115.24 ±3.04 ng/mL) than Caucasians (139.18 ±2.41 ng/mL, p<0.01).

Discussion: SELENOF may have a non-ER function in normal prostate epithelium. Genotype and selenium levels may contribute to the prostate cancer racial disparities observed in African Americans.
O126 - Selenocysteine and the Genetic Code

2. Selenium in the molecular life sciences
2.9 Additional and emerging topics of selenium in molecular life science

Keywords: genetic code, selenoprotein synthesis, synthetic biology, recoding

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Introduction: At the time of its elucidation the genetic code was suggested to be universal in all organisms, and the result of a ‘frozen accident’ unable to evolve further (1).

Method: Today we know 22 natural amino acids (2): selenocysteine, and pyrrolysine are directly inserted into growing poly-peptides during translation.

Result: The incorporation of selenocysteine directed by UGA requires the action of specific RNA and protein elements, a fact that has restricted engineer-ing of selenoproteins. Based on the realization that protein plasticity is a feature of living cells (3), man-made expansion of the genetic code based on orthogonal transla-tion systems (OTSs) is an active research field (4-6).

Discussion: Although the design of in vivo specific and highly active OTS systems is still not ideal (7,8), the increasing number of successful recoding strategies promises a bright future for genetic code expansion (9-12).

Selected references
O127 - Cysteine polysulfidation governed by cysteinyl-tRNA synthetases (CARSs)
2. Selenium in the molecular life sciences
2.9 Additional and emerging topics of selenium in molecular life science
Keywords: Cysteine persulfide, CARS, cysteine persulfide synthase, CPERS, bioenergetics

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Introduction: Cysteine hydropersulfide (CysSSH) occurs in abundant quantities in various organisms including mammals, yet little is known about its biosynthesis and physiological function(s). Moreover, extensive persulfide formation is apparent in Cys-containing proteins in *Escherichia coli* and mammalian cells, and believed to be the result of post-translational processes involving hydrogen sulfide-related chemistry.

Method: Prokaryotic and mammalian cysteinyl-tRNA synthetases (CARSs) were investigated for their potential CysSSH production both in vitro and in vivo.

Result: Here, we demonstrate effective CysSSH synthesis from the substrate L-cysteine, with the synthesis being catalyzed by prokaryotic and mammalian CARSs. Targeted disruption of the genes encoding mitochondrial CARSs in mice and human cells revealed the major role of CARSs in endogenous CysSSH production, which suggests that CARSs serve as the principal cysteine persulfide synthase (CPERS) in vivo. CARSs also catalyze co-translational protein polysulfidation and are involved in the regulation of mitochondrial biogenesis and bioenergetics. Preliminary results furthermore suggest that the selenium-dependent mammalian thioredoxin system may be involved in regulation of CysSSH homeostasis.

Discussion: Investigating translation-coupled persulfide production may thus lead to evolution of the central dogma of molecular biology, unravel mechanisms of aberrant redox-regulation, and suggest novel treatment options for mitochondrial dysfunction.

Selected references
O128 - Direct observation of methylmercury and auranofin binding to selenocysteine in thioredoxin reductase

2.9 Additional and emerging topics of selenium in molecular life science

Keywords: thioredoxin reductase, EXAFS, mercury, methylmercury, auranofin

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Introduction: Methylmercury species are well known as neurotoxins. However, the specific mechanisms of methylmercury toxicity are not clear. One theory connects the toxicity of mercury with its strong affinity for selenium, especially the selenocysteine site of selenoenzymes. Methylmercury has been shown to inhibit the activity of the selenoenzyme thioredoxin reductase (TrxR). The gold drug auranofin, used to treat rheumatoid arthritis but also showing anticancer activity, also is known to inhibit TrxR. Direct structural observations of the binding of these inhibitors to TrxR have been few to date.

Method: Recombinant rat TrxR, along with mutants U498C and U498S, were prepared as previously reported [1] and were reacted with stoichiometric methylmercury nitrate and with auranofin. Extended X-ray absorption fine structure (EXAFS) measurements [2] at the Se-K, Hg-L3 and Au-L3 absorption edges were conducted at the Stanford Synchrotron Radiation Lightsource.

Result: EXAFS with full multiple scattering analysis at both the Se and Hg edges demonstrated the direct and complete binding of methylmercury to the selenium site in TrxR with a Hg-Se distance of 2.47 Angstroms. The gold atom of auranofin is also observed to bind directly to the selenium.

Discussion: EXAFS provides a structural tool to investigate the binding of inhibitors with selenoenzymes without the need for crystallization or other extensive preparation. The results comprise direct evidence that both methylmercury and auranofin bind directly to selenium in thioredoxin reductase, thereby inactivating the enzyme. In the case of methylmercury, it is possible that such direct binding contributes to the toxicity of mercury.

Selected references


O129 - X-ray fluorescence imaging and X-ray absorption spectroscopy combined yield insight on mammaliam selenium biochemistry

2. Selenium in the molecular life sciences
2.9 Additional and emerging topics of selenium in molecular life science

Keywords: reductive metabolism, copper, glutathione peroxidase, ovary, female fertility

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Introduction: Our recent work has focused on various selenium species important in the diet and their relationship to disease states involving oxidative stress. Our interest stems from the apparent contradiction that while common forms of dietary selenium likely generate reactive oxygen species in vivo, epidemiology suggests that their intake reduces incidence of many diseases where oxidative stress is implicated.

Method: At the same time, selenium's high atomic number and low background abundance in tissue make it amenable to study using X-ray fluorescence imaging and X-ray absorption spectroscopy to yield spatial and chemical information respectively.

Result: I will discuss the application of these methods to track the chemistry of selenium species in both cell culture and animal disease model settings. This reveals distinct fates for amino acid forms which can be related to their observed biological effect, a reductive metabolism for selenite linked to delayed production of superoxide radical ions and interactions with copper biochemistry, as well as the discovery of a role for selenium in female reproductive function. In several of these cases, the chemistry of selenium in intact tissue samples provides detail about the redox processes that are involved, while in others, redistribution of other heavy elements is informative.

Discussion: We are then able to link these observations to more traditional biochemical data, which provides valuable context.

Selected references
Metallomics 2015, 7, 71-82
O130 - Se-speciation investigations at neural barrier (NB)

3. Selenium in animal and human health and disease
3.7 Additional and emerging topics of selenium in health or disease

Keywords: Selenium-speciation, HPLC-(CE)-ICP-DRC-MS, neural barrier, cerebrospinal fluid, ALS

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Introduction: Se-speciation helps for deeper insight into Se-metabolism and transport, important at NB or under neurological diseases. Analytical set-up of hyphenated speciation techniques with 2-D-identification of Se-compounds is described. Then applications at NB in paired serum and cerebrospinal fluid (CSF) samples are reported, finishing with comparison of Se-speciation of neurologically diseased persons vs. controls.

Method: Serum and CSF samples were subject to speciation analysis by HPLC-ICP-DRC-MS. For improved species identification Se-species were analysed serially after HPLC by CE-ICP-DRC-MS (2D approach).

Result: Paired samples had 58.39 (serum) or 0.86 (CSF , each µg Se/L). Prominent Se-species were selenoprotein-P (SePP), glutathione-peroxidase (GPx), thioredoxinreductase (TrxR), Se(IV) and Se-albumin (Se-HSA).

Relationships between Se-species from serum and CSF allowed evaluating Se-species passage across NB: SePP-serum correlated with total Se-serum when > 65 µg/L. SePP-CSF appeared independent from SePP-serum. For anti-oxidative Se-enzymes higher correlation factors ($r^2$) were calculated: GPx-serum/GPx-CSF: $r^2=0.3837$ and TrxR-serum/TrxR-CSF: $r^2=0.6293$. No correlation for inorganic Se-compounds was found proving limited representativeness of their circulating levels beyond NB[1].

Se-species-ratios (CSF/serum) were 21.4*10$^{-3}$ (TrxR) or 8.3*10$^{-3}$ (GPx), being significantly elevated compared to NB permeability factor 3.8*10$^{-3}$ (HSA).

In a hospital-referred cases-control we investigated Se-species in CSF of patients with amyotrophic lateral sclerosis, compared to reference neurological patients. We found an excess concentration of inorganic Se(IV) and reduced levels of organic Se-compounds among ALS patients[2].

Discussion: ROS-protecting enzymes GPx and TrxR seem to be shuttled across NB to brain/CSF. In ALS etiology overexposure of inorganic Se(IV) together with decreased organic Se-species may be involved in ALS etiology.

Selected references

O131 - Selenium-Binding Protein 1 in serum may signify a heart at risk

3. Selenium in animal and human health and disease
3.7 Additional and emerging topics of selenium in health or disease

Keywords: SELENBP1, diagnostics, sandwich assay, serum, heart, acute myocardial infarction

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Introduction: Selenium-Binding Protein 1 (SELENBP1) is an intracellular protein with unknown function. The objective of this study was to test for its potential presence in serum and its relation to cardiovascular disease.

Method: Monoclonal antibodies were generated and a sandwich luminometric assay for SELENBP1 was established. The assay has a limit of detection (LOD) of 10 µg/L. Serum samples of 77 patients with a cardio pulmonary bypass (CPB) during cardiac surgery as well as 113 patients with suspected acute myocardial infarction (AMI) were analyzed.

Result: Prior to surgery, serum SELENBP1 was below detection limit in all patients. Serum SELENBP1 increased during surgery in 97% of patients to detectable concentrations until 15 min after weaning from the CPB. SELENBP1 concentrations correlated positively with duration of cardioplegia (rho = 0.56, p = 0.0000004). In the patients with suspected AMI, 69% experienced an AMI or a major adverse cardiac event (MACE) until discharge. A time-resolved analysis indicated rising serum concentrations of SELENBP1 (to > 10 ng/ml) in 41% of patients. A significant rise in serum SELENBP1 was associated with a 78% risk of suffering an AMI or MACE.

Discussion: The physiological role of SELENBP1 remains unclear, but its appearance in serum may provide novel diagnostic or prognostic insights in cardiac disease. These findings and the newly established sandwich assay may help to better pinpoint the physiological functions of SELENBP1 and to elucidate the molecular mechanisms leading to its disease-related release into serum.

Selected references
Supported by Charité – Universitätsmedizin Berlin.
O132 - Of dogs and men: Review of translational impact of dog studies on selenium and prostate cancer risk

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Introduction: Dogs, like men, naturally develop prostate cancer during aging. Therefore, it provides an appropriate context to study the effects of selenium on the aging prostate prior to cancer diagnosis.

Method: In this presentation, we summarize the translational significance of research progress gained from a decade of dog studies on selenium and prostate cancer risk.

Result: In a randomized feeding trial in elderly beagles (equivalent to 65 year-old men), we showed daily supplementation with selenomethionine or selenium-yeast significantly reduced prostatic DNA damage. Our discovery of a U-shaped dose-response between toenail selenium and prostatic DNA damage in dogs remarkably paralleled data from men in North America and the Netherlands. Later, this U-curve provided a plausible explanation for the alarming increase in prostate cancers in men with highest baseline selenium that received supplementation in SELECT. Next, this U-curve guided the discovery of a non-antioxidant, anti-carcinogenic mechanism of organic selenium: “homeostatic housecleaning”, the preferential triggering of apoptosis in DNA damaged cells. Finally, we catalogued form-dependent effects of selenium on intraprostatic phenotypes (prostatic androgen levels, cell proliferation, apoptosis and DNA damage), complementing human studies in which analyses are often limited to markers in serum.

Discussion: Over the last decade, we have demonstrated that dogs have advanced our understanding of selenium’s in vivo chemopreventive actions. Dog studies can contribute important insights into dose-dependent and form-dependent effects – aspects of selenium biology that will have to be further elucidated if the burgeoning science of selenium is to be translated into effective strategies for prevention of human disease.

Selected references

O133 - High selenium induces endothelial dysfunction via endoplasmic reticulum stress

Keywords: ER stress, endoplasmic reticulum stress, endothelial dysfunction.

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Introduction: Selenium is associated with insulin resistance and may affect endothelial function, increasing the risk of type II diabetes and associated cardiovascular disease [1]. However, the molecular mechanisms involved are not clear. The endoplasmic reticulum (ER) stress response is a mechanism involved in high selenium-induced apoptosis in some cancer cells [2] and, also in the pathogenesis of insulin resistance and endothelial dysfunction [3]. Thus, we hypothesised that high selenium status causes ED through ER stress.

Method: Endothelial cells (HUVECs) and EA.hy926 cell lines were treated with selenite (0.5–10 µM) for 24 hours in the presence or absence of the ER chemical chaperone, 4-phenylbutyric acid (PBA). ER stress markers were investigated using qPCR and western blot technique. Endothelial function was assessed by the Griess assay, flow cytometry, Matrigel® and colorimetric assays. Data were expressed as S.E.M (p<0.05) vs control.

Result: High-selenium concentration (5–10 µM) compared to physiological concentration (0.5–2.0 µM) enhanced mRNA expression of ER-stress markers: activating transcription factor-4 (ATF4), CAAA/enhanced-binding homologous protein (CHOP) and X-binding box-1 (XBP-1). In addition, high selenite concentration reduced NO production and angiogenic capacity in endothelial cells. Moreover, high selenite treatment significantly (p<0.05) increased ROS production and induced apoptosis through caspase-3/7 activity. Interestingly, PBA completely reversed all the effects of high selenite on endothelial function, indicating the involvement of the ER-stress response.

Discussion: High-selenium treatment caused endothelial dysfunction through the activation of the ER-stress response. This highlights the importance of monitoring the risk of cardiovascular disease in cancer patients supplemented with high selenium as part of their therapy.

Selected references

Summary of mechanism by which high selenium induces endothelial dysfunction mediated by ER stress
O134 - Selenium and sex: competition between brain and testes for selenium results in male-specific consequences in mice and men

Introduction: Selenium (Se) is essential for normal neurological development and male fertility in mammals, setting the stage for competition between brain and testes when Se is limiting or its distribution is perturbed. Se supplementation studies in humans, and genetic knockout (KO) of selenium transport and recycling proteins in mice, unveiled the surprising findings that males exhibit more severe consequences than females when selenium homeostasis is altered. In one study, Se supplementation in humans resulted in an increased risk of type 2 diabetes (T2D) in men with high baseline Se. However, other studies found no correlation between increased Se and T2D susceptibility, and the discrepancies in these human trials are controversial.

Method: Mice with KO of the Se recycling enzyme, selenocysteine lyase (Scly) or double KO of Scly and the Se transport protein, Selenoprotein P (Sepp1) were characterized.

Result: KO mice exhibit metabolic syndrome that is more pronounced in males, whereas DKO resulted in low viability and neurological deficits including audiogenic seizures, also more pronounced in males. Pre-pubescent castration reversed the double KO neurological phenotype and partially attenuated the Scly KO metabolic phenotype. Assessment of brain and testes Se levels in these mice suggests that competition for available Se between these two tissues is a significant contributor to the observed phenotypes.

Discussion: Testosterone replacement studies are underway to investigate whether there are also hormone-dependent mechanisms at play. These studies unveil novel insights into the complex mechanisms involved in Se distribution and utilization, and may be informative regarding sex specific Se requirements in humans.
O135 - Ceramide analog S14 causes a coordinate downregulation of selenoproteins in a murine psoriasis model

3. Selenium in animal and human health and disease
3.7 Additional and emerging topics of selenium in health or disease

Keywords: Psoriasis, Selenoproteins, S14, Ceramide

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Introduction: Psoriasis is a common disorder affecting 1-2 percent of the world population. Currently, the role of selenoproteins in the pathogenesis of psoriasis is not well understood.

Method: We have previously demonstrated that ceramide analogs based upon the ant venom alkaloid, solenopsin, have anti-inflammatory effects on a well-established murine model, the KC-Tie-2 model of psoriasis. In an effort to better understand the role of selenoproteins and other proteins involved in glutathione metabolism, we compared vehicle treated skin versus S14 treated skin by gene array analysis. S14 is a 14 carbon chain containing solenopsin analog, which has potent anti-psoriatic activity in this model.

Result: Surprisingly, a coordinate downregulation of selenium containing and glutathione metabolizing proteins was observed in treated mice compared with controls. Notably, these include glutathione peroxidase 1 (Gpx1), superoxide dismutase 1 and 2, glutathione peroxidase 4 (Gpx4), and thioredoxin domain containing proteins 15 and 17.

Discussion: Gpx4 is a major regulator of ferroptosis, an iron mediated mechanism of cell death involving oxidation of unsaturated fatty acids. While psoriasis has traditionally been thought to be mediated by excess reactive oxygen, our data suggests that novel upregulation of reactive oxygen species by the S14 ceramide derivative may cause therapeutic benefit by causing ferroptosis and perhaps induction of regulatory T cells by reactive oxygen.

Selected references
O136 - Maternal nutrition and transcript abundance of selenium related genes in fetal bovine hepatic tissues at d 50 of gestation

Keywords: developmental programming, fetus, nutrition, RNA-Seq

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Method: Fourteen Angus-cross heifers were estrus synchronized and assigned at breeding to one of two dietary treatments (CON- 100% of nutrient requirements to gain 0.45 kg/d; RES- 60% of CON). At d 50 of gestation, heifers were ovariohysterectomized, and fetal livers were collected. Analysis by RNA-seq was conducted on the Illumina HiSeq 2500 platform using 50-bp paired-end reads at a depth of $2 \times 10^6$ reads/sample. Transcriptome analysis was performed using the Tuxedo Suite, and ontological analysis with DAVID 6.8. Significance was set at $P < 0.01; q \leq 0.19$.

Result: For fetal liver, a total of 80 selenium-related genes were detected and grouped into glutathione metabolism (n = 42), glutathione peroxidase (n = 11), and selenoprotein metabolism (n = 27). Of the glutathione metabolism genes, 2 (GSTA2 and GSR) were upregulated in CON, and 2 (ETHE1 and GGT5) were upregulated in RES. For glutathione peroxidase, 1 gene (LTC4S) was upregulated in RES compared with CON. In the selenoprotein metabolism, 1 gene (TRNAU1AP) was upregulated in RES compared with CON.

Discussion: These data suggest that by d 50 of gestation, a large number of genes associated with selenoproteins, glutathione metabolism, and glutathione peroxidase are being transcribed in the bovine fetal liver; of these, 6 genes were responsive to maternal nutritional treatment.
O137 - A ten-year study and practice of functional agriculture in China

Natural Biofortification Program (NBP) satellite meeting

Keywords: Functional agriculture, China, 10 years

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Introduction: Functional agriculture was firstly put forward by Qiguo Zhao, academician of Chinese Academy of Sciences, in “Agricultural Science and Technology in China: A road map to 2050” in 2008[1], and development of selenium-biofortification agriculture is an important part of functional agriculture. Dr. Xuebin Yin’s team from University of Science and Technology of China (USTC) firstly conducted systematic studies and industrialization practices of functional agriculture for 10 years[2].

Method: Hundred to million-ha-trials were conducted in more than 30 different crop varieties in different regions in China. Crop uptake models and fertilizer application specifications were developed.

Result: Functional agriculture developed a new agriculture 3.0 era following high yield agriculture and green agriculture (Fig. 1). Studies on the stability of selenium content in selenium-biofortified agricultural products collected from the demonstration bases in the past years showed that the relative standard deviation can be controlled within 30% in the same plot and the same variety (Table 1). More than 20 product technical standards and 5 local planting standards had been formulated, and 1 industrial standard will be issued in June, 2017.

Discussion: Since functional agriculture products had the biofortified trace mineral elements, it will be sold at a higher price and be beneficial to consumers. Several members of the National People's Congress proposed to develop the functional agricultural industry as a new way to reform China's agricultural supply side. Thus, functional agricultural were expressed in “Document No. 1 of China Central Government in 2017", which highly encouraged functional agricultural industries.

Selected references


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Table 1. Stability of selenium content in selenium-enriched rice collected from demonstration bases

<table>
<thead>
<tr>
<th>Demonstration bases</th>
<th>Year</th>
<th>Area (ha)</th>
<th>Sample numbers</th>
<th>Mean Se content (mg/kg)</th>
<th>Relative standard deviation (RSD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Donglin, Tianjin</td>
<td>2013</td>
<td>2000</td>
<td>52</td>
<td>0.19</td>
<td>29%</td>
</tr>
<tr>
<td>Donglin, Tianjin</td>
<td>2014</td>
<td>2000</td>
<td>52</td>
<td>0.24</td>
<td>26%</td>
</tr>
<tr>
<td>Huailai, Jiangsu</td>
<td>2014</td>
<td>30000</td>
<td>32</td>
<td>0.16</td>
<td>23%</td>
</tr>
<tr>
<td>Huailai, Jiangsu</td>
<td>2015</td>
<td>10000</td>
<td>32</td>
<td>0.18</td>
<td>21%</td>
</tr>
<tr>
<td>Baishu, Anhui</td>
<td>2013</td>
<td>3000</td>
<td>52</td>
<td>0.26</td>
<td>24%</td>
</tr>
<tr>
<td>Baishu, Anhui</td>
<td>2014</td>
<td>3000</td>
<td>52</td>
<td>0.27</td>
<td>27%</td>
</tr>
</tbody>
</table>
O138 - From Atom to Field: How and Why Plants Hyperaccumulate Selenium, and How this Affects Ecosystems

Plenary session I

Keywords: hyperaccumulation, plants, ecology, evolution, transcriptomics

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Introduction: Hyperaccumulators can sequester Se to 1.5% of dry weight from seleniferous soil. We address the questions why and how these plants accumulate so much Se, and how hyperaccumulation affects the local ecosystem.

Method: Stanleya pinnata (Brassicaceae) is our model hyperaccumulator, with S. elata and Brassica juncea as non-hyperaccumulator reference species. We study the plants under controlled lab conditions as well as in natural habitats and from atomic to field level. We employ x-ray microprobe analysis (XANES, XRF), RNA-sequencing, biochemical analysis, whole-plant physiology, plant-microbe, plant, herbivore and pollinator interactions and field surveys.

Result: Unique properties of hyperaccumulators are their preferential uptake of selenate over sulfate, the quantitative Se assimilation to methyl-selenocysteine and its S-independent transport and sequestration. The adaptive significance of hyperaccumulation appears ecological: protection from herbivores and pathogens and inhibition of neighboring plants. No evolutionary constraint of hyperaccumulation was found. Hyperaccumulators live in symbiosis with a variety of Se-tolerant ecological partners: herbivores, pollinators, neighboring vegetation, and a hyperaccumulator-specific rhizomicrobiome. The molecular mechanisms of Se hyperaccumulation in S. pinnata include constitutive high expression of selenate/sulfate transporters and sulfate/selenate assimilation genes, which may be linked to constitutively elevated levels of stress/defense hormones jasmonic acid, salicylic acid and ethylene. Stanleya pinnata also showed higher transcript levels of genes involved in oxidative stress resistance, likely upregulated by the same hormones. This may contribute to Se tolerance.

Discussion: These findings shed light on why and how Se hyperaccumulation evolved, and give insight into the profound ecological implications of hyperaccumulation at the ecosystem level.

Selected references
O139 - Selenium from ocean fish provides protection against mercury toxicity

Plenary session I
Keywords: Mercury, Fish

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Laura Raymond²
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Introduction: Since methylmercury is a highly specific irreversible inhibitor of selenoenzymes, this study examined if selenium from bigeye tuna, swordfish, and mako shark provides biologically active selenium to offset the loss that occurs through mercury-dependent sequestration of intracellular selenium.

Method: Se depleted rats were fed 10 ppm methylmercury in diets augmented with selenium from fish protein isolates from bigeye tuna (3.34 ± 0.28 µmol Se/kg), swordfish (2.17 ± 0.01 µmol Se/kg) or mako shark (2.19 ± 0.12 µmol Se/kg) for a final mercury concentration of 41.34 ± 5.50, 50.03 ± 0.43, and 45.14 ± 1.57 µmol Hg/kg respectively. Toxic effects of high MeHg exposures was assessed by growth inhibition and hind limb crossing scoring. Blood, livers, and brains were analyzed for Hg and Se concentrations.

Result: The only group that developed hind limb crossing were those fed low selenium, high MeHg diets. No effects were observed in any rats on low MeHg diets nor in any other group provided high MeHg diets. Neurotoxicity and growth inhibition was independent of total mercury present in brain, since the rats with the highest mercury bioaccumulations did not show toxic effects on either parameter.

Discussion: This study establishes that the selenium present in proteins of three representative varieties of ocean fish is highly bioavailable and fully effective in preventing onset of signs and symptoms of MeHg toxicity and that neurotoxicity and growth inhibition was independent of total mercury present in brain.
O140 - Selenium at the redox interface of the genome and exposome
Plenary session I
Keywords: Redox metabolic networks, integrated metabolomics and transcriptomics study, lipid metabolism

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Introduction: A critical redox interface exists between a living organism and its environment due to the relatively reducing internal steady state relative to the oxidizing external steady state. Selenium, a redox-active element from the environment, is used biologically to maintain the internal redox steady state but also poses a threat to the redox network structures when present in excess. We performed an integrated metabolomics and transcriptomics study of Se excess in mice to explore the central network responses to variation in this essential nutrient.

Method: C57BL6 male mice were treated with sodium selenate (4 mg Na$_2$SeO$_4$/L, 16 weeks) and liver extracts were analyzed by high-resolution metabolomics and transcriptomics with xMWAS for data integration.

Result: Mice with excess Se had increased body weight and more reduced liver GSH/GSSG redox potential. Metabolomics studies showed significantly decreased levels of bile acids and increased acyl carnitines, with pathway effects linked to b-oxidation. Gene set enrichment of transcriptome data showed significant effects on cholesterol homeostasis, pancreatic b-cell signaling and fatty acid metabolism. Integrated analysis showed central hubs associated with fatty acid metabolism and glucose homeostasis.

Discussion: This global analysis of the liver metabolome and transcriptome shows that excess Se causes disruption of central fatty acid and glucose metabolism. The results emphasize that the definition of adequacy and excess Se intake may be inseparable from central redox metabolic networks controlling energy balance.
O141 - Enzymology and biological functions of glutathione peroxidases

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Introduction: Glutathione peroxidases (GPxs) constitute a diversified family of enzymes, five of which are selenoproteins in humans. They all reduce hydroperoxides by thiols via a common ping-pong mechanism.

Method: Structure and function of the selenium-containing GPxs are reviewed based on published and ongoing research.

Result: GPxs are homotetrameric (GPx1-3 and 6) or monomeric (GPx4) proteins. Discrete structural differences between the subfamilies imply peculiar specificities for both, the oxidizing and reducing substrates. The name-giving glutathione peroxidase activity dominates in GPx1, while GPx4 is least specific even accepting protein thiols and efficiently reducing hydroperoxides of membrane-integrated complex lipids. The individual GPxs also differ in localization and transcriptional control, which further enables them to adopt distinct biological roles. GPx1 is essential for detoxification of H₂O₂ and other water-soluble hydroperoxides, but also regulates insulin signaling and, like others, dampens apoptosis and inflammation. GPx2 is implicated in regulating the balance between proliferation and apoptosis of the gastrointestinal epithelium and plays a Janus-faced role in colon carcinogenesis. The extracellular GPx3 inhibits platelet-dependent thrombosis. The roles of GPx4 vary with the expression forms. The cytosolic one is of vital importance, probably by counteracting COX/LOX-driven forms of programmed cell death. The nuclear form contributes to chromatin compaction and the mitochondrial one is pivotal to male fertility. GPx6 is restricted to the olfactory bulb and is speculated to be involved in related sensory functions.

Discussion: Due to structural diversification, the GPx subfamilies adopted surprising roles far beyond the mere detoxification of hydroperoxides such as redox regulation and differentiation.

Selected references
O142 - Selenium versus Sulfur in GSH Peroxidases

Plenary session II

Keywords: GPx, molecular dynamics, MS, DFT-QM

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1
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Introduction: The family of GPx is spread in nature, encompassing enzymes in prokaryotes, animals and plants. While the majority of enzymes use S in the redox center, in vertebrates Se substitutes for S in all tetrameric GPxs and the monomeric GPx4. Almost all S-GPx, but none of Se-GPx use a C to stabilize the oxidized intermediate and then must use Trx, instead of GSH, as substrate at lower reduction potential. The emerging questions are therefore: Which is the actual advantage of substituting Se for S? Which is the mechanism of the reaction? How is the oxidized intermediate stabilized in enzymes missing the C?

Method: Steady-state kinetic analysis of enzymatic reaction of different enzymes –native or heterologously expressed- integrated with quantum-mechanics studies of the single steps of the catalytic cycle and MS analysis of intact proteins and tryptic fragments.

Result: The advantage of Se on GPx is due to: 1) use energetically cheaper GSH instead of Trx; 2) enzymatic activity and rate constant for individual steps much higher in Se-enzymes; 3) energetic barrier for the formation of the transition state leading to a charge separated intermediate lower for Se-enzymes; 4) evolution, in the presence of limiting reducing substrate, of oxidized species (S-OH and Se-OH) to SO-OH and a selenenylamide (Se-N); 5) regeneration of the reduced form of the enzyme only from Se-N; 6) formation of Dehydroalanine from Se-N by beta-cleavage only under severe denaturing conditions

Discussion: Se accounts for a faster reaction and a more stable oxidized form of the enzyme.
O143 - Nrf2 Improves Leptin and Insulin Resistance Provoked by Selenocysteine-tRNA Knockout in Hypothalamus

Plenary session II
Keywords: Nrf2, leptin, insulin resistance, Trsp, hypothalamus

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Introduction: The relationship between loss of hypothalamic function and onset of diabetes mellitus remains elusive.

Method: We generated a targeted oxidative-stress murine model utilizing conditional knockout of selenocysteine-tRNA (Trsp) using rat insulin promoter-driven-Cre (RIP-Cre).

Result: These Trsp-knockout (TrspRIPKO) mice exhibit deletion of Trsp in both hypothalamic cells and pancreatic b-cells leading to increased hypothalamic oxidative stress and severe insulin resistance. Leptin signals were suppressed and numbers of proopiomelanocortin (POMC) positive neurons in the hypothalamus were decreased, resulting in development of general leptin and insulin resistance, obesity and diabetes mellitus. In contrast, a Trsp-knockout mouse (TrspIns1KO) expressing Cre specifically in pancreatic b-cells, but not in the hypothalamus, did not display such insulin and leptin resistance, demonstrating a critical role of the hypothalamus in the onset of diabetes mellitus.

Discussion: Nrf2 regulates antioxidant gene expression. Conditional Keap1 knockout-driven increase in Nrf2 signaling suppressed the hypothalamic oxidative stress and improved insulin and leptin resistance in TrspRIPKO mice. Our results thus demonstrate that Nrf2 induction by conditional Keap1 knockout prevents oxidative damage of POMC-positive neurons and ameliorates the metabolic abnormalities.
O144 - How to Write a Great Research Paper, and Get it Accepted by a Good Journal
Elsevier Workshop
Keywords: Scientific Publishing, Paper authorship, Ethics, Journal

Anthony Newman
Senior Publisher, Life Sciences Department, Elsevier, Amsterdam, The Netherlands
(a.newman@elsevier.com)

Introduction: Knowing the best way of structuring your paper when writing it, and the most appropriate journal to send it to, really helps in getting your paper accepted. Also understanding how editors and publishers think and what they expect, and knowing how the peer review process works, is invaluable insight into the publishing process.

Method: Workshop with a publisher with discussions on key issues in writing and publishing research articles.

Result: After attending this free 2 to 2.5 hour workshop, one in the Elsevier Publishing Connect Workshop series, participants will have a clear idea of the steps needed to be taken before starting to write a paper. They will also be able to plan writing manuscripts using the logical step sequence – not the sequence in which the paper will be read. Authors are also made aware of what aspects of their papers Editors, Reviewers, and Publishers look at critically, and to ensure that in taking care of these areas, their papers are much more likely to be accepted. Dealing with referees’ comments and the art of polite rebuttal are also described such that these can be used to improve the submitted paper suitably. Sensitive areas such as publishing ethics, plagiarism, duplicate publishing, etc are also clearly explained such that participants have a clear understanding of what their responsibilities are, what is allowed, and what is not permitted.

Discussion: These insights into the publishing process will enable the participants to be more confident as an author in the world of science publishing, and so should help them get their papers published more easily.
Poster Abstracts
P1 - In vitro Generation of Superoxide by Selenofolate in MDA-MB-468 Breast Cancer Cells

1. Selenium chemistry and geochemistry
1.1 Inorganic selenium chemistry

Keywords: Selenofolate, Triple Negative Breast Cancer Cell

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Introduction: Prior studies have shown an association between generation of reactive oxygen species and some selenium compounds. Redox activity of selenium compounds is now recognized as beneficial in averting development of drug resistance, impacting the efficacy of chemotherapeutics. Several selenium compounds have been shown to be cytotoxic against cancer cells. Folate, a water-soluble vitamin that is required for DNA and RNA synthesis, and the receptors of which are upregulated in ovarian, breast, kidney and other cancer cells, was conjugated to selenium to determine if this combination produced anticancer effects.

Method: In this study, synthesized Selenofolate and the unconjugated Folate control were assayed using Lucigenin Chemiluminescence (CL) for quantifying generation of superoxide from Glutathione (GSH) oxidation. At different GSH concentrations, total chemiluminescence generated by Selenofolate was markedly increased over Folate alone which did not generate detectable superoxide. Generation of superoxide by Selenofolate was confirmed to occur intracellularly by treating triple negative breast cancer (TNBC) cells, MDA-MB-468, with Dihydroethidium (DHE), a probe that emits a red florescence indicating the presence of superoxide.

Result: Results were analyzed by fluorescence microscopy and quantitated using a fluorometric microplate reader at an excitation of 520 nm and emission of 610 nm. A difference in superoxide generation was observed with Selenofolate treatment over controls.

Discussion: These data suggest that selenium conjugation may present a strategy for the rational drug design of new anticancer pharmaceuticals.

Selected references


P2 - A preliminary study of selenium species in natural selenium-enriched garlic

1. Selenium chemistry and geochemistry
1.2 Imaging of selenium and analytical methodologies

Keywords: selenium species, Se-enriched foods, garlic

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Introduction: Selenium is one of essential trace elements for hu-man health, which plays an important role in protecting cell membrane from the oxidative damage of oxygen-free radicals as an antioxidant (Dumont et al., 2006). The appearance of Se-enriched foods become an effective way for providing selenium for humans (Hu et al., 2002). However, the toxicity and bioavailability of selenium mainly depends on its chemical form present (Rayman. 2008). Selenium species have been characterized by HPLC-ICP-MS in some studies (McSheehy et al., 2000).

Method: HPLC-ICP-MS was applied for analysis of selenium species including SeCys₂, Se(IV) MeSeCys, SeMet and Se(VI) in natural Se-enriched garlic. 0.2 g of garlic sample was put into a centrifuge tube and 5 mL ultrapure water was added. After sonication extraction at 80 °C for 4 h, the solution was centrifuged. The supernatant was filtered through a 0.45 µm filter membrane. Total selenium content was determined by ICP-MS after microwave assisted acid digestion.

Result: The content of total Se was 5.76 mg/kg in test sample. The contents of SeCys₂, MeSeCys, SeMet and Se(VI) were 0.15, 2.53, 0.24 and 0.81 mg/kg, respectively. MeSeCys was found to be the main Se species, which accounted for 43.8% of total Se. The extraction efficiency of this method was around 65%. Two unknown selenium compounds were detected and shown in Figure 1.

Discussion: This method shows good extraction efficiency in garlic samples. MeSeCys was found to be the main Se species in garlic. In order to identify the unknown selenium compounds, more coupling techniques are needed.

Selected references


P3 - Catalytic redox activity of selenium compounds generating superoxide assessed by chemiluminescence

1. Selenium chemistry and geochemistry
1.2 Imaging of selenium and analytical methodologies
Keywords: Redox Chemistry, Superoxide, Cell detection in vitro, Chemiluminescence

Soni Khandelwal
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¹Texas Tech University, Lubbock, TX

Introduction: Selenium (Se) compounds vary greatly in their equimolar systemic toxicity in both in vivo and in vitro models, with selenite usually cited as being most toxic. Universal selenium toxicity i.e., selenite (Seko, 1989); diselenides (Chaudière, 1992); isoselenocyanates (Crampsie, 2012), associated with the initial catalytic generation of the superoxide anion (O₂⁻), remains under appreciated. Here we show the varying catalytic activities of diverse selenium compounds generating superoxide in vitro. Selenite, selenodiglutathione, absorbed diselenides on polymers, diselenides in solution, and selenomethionine.

Method: The catalytic Se generation of superoxide from thiol oxidation was monitored in vitro by reduction of oxidized Cytochrome c (Lavender, 1973), Methylene Blue (Rhead, 1974), or Lucigenin (Owusu-Ansah, 2008). In vitro, superoxide is Se-generated from reduced glutathione, cysteine, and dithiothreitol. The RSe− radical was oxidized by Lucigenin; (9,9′-Bis(N-methylacridinium nitrate) producing an associated photon of light upon reduction; i.e., Chemiluminescence (CL).

Result: The CL measured is proportional to the amount of Se-generated superoxide. In culture, normal cells and selenium-treated cells intracellularly generated superoxide which was detected by the oxidation of Dihydroethidium (DHE, 2,7-Diamino-10-ethyl-9,10-dihydrophenanthridine). Upon oxidation, DHE intercalates into the DNA, resulting in a bright red fluorescence that can be photographed and/or quantitated at 580 nm (Georgiou, 2008).

Discussion: Selenite and other catalytic Se compounds generate superoxide as measured by CL or DHE. This superoxide is transposed into H₂O₂ possibly attributing to the dose dependency of selenium toxicity.

Selected references
P4 - Quantification of selenoprotein P in human serum using isotope dilution analysis

1. Selenium chemistry and geochemistry
1.2 Imaging of selenium and analytical methodologies

Keywords: Selenoprotein P, Quantification, ICP-MS, Se-peptides, Isotope Dilution

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Introduction: Measurement results which are accurate, species-specific and comparable between laboratories are a requirement within the clinical community. A novel absolute quantification method for Selenoprotein P (SEPP₁), an important bio-marker for human nutrition and disease(1-2) was developed and validated.

Method: The approach is based on the use of elemental species-specific double isotope dilution mass spectrometry (SSIDA) in combination with HPLC-ICP-MS/MS for the determination of Se in SEPP₁ at the peptide level in a complex serum matrix at clinical concentrations. Naturally and isotopically enriched Se-peptides, were synthesised and fully characterised. Two reference materials, BCR-637 and SRM1950, for which literature data and a reference value for SEPP₁ are available,(3-4) were used to assess the method accuracy.

Result: The method enabled the accurate quantification of SEPP₁ in 200 µl serum without the need of clean-up or pre-concentration. The mass fraction of Se in SEPP₁ in SRM1950 was determined as 60.6 µg kg⁻¹ with a combined relative measurement uncertainty of 5.35%. Different isotope ratio pairs and peptide calibrants provided confirmatory values. Results are in good agreement with a different SSIDA approach based on the quantification of the entire protein.(5-6)

Discussion: This is the first systematic approach for the quantification of selenoprotein (SEPP₁), which is suitable for reference measurements under relevant clinical conditions. This quantification approach of selenium associated with proteins will be invaluable for the certification of reference materials and the provision of reference values to clinical measurements and clinical trials. Future work will be the extension of the current approach to other selenoproteins relevant to health.

Selected references
P5 - Relevancy of using Se speciation for geographical authentication purpose : example of red wines

1. Selenium chemistry and geochemistry
1.2 Imaging of selenium and analytical methodologies

Keywords: speciation, red wines, geographical authentication

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Introduction: Se is an element naturally present in the soils. It can therefore be recovered in the products coming from plants which have grown on these soils. It’s the case for example of wine. Obviously, determining the Se speciation in these wines brings more speciation on its bioavailability, toxicity,… But the objective of our work was to check whether the Se speciation could be linked to the geographical origin of the wines and therefore to evaluate the relevancy of using Se speciation for wines authentication.

Method: Se speciation was performed by HPLC – ICP MS, a sensitive analytical tool able to fit with the levels of Se encountered in the red wines, after proteolytic digestion.

Result: The presentation will first introduce the development and the validation of the analytical methodology allowing Se speciation in red wines. Then the application of the method developed to the analysis of a set of red wines samples from different geographical origins will be presented. Finally the statistical treatment of these data will be set out to conclude on the relevancy of using Se speciation to allow the geographical origin discrimination of these red wines.

Discussion: Even if an incomplete discrimination can be achieved, Se speciation appears to be more discriminant than the total Se content.
P6 - Selenium Bicentennial: Two Hundred Years of Selenium Discovery

1. Selenium chemistry and geochemistry
1.2 Imaging of selenium and analytical methodologies

Keywords: Berzelius, Laboratory, History

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Mikael Bjornstedt², Julian Spallholz³
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Introduction: Jöns Jacob Berzelius is perhaps the greatest Swedish chemist, and a physician of his day. He discovered Selenium in 1817 for which he is most famous. He developed the modern 2 symbol elemental notation, i.e.; Se, used throughout the Periodic Table while also discovering the elements; Cerium, Thorium and Silicon. Authoring books on chemistry he developed the notation for water, H2O, and first introduced into literature the words, catalysis, isomer, polymer and protein. He is the founder of and became Professor of Chemistry and Pharmacology at the Karolinska Institute in 1807. Along with John Dalton, Antoine Lavoisier, and Robert Boyle he is considered one of the initial "Fathers of Modern Chemistry" and the "Father of Swedish Chemistry". Following his discovery of Selenium he was Secretary of the Academy of Sweden until his death in 1848.

Method: Photographs of the Berzelius Exhibition by Mallory Boylan with Permission for non-commercial display; Stockholm 2011.

Result: It is because of Berzelius that we are here today in Stockholm to commemorate the Bicentennial Publication on the Discovery of Selenium in 1817. This poster shows some of his self-designed laboratory equipment and chemicals used by Berzelius. It is but a small glimpse into his genius, laboratory and scientific life.

Discussion: Born: August 20, 1779, Linköping, Sweden
Died: August 7, 1848, Stockholm, Sweden
Discovered: Silicon, Selenium, Cerium and Thorium
Scientific Papers Over 250: The Use of the Blowpipe in Chemical Analysis, and in the Examination of Minerals
Education: Uppsala University, Katedralskolan, Linköping
Awards: Copley Medal

Selected references
https://en.wikipedia.org/wiki/J%C3%B6ns_Jacob_Berzelius)
P7 - Simultaneous determination of selenium and sulfur species in biological samples

1. Selenium chemistry and geochemistry
1.2 Imaging of selenium and analytical methodologies

Keywords: sulfur, HPLC, mass spectrometry, biological samples

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Introduction: The micronutrient selenium and the macronutrient sulfur are essential for life and show strong similarity in respect to their chemical properties and naturally occurring compounds. To further elucidate similarities and differences of selenium and sulfur, simultaneous determination of their species in biological samples is of interest. While HPLC/ICPMS is the most common technique for selenium speciation analysis in biological samples, polyatomic interferences on the main sulfur isotope hamper the determination of sulfur species using this technique.

Method: Simultaneous determination of selenium and sulfur species in biological samples was performed by HPLC coupled to ICP-triple quadrupole-MS (ICP-QQQ-MS) with oxygen as a reaction gas.

Result: We present a method for the simultaneous determination of selenium and sulfur species in biological samples by HPLC/ICP-QQQ-MS. The oxygen reaction mode is used for the detection of selenium and sulfur as SeO$^+$ and SO$^+$, respectively, thereby overcoming interferences typically hindering the ICPMS detection of these two elements. When employing reversed-phase HPLC as the separation step before ICP-QQQ-MS, limits of detection in the low to sub $\mu$g per liter range were obtained, for example, for selenoneine and its sulfur analogue ergothioneine, which have both been ascribed antioxidative properties and are, hence expected to be of considerable health relevance.

Discussion: HPLC/ICP-QQQ-MS offers the possibility of the simultaneous detection of selenium and sulfur species in biological samples. This approach is a promising tool for further investigation of similarities and differences of these two elements.
P8 - The fractions and distribution of soil selenium in Heilongjiang province and its' impacting factors

1. Selenium chemistry and geochemistry
1.3 Local geological selenium sources and global cycling

Keywords: Soil selenium; Fractions; Physical and chemical properties; Heilongjiang province

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Introduction: The contents, fractions, distribution of soil selenium (Se) and its' influencing factors in different soil types in Heilongjiang Province (China) were surveyed and analyzed.

Method: It was investigated 19 soil samples were selected by above five natural geographical areas were collected by 0-20cm, including black soil, albic soil, chernozem, dark brown soil, volcanic ash soil, meadow soil, aeolian sandy soil and saline-alkali soil for the Se fractions analysis and the measurement of physical and chemical properties.

Result: The topography of terrain in Heilongjiang Province is vast and complex, and the geochemical conditions are obviously different, which leads to a great change in the total Se content in this study, ranging from 0.0689 to 0.4628 mg kg⁻¹, was generally low. Residual Se and organic Se were the main Se fractions in low Se soils, followed by acid soluble Se. The water soluble Se and exchangeable Se contents which plant could uptake was very low, which may be main reason resulting in low Se bioavailability content in soils in Heilongjiang. There was a significant correlation among soil acidic Se, organic Se and residual Se and total Se content in soil, but the correlation between water soluble and exchangeable selenium was not significant.

Discussion: The personal correlation analysis showed that the contents of metal oxides, clay and sand were the main factors affecting soil Se capacity, which restricted the distribution of total Se in soil. pH, soil organic carbon (SOC) and other factors could not be ignored in impacting the content of Se.
P9 - Zoning pollution-free and selenium-rich land resources with geochemistry

1. Selenium chemistry and geochemistry
1.3 Local geological selenium sources and global cycling

Keywords: cadmium, safe utilization

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Introduction: It is difficult to maintain a safe level of selenium (Se) intake in adults, owing mainly to the relatively narrow gap between Se deficiency and excessive Se dose. Thus it’s meaningful for zoning safe and Se-rich soils.

Method: A typical Se-rich area covering 50 000 ha in Enshi (located in Hubei, China) was investigated. Hundreds of samples as rhizosphere soils, corns, irrigation water, soil profiles, rocks, fertilizers and atmospheric depositions were collected. All of the samples were well prepared and analyzed with recommended methods.

Result: It was inferred that the status of Se in crops showed the strong dependence on the total Se in soils that were similar to the status of Cd. The average total Se and Cd concentrations in corn topsoil were 1.88 and 2.12 µg•g⁻¹, respectively. Higher Se and Cd concentrations in soil were distributed with the Permian or Triassic strata which were stretched with the black shale. The geochemical model of the Se cycle model implied that parent rocks were the main source of soil Se contents.

Discussion: The average total Se and Cd concentrations in corn were 0.39 and 0.24 µg•g⁻¹, respectively. In the study area, the concentrations of Se and Cd in corns were highly correlated with Se, Cd, Hg, Cu, Zn, K and Ca in soils. Considering the safe intake level of Se and Cd by corns, the models of Se and Cd concentrations in crop-soil system were built by multiple regression analysis. With these, the areas where were favorable to utilizing safe Se-rich crops were zoned.

Selected references
P10 - Selenium, Nitrogen and Carbon remobilization study following litter decomposition during 10 months

1. Selenium chemistry and geochemistry
1.4 Relationships of selenium between soils, water, and vegetation

Keywords: isotope labelled biomass, litter, decomposition, speciation

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Introduction: In the soils, effective sink for Se, a significant part of Se is associated with the organic pool (Tolu et al., 2014) but some organo-Se compounds still remains unidentified (Di Tullo et al., 2015). Although the biogeochemical Se cycle is well documented and studied, the mobilization processes in this organic Se pool are still poorly understood as well as the biotic and abiotic factors controlling this mobilization and the temporal scale involved.

Method: We propose an original method that consists 1) in labelling a non-accumulator ryegrass with 4 isotopes (\(^{78}\)Se\(^{IV}\), \(^{82}\)Se\(^{VI}\), \(^{15}\)N and \(^{13}\)C) to study uptake, assimilation and speciation of selenite and selenate species in this plant and 2) in examining during 10 months the decomposition of this isotopically labelled biomass added to a grassland soil either as shoot litter or as root litter. This original method allows us to monitor selenium, carbon and nitrogen in the same fractions extracted from the first centimeters of soils, the residual litter and the new vegetation.

Result: Our work conclude that the rye-grass could develop the same strategies as hyperaccumulator plants to limit selenium toxicity: selenate transfer to the shoot compartment and production of methylated Se species considered as precursor compounds for Se volatilization (Versini et al., 2016).

Discussion: The second experimentation showed that the patterns of C and N releases differ from that of Se; the distributions of the tracers are low in the underlying horizons and the amounts of Se-MO pools depend on the litter sources but are almost constant during 10 months.

Selected references


P11 - The Precipitation Impact on the Selenium Speciation in Surface Soil

1. Selenium chemistry and geochemistry
1.4 Relationships of selenium between soils, water, and vegetation

Keywords: speciation, precipitation, soil, infiltration

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Introduction: The bioavailability of selenium(Se) to plants is related to the speciation of Se in soil. The sources and budget of selenate and selenite in soil are the control factors for the evaluation of Se deficiency.

Method: Infiltration waters were collected under 20cm of surface soil with a self-designed devise in field. Rain waters were also collected nearby within the same time-span. The water samples were filtered and determined for Se species and the budget of the element in surface soil were calculated

Result: The average total Se in precipitation ranged from 124-588 ng/L in Se-rich and deficiency areas sampled(Table 1). Selenate, as high as 6.8times that of the selenite was the main spices in rain water. The Se concentration of infiltration water was from 247ng/L to 1.43µg/L, all higher than that of Se in precipitation. This indicated that the process of rainfall to surface soil may lead to a net loss of Se. There is no big difference on the ratio of selenate, selenite and humic-bond Se in surface soil, but in three severe Kaschin-Beck’s Disease areas the selenate concentration has a narrow range of 242-249µg/kg. The calculated Se influx from precipitation to surface soil is at most 9.5% of the total annually, by species it could be as much as 65.3% of selenate in Yongshou.

Discussion: Precipitation is one of the important sources of Se in surface soil, and leaching of humic-bond Se is the main cause of Se loss. Se deficiency area in China has a typical selenate concentration of 250µg/kg.

Selected references
P12 - Effect of pH, iron plaque and phosphorus on selenium uptake by rice seedling

1. Selenium chemistry and geochemistry
1.4 Relationships of selenium between soils, water, and vegetation

Keywords: pH levels, iron, phosphorus, root, shoot

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Introduction: Plants are the dominant source of Se for animals and humans. Rice is the staple food for people in many parts of the world. Rice plays a crucial role in the food supply in China, and it is also an efficient way to provide Se to consumers. Thus, Se uptake by rice plants is a significant component of the food chain for satisfying human nutrition requirements. The aim of the current study was to explore the effect of iron plaque and phosphorus on selenium uptake by rice seedling in response to different pH levels.

Method: A hydroponic experiment was carried out.

Result: The results showed that the content of selenium in shoots and roots of rice seedling was significantly affected by pH level. The highest content of selenium in the shoots and roots were at pH 5 and pH 3, respectively. The selenium uptake by rice seedling shoots was inhibited by iron plaque, but this effect on roots was not significantly. Phosphorus significantly promoted the selenium uptake by rice seedling shoots. When pH changed from 3 to 8, the distribution rate of selenium was increased at first and then decreased in the shoots, while the opposite tendency in the roots.

Discussion: Rice absorption and transport of selenium affected by pH, iron plaque and phosphorus composite. In the rice cultivation, should be paid attention to the comprehensive adjustment of soil pH, phosphorus and iron fertilization conditions, to increase the selenium accumulation in rice.

Selected references


P13 - Analysis of Se (IV) and Se(VI) absorption kinetics of different genotypes of Nicotiana tabacum L.

1. Selenium chemistry and geochemistry
1.4 Relationships of selenium between soils, water, and vegetation

Keywords: absorption kinetics; Se (IV); Se(VI); Nicotiana tabacum L.

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Introduction: Selenium (Se) is an essential element for human beings and animals. Through methods of soil Se-supplement and foliar spraying could improve the plant Se content obviously, and the applying forms of Se inorganic were more inorganic Se. However, the absorption mechanism of selenite and selenate by tobacco was unclear.

Method: A hydroponic experiment was conducted. N. tabacum K326 and N. tabacum yunyan87 were used, Se were added in the form of Na$_2$SeO$_4$ and Na$_2$SeO$_3$, respectively. The ion depletion method was adopted. The content of Se was determined with a hydride generation atomic fluorescence spectrometer.

Result: Compare to Se (IV), the Se (VI) affinity of roots was stronger. Under the condition of different Se source, Se accumulation of cultivar yunyan87 was significantly higher than those of cultivar K326 (Figure 1). Under the condition of different pH, Se (IV) and Se (VI) affinity of flue-cured tobacco were strongest at pH 6 (Figure 2). The addition of PO$_4^{3-}$ suppressed the Se (IV) uptake of flue-cured tobacco more than the Se (VI) absorption; while the addition of SO$_4^{2-}$ had a stronger inhibition of the Se (VI) absorption (Figure 3). The application of uncoupling metabolism inhibitors (CCCP) and calcium ion channel inhibitors (LaCl$_3$) could evenly inhibit the Se (IV) and Se (VI) absorption of flue-cured tobacco, and the effect of CCCP was stronger (Figure 4).

Discussion: Selenate was absorbed by the plant roots through the high-affinity phosphate transporters, while selenite was assimilated by high-affinity sulfate transporters (Zhu et al. 2009). Results fully confirmed that there existed antagonism effects between SeO$_4^{2-}$ and SO$_4^{2-}$ as well as Se$_2$O$_3^{2-}$ and PO$_4^{3-}$.

Selected references
P14 - Agronomic biofortification of carrots with selenium in oxidic soil

1. Selenium chemistry and geochemistry
1.6 Strategies to improve selenium accumulation and biofortification

Keywords: tuber, selenate, Oxisol, greenhouse, nutrition

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Introduction: Selenium is an important element linked to physiological processes in plants, microorganisms, animals, and humans (Rayman, 2012). The addition of selenium to crops via soil application of Se i.e. agronomic biofortification is recommended to overcome dietary deficiencies (Chilimba et al., 2012; Bañuelos et al., 2015; Winkel et al., 2015). This work tested agronomic biofortification strategies to increase Se to nutritionally ideal levels inside the edible parts of carrots growing in oxidic soils.

Method: The experiment was conducted in a greenhouse in Lavras - MG, Brazil. The plots comprised one carrot plant (Daucus carota L.) per pot, filled with 1.6 kg of a clayey-textured Oxisol. The experimental design was completely randomized using seven doses of Se as Na₂SeO₄: 0.0; 0.1; 0.2; 0.4; 0.8; 1.6 and 3.2 mg kg⁻¹. Total selenium content in carrots was determined after acid digestion with HNO₃ + HCLO₄ (ratio 2:1, 50-200°C over 2 h) followed by analysis with GFAAS.

Result: Plants were highly responsive to Se application as selenate in the soil up to the highest dose (3.2 mg kg⁻¹), which resulted in a total Se accumulation of 76.45 mg kg⁻¹ in tubers and 178.89 mg kg⁻¹ by dry weight (DW) in leaves.

Discussion: Selenium content was higher in the leaves compared to the storage roots for all doses. The results showed that Se-enriched carrots can be obtained via agronomic biofortification. Consumption of 100 g f.w. of carrots fertilized with 0.4 mg kg⁻¹ Na₂SeO₄ can supply 100% (80.37 µg) of the Recommended Daily Allowance (RDA) for Se.

Selected references


P15 - Effects of different selenium application on selenium accumulation in Lentinula edode

1. Selenium chemistry and geochemistry
1.6 Strategies to improve selenium accumulation and biofortification

Keywords: Lentinula edodes; selenium biofortification

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Introduction: Many edible fungi have been demonstrated to absorb inorganic Se from the substrate and convert it into more efficient forms (Maseko et al., 2013), but little study is conducted on the effect of exogenous Se species on the growth and enrichment of Se in Lentinula edodes. Therefore the difference of selenite, selenate and Se-enriched yeast on Se accumulation was studied to explore the absorption of Se in Lentinula edodes.

Method: Na$_2$SeO$_3$, Na$_2$SeO$_4$ and Se-enriched yeast (Se content: 510 mg/kg) were injected into base material for mushroom one week before fruiting. The mature fruiting bodies were harvested at 7, 14, 21, 28, 49 and 70 days after injection, respectively. All samples were collected, oven-dried at 55˚C, weighed and ground. After digestion with HNO$_3$-HClO$_4$ (4:1, v/v) and reduction in 6 mol/L HCl, Se concentrations were determined by HG-AFS.

Result: All treatments didn’t cause visible impact for the growth of L.edodes. The concentration in L.edodes fruit bodies, which was applied with Se, considerably increased and it was proportional to the amount of Se added to the substrate (Fig. 1). Se contents of mushrooms increased firstly and then decreased with the increase of harvest times. The highest content was occurred in 21 (selenite) and 49 (selenate and Se-yeast) days after injection, respectively. When the same concentration was added into the substrate, Se contents of mushrooms were in the order of Na$_2$SeO$_3$ > Na$_2$SeO$_4$ > Se-yeast.

Discussion: L.edodes is a good Se accumulator. Among investigated Se species sodium selenite is the best sources in Se biofortification of L.edodes.

Selected references

Introduction: Apples are the most important fruit crop grown in Germany. As with most food of plant origin, the selenium (Se) content of apples is relatively low and thus the fruits contribute little to human Se intake. The objective of this study was to investigate whether apples are suitable for Se biofortification programs, both from an agronomical and a consumer point of view.

Method: Apple trees were sprayed with different doses of sodium selenite and sodium selenate, respectively. The Se content in fruits was analyzed by graphite furnace AAS. The consumer survey, including 384 respondents, was performed online and based on a questionnaire with 32 items.

Result: The targeted selenium content in apple fruits (10 – 20 µg Se/100 g fm) was achieved by a single foliar spray of 100 – 150 g Se/ha both with selenite and selenate. At this enrichment level of Se, the use of nutritional and health claims is approved according to regulations (EC) No 1924/2006 and (EU) No 432/2012, respectively. About 52% of all consumers questioned expect selenium in food to have a positive impact on human health and 56% would appreciate apples offered with the claim “rich in selenium”. Among the health claims, the statement “Se contributes to the normal functioning of the immune system” was most attractive for those questioned (76%).

Discussion: With regard to the promising results of our preliminary spray trials, and the predominantly positive consumer perception of Se enriched fruit produce, it seems to be worthwhile to develop an appropriate Se biofortification technique for apples.
P17 - Optimising fertiliser formulations for selenium biofortification of wheat grain

1. Selenium chemistry and geochemistry
1.6 Strategies to improve selenium accumulation and biofortification
Keywords: biofortification, macronutrient carrier, application method, wheat

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**Introduction:** Currently, about 15% of the global population is at risk of health problems due to inadequate consumption of the essential micronutrient selenium (Se). Although biofortification of staple crops through soil or foliar application of Se has proven to be very successful, few studies have investigated crop response to the application of different Se-enriched macronutrient fertiliser matrices, which could reduce labour and application costs.

**Method:** Hence, pot trials were set up to determine crop uptake of Se by different macronutrient carriers and methods of application in 3 soils with contrasting properties. Four macronutrient fertilisers – urea, di-ammonium phosphate, muriate of potash and sulfate of ammonia – were enriched with Se by mixing the macronutrient powder with sodium selenate solution and pressing the mixture into 4 cm pellets, prior to cutting them into tablets. The granules were then soil applied just before planting the seedlings. For the foliar fertilisation, Se was applied to leaves with a micropipette, as sodium selenate with either urea or urea ammonium nitrate at ear emergence. In both experiments, Se was supplied at a constant rate of 3.33 \(\mu\)g kg\(^{-1}\) soil (a rate equivalent to 10 g ha\(^{-1}\)). Wheat was grown under controlled conditions until maturity. After 12 weeks, grains were harvested and analysed for Se concentrations by ICPMS.

**Result:** Uptake data, which show differences in Se accumulation due to macronutrient carrier and/or application method, will be presented at the conference.

**Discussion:** To our current knowledge, this is the first study undertaken to determine the optimal fertiliser matrix to supplement wheat with Se.
P18 - Effects of foliar spraying selenite or selenate at different stage on the selenium uptake and distri

1. Selenium chemistry and geochemistry
1.6 Strategies to improve selenium accumulation and biofortification

Keywords: Foliar spray, Selenite, Selenate, Organic selenium, Protein selenium

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Introduction: Organic Se, which is mainly originated from plant and animal sources, showing more bioavailability than inorganic Se. However, the organic Se in the crop is primarily bound to the protein (Wang et al., 2013).

Method: A field experiment which has three replicates was conducted at the Shekou village of Qianjiang city, Hubei Province, China (30°33′17″ N, 112°53′23″ E). In this experiment, five treatments consisted spraying sodium selenite (Na2SeO3) and sodium selenate (Na2SeO4) at the tillering stage (T) and heading (H) stage respectively and one control.

Result: Foliar of selenite and selenate at heading stage, selenium concentration in brown rice was 2.6 times and 3.0 times than that at the tillering stage; Whichever period of spraying. The brown rice sprayed by selenate in which Se concentration was about 2 times of that sprayed by selenite (Figure 1). The results of this study also showed that the organic Se accounting for 79.49%-88.70% of total Se. The concentration of protein Se in brown rice accounting for 43.25%-51.16% of the total Se (Table 1).

Discussion: Spraying of selenite or selenate at heading stage, the enrichment ability of Se in rice was 2-3 times of that at the tillering stage. The organic Se in brown rice could reach 79.5%-88.7% and the protein Se could reach 43.2%-51.2% in all treatments. Overall, spraying selenate at heading stage is more favorable for the enrichment of total Se, organic Se and protein Se in grains.

Selected references
P19 - Se and its antagonists Hg, As, Cd in hair of Taiwanese and Russian residents

1. Selenium chemistry and geochemistry
1.7 Selenium interactions with other elements in the environment

Keywords: Key words: Hg, As, Cd, Russia, Taiwan, hair

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Introduction: An antagonism between Se and Hg, As, Cd is well-known, and Se administration has the protective effects against intoxications by these trace elements. Hair levels of Se, Hg, As, Cd were studied in residents of territories differ in climate and life patterns.

Method: Totally 178 Taiwanese and 41389 Russians (39162 Moscow, 1972 Novosibirsk, 255 Sakhalin) 20-50 y/o males and females were investigated. Hair samples were analyzed by ICP-MS.

Result: For Se in men there was no significant difference for Russian vs Taiwanese (median 0.44 vs 0.50 µg/g, p > 0.1), while Russian women had a difference vs Taiwanese (0.33 vs 0.43 µg/g, p < 0.05). Hair As for women and men in Moscow (0.021, 0.046) was significantly lower vs Taiwan (0.037, 0.065). Hair Hg for women and men was lower in Moscow (0.55, 0.60), Novosibirsk (0.40, 0.43) and Sakhalin for men only (1.16) vs Taiwan (0.84, 1.71). Hair Cd for women and men was higher in all Russian cities (0.0105, 0.0140) vs Taiwan (0.0065, 0.0057).

Discussion: Generally, Sakhalin residents were similar to Taiwanese in hair As (males and females), Hg (females), Se (males) due to geographical and nutritional similarity while continental Russians had higher Cd and lower Se. There were no distinct correlations between Se and other elements in hair, while most subjects had their concentrations within normal ranges.
P20 - Selenium and mercury interactions in the apex predators from the Gulf of Trieste (northern Adriatic)

1. Selenium chemistry and geochemistry
1.7 Selenium interactions with other elements in the environment
Keywords: Mercury, Fish, Rays, Northern Adriatic

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Introduction: It was suggested that environmental Se moderates the bioaccumulation and toxicity of Hg in marine organisms.

Method: Interactions between Se and Hg were studied in waters, plankton, and benthic (Pteromylaeus bovinus, Myliobatis aquila) and pelagic (Dasyiatis violacea, Dasyiatis pastinaca) rays, as apex predators, in the Gulf of Trieste (northern Adriatic Sea), one of the most severely Hg polluted area in the Mediterranean and worldwide due to the input of Hg from the former mine in Idrija (NW Slovenia) by the Soča/Isonzo River.

Result: Seawater contains 0.32-2.13 µg/L dissolved Se. Dissolved Hg ranges between 0.18-4.9 ng/L with higher concentrations in area near the river discharge. In net-zooplankton (>200 µm), Se ranged between 2.2-3.8 µg/g (d.w.) and Hg between 130-370 ng/g. Se in ray species varied between 1.0-4.41 µg/g (d.w.) in male and female muscle. Pelagic species contained higher Se in muscle. Slightly lower levels were in liver ranging between 0.87-2.54 µg/g (d.w.) in both genders. The highest Hg (mostly as MeHg) levels were in muscle of D. violacea (1.17-4.40 µg/g d.w.), followed by P. bovinus (0.13-3.05 µg/g d.w.), and D. pastinaca (0.07-0.47 µg/g d.w.).

Discussion: In benthic ray species, a parallel increase of Se and Hg in muscle was found, so greater Hg bioaccumulation resulted in Se coaccumulation. Hg/Se ratios (molar) in muscle of pelagic nad benthic species were ≤1 while in zooplankton and liver were <0.01 and <0.7, respectively. Low Hg contents in muscle and liver of rays corresponded to low Hg/Se ratios increasing in muscle to 1 at about 6 µg/g (d.w.).
P21 - Environmental Selenium Influences Fish Methylmercury Bioaccumulation and Risks

1. Selenium chemistry and geochemistry
1.7 Selenium interactions with other elements in the environment
Keywords: Mercury, Freshwater Fish

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Introduction: Environmental mercury can either bioaccumulate within an aquatic food chain as the readily absorbed and highly toxic form methylmercury (MeHg) or biologically retire as insoluble and inert forms such as mercury sulfide (HgS) and mercury selenide (HgSe).

Method: Selenium analyses were performed on freshwater fish collected from across North America states. Fish included piscivores and non-piscivores.

Result: The HBV of all fish was inversely associated with size and age. While the HBV of more than 95% of piscivores was positive, the incidence of large predatory species with negative HBVs was highly correlated with their occurrence in fresh water bodies with poor environmental Se availability. While Hg accumulation of non-piscivores was also inversely associated with environmental Se, none were found to have negative HBVs.

Discussion: MeHg bioaccumulation was found to be inversely related to environmental Se availability. Since MeHg toxicity is proportional to both its absolute as well as relative amounts (in relation to Se), consumption of such fish is likely to be associated with far greater risks than is currently recognized.
P22 - Study of geochemical characteristics and influencing factors of soil selenium in the typical soil

1. Selenium chemistry and geochemistry
1.7 Selenium interactions with other elements in the environment

Keywords: profiles, influencing factors

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Introduction: Selenium was discovered by a Swedish chemist, in 1817. After two hundred years, people are increasingly concerned about this marvelous element. Many Se-rich soils have been investigated around China, but the mechanism of the migration and transformation of selenium in soil is still hot issues in the field of geochemistry.

Method: The typical soil profiles of carbonate rock and terrigenous clastic rock in Wuming County of Guangxi were chosen as the investigative objects. Chemical elements, pH and total organic carbon (TOC) were analyzed by recommended methods.

Result: The results showed that the soil selenium content was inherited from the parent materials, with average values of 0.55 mg/kg and 1.43 mg/kg respectively; and the selenium content of the carbonate rock was enriched significantly in the surface soil, but relatively stable in the terrigenous clastic rock. The total water-soluble selenium content of the carbonate rock was higher than that in the terrigenous clastic rock, with arithmetic average values of 3.48 µg/kg and 1.81 µg/kg respectively, and they were both given priority to with selenate, followed by selenite and humic acid combined with selenium.

Discussion: Different factors which may control the contents of selenium were considered. It was inferred that the contents and chemical speciations of selenium in soil profiles originated from the carbonate rock were mainly dominated by the total organic carbon (TOC) contents and pH values, while of which originated from the terrigenous clastic rock were controlled by the pH values, the contents of TOC, aluminum oxide, iron oxide and soil texture.

Selected references
P23 - Interaction of selenium and cadmium in soil-corn system in natural selenium and cadmium rich area

1. Selenium chemistry and geochemistry
1.7 Selenium interactions with other elements in the environment
Keywords: cadmium, corn, bioaccumulation factors, interaction

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Introduction: A large number of studies have confirmed that selenium-cadmium (Se-Cd) interaction is a widespread phenomenon. However, little attention has been paid to plants especially in natural Se-Cd rich area. To investigate the influence of Se on the uptake and translocation of Cd in the soil-corn system, the levels of Se and Cd in different parts of corn plants and corresponding soils of root zones collected from a Se-Cd rich area were determined.

Method: Se and Cd speciations in soil were treated by sequential extraction protocol. All treated samples were acid-digested and analyzed for total Se by hydride generation atomic fluorescence spectrometry (HG-AFS), and Cd by atomic absorption spectroscopy (AAS)

Result: Molar ratios of bioavailable forms of Se and Cd (Se:Cd) in the soil and the bioaccumulation factors (BAFs) of Cd were used to elucidate interaction effects of Se-Cd between soil and corn. A reduction in translocation of Cd in the aerial parts with increasing Se levels in the soils was observed. However, the levels of bioavailable forms of Se in the soils were positively correlated with the BAFs of Cd in the roots when molar ratios of bioavailable forms of Se and Cd (Se:Cd) in the soil over ten.

Discussion: The pronounced role of Se in protecting corn plants from Cd bioaccumulation and its potential use for reducing Cd concentration in different tissues of corn plants, and the root may act as an effective buffer or barrier for the suppression of Cd absorption or translocation to the aerial tissues.

Selected references
P24 - Impact of Initial Se Status and Gpx1 Genotype on Selenoenzyme and Transcript Expression in Mice

2. Selenium in the molecular life sciences
2.1 Metabolism of selenium in living cells
Keywords: glutathione peroxidase, metabolism, mice, transcript

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Introduction: Key components of Se metabolism in intact animals include relative rates of Se incorporation into selenoproteins, underlying regulation of selenoprotein transcripts, and Se status itself. At steady state, just 0.1 and 0.05 µg Se/g diet are required to raise liver glutathione peroxidase-1 (Gpx1) and plasma Gpx3 activities, respectively, to plateau levels in wildtype mice. Knockout of Gpx1 both reduces body Se stores but also blocks flux of Se into Gpx1.

Method: Here, we studied the short-term availability (flux) of Se in male Gpx1 wildtype (Gpx1+/+), heterozygote (Gpx1+/-t), and knockout (Gpx1-/-) mice, that had been fed a basal Se-deficient diet for 17 wk after weaning, and then were repleted with 0, 0.05, 0.1, 0.2, and 0.3 µg Se/g diet as Na2SeO3 for only 7 days.

Result: Se response curves for liver Gpx1 activity were sigmoidal, reaching the plateau at 0.15 µg Se/g for Gpx1+/+ mice but at 0.20 µg Se/g diet for Gpx1+/- mice, and with no effect on liver Gpx1 activity in Gpx1-/- mice (as expected). In contrast, plasma Gpx3 activity increased only in mice fed 0.3 g Se/g diet for all genotypes, and there was little effect on RBC Gpx1 activity regardless of genotype or Se supplementation.

Discussion: These results clearly show that the flux of Se metabolism is altered, at least in the short-term, when initially Se-deficient mice are Se repleted, and this is further modified by Gpx1 genotype. Ongoing analysis of selenoprotein transcripts indicates that differential and/or delayed upregulation of Gpx1 transcripts underlies the observed shifts in selenoenzyme activity.
P25 - Insight into the microbial respiration of selenium. A metagenomics study

2. Selenium in the molecular life sciences
2.1 Metabolism of selenium in living cells

Keywords: Biogenic Selenium, Metagenomics, Microbial Selenium Respiration, Nanoparticles, 16S rRNA Sequencing

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Introduction: Selenium oxyanions are effective electron acceptors, and essential trace elements, in microbial respiration [1]. Selenium cycling is highly dependent on microbial communities. A deeper understanding of microbially-mediated selenium cycling is required to exploit selenium microbiology and develop new environmental biotechnologies.

This study investigated the phylogeny of microbial populations underpinning selenium reduction in different environments, including soils and waste-conversion biofilms, to expand our understanding of the microbial ecology of selenium cycling.

Method: Batch incubations of loam soil; Atlantic peatland soil; and both intact and physically-homogenised, anaerobic sludge granules were used to enrich for selenite- and selenate-reducers by serially sub-culturing enrichments to fresh media. Reductants were lactate:acetate (16mM:4mM) or H₂:CO₂ (80:20v/v). Population dynamics and community succession were monitored by sequencing 16S rRNA genes in temporal sub-cultures, and to identify key selenium-reducing taxa. Metagenomics were used to comprehensively characterise highly enriched selenium-reducing cultures. 16S rRNA and functional mRNA targets were probed in cross-sections (10um-thick) of intact sludge granules (diameter 0.5-2mm) using fluorescence in situ hybridisations (FISH).

Result: The formation of elemental selenium nanoparticles, including on the surface of spherical sludge granules, indicated selenate and selenite reduction, even in autotrophic enrichments fed only with H₂ and CO₂. Sequencing indicated distinct taxa associated with selenium metabolism in each of the soil and sludge enrichments. FISH revealed highly organised distributions of selenium cyclers across the architecture of sludge granule biofilms.

Discussion: The results indicate widely distributed potential for selenium-reduction across an expansive phylogeny in previously unexposed samples.

Selected references

P26 - Selenium subcellular distribution, speciation and antioxidant response in rice booting stages

2. Selenium in the molecular life sciences
2.1 Metabolism of selenium in living cells
Keywords: Selenium speciation, photosynthesis, antioxidase, subcellular distribution, panicle initiation stage

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Introduction: Selenium is an essential element of animal and human being, and Se deficiency may cause disease like cardiac disease. Production of Se-enriched rice is one of the most important way to supply Se in human body, and thus understanding of the mechanisms of Se-enriched rice is of great importance.

Method: A rice soil pot experiment was employed to study the effects of selenium addition on the growth, photosynthesis, anti-oxidation, Se uptake and distribution, and selenium speciation in three different stages of panicle initiation stage and maturity stage.

Result: The results showed that soil selenium application increased Se uptake in rice. Low rates of selenium (<5mg/kg) application was good for the improvement of rice photosynthesis, growth and yield. Assay of Se speciation showed that SeCys and SeMet were the two main forms in rice, of which SeMet accounted for 65.5%-100% in ear and leaves, but SeCys accounted for 61.4%-75.6% in rice grain. SeMet were also the main forms in different subcellular parts at panicle initiation stage. However, when soil Se application was more than 5 mg/kg, inorganic Se was existed in rice grain, mainly Se(VI). And in the meantime, activity of SOD, POD, and CAT decreased and the content of MDA increased.

Discussion: The results indicated that Se effects depended on the Se levels of application, and the optimum amount of Se application for rice would be less than 5 mg kg⁻¹.

Selected references
P27 - Explore the subcellular distribution and speciation of Se in pakchoi (Brassica chinensis L.)

2. Selenium in the molecular life sciences
2.1 Metabolism of selenium in living cells
Keywords: subcellular distribution, speciation, pakchoi

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Introduction: Deposition on cell wall and vacuolar compartmentation play important roles in heavy metal(loid) detoxification, tolerance, and hyperaccumulation in plants (Zhou et al. 2016), but subcellular fractionation of Se has been seldom reported so far. Thus, the distribution subcellular of pakchoi and speciation of Se was determined to better understand Se metabolism in plants.

Method: Pot experiment was carried out with selenate and selenite applied in soil. After two months’ growth, the pakchoi fresh leaves were fractionated into cell wall (F1), organelle (F2) and soluble fraction (F3), using the differential centrifugation method modified from Chen et al. 2014. Se speciation was analyzed by LC-UV-HG-AFS after enzymatic extraction by Proteinase E for both intact plant tissues and subcellular fractions.

Result: Se subcellular distribution of pakchoi was in sequence as: soluble fraction > organelles > cell wall. Over 50% of the plant-accumulated Se was found in F3, and the speciation determination (Fig. 1) showed that selenate was the dominated species. Selenomethionine, which was the major organic Se species after enzymolysis and accounted for 4.4 - 29.2 % of total Se, was located in F2. More information can be found by comparing different treatments and plant tissues.

Discussion: Massive selenate found in F3 indicates that pakchoi can compartment excessive Se in vacuole to avoid its toxicity. The selenomethionine detected in F2 echoes its derivation from selenoproteins in organelles. F1 contained the least amount of Se, suggesting that cell wall plays little role in Se metabolism. The different speciation in these fractions confirmed their different roles in Se metabolism.

Selected references
P28 - Comparison in bioavailability of nine bioselenocompounds

2. Selenium in the molecular life sciences
2.1 Metabolism of selenium in living cells

Keywords: bioavailability, Caco-2, ICP-MS, speciation, selenosugar

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Introduction: Biological effects of selenium (Se) are dependent on its chemical form. In this study, we evaluated the bioavailability of nine naturally occurring Se compounds, or the so-called bioselenocompounds, in vivo and in vitro.

Method: The bioselenocompounds such as selenite, selenate and selenocyanate (SeCN⁻), L-selenomethionin (SeMet), Se-methylseleno-L-cysteine (MeSeCys), L-selenohomolanthionine (SeHLan), selenocystine (SeCys₂), Se-methylseleno-N-acetylgalactosamine (SeSug₁) and trimethylselenonium ion (TMSe⁺) were used. Caco-2 cells were used for in vitro membrane permeability study, and in vivo bioavailability was evaluated by the recovery of serum selenoproteins in Se-deficiency rats.

Result: In in vitro experiments, SeMet and MeSeCys were more efficiently permeated through Caco-2 monolayer than other bioselenocompounds. Although SeHLan and SeCys₂ was other selenoamino acids, SeHLan and SeCys₂ were less efficiently transported than SeMet and MeSeCys. Contrary to the results of in vitro study, Quantitative methods indicated no significant differences in bioavailability among the bioselenocompounds except TMSe⁺ in vivo.(1) These results indicate that animals can equally assimilate both inorganic and organic bioselenocompounds except TMSe⁺, which is one of the urinary Se metabolites.

Discussion: The discrepancy between in vitro and in vivo results is interesting and important for the consideration of the nutritional availability of bioselenocompounds. We offer two explanations for the discrepancy. First, although monomeric selenoamino acids such as SeMet and MeSeCys are more rapidly absorbed in the gastrointestinal tract than other Se species, methylated Se in the selenoamino acids is less efficiently used for the synthesis of selenoproteins than non-methylated Se species. Second, the bioselenocompounds are metabolized/decomposed before absorption in the gastrointestinal tract.

Selected references
P29 - Characterization of rhodanese-like protein from Geobacter sulfurreducens

2. Selenium in the molecular life sciences
2.1 Metabolism of selenium in living cells
Keywords: rhodanese, selenoprotein, selenite, Geobacter

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Introduction: Geobacter sulfurreducens is a metal-reducing anaerobic bacterium. The novel selenoprotein MHSEP involved in selenite/tellurite reduction was found in our previous research. The mhsep gene forms an operon-like gene cluster (gsu2940-2930) on the genome. The gsu2940 gene encodes a protein with a sequence similarity to a sulfurtransferase rhodanese, which catalyzes the detoxification of toxic substances by the relay of sulfur atom through its active site Cys residue. In this study, GSU2940 was produced in E. coli, purified, and characterized.

Method: gsu2940 was cloned into pColdI and expressed in E. coli BL21(DE3). GSU2940 was purified as a His-tag fusion protein. A portion of gsu2940 was replaced with a kanamycin-resistance gene to produce the gsu2940-disrupted mutant strain.

Result: The purified GSU2940 exhibited a rhodanese activity with the optimum pH at 7.5 and the optimum temperature at 55°C. Thermal stability (T_m) of GSU2940 was determined to be 40°C. Furthermore, the rhodanese activity in the gsu2940-disrupted strain, was remarkably decreased as compared with that in the wild-type strain, suggesting that GSU2940 is the dominant sulfurtransferase in G. sulfurreducens.

Discussion: Although GSU2940 exhibited a rhodanese activity, it shows only a slight sequence identity with already-known rhodaneses. GSU2940 is proposed to cooperate with MHSEP and other proteins encoded by the same gene cluster.
**P30 - A novel Geobacteraceae-specific outer membrane protein required for selenite and tellurite reduction**

**2. Selenium in the molecular life sciences**

**2.1 Metabolism of selenium in living cells**

Keywords: Porin, selenoprotein, selenite and tellurite reduction, porin-cytochrome complex, Geobacter

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**Introduction:** A putative outer membrane channel protein, GSU2939, was found in a metal-reducing bacterium, *Geobacter sulfurreducens*. The *gsu2939* gene is part of an operon together with 10 other genes including the multi-heme-containing selenoprotein, but the function of GSU2939 remains unexplored. GSU2939 showed only less than 20% homology with phosphate selective porins belonging to the porin P_O family, suggesting that GSU2939 is a Geobacteraceae-specific novel type of porin. Herein, we investigated the characteristics of the putative outer membrane protein GSU2939.

**Method:** Expression level of *gsu2939* was examined under various growth conditions. A *gsu2939*-deficient mutant strain was constructed to investigate its phenotype.

**Result:** The *Geobacteraceae*-specific putative outer membrane porin, GSU2939, showed impermeability in β-lactam antibiotic transportation. Investigation on the expression levels of *gsu2939* at different growth phases in the wild-type cells showed that GSU2930 is a log-phase specific protein. The expression of *gsu2939* was neither affected by phosphate-starving conditions nor osmotic-stressed conditions. However, in the presence of selenite and tellurite, a *gsu2939*-deficient mutant showed more susceptibility than the wild-type strain, suggesting that the protein may not function as a typical outer membrane channel. Furthermore, the *gsu2939*-deficient mutant showed lower reducing ability for ferric citrate, selenite, and tellurite as an electron acceptor, compared with the wild type.

**Discussion:** Our data suggest that GSU2939 may function as a porin-cytochrome protein complex together with the multi-heme-containing selenoprotein and c-type cytochrome proteins encoded in the same gene cluster and facilitate a trans-outner membrane electron transfer during extracellular reduction of ferric citrate, selenite, and tellurite.
P31 - Effect of inorganic and organic selenium compounds on tumor infiltrating lymphocytes

2. Selenium in the molecular life sciences
2.2 Molecular mechanisms of selenium toxicity

Keywords: ovarian cancer, tumor infiltrating lymphocytes

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Introduction: Ovarian cancer is characterized by its unique immunosuppressive tumor microenvironment which alters many cellular processes including immune surveillance (1). This impaired functionality of tumour infiltrating lymphocytes (TILs) can lead to defective cytotoxic activity, diminished secretion of lymphokines and failure to proliferate in response to stimulation. Selenium (Se) has known anticancerous effects with many proposed mechanisms including enhanced immune surveillance (2). Little is however known about the different effects between various Se compounds and their specific effect on immune cells, which we aimed to study herein.

Method: Immune cells were isolated using density gradient centrifugation and were further selected by magnetic beads. Se cytotoxicity was then compared between tumour and immune cells. Following activation of immune cells and preincubation of organic or inorganic selenium compounds with tumour cells, lysis assay was performed. ELISA, qPCR and flow cytometry based analyses were used to analyse markers of the immune response.

Result: Tumour cells were found to be more sensitive to the cytotoxic effects exerted by the Se compounds compared to the immune cells from both healthy donors and TILs from ovarian cancer patients. Our preliminary experiments have also shown effects of Se on immune cell activation, with changes in expression of central genes.

Discussion: In conclusion, Se compounds have the potential to act as anticarcinogenic agents in ovarian cancer at concentrations that do not kill or inhibit the immune cells, and concurrently facilitate the immune response.

Selected references
P32 - Small Selenium Species: Toxicity, Bioavailability and Modes of Action in Caenorhabditis elegans

2. Selenium in the molecular life sciences
2.2 Molecular mechanisms of selenium toxicity

Keywords: selenite, selenomethionine, Se-methylselenocysteine, Caenorhabditis elegans, ICP-QQQ-MS

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Introduction: Small selenium species play a major role in selenium metabolism, but their either toxic or protective potential, bioavailability and metabolic transformations are not fully understood. The roundworm Caenorhabditis elegans is an emerging model in toxicological research and enables to study these endpoints in a whole metabolizing organism. The simplicity of the roundworm having only one selenoprotein (TrxR-1) might give deeper insight into selenium metabolism.

Method: Toxicity of selenite, selenomethionine and Se-methylselenocysteine was assessed by lethality and development assays. Total selenium quantification and selenium speciation analysis were performed by (HPLC)-ICP-QQQ-MS based methods. Reactive oxygen and nitrogen species (RONS) and TrxR activity were measured using fluorimetric and colorimetric assays.

Result: Selenite exerted strongest toxic effects (LD₅₀ = 12 mM after 30 min incubation in the first larval stage) and caused a developmental delay of worm larvae. In contrast, selenomethionine and Se-methylselenocysteine were only slightly toxic (survival rate decreased by 20% at 50 mM). Total selenium quantification revealed that all three species were bioavailable, with an inverse relationship between toxicity and bioavailability. Speciation analysis indicated that selenomethionine and Se-methylselenocysteine are partially metabolized to other small selenium species. Protection against RONS induction was observed at a low dose range (1 – 100 µM) for all three selenium compounds.

Discussion: Our findings indicate that selenomethionine and Se-methylselenocysteine seem to be utilized by Caenorhabditis elegans and undergo, at least in part, metabolism processes. Independently, all three selenium species shared a protective potential against oxidative stress, while the underlying mechanism remains to be clarified.
P33 - Diabetic fish on selenium supplements

2. Selenium in the molecular life sciences
2.2 Molecular mechanisms of selenium toxicity
Keywords: selenomethionine, toxicology, fish, metabolism, cardiac function

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Introduction: A variety of anthropogenic activities increase loading of selenium into aquatic ecosystems, which poses an extreme toxicological hazard to fishes. Although many studies have reported developmental toxicities in larval fishes, fewer studies have investigated sublethal toxicological effects that may occur following dietary selenium exposure in adult fishes.

Method: Adult zebrafish were exposed to dietary selenomethionine (SeMet) at Se-normal levels (1.3 \(\mu\)g Se/g food, dry mass) and supraphysiological levels (3.4, 9.8, or 27.5 \(\mu\)g/g) for 90 days. Swimming performance, \(\text{O}_2\) consumption and metabolic rates were determined using a swim tunnel respirometer. Cardiac function was assessed using high resolution (30 \(\mu\)m) ultrasound biomicroscopy. Energy stores (triglycerides and glycogen) and mRNA transcript abundance of selected genes were determined.

Result: Compared to controls, zebrafish exposed to elevated dietary SeMet exhibited impaired swimming performance (lower fatigue velocity, or \(U_{\text{crit}}\)). This was associated with elevated basal metabolic rate and reduced aerobic scope, indicating impaired aerobic capacity. Triglycerides (the primary fuel for aerobic swimming) were elevated in a dose-dependent manner, which was associated with altered transcript abundance of several genes involved in lipid homeostasis. Ultrasonography revealed decreased cardiac output, which was associated with increased echodensity at the atrial-ventricular junction and reduced mRNA expression of the collagenase, MMP2.

Discussion: These results suggest significant ecophysiological effects that may impair the fitness of fishes exposed to elevated dietary Se in contaminated ecosystems. From a comparative biomedical viewpoint, these results highlight the utility of zebrafish as a model to investigate mechanisms of metabolic, energetic, and cardiovascular toxicities caused by excessive dietary Se exposure.

Selected references


P34 - Biochemical characterization of novel methylseleno derivatives

2. Selenium in the molecular life sciences
2.2 Molecular mechanisms of selenium toxicity

Keywords: methylselenol, pancreatic cancer cells, chemotherapeutic drugs.

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Introduction: Redox active selenium (Se) compounds at subtoxic doses, act as pro-oxidants and are highly cytotoxic to tumor cells and are foreseen as promising candidates as chemotherapeutic agents (1,2). Methylselenol, a key metabolite of Se compounds, is known to be involved in apoptosis and anti-mitogenic processes (2). However, the high reactivity of methylselenol limits its use as a therapeutic drug. In the present study, we aim to test the effects of novel compounds containing a methylselenol moiety.

Method: These novel compounds were tested and characterized in a pancreatic cancer cell line (Panc-1). Methylseleninic acid (MSA), a penultimate precursor of methylselenol, was used as a positive control (3). Experiments were performed in 2D cultures and 3D spheroids using viability assays, clonogenic assay, immunofluorescent (IF) staining and western blotting against a panel of cell death markers. Expression of cellular adhesion markers was analyzed using flow cytometry and IF staining.

Result: Our compounds induced cell death, while analogues lacking the methylselenol moiety were inert. Clonogenic assay revealed loss of colony formation ability of live cells post treatment. Cell death was associated with loss of cellular adhesion and plasma membrane integrity, and was observed with a change in expression of different cellular adhesion markers like integrin beta 1 and cadherins.

Discussion: The novel Se compounds induce cytotoxicity in Panc-1 cells, with one compound being more efficient than MSA. Both MSA and our new compounds display similar mechanism of action, hence suggesting that the key player in inducing cellular death is the methylselenol moiety.

Selected references


P35 - Metabolomics Profiling of the Liver of the Selenocysteine Lyase Knockout Mouse

2. Selenium in the molecular life sciences
2.3 Molecular consequences of selenium deficiency
Keywords: selenocysteine lyase, metabolomics, glycine

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Introduction: The role of the micronutrient selenium in metabolic disorders remains controversial, with epidemiological studies pointing to either a protective or a deleterious correlation. In cells, selenium is processed into selenocysteine (Sec), an amino acid present at the core of selenoproteins. Selenoproteins help to cope with oxidative stress, a hallmark of disorders affecting energy metabolism. The enzyme Sec lyase (Scly) decomposes Sec into alanine and selenide, the latter being reutilized in selenoprotein translation. We previously demonstrated that mice lacking Scly (Scly−/−) develop all characteristics of a metabolic syndrome (MS) phenotype1, including insulin resistance, obesity, glucose intolerance and fatty liver. We thus investigated the role of Scly in hepatic energy metabolism.

Method: We performed metabolomics profiling analysis in the livers of Scly−/− mice fed a selenium adequate and selenium deficient diets to unveil metabolic pathways potentially involved in the Scly−/− metabolic phenotype.

Result: Our preliminary analysis revealed enriched metabolites for amino acid metabolism, specifically glycine, serine, threonine, and creatine, in Scly−/− livers. We further detected elevated expression of dimethylglycine dehydrogenase, a key enzyme in glycine metabolism, in Scly−/− mice. Moreover, the transsulfuration pathway that degrades selenomethionine, and cysteine and methionine metabolism, were affected in a sex-specific manner in the Scly−/− mouse.

Discussion: Our results point to an additional role of Scly in the regulation of glucogenic amino acid metabolism. Ultimately, these results will help to improve our knowledge of the complex relationship between dietary selenium and metabolic disorders.

Selected references
P36 - Cardiac muscle necroptosis induced by selenium deficiency implicates miR-200a activating

2. Selenium in the molecular life sciences
2.3 Molecular consequences of selenium deficiency

Keywords: Necroptosis; Myocardial cells; MiR-200a; Chick; RNF11

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Introduction: Necroptosis have characterized new forms of the paradigm of cell death. At the outset, we found necroptosis may play a key role in heart disease\(^1\), and additionally with the selenium deficiency. However, the related of necroptosis with selenium deficiency in heart disease are still largely unknown.

Method: We chose miR-200a for our study and explored the role of miR-200a in selenium-deficient myocardial necroptosis by dual luciferase reporter assay and RT-PCR. And then we detected endogenous expression of miR-200a by RT-PCR method and transfected miR-200a mimic and inhibitor in myocardial cells to examine the expression of necroptosis related genes and inflammation related genes by quantitative real-time PCR and Western blotting. We observed ultrastructure in myocardial induced by selenium deficiency and transfected myocardial cells, meantime, we performed AO/EB staining and flow cytometry.

Result: The mRNA expression of the inflammation related genes and necroptosis related genes in vivo and in vitro showed significant differences, miR-200a mimic and selenium deficiency cardiac muscle revealed inflammation and necroptosis. According to the results of electron microscope observation and AO / EB staining and flow cytometry, we confirmed that selenium deficiency induced myocardial necroptosis in chicken possible in part by activiting miR-200a and inhibiting RNF11 (Fig1).

Discussion: Based on the biological function identified target genes RNF11, we hypothesized that mir-200a suppress RNF11 to induce necroptosis in selenium deficiency cardiac muscle and may drive the dysfunction of heart.

Selected references

P37 - Selenophosphate synthetase 1 is an essential protein with roles in regulation of redox homeostasis

2. Selenium in the molecular life sciences
2.4 Selenoprotein synthesis pathways

Keywords: cancer, reactive oxygen species, redox regulation, selenophosphate synthetase 1

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Introduction: Selenophosphate synthetase (SPS) was initially detected in bacteria and was shown to synthesize selenophosphate, the active selenium donor. However, mammals have two SPS paralogues, which are designated SPS1 and SPS2. Although it is known that SPS2 catalyses the synthesis of selenophosphate, the function of SPS1 remains largely unclear.

Method: To examine the role of SPS1 in mammals, a Sps1 knocked-out mouse line was generated, and microarray analysis was performed to pinpoint the genes affected by SPS1-deficiency. Several assays such as cell growth rate assay, Sps1-knockdown/rescue vector overexpression assay in cell line, and reactive oxygen species (ROS) detection assay were also performed to validate alterations in gene expression and their phenotypic influences due to the absence of SPS1.

Result: SPS1 tKO mice revealed some differences from WT mice in size of embryo and organ morphology. Loss of Sps1 in the liver showed differences in stress-related selenoproteins and cellular level of iron and manganese. And gene expression analysis of Sps1 suggested linkage between Sps1, GSH system, and cancer properties. Loss of Sps1 in F9 cells showed reduced expression level of Glrx1, Gsto1, and SelW. And loss of Sps1 in F9 cells also showed different level of hydrogen peroxide due to down-regulation of Glrx1 and also showed reverse cancer characteristics. Loss of Sps1 in drosofila cells showed decreased level of PLP.

Discussion: Our observations suggest that SPS1 influences levels of hydrogen peroxide and hydrogen peroxide-related selenoproteins, regulates redox homeostasis-related gene expression, and reverses cancer characteristics in cancer cell line.

Selected references
P38 - Influences of TRIT1 catalysed tRNA-modification i6A37 on translation efficiency of selenoproteins

2. Selenium in the molecular life sciences
2.4 Selenoprotein synthesis pathways

Keywords: TRIT1, tRNA modification, i6A, ribosome profiling, translation efficiency

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Introduction: tRNA modifications in the anticodon loop are crucial for translation efficiency, fidelity and mRNA stability. The enzyme tRNA-isopentenyltransferase 1 (TRIT1) modifies adenosines at position 37 in tRNAs with an isopentenyl moiety at N⁶ (i⁶A₃₇). We want to focus on the defined group of selenoproteins, which are depending on one specific tRNA, the tRNA^[Ser]Sec. It is known that selenoprotein expression is reduced, when tRNA^[Ser]Sec lacks i⁶A₃₇ [1][2]. The i⁶A₃₇ modification is found in cytosolic and in mitochondrial tRNAs. A patient with homozygous TRIT1 p.Arg323Gln point mutation, which causes mitochondrial disorder, was identified [3].

Method: Western blot analysis of selenoproteins and ⁷⁵Se labeling was performed on patient fibroblasts. In order to find out whether the mutation in TRIT1 alters interaction with tRNA^[Ser]Sec, we have set up an in vitro assay for tRNA isopentenylation reaction. We will examine direct influences of i⁶A₃₇ on mRNA abundance using RNA-seq data derived from liver samples of our newly generated Trit1-deficient mice. Following, translation efficiency at single codon resolution will be measured by focusing on i⁶A₃₇ depended codons with ribosome profiling in cytosol and mitochondria [4].

Result: Patient fibroblasts showed no reduced selenoprotein expression. The point mutated TRIT1 enzyme seems to have a decreased activity compared to wildtype TRIT1 in vitro. First experiments on mitochondria derived from our Trit1-deficient mice suggested, that the membrane potential of mitochondria seems to be affected. Brain specific Trit1-deficient mice have microcephaly and showed seizures.

Discussion: Ribosomal profiling will reveal how individual codons are affected by lack of i6A in tRNAs.

Selected references
P39 - Genetic screening for unknown factors in selenium metabolism in Archaea

2. Selenium in the molecular life sciences
2.4 Selenoprotein synthesis pathways
Keywords: selenocysteine, Archaea, Methanococcus, regulation, random mutagenesis

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Introduction: Selenocysteine (sec) -containing proteins (selenoproteins) are found in all three domains of life Archaea, Bacteria, and Eukarya. Most selenoproteins of the archaeal model organism Methanococcus maripaludis are involved in its primary metabolism, hydrogenotrophic methanogenesis. Interestingly, cysteine (cys) -containing isoforms of all selenoproteins (with the exception of formate dehydrogenase), are formed when selenium is scarce or when the path of sec synthesis is disrupted. While formate-dependent growth is impaired under such conditions, the organism can still grow on H₂ + CO₂. We searched for unknown factors involved in selenium metabolism of M. maripaludis, like those sensing the selenium status of the cell or those transducing the information to effect antagonistic regulation of the selenoprotein genes and their cys-encoding isogenes.

Method: A transcriptional fusion comprising the promoter of frcA (encoding a subunit of the selenium-free, F₄₄₂₀-reducing, hydrogenase Frc), PfccA, and bla (encoding beta-lactamase from E. coli) was placed on the chromosome of M. maripaludis. In order to identify factors involved in selenium metabolism, the reporter strain was subjected to random mutagenesis.

Result: For more than 6,000 transposon mutants, the Bla activity and the ability to grow with formate was assessed. For those mutants exhibiting the desired phenotype, the transposon insertion site on the chromosome was mapped. Among other factors, a putative membrane protein was identified, which might serve as a sensor of the exogenous selenium supply.

Discussion: Characterizing new factors involved in the selenium metabolism of M. maripaludis, will aid the understanding of how Archaea cope with fluctuating supply of this important trace element.
P40 - A role of bacterial thioredoxin in selenide delivery to selenophosphate synthetase in vitro

2. Selenium in the molecular life sciences
2.4 Selenoprotein synthesis pathways
Keywords: selenoprotein biosynthesis, bacteria, selenophosphate, thioredoxin, selenide

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Introduction: In bacteria, a key selenium intermediate for selenoprotein biosynthesis, selenophosphate, is synthesized by selenophosphate synthetase (SelD), which has been shown to use selenide (HSe⁻) and ATP as in vitro substrates. However, little is known about the metabolic pathway for HSe⁻ generation and the mechanism of specific supply of HSe⁻ to SelD. In this study, we investigated whether purified TrxA could reduce selenite to provide the HSe⁻ substrate to SelD.

Method: The trxA, selD, and trxB (encoding thioredoxin reductase) genes of Pseudomonas sp. F2a were cloned and expressed as His-tag fusion proteins in E. coli. To elucidate the function of a Trx motif (WCxxC), two mutant TrxAs, C33A and C36A, were generated. Using the purified TrxAs, a selenite reduction assay and a SelD-coupled assay were performed.

Result: The wild-type TrxA showed a selenite reduction activity in the presence of TrxB, whereas significant decreases in the activity were observed in the mutants C33A and C36A compared to the wild-type TrxA. In the SelD-coupled assay, di-thiol reductants, TrxA and DTT, reduced selenite to efficiently provide HSe⁻ to SelD. In contrast, mono-thiol reductants, β-mercaptoethanol and glutathione, did not facilitate the catalytic reaction of SelD.

Discussion: Together with our previous results, TrxA may play a pivotal role in selenoprotein biosynthesis as a HSe⁻ supplier to SelD in bacteria.
P41 - Selenoprotein S is involved in degradation of C99 through ERAD.

2. Selenium in the molecular life sciences
2.6 Selenoprotein function

Keywords: Alzheimer's disease, Amyloid beta, C99, ERAD, Selenoprotein S

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Introduction: In amyloidogenic pathway, C99 is produced from Amyloid beta precursor protein (APP) by β-secretase (BACE1) in the endoplasmic reticulum (ER) membrane. C99 is cleaved to amyloid β which is known to cause Alzheimer’s disease by its aggregation. Selenoprotein S (SelS) is an ER-resident selenoprotein involved in endoplasmic reticulum-associated degradation (ERAD). ERAD has been known as one of the processes to clear C99, but it remains unclear if SelS is required for regulation of C99 through ERAD.

Method: We used γ-secretase inhibitor (DAPT) and mutant C99 construct, because C99 is rapidly cleaved to Aβ. To investigate whether C99 is regulated by SelS, we used SelS knock-downed N2a cells. To identify that C99 is the substrate of SelS-dependent ERAD, we performed cycloheximide chase assay, immunoprecipitation, and ubiquitination assay.

Result: While the SelS was increased by ER stress, C99 level was decreased. However, there was no change of C99 level in SelS knock-down cells, despite the induction of ER stress. C99 was proved to be the substrate of SelS-dependent ERAD by confirming that the interaction of C99 with p97 (VCP) did not occur in SelS knock-down cells. Moreover, the ubiquitination of C99 was decreased in SelS knock-down cells. We also found that the extracellular level of Aβ1-42 was increased by SelS knock-down.

Discussion: Triggering the C99 degradation before Aβ formation is a potential therapeutic target for preventing the disease. We demonstrated that SelS is involved in C99 clearance through ERAD, resulting in reduction of Aβ production.

Selected references

P42 - Computational identification of the selenocysteine tRNA (tRNA-Sec)

2. Selenium in the molecular life sciences
2.4 Selenoprotein synthesis pathways
Keywords: tRNA-Sec

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Introduction: The selenocysteine tRNA (tRNA^Sec) drives the recoding of UGA codons from stop signal to Sec in selenoprotein transcripts. Although found in organisms from the three domains of life, Sec is not universal. Many organisms are completely devoid of selenoprotein genes and lack the ability to synthesize Sec. Since tRNA^Sec is a key component in selenoprotein biosynthesis, its efficient identification in genomes is instrumental to characterize the utilization of Sec across lineages. Available tRNA prediction methods fail to accurately predict tRNA^Sec due to its unusual structure.

Method: We present Secmarker [1], a method based on manually curated covariance models capturing the specific tRNA^Sec structure in archaea, bacteria and eukaryotes. We exploited the non-universality of Sec to build a proper benchmark set for tRNA^Sec predictions, which is not possible for other tRNAs. We show that Secmarker greatly improves the accuracy of previously existing methods.

Result: The analysis of a large set of fully sequenced genomes revealed new insights in the biology of tRNA^Sec, led to the discovery of a novel bacterial selenoprotein family, and shed additional light on the phylogenetic distribution of selenoprotein containing genomes. Secmarker is freely accessible for download, or online analysis through a web server at http://secmarker.crg.cat.

Discussion: We describe here the development and validation of Secmarker, a tool to predict tRNA^Sec in genomes.

Selected references
P43 - The use of dimedone to study redox states of selenoproteins

2. Selenium in the molecular life sciences
2.6 Selenoprotein function
Keywords: selenocysteine, dimedone, selenenic acid, seleninic acid

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Introduction: Dimedone is a widely used reagent to assess the redox state of cysteine-containing proteins as it will alkylate sulfenic acid residues, but not sulfenic acid residues. While it has been reported previously that dimedone can label selenenic acid residues in selenoproteins, we investigated the stability, and reversibility of this label in a model peptide system. We also wondered whether dimedone could be used to detect seleninic acid residues due to the much stronger electrophilic character of selenium relative to sulfur.

Method: We used model chemical compounds, benzenesulfinic acid, benzeneselenenic acid, and a model selenocysteine-containing peptide to investigate possible reactions with dimedone. For the peptide, the reaction was initiated by reaction with H₂O₂. The reactions were followed by ¹H-NMR, ⁷⁷Se-NMR, and ESI-MS.

Result: As expected, benzenesulfinic acid did not react with dimedone. However, benzeneselenenic acid reacted with dimedone to yield diphenyldiselenide and a dimedone dimer. The model peptide could be labeled with dimedone at low concentrations of H₂O₂, but the reaction was reversible by addition of thiol, disulfide, or diselenide. At high concentrations of H₂O₂ selenium was eliminated from the peptide and the dimedone dimer could be detected.

Discussion: Dimedone is not a good reagent for detecting selenenic acids in selenoproteins due to the reversible nature of this alkylation. However, dimedone can potentially be used to detect seleninic acid residues. The reaction of dimedone with oxidized cysteine residues is quite different from the same reaction with oxidized selenocysteine residues.
P44 - Site specific acetylation of thioredoxin reductase 1

2. Selenium in the molecular life sciences
2.6 Selenoprotein function
Keywords: orthogonal translation, post-translational modification, redox, selenocysteine, thioredoxin reductase

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Introduction: Thioredoxin reductase 1 (TrxR1) is an essential human selenoprotein involved in ROS defense. Overactive TrxR1 is a diagnostic marker for some cancers (1), and a drug target for some drug resistant cancers (2). TrxR1 acetylation sites have been experimentally characterized, but the effect of acetylation at specific sites on the activity of TrxR1 has not been determined. TrxR1 exists as a catalytically active dimer, but can form low activity tetramers or higher order multimers (3). The acetylation sites occur on the surface of the dimer, at the dimer dimer interface in the tetramer (3).

Method: We employed genetic code expansion to produce site specifically acetylated TrxR1 in E. coli. This was accomplished by reassigning UAG stop codons to acetyl-Lysine (acK) using an orthogonal tRNA, tRNA acK synthetase pair, as well as using E. coli’s endogenous selenocysteine insertion system to recode a single UGA codon to selenocysteine.

Result: We found that site specific acetylation of TrxR1 increased TrxR1 activity on DTNB. Acetylated TrxR1 variants eluted in later fractions from a size exclusion column under native conditions compared to unacetylated TrxR1, indicating a shift towards lower molecular weight oligomers. The effect of acetylation on TrxR1 activity was reversed following incubation of acetylated TrxR1 with histone deacetylase 3.

Discussion: We discovered that acetylation increases TrxR1 activity, likely by destabalizing low activity tetramers and promoting the formation of catalytically active dimers (Figure 1). This provides new insight into how TrxR1 activity is regulated in the cell, and could be relevant to diseases involving TrxR1.

Selected references


Figure 1. Acetylation-dependent TrxR1 activation mechanism. (A) Acetylation of K141 (Blue), K200 (Purple) or K307 (Red) on the surface of TrxR1 dimers breaks key salt bridges and hydrogen bonds (B-C) in the TrxR1 tetramer interface. Close up view of key interactions at the tetramer interface by (B) K141, (C), K200 and K307 (PDB: 4KPR (3)).
P45 - Thioredoxin reductase 1 and NADPH directly protect protein tyrosine phosphatase 1B from inactivation

Abstracts

2. Selenium in the molecular life sciences
2.6 Selenoprotein function

Keywords: PTP1B, TrxR1, Prx2, redox regulation

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Introduction: Protein tyrosine phosphatases (PTPs) are regulated by oxidation of their active site cysteine during growth factor signaling (1). Recent work by us has identified thioredoxin reductase (TrxR1), in combination with thioredoxin (Trx1) as activators of PTP1B both in vitro as well as during cellular signaling (2). How PTP1B can be oxidized in the presence of peroxiredoxins and the thioredoxin system is however insufficiently unknown. Here we have characterized the influence of different components of the Trx system on hydrogen peroxide-mediated PTP inactivation.

Method: Recombinant PTP1B, TrxR1, Trx1 and Prx2 were used to mimic cellular growth factor dependent PTP regulation in vitro under different conditions of oxidative bursts, PTP activity was assessed using a chromogenic substrate (pNPP).

Result: We found that H₂O₂ cannot efficiently inactivate PTP1B in the presence of complete TrxR1 / Trx1 / Prx system. Our results are against a transfer of oxidative equivalents from Trx1 or Prx2 to PTP1B upon burst of H₂O₂. We found that TrxR1 and NADPH (without Trx) was able to protect reduced PTP1B if present during exposure to hydrogen peroxide. The protective effect was dependent on TrxR1 concentration, required an intact selenocysteine residue, and was blocked by the TrxR1 inhibitor auranofin.

Discussion: TrxR1 was shown to prevent H₂O₂ dependent PTP1B inactivation. We found that Trx or Prx2 also protect PTP1B from inactivation (3). Mechanisms of PTP1B inactivation by H₂O₂ still need to be characterized.

Selected references


Dagnell et al. "Manuscript in preparation" Thioredoxin reductase 1 and NADPH directly protect protein tyrosine phosphatase 1B from inactivation during exposure to H₂O₂
P46 - Details in the catalytic mechanism of mammalian selenoprotein thioredoxin reductase 1 revealed using point mutations and juglone reducing activities

Abstracts

2. Selenium in the molecular life sciences
2.6 Selenoprotein function

Keywords: Redox Cycling, Superoxide, NADPH Oxidase Activity, Electron Transfer, Thioredoxin Reductase, Juglone

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Introduction: The mammalian selenoprotein thioredoxin reductase 1 (TrxR1) is a key enzyme in redox regulation, antioxidant defense, and cellular growth. TrxR1 catalyzes efficient reduction of juglone (5-hydroxy-1,4-naphthoquinone; walnut toxin), however, the details in the catalytic mechanism of TrxR1-mediated reduction of juglone and most other substrates, remain unclear.

Method: Using a number of TrxR1 mutant variants, we here found that a sole Cys residue at the C-term tail of TrxR1 is required for high-efficiency juglone-coupled NADPH oxidase activity of Sec-deficient enzyme, occurring with mixed one- and two-electron reactions producing superoxide. The activity also utilizes the FAD and the N-terminal redox active disulfide/dithiol motif of TrxR1.

Result: If a sole Cys residue at the C-term tail of TrxR1, in the absence of Sec, was moved further towards the C-term end of the protein compared to its natural position at residue 497, juglone reduction was, surprisingly, further increased. Four residues of Sec-deficient TrxR1 were found to be easily arylated by juglone, including the Cys residue at position 497.

Discussion: Based upon our observations, we suggest a model for involvement of the juglone-arylated C-term motif of TrxR1 to explain its high activity with juglone. This study thus provides novel insights into the catalytic mechanisms of TrxR1. One-electron juglone reduction by TrxR1 producing superoxide should furthermore contribute to the well-known prooxidant cytotoxicity of juglone.

Selected references
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P47 - Probing the role of TRP14 (TXNDC17) in cellular signaling pathways

2. Selenium in the molecular life sciences
2.6 Selenoprotein function

Keywords: TRP14, pTRAF, Nrf2, NFkB, HIF

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Introduction: Thioredoxin related protein of 14 kDa (TRP14, also named TXNDC17) is a cytosolic protein of the thioredoxin fold family expressed in all tissues, and is an efficient substrate of the selenoprotein thioredoxin reductase 1 (TrxR1, TXNRD1). TRP14 can not support the activities of classical Trx1 substrates such as peroxiredoxins, methionine sulfoxide reductases or ribonucleotide reductase, but affects NFκB signaling (1). TRP14 is however efficient in reducing L-cystine and support S-denitrosylation reactions (2), reduce protein persulfides (3), and can reactivate oxidized protein tyrosine phosphatase PTP1B (4). These and other observations suggest that TRP14 may have dedicated roles in cellular signaling pathways, which was studied here.

Method: Using our previously developed (5) pTRAF tool (plasmid for transcription factor reporter activation based upon fluorescence), we assessed activation patterns of the transcription factors Nrf2, NFκB and HIF in HEK293 cells with or without modulated TRP14 levels.

Result: Our preliminary results suggest that TRP14 is an unusual member of the Trx system, in that TRP14 may act as a repressor of NRF2, modulate HIF activation under hypoxia as well as acting as a weak repressor of NFκB activation. Further analyses are needed to fully evaluate the pTRAF readout (see Figure) also including additional methodologies.

Discussion: Our results point towards the role of TRP14 as a dedicated fine-tuning modulator of redox regulated signaling pathways. The many observations suggesting that altered levels or activities of the selenoprotein TrxR1 can affect cellular phenotype should thereby consider the possibility that TRP14 may be part of the molecular mechanisms explaining such effects. We are currently further probing this hypothesis.

Selected references

P48 - Apatone inhibited GPx activity and triggered AIF-mediated cell death pathway in cancer cells

2. Selenium in the molecular life sciences
2.6 Selenoprotein function

Keywords: Apatone, oxidative stress, replicative stress, glutathione peroxidase, cell death

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Introduction: Apatone (Vitamin C and Vitamin K3), is an investigational drug under clinical trials for cancer treatment. Extracellular VC is quickly oxidized to dehydroascorbic acid (DHA) and taken up by cells via glucose transporters, over-expressed in many cancer cells. It has been shown that Apatone kills cancer cells by inducing intracellular hydrogen peroxide via a redox cycling reaction \(^1\) and caused oxidative and replicative stress in different cancer cell lines. However, the exact cell death mechanism has not been fully understood yet.

Method: Neuroblastoma cell line A172 and prostate cancer cell line LNCaP were treated with Apatone (VC: K3=500 µM : 5 µM), cellular lipid peroxidation was measured by thiobarbituric acid (TBA) assay. Glutathione peroxidase (GPx) activity was measured by a glutathione-glutathione reductase (GR)-NADPH -coupled method. Nuclear translocation of the apoptosis-inducing factor (AIF) was measured by western blot.

Result: A172 and LNCaP cell lines showed different sensitivity to Apatone treatment. Apatone treatment enhanced lipid peroxidation in both cell lines, however, A172 exhibited a more significant increase than LNCaP cells, which is consistent with the viability data. GPxs are the main seleno-enzymes reducing lipid peroxides. We found Apatone inhibited GPx activity both on pure protein level and cellular level. AIF has been reported to be a lipid peroxide sensitive cell death mediator \(^2\). Apatone treatment promoted AIF nuclear translocation which contributed to cell death.

Discussion: In this study, we elucidated an AIF-mediated cell death pathway which is triggered by lipid peroxidation and GPx inhibition caused by Apatone.

Selected references


P49 - The significance of selenoprotein P expression in pancreatic beta-cell line MIN6

2. Selenium in the molecular life sciences
2.6 Selenoprotein function

Keywords: Selenoprotein P, Antioxidant, MIN6, Insulin, Cell death

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Introduction: Selenoprotein P (SeP), a selenocysteine (Sec) -containing protein, is mainly synthesized in the liver and functions as a selenium (Se) -supply protein to maintain Se levels in several tissues 1). Se is essential for the synthesis of antioxidative selenoproteins, such as glutathione peroxidases (GPxs) and thioredoxin reductases (TRs). Hence, SeP contributes to cellular antioxidative system via the maintenance of selenoproteins. Several studies have reported that pancreatic beta-cells are vulnerable to oxidative stress and antioxidant system is essential for beta-cells 2). Recently, we found that MIN6 cells express SeP. In the present study, we investigated the significance of SeP expression in MIN6 cells.

Method: We established SeP-deficient MIN6 cells by using siRNA. We used western blotting to evaluate expression levels of proteins such as insulin and selenoproteins (GPx1, GPx4 and TR1). Real-time PCR was used to measure these mRNA expression levels. WST-8 assay was used to evaluate cell viability.

Result: SeP-siRNA significantly decreased SeP mRNA levels in MIN6 cells. Knockdown of SeP resulted in significant reduction in protein levels of selenoproteins and insulin in MIN6 cells. The decrease in cell viability was also observed by WST-8 assay. Addition of Sec to SeP-deficient MIN6 cells recovered these protein levels and cell viability. Alpha-Tocopherol, a classical lipophilic antioxidant, also recovered cell viability.

Discussion: This report provides the significance of SeP expression in MIN6 cells for maintaining levels of selenoprotein, insulin and cell viability. The effect of alpha-tocopherol and possible cell death signaling will be discussed.

Selected references
P50 - Interaction between selenoprotein W and 14-3-3 is regulated by oxidative stress.

2. Selenium in the molecular life sciences
   2.6 Selenoprotein function

Keywords: SelW, 14-3-3, Oxidative stress

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Introduction: Mouse Selenoprotein W (SelW) contains a selenocysteine (sec) and four cysteines. N-terminal of SelW contains a CXXU motif, corresponding to the thioredoxin active site CXXC motif. Thus SelW has been none seen as a putative antioxidant protein that interaction with other redox regulators. We have previously reported that the binding of 14-3-3 protein to its target proteins, including CDC25B, Rictor and TAZ, is inhibited by the interaction of 14-3-3 protein with SelW. In this study, we sought to determine the binding site of SelW to understand the regulatory mechanism of the interaction between SelW and 14-3-3 and its biological effects.

Method: The interaction between SelW and 14-3-3 was determined by both immunoprecipitation assay and GST-pull-down assay using various purified GST-SelW mutants. To understand the reguratory mechanism, the interaction was also determined in oxidative stress condition. Cell cycle rescue assay was performed to examine the biological effect of the interaction using SelW-knockdown MCF7 cells.

Result: A mutant SelW in which Cys (U13C) is changed to Ser (U13S) was unable to interact with 14-3-3 protein and thus did not inhibit the interaction of 14-3-3 to other target proteins. However, other Cys mutants of SelW (C10S, C33S and C37S) normally interacted with 14-3-3 protein. Also, the interaction of SelW to 14-3-3 protein was enhanced by oxidative stress. We also found that SelW (U13S) was not able to rescue the effect of SelW depletion.

Discussion: Thus, SelW may have a regulatory function in redox cell signaling by interacting with 14-3-3 protein.

Selected references


P51 - Immortalized human prostate cell line RWPE-1 as a model to study selenoprotein regulation and function in normal prostatic tissue

2. Selenium in the molecular life sciences
2.6 Selenoprotein function
Keywords: selenoproteins, GPX1, SELENOF, SBP1, RWPE1

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Introduction: While epidemiology has indicted a likely impact of selenium status on prostate cancer risk, the role of selenoproteins in this disease remains to be determined. Many studies have investigated selenoprotein levels and function in prostate cancer cell lines, but little is known about the function of several of these proteins in normal prostate epithelia. Towards this goal, studies have focused on an immortalized but not transformed human prostate cell line, RWPE-1.

Method: GPX1, SELENOF and SBP1 levels were determined by western blotting with specific antibodies with and without treatment with 100 nM sodium selenite. The cellular localization of these proteins in RWPE-1 cells was evaluated by confocal microscopy and compared to that seen in human prostate cancer cell lines and primary prostate epithelia cells using the same techniques, as well as immunohistochemistry of human tissue.

Result: SELENOF was predominantly located in the outer membrane of RWPE-1 cells as was observed for primary cells and benign human prostate tissue. Unlike what is typically seen in prostate cancer cell lines, the levels of GPX1 and SELENOF were either unchanged or minimally changed by the addition of selenium to the culture media, while SBP1 was undetectable in these cells.

Discussion: Selenoprotein location and levels in RWPE-1 cells closely resemble that seen in human tissues. These cells will therefore be a useful model for further research on the function of selenoproteins in normal prostatic tissue biology and the progression to cancer.
P52 - Glutathione depletion in HEK293T adresses cytosolic GPx4 to the membrane

2. Selenium in the molecular life sciences
2.6 Selenoprotein function
Keywords: PHGPx, glutathione, ferroptosis

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Introduction: Due to its important role in catalysing reactions inhibiting lipid peroxidation, glutathione peroxidase (GPx4) stands out as the key regulator of the cell death pathway named ferroptosis. In order to study the molecular mechanism of GPx4 inactivation by ferroptosis inducing agents, a massive depletion of its reducing substrate glutathione was produced using a thiol alkylating agent, diethyl maleate (DEM), together with an inhibitor of GSH synthesis, buthionine sulfoximine (BSO).

Method: For this study the HEK293T cell line was selected. The cell pellet was lysed, sonicated and centrifuged. Supernatants were used for protein quantification, GPx4 enzymatic activity test and Tietze assay. After SDS-Page, Western blot analysis was performed using an anti-GPx4 antibody.

Result: Treatment of HEK293T cells with BSO and DEM yields a decline in the total glutathione content in cytosol. Furthermore, a marked decrease of GPx4 specific activity is observed already after 3h, which is accompanied with a decline of GPx4 protein, as seen by Western blot. As the treatment with proteosomal inhibitor MG132 fails to prevent this phenomenon, the GPx4 content in BSO/DEM treated cells was analysed in the residual pellet after the centrifugation. Results show that, while disappearing from cytosol, GPx4 appears in the membrane fraction.

Discussion: Thus, upon GSH depletion, GPx4 is addressed to the membranous fraction of the cells. This result complies with both, SPR analysis and in silico study indicating that the affinity of GPx4 for the polar head of membrane phospholipids is increased when GSH is absent.

Selected references


P53 - Synthesis and efficacy of conjugated selenium ADC monoclonal antibodies in vitro

2. Selenium in the molecular life sciences
2.8 Selenium based biotechnological applications

Keywords: Monoclonal Antibodies, Triple Negative Breast Cancer Cell, Treatment

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Introduction: Monoclonal antibodies (mAbs) are known for their high specificity and affinity to target antigens. Yet sometimes mAbs lack therapeutic activity; primary resistance (Pohlmann, 2009; Wilkins, 2010) hence the efficacy of mAbs may be improved by drug conjugation (ADCs).

Method: We describe here the attributes of two selenium (Se) monoclonal ADCs; Se-Herceptin and Se-Avastin. Redox selenium (Se) was covalently attached to commercially available and clinically employed Her2+ and VEGF mAbs, Herceptin\textsuperscript{®} and Avastin\textsuperscript{®}. Selenium attachment to the mAbs is made using a Se-modified Bolton-Hunter reagent (Lane, 2011; Bapat, 2015). The Se-Bolton-Hunter reagent was dissolved in tetrahydrofuran and incubated with Herceptin\textsuperscript{®} or Avastin\textsuperscript{®} in a sodium borate buffer at 4\textdegree{}C for 72 hr and then exhaustively dialyzed against PBS. Control Herceptin\textsuperscript{®} and Avastin\textsuperscript{®} mAbs were treated identically but without selenium attachment; Figure.

Result: After dialysis, Se-ADCs were quantitated using ICP-MS. Chemiluminescence activity of the Se-ADCs was determined for the detection of superoxide generation following the procedures of Chen (Chen, 2008). Protein concentrations of the Se-ADCs were determined by a BCA assay. All mAbs were subjected to PAGE gel electrophoresis under reducing and non-reducing conditions and stained with Coomassie Blue R250.

Discussion: Human Triple Negative Breast Cancer (TNBC) cell lines MDA-MB-468 and MDA-MB-231 ex vivo were subjected to time and dose responses to Herceptin\textsuperscript{®}, Avastin\textsuperscript{®}, and their corresponding Se-ADCs. MDA-MB-468 and MDA-MB-231 cells at all time and concentrations were unaffected by Herceptin\textsuperscript{®} or Avastin\textsuperscript{®} treatments, whereas cytolyis of these TNBC lines were highly responsive to the time and concentrations of the redox (Stewart, 1999) Se-ADC treatments.

Selected references
P54 - Antifungal Activity of Selenium Nanoparticles Synthesized by B. subtilis Against P. syringae

2. Selenium in the molecular life sciences
2.8 Selenium based biotechnological applications

Keywords: Antifungal Activity, Nano Selenium, Biosynthesis, Bacillus subtilis, Pseudomonas syringae

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Introduction: The low toxicity of Se nanoparticles (NPs) and properties such as antibacterial, antiviral, and antioxidant activities have attracted research attention. Physicochemical techniques, biological methods, and the synthesis of nanostructures with bacterial and fungal strains as well as several plant extracts offer novel, clean, non-toxic, and eco-friendly methods for the production of Se NPs.

In the present study, biogenic Se NPs were purified from the whole-cell lysate of Bacillus subtilis Bs-217 and characterized. Then, the antibacterial efficacy of these novel NPs against Pseudomonas syringae pv. actinidiae was studied.

Method: Se NPs were prepared by reducing Se$^{4+}$ ions with the B. subtilis Bs-217, which was previously identified with a 16S ribosomal DNA method. The antibacterial activities of Se NPs against P. syringae pv. actinidiae were determined by using the MIC method.

Result: Se NPs were successfully synthesized with B. subtilis Bs-217 as evidenced by a significant change in the cultivation medium from colorless to insoluble orange-red, indicating the presence of elemental Se (Se$^{0}$), after 24 hours.

The antibacterial activities of the Se NPs against P. syringae pv. actinidiae is shown in Table 1. The measured MICs for P. syringae pv. actinidiae (60 µg/mL) showed that the biogenic Se NPs had good antibacterial activity.

Discussion: Use of NPs against pathogenic bacteria is yet another novel approach. The results of the present study suggest that nanoscale biogenic elemental Se has antibacterial activity against P. syringae pv. actinidiae. The mechanism of this antibacterial effect is unknown and merits further in vivo and in vitro studies.

Selected references

Table 1. Antifungal Activities of Biogenic Selenium Nanoparticles (Se NPs) against P. syringae pv. actinidiae.

<table>
<thead>
<tr>
<th>Strain</th>
<th>Antifungal Activity of Se NPs MIC$^a$</th>
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<tr>
<td>P. syringae</td>
<td>60</td>
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$^a$ MIC, minimum inhibitory concentration.
P55 - Exosome-mediated methylmercury detoxification accelerated by selenium compound, selenoneine in aquatic organism

2. Selenium in the molecular life sciences
2.9 Additional and emerging topics of selenium in molecular life science

Keywords: exosome, methylmercury, selenoneine, zebrafish

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Introduction: Exosomes are extracellular small granule vesicles produced by cells, and might mediate a signal transduction mechanisms to transfer the information between cells and organisms. They contain various molecular constituents, such as proteins, mRNA and miRNA, and transfer molecules from cells through membrane vesicle trafficking. We found exosome-mediated methylmercury (MeHg) excretion pathway accelerated by 2-selenyltrimethylhistidine, selenoneine, in zebrafish embryos. Here, we report evidence that a selenoneine accelerates the excretion of MeHg though exosomal secretion, and dynamics of secreted exosomes following MeHg exposure in zebrafish models.

Method: The embryos (8 hpf) were microinjected with MeHg-Cys (10−200 µg Hg/ml) into yolk sac, and cultured for 16 h. Exosomes were purified by ultracentrifugation at 100,000g for 1-3 h. Exosomal markers (protein content, AChE activity and CD63 content) were measured. To visualize the production of exosomes in vivo, cd63-GFP expressing zebrafish lines were generated.

Result: Exosomes were released by MeHg exposure in a dose dependent manner. Scanning probe and electron microscopic observation revealed that the exosomes were vesicles of 20-50 nm in diameter. Hg (0.05-0.1 ng Hg/embryo) was contained in the exosomes. Selenoneine (3 µM) treatment accelerated exosomal secretion and Hg excretion. Exosomal secretion pathway was accelerated by selenoneine incorporated into cells and played an essential role for methylmercury detoxification. In cd63-GFP transgenic zebrafish lines, the fluorescent signals was observed in central nervous system, yolk sac and somite, and enhanced signals were detected by MeHg injection and selenoneine treatment.

Discussion: Therefore, induced production of exosomes was visualized in living animals, and indicated that selenoneine involve in MeHg detoxification.
P56 - Studying selenoproteome regulation using selenium stable isotope labeling

2. Selenium in the molecular life sciences
2.9 Additional and emerging topics of selenium in molecular life science

Keywords: Stable isotope, ICP-MS, selenoproteins

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Introduction: Selenium (Se) is an essential trace element, which is incorporated as a rare aminoacid, selenocysteine, in twenty five selenoproteins, to constitute the selenoproteome. Selenoprotein family is one of the most important bioactive form of selenium in human health. To understand the function and regulation of human selenoproteome, which is expressed at a trace levels, it appears critical to develop innovative strategies. The use of isotopically enriched selenium also allows cellular labelling and tracing of selenoproteins and other seleno-compounds.

Method: Selenium has a particular isotopic profile with six stable isotope (74Se, 76Se, 77Se, 78Se, 80Se and 82Se) used as a signature in our analysis with ICP-MS or ESI-MS/MS. When selenium is added as pure isotope, its incorporation into selenoproteins can be followed by ICP-MS.

Result: We are able to (i) follow the selective labeling of the cellular selenocompounds with non radioactive Se-enriched isotopes (76Se, 77Se) and (ii) detect the Se-isotopic ratio of the selenoproteins using size-exclusion chromatography followed by inductively coupled plasma mass spectrometry detection (SEC ICP-MS). The reliability of our strategy is further confirmed by western blots with distinct selenoprotein-specific antibodies. Dose response and kinetic experiments of selenium supplementation or exchange have been used to study the hierarchy of selenoproteome regulation.

Discussion: Here, we describe the use of isotopically-enriched forms of Se as an alternative stragegy to radioactive 75Se, for the labeling and tracing of selenoproteins in cultured cell lines.

Selected references

Anne-Laure Bulteau and Laurent Chavatte (2015) Update on selenoprotein biosynthesis, Antioxidants and Redox Signaling, 23, 775-94.


P57 - Progress in selenium nutrition in China during the last thirty years

3. Selenium in animal and human health and disease
3.1 Selenium supplementation for animal and livestock health

Keywords: soil, china, human intake

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Introduction: From the perspective of soil-plant nutrition science, this paper summarized the progress and achievements of Se study in agriculture science in China during the last 30 years.

Method: The literatures from China Academic Journal Network Publishing Database were analyzed using bibliometrics method.

Result: We presented temporal and spatial variations of Se in agricultural soils and its availability, temporal and spatial variations of Se in main crops, the role of Se on crop growth and its influence factors, and development of Se-rich agricultural products. In view of the two sides of Se (nutrition and toxicity), it is put forward that Se effective state and critical threshold research needs to be strengthened to meet the safe use in soil-crop-human body system, and the development of Se related industry standards and quality supervision system should be accelerated to meet the development of Se-enrich products. In this regard, Chinese authorities have made a draft standard of Se-rich agricultural products, and are promoting it as a national standard.

Discussion: Interestingly, although China has made rapid development in its economy during the recent thirty years, the resident daily selenium intake showed decreasing trend. Thus, appropriate supplementary selenium is very necessary.

Selected references
P58 - Protective effect of organic selenium-enriched extract from cardamine violifolia on carbon tetrachloride–induced hepatic damage in mice

3. Selenium in animal and human health and disease
3.1 Selenium supplementation for animal and livestock health

Keywords: Cardamine violifolia, Organic selenium-enriched extract, Liver protection

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Introduction: Cardamine violifolia is a kind of wild plant in western China and found to have strong selenium-enriching ability in World Selenium Capital-Enshi, China. We has extracted a kind of organic selenium-enriched extract containing 4800 µg/g of selenium. The aim of this study was to evaluate the protective effect of organic selenium-enriched extract on carbon tetrachloride-induced hepatic injury in mice.

Method: The mice of each group were given different doses of the extract (calculated in selenium: 60, 180, 540 µg/kg·d) or sodium selenite (calculated in selenium: 180 µg/kg·d) for 15 days. Then the mice were injected intraperitoneally with CCl₄ (0.3% in olive oil, 10 ml/kg of olive oil solution). After 24 hours, the mice were sacrificed to detect liver injury, oxidation and inflammation related index.

Result: The medium (180 µg/kg·d) and high dose (540 µg/kg·d) of the extract significantly alleviated the pathological injuries in liver and significantly decreased the release of ALT and AST from the liver compared with the CCL₄ group. Compared with the CCL₄ group, the activities of SOD and GSH-Px were elevated, and TNF-α, iNOS and COX-2 were reduced in livers from mice treated with the extract. We will further separate some pure selenium compounds to determine the active ingredient.

Discussion: The results suggest that the pre-supplementation of organic selenium-enriched extract from cardamine violifolia alleviate CCl₄-induced acute liver injury, which is likely due to the improvement of antioxidant enzymes and down-regulation of pro-inflammatory mediators. Cardamine violifolia may be a potential healthy organice selenium source.

Selected references
P59 - Comparison of Trace Elements and Oxidant Status in Dairy Cows at Different Physiological Stages

3. Selenium in animal and human health and disease
3.7 Additional and emerging topics of selenium in health or disease

Keywords: Cows, Trace elements, Oxidant Status, Different Physiological Stages

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Introduction: Hardly shall we find specific directions for trace element additives, preventing diseases caused by oxidative stress in cows at different physiological stages for dairy husbandry. The aim of the study is to compare the serum concentrations of trace elements and oxidative status in dairy cows at different physiological stages.

Method: 20 calves (3-4 month old), replacement heifers, prenatal cows (30 days before delivery), postnatal cows (30 days after delivery) and late-lactation cows were randomly selected into 5 groups. Serum concentrations of selenium, zinc, iron, copper and manganese of each group were determined by ICP-MS. The parameters including glutathione peroxidase (GSH-Px), superoxide dismutase (SOD), catalase (CAT) activity and malondialdehyde (MDA) content were measured by commercial ELISA kits.

Result: The results showed that except for zinc, other trace element levels were significant lower in calf group (\(P<0.05\) or \(P<0.01\)) than others. There is no significant difference (\(P>0.05\)) between replacement heifers and adult cows. The level of selenium was significant lower (\(P<0.05\) or \(P<0.01\)) in groups of prenatal and late-lactation cows. Parameters indicative of oxidative status was not related to ages, however the levels of GSH-Px, SOD, CAT activity and MDA were significant higher (\(P<0.05\) or \(P<0.01\)) in prenatal and postnatal cows. However the mechanism of the difference of oxidative status in dairy cows at different physiological stages has been understood poorly.

Discussion: The results above confirmed that the concentrations of trace elements and oxidative status in calves were the lowest, and imbalance of oxidative state was associated with late pregnancy and early lactation.

Selected references
P60 - Relative bioavailability of selenium sources for beef cattle using glutathione peroxidase activity in liver

3. Selenium in animal and human health and disease
3.1 Selenium supplementation for animal and livestock health

Keywords: beef cattle, feedlot, liver, selenium yeast, sodium selenite

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Introduction: Most of Selenium used in animal nutrition is in inorganic form, but organic sources have better bioavailability. To compare the bioavailability of organic and inorganic selenium sources using glutathione peroxidase activity (GPx) in beef cattle liver.

Method: It was used 45 Nellore cattle with 24 months of age and 392 kg live weight, in a feedlot study during 84 days, in individual pens. The animals (nine per treatment) were submitted to one of the five diets: control diet without additional selenium supplementation; control + 0.3 mg Se kg⁻¹ DM in the form of sodium selenite or selenium yeast; control + 0.9 mg Se kg⁻¹ DM in the form of sodium selenite or selenium yeast. The control diet was deficient and had 0.065 mg of Se/kg of DM. Diets had a roughage: concentrate ratio of 30:70, based on corn silage, corn grain and soybean meal. At the end of the experiment the animals were slaughtered and liver samples were taken for glutathione peroxidase activity according Paglia & Valentine (1967). The bioavailability was calculated by technique of slope ratio assay (AMMERMAN et al., 1995). Linear regression was performed with the Proc GLM of SAS to characterize the glutathione peroxidase activity.

Result: The difference between the slopes was significant (P<0.0001), being that estimation with standard errors were: 37.6 ± 16.7 for the sodium selenite and 146.0 ± 16.7 for the yeast source.

Discussion: The relative bioavailability estimated by GPx for the yeast selenium in relation to the sodium selenite (100) was 3.89 or 389%.

Selected references

P61 - Organic Se has better protective effects than inorganic Se against AFB1/OTA-induced immunotoxicity

3. Selenium in animal and human health and disease
3.1 Selenium supplementation for animal and livestock health

Keywords: Selenomethionine, Sodium selenite, Aflatoxin B1, Ochratoxin A, GPx1

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Introduction: Aflatoxin B1 (AFB1) and ochratoxin A (OTA) frequently contaminate a wide variety of food and feeds and decrease immune function of humans and animals. Whether Se alleviates the immune toxicity induced by these mycotoxins and the underlying mechanism remain unclear.

Method: Advanced technologies such as flowcytometry, western blotting, confocal laser, and RNA interference were used in the study.

Result: Our present study demonstrated that AFB1 or OTA induced immunotoxicity in porcine primary splenocytes or porcine alveolar macrophages (PAMs) cell line 3D4/21 respectively in a dose-dependent manner, as demonstrated by decreasing cell viability, and increasing LDH activity, Annexin V-binding, Bcl-2/Bax mRNA ratio, and pro-inflammatory cytokines expression. AFB1 or OTA increased intracellular ROS levels. NAC, a free radical scavenger, alleviated the AFB1 or OTA-induced ROS production and cell toxicity. Further, organic Se (selenomethionine, Se-Met) and inorganic Se (sodium selenite, SS) are supplemented to study their effects on toxicity of AFB1 or OTA. The results showed that both Se sources have significant effects on alleviating AFB1 or OTA -induced immunotoxicity in porcine primary splenocytes or PAMs. Se-Met and SS supplementation significantly improved GPx1, SelS and TR1 expression. However, Se-Met supplementation has a more dramatic effect on alleviating the toxicity of AFB1 and OTA, and up-regulating of GPx1 expressions than SS supplementation. Knockdown of GPx1 by GPx1-specific siRNA diminished the protective effects of SeMet and SS against AFB1 or OTA -induced immunotoxicity.

Discussion: These results suggest that SeMet and SS could alleviates AFB1 and OTA -induced immunotoxicity by improving GPx1 expression, in which organic Se is better than inorganic Se.
P62 - Dietary selenium forms influence selenogenome expression in broiler chickens

3. Selenium in animal and human health and disease
3.1 Selenium supplementation for animal and livestock health

Keywords: broiler chicken, dietary selenium, hydroxy-selenomethionine

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Introduction: Poultry diets can be supplemented with either mineral or organic Se forms (Briens et al., 2013). Thus, this study aimed to compare sodium selenite and hydroxy-selenomethionine on chicken selenoprotein gene expression.

Method: Chickens were fed from 1 to 21 days old three dietary treatments. The dietary treatments correspond to a standard diet either unsupplemented in Se: negative control (NC); NC supplemented with: sodium selenite (SS-0.2) and NC supplemented with hydroxy-selenomethionine (SO-0.2), both at 0.2 mg Se/kg feed. On day 21, eight birds per treatment were selected for breast muscle and liver collection for total RNA extraction and RT q-PCR measures of the 26 chicken selenoprotein genes.

Result: The three dietary treatments were analyzed for endogenous Se content and revealed the following concentrations of Se, NC: 0.13 mg/kg; SS-0.2: 0.32; SO-0.2: 0.32. In the muscle, Selenom expression was significantly increased in Se supplemented groups compared to NC by >1.3 fold. Only SO-0.2 treatment significantly increased gene expression of Gpx1, Gpx3, Txnrd1, Txnrd2, compared to SS-0.2 (1.15 to 1.35 fold). In the liver, only SO-0.2 treatment significantly increased gene expression of Dio2, Dio3, Gpx1, Selenop, compared to SS-0.2 (1.31 to 1.61 fold).

Discussion: Those results indicate that not only the selenium level in the diet but also the chemical form of Se can influence selenoprotein gene expression. Several genes with known redox functions were upregulated when hydroxy-selenomethionine was given. Further studies are needed to measure the benefits of those enhanced selenoprotein expressions on the antioxidant capacities of the animals and their capacity to fight oxidative stress conditions.

Selected references

P63 - Eisenia fetida - a novel organic selenium-biofortified soil animal

3. Selenium in animal and human health and disease
3.1 Selenium supplementation for animal and livestock health

Keywords: Eisenia fetida, organic selenium, speciation, biofortification

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Introduction: Selenium (Se) is a crucial element for human health [1]. Fungi, vegetables and cereals are typical Se-biofortified materials. We here recommend an organic Se-biofortified animal - *Eisenia fetida*, which could enrich kinds of metals efficiently [2] including Se. This study will provide the foundation for further application of the Se-biofortification strategy.

Method: *Eisenia foetida* were put in the mixed-matrix of cow manure and mushroom residue (m:m=1:1, Se (Na₂SeO₃) concentrations were 0 and 40 mg/kg) and bred 45 days. All the simples were acid-digested (HNO₃:HClO₄=8:2, v:v), Se concentration and speciation were determined by AFS and HPLC-UV-HG-AFS respectively.

Result: Se Concentrations are shown in Table 1. In the experimental group, the value reached up to 133 mg/kg, which was about 21 times more than the control group (5.97 mg/kg).
For Se speciation (Table 2.), the proportion of SeMet compound was 54.67% in the experimental group, SeCys₂ and SeMeCys accounted for 42.23%, the percentage of Se⁴⁺ was 1.61%, while an unknown organic Se form was 1.59%, no Se⁶⁺ found. Over all, organic Se accounted for 98.39% of total selenium. This study indicated that *Eisenia foetida* is a good material for Se-biofortification.

Discussion: Yeast (3000 mg Se/kg) is probably the best Se enriched material [3], but *Eisenia foetida* may be a good terrestrial animal for Se-biofortification owing to the Se concentration and organic Se conversion efficiency are extremely high. Se biological effects are highly dependent on organic Se [4], therefore, *Eisenia foetida* would be an excellent source of Se supplement.

Selected references
P64 - Dietary selenium supplies in Malawi

3. Selenium in animal and human health and disease
3.2 Epidemiology of selenium related health and disease

Keywords: Malawi; household survey; spatial; socioeconomic

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Introduction: Dietary micronutrient deficiencies (MNDs) including selenium (Se) are widespread in Malawi causing a large disease burden. Nationally-representative information on the prevalence of dietary deficiencies, with appropriate spatial resolution and stratification by socioeconomic factors, can inform national policies to address MNDs.

Method: This study combined food supply data from the Third Integrated Household Survey of Malawi with locally-generated food crop composition data to derive estimates of dietary Se supply in Malawi.

Result: We estimated that 74% of households in Malawi have inadequate dietary Se supplies. Hidden hunger is likely to be widespread: among households with adequate energy supplies, 56% had inadequate Se supplies. Soil factors are a major underlying factor, with inadequate dietary Se supplies for >80% of rural households living on low-pH soils but 55% living on calcareous soils. Inadequate dietary Se supplies occur alongside inadequate supplies of Ca and Zn in >80% of the poorest rural households living on low-pH soils. An agronomic biofortification strategy could reduce the prevalence of inadequate dietary Se supplies from 82% to 14% of households living in areas with low-pH soils, including from 95% to 21% for the poorest subset of those households, at a cost per alleviated case of dietary Se deficiency of ~ US$ 0.36 year⁻¹.

Discussion: Dietary consumption data from household surveys in conjunction with high resolution crop and geochemical data can provide useful insights into the prevalence and underlying causes of dietary mineral deficiencies, allowing disaggregation by spatial and socioeconomic criteria. Furthermore, impacts of potential interventions can be quantified.

Selected references


P65 - Selenium status in healthy elderly from the Northwest of Algeria

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Introduction: Selenium is a key component of several major metabolic pathways, including thyroid hormone metabolism, antioxidant defense systems, and immune function [1]. Selenium inadequacy is common in older people [1]. The observation that Se status is low in older individuals suggests that requirements may increase with age [1]. The purpose of this study is to determine plasma Se level and dietary selenium intake of 80 healthy elderly subjects living in Tlemcen city situated in Northwest of Algeria. This group is over 60 years.

Method: Selenium was measured using differential pulse cathodic stripping voltammetry. Data of selenium intake were collected by weighted 3-days diet records. The results are compared by t test with healthy individuals who are between 20-50 years of age.

Result: The mean of plasma selenium concentrations is 68.10±20.10 µg/L. This concentration doesn’t vary in relation to the gender of the subject (67.60±20.42 µg/L in women versus 69.70±19.30 µg/L in men, P>0.05) but is significantly lower than Algerians 20-50 years old (78.65±20.80 µg/L, P<0.05).

The mean of Se dietary intake is 45.21±42 µg/day. It is not significantly different from that of younger individuals (48.12±12 µg/day).

Discussion: Similar results were observed in men who took part in the National Health and Nutrition Examination Survey (NHANES III) carried out in the United States [2]. Selenium level observed in healthy elderly of Northwest Algeria is lower than that of younger subjects. All groups consume less Se than the U.S. Recommended Dietary Allowance values [3].

Selected references

P66 - Association between Selenium Levels and Antioxidant Capacity in An Elderly Chinese Population

Introduction: Selenium (Se) has been hypothesized to prevent geriatric diseases for the antioxidant properties of selenoproteins. However, selenoprotein levels and the antioxidant capacity of elderly population were rarely measured in published studies. This study aims to explore the association between Se levels and the antioxidant capacity of an elderly Chinese population with life-long natural selenium exposure.

Method: A cross-sectional study of 433 elderly people aged 65 and older (72.4±5.3a) from four rural counties in Enshi with diverse environmental Se levels was conducted. Whole blood total Se, serum Selenoprotein P1 (SEPP1) and Malondialdehyde (MDA) concentration, serum Glutathione peroxidase (GPx) and Superoxide dismutase (SOD) activity were measured. ANCOVA models and logistic regression models were used to examine the relationships.

Result: The mean measurements for the whole blood Se, SEPP1, GPx, SOD, MDA were 0.159µg/L, 60.67pg/ml, 211.36µmol/L, 50.68 U/ml, 4.24 nmol/ml, respectively. Significant associations between whole blood Se and SOD, whole blood Se and SEPP1 were observed adjusting for age, gender, education, alcohol consumption and smoking. Although not statistically significant, a negative association between SEPP1 and MDA was observed. The mean SEPP1 and GPx were higher and MDA was lower in higher blood Se quintile groups. No significant difference in SOD was observed among five blood Se quintile groups.

Discussion: Higher whole blood Se levels were associated with higher selenoprotein levels and lower lipid peroxidation levels in the elderly population. Selenoproteins should be measured in future studies concerning on Se and geriatric diseases.

Selected references
P67 - Selenium and Alzheimer’s disease: facts and effects

3. Selenium in animal and human health and disease
3.3 Nutritional selenium intervention studies in human

Keywords: Alzheimer's disease, neurodegeneration, cognitive decline

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Introduction: Deficient status of antioxidants has been associated with cognitive decline and risk for dementia.

Method: We first aimed to associate selenium status and cognitive performance in Alzheimer’s disease (AD) and mildly cognitive impaired (MCI) people in two different populations (Brazilian and Australian). Moreover, we conduct two independent pilot trials that used different selenium sources for 6 months aiming to improve selenium status and cognition: i) a Brazilian population with MCI received one Brazil nut, the richest selenium food source, daily (~288 µg Se/d); ii) an Australian AD group received 10 mg of sodium selenate. In both studies, a control (non-treated) group was included.

Result: More than 90% of Brazilians were selenium deficient, while in Australian group only one individual did not present sufficient selenium status. AD Brazilian patients presented lower plasma and erythrocyte selenium levels when compared with healthy controls, and in the Australian population only erythrocytes were statistically lower in AD group. Such discrepancy between these populations raises the discussion about the best biomarkers for selenium. MCI elderly who received Brazil nuts improved cognition performance measured by constructional praxis and verbal fluency tests. Moreover, AD patients treated with sodium selenate presented higher selenium concentration in CSF, which was associated with better performance on Mini Mental State Exam.

Discussion: The discrepancy between Brazilian and Australian populations regarding selenium status raises the discussion about the best biomarkers for selenium. Additionally, our data shows the relevance of selenium as strategy to slow the onset of dementia, and encourages larger studies that consider baseline selenium status.

Selected references


P68 - Selenomethionine improves synaptic deficit in an Alzheimer’s disease mouse model

3. Selenium in animal and human health and disease
3.1 Selenium supplementation for animal and livestock health
Keywords: Alzheimer’s disease, Synaptic dysfunction, Synaptic plasticity, Dendritic spine

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Introduction: Alzheimer’s disease (AD) is the most common type of dementia in the aged people. Synaptic dysfunction induced by the accumulation of extracellular senile plaques (SPs) and intracellular neurofibrillary tangles (NFTs) is believed to be the basis for memory loss in early stages of AD. Studies have showed that the increased basal intracellular Ca$^{2+}$ concentration can block the neuronal signaling and impair the generation and maturation of dendrite spines which directly regulate the synaptic plasticity, thereby leading to the decline in learning and memory in AD. Selenomethionine (Se-Met), a major form of dietary selenium (Se) with powerful antioxidant capacity, reduced Aβ deposition and tau hyperphosphorylation, improved synaptic proteins levels and ameliorated cognitive deficits in the triple transgenic AD mice. However, the specific mechanism underlying the synapse- or memory-improving effects of Se-Met remains unknown.

Method: In the present study, a 3xTg-AD mouse model with YFP labeled neurons was used to dynamically investigate the effect of Se-Met on dendritic spine formation in vivo.

Result: We found 3-month treatment with Se-Met lead to a rapid outgrowth of dendritic protrusions and an increase in spine generation in AD mice brain. Meanwhile, Se-Met significantly attenuated the basal intracellular Ca$^{2+}$ concentrations and stimulated long-term potentiation (LTP) between perforant path and DG circuit. Se-Met treatment also decreased the levels of NMDAR, Fyn and Src and activated ERKs and CREB.

Discussion: Our results demonstrated for the first time that Se-Met improved the synaptic function and neurons signal transmission, and maintained the stability of dendritic spines in AD mice.

Selected references
P69 - Effect of selenium supplementation in Hungarian patients with autoimmune thyroiditis and endocrine orbitopathy

Keywords: autoimmune thyroiditis, orbitopathy

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Introduction: In the last decade selenium (Se) supplementation has been adopted as an adjuvant treatment for patients with autoimmune thyroiditis (AIT) or in patients with mild endocrine orbitopathy. The dosage and length of treatment with Se is still a matter of debate. The aim of this small study was to follow up patients, to evaluate the effects of treatment with Se, and to register possible side effects.

Method: Altogether 150 patients with autoimmune thyroiditis and 20 patients with orbitopathy participated in the study. The patients received 100 or 200 µg organic Se daily depending on the activity of the autoimmune process. The relevant disease markers were determined before the beginning of treatment and at regular check-up appointments.

Result: Patients with orbitopathy benefitted from treatment with Se as symptoms and clinical activity scores improved significantly. The effect of Se treatment in patients with autoimmune thyroiditis was less evident. Elevated thyroglobulin antibody levels generally decreased significantly. Anti-thyroid peroxidase antibodies decreased in less than 40% of patients. In case of treatment with selenium-enriched yeast, serum Se concentrations exceeded the reference range in less than the recommended treatment time of six months. No side effects were registered. The patients did not develop type 2 diabetes mellitus during treatment.

Discussion: Treatment with Se is effective in Hungarian patients with endocrine orbitopathy. In patients with autoimmune thyroiditis, Se may contribute to a decrease in thyroglobulin antibody titers but elevated anti-thyroid peroxidase antibodies may not improve at all, the cause of which remains unclear and should prompt further investigations.
P70 - Selenium supplementation as a defense method against posttraumatic stress disorder development in combats

3. Selenium in animal and human health and disease
3.3 Nutritional selenium intervention studies in human

Keywords: Posttraumatic Stress Disorder, Oxidative Stress, Antioxidants, Combats

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Introduction: The combats of Contingent of the International Operations (CIO) are a subject of various extreme factors action, which can cause Posttraumatic Stress Disorder (PTSD). Oxidative stress (OS) level considerably changes because of reactive oxygen species excess accumulation. Neuronal membranes phospholipids are especially vulnerable to damage, the injury leads to the receptor-mediated signal transduction and information processing disorders. Selenium (Se) is recognized to be a potentially protective factor because of scavenging endogenous and environmental oxidant sources. Study aim: to assess PTSD, OS and Se levels, their correlation in CIO, defense by Se ordination efficacy.

Method: Prospective study. Totally 143 participants in patients and control groups: Latvian CIO, regular personnel, males, Europeans, average age of 27.4, before and after Peace Support Mission (PSM) in Afghanistan examined. Questionnaires PCL-M, the valid Latvian language version were used for PTSD prevalence rate (PR) evaluation. Neuronal tissue specific OS indicator - Intensity of lipid peroxidation - Malondialdehyde (MDA) and Se in plasma (SeP) were determined. Patient group received 200 mcg of Se per day during mission.

Result: Before PSM: response rate (RR) 0.98, PR 0.04, MDA level 2.56 µM, SeP 85.25 µg/L. After PSM - patients: RR 0.94, PR 0.05, MDA 2.41 µM, SeP 103.24 µg/L. After PSM - controls: RR 0.92, PR 0.09, MDA 3.18 µM, SeP 88.79 µg/L. There is positive correlation between increase of SeP, OS and PTSD in CIO.

Discussion: PTSD PR after PSM reduced for 46.03% in patients compared with controls. Se ordination is the effective defense method against both OS and PTSD development.
P71 - Selenium as a mercury antidote

3. Selenium in animal and human health and disease
3.3 Nutritional selenium intervention studies in human

Keywords: SelenoPrecise, Mercury, heavy metals

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Introduction: There is a complex biochemical relationship between selenium and mercury and other heavy metals. Because the bonding is stronger than a similar sulfur bond, selenium seems to be able to extract mercury – including the methylmercury ion - from protein structures in the cell (A-402). Selenium binds to mercury in an inert selenid complex, that prevents the heavy metal damage and the normal toxicity symptoms are not observed (A-402). In addition, selenium in adequate amounts is metabolized to about 35 different selenoenzymes that protect different cellular structures or vital biological transformations (A-6149). If the diet contains inadequate amounts of selenium, the detoxification only operates incomplete. This would increase the risk of chronic disease by weakening the immune response further (A-402).

Method: A well-controlled, human study examined this relationship in 103 persons exposed to long-term heavy metal pollution in the Wanshan mercury mining area in China. Half of the persons received 100 mcg selenium yeast (SelenoPrecise, Pharma Nord, Denmark) per day for 3 months, while the remaining half received placebo.

Result: Compared to placebo, urinary excretion rate of mercury increased significantly by about 150% after 30 days until the 90 days end of the study period. Similarly biochemical markers supported the main observation of a mercury reduction in the body (A-9287).

Discussion: The current evidence shows that increased intakes of selenium relative to mercury have at least three beneficial effects; binding bio-active mercury in an inert complex, increasing the protection from toxicity through selenoenzymes, and increasing the urinary excretion rate of mercury.

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Li YF, Dong Z, CHEN C, LI B, et al.: Organic selenium supplementation increases mercury excretion and decreasesoxidative damage in long-term mercury-exposed residents from Wanshan, China; Environ Sci Technol 2012 46(20), 11313-8
P72 - Gender specific differences in urinary level of 8-oxodG in selenium supplemented subjects

3. Selenium in animal and human health and disease
3.3 Nutritional selenium intervention studies in human

Keywords: selenium supplementation, gender, 8-oxodG

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Introduction: It is supposed that biological effects of selenium (Se) supplementation as well as its association with risk of cancer, are sex-specific. However, this issue has not been widely investigated. In this study we aimed to analyze gender differences in the urinary level of 8-oxo-7,8-dihydro-2'-deoxyguanosine (8-oxodG) in the Se supplemented individuals. 8-oxodG is a predominant form of oxidative lesion in DNA and its urinary level is a common biomarker of oxidative stress and carcinogenesis.

Method: 52 individuals (26 males and 26 females) were supplemented with 200 mcg of Se/day (as Se yeast) for six weeks. Blood and urine were collected at baseline, after 2 weeks/6 weeks of supplementation and 4 weeks after washout. Urinary concentration of 8-oxodG was analyzed using high-performance liquid chromatography with tandem mass detection (HPLC-MS/MS). We determined also two markers of Se status: plasma Se concentration (using atomic absorption spectroscopy) and glutathione peroxidase 1 (GPx1) activity in erythrocytes (by method of Paglia and Valentine). Differences with respect to time and sex were analyzed using two-way repeated measures ANOVA.

Result: At baseline, males and females were not different in terms of plasma Se and urinary 8-oxodG, however females had significantly higher baseline GPx1 activity as compared to males. Significant sex-specific differences were observed upon Se supplementation in 8-oxodG concentration ($p$ for time*sex interaction = 0.008). The marker was increased by 69% in males and decreased by 27% in females after 6 weeks of supplementation.

Discussion: This study supports the modifying role of sex in determining oxidative stress response to Se supplementation.
P73 - Cystine-glutamate antiporter expression as a potential target for cancer therapy using redox active selenium compounds

3. Selenium in animal and human health and disease
3.4 Selenium based medical therapeutics
Keywords: Drug resistance, Cancer chemotherapy, Selenium compounds, Cystine-glutamate antiporter

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Introduction: One of the major drawbacks in cancer treatment is the development of resistance against cytostatic drugs. Chemotherapy resistance has been positively correlated with overexpression of the antiporter system xCT, cystine-glutamate transporter (gene name: SLC7A11). This transporter plays a key role in protecting cells against oxidative stress via uptake of cystine. However, xCT overexpression is associated with enhanced sensitivity towards certain selenium compounds. In the present study, we aim at investigating the involvement of xCT antiporter in mediating cytotoxicity of selected redox-active selenium compounds.

Method: We used multiple cancer cell lines in which ectopic overexpression and siRNA-mediated knock-down of xCT were performed and validated. Additionally, we adopted chemical inhibition approach to test the functional roles of this transporter. Western blot and qPCR were used to quantitate the expression of xCT.

Result: The basal levels of xCT expression in the tested cancer cell lines correlated well with the sensitivity towards the tested selenium compounds. When we used siRNA targeted against xCT, cytotoxic effects of these selenium compounds were markedly inhibited. Conversely, ectopic overexpression of xCT resulted in dramatic increase in cytotoxicity, especially in those cells that were resistant to the tested selenium compounds. Small molecule-mediated inhibition of xCT transporter activity resulted in loss of sensitivity towards these selenium compounds.

Discussion: Our results demonstrate the functional role of xCT in mediating cytotoxicity of the studied selenium compounds. These findings suggest the potential cancer chemotherapeutic applications of selected selenium compounds, specifically in xCT over-expressing tumors.
P74 - Finding What’s Important In Selenium Drug Development

3. Selenium in animal and human health and disease
3.4 Selenium based medical therapeutics

Keywords: ebselen, LC-ICP-MS, ototoxicity, hearing, Meniere's

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Introduction: Ebselen, a seleno-organic drug substance (DS), is being evaluated under four investigational new drug (IND) applications: prevention of noise-induced hearing loss, treatment of Meniere’s disease, and prevention of chemotherapy- or aminoglycoside-induced ototoxicity. Comprising approximately 29% of ebselen by weight, Se may also exist in various inorganic species, e.g., selenate and selenite, considered impurities in DS and drug product (DP) (ebselen, 200 mg, SPI-1005 capsules). Tentative acceptance criteria and a novel method of control were justified to the U.S. Food and Drug Administration (FDA) in the INDs.

Method: Pharmaceutical regulatory guidance (ICH Q3D) addresses elemental impurities in general, however, analytical methodology and limits must be justified by Sponsor to FDA. A validated liquid chromatography-inductively coupled plasma-mass spectrometry (LC-ICP-MS) method was developed and validated to quantitate Se species in DS and DP.

Result: An LC-ICP-MS method was developed (Limit of detection (LOD)=0.2 ppm) to quantitate Se in DS and DP. Levels of inorganic Se (up to 1.3 ppm selenite and no selenate in several lots of DS; up to 0.7 µg selenite/capsule in early lots, but no selenate or selenite in recent lots of DP) were low or below the LOD, with no significant changes in DP over 18 months stability storage (as per ICH). Inorganic Se < 10 ppm in DS ensures < 2 µg/SPI-1005 capsule.

Discussion: Control strategy and tentative acceptance criterion of inorganic Se < 2 µg/SPI-1005 capsule were agreed upon with FDA. A novel LC-ICP-MS method was developed, validated and found sufficient for this clinical purpose.
P75 - Seleno heterocyclic compounds as antitumoral and radical scavenging agents

3. Selenium in animal and human health and disease
3.4 Selenium based medical therapeutics

Keywords: Heterocycle, cancer, radical scavenging

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Introduction: Adequate levels of selenium (Se) have been associated with low overall mortality. Accordingly, heterocyclic compounds with Se atom in endo position, such as ebselen (EBS), have achieved very promising activity against cancer and excellent pharmacological profile, currently being under Phase II clinical trial for the evaluation of its therapeutic safety and efficacy (1). We decided to make structural modifications over this scaffold with the aim of improving its characteristics (2).

Method: Twenty-eight novel benzo[c][1,2,5]selenodiazole-5-carboxamide derivatives were designed and synthesized. Cytotoxicity of these structures was tested in vitro against a panel of seven human cancer and non-malignant cell lines (PC-3, HT-29, CCRF-CEM, HTB-54, MCF-7, 184B5 and BEAS-2B) by the MTT assay. For MCF-7 cells, the apoptotic status and cell cycle analysis of the cells were based on the TUNEL technique. Likewise, their radical scavenging activity was determined using the DPPH assay.

Result: The GI₅₀ values for every tested product were lower than EBS. The cytotoxic activity of compound 8 was remarkable in MCF-7 cells. However, the induction of cell death was independent of the apoptotic process and do not either effect the cell cycle progression. The radical scavenging capacity of compounds 1, 2, 9 and 10 was greater than EBS.

Discussion: The toxicity of 8 on the nontumoral line 184B5 was low and it exhibited promising antiproliferative values in MCF-7 cells. Four compounds showed potent radical scavenging activity.

Selected references

P76 - Selenocyanate and diselenide amides: A new class of potent antichagasic agents
3. Selenium in animal and human health and disease
3.4 Selenium based medical therapeutics
Keywords: Selenocyanate, diselenide, Chagas

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Introduction: American trypanosomiasis or Chagas disease is an infectious and neglected tropical disease caused by different parasite species of Trypanosoma cruzi. Over six million people are infected around the world. The chemotherapy drugs used nowadays are not effective enough for the treatment of chronic patients. Therefore, there is an urgent need to develop more effective, safe and affordable drugs. (1, 2)

Method: In this work, 28 novel amides containing selenocyanate and diselenide moieties have been synthesized. Compounds have been tested in vitro in three different epimastigotes strains (SN3, Thulawen and Arequipa) and in Vero cells afterwards in order to determine their effectiveness and toxicity. After these preliminary studies and considering the results obtained, two compounds were selected as the most promising structures. The lead compounds were tested in both amastigote and tripomastigote forms in three strains.

Result: On the basis of their potent activity and high selectivity index (<40), these two compounds were selected for further in vivo evaluation

Selected references

P77 - Cytotoxicity, oxidative stress and antioxidant enzyme activity in pancreatic cancer cells treated with organic selenium compounds

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Introduction: Pancreatic cancer (PC) has a poor prognosis and limited chemotherapeutic efficacy. Recently, a key role for K-ras mutation has been identified in PC carcinogenesis, ultimately favouring glycolysis and altering the redox environment. Although cancer cells adapt to regulate oxidative stress, this capacity is considered to be maximised. Since supranutritional selenium is a potent pro-oxidant, targeting antioxidant pathways with organic selenium (Seleno-L-cystine, methylseleninic acid and Se-(methyl)selenocysteine) is a promising therapeutic approach.

Method: All experiments were performed in parallel in PSN1 (K-ras mutated PC) and BxPC3(K-ras wildtype PC) cells. Cell viability was assayed by both MTT and SRB. Cellular selenium uptake was examined by GF-AAS. ROS production and mitochondrial membrane potential assays were performed. Antioxidant enzyme activity was determined in vitro and in cell lysates to measure the direct effects of and cellular responses to selenium treatment, respectively.

Result: As determined by MTT, all compounds displayed IC$_{50}$ values >10x lower in PSN1 than in BxPC3 cells, correlating with cellular selenium uptake. Conversely, comparable IC$_{50}$ values across cell lines were determined by SRB. Both ROS production and mitochondrial membrane depolarisation was higher in PSN1 than in BxPC3 cells. Congruently, the induction of antioxidant enzyme activity was substantially higher in BxPC3 than PSN1 cells.

Discussion: The higher susceptibility of PSN1 than BxPC3 cells is likely explained by their already maximised antioxidant defense, rendering them incapable of combating selenium induced stress. The different redox environments across cells is partly related to K-ras mutation, which is an important consideration given the high prevalence of K-ras mutation in PC.

Selected references

P78 - Altered Chemoresistance in an Ex Vivo Pancreatic Cancer Model Compared to In Vitro Cell Culture

3. Selenium in animal and human health and disease
3.4 Selenium based medical therapeutics
Keywords: pancreas, cancer, ex vivo, selenocompounds, spheroid

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Introduction: In vitro cell culture models are well-established and versatile tools in drug testing. However, their limitations make experimental results difficult to translate to humans. Thus, the development of new experimental models mimicking an in vivo setting is of greatest importance within translational research. In this study, we set out to compare toxicity of selenium compounds and chemotherapeutics in a 2D, 3D and an ex vivo organotypic model of pancreas cancer.

Method: Monolayer culture of PANC-1, a human pancreatic carcinoma of ductal origin, was used as the cognate 2D model. The formation of 3D spheroids was aided by the addition of methylcellulose in the media of the PANC-1 cells. Pancreatic tumor tissues were sliced using a vibrating microtome and cultured for up to 7 days. Drug-induced changes in viability of the cells and tissue were measured and compared.

Result: The 3D cell culture model and the ex vivo pancreatic tumor tissue model showed decreased sensitivity to various selenium compounds as well as to conventional chemotherapeutic drugs when compared to the 2D model.

Discussion: Consideration of the complex environment of a cancerous tissue is essential to implement successful drug efficacy screening. Development and the use of an ex vivo model of pancreatic tumor tissue would allow for better drug screening as it closely resembles the in vivo tissue architecture, with increased chemoresistance compared to customary in vitro monolayer cell culture models.
P79 - Novel selenocyanate and diselenide phosphoramides as potent anticarcinogenic agents

3. Selenium in animal and human health and disease
3.4 Selenium based medical therapeutics
Keywords: Selenocyanate, diselenide, cancer

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Introduction: Cancer is one of the leading causes of death worldwide and the efficacy of current chemotherapeutic agents is limited by the development of tumor resistance and the appearance of undesired side effects (1, 2). In the last years, our group has demonstrated the potent activity of selenium containing compounds against this illness. (3)

Method: In this work, twelve novel phosphoramides containing selenocyanate and diselenide moieties have been synthesized. Compounds were screened at two concentrations against three human cancer cells lines (MCF-7, HT-29 and CCRF-CEM). Compounds showing GI50 values below 10 µM, where tested to evaluate dose response activity against five human cancer cells lines (MCF-7, HT-29, PC-3, HTB54 and CRF-CEM) and a normal human mammary and lung epithelial cell line (184B5 and BEAS-2B) in order to determine their activity and selectivity against malignant cells.

Result: Four compounds showed GI50 values below 10 µM in more than one cancer cell line and two of them exhibited a selectivity index higher than 6. On the basis of their potent activity and high selectivity index, one of the compounds was selected for further biological evaluation of cycle arrest in MCF-7 and CCRF-CEM cell lines and apoptosis mechanism.

Selected references
P80 - Inducing Change in Selenium Drug Development

3. Selenium in animal and human health and disease

3.4 Selenium based medical therapeutics

Keywords: cochlea, ebselen, hearing, microarray, selenoprotein

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Introduction: Ebselen, an organo-selenium drug, mimics activity of selenoprotein isoforms of Glutathione Peroxidase (GPx) and increase cochlear levels of selenoprotein GPx1, is being evaluated under four investigational new drug (IND) applications, for prevention of sensorineural hearing loss, treatment of Meniere’s disease and prevention of chemotherapy- or aminoglycoside-induced ototoxicity. To understand effects on gene expression in the ear, we performed micro-array analysis with an emphasis on expression changes for known selenoproteins.

Method: Three-month-old F-344 rats with normal hearing by ABR, were noise exposed to octave band noise (OBN) at 115 dB for 4 hours which caused temporary and permanent hearing loss. Ebselen (8mg/kg) or vehicle was orally delivered for 3 or 14-days, beginning 1 day pre-noise. Cochlear RNA was extracted at 1-hr, 1-day, 1-week and 1-month post-noise. Total RNA was labeled and hybridized to Affymetrix microarrays. Data were analyzed using Genesifter software. Three cochlea for each time and condition were analyzed for temporal and conditional change in gene expression following noise exposure in the presence or absence of the electrophilic Nrf2 inducer ebselen (Sakurai, et al., 2006, Kil, et al., 2007).

Result: Spiral ligament and stria vascularis were most affected structures within noise-exposed cochlea and had highest level of GPx1 protein. Changes in mRNA expression at 1 hour post-noise returned to near baseline within 24 hours in vehicle- or ebselen-treated rats exposed to OBN.

Discussion: Microarray analysis of rat cochlear RNA provided a sensitive measure of environmentally-induced changes in gene expression and can be used to provide new insights into functional mechanisms of otoprotective agents.

Selected references


Ebselen Kil J, Pierce C, Tran H, Gu R, Lynch ED. Hear Res. 2007 Apr;226(1-2):44-51. PMID: 17030476
P81 - Metabolic syndrome induced oxidative stress and skin premature ageing beyond topical use of seleno-L-methionine.

Introduction: Metabolic syndrome (MetS) is a combination of changes linked to disruption of metabolism, it also features a latent inflammation that activates peroxidation. Accumulation of lipid peroxidation and oxidative lipoprotein and lipid layer damage lead to morphologic changes of skin structural components.

Method: The prospective controlled study of random, age- and sex-stratified population sample was conducted during 2015-2016 at the Department of Internal Medicine, Riga Stradiņš university, Latvia. Detailed examination of the skin for the presence of MetS associated skin changes was performed. Serum MDA and SOD were determined. A specific cream composition, which contains seleno-L-methionine was being applied in thin layer 2 times per day (in the morning and in the evening) on clean facial skin, neck and decollete areas. Clinical evaluation was repeated in 14, 30 and 60 days from the start of cream use.

Result: Statistical analysis (SPSS 20.0 for Windows) showed a direct linear relationship (Positive Pearson correlation, Chi-square test, p <0.05) between skin alterations and intensity of MDA and SOD. In 60 days improvement of the barrier function of the skin was ascertained.

Discussion: Topical use of a specific cream composition, which contains seleno-L-methionine improves turgor and hydration of the skin, restores elasticity and reduces the phenomenon of pigmentaion, inflammatory papilloma.

Selected references


P82 - Whole Blood Selenium Levels and Selenium Supplementation in Patients treated in a Family Practice

3. Selenium in animal and human health and disease
3.4 Selenium based medical therapeutics

Keywords: Whole Blood Selenium Levels, Selenium Supplementation, Family Practice

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Introduction: Se supply varies widely in Germany. Therefore, a laboratory study was carried out in patients treated in a family practice in Brandenburg (Germany). We examined whether there is a general Se deficiency in this area. Specifically, whether Se concentrations differ with age, sex or cancer. Moreover, we tested the effects of a Se supplementation on whole blood Se levels (WBSL).

Method: WBSL were analyzed in 871 patients (496 female, 395 male, median age: 67 years) in the time-period of 2006 to 2013. 143 of them (78 female, 65 male) were cancer patients in the aftercare situation. 317 patients (76 with tumors, 241 without tumors) received continuous Se supplementation with sodium selenite (300 µg per day). WBSL were compared by student's T-test for paired and independent samples.

Result: The initial WBSL of all patients was 97.2 +/- 20.7 µg/l (mean +/- SD). WBSL did not differ with regard to age and sex, but cancer patients had the lowest WBSL (Table 1). Se-supplementation increased mean WBSL both in cancer patients (to 128.5 µg/l) and in patients without tumors (to 119.52 µg/l) (p <0.001).

Discussion: Tumor patients displayed significantly lower WBSL than non-tumor patients, indicating some negative effect of tumors on Se intake, absorption or metabolism. Significant influences of age or sex were not observed. As selenite supplementation was efficiently improving the WBSL to concentrations considered to provide health benefits and protecting from Se-deficiency associated health risk, we consider this measure as a meaningful adjuvant therapeutic option in subjects with proven low WBSL.
P83 - Influences of selenium status on thyroid response of children after iodine repletion

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease

Keywords: thyroid metabolism, children, iodine deficiency

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Introduction: Selenium is integral part of selenoproteins important for thyroid metabolism. The influence of selenium status on thyroid response of previously iodine deficient Ethiopian children, 15 months after iodized salt became available, was investigated.

Method: A total of 624 children at baseline and 555 children at endline participated in the study. Inductively Coupled Plasma Mass Spectrometer for serum selenium and a clinical analyzer for thyroid markers analysis in serum-triiodothyronine (T3), thyroxin (T4), thyroid stimulating hormone (TSH), and thyroglobulin were used. Pearson correlation to study relationships between serum selenium and thyroid markers, and independent t-test to compare thyroid markers by selenium status were used.

Result: The mean age of children at baseline was 56.9±1.8 mo (54-60). There was a significant increase in median urinary iodine at endline (167.1µg/l vs 9.3µg/l; p<0.001). Selenium deficiency was prevalent in 57.8% of children. The participants had median serum selenium concentration of 61.4µg/l at baseline and 70.6µg/l at end line (p=0.07). Selenium concentration was significantly correlated (r=-0.22, p <0.01) with T4 at baseline and with T3 (r=0.38, p<0.001), Td (r=0.15, p<0.001) and TSH (r=-0.205, p<0.001) at endline. Compared to selenium adequate children, children with selenium deficiency had higher T4 (p<0.001) at baseline but lower T4 (p<0.001) and T3 (p<0.001) concentration at endline. At endline, compared to children with normal selenium status, selenium deficient children had higher TSH concentration (p=0.002). At endline, a significant increase in serum T3 in selenium adequate (p=0.004) but not in selenium deficient children (p=0.208) was observed.

Discussion: Selenium deficiency lowers T3 concentration in iodine replete children.
P84 - Thiol-dependent redox systems as antimicrobial drug targets

3. Selenium in animal and human health and disease

3.4 Selenium based medical therapeutics

Keywords: thiol, selenoprotein, bacteria, parasite, thioredoxin

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Introduction: It is urgent to discover new antibiotics and antibiotic principles because of the occurrence multi-drug resistant bacteria and parasite. The difference between the mammalian host and the bacteria or parasite in thiol-dependent antioxidant enzymes emerges as the selective target. Mammalian cells remove ROS very efficiently with various selenoproteins in the thioredoxin and glutathione system (thioredoxin reductases and glutathione peroxidases). In contrast, bacteria and parasites do not possess selenoproteins in their thiol-dependent redox systems. In particular, some infection pathogens exert specific thiol-dependent antioxidant activities. For example, GSH is absent in pathogens such as H. pylori, S. aureus or M. tuberculosis and thus thioredoxin system become an essential unique thiol-dependent pathway for these bacteria. Thioredoxin reductase and glutathione reductase are absent in Trypanosoma brucei and the pathogen employ special trypanothione pathway for the thiol dependent electron transfer. These properties give us an opportunity to treat these pathogen-caused human diseases by targeting microbial thiol-dependent redox system.

Method: We have found that ebselen an organoselenium compound and its sulfur analog ebsulfur have been shown to be promising antimicrobial drug candidates for this antimicrobial concept.

Result: Ebselen acts as a specific inhibitor of bacterial TrxR, whereas it is substrate for the corresponding human enzyme. Bacteria which are bacteria lacking GSH, are susceptible to ebselen. Ebsulfur formed a complex with TryR and caused oxidation and inactivation of the enzyme. Inhibition of TryR produced more intracellular ROS and enhanced by the presence of ROS.

Discussion: These results display a novel antimicrobial strategy to fight with multidrug resistant bacteria and parasites.
P85 - Novel acylselenoureas as antiproliferative agents: design, synthesis and biological evaluation

Keywords: acylselenoureas, cancer, synthesis

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Introduction: Cancer can be categorized as a serious clinical problem as it affects millions of patients worldwide. Only in the United States of America statistics show death rate by cancer will reach the alarming ratio of 1,600 deaths per day1. Our recent reports have proven potent antitumor activity of several moieties containing selenium such as selenocyanate, diselenide and selenourea2.

We have now designed a series of compounds containing both selenourea and diselenide moieties. Structure endings vary from substituted aromatic rings to heterocyclic scaffolds.

Method: The in vitro effects on cell viability was assessed using a colorimetric MTT assay at 72 h in six human cancer cell lines. In order to determine their selectivity index (SI) the assay was also performed on two cell lines derived from non-malignant cells. As a preliminary approach to the mechanism of action, cell cycle distribution and cell death status was studied by flow cytometry by the TUNEL technique.

Result: Fifteen new derivatives were synthesized. Seven compounds exhibited GI50 values under 10 µM in CCRF-CEM, MCF-7 and PC-3. Six structures are highly selective for breast adenocarcinoma with SI values up to 120-fold higher than those exhibited by doxorubicin.

Discussion: Regarding affection to cell cycle distribution, arrest in G2/M was induced by the leader structures in a time and dose dependent manner. Both derivatives increased the percentages of subdiploid cells in a dose- and time-dependent manner in the TUNEL assay.

Selected references
P86 - Role of selenium nanoparticles to dampen the metastatic potential of aggressive cancers

3. Selenium in animal and human health and disease
3.4 Selenium based medical therapeutics

Keywords: Ovarian and prostate cancers cells, Nanoparticles, AFM, Synchrotron X-ray fluorescence microscopy

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Introduction: Selenium is a trace element that is reported to be efficient to prevent transformation of normal cells to malignant cells and could be an effective chemopreventive and chemotherapeutic agent. Se-bioavailability and toxicity (related to its chemical species) limit Se-use as a chemotherapeutic agent. It appears that Se-nanoparticles (Se-NPs) present interesting characteristics such as a higher bioavailability, an excellent low toxicity (towards non-cancerous cells) and anti-proliferative properties. Se-NPs might have potential applications for the management of human cancers and could be used as potential therapeutic-agent and drug-carrier.

Method: Specific human cell lines (OV-90 and TOV-112D (ovarian), PC3 and LNCap clone FGC (Prostate)) with different metastatic potentials are exposed to different types of Se-NPs (either coated with bovine-serum-albumin or chitosan). Cells are cultured in RPMI medium and further exposed or not to ~ 25 µg/mL BSA/chitosan stabilized Se-NPs (~40nm-diameter) for 24h to 48h.

Result: Using synchrotron X-ray spectroscopic methods (X-ray fluorescence nanoimaging/HERFD-XAS) and optical-fluorescence microscopy, detailed quantitative information are collected on Se-NPs intracellular distributions and on their chemical form to unveil targeted their intracellular compartmentalization and biotransformation. These information are vital to investigate the in-situ cellular and matrix-biomechanical effects of Se-NPs on cancer cells and understand how Se-NPs could lead to significant alterations to the physicochemical forces that drive proliferation, differentiation, migration and biomechanics of studied cancer cells.

Discussion: Information on bio-nanomechanical properties of internalized Se-NPs will provide important understanding regarding chemopreventive effects of these nanoparticles. This is major point for designing drugs to inhibit invasion and metastasis, from which the majority of cancer mortalities arise.

Selected references
P87 - Redox-active Selenium as an Anticancer Agent: A Critical Review

3. Selenium in animal and human health and disease
3.4 Selenium based medical therapeutics
Keywords: Fibrinogen, selenite, sulfhydryl groups, tumors

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Introduction: Selenium (Se) occurs in the Earth soil in various organic and inorganic forms, however their distribution is very uneven, and in certain cases can result in a severe Se deficiency resulting in an increased incidence of cancer. Consequently, several organic and inorganic selenium compounds were developed and tested as possible anticancer agents. However, most recent data indicate that the most biologically active is an inorganic sodium selenite with four-valent Se, and not that with six-valent (selenate). This difference in their biological activities is due to their physicochemical properties.

Method: It was documented that the vicinal sulfhydryl groups of tumor protein membranes can react with fibrinogen molecules converting them into an insoluble polymer that coats tumors cells and presents them as ‘self’ to the innate cellular immune system.

Result: As the result, macrophages of the lymphatic system do not recognize neoplastic cells as ‘foreign bodies’ and spare them from the immune destruction. It is argued that redox forms of selenium oxidize proteins sulfhydryl groups expressed on the surface of tumor cells and thus inhibit the modification of fibrinogen molecules.

Discussion: The formation of a protective coat around the cancer cells can explain the failure of various immunotherapies to eliminate completely tumors from human bodies. In conclusion, redox-active forms of selenium can be used as inexpensive drugs of choice in the cancer treatment and prevention.
P88 - New insights into the development and biological evaluation of novel methylselenol precursors

3. Selenium in animal and human health and disease
3.4 Selenium based medical therapeutics

Keywords: methylselenol, cytotoxicity, cell death, cell cycle arrest

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Introduction: Selenium compounds can be potential therapeutic agents in cancer. Importantly, speciation, dose and cell status determine the biological effects, which are exerted by their metabolites. Methylselenol is one of the key executors of Se antitumor activity but given its high reactivity and volatility, the synthesis of precursors is required. Herein we developed a new series of mono- or bifunctionalized methylselenoesters where the core of the molecule modulates the release of methylselenol in aqueous environment.

Method: The compounds were synthesized by reaction of sodium hydrogen selenide with an acyl chloride and consequent methylation. Their ability to release methylselenol was tested by Ellman’s assay and their radical scavenging activity was evaluated by the DPPH and ABTS assays. Cytotoxicity was assessed by the MTT assay in a panel of 6 cancer cell lines. Two compounds were selected to evaluate their interaction with redox-related enzymes and their effect on cell cycle and cell death by flow cytometry.

Result: The compounds presented different rates of methylselenol release without significantly affecting cytotoxicity. Only one compound had radical scavenging activity. Most of the compounds had GI₅₀ values below 10 µM at 72h in most cell lines. The two analyzed compounds were substrates for thioredoxin reductase but not for the glutathione/glutaredoxin system. Both of them induced G2/M arrest in MCF7 cell line and cell death was partially executed by caspases.

Discussion: Even though further characterization is required, we provide new insights into the development of methylselenol precursors with good cytotoxic activity.

General structure of the novel synthesized compounds.
P89 - Genetic interactions in human methylation of selenium and arsenic

Introduction: Toxicological interactions between selenium and arsenic have been proposed, but how the elements interact in humans is unclear. We investigated potential interactions in the methylation of selenium and arsenic.

Method: For 226 pregnant women in a rural Bangladeshi area with elevated arsenic exposure through drinking water, we measured urinary selenium (U-Se) and arsenic (U-As) by ICPMS. Mono- and dimethylated arsenic metabolites (MMA and DMA) and trimethyl selenonium ion (TMSe) in urine were measured by HPLC-vapor generation-ICPMS. Methylation efficiency was assessed by relative amounts (%) of metabolites in urine. Genotyping for the main arsenic and selenium methyltransferases, AS3MT and INMT, was performed with TaqMan probes or Sequenom.

Result: In this sub-cohort with poor selenium status (~60% with plasma selenium <60 µg/L), the median concentrations of U-Se and U-As was 7.1 and 80 µg/L, respectively. Multiple linear regression analyses for %TMSe showed positive associations with %MMA and U-As, but inverse with U-Se. The associations were strongest in women predisposed to produce TMSe. There was an interaction for two of the assessed haplotypes of AS3MT and INMT genotype for their impact on %MMA and %TMSe. In particular, %MMA was only influenced by AS3MT among women with INMT GG genotype (TMSe non-producers). When aligning INMT and AS3MT sequences we found no strong evidence for homology.

Discussion: Our study indicates interactions between methylation of the essential element selenium and the highly toxic arsenic, although the methyltransferases INMT and AS3MT showed no sequence similarity. Importantly, INMT genotype appeared to impact the influence of AS3MT on arsenic methylation.
P90 - Organic selenium supplementation increases mercury excretion and decreases oxidative damage in long-term mercury-exposed residents from Wanshan, China

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease

Keywords: SelenoPrecise, Mercury, excretion, malondialdehyde, 8-hydroxy-2-deoxyguanosine

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Introduction: There are about 15 000 residents around WanShan Hg mine site, China. Due to a long history of extensive mercury mining and smelting activities, local residents are suffering from elevated mercury exposure. The current objective was to study the effects of oral supplementation with selenium-enriched yeast (SelenoPrecise, Pharma Nord, Denmark) in these long-term mercury-exposed populations.

Method: One hundred and three volunteers from Wanshan area were recruited and 53 of them were supplemented with 100 µg of organic selenium daily as selenium-enriched yeast while 50 of them were supplemented with the nonselemite-enriched yeast for 3 months. The effects of selenium supplementation on urinary mercury, selenium, and oxidative stress-related biomarkers including malondialdehyde and 8-hydroxy-2-deoxyguanosine were assessed.

Result: This 90-day selenium supplementation trial indicated that organic selenium supplementation could increase mercury excretion and decrease urinary malondialdehyde (MDA) and 8-hydroxy-2-deoxyguanosine (8-OHdG) levels in local residents.

Discussion: The significantly increased Hg excretion (approximately 1.5-2.5-fold) was seen after 30 days of Se supplementation and it continued to increase even at the end of the supplementation (day 90). The increased Hg excretion can be ascribed to the redistribution of Hg from the body deposit to the kidney and further be excreted through urine. The decreased urinary MDA and 8-OHdG levels indicate that the restored function of the selenoenzymes which lead to the reduced lipid peroxidation and DNA damage.
P91 - Positive association of apolipoprotein E allele ε4 with plasma selenium in Croatian pregnant females

Keywords: selenium status, apolipoprotein E, APOE polymorphism, metal(loid)s

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Introduction: Relation between apolipoprotein E gene (APOE) polymorphisms (allele ε4 carriers versus non carriers) and metal(loid)s (Hg, As, Pb, Cd, Se) in mothers and their newborns exposed to seafood metal(loid)s was investigated. Apo E is a lipid binding glycoprotein involved in lipid and neuronal metabolism, having antioxidative, metal-binding and immunomodulatory/anti-inflammatory effects. Its three isoforms are encoded by alleles ε2, ε3 and ε4, respectively. Allel ε4 is believed to be beneficial early in life and detrimental in age related diseases. ε4 carriers could be more susceptible to oxidative stress and metal toxicity.

Method: Archived DNA from Croatian pregnant females (n=222, aged 19-44 years, 3rd trimester) and newborns (n=176) was genotyped by TaqMan® SNP assay for rs429358 and rs7412 (Applied Biosystems, USA). Results were compared with metal(loid)s in maternal urine, milk, hair, blood, plasma, cord blood, cord plasma, and child urine. Statistics: STATA; projects: EU-PHIME1,2, EU-HEALS.

Result: We identified 17% (n=37) and 20% (n=35) carriers of allele ε4 among mothers and newborns, respectively. Mothers with allele ε4 had higher geometrical means of: i) blood selenium, mercury and arsenic; ii) plasma selenium, iii) hair mercury and iv) cord blood mercury. For selenium only, the associations persisted after taking into account the influence of possible confounders (Table 1). Drawbacks of the study were small number of ε4 carriers and low levels of metals (< 4.5 ngHg/g, < 4 ngAs/g , < 14 ngPb/g, < 1ng Cd/g - maternal & cord blood).

Discussion: Superior selenium status found in healthy pregnant females carrying allel ε4 could be linked to the ‘APOE ε4 beneficial effects early in life’.

Selected references
P92 - Preliminary toxicology evaluation of selenite cataract model

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease

Keywords: Selenite cataract, Toxicology, Oxidative damage, Model evaluation, Drug screening

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Introduction: Selenite cataract, an experimental animal model for simulating human senile cataract, is widely used in investigating mechanism of cataract formation, as well as screening anti-cataract drugs, because it is quick and convenient. As screening anti-cataract drugs, the biological function of candidate drugs have to be affected inevitably by physiological state of the model animal. However, there is no report on the effect of high dosage sodium selenite on the other major organs.

Method: In order to investigate this important topic, we measured the changes of antioxidant status of some tissues in selenite cataract rat, such as liver, kidney and brain.

Result: The results show that, comparing with the control group, injecting sodium selenite resulted in slight decrease of mRNA expression level of GPx1 and MsrA in liver and brain, but increase in kidney; the change trend of GPx activity is also like this. However, mRNA expression level of MsrB1 in liver, kidney and brain all increased. After injection of sodium selenite, MDA content increased significantly in rat liver, remained unchanged in brain, but reduced slightly in kidney. The result of immunohistochemistry analysis displayed the similar change trend.

Discussion: These results indicate significant oxidative damage in the liver of selenite cataract rat; the balance of redox state in liver and kidney has been broken by high dosage sodium selenite. So in the future research, this important change of redox balance should be taken into account when screening anti-cataract drugs by this animal model.

Selected references
P93 - The powerful ameliorating effect of selenium against deltamethrin-induced oxidative stress in lactating rats and their suckling pups

3. Selenium in animal and human health and disease
3.1 Selenium supplementation for animal and livestock health

Keywords: Lactating rats, Offspring, Deltamethrin, Oxidative stress, Amelioration index

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Introduction: Selenium (Se) has shown its protective effects against toxic hazards of methomyl-induced hepatorenal dysfunction in experimental animals [1, 2]. This was referred to its role as an antioxidant agent. Here, we evaluate the ameliorative effect of Se against deltamethrin-induced oxidative stress to lactating rats and their suckling pups.

Method: Deltamethrin (DEL) at doses equivalent to 1/10 and 1/100 LD50 were administered to dams, either with or without Se (6.66 µg/kg bwt), from postnatal day 1 (PN1) until day 20 (PN20) after delivery. One more group was administered Se only, while the sixth one served as water control. On the day 21, both dams and their suckling pups were sacrificed.

Result: Compared with control results, animals treated with DEL recorded high elevation in the activity of ALT, AST, ALP, creatinine and MDA, as well as severe decline in BuChE, urea, SOD and TAC. Also, DEL caused noticeable histopathological changes in liver, kidney and ovary. The dams were affected than their pups. Co-administration of Se in conjunction with DEL normalized the hepatorenal parameters, as well as the antioxidant status especially at the lower dose, and achieved noticeable improvement at the higher one.

Discussion: The efficiency of Se was revealed by estimating “Amelioration Indices” [3]; which ranged between 0.70 and 1.03 for the obtained biochemical parameters in DEL+Se treatments. It was concluded that low doses of DEL can induce oxidative stress to lactating rats and to their offspring which were not treated directly with the pesticide, but Se can alleviate such hazards to a great extent.

Selected references
P94 - Nutritional availability of selenium derived from raw or roast beef

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease
Keywords: Beef, nutritional availability, cooking with heat

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Introduction: It is believed that beef contains selenium (Se) as selenocysteine (Sec) residue in protein. In many cases, beef is eaten after cooking with heat. Because Sec is unstable chemically, a nutritional availability of Se in beef may change with heat. In the present study, we examined the nutritional availability of Se derived from raw or roast beef.

Method: Beef round was sliced to an approximately 8 mm thickness and roasted at 200 to 250°C for 5 minutes. After lyophilization, the raw and roast beef round were powdered and defatted. Protein and Se contents in these beef powder were 88.5% and 610 ng/g, respectively. Three types of experimental diet (casein diet, raw beef diet, roast beef diet) were prepared using casein and sodium selenite, the raw beef powder or the roast beef powder as protein and Se sources. Se content of these experimental diets was 0.11 µg/g. Male 4-week-old Wistar rats were fed these three diets. After four weeks, Se content and glutathione peroxidase (GPX) activity of serum, liver and kidney were measured.

Result: In Se content, rats fed the raw beef diet showed lower values in the serum and liver compared to other two groups. In the kidney Se, the highest value was observed in rats fed the roast beef diet. However, in GPX activity, rats fed the roast beef diet tend to show lower values in liver and kidney.

Discussion: These results indicate that cooking with heat may change the tissue accumulation and nutritional utilization of Se derived from beef.
P95 - The inhibitory effect of selenium nanoparticles on atherosclerosis and the underlying mechanism

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**Introduction:** Atherosclerotic cardiovascular diseases represent one of the greatest threats to human health worldwide. Selenium nanoparticles (SeNPs) will be a novel prospect for nutritional supplementation because of their lower toxicity [1]. In this work, we investigated for the first time the potential of SeNPs to protect against atherosclerosis and the underlying mechanisms.

**Method:** SeNPs were synthesized by the wet chemical reduction method in the presence of bovine serum albumin. Apolipoprotein E-knockout (apoE<sup>−/−</sup>) mice fed with high-fat diet were randomly divided into four groups: control group (untreated with SeNPs), low dose SeNPs group (25 mg·kg<sup>−1</sup>·d<sup>−1</sup>), moderate dose SeNPs group (50 mg·kg<sup>−1</sup>·d<sup>−1</sup>), high dose SeNPs group (100 mg·kg<sup>−1</sup>·d<sup>−1</sup>). SeNPs were administrated by gavage. After 12 weeks, atherosclerotic lesions in the aortic arch were assessed by haematoxylin and eosin staining and transmission electron microscope. Serum lipid levels, serum and hepatic oxidative stress, the expressions of some selenoproteins and the key enzymes of cholesterol synthesis and fatty acid synthesis were measured as well.

**Result:** Administration of SeNPs at different doses significantly decreased serum lipid levels and atherosclerotic lesions in the aortic arch of mice. Furthermore, SeNPs significantly counteracted serum and hepatic oxidative stress. The expressions of some selenoproteins having antioxidant capacity were increased significantly in SeNPs-treated mice. In addition, SeNPs were able to inhibit the expressions of HMG-CoA reductase, fatty acid synthetase, acetyl CoA carboxylase and stearoyl-CoA desaturase-1.

**Discussion:** This study suggested that SeNPs could alleviate hyperlipidemia and atherosclerotic lesions in apoE<sup>−/−</sup> mice, possibly by regulating oxidative stress and lipid metabolism.

**Selected references**

P96 - Effects of Selenium on Protein related-High Density Lipoprotein and Apolipoprotein B-100 Expression in Human Primary Hepatocytes.

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease

Keywords: Apolipoprotein B-100, Glutathione Peroxidase, High Density Lipoprotein, Sodium Selenite

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Introduction: Selenium dependent enzyme, Glutathione Peroxidase 1 (GPx1) is correlated with the activity of nuclear factor-kB (NF-kB). NF-kB is a major transcription factor that regulates the expression of various genes, such as lipid transport-related genes, including apolipoprotein. Its activation can be triggered by reactive oxygen species (ROS). The purpose of this study is to investigate the effect of selenium supplementation on the preβ-HDL formation and apolipoprotein B-100 (apoB-100) expressions in human primary hepatocyte (Hc cells) under a basal state condition.

Method: The Hc cells, representing a healthy human liver, were cultured in a medium supplemented with 0–10 µM sodium selenite. The effects of sodium selenite supplementation on several target proteins and genes expressions were measured by western blot analysis and real-time PCR, respectively.

Result: After treatment with sodium selenite, protein and gene expressions of GPx-1 and apolipoprotein A-I (apoA-I) were upregulated significantly (p<0.05). There were also upregulation of mRNA expressions of selenoprotein P and apoB-100 significantly (p<0.05). The optimum dose of sodium selenite supplementation on protein and mRNA expressions on the same dose (50 nM) and higher concentrations reduced the effect.

Discussion: Sodium selenite supplementation increases the expression of ApoA-1 and apoB-100 in Hc cells at low doses under basal state condition.

Selected references


P97 - Thoracic aortic degeneration and dilatation: a newly-recognised complication of human selenoprotein

Keywords: Human selenoprotein deficiency, aortic dilatation, ROS

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Introduction: 12 individuals from 9 families with selenoprotein deficiency due to homozygous or compound heterozygous mutations in SECISBP2 have been described. Here, we describe a new, potentially life-threatening, vascular phenotype associated with this disorder.

Method: Clinical and imaging surveillance has documented progressive aortic dilatation in three patients (P1, P2, P3). Histological and biochemical studies were undertaken in aortic tissue and vascular smooth muscle cells from P1 following surgery.

Result: Patient P1 (male, age 43yrs), with abnormal thyroid function (1) and compound heterozygous SECISBP2 mutations (K438X, c.1312A>T and c.1894-3C>G), developed ascending aortic dilatation (8cm) age 23yrs, requiring aortic arch and valve replacement; recurrent distal thoracic aortic dilatation age 43yrs, necessitated further surgery. Histological studies show cystic degeneration of the medial aortic layer. Cultured medial smooth muscle cells exhibit reduced selenoprotein expression, raised ROS and increased membrane lipid peroxidation. P2 (male, age 44yrs), with known, compound heterozygous SECISBP2 mutations (2), has a high-arched palate, joint hypermobility, digital vasospasm (Raynaud’s) and varicose veins. Progressive aortic dilatation (5cm) and valvular regurgitation mandate surgery. P3 (male, age 19yrs), with a homozygous SECISBP2 defect (3), has progressive dilatation (Z score >3) of aortic root and sinotubular junction.

Discussion: We have documented progressive thoracic aortic dilatation, requiring surgery in two cases, in three adult, selenoprotein-deficient patients. Clinical features (P2) and aortic histology (P1) resemble other aortopathy-associated syndromes (e.g. Marfan’s, Loeys-Dietz, Ehlers-Danlos). Aortopathy may not have developed as yet (e.g. childhood patients) or been ascertained in other cases. How selenoprotein deficiency mediates this vascular abnormality remains to be elucidated.

Selected references

P98 - The combined use of selen organic compounds with bioslastelin at toxic hepatitis

Keywords: bioslastelin, toxic hepatitis, cytochrome P450, cytochrome b5, antioxidant system

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Introduction: Nowadays despite the many positive attributes of herbal drugs, great attention is paid on the treatment of a disease by the combination of drugs. The aim of this work is to study the effect of new synthetic compound of selenium in combination with bioslastelin on the quantitative content of cytochromes P450, b5 and antioxidant system of the body on the background of toxic hepatitis (TH).

Method: The study was performed on outbred rats of both sexes. Piperidine selenophosphate (PSP) is newly synthesized selenoorganic compound. Its (LD50) was 300 mg / kg. TH of chemical genesis was modeled by intragastric injection water suspension of yellow phosphorus.

Result: The combined use of PSP and bioslastelin at TH significantly improves the studied parameters of antioxidant system of the body and quantitative content of cytochromes P450, b5. Thus the total therapeutic effects of binary composition of PSP and bioslastelin have significantly better therapeutic effect than applied individually. This demonstrates the synergy of their action in a joint application.

Discussion: To sum up, the potentiation of therapeutic effects of the selenium organic compound and liquorice root is explained by their effect on various links of antioxidant system and ability of bioslastelin to bond cytochrome P450, which makes it possible to justify binary combination as a more advanced and recommend using for the treatment and prevention of hepatitis and toxic poisoning.
P99 - In vivo effects of repeated thyronamine (T0AM) administration in male C57BL/6J mice

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease
Keywords: thyroid hormone metabolism, selenoprotein function

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Introduction: Thyronamines are decarboxylated and deiodinated metabolites of thyroid hormone (TH). Of all possible thyronamine variants, only 3-iodothyronamine (3-T₁AM) and thyronamine (T₀AM) have been detected in vivo. While intensive research is done on the (patho-)physiological action of 3-T₁AM, the role of T₀AM is studied less-intensively.

Method: Therefore, we determined whether a single pharmacological dose (50 mg/kg, i.p.) or repeated administration (5 mg/kg/day, i.p., for 7 days) of T₀AM affects metabolism, cardiovascular function or thermoregulation in C57BL/6J male mice.

Result: A single injection of T₀AM had no effect on heart rate, temperature or activity as assessed by radio telemetry. Likewise, daily administration of T₀AM did not alter body weight, food or water intake, heart rate, blood pressure, brown adipose tissue thermogenesis or body temperature, and no significant differences in hepatic glycogen content or mRNA expression of genes involved in cardiovascular function or metabolic control were determined. However, the TH - responsive genes Spot14 and selenocysteine-tRNA synthase were significantly upregulated, whereas hepatic deiodinase 1 and serum total TH concentrations were unaffected albeit hepatic T₀AM was significantly increased. Since selenium is important for proper TH function, we have analyzed its concentration in liver, serum and kidney using total reflection X-ray analysis, but no significant changes were observed.

Discussion: In summary, our data demonstrate that T₀AM elicits no obvious metabolic, cardiovascular, or thermoregulatory activities in mice. As T₀AM does also not interfere with TH metabolism, we conclude that the deiodination of 3-T₁AM to T₀AM constitutes an efficient inactivation mechanism, terminating the actions of the more powerful precursor.
P100 - Dynamic regulation of glutathione peroxidase 4 (GPX4) upon renal ischemia/reperfusion injury (IRI)

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease
Keywords: Glutathione peroxidase 4, ischemia/reperfusion injury, acute tubular necrosis, Ferroptosis

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Introduction: Among the known 24 (25) mammalian selenoproteins, GPX4 has emerged as one of the key selenocysteine containing enzymes being essential for early embryonic development in mice. Moreover, GPX4 has been recently described as a key regulator of ferroptotic cell death, a novel form of lipid peroxidation driven, caspase-independent, non-apoptotic cell death. Conditional ablation of Gpx4 in adult mice was shown to cause lethal acute kidney failure approximately 14 days after induction. Histopathologically, acute tubular necrosis (ATN) is the main cause of death sharing great similarities with human kidneys suffering from IRI.

Method: Kidneys of mice have been collected after transient unilateral renal artery clamping (marginally sublethal) followed by different periods of reperfusion to induce ATN. Immunohistochemistry as well as time course immunoblots of whole tissue lysates of the affected kidneys were performed and compared to its contralateral, non-clamped counterparts to evaluate GPX4 expression.

Result: We observed an early and massive downregulation of GPX4 protein levels in response to reperfusion compared to contralateral kidneys. The decrease in GPX4 protein was even more distinctive when focusing on the ischemic sensitive outer stripe of the outer medulla (OSOM) of the respective kidney.

Discussion: For the first time we provide evidence for the loss of murine GPX4 upon renal IRI. Further studies will reveal whether downregulation of GPX4 as well as ATN can be pharmacologically ameliorated. Additionally, we will evaluate renal GPX4 expression of adult deceased-donor kidney transplant recipients suffering ATN after transplantation.

Selected references
P101 - Assessment of Trace Elements and Systemic Oxidant Status in Dairy Cows during the Perinatal Period

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease

Keywords: Dairy Cows, Perinatal period, Oxidative stress, Trace elements

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Introduction: Oxidative stress is one of the main causes of ketosis and mastitis, diseases easily caught by cows during perinatal period. Some antioxidation enzymes and trace elements are widely used to measure oxidant status. The aim of this study is to investigate the variation and relationship among them in cows during the perinatal period.

Method: Blood samples were taken from 40 prenatal cows every 5 days. GSH-Px, SOD, CAT, GSH and MDA was measured by ELISA kits. Se, Mn, Zn, Cu and Fe was assayed using ICP-MS.

Result: The results showed that the highest value of GSH-Px accured at the 30th day before calving, while SOD at the 15th day. The value of CAT and Zn reduced to the bottom in labor, on the contrary GSH on the top. The MDA value began to rise gradually on the 30th day before delivery and sustained at high level afterwards. At calving day, the content of Se reached the peak and didn’t drop until the 5th day after delivery. The concentration of Mn increased in waves, Cu only elevated moderately from the calving day to the postnatal 10th day, and Fe decreased in the postnatal period. The levels of MDA exhibited a significant positive correlation (P < 0.05 or P< 0.01) with CAT and GSH, while an inverse relationship (P < 0.05 or P< 0.01) was found between Se/Cu and GSH-Px/SOD respectively.

Discussion: Our findings indicated that cows during the perinatal period suffered from oxidative stress, and Se addition might be needed to relieve such situation.

Selected references

P102 - MsrA Protects Hepatocytes against Acetaminophen-Induced Toxicity via TXNRD1 Regulation

Abstracts

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease
Keywords: Methionine sulfoxide, Acetaminophen, Hepatotoxicity, Thioredoxin reductase

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Introduction: Thioredoxin reductase 1 (TXNRD1) is associated with susceptibility to acetaminophen (APAP)-induced liver damage. Methionine sulfoxide reductase A (MsrA) is an antioxidant and protein repair enzyme that specifically catalyzes the reduction of methionine S-sulfoxide residues. We have previously shown that MsrA deficiency exacerbates acute liver injury induced by APAP.

Method: We used primary hepatocytes isolated from MsrA gene-deleted (MsrA<sup>−/−</sup>) and wild-type (MsrA<sup>+/+</sup>) mice to investigate the underlying mechanism of the protective effect of MsrA against APAP-induced hepatotoxicity.

Result: MsrA<sup>−/−</sup> hepatocytes showed higher susceptibility to APAP-induced cytotoxicity than MsrA<sup>+/+</sup> cells, consistent with our previous in vivo results. MsrA deficiency increased APAP-induced glutathione depletion and reactive oxygen species production. APAP treatment increased Nrf2 activation more profoundly in MsrA<sup>−/−</sup> than in MsrA<sup>+/+</sup> hepatocytes. Basal TXNRD1 levels were significantly higher in MsrA<sup>−/−</sup> than in MsrA<sup>+/+</sup> hepatocytes, while TXNRD1 depletion in both MsrA<sup>−/−</sup> and MsrA<sup>+/+</sup> cells resulted in increased resistance to APAP-induced cytotoxicity. In addition, APAP treatment significantly increased TXNRD1 expression in MsrA<sup>−/−</sup> hepatocytes, while no significant change was observed in MsrA<sup>+/+</sup> cells. Overexpression of MsrA reduced APAP-induced cytotoxicity and TXNRD1 expression levels in APAP-treated MsrA<sup>−/−</sup> hepatocytes.

Discussion: Collectively, our results suggest that MsrA protects hepatocytes from APAP-induced cytotoxicity through the modulation of TXNRD1 expression.
P103 - Knockdown of Sep15 modulates the expression of Pax6 induced by glucose in HLE cells

Keywords: Sep15; Pax6; cataract; lens epithelial cells

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Introduction: The 15-kDa selenoprotein (Sep15) is a thioredoxin-like selenoprotein, residenting in the lumen of endoplasmic reticulum and implicated in the quality control of glycoprotein folding through its interaction with UDP-glucose:glycoprotein glucosyltransferase. Although Sep15 KO mice developed a prominent nuclear cataract, the exact function of this protein is not known. Transcription factor Pax6 is highly expression in human lens epithelial (HLE) cells and essential for development of the eye system. The expression level of Pax6 changes with the process of lens fiber differentiation. When epithelial cells differentiate into primary lens fiber cells, the expression of Pax6 will be inhibited. However, effect of Sep15 knockdown on Pax6 remains largely unknown in lens fiber cells differentiation.

Method: Short interfering RNAs (siRNA) was used to knockdown the expression of Sep15 in human lens epithelial cells (SRA01/04). Expressions levels of Sep15, Pax6, αA-crystallin, p-ERK, p-AKT, β-catenin, sfrp2, six3 and cyclin D1 were measured by Western blotting or RT-PCR.

Result: In the present study, effects of knockdown of Sep15 or glucose on expression of Pax6 and related signal pathway were explored. Our results showed that knockdown of Sep15 or 30mM glucose treated alone resulted in the high expression of Pax6, however, combination showed the expression of Pax6 decreased significantly. Expressions of αA-crystallin, cyclin D1, p-ERK , p-AKT and other signal molecules also changed significantly.

Discussion: Our results demonstrate that knockdown of Sep15 impacts expression of Pax6 and related signal molecules, suggesting that Sep15 might play a role in regulating lens fiber cell differentiation.

Selected references
P104 - Roles of the 15kD selenoprotein in lens epithelial differentiation

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease

Keywords: 15kD selenoprotein, lens, cataract

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Introduction: Selenium is an essential micronutrient in mammals mainly presented in the form of selenoprotein. The 15kD selenoprotein (Sep15) is an endoplasmic reticulum-resident selenoprotein while its precise function still remains elusive. It was reported that Sep15 knockout mice developed nuclear cataracts at an early age, indicated the essential role of Sep15 in lens development. So we speculate that Sep15 might participate in regulation of differentiation in lens epithelial cells based on its structure and function.

Method: Effects of Sep15 knockdown by RNAi on some differentiation biomarkers were studied in bFGF-induced human lens epithelial cells (HLEC). Sep15 knockout mice were established by CRISPR/Cas9. Westernblot, RT-PCR and immunohistochemical were used to measure expression levels of differentiation biomarkers or some signaling molecules in HLEC and mice lens. HE stain and TEM were used to detect the pathology phenotypes of mice lens.

Result: Several biomarkers of lens cell differentiation such as β-crystallin and N-cadherin were significantly influenced by Sep15 knockdown in bFGF induced HLEC. That was also verified in lens of Sep15 knockout mice. The difference of lens lesion including some parameters between knockout and wildtype mice increased with age of mouse.

Discussion: The current results show that lens epithelial cell differentiation was influenced by Sep15 knockdown or knockout, suggesting that Sep15 might participate in regulating differentiation of lens epithelial cell. Further researches are needed to observe the lens abnormalities and more parameters in mice of different ages. We hope this work could contribute to reveal the relationship between Sep15 and cataract formation.

Selected references


P105 - Role of the 15kDa selenoprotein in colorectal inflammation

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease
Keywords: inflammation, colitis, 15kDa selenoprotein, gut microbiome

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Introduction: Previously, systemic knockout (KO) of the 15 kDa selenoprotein (SELENOF) protected mice against the formation of chemically-induced pre-neoplastic lesions and suggested modulation of inflammatory pathways [1]. We hypothesized that SELENOF expression is inversely correlated with a protective pro-inflammatory response in a colitis-associated mouse model.

Method: The potential role of SELENOF in inflammation was examined by assessing gene expression of KO mice and littermate controls (WT) that were either treated with 2% dextran sulfate sodium (DSS) to induce colitis or given water (control).

Result: Microarray analyses of colonic mucosa suggested that lack of SELENOF expression itself changed gene expression more significantly than DSS-induced inflammation. Changes in colonic gene expression in SELENOF-KO mice, as validated with qPCR, included up-regulation of inflammation-related genes, such as the interferon (IFN)-γ-regulated guanylate-binding protein 1 (p=0.004), compared to WT mice regardless of treatment. Because intestinal microbiota are known to influence inflammation of the host, bacterial DNA was isolated from feces of both WT and KO mice. Subsequent amplification of the 16S rRNA gene and sequencing with Illumina MiSeq revealed increased abundance of Proteobacteria, particularly Enterobacteriaceae, in untreated WT compared to KO mice.

Discussion: We evaluated the potential influence of SELENOF on the composition of the intestinal microbiome. Our preliminary analyses indicate that SELENOF may influence the expression of inflammatory genes in the colon. Its contribution to the regulation of colitis, which may be modulated by the microbiome, will be further examined.

Selected references
P106 - Low selenoprotein P status in patients with traumatic spinal cord injury

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease

Keywords: SELENBP1, diagnostics, sandwich assay, serum, neuron, paraplegia

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Introduction: Traumatic spinal cord injury is associated with neuronal loss and bears the risk for paraplegia. As selenium is an essential factor for neuronal development and protects from neuron degeneration, and traumatic injury constitutes a major inflammatory stimulus potentially down-regulating selenium metabolism, we aimed to monitor selenium status upon traumatic spinal cord injury during remission or non-remission by a time-resolved analysis.

Method: Selenoprotein P was analyzed by sandwich ELISA. Total serum selenium concentrations were measured by total reflection X-ray fluorescence. Concentrations of selenium-binding protein 1 were determined by a newly developed sandwich assay.

Result: A set of 20 subjects with traumatic spinal cord injury were analyzed, 10 of which went into remission and 10 developed paraplegia. In addition, 10 subjects with an unrelated injury were analyzed. Selenoprotein P showed a wide spectrum of concentrations within a given group, as well as between the three patient groups (mean±SD; 4.7±1.9; 4.9±1.6; 4.6±1.4 mg/L). Especially in one group of patients, selenium binding protein 1 was detectable in the early time points analyzed, decreasing with time. De-blinding of the patient characteristics for clinical associations has not yet taken place.

Discussion: Selenoprotein P is a cell-death inhibitory factor for neurons, which may be of high clinical importance in traumatic spine injuries. Some patients displayed extremely low selenoprotein P concentrations (< 1 mg/L) at early time points post injury, which may impair regenerative processes, causing excessive neuronal loss and may warrant selenium supplementation as promising adjuvant therapeutic measure in order to reduce paraplegia risk.

Selected references
Supported by Charité – Universitätsmedizin Berlin.
P107 - Redox regulated transcription factors in 3D spheroids enriched for cancer stem cells

Keywords: Redox regulated transcription factors, Nrf2, HIF, NF-kB, cancer stem cells

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Introduction: Previously, it is shown that specific types of cells, e.g. cancer stem cells (CSCs), are responsible for resistance, migration and invasion in many different cancers. Recently it has also been shown that the redox status and redox regulated transcription factors are highly connected with stem cell and CSC regulation and differentiation.

Method: Herein, we study the redox regulated transcription factors Nrf2, HIF and NF-κB during the initiation, resistance and maintenance of 3D spheroid cultures enriched for CSCs. To do this we have developed HCT116 cell lines that stably express, by us previously developed, reporter plasmid (pTRAF). The pTRAF system can be used to monitor the simultaneous activity of Nrf2, HIF and NF-κB and we here investigate the importance of these transcription factors in both adherent 2D and spheroid 3D colon cancer cultures.

Result: We demonstrate a unique time-and cell-resolved activation of these transcription factors that were coordinated during spheroid formation. We found that the Nrf2, HIF and NF-κB were activated in concurrence with specific markers of CSC. In growing spheroids Nrf2 and HIF were especially activated within the core of the spheroids. Additionally, pharmaceutically triggered induction of HIF promoted a quiescent stem cell like phenotype that were highly resistant to otherwise toxic insults.

Discussion: We believe that this unique platform can be used to further increase our knowledge of the redox regulatory systems, especially in resistant CSCs. This new knowledge can thereby be used, to specifically target these highly resistant CSCs, as a new strategy to combat cancer.

Selected references


P108 - Differential Impact of Two Secisbp2 Mutations on Selenoprotein Expression in Liver and Brain

Keywords: Secisbp2, missense mutation, R543Q, C696R, selenoprotein

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Introduction: Secisbp2 (SECIS-binding protein 2) interacts with SECIS (selenocysteine insertion sequence) element, located in the 3′-untranslated region of eukaryotic selenoprotein mRNAs. Secisbp2 is essential for the incorporation of selenocysteine, the 21st amino acid, into protein [1]. Recently, several SECISBP2 mutations were reported in human patients [2,3,4]. These mutations lead to reduced selenoprotein expression in many organs. Most patients suffer from thyroid hormone resistance, some also show neurological and muscular impairments [4,5].

Method: We have generated, liver and brain-specific Secisbp2 missense mutations in mice (C696R [2], R543Q [3]), in addition to Secisbp2 knockout mice which we have already published [6,7]. We also have employed ribosomal profiling to analyze in detail the impact of the R543Q mutation on selenoprotein expression in cerebral cortex.

Result: Our data show that both mutations significantly reduce the levels of several selenoproteins in liver – indistinguishable from the knockout mice. In neuron-specific transgenic mice, selenoprotein expression is significantly reduced in C696R mutant mice resembling Secisbp2 knockout mice. In contrast, the R543Q mutation is milder and does not lead to overt neurological dysfunction in mice in line with mild reduction of several selenoproteins.

Discussion: Our findings indicate that the distinct phenotypes in two mutations are caused by differential impact on selenoprotein expression. Next we will find out how two mutations affect the translation status of selenoprotein biosynthesis.

Selected references
P109 - A suppressive effect of selenium on amyloid-β plaque deposition in Tg2576 transgenic mice brain

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease

Keywords: Alzheimer's disease, selenium deficiency, GPx

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Introduction: Alzheimer’s disease (AD) is a chronic neurodegenerative disease and characterized by deposition of amyloid-b (Ab) peptide in the brain. Reactive oxygen species (ROS) are thought to be associated with the onset and/or progression of AD. Selenium-dependent glutathione peroxidases (GPxs) play a critical role in the brain in the extinction of ROS. The selenium concentration in the brain is kept higher than those of other organs/tissues even when dietary selenium is limited, which suggests the importance of this element in the brain. In this paper, we investigated the effect of dietary selenium on the Ab plaque deposition in the Tg2576 transgenic mice brain.

Method: Female 16-week-old Tg2576 mice (Taconic Farms, Inc.) were fed diets with different selenium-status for 17-month. Selenium concentrations were fluorometrically determined with 2,3-diaminonaphthalene after acid digestion. GPx activity was determined by measuring a decrease in the absorbance of NADPH at 340 nm. Ab plaques in brain sections were fluorescently detected using amyloid-specific thioflavin T.

Result: Selenium concentration and GPx activity in the brain of the selenium-deficient diet-fed mice was lower than those in the selenium-adequate conventional diet-fed mice. The number of Ab plaques in the brain sections of the selenium-deficient diet-fed mice was larger than those in the selenium-adequate conventional diet-fed mice. A similar trend was also observed between the selenium-deficient and the selenomethionine (SeMet)-supplemented diet-fed mice.

Discussion: The Ab plaque deposition in the brain of the Tg2576 mice appears to associate with its GPx activity.
P110 - Homozygous mutation p.P190L in TXNRD1 is associated with genetic generalized epilepsy

3. Selenium in animal and human health and disease
3.6 Clinical genetics of selenium or selenoprotein-encoding genes

Keywords: Generalized seizures, Epilepsy, Oxidative stress, TXNRD1

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Introduction: Growing evidence points to oxidative stress is connected to epilepsy pathogenesis. Increased production of ROS has been reported during seizures. Loss-of-function mutations in brain selenoproteins lead to epileptic phenotypes in mice and humans.

Method: Here, we report a homozygous mutation in TXNRD1 (thioredoxin reductase 1) in a family with genetic generalized epilepsy identified via whole exome sequencing.

Result: The TXNRD1 mutation p.Pro190Leu affecting a highly conserved amino acid residue co-segregates with the phenotype within the family. TXNRD1 activity was determined in subcellular fractions from a skeletal muscle biopsy and skin fibroblasts of the index patient. As result of the mutation, the activity of TXNRD1 was reduced in the patient's fibroblasts and skeletal muscle (to 34±3% and 16±8% of controls, respectively). Expression of the mutated protein was decreased as judged by 75Se labeling and Western blot analysis. We detected reduced 75Se-labeling of the enzyme (41±3% of controls) in patient fibroblasts. An in-depth in vitro kinetic analysis of the recombinant enzyme indicated 30–40% lowered kcat/Se values. Moreover, the patient fibroblasts were less resistant to a hydrogen peroxide challenge than controls.

Discussion: Therefore, a reduced activity of the enzyme in the patient's tissue samples is explained by (i) lower enzyme turnover and (ii) reduced abundance of the mutated enzyme. Our data agree with a potential role of insufficient ROS detoxification in epilepsy. They show that findings in genetic models deficient in cerebral selenoprotein expression are relevant to human disease.
P111 - Comparative analysis of selenium status in common neuropsychiatric disorders in children

Keywords: autism, cerebral palsy, attention deficit hyperactivity disorder, epilepsy

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Introduction: Selenium plays a significant role in various metabolic processes through its structural role in selenoproteins. At the same time, both selenium deficiency and excess was shown to be associated with increased oxidative stress and different pathologies, including neurodegenerative diseases.

Method: Therefore, the objective was to assess the level of Se in hair and serum of children (aged 0-5 and 5-15 years) with autism spectrum disorders (ASD), cerebral palsy (CP), communication disorders (CD), attention deficit hyperactivity disorder (ADHD), and epilepsy (Ep) in comparison with the control ones using ICP-DRC-MS.

Result: The obtained data demonstrate that children (< 5 y.o.) with Ep > ADHD > ASD > CD were characterized by significantly (p < 0.001) increased hair Se levels, whereas no significant difference was detected between the control and CP group values. More profound difference was detected in the older children (5-15 y.o.) due to significantly increased hair Se levels in all examined pathologies (p < 0.001). It is notable that serum Se levels were more stable in both groups of children. In a group of < 5 y.o. children hair Se levels decreased in the following order Ep > CP > ADHD > CD > ASD, whereas the difference from the control levels was significant only in Ep and CP groups (p < 0.05). In older children serum Se levels did not differ significantly from the control levels.

Discussion: So, Se metabolism is significantly deranged in children with common neuropsychiatric diseases. The observed difference in hair and serum Se levels may be indicative of increased Se excretion and changes in Se species.

Selected references


P112 - Does selenium intake relate to longevity: Case study in Shitai, Anhui, China

3. Selenium in animal and human health and disease
3.7 Additional and emerging topics of selenium in health or disease

Keywords: Selenium intake, Human longevity, Shitai

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Introduction: McCann found that age-related diseases were prospectively associated with modest Selenium(Se) deficiency and genetic dysfunction of nonessential selenoproteins [1]. A viewpoint that Se at low levels may be considered a hormetic chemical and decouple healthspan and longevity, was observed in a mice experiment [2]. Does Se play a vital role in human longevity? The understanding of its mechanism is far from being completed on human beings.

Method: In order to exclude the impact of climate, socio-economy, lifestyle and other variable factors, two different villages, Dashan as a longevity village and Kushan, 10 kilometers apart, as a common village, were selected in Shitai, Anhui, China. Soil, rice, hair and plasma samples were collected to determine Se concentrations. All treated samples were acid-digested and analyzed for total Se by atomic fluorescence spectrometry (AFS).

Result: As shown in Table 1, soil, rice, hair and plasma Se concentrations in Dashan were significant higher than those in Kushan ($P<0.05$)

Discussion: In this study, residents in Dashan have longer lifespan than those in Kushan (unpublished data). Except for the Se contents in soil, these two villages have similar conditions. Since the residents only eat foods locally, Se could be intaken by human beings via soil to rice, which is reflected by the Se contents in hair and plasma. To elucidate the mechanism of Se on human health, more investigations about human health indicators, such as GPX, SOD, will be carried out in future.

Selected references


Table 1 Se concentrations in soil, hair and plasma in two villages of Shitai County ($\mu$g/kg or $\mu$g/L)

<table>
<thead>
<tr>
<th>Villages</th>
<th>Soil Average</th>
<th>SD</th>
<th>Rice Average</th>
<th>SD</th>
<th>Hair Average</th>
<th>SD</th>
<th>Plasma Average</th>
<th>SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dashan</td>
<td>1607$^a$</td>
<td>242</td>
<td>699$^a$</td>
<td>/</td>
<td>351$^a$</td>
<td>179</td>
<td>115$^a$</td>
<td>26</td>
</tr>
<tr>
<td>Kushan</td>
<td>258$^b$</td>
<td>25</td>
<td>43$^b$</td>
<td>6</td>
<td>274$^b$</td>
<td>103</td>
<td>83$^b$</td>
<td>25</td>
</tr>
</tbody>
</table>

Average followed by different letters are significantly different at $P < 0.05$, SD=Standard Deviation
P113 - Influence of statins in selenium status and inflammatory profile considering creatine kinase levels

3. Selenium in animal and human health and disease
3.7 Additional and emerging topics of selenium in health or disease
Keywords: statin, inflammatory profile

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Introduction: Statins are the most frequently used medication for the treatment of hypercholesterolemia and prevention of cardiovascular events. However, their use has been associated with alterations in bioavailability of selenium, an essential micronutrient that plays a role in anti-inflammatory and antioxidant processes. The aim of this study was to assess the influence of the use of statins in selenium status and inflammatory profile in patients with normal or increased creatine kinase (CK) concentration.

Method: Individuals, using statin for less than a year, were classified as: Group 1 (G1): n=24, with changes in the CK levels, and Group 2 (G2): n=31, without changes in CK levels. Erythrocyte selenium, C-reactive protein (CRP), Interleukin 6 (IL-6), tumor necrosis factor alpha (TNF-α) and CK were evaluated.

Result: The average age of participants was 56.5±14.4 years. The sample consisted of 35 males (63.6%) and 20 females (36.4%). Comparison of CK, erythrocyte selenium, CRP, IL-6 and TNF-α between study groups are displayed in Table 1.

Discussion: Our study demonstrated that individuals using statin presented a deficiency in erythrocyte selenium. The higher levels of IL-6 in both groups indicate a pro-inflammatory profile that did not seem related with CK changes but could be associated with the deficiency of selenium. Interestingly, the concentrations of TNF-α were increased in both groups and was higher in G1 (increased CK levels) indicating a relative risk for cardiovascular events. Finally, more studies are required to further address the effects (or side effects) of statins in selenium status and metabolism.

Selected references

b VOLP, ACP et al. Inflammation biomarkers capacity in predicting the metabolic syndrome. Arq Bras Endocrinol Metab. 52(3): 537-549, 2008.

Table 1: Concentrations of CK, erythrocyte selenium, CRP, IL-6 and TNF-α of participants according to the study group

<table>
<thead>
<tr>
<th>Variable</th>
<th>G1</th>
<th>G2</th>
<th>P-value</th>
<th>Reference values</th>
</tr>
</thead>
<tbody>
<tr>
<td>N</td>
<td>24</td>
<td>31</td>
<td></td>
<td></td>
</tr>
<tr>
<td>CK (U/L)</td>
<td>205±5.6</td>
<td>62.2±24.3</td>
<td>&lt;0.001*</td>
<td>24–195</td>
</tr>
<tr>
<td>Erythrocyte Se (pg/L)</td>
<td>71.4±20.7</td>
<td>77.3±51.0</td>
<td>0.59</td>
<td>95</td>
</tr>
<tr>
<td>CRP (mg/dL)</td>
<td>1.1±3.6</td>
<td>0.66±0.7</td>
<td>0.50</td>
<td>≤1.0*</td>
</tr>
<tr>
<td>IL-6 (pg/mL)</td>
<td>4.5±5.0</td>
<td>5.33±4.4</td>
<td>0.53</td>
<td>1.35–3.22*</td>
</tr>
<tr>
<td>TNF-α (pg/mL)</td>
<td>11.5±4.5</td>
<td>8.6±3.4</td>
<td>&lt;0.01*</td>
<td>6.7*</td>
</tr>
</tbody>
</table>

Note: G1 and G2 values expressed as mean ± standard deviation. P-values obtained from Student’s t-tests.

*Significant difference at p<0.05.
P114 - The role of selenium in an in vitro invasive model of pancreatic cancer

3. Selenium in animal and human health and disease
3.7 Additional and emerging topics of selenium in health or disease

Keywords: pancreatic cancer invasion

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Introduction: Pancreatic cancer patients have been found to have lower levels of serum selenium compared to healthy controls, and higher levels of serum selenium have been correlated with longer survival of patients with pancreatic cancer¹. These findings suggest an important role for selenium intake and potential therapeutic supplementation in pancreatic cancer. However, less is known about the role selenium may play in pancreatic cell invasion and motility. This work focuses on the effect of pre-treatment and co-treatment with an organic and an inorganic selenium source on motility and invasion in pancreatic cancer cells in vitro.

Method: A panel of pancreatic cell lines with varying invasive profiles were tested for their sensitivity to two selenium compounds using acid phosphatase as an end-point to toxicity assays.

Pancreatic cancer cells were pre-treated or co-treated with two selenium compounds before being set up for in vitro invasion assays using the Boyden chamber method. Motility (without matrigel) and invasion (with matrigel) was assessed by crystal violet staining of migrated or invaded cells.

Result: Selenium tolerance varied greatly in the panel of pancreatic cell lines.

Pre-treatment and co-treatment with the organic and inorganic compounds gave differential effects on the migrating and invading capacity of the cells.

Discussion: This work will aid in our understanding of any potential role for selenium in pancreatic cancer metastasis, and whether that role is affected by time or by the chemical form of selenium used.

Selected references

P115 - The role of environmental metal ions in human health and diseases

3. Selenium in animal and human health and disease
3.7 Additional and emerging topics of selenium in health or disease

Keywords: Excess selenium, cadmium toxicity, metal interactions, lipid metabolism, liver disease

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Introduction: Multiple metal ions are present in the human diet, with some being required as essential nutrients and all causing toxicity at high intakes. Low level environmental cadmium (Cd) occurs in human diet and accumulates in vivo. Selenium (Se) is an essential micronutrient, and a component of 25 selenoproteins responsible for physiological functions. However, excess Se increases the risk of diseases, therefore, determination of adequate Se amount for human health is critical. Impacts of metals and their interaction on cells and organs need to be addressed.

Method: C57BL6 male mice were treated with Se (4 mg Na₂SeO₄/L, 16 weeks), Cd (3.3 mg CdCl₂/L, 16 weeks) and mixture. Metals, redox states, metabolites and phenotypic markers for lung and liver functions were analyzed.

Result: Urinary Se was 4-6 folds higher in Se-treated group while no changes were observed in lung and liver. Cd levels were elevated in urine, lung and liver in Cd mice. Se-treated mice significant increased body mass than control or Cd groups. Cd oxidized Prx3 in lung but not in liver, and co-treatment showed that Se inhibited Prx3 oxidation. Liver metabolomics of Se-treated mice showed that altered metabolome was significantly associated with weight gain, and accompanied by decreased levels of bile acids and acyl carnitines.

Discussion: Excess Se disrupts lipids homeostasis. Our study showed that mice treated with Se caused higher Cd accumulation in lung and liver, suggesting that excess Se could elevate Cd toxicity. The results suggest that presence of multiple metal ions in diet could affect human health potentiating metal toxicity.
P116 - Selenoproteins regulate B cell functions by targeting B cell receptor (BCR)-mediated antigen present

3. Selenium in animal and human health and disease
3.5 Selenium metabolism or selenoprotein function in health and disease

Keywords: selenoprotein, B cell, ROS, antigen processing

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Introduction: The role selenoproteins in B cells remains poorly understood. We sought to determine the role of selenoproteins in BCR-mediated effects because BCR-mediated signaling is critical in B cell development, activation, and functions and is highly sensitive to ROS.

Method: BCR endocytosis, calcium signaling, antigen trafficking, and antigen processing were measured in B cells that were maintained at various concentrations of sodium selenite (Na₂Se₃, 0-500 nmol) or B cells isolated from the mice maintained on Se-deficient, -adequate or -supplemented diets. The effect of ROS on BCR-mediated functions was determined by treating the cells with N-acetyl Cysteine (NAC). The role of selenoproteins were determined by generating mice that lack selenoproteins in B cells.

Result: The kinetics of BCR endocytosis, Ca signaling, antigen trafficking and processing were significantly increased in B cells treated with >200 nmol Na₂Se₃ compared to cells treated with 0-100 nmol. Similarly, BCR-mediated signaling and antigen processing were significantly increased in B cells from Se-supplemented compared to that of Se-deficient or -adequate mice. The ROS levels were significantly higher in B cells maintained at deficient or adequate levels of Se. However, upon treatment with the NAC, the BCR-mediated functions were significantly enhanced in Se-deficient and Se-adequate B cells. It was further determined that BCR-mediated effects were regulated by selenoproteins.

Discussion: This study demonstrates that selenoproteins regulate BCR-mediated effects by controlling ROS levels in the cells. Interestingly, the functions of B cells were significantly enhanced by supplementing with higher levels of Se. Thus, selenoproteins could be the new targets for optimizing humoral immunity.
P117 - Effect of anti-rheumatic treatment on selenium levels in rheumatoid arthritis, psoriatic arthritis

3. Selenium in animal and human health and disease
3.7 Additional and emerging topics of selenium in health or disease

Keywords: Rheumatoid arthritis, Psoriatic arthritis, Ankylosing arthritis, inflammation

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Introduction: The reason for increased cardiovascular (CV) risk in inflammatory rheumatic diseases (IRDs) is unclear. Selenium deficiency has been proposed to contribute to development of CV disease. Although the reference range is 50-120 µg/L, there are indications that levels <80 µg/L might be insufficient [1]. Our aim was to measure serum selenium (s-Se) levels in IRD patients, and to evaluate the effect of anti-rheumatic treatment on s-Se levels.

Method: We examined samples from 130 patients with IRDs (64 with Rheumatoid arthritis (RA), 40 with Psoriasis arthritis (PsA) and 26 with Ankylosing spondylitis (AS)) starting with methotrexate monotherapy or anti-tumor necrosis factor treatment (anti-TNF) with or without methotrexate. s-Se and serologic inflammatory biomarkers were measured. The patients were evaluated at baseline, and after 6 weeks and 6 months of therapy.

Result: The baseline median s-Se level in the total IRD group was 72µg/L. The s-Se levels increased in all groups (total, RA, PsA and AS) after therapy, but the differences from baseline were not significant. Changes in s-Se were significantly negatively related to changes in C-reactive protein and erythrocyte sedimentation rate. s-Se increased at 6 weeks (p=0.012) and 6 months (p=0.038) in patients treated with methotrexate. Anti-TNF was not associated with any significant s-Se changes.

Discussion: IRD patients appeared to have suboptimal s-Se levels, but the levels significantly increased with methotrexate treatment. Further studies are needed to determine if methotrexate and other anti-rheumatic drugs might reduce CV risk, and to determine if Se supplementation might reduce CV risk in IRD [2].

Selected references

P118 - Selenium, prostate cancer, and U-shaped thinking: A paradigm shift in public health research

3. Selenium in animal and human health and disease
3.7 Additional and emerging topics of selenium in health or disease
Keywords: SELECT, Cancer prevention, Prostate, U-shaped dose response, Human health

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Introduction: The perception pervasive among the public is that, when it comes to taking dietary supplements, “more is better”. A growing body of scientific evidence, however, suggests that the dose response between intake of selenium and other micronutrients and the cellular damage that accompanies age-related disease is U-shaped. As a consequence, more selenium may not always be better for overall health or disease prevention.

Method: Our challenge as health professionals will always be one of integrating our fragmented clinical experience. In this presentation, we transform this challenge into opportunity by looking at the relationship between selenium and prostate cancer risk as a potential paradigm shift in public health research — seeing prevention through the lens of U-shaped thinking, using 4 different angles of vision (Figure).

Result: A recent meta-analysis by Hurst et al showed a U-shaped dose response relationship between toenail selenium and risk for prostate cancer. Landing in the trough of the U — achieving mid-range selenium status — is better than being too low or too high. This stance is bolstered by an extensive review of the scientific literature by Rayman. This kind of U-shaped thinking sits at the core of personalized (precision) medicine.

Discussion: Promoting health in a more-is-not-necessarily-better world poses distinctive challenges. But U-shaped thinking can move us closer to defining risk-benefit profiles for nutrients — moving toward conceptualizing zones of selenium status where supplementation might be beneficial, as well as identifying detrimental zones of selenium status where supplementation would be contraindicated.

Selected references
P119 - Development of a point-of-care test for selenoprotein P

3. Selenium in animal and human health and disease
3.7 Additional and emerging topics of selenium in health or disease

Keywords: POCT, lateral-flow, sandwich, ELISA, blood, bedside test

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Introduction: Selenoprotein P (SELENOP) constitutes a relevant and reliable biomarker of selenium intake and status in humans. Both genetic and analytical studies provided evidence linking SELENOP genotype and expression levels with human disease, including cancer, autoimmune or inflammatory diseases. Concentrations of SELENOP can be determined by different analytical methods, yet a point-of-care test (POCT) for bedside analysis is missing.

Method: Two SELENOP-specific monoclonal antibodies (mAb) have been generated in mice, intensively characterized and used to establish a reliable sandwich ELISA (1). One of these mAb was labeled by colloidal-gold and the other was immobilized as catcher line on a lateral flow matrix. Different labeling methods, mAb concentrations and ways of storage and pre-dilution of human serum samples are compared for optimizing signal generation.

Result: Taking advantage of the same mAb used for the sandwich ELISA, a reliable detection of SELENOP from human serum samples by a POCT is feasible. Serial dilutions of a standard sample yielded signal intensities correlating to the SELENOP amount in a concentration range of 10 – 500 ng/ml. Inter-assay POCT coefficients of variation were not consistently below 30%, indicating that further optimization steps are necessary.

Discussion: A SELENOP-specific POCT will enable the determination of SELENOP concentrations from drops of blood at the bed side or in field studies in a non-invasive way without the need for lab equipment or a cooling chain. Unfortunately, the requirements for a highly reliable systems are not yet fully reached, and some parameters need to be further optimized before it can be used in clinical studies.

Selected references
P120 - Age-dependent protective effect of Selenium against UVA irradiation in primary human keratinocytes

Abstracts

3. Selenium in animal and human health and disease
3.7 Additional and emerging topics of selenium in health or disease

Keywords: Skin, DNA repair, aging

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Introduction: Few studies have focused on the protective role of selenium (Se) on skin aging and photoaging although selenoproteins are essential for keratinocyte function and skin development.

Method: Our study aimed at studying the effect of selenium supplementation on primary human keratinocytes obtained from normal skin biopsies of two age-groups of donors elderly (60-70 years old) or young (20-30 years old), which could reflect different genetic background and physiological conditions, at baseline or after exposure to UVA irradiations. Moreover, using our multiplexed DNA repair assay, we assessed the DNA repair signature in primary keratinocytes coming from the age-groups of donors with or without Se supplementation.

Result: Low doses of Se were very potent protector against UVA-induced cytotoxicity on young keratinocytes, whereas aged keratinocytes require four times more of Se than young keratinocytes in order to be protected from UVA-induced cytotoxicity. Moreover, we showed a drastic fall in DNA repair capacities on old keratinocytes versus young one at basal level and Se supplementation enhances significantly DNA repair of 8oxoG, only on keratinocytes isolated from young donors.

Discussion: These original data strongly suggest an increased vulnerability of aged keratinocytes to oxidative damages with age and should be taken into account regarding Se needs in elderly. Strengthening of DNA repair activities by selenium may represent a new strategy to fight against aging and skin photo-aging. These results will highlight the protective mechanism of selenium, and therefore, could identify new targets for UVA exposure protection.

Selected references


P121 - Research and practice on the standard “Selenium-enriched agricultural products”

3. Selenium in animal and human health and disease
3.7 Additional and emerging topics of selenium in health or disease

Keywords: standardization, China, agricultural product

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Introduction: Selenium-enriched agricultural products has no corresponding national standards or industrial standards in China. Only a minority provinces in China established relevant local standards about selenium-enriched food, while organic selenium contents were not provided yet.

Method: From 2013 to 2015, selenium content and Se amino acid content in 401 agricultural products samples collected from 17 counties from 6 provinces of China were determined. The setting values of this standard were mainly considered from the following aspects: daily selenium intake should be between 60 and 250 g/d recommended by Chinese Nutrition Society (2013 Edition), and the maximum dietary intake does not exceed 400 g/d. Reference to the existing selenium standards, this national standard “Selenium-enriched agricultural products” was established for supply and marketing cooperatives in China.

Result: This standard requests total selenium and organic selenium contents in 8 kinds of selenium-enriched agricultural products, including cereals, beans, potatoes, vegetables, edible fungus, meat, eggs, and teas.

Discussion: This standard “Selenium-enriched agricultural products” was compared with the existing standards of selenium-enriched agricultural products, such as Hubei provincial standard “Label of Selenium-enriched food” (DB42/211-2002), Ankang local standard “Selenium content classification standard for selenium-enriched food” (DB6124.01-2010), Shanxi provincial standard “Selenium content standard for selenium-enriched food and related products” (DB61/T556-2012), Guangxi provincial standard "Selenium content classification standard for selenium-enriched agricultural products" (DB45/T1061-2014), and Hubei provincial standard for food safety "Selenium content of organic-selenium-enriched foods" (DBS42/002-2014), the data of the samples collected in this standard was basically the same as the above standards.
P122 - Micro-spectroscopic investigation of selenium speciation in Se-rich rocks

Introduction: Selenium (Se) speciation in rocks is critically important for understanding the solubility, mobility, bioavailability, and toxicity of Se in the environment. However, Se speciation analysis in rocks using X-ray absorption fine structure (XAFS) technique is oftentimes limited due to its relatively low Se content.

Method: In this study, the distribution and chemical speciation of Se in Se-rich rocks from seleniferous areas were investigated using the µ-XRF-XAFS techniques.

Result: The µ-XRF result showed that Se distribution in rocks is highly heterogeneous, and elevated intensity of Se was generally found in the areas with higher intensity of Fe. Se K-edge µ-XAFS spectra for Se hotspots in weathered rocks were very similar to that of Se(IV), but reduced Se species was dominant in relatively fresh rocks.

Discussion: These results suggest that Fe-containing minerals are likely to be the main host phases of Fe in Se-rich rocks. Se in rocks can be transformed from Se(-II) to Se(IV) during its weathering process, resulting in the release of Se into soil and aqueous environment.
P123 - Selenium-enriched yeast inhibits Aβ generation and modulates autophagy in the 3xTg AD mouse

3. Selenium in animal and human health and disease
3.1 Selenium supplementation for animal and livestock health

Keywords: Selenium-enriched yeast, Alzheimer’s disease, Aβ, autophagy

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Introduction: As a progressive, degenerative brain disease that destroys memory and thinking skills in aged people, Alzheimer’s disease (AD) is pathologically featured by numerous amyloid plaques surrounded by dystrophic neurites, profound synaptic loss, and neurofibrillary tangles. The amyloid plaques are mainly composed of amyloid β-peptide (Aβ), a primary factor contributing to the pathogenesis of AD. Elimination or reducing the level of Aβ is considered an important strategy for AD treatment. The therapeutic effect of selenium has been confirmed in a number of experimental models of neurodegenerative diseases. Selenium-enriched yeast (Se-yeast) is a common used selenium-containing nutritional supplement.

Method: The objective of our study was to investigate the effect and underlying mechanisms of Se-yeast on Aβ pathology in a 4-month-old triple transgenic mouse model of AD (3×Tg AD mice).

Result: Administration of Se-yeast attenuated Aβ deposition in the brain of AD mice and decreased the ratio of LC3II/LC3I. Se-yeast treatment also decreased the amyloid protein precursor (APP) level, downregulated the activity of AMP-activated protein kinase (AMPK) and upregulated the activities of Akt, mTOR, and p70S6K. Furthermore, concomitant with the increased turnover of Aβ and APP, the level of p62 also significantly decreased and cathepsin D level increased in Se-yeast-treated AD mice.

Discussion: These results suggest that besides decreasing the generation of Aβ, Se-yeast also inhibited the initiation of autophagy through modulating the AMPK/AKT-mTOR-p70S6K signaling pathway and enhanced autophagic clearance, thus reduced the burden of Aβ accumulation in the brains of AD mice. Our results further highlight the potential therapeutic effects of Se-yeast on AD.
P124 - Metalloomic quantifying metals in metalloproteins: The levels of Hg and Se binding to serum selenoproteins

1. Selenium chemistry and geochemistry
1.7 Selenium interactions with other elements in the environment

Keywords: metalloomics, methylmercury, selenoproteins, rats, isotope dilution

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Introduction: It has been known for decades that Se could antagonize the toxicity of Hg, both in inorganic (Hg2+) and organic (i.e. methylmercury, MeHg) forms in different organisms when co-exposed. It is not known if such kind of complex will also form when selenite meet methylmercury (MeHg) in blood stream. Besides, it is not known how much Hg and Se in serum were bound to selenoproteins.

Method: Selenite was supplemented for 30 days in methylmercury-poisoned rats. Blood samples were collected and the contents of Se and Hg in serum were measured using ICP-MS. AF coupled to ICP-MS was used to find out the Hg, Se binding selenoproteins in serum samples. Post-column isotope dilution analysis using both enriched 78Se and 199Hg as spike was applied to quantify the levels of Hg and Se binding to selenoproteins.

Result: It was found that 93.6% of total Se and 73% of Hg were bound to SeIP in serum after Se treatment. Our results showed that Se treatment can reduce the level of Hg in serum, and increase the proportion of Hg bound to SeIP, indicating that interactions between Hg/Se in serum related to accumulation and detoxification processes.

Discussion: These results provide information about elements distribution and interactions and reveal the potential of metalloomic approaches in such kind of studies. This quantitative methodology to the same protein could be very useful for investigating the interaction between Se and other heavy metals in living organisms to understanding the role of these elements in health and disease.

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