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The selenium content of SEPP1 versus selenium requirements in vertebrates

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Selenoprotein P (SEPP1) distributes selenium (Se) throughout the body via the circulatory system. The Se content of SEPP1 varies from 7 to 18 Se atoms depending on the species, but the reason for this variation remains unclear. Herein we provide evidence that vertebrate SEPP1 Sec content correlates positively with Se requirements ($R^2=0.88$). As the Se content of full length SEPP1 is genetically determined, this presents a unique case where a nutrient requirement can be predicted based on genomic sequence information.

1 **The selenium content of SEPP1 versus selenium requirements in vertebrates.**

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13 Abstract

14 Selenoprotein P (SEPP1) distributes selenium (Se) throughout the body via the circulatory
15 system. The Se content of SEPP1 varies from 7 to 18 Se atoms depending on the species, but the
16 reason for this variation remains unclear. Herein we provide evidence that vertebrate SEPP1 Sec
17 content correlates positively with Se requirements ($R^2=0.88$). As the Se content of full length
18 SEPP1 is genetically determined, this presents a unique case where a nutrient requirement can be
19 predicted based on genomic sequence information.

20 Introduction

21 Selenium (Se) is an essential trace element required for the selenocysteine (Sec) residues inserted
22 during mRNA translation into Se dependent proteins, termed selenoproteins ([Brigelius-Flohé](#)
23 [1999](#)). The number of selenoprotein coding genes differs among vertebrate species, ranging from
24 the 24 to 25 found in mammals up to the 35 to 38 found in bony fish. Most selenoproteins are
25 redox enzymes containing a single Sec residue involved in catalytic activity ([Papp et al., 2007](#)).
26 An exception is the Se rich glycoprotein, selenoprotein P (SEPP1; aka SeP, SEPP, SEPP1a),
27 which in vertebrates contains 7 to 18 Sec residues, depending on the species ([Lobanov et al.,](#)
28 [2008](#)). The high Se content of SEPP1 is thought to facilitate Se distribution throughout the body.
29 In mammals, the liver is a major site of SEPP1 expression, where it is synthesised utilising food
30 derived Se. Hepatic SEPP1 is then secreted into the blood plasma ([Kato et al., 1992](#)), where
31 SEPP1 accounts for between 40 and 80% of the total Se ([Hill et al., 1996](#); [Hill et al., 2007](#); [Read](#)
32 [et al., 1990](#)). Tissues utilise a combination of receptor mediated endocytosis and pinocytosis to
33 obtain SEPP1 from the plasma, where it is then catabolised to release Se for *de nova*
34 selenoprotein synthesis ([Burk and Hill 2009](#); [Burk et al., 2013](#)).
35 Several features of SEPP1 are conserved among vertebrates including, i) a single N-terminal
36 domain Sec residue present within a thioredoxin like motif (UXXC, where U is Sec), ii) a
37 histidine rich region in the mid region of the protein), and iii) an apolipoprotein E receptor-2
38 (APOER2; aka LRP8) binding site followed by five Sec residues in proximity to the C-terminal
39 (Figure 1) ([Lobanov et al., 2008](#)). APOER2 is widely expressed in human tissues
40 (www.humanproteomemap.org; ([Kim et al., 2014](#)). APOER2 facilitated uptake of plasma SEPP1
41 is an essential (testes) or important (brain and foetus) pathway in some, but not all (muscle,
42 kidney, liver or whole body) tissues for maintaining Se homeostasis *in vivo* ([Burk et al., 2007](#);
43 [Burk et al., 2013](#); [Hill et al., 2012](#); [Olson et al., 2007](#)). In contrast, the histidine rich regions of

SEPP1 presumably interact with multiple receptors, including megalin (LRP2). A megalin facilitated uptake pathway minimises excretion of Se by binding SEPP1 fragments in the kidney (Kurokawa et al., 2014; Olson et al., 2008) and plays a role in maintaining tissue Se homeostasis (Chiu-Ugalde et al., 2010; Steinbrenner et al., 2006). Additionally, the histidine rich regions are associated with the heparin binding properties of SEPP1. It is postulated that the heparin binding properties of SEPP1 allow the N-terminal Sec of SEPP1 to provide antioxidant protection for endothelial cells at sites of inflammation (Hondal et al., 2001; Saito et al., 2004).

In contrast, other domains in SEPP1 have low conservation among species. For instance, single base mutations in genomes have led to many cases of Sec to cysteine (Cys) substitution within the vertebrate SEPP1 C-terminal domain upstream and including the APOER2 binding site (Figure 1) (Lobanov et al., 2008). The reason why Sec content plasticity is observed only within this region of SEPP1 is unclear, but it is responsible for most of the variation between the SEPP1 Sec content among vertebrates (Lobanov et al., 2008). Furthermore, why SEPP1 Sec content differs among species also remains unknown. Several lines of evidence suggest vertebrate SEPP1 Sec number may be a direct function of Se utilisation. For instance, vertebrate SEPP1 Sec content correlates positively with selenoproteome size, tissue Se levels, and Se bioavailability in the environment (Lobanov et al., 2008).

If a direct relationship between SEPP1 Sec content and Se requirements exists, the SEPP1 Sec content of a species could predict its Se requirements, or vice versa. In doing so, this would provide a new insight into how the genome effects nutrient utilisation. Additionally, such a relationship would allow considerable scope for implementing the 3R's (replace, reduce, refine). For example, this relationship would indicate the dietary Se levels to focus on when investigating the Se requirements for novel species. Such knowledge would reduce both the number of animals required and the risk of exposure to Se levels that may compromise animal welfare in such experiments.

70 In the following work, we compared the Sec content of mammalian and bony fish SEPP1's
 71 predicted *in silico* with their Se requirements determined *in vivo*. We found a strong positive non-
 72 linear correlation (0.88) between the two, suggesting Se requirements can be predicted from the
 73 *Sepp1* gene sequence. The correlation was dictated by the Sec content within the C-terminal
 74 domain upstream and including the APOER2 binding site. The model was limited, whereby
 75 further analysis suggested it could not predict Se requirements in species whose SEPP1 Sec
 76 content was >16 residues, as found in many bony fish species. The predicted Se requirements for
 77 vertebrate species based on their SEPP1 Sec content are provided.

Materials and methods

The *in silico* predicted species specific Sec content of SEPP1 (SEPP1a in fish) were obtained from Lobanov et al. (2008), the open access selenoprotein database (selenodb.org; (Romagné et al., 2014)) or by analysing genomic *Sepp1* sequences (NCBI) for Sec content (<http://sebastian.crg.es/>), an open access online software for this purpose (Mariotti et al., 2013). The SEPP1 Sec content of five bony fish species; loach (*Paramisgurnus dabryanus*), cobia (*Rachycentron canadum*), grouper (*Epinephelus malabaricus*), gibel carp (*Carassius auratus gibelio*) and yellowtail kingfish (*Seriola lalandi*); were assumed to be within the 15 to 17 residue range found for fish in general (Lobanov et al., 2008)(See Supp. Table 2). The species specific Se requirement data were obtained from published studies and from the National Research Council of the USA (NRC) nutrient requirement reports (Gatlin and Wilson 1984; Han et al., 2011; Hao et al., 2014; Hilton et al., 1980; Jensen and Pallauf 2008; Le and Fotedar 2013; Lei et al., 1998; Lin and Shiau 2005; Liu et al., 2010; NRC 1963; 1985; 1995; 1997; 2011; Penglase et al., 2014; Sunde et al., 2009; Wedekind et al., 2004; Weiss et al., 1996; 1997). See Supp. Table 1 for further information regarding these animal Se requirement studies. Where multiple Se requirement studies for a species were available, the dietary Se requirements to fulfil the requirements of the actively growing juvenile stage was selected. Data were analysed in GraphPad Prism (GraphPad Software, San Diego, CA, USA, V. 5.04). Data were fitted with a horizontal line (null hypothesis) and then tested against more complex models in the following sequence; first order polynomial, second order polynomial and five parameter logistic equation (5PL) asymmetric sigmoidal; until the simplest model that explained the data was found ($p < 0.05$). Model parameters were optimised to reflect current knowledge; vertebrates with seven Sec SEPP1; guinea pigs (*Cavia porcellus*) and naked mole rats (*Heterocephalus glaber*); have a Se requirement (Jensen and Pallauf 2008; Kasaikina et al., 2011). Thus the y intercept (no Se requirement) of models was constrained to ≤ 6

102 Sec for whole SEPP1 (Figure 2). Other vertebrate classes were excluded from the analyses
103 because of limited data.

104 **Results and Discussion**

105 ***The selenocysteine content of Selenoprotein P correlates strongly with selenium requirements.***

106 The Sec content in SEPP1 were identified for a total of 11 species; three bony fish and eight
107 mammals; for which the Se requirements are also published (Supp. Table 1). Using this data, a
108 positive non-linear correlation ($R^2=0.88$) was found between the Se requirements and SEPP1 Sec
109 number (Figure 2). Similarly, a positive correlation also occurs between SEPP1 Sec content and
110 selenoprotein number in vertebrates ([Kryukov and Gladyshev 2000](#); [Lobanov et al., 2008](#)). All
111 fish annotated to date have SEPP1 (aka SEPP1a in fish) with 15 to 17 Sec residues (See Supp.
112 Table 2). Based on this, an additional five bony fish species with known Se requirements were
113 assumed to have SEPP1's with 17 Sec residues and added to the data set, which was then re-
114 analysed. This resulted in an asymmetric sigmoidal trend with a plateau at 17.0 (Figure 2),
115 suggesting that a species SEPP1 is only useful for predicting Se requirements prior to this plateau
116 (≤ 16 Sec residues). When a species SEPP1 has >16 Sec residues, as is found in many fish
117 species, the curve predicts a minimum requirement (0.24 mg/Se kg dry matter (DM)) but not a
118 maximum (there is no correlation between SEPP1 Sec content and Se requirements above this
119 level). Modelling the data with alternative SEPP1 Sec content (15 or 16 Sec) for these five fish
120 species shifts the plateau height towards those values, but retains the general features of the
121 model (data not shown).

122 We then used this regression model (Figure 2) to predict the Se requirements of a species based
123 on its SEPP1 Sec residue number predicted *in silico* (Table 1). As discussed before, there is a
124 broad range of Se requirements found for bony fish that occurs in absence of an equally large
125 distribution of SEPP1 Sec content. The reason for this occurrence is unknown. Perhaps the

relatively straightforward single base mutations of Sec to Cys codons ([Lobanov et al., 2008](#)) allowed mammalian SEPP1 Sec content to decrease in comparison to the ancestral vertebrate SEPP1 and in line with Se requirements, while fish utilised other regulatory mechanisms to increase Se supply to peripheral tissues. For example *Sepp1* mRNA expression is elevated in fish, particularly in the kidneys, in comparison to mammals ([Lobanov et al., 2008](#)). This suggests plasma SEPP1 in fish may be replenished by SEPP1 synthesised from Se scavenged in the kidneys. As expected, the model (Figure 2) generally reflects the Se requirements of the species used to construct it, i.e. rat Se requirements are 0.1 mg Se/kg DM ([Weiss et al., 1996; 1997](#)) and it has a SEPP1 Sec content of 10 (Supp. Table 1), falling within the 0.08 ± 0.02 range predicted by the model (Table 1).

A hypothesis for the Sec number plasticity or conservation in different domains of vertebrate SEPP1.

As discussed, most of the difference in the SEPP1 Sec content between species is a result of differences in the Sec content found upstream and including the APOER2 binding site within the C-domain of SEPP1 (Figure 1 and Supp. Table 2). Thus as expected, when we analysed the Sec content in this region in relation to a species Se requirement (Supp. Figure 1), we found a similar positive correlation as found for full length SEPP1 and Se requirements (Figure 2). Recently it was found that SEPP1 Sec residues closer to the C-terminal are translated with greater efficiency than those towards the N-terminal ([Shetty et al., 2014](#)). Premature termination of SEPP1 translation at Sec codons appears to be a common event. For instance, four rat SEPP1 isoforms have been identified in plasma, whereby in addition to the full length protein, shorter variants are synthesised when translation is terminated at the second, third or seventh Sec codon ([Ma et al., 2002](#)). Thus, on average each plasma SEPP1 in mice contains 5 Sec residues, not the 10 Sec residues expected if only the full length protein is present ([Hill et al., 2007](#)). As a consequence of

this, a proportion of translated SEPP1 proteins will not contain the APOER2 binding site (Figure 1).

Taking in mind the above, we first hypothesise that decreases in Se requirements are an evolutionary adaption to Se availability. For instance, guinea pigs and naked mole rats both have low Se requirements ([Jensen and Pallauf 2008](#); [Kasaikina et al., 2011](#)), and inhabit the Andes or East Africa respectively, both regions of low Se status ([FAO 1992](#); [Rachel et al., 2013](#)). Secondly, we hypothesise that the Se requirements of the brain among species is similar on a weight basis, despite differences in the Se requirements of the whole body. For instance, compared to mice, naked mole rats have lower levels (-30 to -75%) of Se in most tissues except the brain ([Kasaikina et al., 2011](#)). And lastly, low Se availability can stall translation of selenoproteins at Sec codons ([Weiss Sachdev and Sunde 2001](#)), and perhaps results in the truncated forms of SEPP1 translated *in vivo*. We therefore hypothesise that Sec to Cys substitutions in SEPP1 occurred specifically in the region downstream and including the APOER2 binding site as it aids the translation of full length protein under Se limiting conditions, such as those faced by naked mole rats and guinea pigs. The subsequent retention of the APOER2 binding site would allow the continuation of a controlled Se supply to critical organs utilising APOER2 mediated uptake of SEPP1, such as the brain.

Conclusion

The Sec content of SEPP1 correlates with Se requirements in vertebrates when Sec residue number is ≤ 16 . There was no correlation between SEPP1 Sec content and Se requirements for species with SEPP1's with >16 Sec residues, as is the case for many bony fish species. However, for those species with SEPP1's with >16 Sec residues, a minimum Se requirement of 0.24 mg Se/kg DM was predicted. This study suggests that genome evolution is affected directly by

173 nutrient availability in the environment, and provides novel evidence that the genomic sequence
174 can be used to predict a nutrient requirement.

175 References

176 Brigelius-Flohé R. 1999. Tissue-specific functions of individual glutathione
177 peroxidases. *Free Radic Biol Med* **27**:951-965. doi:10.1016/s0891-
178 5849(99)00173-2.

179 Burk RF, Hill KE. 2009. Selenoprotein P - Expression, functions, and roles in
180 mammals. *Biochim Biophys Acta* **1790**:1441-1447.
181 doi:10.1016/j.bbagen.2009.03.026.

182 Burk RF, Hill KE, Olson GE, Weeber EJ, Motley AK, Winfrey VP, Austin LM.
183 2007. Deletion of apolipoprotein E receptor-2 in mice lowers brain selenium
184 and causes severe neurological dysfunction and death when a low-selenium
185 diet is fed. *J Neurosci* **27**:6207-11. doi:10.1523/JNEUROSCI.1153-07.2007.

186 Burk RF, Olson GE, Hill KE, Winfrey VP, Motley AK, Kurokawa S. 2013.
187 Maternal-fetal transfer of selenium in the mouse. *FASEB J* **27**:3249-3256.
188 doi:10.1096/fj.13-231852.

189 Chiu-Ugalde J, Theilig F, Behrends T, Drebes J, Sieland C, Subbarayal P, Köhrle
190 J, Hammes A, Schomburg L, Schweizer U. 2010. Mutation of megalin leads to

191 urinary loss of selenoprotein P and selenium deficiency in serum, liver,
192 kidneys and brain. *Biochem J* **431**:103-111. doi:10.1042/bj20100779.

193 FAO. 1992. *Status of Cadmium, Lead, Cobalt and Selenium in Soils and Plants*
194 *of Thirty Countries*. Forssa, Finland.

195 Gatlin DM, III, Wilson RP. 1984. Dietary selenium requirement of fingerling
196 channel catfish. *J Nutr* **114**:627-633.

197 Han D, Xie S, Liu M, Xiao X, Liu H, Zhu X, Yang Y. 2011. The effects of dietary
198 selenium on growth performances, oxidative stress and tissue selenium
199 concentration of gibel carp (*Carassius auratus gibelio*). *Aquaculture Nutr*
200 **17**:e741-e749. doi:10.1111/j.1365-2095.2010.00841.x.

201 Hao X, Ling Q, Hong F. 2014. Effects of dietary selenium on the pathological
202 changes and oxidative stress in loach (*Paramisgurnus dabryanus*). *Fish*
203 *Physiol Biochem* doi:10.1007/s10695-014-9926-7.

204 Hill K, Wu S, Motley A, Stevenson T, Winfrey V, Capecchi M, Atkins J, Burk R.
205 2012. Production of selenoprotein P (Sepp1) by hepatocytes is central to
206 selenium homeostasis. *J Biol Chem* **287**:40414-40424.
207 doi:10.1074/jbc.M112.421404.

208 Hill KE, Xia Y, Akesson B, Boeglin ME, Burk RF. 1996. Selenoprotein P
209 concentration in plasma is an index of selenium status in selenium-deficient
210 and selenium-supplemented Chinese subjects. *J Nutr* **126**:138-145.

211 Hill KE, Zhou J, Austin LM, Motley AK, Ham AJL, Olson GE, Atkins JF, Gesteland
212 RF, Burk RF. 2007. The selenium-rich C-terminal domain of mouse
213 selenoprotein P is necessary for the supply of selenium to brain and testis but
214 not for the maintenance of whole body selenium. *J Biol Chem*
215 doi:10.1074/jbc.M700436200.

216 Hilton JW, Hodson PV, Slinger SJ. 1980. The requirement and toxicity of
217 selenium in rainbow trout (*Salmo gairdneri*). *J Nutr* **110**:2527-2535.

218 Hondal RJ, Ma S, Caprioli RM, Hill KE, Burk RF. 2001. Heparin-binding histidine
219 and lysine residues of rat selenoprotein P. *J Biol Chem* **276**:15823-15831.
220 doi:10.1074/jbc.M010405200.

221 Jensen C, Pallauf J. 2008. Estimation of the selenium requirement of growing
222 guinea pigs (*Cavia porcellus*). *J Anim Physiol Anim Nutr* **92**:481-491.
223 doi:10.1111/j.1439-0396.2007.00738.x.

224 Kasaikina MV, Lobanov AV, Malinouski MY, Lee BC, Seravalli J, Fomenko DE,
225 Turanov AA, Finney L, Vogt S, Park TJ, Miller RA, Hatfield DL, Gladyshev VN.
226 2011. Reduced utilization of selenium by naked mole rats due to a specific

227 defect in Gpx1 expression. *J Biol Chem* **286**:17005-17014.
228 doi:10.1074/jbc.M110.216267.

229 Kato T, Read R, Rozga J, Burk RF. 1992. Evidence for intestinal release of
230 absorbed selenium in a form with high hepatic extraction. *Am J Physiol*
231 **262**:G854-G858.

232 Kim M-SS, Pinto SM, Getnet D, Nirujogi RS, Manda SS, Chaerkady R,
233 Madugundu AK, Kelkar DS, Isserlin R, Jain S, Thomas JK, Muthusamy B, Leal-
234 Rojas P, Kumar P, Sahasrabuddhe NA, Balakrishnan L, Advani J, George B,
235 Renuse S, Selvan LD, Patil AH, Nanjappa V, Radhakrishnan A, Prasad S,
236 Subbannayya T, Raju R, Kumar M, Sreenivasamurthy SK, Marimuthu A, Sathe
237 GJ, Chavan S, Datta KK, Subbannayya Y, Sahu A, Yelamanchi SD, Jayaram S,
238 Rajagopalan P, Sharma J, Murthy KR, Syed N, Goel R, Khan AA, Ahmad S, Dey
239 G, Mudgal K, Chatterjee A, Huang T-CC, Zhong J, Wu X, Shaw PG, Freed D,
240 Zahari MS, Mukherjee KK, Shankar S, Mahadevan A, Lam H, Mitchell CJ,
241 Shankar SK, Satishchandra P, Schroeder JT, Sirdeshmukh R, Maitra A, Leach
242 SD, Drake CG, Halushka MK, Prasad TS, Hruban RH, Kerr CL, Bader GD,
243 Iacobuzio-Donahue CA, Gowda H, Pandey A. 2014. A draft map of the human
244 proteome. *Nature* **509**:575-581. doi:10.1038/nature13302.

245 Kryukov GV, Gladyshev VN. 2000. Selenium metabolism in zebrafish:
246 multiplicity of selenoprotein genes and expression of a protein containing 17
247 selenocysteine residues. *Genes Cells* **5**:1049-1060. doi:10.1046/j.1365-
248 2443.2000.00392.x.

249 Kurokawa S, Eriksson S, Rose KL, Wu S, Motley AK, Hill S, Winfrey VP,
250 McDonald WH, Capecchi MR, Atkins JF, Arnér ES, Hill KE, Burk RF. 2014.
251 Sepp1(UF) forms are N-terminal selenoprotein P truncations that have
252 peroxidase activity when coupled with thioredoxin reductase-1. *Free Radical*
253 *Bio Med* **69**:67-76. doi:10.1016/j.freeradbiomed.2014.01.010.

254 Le KT, Fotedar R. 2013. Dietary selenium requirement of yellowtail kingfish
255 (*Seriola lalandi*). *Agri Sci* **4**:68-75. doi:10.4236/as.2013.46A011.

256 Lei XG, Dann HM, Ross DA, Cheng WH, Combs GF, Roneker KR. 1998. Dietary
257 selenium supplementation is required to support full expression of three
258 selenium-dependent glutathione peroxidases in various tissues of weanling
259 pigs. *J Nutr* **128**:130-135.

260 Lin YH, Shiau SY. 2005. Dietary selenium requirements of juvenile grouper,
261 *Epinephelus malabaricus*. *Aquaculture* **250**:356-363.
262 doi:10.1016/j.aquaculture.2005.03.022.

263 Liu K, Wang XJ, Ai QH, Mai KS, Zhang WB. 2010. Dietary selenium
264 requirement for juvenile cobia, *Rachycentron canadum* L. *Aquaculture Res*
265 **41**:e594-e601. doi:10.1111/j.1365-2109.2010.02562.x.

266 Lobanov A, Hatfield D, Gladyshev V. 2008. Reduced reliance on the trace
267 element selenium during evolution of mammals. *Genome Biol* **9**:R62.
268 doi:10.1186/gb-2008-9-3-r62.

269 Ma S, Hill KE, Caprioli RM, Burk RF. 2002. Mass spectrometric characterization
270 of full-length rat selenoprotein P and three isoforms shortened at the C
271 terminus. Evidence that three UGA codons in the mRNA open reading frame
272 have alternative functions of specifying selenocysteine insertion or
273 translation termination. *J Biol Chem* **277**:12749-12754.
274 doi:10.1074/jbc.M111462200.

275 Mariotti M, Lobanov AV, Guigo R, Gladyshev VN. 2013. SECISearch3 and
276 Seblastian: new tools for prediction of SECIS elements and selenoproteins.
277 *Nucleic Acids Res* **41**:e149. doi:10.1093/nar/gkt550.

278 NRC. 1963. Nutrient requirements of beef cattle. National Academy Press,
279 Washington DC.

280 NRC. 1985. Nutrient requirements of sheep. National Academy Press,
281 Washington DC.

282 NRC. 1995. Nutrient requirements of laboratory animals. National Academy
283 Press, Washington DC.

284 NRC. 1997. Nutrient requirements of horses. National Academy Press,
285 Washington DC.

286 NRC. 2011. Nutrient requirements of fish and shrimp. National Academy
287 Press, Washington DC.

288 Olson GE, Winfrey VP, Hill KE, Burk RF. 2008. Megalin mediates selenoprotein
289 P uptake by kidney proximal tubule epithelial cells. *J Biol Chem* **283**:6854-
290 6860. doi:10.1074/jbc.M709945200.

291 Olson GE, Winfrey VP, NagDas SK, Hill KE, Burk RF. 2007. Apolipoprotein E
292 receptor-2 (ApoER2) mediates selenium uptake from selenoprotein P by the
293 mouse testis. *J Biol Chem* **282**:12290-12297. doi:10.1074/jbc.M611403200.

294 Papp LV, Lu J, Holmgren A, Khanna KK. 2007. From selenium to
295 selenoproteins: synthesis, identity, and their role in human health. *Antioxid*
296 *Redox Signal* **9**:775-806. doi:10.1089/ars.2007.1528.

297 Penglase S, Hamre K, Rasinger JD, Ellingsen S. 2014. Selenium status affects
298 selenoprotein expression, reproduction, and F1 generation locomotor activity
299 in zebrafish (*Danio rerio*). *Brit J Nutr* **111**:1918-1931.
300 doi:10.1017/S000711451300439X.

301 Rachel H, Edwin WPS, Scott DY, Allan DCC, Edward JMJ, Colin RB, Ander EL,
302 Michael JW, Benson C, Jellita G, Dalitso Ko, Alexander JS, Susan JF-T, Rosalind
303 SG, Alexander AK, Martin RB. 2013. Soil-type influences human selenium
304 status and underlies widespread selenium deficiency risks in Malawi. *Sci Rep*
305 doi:10.1038/srep01425.

306 Read R, Bellew T, Yang JG, Hill KE, Palmer IS, Burk RF. 1990. Selenium and
307 amino acid composition of selenoprotein P, the major selenoprotein in rat
308 serum. *J Biol Chem* **265**:17899-17905.

309 Romagné F, Santesmasses D, White L, Sarangi GK, Mariotti M, Hübler R,
310 Weihmann A, Parra G, Gladyshev VN, Guigó R, Castellano S. 2014. SelenoDB
311 2.0: annotation of selenoprotein genes in animals and their genetic diversity
312 in humans. *Nucleic Acids Res* **42**:D437-43. doi:10.1093/nar/gkt1045.

313 Saito Y, Sato N, Hirashima M, Takebe G, Nagasawa S, Takahashi K. 2004.
314 Domain structure of bi-functional selenoprotein P. *Biochem J* **381**:841-846.
315 doi:10.1042/BJ20040328.

316 Shetty SP, Shah R, Copeland PR. 2014. Regulation of selenocysteine
317 incorporation into the selenium transport protein, selenoprotein P. *J Biol*
318 *Chem* **289**:25317-25326. doi:10.1074/jbc.M114.590430.

319 Steinbrenner H, Bilgic E, Alili L, Sies H, Brenneisen P. 2006. Selenoprotein P
320 protects endothelial cells from oxidative damage by stimulation of
321 glutathione peroxidase expression and activity, **40**:936-943.
322 doi:10.1080/10715760600806248.

323 Sunde RA, Raines AM, Barnes KM, Evenson JK. 2009. Selenium status highly
324 regulates selenoprotein mRNA levels for only a subset of the selenoproteins

325 in the selenoproteome *Bioscience Rep* **29**:329-338.
326 doi:10.1042/BSR20080146.

327 Wedekind KJ, Yu S, Combs GF. 2004. The selenium requirement of the puppy. *J*
328 *Anim Physiol Anim Nutr* **88**:340-347. doi:10.1111/j.1439-0396.2004.00489.x.

329 Weiss Sachdev S, Sunde R. 2001. Selenium regulation of transcript
330 abundance and translational efficiency of glutathione peroxidase-1 and -4 in
331 rat liver. *Biochem J* **357**:851-858.

332 Weiss SL, Evenson JK, Thompson KM, Sunde RA. 1996. The selenium
333 requirement for glutathione peroxidase mRNA level is half of the selenium
334 requirement for glutathione peroxidase activity in female rats. *J Nutr*
335 **126**:2260-2267.

336 Weiss SL, Evenson JK, Thompson KM, Sunde RA. 1997. Dietary selenium
337 regulation of glutathione peroxidase mRNA and other selenium-dependent
338 parameters in male rats. *J Nutr Biochem* **8**:85-91. doi:10.1016/S0955-
339 2863(96)00178-7.

Table 1 (on next page)

The Se requirements (mg Se/kg DM) predicted by the model (Figure 2) versus those determined in feeding trials for vertebrates as the selenocysteine content (Sec no.) of selenoprotein P (SEPP1) increases.

Predicted Se requirement data are mean $\pm$ 95% confidence interval. ¹ mg Se/kg feed DM, mean ($\pm$ 95% confidence interval, when shown) ² Further information on the representative species used for the determined Se requirement data can be found in Supp. Table 1. ³ There are currently no known species with full length SEPP1 containing 6 Sec residues.

Table 1. Table of the Se requirements (mg Se/kg DM) predicted by the model (Figure 2) versus those determined in feeding trials for vertebrates as the selenocysteine content (Sec no.) of selenoprotein P (SEPP1) increases. Predicted Se requirement data are mean $\pm$ 95% confidence interval.

Class	Sec no.	Predicted Se requirement ¹	Determined Se requirement ¹	Representative species ²
? ³	6	0.02 $\pm$ 0.03	ND	-
Mammals	7	0.04 $\pm$ 0.03	0.06	Guinea pig
	8	0.05 $\pm$ 0.02	ND	-
	9	0.07 $\pm$ 0.02	ND	-
	10	0.08 $\pm$ 0.02	0.10	Rat
	11	0.10 $\pm$ 0.02	ND	-
	12	0.12 $\pm$ 0.03	0.10	Cow
	13	0.14 $\pm$ 0.04	0.10	Horse
	14	0.17 $\pm$ 0.05	0.20	Pig
	15	0.20 $\pm$ 0.06	0.21	Dog
Bony	16	0.24 $\pm$ 0.06	0.25	Catfish
fish	17+	>0.24	0.30	Zebrafish

¹ mg Se/kg feed DM, mean ($\pm$ 95% confidence interval, when shown)

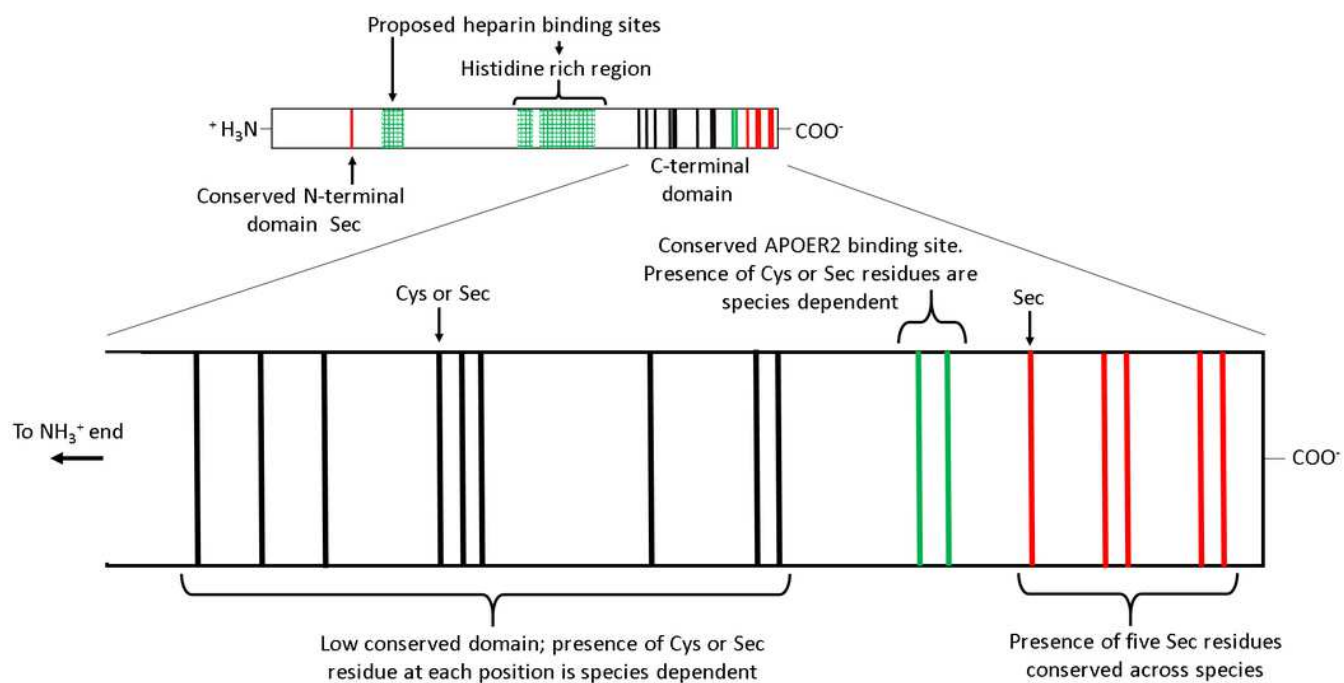
² Further information on the representative species used for the determined Se requirement

data can be found in Supp. Table 1.

³ There are currently no known species with full length SEPP1 containing 6 Sec residues.

The receptor binding sites and selenocysteine (Sec) residues of vertebrate selenoprotein P (SEPP1)

From the N-terminal side, SEPP1 is comprised of a conserved N-terminal domain Sec residue, followed by several proposed heparin binding sites which include a histidine rich region. Following this, there is the shorter Sec residue rich C-terminal domain which contains an APOER2 binding site. The C-terminal domain can be further divided into two subdomains. The first subdomain exists on the N-terminal side of the APOER2 binding site and contains a region with a low conservation of Sec residues among vertebrates (mainly due to Sec to cysteine (Cys) conversions (Lobanov et al., 2008)). The second subdomain is located downstream of the APOER2 binding site and contains five Sec residues that are conserved across vertebrate species. Several species of amphibians also have an additional Sec residue in the C-terminal end of this region (Lobanov et al., 2008) . The proposed heparin binding sites/histidine rich regions are based on rat SEPP1 found by Hondal et al. (2001) . Similar histidine rich regions are found in the SEPP1's of other species (selenodb.org). Cys residues outside the C-terminal domain are not shown. Red lines = conserved Sec residues; Black lines = Cys or Sec residues; Green lines = Cys/Sec residues within the APOER2 binding site; Green box grids = proposed heparin binding sites.



The relationship between the selenocysteine content of selenoprotein P and selenium requirements in vertebrates.

The solid line with the solid circles (●) is the best fit model for the SEPP1 Sec content versus Se requirements (mg Se/kg dry matter (DM)) from 11 species with representatives from the mammalian and bony fish classes where the genome sequences were available (Second order polynomial, $R^2 = 0.88$, $y = 4.3 + 78x - 122x^2$). The broken line represents the same data modelled with an additional five bony fish species with known Se requirement levels (○), but unannotated genomes. SEPP1 Sec content in these fish were assumed to be within the likely range of 15-17 Sec residues found for fish in general (5PL Asymmetric sigmoidal, $R^2 = 0.92$, $y = 5.13 + (11.9 / ((1 + 10^{(-1.6410 - X) \times 6.391}))^{9.611^{10}}))$). Shaded boxes group animals within classes. The X axis is log transformed.

