- **1** Ophiuroid phylotranscriptomics enables discovery of novel echinoderm representatives
- 2 of bilaterian neuropeptide families and reconstruction of neuropeptide precursor
- 3 evolution over ~270 million years.
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#### 27 Abstract

#### 28 Background:

Neuropeptides are a diverse class of intercellular signaling molecules that mediate neuronal regulation of many physiological and behavioural processes, including feeding, reproduction and locomotion. Recent advances in genome/transcriptome sequencing are enabling identification of neuropeptide precursor proteins in species from a growing variety of animal taxa, providing new insights into the evolution of neuropeptide signaling. Here we report a phylo-transcriptomic analysis of neuropeptide precursors in over fifty species of brittle stars (Class Ophiuroidea; Phylum Echinodermata).

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#### 37 **Results:**

38 Detailed analysis of transcriptome sequence data from three brittle star species, 39 Ophionotus victoriae, Amphiura filiformis and Ophiopsila aranea, enabled the first 40 comprehensive identification of neuropeptide precursors in ophiuroids. Representatives of 41 over thirty bilaterian neuropeptide precursor families were identified, some of which occur as 42 paralogs thyrotropin-releasing hormone, (e.g. corticotropin-releasing hormone, 43 cholecystokinin, somatostatin and pedal peptide). Furthermore, homologs of 44 endothelin/CCHamide, eclosion hormone, neuropeptide-F/Y and nucleobinin/nesfatin were discovered here in a deuterostome/echinoderm for the first time. The majority of ophiuroid 45 46 neuropeptide precursors contain a single copy of a neuropeptide, but several precursors 47 comprise multiple copies of identical or non-identical, but structurally-related, neuropeptides. 48 Here we performed an unprecedented investigation of the evolution of neuropeptide copy-49 number over a period of  $\sim 270$  million years by analysing sequence data from over fifty 50 ophiuroid species, with reference to a robust phylogeny. Interestingly, the number of 51 neuropeptide copies in the majority of precursors was constant across all the species 52 examined, but examples of clade-specific losses/gains of neuropeptides were also observed.

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#### 54 **Conclusions:**

We report here the most comprehensive analysis to date of neuropeptide precursors in the phylum Echinodermata, with novel representatives of several bilaterian neuropeptide families discovered for the first time in echinoderms. Furthermore, analysis of precursor proteins comprising multiple copies of identical or related neuropeptides across ~270 million years of ophiuroid evolution indicates that the composition of neuropeptide "cocktails" is functionally important, but with plasticity over long evolutionary time scales.

61

# 62 Keywords (3 to 10):

- 63 Neuropeptide; echinoderm; Ophiuroidea; eclosion hormone; CCHamide; neuropeptide-Y;
- 64 evolution
- 65

#### 66 Introduction

67 The nervous systems of animals utilize a wide variety of chemicals for neuronal 68 communication. These include amino acids (e.g. glutamate), biogenic amines (e.g. serotonin), 69 and neuropeptides (e.g. vasopressin) amongst others. Neuropeptides are by far the most-70 diverse and they control many physiological/behavioural processes, including feeding, 71 reproduction and locomotion [1-3]. Recent advances in genome/transcriptome sequencing are 72 enabling identification of neuropeptide precursor proteins in species from a growing variety 73 of animal taxa, providing new insights into the evolution of neuropeptide signaling [4-8]. The 74 echinoderms are notable in this regard because as deuterostomian invertebrates they occupy 75 an "intermediate" phylogenetic position with respect to the vertebrates and intensely studied 76 protostomian invertebrates such as insects (e.g. Drosophila melanogaster) and nematodes 77 (e.g. Caenorhabditis elegans). Accordingly, characterisation of neuropeptide signaling 78 systems in echinoderms has recently provided key "missing links" for determination of 79 neuropeptide relationships and reconstruction of neuropeptide evolution [8-10].

80 The phylum Echinodermata comprises five extant classes: Echinoidea (sea urchins 81 and sand dollars), Holothuroidea (sea cucumbers), Asteroidea (starfish), Ophiuroidea (brittle 82 stars and basket stars) and Crinoidea (sea lilies and feather stars). Recent molecular 83 phylogenetic studies support the hypothesis that Echinoidea and Holothuroidea are sister 84 groups (Echinozoa) and Asteroidea and Ophiuroidea are sister groups (Asterozoa), with the 85 Crinoidea basal to the Echinozoa + Asterozoa clade (Eleutherozoa) [11, 12]. Echinoderms 86 are marine organisms that have several unique features including pentaradial symmetry as 87 adults, a remarkable ability to autotomise and regenerate body parts, and neurally-controlled 88 mutable collagenous tissue [13, 14]. Previous transcriptomic analyses have identified 89 neuropeptide precursor complements in Strongylocentrotus purpuratus (purple sea urchin), 90 Apostichopus japonicus (Japanese sea cucumber) and Asterias rubens (common European 91 starfish) [8, 15, 16]. Furthermore, the identification of neuropeptides in these species has 92 facilitated investigation of the evolution and physiological roles of various neuropeptide 93 signaling systems [8-10, 17-21].

The recent progress in transcriptomic/genomic characterization of echinoderm neuropeptide systems has hitherto not been extended to ophiuroids or crinoids. The Ophiuroidea constitutes the largest class among extant echinoderms [22] with a long evolutionary history that extends back to the early Ordovician (around 480 million years ago) [23], whilst extant families date from the mid-Permian (~ 270 million years ago) [12]. Available molecular data for ophiuroids has increased significantly in recent years with the emergence of numerous transcriptomic studies [20, 24-29]. Here, we utilize transcriptome

sequence data from three brittle star species, *Ophionotus victoriae, Amphiura filiformis* and *Ophiopsila aranea* to perform the first comprehensive identification of neuropeptide precursors in ophiuroids. We identify representatives of over thirty neuropeptide families including homologs of endothelin/CCHamide, eclosion hormone (EH), neuropeptide-F/Y (NPF/NPY) and nucleobinin (NUCB)/nesfatin, which are the first to be discovered in a deuterostome/echinoderm.

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108 Transcriptomes have also been employed to investigate the phylogenetic relationships 109 of the ophiuroids, utilising data from fifty-two species [12]. In this the most comprehensive 110 molecular analysis of ophiuroid phylogeny to date, previous morphology-based classification 111 schemes [30] were rejected in favour of a new phylogeny comprising three primary ophiuroid 112 clades [12, 31, 32]. This landmark study and the associated large dataset has provided a 113 unique opportunity to investigate the conservation and diversification of neuropeptide 114 precursor structure over a period of ~270 million years of ophiuroid evolution. Our analysis 115 reveals that the majority of ophiuroid neuropeptide precursors contain a single copy of a 116 neuropeptide, but several precursors comprise multiple copies of identical or non-identical, 117 but structurally-related, neuropeptides. Interestingly, the number of neuropeptide copies in 118 the majority of precursors is constant across all the ophiuroid species examined, but examples 119 of clade-specific losses/gains of neuropeptides are also observed. This remarkable 120 conservation in neuropeptide copy number across ~270 million years of ophiuroid evolution 121 indicates that the composition of neuropeptide "cocktails" is functionally important, but with 122 plasticity over long evolutionary time scales.

#### 123 Results and discussion

124 Here we have identified ophiuroid homologs of neuropeptide precursors that have 125 been identified previously in other echinoderms and these include, alphabetically: AN 126 peptides, bursicon ( $\alpha$  and  $\beta$ ), calcitonin, cholecystokinin (CCK), corazonin, corticotropin-127 releasing hormone (CRH), glycoprotein hormones ( $\alpha 2$  and  $\beta 5$ ), gonadotropin-releasing 128 hormone (GnRH), insulin-like peptide (ILP), kisspeptin (KP), lugin, melanin-concentrating 129 hormone (MCH), NG peptides (neuropeptide-S), orexin, pedal peptides, pigment-dispersing 130 factor (PDF), relaxin-like peptide, SALMFamides (L-type and F-type), somatostatin, 131 tachykinin, thyrotropin-releasing hormone (TRH) and vasopressin/oxytocin. Identification of 132 ophiuroid representatives of these neuropeptide precursor types has in some cases provided 133 new insights into neuropeptide precursor structure and evolution, as discussed in more detail 134 below. First, however, we will highlight representatives of bilaterian neuropeptide precursor 135 families that have been identified here for the first time in an echinoderm species.

136

#### 137 Discovery of the first echinoderm representatives of bilaterian neuropeptide families

Comprehensive analysis of transcriptome sequence data from three ophiuroid species, *O. victoriae, A. filiformis and O. aranea,* has enabled the discovery of the first echinoderm representatives of four bilaterian neuropeptide families. Specifically, we have discovered the first deuterostomian homologs of eclosion hormone (**Figure 2**), the first ambulacrarian homolog of CCHamide/endothelin-type peptides (**Figure 3A**), and the first echinoderm homologs of neuropeptide-Y/neuropeptide-F (**Figure 3B**) and NUCB/nesfatin (**Figure S1**), as discussed in detail below.

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## 146 <u>Eclosion hormone</u>

147 Eclosion hormone (EH) was first isolated and sequenced in the insects Manduca sexta 148 (tobacco hornworm) and *Bombyx mori* (silk moth) and shown to alter the timing of adult 149 emergence [33, 34]. EH is one of the main peptide/protein hormones involved in control of 150 ecdysis (*i.e.* shedding of the cuticle) behavior in insects [35]. It binds to and activates a 151 receptor guanylyl cyclase that is expressed in epitracheal Inka cells and causes the secondary 152 release of ecdysis-triggering hormone (ETH) that is also expressed in Inka cells [36, 37]. In 153 Drosophila, EH is important but not essential for ecdysis as some flies lacking EH are able to 154 undergo ecdysis [38]. Insect EHs have six conserved cysteine residues that form three 155 disulfide bridges [36]. EHs have not been discovered previously outside of arthropods. 156 Interestingly, four EH-like precursors were identified in A. *filiformis* and O. aranea and two 157 in O. victoriae (Figure S2-S4). The ophiuroid EH-like precursors are orthologous to

158 neuropeptide precursors previously identified in the sea-urchin S. purpuratus (Spnp11 and 159 Spnp15, which we now rename as Spur EH1 and Spur EH2, respectively) [16] and the 160 starfish A. rubens (Arnp11, Arnp15 and Arnp15b renamed as Arub EH1, Arub EH2a and 161 Arub EH2b, respectively) [8]. The positions of cysteine residues are conserved across all 162 echinoderm and insect EHs, but aside from this there is little sequence conservation (Figure 163 2A). The echinoderm EH-like precursor sequences were also analysed using a sequence-164 similarity-based clustering approach based on BLASTp e-values using CLANS software 165 [39]. The analysis shows that echinoderm EH-like precursors (i) cluster in two compact 166 subgroups (echinoderm EH-like precursor 1 and EH-like precursor 2 and (ii) have strong 167 positive BLAST results with arthropod EHs and, to a lesser extent, with arthropod ion 168 transport peptide (ITP) and vertebrate atrial natriuretic peptide (ANP) (Figure 2B). ITP 169 precursors also possess six cysteine residues; however, the position of these residues is not 170 conserved with cysteine residues found in echinoderm EH-like precursors (not shown).

171 To obtain further evidence for the presence of an EH-like signaling system in 172 echinoderms, we performed a phylogenetic analysis of EH-type receptors. Insect EHs 173 mediate their effects by binding to membrane guanylyl cyclase receptors [37]. EH receptors 174 are closely related to vertebrate ANP receptors and various orphan receptors [40]. Specific 175 BLAST searches enabled identification of transcripts in O. victoriae, A. filiformis and O. 176 aranea that encode proteins similar to arthropod EH receptors. Maximum likelihood and 177 Bayesian phylogenetic analyses confirmed that these sequences group with the receptor 178 cluster containing EH receptors (Figure 2C). The discovery of the first deuterostomian EHs 179 suggests an ancient bilaterian origin of EHs and indicates that these hormones may have other 180 functions in invertebrates aside from their role in ecdysis.

181

#### 182 <u>CCHamide</u>

183 CCHamides are neuropeptides that were discovered relatively recently in the 184 silkworm *Bombyx mori* [41]. Later, it was found that insects have two CCHamide genes, 185 CCHamide-1 and CCHamide-2, each encoding a single copy of the mature peptide [42]. 186 These peptides are referred to as CCHamides because they contain two cysteine residues and 187 a characteristic histidine-amide C-terminal motif. There are two CCHamide receptors in 188 insects: CCHamide-1 specifically activates one receptor and CCHamide-2 specifically 189 activates the second receptor [42, 43]. CCHamide-1 has a physiological a role in starvation-190 induced olfactory modifications [44] whereas as CCHamide-2 regulates feeding, growth and 191 developmental timing in flies [43, 45]. Recent studies examining the evolution of 192 neuropeptides in the Bilateria have shown that protostomian CCHamides are related to 193 elevenin (another protostomian neuropeptide originally discovered from the mollusc Aplysia 194 californica L11 neuron), lophotrochozoan GGNG peptides, endothelins and gastrin-releasing 195 peptides (GRPs) [6, 7, 46, 47]. The latter two are neuropeptide types that have not been found 196 outside chordates. Furthermore, the degree of sequence/structural conservation varies across 197 these different peptide families. Hence, CCHamides are amidated and have a disulphide 198 bridge, elevenins and endothelins have a disulphide bridge but are non-amidated and GRPs 199 are amidated but lack the disulphide bridge. Furthermore, CCHamide-1 is located 200 immediately after the signal peptide whereas there is a dibasic cleavage site separating the 201 signal peptide and CCHamide-2 [42].

202 Here we have identified two neuropeptide precursors in brittle stars whose sequence 203 and precursor structure resembles those of lophotrochozoan GGNG peptides and insect 204 CCHamide-1 (Figure 3A). The CCHamide-like precursor 1 identified in O. victoriae is 205 orthologous to an uncharacterized neuropeptide precursor (Arnp25) identified previously in 206 the starfish A. rubens [8], whereas the CCHamide-like precursor 2 was only found in brittle 207 stars. Both CCHamide-like precursors in O. victoriae comprise a single copy of a putative 208 cyclic amidated peptide that is flanked by a signal peptide at the N-terminus and a dibasic 209 cleavage site at the C-terminus. Interestingly, both of these peptides lack a penultimate 210 histidine residue, just like the lophotrochozoan GGNG peptides (Figure 3A) [46, 47].

211

#### 212 <u>Neuropeptide-Y/Neuropeptide-F</u>

213 Neuropeptide-Y (NPY) was first isolated and sequenced from the porcine 214 hypothalamus in 1982 [48, 49]. Although the NPY/NPF family of peptides are pleiotropic in 215 nature [50], they are mainly known for their roles in regulation of feeding and stress [3, 51, 216 52]. The discovery of Neuropeptide-F (NPF) in the tapeworm Monieza expansa in 1991 217 demonstrated for the first time the occurrence of NPY homologs in invertebrates [53]. Here, 218 we have identified the first echinoderm representatives of the NPY/NPF family in brittle stars 219 and starfish (Figure 3B). The brittle star precursors contain a peptide with a C-terminal 220 RYamide, in common with NPY in vertebrates and an ortholog in the starfish Patiria 221 miniata. In contrast, an ortholog in the starfish A. rubens has a C-terminal RFamide, a feature 222 that it shares with NPY/NPF-type peptides in the hemichordate S. kowalevskii and in 223 protostomes. Thus, our findings have revealed that NPY/NPF-type peptides with a C-terminal 224 Yamide motif are not restricted to vertebrates. Echinoderm NPY/NPF-type peptides are 225 located immediately after the signal peptide in the precursor proteins, as is the case in other 226 bilaterian species. Surprisingly, we did not find NPY/NPF-type precursors in the sea urchin S. 227 purpuratus or the sea cucumber A. japonicus. However, we suspect that this may reflect

sequence divergence rather than gene loss because a gene encoding a NPY/NPF-type receptor

can be found in the *S. purpuratus* genome [54].

230

#### 231 <u>NUCB</u>

Nucleobindins (NUCB1 and NUCB2) are multidomain  $Ca^{2+}$  and DNA binding 232 233 proteins. NUCB1 was first discovered in 1992 and thought to play a role in apoptosis and 234 autoimmunity [55]. Interestingly, the NUCB1 precursor has both a signal peptide and a 235 leucine zipper structure suggesting that it can bind DNA and act as an endocrine factor [56]. 236 NUCB2 is a homolog of NUCB1 and was named based on high sequence similarity between 237 the two precursors [57]. In 2006, an 82 amino acid peptide located in the N-terminal region of 238 NUCB2 was reported. This peptide, Nesfatin-1 (Nucleobindin-2-Encoded Satiety and FAT-239 Influencing proteiN-1), was discovered as a satiety inducing factor in the rat hypothalamus 240 [58]. Its role in inhibiting food intake in vertebrates is now well-established [57, 59]. 241 Moreover, this pleiotropic peptide also modulates other processes including glucose and lipid 242 metabolism, and cardiovascular and reproductive functions. Recently, nesfatin-1-like peptide 243 derived from NUCB1 was shown to be anorexigenic in goldfish [60]. Surprisingly, the 244 presence of NUCBs in invertebrates had not been reported, in spite of the potential 245 therapeutic applications of these molecules in obesity related disorders. Here, we show that 246 NUCB-type precursors are present in echinoderms (Figure S1A). Phylogenetic analysis of 247 NUCB precursors reveals that a single copy of the NUCB precursor is found in invertebrate 248 species and gene duplication in the vertebrate lineage gave rise to NUCB1 and NUCB2 249 (Figure S1B). In chordates, the NUCB precursors are predicted to generate three peptides 250 (Nesfatin-1, 2 and 3); however, no biological role has been attributed specifically to nesfatin-251 2 and nesfatin-3. Interestingly, the prohormone convertase cleavage sites expected to 252 generate Nesfatin-1, 2 and 3 are conserved between echinoderm and chordate NUCBs. 253 Moreover, the O. victoriae precursor has an additional predicted cleavage site within the 254 Nesfatin-1 containing region, which is not present in other species (except for *Drosophila* 255 *melanogaster*). However, it remains to be determined whether or not this cleavage site in the 256 O. victoriae precursor is functional.

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#### 8 First comprehensive identification of neuropeptide precursors in ophiuroids

We have identified neuropeptide precursors belonging to 32 families, which represents the first comprehensive analysis of neuropeptide precursors in ophiuroids (**Figure 4; Figure S2-S4**). Several of these neuropeptide families have been identified previously in echinoderms and include homologs of AN peptides, bursicon ( $\alpha$  and  $\beta$ ), calcitonin, CCK

263 [15], corazonin [10], CRH, glycoprotein hormones ( $\alpha 2$  and  $\beta 5$ ) [61], GnRH [10], ILP [61], 264 KP [8], luqin [7], MCH [8], NG peptides (neuropeptide-S) [9, 62], orexin [6, 8], pedal 265 peptides [16], PDF [8], relaxin-like peptide [63], SALMFamides (L-type and F-type) [19, 20, 266 64], somatostatin [8], tachykinin [8], TRH [16] and vasopressin/oxytocin [61, 62] (Figures 5-267 7 and S5-S9). With the exception of MCH (which may be unique to deuterostomes) [6, 8], 268 AN peptides and SALMFamides (which thus far have only been identified in echinoderms), 269 the origins of all of the neuropeptide precursors identified here in ophiuroids predate the 270 divergence of protostomes and deuterostomes [6, 7]. Of the three species examined here, the 271 neuropeptide precursor complement of O. victoriae was the most complete (Figure 4) and 272 therefore this species is used as a representative ophiuroid for sequence alignments, except in 273 a few cases where a neuropeptide precursor was not found in O. victoriae. Below we 274 highlight several interesting and/or unusual features of ophiuroid neuropeptides and 275 neuropeptide precursors.

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#### 277 Neuropeptide precursors that occur in multiple forms in O. victoriae

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279 Thyrotropin-releasing hormone (TRH)-type precursors

280 TRH (also known as thyrotropin-releasing factor or thyroliberin) was first isolated and 281 sequenced in the 1960s [65-67]. In mammals, TRH is produced in the hypothalamus and 282 stimulates the release of thyroid-stimulating hormone (TSH) and prolactin from the anterior 283 pituitary [68, 69]. The recent discovery of a TRH receptor in the annelid *Platynereis* 284 *dumerilii* indicates that the evolutionary origin of this neuropeptide signaling system predates 285 the divergence of protostomes and deuterostomes [70].

286 The human TRH precursor contains six copies of the tripeptide pQHPamide [71]. 287 Precursor proteins comprising multiple copies of TRH-like peptides have been identified 288 previously in the sea urchin S. purpuratus, the sea cucumber A. japonicus and the starfish A. 289 rubens [8, 15, 16], with a single TRH-type precursor found in each of these species. 290 Interestingly, here we identified two TRH-type precursors (OvTRHP1 and OvTRHP2) in O. 291 victoriae (Figure S2 and 6A). OvTRHP1 comprises 21 copies of putative TRH-like 292 tetrapeptides with the motif pQXXXamide (where X is variable). OvTRHP2, on the other 293 hand, comprises two copies of the putative tetrapeptide pQGPRamide and two longer 294 peptides that also have a C-terminal GPRamide motif but lack the N-terminal pyroglutamate.

295

#### 296 Cholecystokinin (CCK)-type precursors

297 A CCK-type peptide (formerly pancreozymin) was first sequenced in the 1960s [72]. 298 CCK-type peptides play numerous roles in feeding and digestion related physiology. CCK 299 mediates satiety, stimulates the release of digestive enzymes and gall bladder contractions 300 [73-75]. CCK-type peptides are involved in mechanisms of learning and memory, and 301 analgesia [76]. A neuropeptide precursor comprising two CCK-like peptides was recently 302 identified in the starfish A. rubens [8]. Here we have identified two CCK-type precursors in 303 O. victoriae (OvCCKP1 and OvCCKP2) and orthologs of both of these precursors were also 304 identified in the sea urchin S. purpuratus (Figure S2) [16]. The CCK-type precursor 1 305 comprises three CCK-like peptides in both O. victoriae and S. purpuratus and this precursor 306 is similar to the A. rubens CCK-type precursor, which comprises two CCK-like peptides. In 307 contrast, the CCK-type precursor 2 comprises a single CCK-like peptide in both O. victoriae 308 and S. purpuratus. Interestingly, the sequence of the S. purpuratus CCK-type precursor 2 was 309 reported previously as part of a genome-wide search for neuropeptides [77], but the authors 310 of this study did not identify it as a CCK-type precursor. However, based on the presence of a 311 conserved tyrosine residue and a C-terminal F-amide motif in the predicted neuropeptide 312 derived from this protein, it is evident that it belongs to the family of CCK-type precursors 313 (Figure 6B). A search of a preliminary genome assembly of the starfish *Patiria miniata* 314 (http://www.echinobase.org) [78] did not reveal a gene encoding a CCK-type precursor 2. 315 Therefore, it appears that this neuropeptide precursor type may have been lost in the 316 Asteroidea; nevertheless, further analysis of a wider range of starfish species will be required 317 to draw definitive conclusions. With a broader evolutionary perspective, CCK-type peptides 318 in deuterostomes are orthologs of sulfakinin (SK)-type neuropeptides found in insects [6, 7]. 319 Interestingly, insects have a single SK precursor, which comprises two neuropeptides, SK-1 320 and SK-2 [79], and this may reflect the ancestral condition in the common ancestor of 321 protostomes and deuterostomes. Thus, the occurrence of two CCK-type peptides on a single 322 precursor in A. rubens and insects may be an ancestral characteristic and the occurrence of 323 two CCK-type precursors that comprise one and three CCK-type peptides appears to be a 324 derived characteristic.

325

#### 326 <u>Somatostatin-type precursors</u>

Somatostatin was first isolated and sequenced from sheep hypothalamus in 1973 [80]. This peptide inhibits the release of pituitary hormones such as growth hormone, prolactin and thyroid-stimulating hormone [81]. Moreover, it also inhibits the release of gastrointestinal (cholecystokinin and gastrin amongst others) and pancreatic (insulin and glucagon) hormones [82-84]. Aside from its effects on release of hormones, somatostatin also has central actions

332 that influence motor activity [82]. Here, we have identified two somatostatin-type precursors 333 (OvSSP-1 and OvSSP-2) in O. victoriae. (Figure S2 and 6C). Homologs of both of these 334 precursors are present in the sea urchin S. purpuratus (Figure S2 and 6C), one of which was 335 previously referred to as Spnp16 [16]. By comparison, only a single somatostatin-type 336 precursor has been found in the starfish A. rubens, which is an ortholog of OvSSP-1 [8]. All 337 somatostatin-type precursors comprise a single copy of the bioactive neuropeptide, which is 338 located in the C-terminal region of the precursor [85, 86]. Interestingly, the type-1 339 somatostatins in echinoderms have a phenylalanine residue located in the middle part of the 340 peptide and this conserved feature is found in human somatostatin. Conversely, type-2 341 somatostatins in echinoderms lack the phenylalanine residue but have a neighbouring 342 tryptophan-lysine (WK) motif that is also conserved in human and B. floridae somatostatins 343 (Figure 6C). The deuterostomian somatostatins are orthologous to the allatostatin-C 344 neuropeptide family in arthropods [7]. This family of peptides comprises three precursor-345 types: allatostatin-C, allatostatin-CC and the recently discovered allatostatin-CCC [86, 87]. 346 Both allatostatin-C and allatostatin-CC are non-amidated, like somatostatins; however, 347 allatostatin-CCC has a C-terminal amide. Hence, non-amidated peptides may be 348 representative of the ancestral condition in the common ancestor of protostomes and 349 deuterostomes, with the amidated allatostatin-CCC probably having evolved only within the 350 arthropod lineage [87]. It remains to be determined whether or not the duplication of 351 somatostatin-type precursors in echinoderms and the duplication of allatostatin C (to give rise 352 to allatostatin-CC) represent independent duplications. Further insights into this issue may be 353 obtained if the receptors for somatostatin-type peptides in echinoderms are deorphanised.

354

#### 355 <u>Corticotropin-releasing hormone (CRH)-type precursors</u>

356 CRH-type peptides are a family of related neuropeptides that include CRH, urocortins 357 and urotensin-I in chordates, egg-laying hormone (ELH) in lophotrochozoans and diuretic 358 hormone 44 (DH<sub>44</sub>) in arthropods [6, 7]. Arthropods usually have a single DH<sub>44</sub> precursor, 359 which comprises a single copy of the mature peptide. In some insects, such as *Tribolium* 360 *castaneum* and *Bombyx mori*, alternative splicing of DH<sub>44</sub> transcripts results in multiple 361 mature peptide isoforms of varying lengths [41, 88]. The situation in lophotrochozoans is 362 more complex, with several species having multiple precursors and some of these precursors 363 comprising multiple ELH mature peptides [4, 89]. A single CRH-type precursor was found 364 previously in the starfish A. rubens, whereas here we have identified four CRH-type 365 precursors in O. victoriae (Figure S2 and 6D). Thus, expanded families of CRH-type

366 peptides and receptors appear to have evolved independently in multiple animal lineages,

including chordates and ophiuroid echinoderms [90, 91].

368

#### 369 Diversity in neuropeptide precursor structure: new insights from ophiuroids

370

#### 371 <u>Tachykinins</u>

372 The mammalian neuropeptide substance P was the first tachykinin-type peptide to be 373 isolated and sequenced [92-94]. Subsequently, tachykinin-type peptides were discovered in 374 other animals including tunicates [95], insects [96, 97], annelids [98] and molluscs [99]. 375 Tachykinin-type peptides regulate various physiological processes including muscle 376 contractility [100], nociception [101] and stress responses [102] amongst others [103]. 377 Analysis of genomic/transcriptomic sequence data from the sea urchin S. purpuratus and the 378 sea cucumber A. japonicus did not identify candidate tachykinin-type precursors [6, 7, 15, 379 16]. However, recently a putative tachykinin-type precursor was discovered in the starfish A. 380 rubens (ArTKP), indicating that this signaling system does occur in some echinoderms [8]. 381 Here we have identified orthologs of ArTKP in O. victoriae and other ophiuroids (Figure 4 382 and 7A). Collectively, these findings indicate that this signaling system has been retained in 383 the Asterozoa but lost in the Echinozoa. Comparison of the structure of the asterozoan 384 tachykinin-type precursors reveals that the A. rubens precursor (ArTKP) comprises two 385 putative mature peptides, whereas the O. victoriae precursor comprises four mature peptides 386 (Figure 7B). It remains to be determined, however, which of these two conditions represents 387 the ancestral state in the common ancestor of the Asterozoa. Further insights into this issue 388 may be obtained if sequence data from a variety of starfish species are analysed.

389

#### 390 <u>Kisspeptins (KP)</u>

391 Kisspeptin (formerly known as metastin) is encoded by the KiSSI gene in humans. 392 KiSS1 was originally discovered as a gene that may suppress the metastatic potential of 393 malignant melanoma cells [104]. Subsequently, it was found to play a vital role in regulating 394 the onset of puberty. Thus, in vertebrates kisspeptin binds to its receptor GPR54 to stimulate 395 pituitary release of gonadotropin-releasing hormone (GnRH) [105]. The first KP-type 396 precursors to be identified in non-chordates were discovered recently in ambulacrarians - the 397 echinoderms A. rubens and S. purpuratus and the hemichordate S. kowalevskii [8]. 398 Accordingly, here we have identified KP-type precursors in O. victoriae and other 399 ophiuroids. All of the ambulacrarian precursor proteins comprise two KP-type peptides and 400 the first putative neuropeptide in the echinoderm precursors has two cysteine residues at the

401 N-terminus, which could form an N-terminal disulphide bridge similar that of calcitonin-type 402 peptides (see below). In contrast, the second putative neuropeptide does not contain any 403 cysteine residues and is typically shorter than the first peptide (Figure 7C and D). 404 Interestingly, comparison of the sequences of the first (long) and second (short) KP-type 405 peptides in echinoderms reveals that the long and short peptides share less sequence 406 similarity with each other within a species than they do with respective peptides in other 407 species (Figure 7C). This indicates that the duplication event that gave rise to the occurrence 408 of the long and short peptides occurred before the divergence of the Asterozoa and 409 Echinozoa. Interestingly, previous studies have revealed that there has been an expansion of 410 KP-type receptors in ambulacraria (S. purpuratus and S. kowalevskii) and in the 411 cephalochordate, *Branchiostoma floridae*, with 16 KP receptors present in the latter [6, 54]. 412 Further studies are now needed to identify the proteins that act as receptors for the KP-type 413 peptides identified here in ophiuroids and previously in other echinoderms [8].

414

#### 415 <u>Calcitonin</u>

416 Calcitonin was first discovered in 1962 by Copp and Cheney [106]. The sequencing of 417 the porcine calcitonin in 1968 revealed that this polypeptide is composed of 32 amino acids [107]. In vertebrates, calcitonin is produced by the thyroid gland [108] and regulates calcium 418 419  $(Ca^{2+})$  levels in the blood, antagonizing the effects of parathyroid hormone [109, 110]. The 420 evolutionary antiquity of calcitonin-related peptides was first revealed with the discovery that 421 a diuretic hormone in insects  $(DH_{31})$  is a calcitonin-like peptide [111]. However,  $DH_{31}$  shares 422 modest sequence similarity with vertebrate calcitonins and lacks the N-terminal disulphide 423 bridge that is characteristic of calcitonin-type peptides in vertebrates. More recently, it has 424 been discovered that both  $DH_{31}$ -type and vertebrate calcitonin-type neuropeptides occur in 425 some protostomian invertebrates, including the annelid *Platynereis dumerilii* and the insect 426 Locusta migratoria [4, 112]. Hence, it is proposed that an ancestral-type calcitonin precursor 427 gene duplicated in the common ancestor of protostomes to give rise to  $DH_{31}$ -type and 428 calcitonin-type peptides, but with subsequent loss of calcitonin-type peptides in some 429 protostomes. Consistent with this hypothesis, calcitonin-type precursors but not  $DH_{31}$ -type 430 precursors have been identified in deuterostomian invertebrates, including echinoderms [8, 431 <u>15, 16, 113</u>].

An interesting feature of calcitonin/ $DH_{31}$  precursors is the occurrence of multiple splice variants. In vertebrates, alternative splicing of the calcitonin gene results in two transcripts: one transcript encodes calcitonin and the other transcript encodes calcitonin generelated peptide [114]. Furthermore, a complex interplay of receptors and accessory proteins

determines the pharmacological profile of these peptides [115, 116]. Alternative splicing of DH<sub>31</sub> and calcitonin precursors in insects has also been previously reported [112, 117, 118]. Interestingly, alternative splicing of insect calcitonin genes also generates variants that give rise to different mature peptides [112]. However, unlike the calcitonin gene, DH<sub>31</sub> splice variants all produce an identical mature peptide [117, 118].

441 Our analysis of the ophiuroid transcriptomes also identified two transcript variants for 442 calcitonin (Figure 7E and F). Based on our analysis of transcript sequences, ophiuroid 443 calcitonin genes comprise at least three putative coding regions or 'exons'. It is unclear if 444 these three coding regions represent three or more exons due to the lack of genomic data, but 445 for the sake of simplicity, we refer to them here as 'exons'. Transcript variant 1 comprises 446 'exons' 1 and 3 but lacks 'exon' 2 whereas transcript variant 2 contains all 3 'exons'. 447 Interestingly, 'exons' 2 and 3 both encode a calcitonin-type peptide. Hence, transcript variant 448 1 encodes a precursor that produces one calcitonin-type peptide and transcript variant 2 449 encodes two non-identical calcitonin-type peptides. These alternatively spliced transcripts 450 were found in several brittle star species (Figure 8) and thus this may represent an ancient 451 and conserved feature, although transcript variant 1 was not found in O. victoriae.

452 Previous studies have identified precursors comprising a single calcitonin-type 453 peptide in the starfish A. rubens and the sea urchin S. purpuratus [8, 16], and a precursor 454 comprising two calcitonin-type peptides in the sea cucumber A. *japonicus* [15]. Informed by 455 the identification here of two transcript types in ophiuroids (transcript variant 1 and 2), we 456 have now discovered that two transcript types also occur in A. japonicus transcriptome. 457 Hence, alternative splicing of calcitonin-type precursor genes can be traced back in the 458 echinoderm lineage to the common ancestor of the Asterozoa and Echinozoa, but with 459 subsequent loss of this characteristic in some lineages.

460

#### 461 GPA2 and GPB5

The vertebrate glycoprotein hormone family comprises luteinizing hormone (LH) 462 463 follicle-stimulating hormone (FSH), chorionic gonadotropin (CG), thyroid-stimulating 464 hormone (TSH) and the recently discovered thyrostimulin (TS) [119, 120]. Thyrostimulin is a 465 heterodimer composed of two subunits, glycoprotein alpha 2 (GPA2) and glycoprotein beta 5 466 (GPB5). Orthologs of GPA2 and GPB5 have been identified and characterized in the insect 467 *Drosophila melanogaster* [121] and in other invertebrates, including echinoderms [122]. 468 Insect GPA2 and GPB5 both contain 10 conserved cysteine residues that are important in 469 forming a heterodimeric cysteine-knot structure. Surprisingly, A. japonicus GPA2 contains 470 only 7 cysteine residues (having lost residues 7, 8 and 9) while O. victoriae GPB5.1, A.

471 *rubens* GPB5.1 and *S. purpuratus* GPB5 all contain 8 cysteine residues (having lost the final 472 two cysteine residues) (**Figure S5**). It is difficult to predict the structural differences that may 473 arise in the heterodimer due to this variability in the number of cysteine residues. The 474 possibility of GPA2 and/or GPB5 monomers or homodimers exerting their own biological 475 functions has not been ruled out [123]. Additional investigations are needed to investigate if 476 GPA2 and GPB5 are co-localized in echinoderms and if the monomers and dimers (both 477 homo and hetero) exert different effects.

478

#### 479 Uncharacterized neuropeptides

In addition to the neuropeptides discussed above, we have also identified three neuropeptide precursors that could not be classified into any known neuropeptide families. These include *O. victoriae* neuropeptide precursor (Ovnp) 18 (*O. victoriae* ortholog of Spnp18 in *S. purpuratus*) [16], Ovnp26 and Ovnp27, with the latter two identified for the first time in echinoderms. The choice of nomenclature for Ovnp26 and Ovnp27 is based on a previously used numerical nomenclature in *S. purpuratus* and/or *A. rubens*, which goes up to Arnp25 in *A. rubens*.

487

#### 488 <u>Ovnp18</u>

Ovnp18 comprises four copies of a predicted mature peptide with the sequence LFWVD and the C-terminal region of the precursor (partial sequence) contains at least four cysteine residues (**Figure 5F**). Interestingly, this precursor type only comprises a single mature peptide in *A. rubens, S. purpuratus* and *A. japonicus* and the C-terminal region contains 9, 8 and 8 cysteine residues, respectively (data not shown) [8, 15, 16].

494

#### 495 <u>Ovnp26</u>

496 Ovnp26 was identified following an analysis of O. victoriae transcriptome sequence 497 using NpSearch [8]. Orthologs of Ovnp26 were identified in other brittle stars but not in other 498 echinoderms (Figure S2-S4). Ovnp26 comprises seven copies of peptides with a conserved 499 C-terminal GW motif, whereas orthologs in O. aranea and A. filiformis are predicted to 500 generate eight copies of the mature peptide. Some of the mature peptides have a C-terminal 501 SGW motif, which is similar to the C-terminus of predicted mature peptides derived from O. 502 victoriae pedal peptide precursor 3 (Figure S7). However, the lack of sequence similarity in 503 other parts of the peptide suggests that the C-terminal similarity may reflect convergence 504 rather than homology.

505

#### 506 <u>Ovnp27</u>

507 Ovnp27 was identified following a HMM-based search for SIFamide-type peptides 508 [124, 125], albeit with a high E-value. This neuropeptide precursor comprises two putative 509 amidated mature peptides that are located immediately after the signal peptide (Figure S2-510 **S4**), as seen in SIFamide precursors  $[\underline{126}]$ . The first peptide of the *O. victoriae* precursor has 511 a C-terminal IFamide motif just like in insect SIFamides (Figure S9). However, there is no 512 sequence similarity with SIFamides in the rest of the peptide. This coupled with the fact that 513 SIFamide-type receptors have not been identified in echinoderms [6] suggests that the 514 sequence similarity that peptides derived from Ovnp27-type precursors share with SIFamides 515 may reflect convergence rather than homology.

516

#### 517 Neuropeptide precursors not found in brittle stars

518 Our analysis of ophiuroid transcriptome sequence data did not reveal orthologs of the 519 Spnp9 precursor from *S. purpuratus* or the Arnp21, Arnp22, Arnp23 and Arnp24 precursors 520 from *A. rubens* [8, 16]. An Spnp9 ortholog is found in *A. japonicus* but not in *A. rubens* [15] 521 and therefore this neuropeptide precursor type may be restricted to the Echinozoa. Orthologs 522 of Arnp21-24 have not been found in *O. victoriae, S. purpuratus* or *A. japonicus*, which 523 suggests that these may be Asteroidea-specific precursors.

Previous studies have shown that receptors for leucokinin, ecdysis-triggering hormone, QRFP, parathyroid, galanin/allatostatins-A and Neuromedin-U/CAPA are present in ambulacraria [6, 7, 15]. The presence of these receptors suggests that their cognate ligands should also be present in ambulacraria. However, our search approaches failed to identify any proteins in ophiuroids that resemble precursors of these neuropeptides.

- 529
- 530

#### <u>Evolutionary conservation and variation of neuropeptide copy number in the Ophiuroidea</u>

531 Many neuropeptide precursors comprise several structurally similar but non-identical 532 bioactive peptides – i.e. the precursor protein gives rise to a neuropeptide "cocktail". This 533 feature of neuropeptide precursors occurs throughout metazoans. But how do these 534 "cocktails" of neuropeptides evolve and what is their functional significance? Are the copies 535 of mature peptides functionally redundant or do they have their own specific functions? 536 These are important questions in neuroendocrinology for which answers remain elusive.

Evidence that neuropeptide copy number may be functionally important has been obtained from comparison of the sequences of neuropeptide precursors in twelve *Drosophila* species, the common ancestor of which dates back ~50 million years [127]. The number of peptide copies in each neuropeptide precursor was found to be identical (except for the

541 FMRFamide precursor) when compared between the twelve species, suggesting that 542 stabilising selection has acted to conserve neuropeptide "cocktails" in the *Drosophila* lineage.

- 543 Here, a comparison of O. victoriae, A. filiformis and O. aranea neuropeptide 544 precursors and their putative mature peptides revealed that fourteen neuropeptide precursors 545 comprised multiple neuropeptide copies. In certain cases, the number of the mature peptides 546 derived from a particular precursor varied across species, whereas in other cases the numbers 547 remained constant (Figure 4). Interestingly, these three species belong to two of the three 548 major clades of brittle stars that evolved  $\sim 270$  million years ago [12]. While O. victoriae 549 belongs to the Chilophiurina infraorder (clade A), A. filiformis and O. aranea belong to the 550 Gnathophiurina infraorder (clade C). Hence, this prompted us to examine the evolution of 551 neuropeptides and neuropeptide copy number variation at a higher level of phylogenetic 552 resolution. To do this, we utilized a unique dataset comprising 52 ophiuroid transcriptomes. 553 These transcriptomes were recently used as part of a phylotranscriptomic approach to 554 reconstruct the phylogeny of ophiuroids, generating a robust phylogenetic tree that comprises 555 three major clades [12]. Hence, this dataset allowed us to explore the evolution of 556 neuropeptide precursors in the context of an established phylogenetic framework spanning 557 over an unprecedented timescale of ~270 million years.
- 558 We selected for analysis neuropeptide precursors comprising more than a one putative 559 mature neuropeptide, which include AN peptide, calcitonin, cholecystokinin 1, kisspeptin, 560 np18, np26, np27, NG peptide, PDF, SALMFamide (L-type and F-type), tachykinin and TRH 561 (1 and 2). Pedal peptide precursors (1, 2 and 3) were excluded from the analysis because 562 orthology relationships between these precursors could not be established with confidence 563 across all species (data not shown). We used O. victoriae representatives of these 564 neuropeptide precursor families and the A. filiformis AN peptide precursor to mine 52 565 ophiuroid transcriptomes using BLAST. Multiple sequence alignments were generated based 566 on the search hits (Figure S10) and the number of predicted mature peptides were compared 567 (Figure 8). Interestingly, the number of peptides within the majority of precursors remained 568 constant across all the species examined, which share a common ancestor estimated to date 569 from  $\sim 270$  million years ago [12].
- 570 Some studies that have investigated the physiological significance of neuropeptide 571 "cocktails" indicate that neuropeptides derived from the same precursor protein are 572 functionally redundant. For example, this was found for myomodulin neuropeptides in the 573 mollusk *Aplysia californica* using the accessory radula closer muscle preparation as a 574 bioassay [128] and for FMRFamide-related neuropeptides in *Drosophila melanogaster* when 575 analysing effects on nerve-stimulated contraction of larval body-wall muscles [129].

576 However, the authors of the latter study cautiously highlighted the need to "search for 577 additional functions or processes in which these peptides may act differentially". Importantly, 578 studies employing use of multiple bioassays have obtained data indicating that neuropeptides 579 derived from a single precursor protein are not functionally redundant. For example, when 580 the actions of fourteen structurally related neuropeptides derived from a precursor of Mytilus 581 Inhibitory Peptide-related peptides in *Aplysia* were tested on three organ preparations (oesophagus, penis retractor, body wall) it was found that the rank order of potency for the 582 583 peptides differed between preparations [130]. Similarly, when assaying the effects of 584 allatostatin neuropeptides in cockroaches, tissue-specific differences in potency were 585 observed [131]. The conservation of peptide copy number across a timescale of  $\sim 270$  million 586 years in the Ophiuroidea supports the idea that the occurrence of multiple copies of identical 587 or structurally related neuropeptides is functionally important.

588 For those neuropeptide precursors that did exhibit variation in neuropeptide copy 589 number, TRH-type precursors exhibited the highest variation, with numbers ranging from 16 590 to 20 copies (Figure 9). F-type SALMFamide precusors also showed variation in copy 591 numbers (Figure 10) but loss of peptides was more frequent in F-type SALMFamide 592 precursors than in TRH-type precursors. Furthermore, detailed analysis of sequence 593 alignments for these precursors revealed that loss of neuropeptide copies is usually a 594 consequence of non-synonymous mutations in codons for residues that form dibasic cleavage 595 sites or for glycine residues that are substrates for the C-terminal amidation. This is not 596 surprising since the C-terminal amide in smaller-sized peptides is usually important for 597 receptor binding and activation. What is unclear at the moment is how the peptide copy 598 number increases within a given precursor. Perhaps the increase in peptide copy number 599 occurs as a result of unequal crossing-over during recombination [127].

600 The number of peptides within the F-type SALMFamide precursors appear to be clade 601 specific. Thus, the average/median number of F-type SALMFamides in precursors from clade 602 A is 13, clade B is 12 and clade C is 11, with a few exceptions (Figure 8). Similarly, the 603 number of peptides within NP26-type precursors also appears to be clade specific. Hence the 604 number of peptides is highly stable at 7 peptides within clades A and B but a high variation in 605 peptide copy number is observed in clade C. When examining peptide copy number within 606 clades, there are a few cases where the number of peptides within a given precursor for 607 certain species appears to be an exception/outlier. For instance, 16 copies of the mature 608 peptide in Ophioplax lamellosa TRH-1 precursor is distinctly different to the 19 copies found 609 in other species within that clade (clade C). Likewise, Ophiactis savignyi only has 3 copies of 610 kisspeptin-type peptides compared to 4 copies found in other species of that clade (Figure 8).

611 It could be argued that misalignments during transcriptome assembly may have 612 influenced the number of predicted peptides found in a given precursor. However, it is 613 unlikely that misalignments have affected the predicted sequences of neuropeptide precursors 614 comprising multiple copies of peptides that are similar but non-identical, which applies to the 615 majority of the precursor proteins analysed here in ophiuroids. The only exception to this are 616 the TRH-type precursors, where the encoded peptide sequences are short and often identical, 617 even at the nucleotide level (data not shown), Another limitation of using transcriptome data 618 is that the sequences of neropeptide precursors may be partial or unknown for some species 619 and where this applies a peptide copy number is not shown in Fig. 8. An extreme example of 620 this is the AN peptide precursor, where complete precursors sequences were only obtained 621 from the three reference species and three other species. However, for the majority of 622 precursor types, sequence data was obtained from a variety of species from each of the three 623 clades of ophiuroids. For example, complete F-type SALMFamide precursor sequences were 624 found in most of the investigated species (39 species + 3 reference species).

625

#### 626 <u>Conclusion</u>

627 Here we report the first detailed analysis of the neuropeptide precursor complement of 628 ophiuroids and the most comprehensive identification of echinoderm neuropeptide precursors 629 to date. We have identified novel representatives of several bilaterian neuropeptide families 630 in echinoderms for the first time, which include orthologs of endothelin/CCHamide, eclosion 631 hormone, neuropeptide-F/Y and nucleobinin/nesfatin. Furthermore, analysis of precursor 632 proteins comprising multiple copies of identical or related neuropeptides across ~270 million 633 years of ophiuroid evolution indicates that the precise composition of neuropeptide 634 "cocktails" is functionally important as evident from the conservation of neuropeptide copy 635 number for multiple precursors.

636

#### 637 Methods

#### 638 Sequencing and assembly of transcriptomes

Ophiuroid transcriptomes used in this study were sequenced and assembled as
reported previously [<u>12</u>, <u>20</u>, <u>24</u>].

641

#### 642 Identification of neuropeptide precursors in ophiuroids

In order to identify neuropeptide precursors in *O. victoriae*, *A. filiformis* and *O. aranea*, sequences of neuropeptide precursors identified previously in other echinoderms (including the starfish, *A. rubens*, the sea urchin *S. purpuratus* and the sea cucumber, *A.* 

japonicus) were used as queries for tBLASTn analysis of a transcriptome database, using an e value of 1000. Sequences identified as potential neuropeptide precursors by BLAST were translated using the ExPASy Translate tool (<u>http://web.expasy.org/translate/</u>) and then analysed for features of neuropeptide precursors. Specifically, sequences were evaluated based on 1) the presence of an N-terminal signal peptide (using Signal P v 4.1 with the sensitive cut-off of 0.34) and 2) the presence of monobasic or dibasic cleavage sites flanking the putative bioactive peptide(s).

To identify novel neuropeptide precursors or highly-divergent precursors with low sequence similarity to known precursors, we utilized two additional approaches. In the first approach, we used NpSearch [8], software that identifies putative neuropeptide precursors based on various characteristics (presence of signal peptide and dibasic cleavage sites amongst others). In the second approach, NpHMMer (<u>http://nphmmer.sbcs.qmul.ac.uk/</u>), a Hidden Markov Models (HMM) based software was used to identify neuropeptides not found using the above approaches.

660 Neuropeptide precursors identified in O. victoriae (which represented a more 661 comprehensive neuropeptide precursor repertoire compared to A. filiformis and O. aranea) 662 were then submitted as queries for BLAST analysis of sequence data from 52 Ophiuroidea 663 species, using an E-value of 1e-06. BLAST hits were then further analysed using an 664 automated ruby script (available on Github). Each BLAST hit was translated using BioRuby 665 and the open reading frame (ORF) containing the BLAST high-scoring segment pair was 666 extracted. These ORFs were then examined for the presence of a signal peptide using Signal 667 P 4.1 using a sensitive cut-off of 0.34. All sequences were then aligned using MAFFT, with 668 the number of maximum iterations set to 1000 to ensure an optimal alignment. These 669 alignments were then further optimized by manually adjusting the location of the bioactive 670 peptide and cleavage sites. Finally, the alignments were annotated using different colours for 671 the signal peptide (blue), the bioactive peptide(s) (red) and cleavage sites (green).

672

## 673 Phylogenetic and clustering analyses of sequence data

Phylogenetic analysis of membrane guanylyl cyclase receptors and nucleobindins was performed using maximum likelihood and Bayesian methods. Prior to these analyses, corresponding multiple alignments were trimmed using BMGE [132] with the following options: BLOSUM30, max -h = 1, -b = 1, as described previously [10, 91]. The maximum likelihood method was implemented in the PhyML program (v3.1/3.0 aLRT). The WAG substitution model was selected assuming an estimated proportion of invariant sites (of 0.112) and 4 gamma-distributed rate categories to account for rate heterogeneity across sites.

681 The gamma shape parameter was estimated directly from the data. Reliability for internal 682 branch was assessed using the bootstrapping method (500 bootstrap replicates). The Bayesian 683 inference method was implemented in the MrBayes program (v3.2.3). The number of 684 substitution types was fixed to 6. The poisson model was used for substitution, while rates 685 variation across sites was fixed to "invgamma". Four Markov Chain Monte Carlo (MCMC) 686 chains were run for 100000 generations, sampling every 100 generations, with the first 500 687 sampled trees discarded as "burn-in". Finally, a 50% majority rule consensus tree was 688 constructed.

689 CLANS analysis was performed on echinoderm EH-like, arthropod EH, arthropod 690 ITP and vertebrates ANP precursors based on all-against-all sequence similarity (BLAST 691 searches) using BLOSUM 45 matrix (<u>https://toolkit.tuebingen.mpg.de/clans/</u>) [39] and the 692 significant high-scoring segment pairs (HSPs). Neuropeptide precursors were clustered in a 693 three-dimensional graph represented here in two dimensions.

694

## 695 Competing Interests

- 696 The authors declare that no competing interests exist.
- 697

#### 698 <u>Author contributions</u>

699 M.Z., T.D.O. and M.R.E.: designed the research; I.M.: generated HMM models; M.Z., I.M.,

700 L.A.Y.G., J.D., N.A. and A.F.H: identified the neuropeptide precursors; M.Z., I.M.,

701 L.A.Y.G., J.D. and N.A.: analysed the data; M.Z., J.D. and M.R.E. wrote the manuscript with

input from other authors. M.Z. and M.R.E: supervised the study.

703

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#### 1097 **Figure captions**

**Figure 1:** Bilaterian animal phylogeny. The diagram shows i). the phylogenetic position of the phylum Echinodermata in the ambulacrarian clade of the deuterostomes and ii) relationships between the five extant classes of echinoderms, which include the focal class for this study – the Ophiuroidea (e.g. *Ophionotus victoria*e).

1102

1103 Figure 2: Eclosion hormone (EH)-type peptides and receptors in echinoderms A) Partial 1104 multiple sequence alignment of eclosion hormone-type precursor sequences, excluding the N-1105 terminal signal peptide; B) Cluster analysis of arthropod EH precursors, echinoderm EH-like 1106 precursors, arthropod ion transport peptides (ITPs) and vertebrate atrial natriuretic peptides shows that echinoderm EH-like precursors are more closely related to arthropod EH than ITP 1107 1108 C) Phylogenetic analysis of membrane guanylate cyclase receptors shows that EH-like 1109 receptors are found in echinoderms but are absent in vertebrates as seen for the EH-like 1110 precursors. Species names: Ophionotus victoriae (Ovic), Asterias rubens (Arub), 1111 Strongylocentrotus purpuratus (Spur), Drosophila melanogaster (Dmel), Bombyx mori 1112 (Bmor) and *Pediculus humanus corporis* (Pcor).

1113

Figure 3: Multiple sequence alignments of A) CCHamide-type and B) Neuropeptide-F/Ytype peptides. Species names: *Ophionotus victoriae* (Ovic), *Asterias rubens* (Arub), *Apostichopus japonicus* (Ajap), *Drosophila melanogaster* (Dmel), *Apis mellifera* (Amel), *Lottia gigantea* (Lgig), *Aplysia californica* (Acal), *Homo sapiens* (Hsap), *Ophiopsila aranea*(Oara), *Amphiura filiformis* (Afil), *Patiria miniata* (Pmin), *Saccoglossus kowalevskii* (Skow), *Branchiostoma floridae* (Bflo) and *Daphnia pulex* (Dpul).

1120

Figure 4: Summary of neuropeptide precursors identified in *Ophionotus victoriae, Amphiura filiformis* and *Ophiopsila aranea*. Neuropeptide precursors are classified based on the type of G-protein coupled receptor (GPCR) their constituent peptides are predicted to activate (see Mirabeau and Joly, 2013). Some peptides bind to receptors other than GPCRs and these are grouped with peptides where the receptor is unknown. Ophiuroids have neuropeptide precursors from up to 32 families. The number of putative mature peptides derived from each precursor has been indicated along with the presence of amidation and pyroglutamation.

1128

Figure 5: Multiple sequence alignments of mature peptides belonging to selected
neuropeptide families. A) corazonin alignment; B) gonadotropin-releasing hormone (GnRH)
alignment; C) orexin alignment; D) luqin alignment; E) vasopressin/oxytocin (VP/OT)

alignment; F) Ovnp18 alignment; G) melanin-concentrating hormone (MCH) alignment; H)
NP peptide alignment; I) pigment dispersing factor (PDF) alignment. Species names: *Ophionotus victoriae* (Ovic), *Asterias rubens* (Arub), *Strongylocentrotus purpuratus* (Spur), *Apostichopus japonicus* (Ajap), *Saccoglossus kowalevskii* (Skow), *Branchiostoma floridae*(Bflo), *Anopheles gambiae* (Agam), *Daphnia pulex* (Dpul), *Strigamia maritima* (Smar),

- 1137 Lottia gigantea (Lgig) and Homo sapiens (Hsap).
- 1138
- Figure 6: Alignments of neuropeptides derived from precursors that exist in multiple forms
  in ophiuroids. A) thyrotropin-releasing hormone (TRH) alignment; B) cholecystokinin
  alignment; C) somatostatin alignment; D) corticotropin-releasing hormone (CRH) alignment.
  Species names: *Ophionotus victoriae* (Ovic), *Asterias rubens* (Arub), *Strongylocentrotus purpuratus* (Spur), *Apostichopus japonicus* (Ajap), *Branchiostoma floridae* (Bflo), *Homo*sapiens (Hsap), *Drosophila melanogaster* (Dmel) and *Lottia gigantea* (Lgig).
- 1145

1146 Figure 7: Comparative analysis of ophiuroid tachykinin, kisspeptin and calcitonin-type 1147 precursors and neuropeptides. A) Alignment of tachykinin-type peptides in O. victoriae 1148 (Ophiuroidea) and A. rubens (Asteroidea); B) Schematic diagrams of the O. victoriae and A. 1149 rubens tachykinin precursors showing the location of the signal peptide (SP) and predicted 1150 neuropeptides (labelled 1 to 4); C) Alignments of the long and short forms of kisspeptin-type 1151 neuropeptides in O. victoriae, A. rubens and S. purpuratus (Echinoidea) D) Schematic 1152 diagrams of the O. victoriae and A. rubens kisspeptin precursors showing the locations of the 1153 SP, short and long orthocopies and cysteine (C) residues; E) Alignment of calcitonin-type 1154 peptides from O. victoriae, A. rubens, S. purpuratus and A. japonicus (Holothuroidea); F) 1155 Predicted alternative splicing of the calcitonin gene in ophiuroids, with the location of the SP 1156 and neuropeptides (CT1 and CT2) labelled. Species names: Ophionotus victoriae (Ovic), 1157 Asterias rubens (Arub), Strongylocentrotus purpuratus (Spur) and Apostichopus japonicus 1158 (Ajap).

1159

Figure 8: Comparison of neuropeptide copy numbers across the Ophiuroidea for precursors
comprising multiple copies of neuropeptides. Neuropeptide precursors were mined from 52
ophiuroid transcriptomes, with the phylogeny adapted from O'Hara et al. (2014) [12].
Am\_laud: Amphiophiura laudata, Am\_spat: Amphiophiura spatulifera, Am\_cipu:
Amphioplus cipus, Am\_cten: Amphioplus ctenacantha, Am\_squa: Amphipholis squamata,
Am\_cons1: Amphiura constricta 1, Am\_cons2: Amphiura constricta 2, As\_love: Asteronyx
loveni, As\_bidw: Asteroschema bidwillae, As\_tubi: Asteroschema tubiferum, Ba\_hero:

1167 Bathypectinura heros, Cl cana: Clarkcoma canaliculata, Gl sp no: Glaciacantha sp nov, 1168 Go\_pust: Gorgonocephalus pustulatum, Mi\_grac: Microphiopholis gracillima, Op\_fune: 1169 Ophiacantha funebris, Op abys: Ophiactis abyssicola, Op resi: Ophiactis resiliens, Op savi: 1170 Ophiactis savignyi, Op vall: Ophiernus vallincola, Op pilo: Ophiocentrus pilosus, Op\_wend: Ophiocoma wendtii, Op\_oedi: Ophiocreas oedipus, Op\_tube: Ophiocypris 1171 1172 tuberculosis, Op\_appr: Ophioderma appressum, Op\_bisc: Ophiolepis biscalata, Op\_impr: 1173 Ophiolepis impressa, Op\_brev: Ophioleuce brevispinum, Op\_perf: Ophiolimna perfida, Op\_prol: Ophiologimus prolifer, Op\_obst: Ophiomoeris obstricta, Op\_lyma: Ophiomusium 1174 1175 lymani, Op\_aust: Ophiomyxa australis, Op\_vivi: Ophiomyxa sp cf vivipara, Op\_fasc: Ophionereis fasciata, Op\_reti: Ophionereis reticulata, Op\_scha: Ophionereis schayeri, 1176 Op\_filo: *Ophiophragmus filograneus*, 1177 Op\_cyli: *Ophiopeza* cylindrica, Op\_wurd: 1178 Ophiophragmus wurdemanii, Op\_liod: Ophiophrura liodisca, Op\_john: Ophiophycis johni, 1179 Op\_lame: Ophioplax lamellosa, Op\_iner: Ophiopleura inermis, Op\_plic: Ophioplinthaca 1180 plicata, Op\_bisp: Ophioplocus bispinosus, Op\_macu: Ophiopsammus maculata, Op\_angu: 1181 Ophiothrix angulata, Op\_caes: Ophiothrix caespitosa, Op\_exim\_1: Ophiotreta eximia 1,

1182 Op\_exim\_2: *Ophiotreta eximia* 2, Op\_sp\_no: *Ophiura sp nov*.

1183

**Figure 9:** A partial multiple sequence alignment of ophiuroid thyrotropin-releasing hormone (TRH) precursors showing clade-specific gain/loss of neuropeptide copies. Mono- and dibasic cleavage sites are highlighted in green, mature peptides in red with the glycine residue for amidation in pink. Species have been grouped and coloured (clade A in purple, clade B in blue and clade C in orange) based on the phylogeny determined by O'Hara et al. (2014) [12].

1189

**Figure 10:** A partial multiple sequence alignment of ophiuroid F-type SALMFamide precursors showing clade-specific gain/loss of neuropeptide copies. Di-basic cleavage sites are highlighted in green, mature peptides in red with the glycine residue for amidation in pink. Species have been grouped and coloured (clade A in purple, clade B in blue and clade C in orange) based on the phylogeny determined by O'Hara et al. (2014) [12].

1195

#### 1196 Supplementary files

Figure S1: Alignment and phylogenetic analysis of nucleobindins (NUCB). A) Partial sequence alignment (excludes the signal peptide) of NUCB precursors. The locations of *Homo sapiens* nesfatin-1, 2 and 3 are indicated. A dibasic cleavage site in *O. victoriae* nesfatin-1 is marked in red. B) Phylogenetic analysis of NUCB precursors. Species names: *Ophionotus victoriae* (Ovic), *Amphiura filiformis* (Afil), *Ophiopsila aranea* (Oara),

1202 Apostichopus japonicus (Ajap), Strongylocentrotus purpuratus (Spur), Homo sapiens (Hsap),

1203 Mus musculus (Mmus) and Drosophila melanogaster (Dmel).

1204

- 1205 Figure S2: Ophionotus victoriae neuropeptide precursor repertoire.
- 1206

1207 Figure S3: Amphiura filiformis neuropeptide precursor repertoire.

1208

1209 Figure S4: *Ophiopsila aranea* neuropeptide precursor repertoire.

1210

1211 Figure S5: Partial multiple sequence alignments of echinoderm representatives of A)

1212 glycoprotein alpha 2 (GPA2)-type subunits and B) glycoprotein beta 5 (GPB5)-type subunits.

1213 Species names: Ophionotus victoriae (Ovic), Asterias rubens (Arub), Strongylocentrotus

1214 *purpuratus* (Spur) and *Apostichopus japonicus* (Ajap).

1215

Figure S6: Partial multiple sequence alignments of echinoderm representatives of large protein hormones. A) insulin/insulin-like growth factor; B) relaxin-like peptide; C) bursicon (bursicon alpha); D) partner of bursicon (bursicon beta). Species names: *Ophionotus victoriae* (Ovic), *Asterias rubens* (Arub), *Strongylocentrotus purpuratus* (Spur) and *Apostichopus japonicus* (Ajap).

1221

Figure S7: Multipe sequence alignment of echinoderm pedal peptides. Species names: *Ophionotus victoriae* (Ovic), *Asterias rubens* (Arub), *Strongylocentrotus purpuratus* (Spur)
and *Apostichopus japonicus* (Ajap).

1225

Figure S8: Multiple sequence alignments of echinoderm neuropeptide families. A) F-type
SALMFamide alignment; B) L-type SALMFamide alignment; C) AN peptide. Species
names: *Ophionotus victoriae* (Ovic), *Asterias rubens* (Arub), *Strongylocentrotus purpuratus*(Spur) and *Apostichopus japonicus* (Ajap).

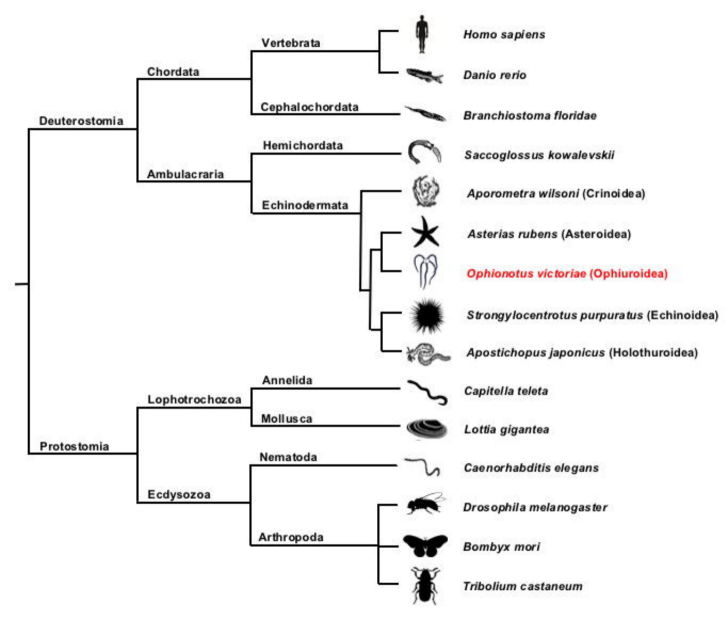
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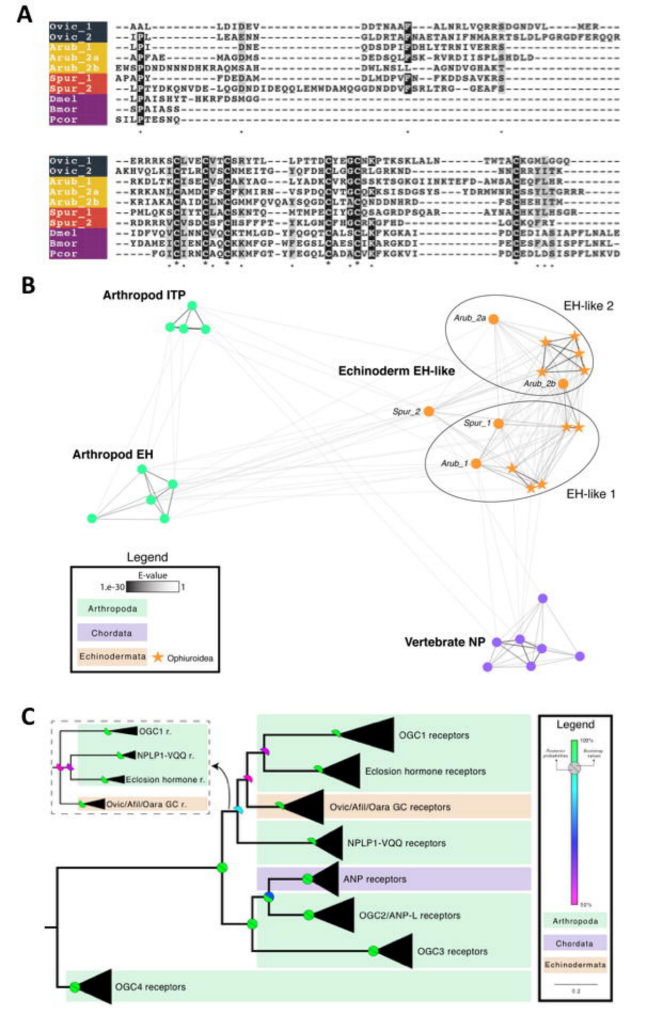
Figure S9: Multiple sequence alignment of predicted peptides derived from neuropeptide
precursor 27 in *Ophionotus victoriae* (Ovic), *Amphiura filiformis* (Afil), *Ophiopsila aranea*(Oara) and *Apostichopus japonicus* (Ajap).

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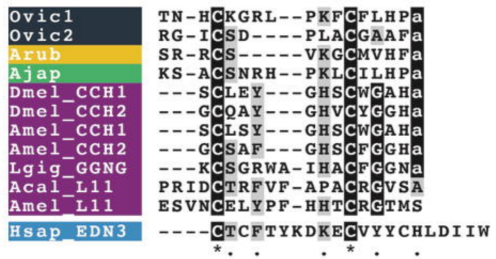
Figure S10: Multiple sequence alignments of neuropeptide precursors used to generateFigure 8.

1237









В

А

## Neuropeptide-F/Y

Oara	RTTGDKALDAILSGQY-RSHLRYa
Afil	RTTGDKALDAILSGQY-RHHLRYa
Arub	pQDRSKAMQAERTGQLRRLNPRFa
Pmin	pQSDMRDKAMQAITTGQINRNHARYa
Skow	DASDYQAPTAPSRGASLAEWDRYLRELSLYRQYADIQRFa
Bflo	pQEEEDVEAPEEGKYYKNLANYLRLLTRQRYa
Hsap	YPSKPDNPGEDAPAEDMARYYS <mark>AL</mark> RHYINLITRQRYa
Dmel	SNSRPPRKNDVNTMADAYKFLQDLDTYYGDRARVRFa
Dpul	DGGDVMSGGEGGEMTAMADAIKYLQGLRRYDNSLVRPRFa
Lgig	pQDSMLAPPDRPSEFRSPDELRRYLKALNEYYAIVGRPRFa
	*.*

96				O. victoriae				A. filiformis				O. aranea			
Receptor type			Precursor	Predicted peptides	Amidated	Pyroglutamate	Precursor	Predicted peptides	Amidated	Pyroglutamate	Precursor	Predicted peptides	Amidated	Pyroglutamate	
		Neuropeptide family				٤.				٤.				5	
	1	CCHamide-like 1		1				1		Щ		1			
		CCHamide-like 2		1	_			1		Щ		1			
	2	<i>Cholecystokinin</i> 1 bioRxiv preprint doi: https://doi.org/10	110	3 1/12978	······································	nis ve	ersion	3 nosted	May	20	2017	1* The co	nyrio	ht h	older f
		bioRxiv preprint doi: https://doi.org/19 certified by peer review) is the author/fu	nder		has g	rante	ed bioF	Rxiv <sup>+</sup> a I 4.Q Int	icens	se to	displa	y the p	repri	nt in	perpe
	3	Corazonin		-	100-			-		lione		<sup>30</sup> .1			
-	4	Gonadotropin-releasing hormone	111.	1	_			1					$\vdash$		
Rhodopsin <b>β</b>	5	Luqin Neuropeptide-F/Y 1		1				1				1			
dop	6	Neuropeptide-F/Y 2		1*		H		1				1			-
Rho	-	NG peptide / Neuropeptide-S				Н		1	-				$\mathbf{H}$	$\leftarrow$	
	7 8	Orexin 1		2				2		Н		2			
	0	Orexin 2		1	-			1	—			1		-	
	9	Tachykinin		1				1 4				1 4		777.	
	10	Thyrotropin-releasing hormone 1		4 21	-			4 14*		<i></i>					
	10	Thyrotropin-releasing hormone 2		4		77		14 · 4*		77		17			
	11	Vasopressin / Oxytocin		1				4			77	1	H	F	
>	12	Kisspeptin		2				1*		$\vdash$		1			
Rhodopsin y	13	Melanin-concentrating hormone		1		····		1		Н		1		F	
topo	14	Somatostatin 1		1				1		H		-			
Rhe		Somatostatin 2		1				1		$\square$		1	Ħ	É	
	15	Bursicon alpha		1				-				-			
	16	Bursicon beta		1				1	Ħ	Ħ					
inδ	17	Glycoprotein hormone alpha 2.1						1				1			
hodopsin <b>ð</b>		Glycoprotein hormone alpha 2.2		1		Ē		1				1			
Rhoc	18	Glycoprotein hormone beta 5.1		1			11	1							
-		Glycoprotein hormone beta 5.2		1				1							
	19	Relaxin-like peptide		а				а				а			
	20	Calcitonin		2				1/2				1/2			
	21	Corticotropin-releasing hormone 1		1				1					$\square$		
Secretin		Corticotropin-releasing hormone 2		1				1				1			
Seci		Corticotropin-releasing hormone 3		1*				1							
		Corticotropin-releasing hormone 4		1*				1*							
	22	Pigment-dispersing factor		2				2				2			
	23	AN peptide						5*				7			
	24	Eclosion hormone 1.1				$\square$		1				1			
		Eclosion hormone 1.2		1		Ц		1	Ц	Ц		1	Ш		
		Eclosion hormone 2.1			$\square$	4		1		Ш		1			
ers		Eclosion hormone 2.2		1				1	Щ	Щ		1			
Oth	25	Insulin-like peptide		a				а		Щ			А	K	
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cnov	27	Pedal peptide 1 Pedal peptide 2	11	o 4*		$\vdash$		L	$ \blacksquare $			1*			
Unk		Pedal peptide 3		8*		$\vdash$	7/	С		Η		c			
	28	SALMFamide (L-type)		4				4			11	4*			
	29	SALMFamide (F-type)		12				11		111.		11			
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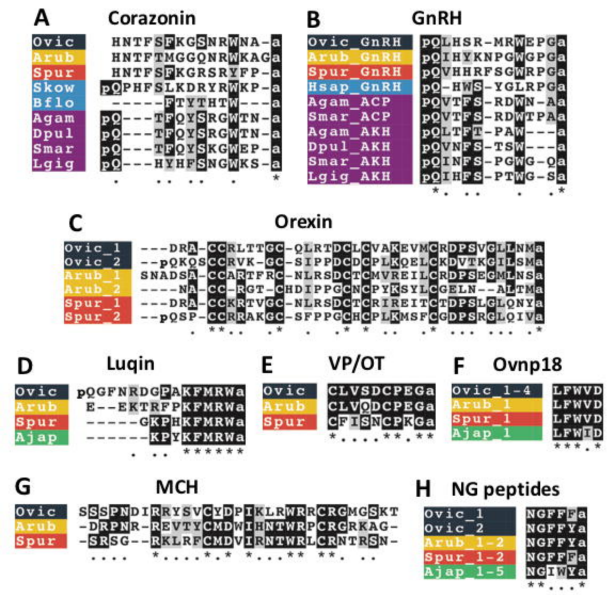
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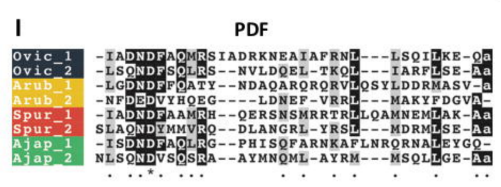
Partial / some mature peptides

Absent

Cannot be determined

- a Heterodimer of A-chain and B-chain
- b Number of mature peptides unknown
- c Multiple partial precursors





# TRH

в

Cholecystoki	nin
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.\*..\*.

Ovic1_1	pQESPa
<b>Ovic1</b> 2-17	pOFSAa
Ovic1_18-21	POFAAa
Ovic2_3-4	pQGPRa
Arub_1-12	POWYTa
Spur_1-10	pQYPGa
Spur_11	pQFPAa
Spur_12-16	pQWPGa
Spur_17	pQFPGa
Ajap_1-10	pQYFAa
Ajap_11	pQLPGa
Ajap_12-15	pQFFQa
Ajap_16	pQHFVa
Ajap_17	pQHFAa
Ajap_18	pOHFLa

Ovic 1 Ovic<sup>2</sup>

1

2

2

C

Spur

Spur

Ajap

Bflo Hsap\_SI Hsap

5 0	nolecystokinin
Ovic1	1SKDYGWGMAFa
Ovic1	2NKDYGWGMAFa
Ovic1_	3NEYGWGHMFa
Ovic2	SLDYGFGMGFa
Arub1_	VDDYGHGLFWa
Arub1	2 GGDDQYGFGLFFa
Spur1_	1DYGHGMFFa
and the second se	2PDDYNWGMWFa
the state of the s	3 – – DKADLYGWGGFFa
Spur2	DAGPHAWYGTGM-Fa
Ajap1_	1MNGWY-TGM-Fa
Ajap1_	2NIPQTYLSGDYFa
	.***

Q	F	s	P	а		0	v	i
Q	F	s	A	a		0	v	i
Q	F	A	A	a		0	v	i
Q	G	P	R	а		0	v	i
Q	W	Y	т	a		A	r	u
Q	Y	P	G	a		А	r	
Q	F	P	A	a		s	p	u
Q	W	P	G	a		s	p	u
Q	F	P	G	а		s	p	u
Q	Y	F	A	a		s	p	u
Q	L	P	G	a		A	j	a
Q	F	F	Q	a		A	j	a
Q	H	F	V	a				
	н							

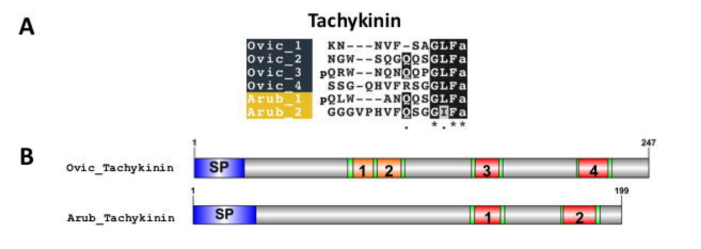
	-	-	
4	۲	-	
	Γ.		
٩		-	
	-		

# Somatostatin

	GK	-VG	REVP	-YM-MN	-
	PG	-VY	DIWKGR	GLSR	т
				-FS-MP	
	GK	-MG	RFGP	-YM-LN	- 2
	PARKI	-IN	DIWKGR	GGG-LR	N
				GGSNHR	
	AKG	-AR	FYWKMP	ATA-MS	- 2
MS				-TF-TS	
DRT	-DRMP	-RN	FWK	-TF-SS	- 2
	3	* .			R:

C	DI	
L	ΓI	

Ovic_1	-TGSPIALNPGLVVLDIERSTIDNDRRR-QQMSEAAAMSEEFTRVA
Ovic_2	- pQMN DLFTTFSVIREAFESAKNE-RDRASALAANGRIFAAGa
Ovic_3	- pQMTVDPFTTMQIIRDHQTAEKE-RQRQKAIDINGRIFAAGA
Ovic_4	-DNFEFGLFTSLDIMRDAFQSKSE-RERADALAANEDLAAAB
Arub	pQGLSVSPIFPIQRIR-ENAIERDR-QDQVDQAEANQGLFQIA
Hsap_CRH	SEEPPISIDLTFHLIREVLEMARAEQLAQQAHSNRKIMEIIE
Hsap_UCN1	-DNPSISIDLTFHLIRTILELARTQSQRERAEQNRIIFDSV
Hsap_UCN2	IVLSLDVPIGLLQILLEQARARAAREQATTNARILARVCHC
Hsap_UCN3	FTLSLDVPTNIMNLLFNIAKAKNLRAQAAANAHLMAQI3
Dmel_DH44	-NKPSLSIVNPLDVLRQRLLLEIARRQMKENSRQVELNRAILKNVa
Lgig_ELH1	SRLSINQELKSLANLLVLRENK-RREAQKTKLRSKL-LSIG
Lgig_ELH2	AGRLSINGALSSLADLLVSENQR-RDRLESMELRQRL-QYLE

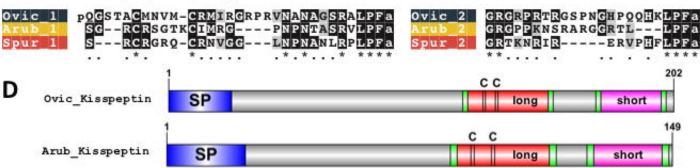


#### С Kisspeptin (long)

CT variant 1

SP

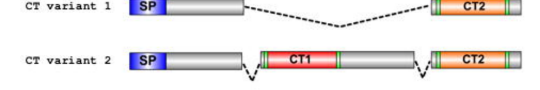
# **Kisspeptin** (short)

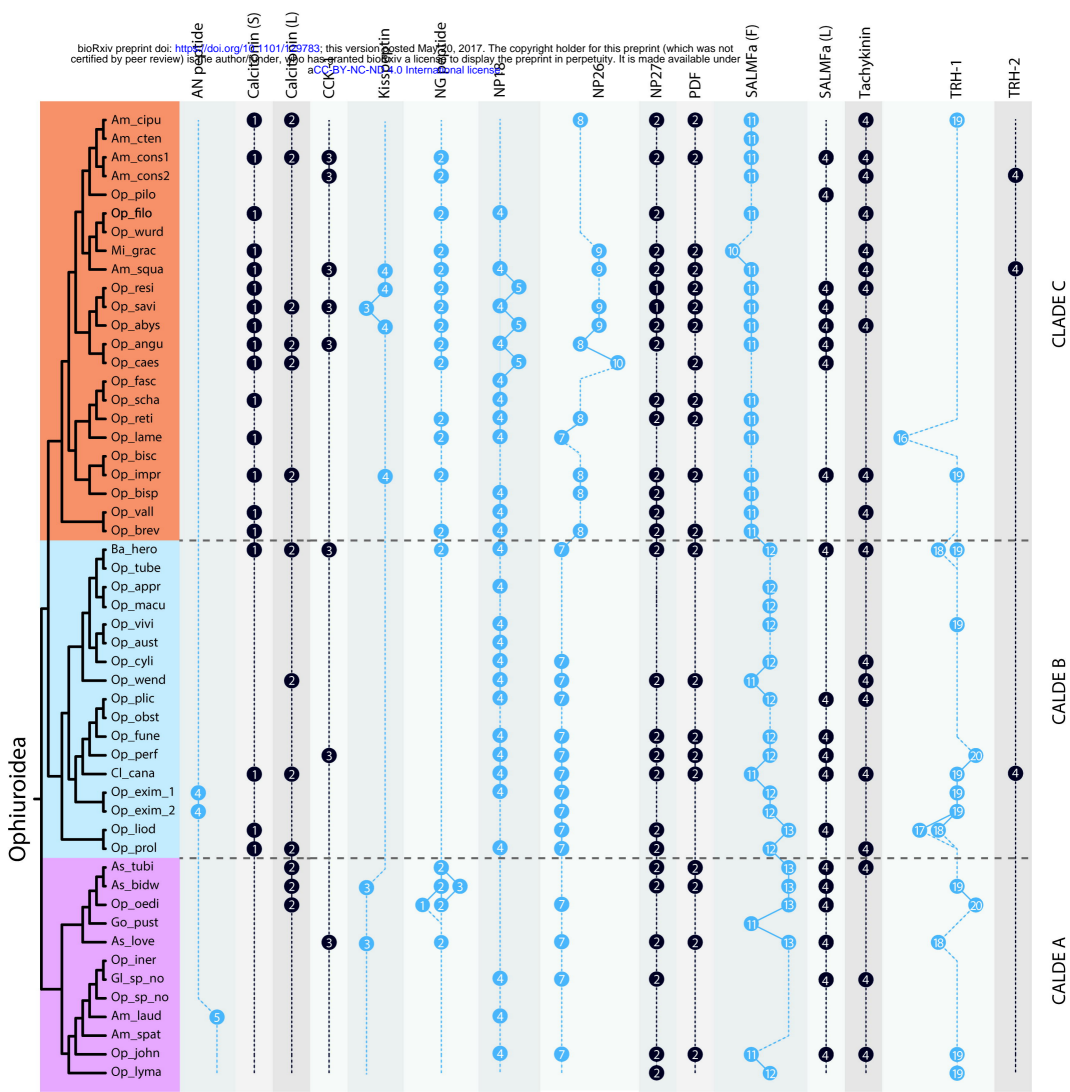


Ε

F

#### Calcitonin S-GNGGCAG-FTGCAQLAAGQNALRNFMHSNRASLFTGASGPA N-GNGGCAG-FTGCAQLAAGQSALQAMIHSGRASLF-GSGGPA NGESRGCSG-FGGCGVLTIGHNAAMRMLAESNSP-F-GASGPA ---SKGCGS-FSGCMQMEVAKNRVAALLRNSNAHLF-GLNGPA ----SCSNKFAGCAHMKVANAVLKQNSRGQQQFKF-GSAGPA --RVGGCGD-FSGCASLKAGRDLVRAMLRPSK---F-GSGGPA Ovic 1 Ovic 2 Spur ap Ajap \* . \* . \* \* . Putative calcitonin splicing in Ophiuroidea Exon 3 Exon 1 Exon 2 CT2 SP CT1





# <u>TRH-1</u>

Am cipu	SDDPFSPD <mark>KRQFSAGKRQFSAGKRQFSAGKRQFSAGKRQFSAGKR</mark> QWLGGEEEYDPEENLNMET <mark>RQFSAGKR</mark> QFSA <mark>GKR</mark> QFSAGKR
Op_angu	VDMPET <mark>R</mark> QFSAGKRQFSAG <mark>KR</mark> QFSAC <mark>KR</mark> QFSAGKRQFSAGKRQWVGGEEDDGLEENDDM <mark>KRQFSAGKR</mark> QFSAC <mark>KR</mark> QFSAGKRQFSAGKR
Op_lame	VDMPET <mark>RQFSAGKRQFSAGKR</mark> QFSAC <mark>KR</mark> QFSAGKRQWVGGEPEEWEDEDM <mark>KRQFSAGKR</mark> QFSACKRQFSACKR
Op_impr	DDM <mark>kr</mark> qfsag <mark>kr</mark> qfsa <mark>gkr</mark> qfsagkrqfsagkrqfsagkrqwvggfplefededv <mark>krqfsagkr</mark> qfsagkrqfsagkrqfsagkr
Ba_hero_a	VDMPET <mark>RQFSAGKR</mark> QFSAG <mark>KR</mark> QFSAC <mark>KR</mark> QFSA <mark>GKR</mark> QWVGGEPDVLNQDE <mark>KR</mark> QFSAG <mark>KR</mark> QFSAC <mark>KR</mark> QFSA <mark>GKR</mark> QFSAGKRQFS
Ba_hero_b	VDMPET <mark>RQFSAGKRQFSAGKR</mark> QFSA <mark>GKR</mark> QFSA <mark>GKR</mark> QWVGGEPDVLNQDE <mark>KRQFSAGKR</mark> QFSA <mark>GKR</mark> QFSA <mark>GKR</mark> QFSAGKR
Op_vivi	VDMPET <mark>RQFSAGKRQFSAGKRQFSAGKRQFSAGKRQFAAGKR</mark> QWVGGEPDEFD-EAQ <mark>KRQFSAGKR</mark> QFAA <mark>GKR</mark> QFAAG <mark>KR</mark> QFAAGKR
Op_perf	VDMPET <mark>RQFSAGKR</mark> QFSAG <mark>KR</mark> QFSAGKRQFSAGKRQFSAGKRQWVGGEPDEEEE <mark>KRQFSAGKR</mark> QFSAG <mark>KR</mark> QFSAGKRQFSAGKR
Op_exim_1	VDMPET <mark>RQFSAGKR</mark> QFSAG <mark>KR</mark> QFSAGKRQFSAGKRQWVGGQPDLLDDEEE <mark>KRQFSAGKR</mark> QFSAG <mark>KR</mark> QFSAGKRQFSAGKR
Op_liod_a	VDMPET <mark>R</mark> QFSP <mark>GKR</mark> QFSPC <mark>KR</mark> QFSPC <mark>KR</mark> QFSPGKRQFSPGKRQWVGGESDEFEDEEE <mark>KR</mark> QFSPC <mark>KR</mark> QFSPCKRQFSPGKRQFSPGKR
Op_liod_b	VDMPET <mark>RQFSPGKRQFSPGKRQFSPCKRQFSPGKR</mark> QWVGGESDEFEDEEE <mark>KRQFSPGKRQFSPCKRQFSPGKR</mark> QFSP <mark>GKR</mark> QFSP <mark>GKR</mark> QFSP <mark>GKR</mark>
As_bidw	VDMPETRQFSAGKRQFSAGKRQFSAGKRQFSAGKREWMDDGPDMLEEEDEKRQFSAGKRQFSAGKRQFSAGKRQFSAGKR
Op_oedi	VDMPETRQFSAGKRQFSAGKRQFSAGKRQFSAGKREWMDDGPNMLEEEDEKRQFSAGKRQFSAGKRQFSAGKRQFSAGKR
As_love	VDMPETROFSAGKROFSACKROFSACKROFSAGKREWM-DEPDMLDEEDAKROFSAGKROFSACKROFSAGKROFSAGKROFSAGKR
Op_john	VDMPQTROFSAGKRQFSAGKRQFSAGKRQFSAGKROFSAGKROFSAGKRQFSAGKRQFSAGKRQFSAGKRQFSAGKRQFSAGKRQFSAGKR
Op_lyma	VDIPQT <mark>RQFSAGKRQFSAGKR</mark> QFSA <mark>GKR</mark> QFSA <mark>GKR</mark> QWIGGEDDANEEA <mark>KRQFSAGKR</mark> QFSA <mark>GKR</mark> QFSA <mark>GKR</mark> QFSA <mark>GKR</mark> QFSAGKR
Am_cipu	<mark>QFSAGKR</mark> DWEEE-LTPEELMDMFQAPET <mark>RQFSAGKR</mark> QFSAG <mark>KR</mark> QFSAG <mark>KR</mark> QWVGGEEEYDPEEMLNMAT <mark>RQFSAGKR</mark>
Op_angu	<mark>QFSAGKR</mark> DWEETELTPEEFMDMIPLPET <mark>RQFSAGKR</mark> QFSAG <mark>KR</mark> QFSAG <mark>KR</mark> QFSAGKRQWVGGDLEYEPEEDLDMET <mark>RQFSAGKR</mark> QFS
Op_lame	QFSAG <mark>KR</mark> DWEDE-LTPEDLMDILPAPET <mark>RQFSAGKRQFSAGKRQFSAGKR</mark> QWVGGEYNPDDMLDMET
Op_impr	<mark>QFSAGKR</mark> DWEELTPEDLSDIVAAPET <mark>RQFSAGKR</mark> QFSAG <mark>KR</mark> QFSAG <mark>KR</mark> QWVGGMENPDDMLDMET <mark>RQFSAGKR</mark>
Ba_hero_a	AC <mark>KR</mark> QFSAG <mark>KR</mark> DWEEENLTPQDLLALDMLPLPET <mark>RQFSAGKR</mark> QFSAC <mark>KR</mark> QWVGGELEYDPNEMLDMET <mark>RQFSAGKR</mark>
Ba_hero_b	QFSAG <mark>KR</mark> DWEEENLTPQDLLALDMLPLPET <mark>RQFSAGKRQFSAGKR</mark> QWVGGELEYDPNEMLDMET <mark>RQFSAGKR</mark>
Op_vivi	QFSAGKRDWEEEELTPEDLLALDMLPVPETRQFSAGKRQFSAGKRQFSAGKRQWVGGDLEYNPEEMLDMETRQFSAGKR
Op_perf	QFSAG <mark>KR</mark> DWEEDNLTPQDLLALGMLPIPET <mark>RQFSAGKRQFSAGKRQFSAGKR</mark> QWVGGEQEYDPEDMLDMET <mark>RQFSAGKR</mark>
Op_exim_1	QFSAGKRDWEEEDLTPQDLLALEMLPLPETRQFSAGKRQFSACKRQFSACKRQWVGGEQEYNPEDMLDMETRQFSAGKR
Op_liod_a	QFSPCKREWDND-LTPEDLLAMGLLPAPETROFSPCKRQFSPCKRQFSPCKRQWVGGELEYNPDDMLEMEARQFSPCKR
Op_liod_b	QFSPCKREWDND-LTPEDLLAMGLLPAPETROFSPCKRQFSPCKRQFSPCKRQWVGGELEYNPDDMLEMEARQFSPCKR
As_bidw Op oedi	<mark>QFSACKR</mark> DWEQD-LTPEDYLAMEMLPAPET <mark>RQFSACKRQFSACKRQFSACKRQFSACKR</mark> QWVGGDYDPEELLDMET <mark>RQFSACKR</mark> <mark>QFSACKR</mark> DWEQD-LTPEEYLAMEMLPAPET <mark>RQFSACKRQFSACKRQFSACKRQFSACKR</mark> QWVGGDYDPEELLDMET <mark>RQFSACKR</mark>
As love	DFSACKRDWEQD-LIFEEILAMEMIFAFEIRQFSACKRQFSACKRQFSACKRQWVGGDIDFEELLDMEIRQFSACKR DWRQD-LTPEELLAMEMIFAFEIRQFSACKRQFSACKRQFSACKRQFSACKRQWVGGEYDPEELLNMEARQFSACKR
Op john	QFSAGKRDWEEH-LTPEEYLAMEMMPAPETRQFSAGKRQFSAGKRQFSAGKRQWIGGQEEQEYNPDDFLDMETRQFSAGKR
Op_John Op_lyma	QFSACKADWEEN-DIFEETDAMEMITAFEINQFSACKRQFSACKRQFSACKRQWIGGDEGQEYNPDDFDMEINQFSACKR
op_ryma	
Am_cipu	QFSAGKRQFSAGKRQWVGGEEAFLPEMDTRQFSAGKRQFSAGKRQFSAGKRQFSAGKRDDGETNILDEILEAEPDLAEAE
Op_angu	AG <mark>KRQFSAGKRQFSAGKR</mark> QWVGGDVLPEMETRQFSAGKRQFSAG <mark>KRQFSAGKR</mark> QFSAG <mark>KR</mark> D-ADTDILDQILNADTTEEE RQFSAGKRQFSAGKRQFSAGKRQFSAGKRQFSAGKRQFSAGKR
Op_lame Op impr	<mark>QFSACKRQFSACKR</mark> QWVGGMENPDDMLDMET <mark>RQFSACKRQFSACKROFSACKR</mark> DETNILDEILDPAADDALAE
Ba hero a	QFSACKRQFSAGKRQWVGGMENFDDMLDMEIRQFSAGKRQFSAGKRQFSAGKRDEINILDEILEADFAGEDALAE QFSAGKRQFSAGKRQWVGGDVLPEMDTRQFSAGKRQFSAGKRQFSAGKRDEINILDEILEADFAGEDALAE
Ba hero b	QFSACKRQFSACKRQWVGGDVLPEMDIRQFSACKRQFSACKRQFSACKR QFSACKRQFSACKRQWVGGDVLPEMDTRQFSACKRQFSACKRQFSACKR
Op vivi	QFSAGKRQFSAGKRQWVGGDALPEMETRQFSAGKRQFSAGKRQFSAGKRDETDILDEILQAEPEAEDAFSE
Op perf	QFSAGKRQFSAGKRQWVGGDVLPEMDTRQFSAGKRQFSAGKRQFSAGKRDETNILDEILDAEPAAANALSE
Op exim 1	<mark>QFSAGKR</mark> QFSAGKRQWVGGDVLPEMDT <mark>RQFSAGKRQFSAGKR</mark> QFSAGKR
Op liod a	D-ETNILDEILEAEPAAENALSE
Op liod b	DETNILDEILEAEPAAENALSE
As bidw	<mark>QFSAGKR</mark> QISAGNRQWVGGEALPEMET <mark>RQFSAGKRQFSAGKRQFSAGKR</mark> DESNILHEILNAEPAAANSLSE
Op oedi	QFSAGKRQFSAGKRQWVGGEALPEMETRQFSAGKRQFSAGKRQFSAGKR
As love	DETNILDEILAAEPAVANALSE
Op_john	QWIGGDVIPDMETROFSAGKROFSAGKROFSAGKROFSAGKROFSAGKROFAGKRDDTNILDEFLEANPAENDALSE
Op_lyma	<mark>QFNPGKRQFSAGKR</mark> QWIGGDAIPNMET <mark>RQFSAGKRQFSAGKRQFSAGKR</mark> DETNILDEILENDPAAENALSE

#### F-type SALMFa

Am_cipu				G <mark>KR</mark> RDPSALSAFSFG <mark>KR</mark> RDPM-GLNALTF <mark>GKR</mark> -GMN
Op_filo				G <mark>KR</mark> RDPSGLTAFSFG <mark>KR</mark> RDPL-GLNALTF <mark>GKR</mark> MS
Mi_grac				G <mark>KR</mark> RDPSGLSAFSFG <mark>KR</mark> RDPT-RLSALTFG <mark>KR</mark> -GMS
Am_squa	PLV <mark>RR</mark> <mark>-SAQ</mark> -	-SKPVKLAGFAF <mark>GKR</mark> -GQLE <mark>KR</mark> S	SADDKLMEEDETE <mark>KR</mark> ALSS-AFTF	G <mark>KR</mark> RDPSGLSALTF <mark>G</mark> KRRDPM-GLSALTF <mark>GKR</mark> -GMN
Op_resi	QLV <mark>RR</mark> <mark>-SASS</mark>	GAKPVKLAGFAF <mark>G</mark> KRAGQLV <mark>KR</mark> S	SSDDQLVEEDGAE <mark>KR</mark> AAMD-AFTF	G <mark>KR</mark> YDPSGLSAFSF <mark>GKR</mark> RDPL-GLSALTF <mark>GKR</mark> -GMN
Op abys	SLV <mark>RR</mark> SASSO	GSKPVKLAGFAF <mark>G</mark> KR-GQLV <mark>KR</mark> S	SSDDQLLEEDSTE <mark>KR</mark> AAMD-AFTF	G <mark>KR</mark> MSDPSGLSAFSF <mark>GKR</mark> RDPM-GLSALTF <mark>GKR</mark> -GMT
Op angu	QLV <mark>RR</mark> SAKSG	GDKPVKLAGFAF <mark>GKR</mark> -GOPV <mark>KR</mark> S	STNDELEEDGEE <mark>KR</mark> AAMD-AFTF	G <mark>KR</mark> ISDQE-LSPFSFE <mark>KR</mark> RDPT-GLSALTF <mark>GKR</mark> -GMH
Op_scha	QLVRR	GSKPVKLAGFAF <mark>GKR</mark> -GOLV <mark>KR</mark> S	SSDDQLEEEDEAE <mark>KR</mark> AAMD-AFTF	G <mark>KR</mark> LSKDPSALSAFNFCKRRDPM-GLSALTFCKR-GMD
Op_lame				G <mark>KR</mark> LSNDPSALSAFSF <mark>GKR</mark> RDPM-GLSALTFG <mark>KR</mark> -GMN
Op bisp				G <mark>KR</mark> PSGDPTGLSAFSF <mark>GKR</mark> RDPM-SLSALTF <mark>GKR</mark> -GMD
Op_brev				GKRKAGDLSAFSF <mark>GKR</mark> RDPLSALTF <mark>GKR</mark> -GMK
Ba hero				GKRPSGNPTGLSAFSFGKRREPVGSLSALTFCKR-GMD
Op appr				GKRPSGNPSGLSAFSFGKRREPLGSLSALTFGKR-GTD
Op vivi			SSDDKVEEQDDKRGAMD-AFTF	
Op wend				AKRPSGDPSGLSAFSFGKRRDPVGSLSALTFGKR-AME
Op plic				GKRLSGDPSALSAFSFGKRRDPVSSLSALTFGKR-GMD
Op perf				GKRRSGDPSGLSAFSFGKRRDPASSLSALTFGKR-GMD
Cl cana				GKRLSGGKSALSAFSFGKRRDPVGSLSALTFGKR-GMD
Op exim 1	~ _			GKRLPGDPSALSAFSFGKRRDPVSSLSALTFGKR-GMD
Op liod			~ ~	GKRLSNDPSGLSAFSFGKR-EPMGSLSGLTFGKR-GMD
				GKRLSNDPSGLSAFSFGKR-EPMGSLSGLFFGKR-GMD GKRLSSDPLSAFNFGKRREPVSSLSALTFG <mark>KR</mark> -GMD
Op_prol				
As_tubi				GKRLSGDPSGLSTFSFGKRRNPGTSLSALTFCKR-GMY
Op_oedi				GKRLSGDPSGLSTFSFGKRRNPGTSLSALTFCKR-GMY
Go_pust				GKRLSSDPAAVTFEKR-GMN
As_love				GKRLSGNPSALSAFSFGKRREPGSALSALTFGKR-GMN
Op_john				G <mark>KR</mark> PSGDPTGLSAFSF <mark>GKR</mark> RDPMSSLSALAFG <mark>KR</mark> -GMD
Op_lyma	PLV <mark>RR</mark> <mark>-SAG</mark> A	<mark>GSKPVKLAGFAF<mark>GKR</mark>NPV<mark>KR</mark>S</mark>	SSDNEANDKEE <mark>KRVPMD-AFAF</mark>	G <mark>KR</mark> PSGDPTGLSAFSF <mark>GKR</mark> RDPLSSLSALAF <mark>GKR</mark> -GMD
Am cipu	PASGYSAFTFCKRGOMDNLHAFSFCKR-GMDPSGLS2	FSF <mark>GKR</mark> GRDPSALSAFSF <mark>GKR</mark>		KREGLE-EDGAFE-EENDDEKRNOLSSLUGAUECKR
Am_cipu Op_filo	PASGYSAFTEC <mark>KR</mark> GOMDNLHAFSFC <mark>KR</mark> -GMDPSGLSA	FSFGKRGRDPSALSAFSFGKR		KREGLE-EDGAFE-EENDDEKRNQLSSLTGYTFGKR
Op_filo	PASGYSAFTFC <mark>KR</mark> GOMDNLHAFSFG <mark>KR</mark> -GMDPSGLSA P-SGYSAFTFC <mark>KR</mark> GOMDNLHAFSFC <mark>KR</mark> -GMDPSSLSA P-SGYSAFTFCKRGRMDNLNAFSFCKR-GMDPSTLSA	FSFGKRGRDPSALSAFSFCKR LTFGKRGRDPSSLSAFSFCKR FSFGKRGRDPSALSAFSFCKR		KREGLE-EDGAFE-EENDDEKRNQLSSLTGYTFGKR KRDELE-EDGAFE-DENDDEKRSRLSSLTGYTFGKR KRDELE-EDGAFE-EENDDEKRSYSKR
Op_filo Mi_grac				KREGLE-EDGAFE-EENDDEKRNQLSSLTGYTFGKR KRDELE-EDGAFE-DENDDEKRSRLSSLTGYTFGKR KRDELE-EDGAFE-EENDDEKRSYSKR KRDEEDGAFE-EENVDEKRSRIGALTGLTYGKR
Op_filo Mi_grac Am_squa	P-SGYSAFTFG <mark>KR</mark> GRMDNLNAFSF <mark>GKR</mark> -GMDPSGLS#	.FSF <mark>GKR</mark> GRDPSALSAFSF <mark>GKR</mark>	PAFTF <mark>G</mark>	KRDEEDGAFE-EENYDE <mark>KR</mark> SRIGALTGLTYGKR
Op_filo Mi_grac Am_squa Op_resi	P-SGYSAFTF <mark>CKR</mark> GRMDNLNAFSF <mark>CKR-</mark> GMDPSGLSA P-SGMSAFSF <mark>CKR</mark> -RMEPLSAFSF <mark>CRKR</mark> GMDPSGLSA	FSFC <mark>KR</mark> GRDPSALSAFSFC <mark>KR</mark> FSFC <mark>KR</mark> GMDPSGLSAFSFC <mark>KR</mark>	PAFTFG MG-M-NAFTFG	<mark>KR</mark> DEEDGAFE-EENYDE <mark>KR</mark> SRIGALTGLTYCKR KREGGEEEDPAFE-EENNN-EE <mark>KRAGYNGLSQFTFC</mark> KR
Op_filo Mi_grac Am_squa Op_resi Op_abys	P-SGYSAFTFC <mark>KR</mark> GRMDNLNAFSFC <mark>KR-</mark> GMDPSGLS& P-SGMSAFSFC <mark>KR</mark> -RMEPLSAFSFC <mark>KKR</mark> GMDPSGLS& P-SGMSAFSFC <mark>KR</mark> -RMEPLSAFSFC <mark>RKR</mark> GMDPSGLS&	FSF <mark>CKR</mark> GRDPSALSAFSF <mark>CKR</mark> FSF <mark>CKR</mark> GMDPSGLSAFSF <mark>CKR</mark> FSF <mark>CKR</mark> GMDPLGLNAFSF <mark>CKR</mark>		KRDEEDGAFE-EENYDEKRSRIGALTGLTYGKR KREGGEEEDPAFE-EENNN-EEKRAGYNGLSQFTFGKR KREGLEEEDAALE-EEDNNDDEKR <mark>AGYNGLSQFTFGKR</mark>
Op_filo Mi_grac Am_squa Op_resi Op_abys Op_angu	P-SGYSAFTFC <mark>KR</mark> GRMDNLNAFSFC <mark>KR-</mark> GMDPSGLSA P-SGMSAFSFC <mark>KR</mark> -RMEPLSAFSFC <mark>KKR</mark> GMDPSGLSA P-SGMSAFSFC <mark>KR</mark> -RMEPLSAFSFC <mark>KKR</mark> GMDPSGLSA P-SSMSAFSFC <mark>KR</mark> -RMDPLSAFSFC <mark>KKR</mark> AMDPAGLSA	FSF <mark>GKR</mark> GRDPSALSAFSFG <mark>KR</mark> - FSF <mark>GKRGMDPSGLSAFSFGKR-</mark> FSF <mark>GKRGMDPLGLNAFSFGKR-</mark> FSF <mark>GKRGMDPSALSAFSFG</mark> KR <mark>G</mark> 1	PAFTFG MG-M-NAFTFG 	KRDEEDGAFE-EENYDEKRSRIGALTGLTYGKR KREGGEEEDPAFE-EENNN-EEKRAGYNGLSQFTFGKR KREGLEEEDAALE-EEDNNDDEKRAGYNGLSQFTFGKR KREGEE-EETAFKKNTNDDEKRAGYNGLSQFTFGKR
Op_filo Mi_grac Am_squa Op_resi Op_abys Op_angu Op_scha	P-SGYSAFTFC <mark>KR</mark> GRMDNLNAFSFC <mark>KR-</mark> GMDPSGLSA P-SGMSAFSFC <mark>KR-RMEPLSAFSFCRKR</mark> GMDPSGLSA P-SGMSAFSFC <mark>KR-RMEPLSAFSFCKRGMDPSGLSA P-SSMSAFSFCKR-RMDPLSAFSFCKRAMDPAGLSA P-SGFSAFSFC<mark>KR-R-EPYSAFSFCKR-GMDPSALS</mark>A</mark>	FSF <mark>GKR</mark> GRDPSALSAFSFG <mark>KR</mark> FSF <mark>GKR</mark> GMDPSGLSAFSFG <mark>KR</mark> FSF <mark>GKR</mark> GMDPLGLNAFSFG <mark>KR</mark> FSF <mark>GKR</mark> GMDPSALSAFSFG <mark>KR</mark> G <sup>T</sup> FSF <mark>GKR</mark> ARDPSALSAFNF <mark>G</mark> K <mark>R</mark> -	PAFTFG 	KRDEEDGAFE-EENYDEKRSRIGALTGLTYGKR KREGGEEEDPAFE-EENNN-EEKRAGYNGLSQFTFGKR KREGLEEEDAALE-EEDNNDDEKRAGYNGLSQFTFGKR KREGEE-EETAFKKNTNDDEKRAGYNGLSQFTFGKR KREGLEEDGAFE-EENQDEEEK <mark>RGGYNGIAGYTFGKR</mark>
Op_filo Mi_grac Am_squa Op_resi Op_abys Op_angu Op_scha Op_lame	P-SGYSAFTFCKRGRMDNLNAFSFCKR-GMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SSMSAFSFCKR-RMDPLSAFSFCKRAMDPAGLSA P-SGFSAFSFCKR-R-EPYSAFSFCKR-GMDPSALSA P-SGFSAFTYCKR-R-EPLSAFSFCKR-GMDPSALSA	FSF <mark>GKR</mark> GRDPSALSAFSFG <mark>KR</mark> FSF <mark>GKRGMDPSGLSAFSFGKR</mark> FSF <mark>GKRGMDPLGLNAFSFGKR</mark> FSF <mark>GKRGMDPSALSAFSFGKRG</mark> FSF <mark>GKR</mark> ARDPSALSAFNFG <mark>KR</mark> FSF <mark>GKR</mark> GRDPSALSAFNF <mark>GKR</mark> -	PAFTFG 	KRDEEDGAFE-EENYDEKRSRIGALTGLTYGKR KREGGEEEDPAFE-EENNN-EEKRAGYNGLSQFTFGKR KREGLEEEDAALE-EEDNNDDEKRAGYNGLSQFTFGKR KREGEE-EETAFKKNTNDDEKRAGYNGLSQFTFGKR KREGLEEDGAFE-EENQDEEEKRGGYNGIAGYTFGKR KRDDLEEDGAFE-EEENQEEEKRGGYNGISGYTFGKR
Op_filo Mi_grac Am_squa Op_resi Op_abys Op_angu Op_scha Op_lame Op_bisp	P-SGYSAFTFCKRGRMDNLNAFSFCKR-GMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKKRGMDPSGLSA P-SSMSAFSFCKR-RMDPLSAFSFCKRAMDPAGLSA P-SGFSAFSFCKR-R-EPYSAFSFCKR-GMDPSALSA P-SGFSAFTYCKR-R-EPLSAFSFCKR-GMDPSALSA P-SGFSAFSFCKR-R-DPFSALTFCKR-GMDPSALSA	FSF <mark>GKR</mark> GRDPSALSAFSFG <mark>KR</mark> FSFGKRGMDPSGLSAFSFG <mark>KR</mark> FSFGKRGMDPLGLNAFSFG <mark>KR</mark> FSFGKRGMDPSALSAFSFG <mark>KR</mark> GT FSFGKRARDPSALSAFNFG <mark>KR</mark> FSFGKRGRDPSALSAFNFG <mark>KR</mark> YSFG <mark>KR</mark> GRDPSALSAFNFG <mark>KR</mark> -		KRDEEDGAFE-EENYDEKRSRIGALTGLTYGKR KREGGEEEDPAFE-EENNN-EEKRAGYNGLSQFTFGKR KREGLEEEDAALE-EEDNNDDEKRAGYNGLSQFTFGKR KREGEE-EETAFKKNTNDDEKRAGYNGLSQFTFGKR KREGLEEDGAFE-EENQDEEEKRGGYNGIAGYTFGKR KRDDLEEDGAFE-EEENQEEEKRGGYNGISGYTFGKR KRDDAEEDGAFE-EDNNDEKR-GFNGISGYTFGKR
Op_filo Mi_grac Am_squa Op_resi Op_abys Op_angu Op_scha Op_lame Op_bisp Op_brev	P-SGYSAFTFCKRGRMDNLNAFSFCKR-GMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCRKRGMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCRKRGMDPSGLSA P-SSMSAFSFCKR-RMDPLSAFSFCKRAMDPAGLSA P-SGFSAFSFCKR-R-EPLSAFSFCKR-GMDPSALSA P-SGFSAFTYCKR-R-EPLSAFSFCKR-GMDPSALSA P-SGFSAFSFCKR-R-DPFSALTFCKR-GMDPSALSA P-SAFDAFSFCKR-R-DPLSAFSFCKR-GMDPSALSA	FSFG <mark>KR</mark> GRDPSALSAFSFG <mark>KR</mark> FSFG <mark>KRGMDPSGLSAFSFG</mark> KR FSFG <mark>KRGMDPSGLSAFSFG</mark> KR- FSFG <mark>KR</mark> GMDPSALSAFSFG <mark>KR FSFGKRGRDPSALSAFNFGKR YSFG<mark>KR</mark>GRDPSALSAFNFG<mark>KR FSFG<mark>KR</mark>GRD-NALGAFSFC<mark>KR</mark></mark></mark>	-MGPAFTFG -MG-M-NAFTFG -MG-M-NAFTFG TGPS-GLSAFSFGKR-MG-M-NAFTFG -MGGMTNAFTFG -MGGLTNAFTFG -MGGLTNAFTFG GM-DAFTFG	KRDEEDGAFE-EENYDEKRSRIGALTGLTYGKR KREGGEEEDPAFE-EENNN-EEKRÄGYNGLSQFTFGKR KREGLEEEDAALE-EEDNNDDEKRAGYNGLSQFTFGKR KREGLEEETAFKKNTNDDEKRAGYNGLSQFTFGKR KRDLEEDGAFE-EENQDEEEKRGGYNGLSGYTFGKR KRDDAEEDGAFE-EENNDEKR-GFNGISGYTFGKR KRDDAEEGAFE-EDNNDEKR-GFNGISGYTFGKR
Op_filo Mi_grac Am_squa Op_resi Op_abys Op_angu Op_scha Op_lame Op_bisp Op_brev Ba_hero	P-SGYSAFTFCKR-RMEPLSAFSFCKR-GMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKKRGMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKKRGMDPSGLSA P-SGFSAFSFCKR-RMEPLSAFSFCKR-GMDPSALSA P-SGFSAFSFCKR-R-EPLSAFSFCKR-GMDPSALSA P-SGFSAFSFCKR-R-DPFSALTFCKR-GMDPSALSA P-SAFDAFSFCKR-R-DPLSAFSFCKR-GMDPSALSA P-SAFDAFSFCKR-R-DPLSAFSFCKR-GMDPSALSA P-AGFSAFNFCKR-R-DPLSAFNFCKR-GMDPSGLSA	FSFG <mark>KR</mark> GRDPSALSAFSFG <mark>KR</mark> FSFG <mark>KR</mark> GMDPSGLSAFSFG <mark>KR</mark> FSFG <mark>KR</mark> GMDPSALSAFSFG <mark>KR</mark> FSFG <mark>KR</mark> GRDPSALSAFNFG <mark>KR</mark> FSFG <mark>KR</mark> GRDPSALSAFNFG <mark>KR</mark> YSFG <mark>KR</mark> GRDPSALSAFNFG <mark>KR</mark> FSFG <mark>KR</mark> GRDPSGLSAFSFC <mark>KR</mark>	-MGPAFTFG -MG-M-NAFTFG -MG-M-NAFTFG -MGM-NAFTFG -MGMTNAFTFG -MGMTNAFTFG -MGGLTNAFTFG -MGGLTNAFTFG -MGGLTNAFTFG RVPSLSAFDFC <mark>KR</mark> G-M-DAFTFG	KRDEEDGAFE-EENYDEKRSRIGALTGLTYGKR KREGGEEEDPAFE-EENNN-EEKRAGYNGLSOFTFGKR KREGLEEEDAALE-EEDNNDDEKRAGYNGLSOFTFGKR KREGLEEETAFKKNTNDDEKRAGYNGLSOFTFGKR KREGLEEDGAFE-EENQDEEEKRGGYNGISGYTFGKR KRDDAEEDGAFE-EENNDEKR-GFNGISGYTFGKR KRDDAEEGAFE-DEDEKR-AYNPISAYTFGKR KRDDEEGAFE-DENDDEKR-GFNGISGYTFGKR
Op_filo Mi_grac Am_squa Op_resi Op_abys Op_angu Op_scha Op_lame Op_lame Op_bisp Op_brev Ba_hero Op_appr	P-SGYSAFTFCKR-RMEPLSAFSFCKR-GMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SSMSAFSFCKR-RMEPLSAFSFCKR-GMDPSALSA P-SGFSAFSFCKR-R-EPLSAFSFCKR-GMDPSALSA P-SGFSAFSFCKR-R-DPLSAFSFCKR-GMDPSALSA P-SAFDAFSFCKR-R-DPLSAFSFCKR-GMDPSALSA P-AGFSAFNFCKR-R-DPLSAFNFCKR-GMDPSGLSA P-AGFSAFNFCKR-R-DPLSAFNFCKR-GMDPSGLSA	FSF <mark>G KR</mark> GRDPSALSAFSFG KR FSFG KRGMDPSGLSAFSFG KR FSFG KRGMDPSALSAFSFG KR FSFG KRGMDPSALSAFNFG KR FSFG KRGRDPSALSAFNFG KR FSFG KRGRDPSALSAFNFG KR FSFG KRGRD - NALGAFSFG KR FSFG KRGRDPSGLSAFSFG KR SF FSFG KRGRDPSGLSAFSFG KR SF		KRDEEDGAFE-EENYDEKRSRIGALTGLTYGKR KREGGEEEDPAFE-EENNN-EEKRAGYNGLSQFTFGKR KREGLEEEDAALE-EEDNNDDEKRAGYNGLSQFTFGKR KREGLEEETAFKKNTNDDEKRAGYNGLSQFTFGKR KRDDLEEDGAFE-EENQEEEKRGGYNGIAGYTFGKR KRDDAEEDGAFE-EENNDEKR-GFNGISGYTFGKR KRDDAEEGAFE-DENNDEKR-AYNPISAYTFGKR KREDLD-EEGAFE-DENDDEKR-GFNGISGYTFGKR KREDLD-EEGAFE-DENDDEKR-GFNGISGYTFGKR
Op_filo Mi_grac Am_squa Op_resi Op_abys Op_angu Op_scha Op_lame Op_lame Op_bisp Op_brev Ba_hero Op_appr Op_vivi	P-SGYSAFTFCKR-RMEPLSAFSFCKR-GMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SGFSAFSFCKR-RMDPLSAFSFGKR-GMDPSGLSA P-SGFSAFSFCKR-R-EPLSAFSFGKR-GMDPSGLSA P-SGFSAFTYCKR-R-DPLSAFSFCKR-GMDPSGLSA P-SGFSAFNFCKR-R-DPLSAFSFGKR-GMDPNALGA P-AGFSAFNFCKR-R-DPLSAFNFGKR-GMDPSGLSA P-SGFSAFNFCKR-R-DPLSAFNFGKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFGKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFGKR-GMDASGLSA	FSFG <mark>KR</mark> GRDPSALSAFSFG <mark>KR</mark> FSFG <mark>KR</mark> GMDPSGLSAFSFG <mark>KR</mark> FSFG <mark>KR</mark> GMDPSALSAFSFG <mark>KR</mark> FSFG <mark>KR</mark> GRDPSALSAFNFG <mark>KR</mark> FSFG <u>KR</u> GRDPSALSAFNFG <mark>KR</mark> FSFG <u>KR</u> GRD-NALGAFSFG <mark>KR</mark> FSFG <u>KR</u> GRDPSGLSAFSFG <mark>KR</mark> FSFG <u>KR</u> GRDPSGLSAFSFG <mark>KR</mark>		KRDEEDGAFE-EENYDEKRSRIGALTGLTYGKR KREGGEEEDPAFE-EENNN-EEKRAGYNGLSQFTFGKR KREGLEEEDAALE-EEDNNDD-EKRAGYNGLSQFTFGKR KREGLEEETAF <mark>KK</mark> NTNDDEKRAGYNGLSQFTFGKR KREDLEEDGAFE-EENQDEEEKRGGYNGIAGYTFGKR KRDDLEEDGAFE-EENNDEKR-GFNGISGYTFGKR KRDDEEGAFE-DENNDEKR-AYNFISAYTFGKR KREDLD-EEGAFE-DENDDEKR-GFNGISGYTFGKR KREDLD-EEGAFE-DENDDEKR-GFNGISGYTFGKR KREDLD-EDGAFE-DENDDEKR-GFNGISGYTFGKR
Op_filo Mi_grac Am_squa Op_resi Op_abys Op_angu Op_scha Op_lame Op_lame Op_bisp Op_brev Ba_hero Op_op_vivi Op_vivi Op_vivi	P-SGYSAFTFCKR-RMEPLSAFSFCKR-GMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SGFSAFSFCKR-RMDPLSAFSFCKR-GMDPSGLSA P-SGFSAFSFCKR-R-EPLSAFSFCKR-GMDPSALSA P-SGFSAFSFCKR-R-DPLSAFSFCKR-GMDPSALSA P-SGFSAFFFCKR-R-DPLSAFSFCKR-GMDPSGLSA P-AGFSAFNFCKR-R-DPLSAFNFCKR-GMDPSGLSA P-AGFSAFNFCKR-R-DPLSAFNFCKR-GMDAGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDAGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDAGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDAGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA	FSFG <mark>KR</mark> GRDPSALSAFSFG <mark>KR</mark> FSFG <mark>KRGMDPSGLSAFSFG</mark> KR FSFG <mark>KRGMDPSALSAFSFG</mark> KR FSFG <mark>KR</mark> GRDPSALSAFNFG KR SFG <mark>KRGRDPSALSAFNFG FSFG KRGRDPSALSAFNFG FSFG KRGRDPSGLSAFSFG KRGRDPSGLSAFSFG FSFG KRGRDSGLSAFSFG KRGRDSGLSAFSFG KRGRDSGLSAFSFG KRGRDSGLSAFSFG KRGRDSGLSAFSFG KRGRDSGLSAFSFG KRGRDSGLSAFSFG KRGRDSGLSAFSFG KRGRDSGLSAFSFG KRGRDSGLSAFSFG KRGRDSGLSAFSFG KRGRDSGLSAFSFG KRGRDSGLSAFSFG</mark>		KRDEEDGAFE-EENYDEKRSRIGALTGLTYGKR KREGGEEEDPAFE-EENNN-EEKRAGYNGLSQFTFGKR KREGLEEEDAALE-EEDNNDDEKRAGYNGLSQFTFGKR KREGEE-EETAF <mark>KK</mark> NTNDDEKRAGYNGLSQFTFGKR KRDDLEEDGAFE-EENQDEEEKRGGYNGISGYTFGKR KRDDAEEDGAFE-EENND-EKR-GFNGISGYTFGKR KRDDAEEGAFE-DENND-EKR-GFNGISGYTFGKR KREDLD-EEGAFE-DENDDEKR-GFNGISGYTFGKR KREDLD-EEGAFE-DENDDEKR-GFNGISGYTFGKR KREDLD-EGAFE-DENDDEKR-GFNGISGYTFGKR KREDLD-EGAFE-EENDDEKR-GFNGISGYTFGKR KREELD-DEGAFE-EENEDEKR-GFNGISGYTFGKR
Op_filo Mi_grac Am_squa Op_resi Op_abys Op_angu Op_scha Op_lame Op_bisp Op_brev Ba_hero Op_brev Ba_hero Op_vivi Op_vivi Op_vivi	P-SGYSAFTFCKR-RMEPLSAFSFCKR-GMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SGFSAFSFCKR-RMDPLSAFSFCKR-GMDPSGLSA P-SGFSAFSFCKR-R-EPLSAFSFCKR-GMDPSGLSA P-SGFSAFSFCKR-R-DPLSAFSFCKR-GMDPSGLSA P-SGFSAFNFCKR-R-DPLSAFSFCKR-GMDPSGLSA P-AGFSAFNFCKR-R-DPLSAFNFCKR-GMDPSGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLGAFSFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLGAFSFCKR-GMDASGLSA	FSFG <mark>KR</mark> GRDPSALSAFSFG <mark>KR</mark> FSFG <mark>KRGMDPSGLSAFSFG</mark> KR FSFG <mark>KRGMDPSALSAFSFG</mark> KR FSFG <mark>KR</mark> GRDPSALSAFNFG KR YSFG <del>KR</del> GRDPSALSAFNFG FSFG KRGRDPSGLSAFSFG FSFG KRGRDPSGLSAFSFG FSFG KRGRDPSGLSAFSFG FSFG KRGRDSGLSAFSFG FSFG KRGRDATGLSAFSFG KRGRDAAGLSAFSFG KRGRDAAGLSAFSFG KRGRDAAGLSAFSFG KRGRDAAGLSAFSFG KRGRDAAGLSAFSFG KRGRDAAGLSAFSFG KRGRDAAGLSAFSFG KRGRDAAGLSAFSFG KRGRDAAGLSAFSFG KRGRDAAGLSAFSFG KRGRDAAGLSAFSFG		KRDEEDGAFE-EENYDEKRSRIGALTGLTYGKR KREGGEEEDPAFE-EENNN-EEKRAGYNGLSQFTFGKR KREGLEEEDAALE-EEDNNDDEKRAGYNGLSQFTFGKR KREGEE-EETAF <mark>KK</mark> NTNDDEKRAGYNGLSQFTFGKR KRDDLEEDGAFE-EENQDEEEKRGGYNGISGYTFGKR KRDDLEEDGAFE-EENND-EKR-GFNGISGYTFGKR KRDDAEEGAFE-DENDDEKR-AYNPISAYTFGKR KREDLD-EEGAFE-DENDDEKR-GFNGISGYTFGKR KREDLD-EGAFE-DENDDEKR-GFNGISGYTFGKR KREDLD-EGAFE-DENDDEKR-GFNGISGYTFGKR KRELD-DEGAFE-EENEDEKR-GFNGISGYTFGKR KRELD-DEGAFE-EENEDEKR-GFNGISGYTFGKR KREDLEEEDGAFE-EENEDEKR-GFNGISGYTFGKR KREDLD-EEGAFE-DENDNEKR-GYQGISGYTLGKR KREDLEEEGAFE-DENDNEKR-GYQGISGYTLGKR
Op_filo Mi_grac Am_squa Op_resi Op_abys Op_angu Op_scha Op_lame Op_bisp Op_brev Ba_hero Op_op_vivi Op_wend Op_vivi Op_wend	P-SGYSAFTFCKR-RMEPLSAFSFCKR-GMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SGFSAFSFCKR-RMDPLSAFSFCKRAMDPAGLSA P-SGFSAFSFCKR-R-PPLSAFSFCKR-GMDPSALSA P-SGFSAFSFCKR-R-DPLSAFSFCKR-GMDPSALSA P-SGFSAFSFCKR-R-DPLSAFSFCKR-GMDPSALSA P-AGFSAFNFCKR-R-DPLSAFNFCKR-GMDPSGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLGAFSFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLGAFSFCKRGGMDATGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA	FSFGKRGRDPSALSAFSFGKR FSFGKRGMDPSGLSAFSFGKR FSFGKRGMDPSALSAFSFGKR FSFGKRGRDPSALSAFNFGKR YSFGKRGRDPSALSAFNFGKR 		KRDEEDGAFE-EENYDEKRSRIGALTGLTYGKR KREGGEEEDPAFE-EENNN-EEKRAGYNGLSQFTFGKR KREGLEEEDAALE-EEDNNDDEKRAGYNGLSQFTFGKR KREGEE-EETAFKKNTNDDEKRAGYNGLSQFTFGKR KRDDLEEDGAFE-EENQEEEKRGGYNGISGYTFGKR KRDDLEEDGAFE-EENND-EKR-GFNGISGYTFGKR KRDDAEEDGAFE-DENDD-EKR-GFNGISGYTFGKR KREDLD-EEGAFE-DENDDEKR-GFNGISGYTFGKR KRELL-D-EGGAFE-DENDD-EKR-GFNGISGYTFGKR KRELL-D-DEGAFE-EENEDEKR-FNGISGYTFGKR KRELL-D-DEGAFE-EENED-EKR-FNGISGYTFGKR KRELL-D-EEGAFE-EENED-EKR-FNGISGYTFGKR KRELL-D-EEGAFE-EENED-EKR-FNGISGYTFGKR KRELL-D-EEGAFE-EENED-EKR-FNGISGYTFGKR KREGLEEEGAFE-ENDD-EKR-GYQGISGYTLGKR KREGL-D-EEGAFE-ENDD-EKR-FNGISGLTFGKR
Op_filo Mi_grac Am_squa Op_resi Op_abys Op_angu Op_scha Op_lame Op_bisp Op_brev Ba_hero Op_operv Op_vivi Op_wend Op_vic Op_perf Cl_cana	P-SGYSAFTFCKR GRMDNLNAFSFCKR-GMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCRKRGMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCRKRGMDPSGLSA P-SGFSAFSFCKR-RMDPLSAFSFCKR-GMDPSALSA P-SGFSAFTYCKR-R-EPLSAFSFCKR-GMDPSALSA P-SGFSAFTYCKR-R-DPLSAFSFCKR-GMDPSALSA P-SGFSAFSFCKR-R-DPLSAFSFCKR-GMDPSALSA P-AGFSAFNFCKR-R-DPLSAFNFCKR-GMDPSGLSA P-AGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLGAFSFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLGAFSFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLGAFSFCKRGGMDATGLSA P-SGFSAFNFCKR-R-DPLGAFSFCKRGGMDATGLSA P-SGFSAFNFCKR-R-DPLGAFSFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFGKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFGKRGGMDASGLSA	FSFGKRGRDPSALSAFSFGKR FSFGKRGMDPSGLSAFSFGKR FSFGKRGMDPSALSAFSFGKR FSFGKRGMDPSALSAFNFGKR FSFGKRGRDPSALSAFNFGKR FSFGKRGRDPSALSAFNFGKR FSFGKRGRDPSGLSAFSFGKR FSFGKRGRDPSGLSAFSFGKR FSFGKRGRDSGLSAFSFGKR FNFGKRGRDANGLSAFSFGKR FSFGKRGRDAGLSAFSFGKR FSFGKRGRDASGLSAFSFGKR FSFGKRGRDASGLSAFSFGKR FSFGKRGRDASGLSAFSFGKR		KRDEEDGAFE-EENYD-EKRSRIGALTGLTYGKR KREGGEEEDPAFE-EENNN-EEKRAGYNGLSQFTFGKR KREGLEEEDAALE-EEDNNDDEKRAGYNGLSQFTFGKR KREGLEEDGAFE-EENQDEEEKRGGYNGIAGYTFGKR KRDDLEEDGAFE-EENQDEEEKRGGYNGISGYTFGKR KRDDAEEDGAFE-DENDD-EKR-GFNGISGYTFGKR KRDDAEEGAFE-DENDD-EKR-GFNGISGYTFGKR KREDLD-EEGAFE-DENDD-EKR-GFNGISGYTFGKR KREDLD-EGGAFE-DENDD-EKR-GFNGISGYTFGKR KREDLD-EGGAFE-DENDD-EKR-GFNGISGYTFGKR KREDLD-EGGAFE-DENDN-EKR-GFNGISGYTFGKR KREDLD-EGGAFE-EENDD-EKR-NFNGISGYTFGKR KREDLD-EEGAFE-DENDN-EKR-GYQGISGYTLGKR KREGLD-EEGAFE-ENDD-EKR-FNGISGLTFGKR KREGLDEGGAFL-ENDD-EKR-FNGISGLTFGKR
Op_filo Mi_grac Am_squa Op_resi Op_abys Op_angu Op_scha Op_lame Op_bisp Op_brev Ba_hero Op_op_brev Ba_hero Op_appr Op_vivi Op_wend Op_plic Op_perf Cl_cana Op_exim 1	P-SGYSAFTFCKR-RMEPLSAFSFCKR-GMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCRKRGMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCRKRGMDPSGLSA P-SGFSAFSFCKR-RMDPLSAFSFCKR-GMDPSALSA P-SGFSAFTYCKR-R-EPLSAFSFCKR-GMDPSALSA P-SGFSAFTYCKR-R-DPLSAFSFCKR-GMDPSALSA P-SAFDAFSFCKR-R-DPLSAFSFCKR-GMDPSALSA P-AGFSAFNFCKR-R-DPLSAFNFCKR-GMDPSGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLGAFSFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLGAFSFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA	FSFGKRGRDPSALSAFSFGKR FSFGKRGMDPSGLSAFSFGKR FSFGKRGMDPSALSAFSFGKR FSFGKRGMDPSALSAFSFGKR FSFGKRGRDPSALSAFNFGKR YSFGKRGRDPSALSAFNFGKR FSFGKRGRDPSGLSAFSFGKRGF FSFGKRGRDSGLSAFSFGKRGF FSFGKRGRDSGLSAFSFGKRGF FSFGKRGRDAGLSAFSFGKRGF FSFGKRGRDAGLSAFSFGKRGF FSFGKRGRDAGLSAFSFGKRGF FSFGKRGRDAGLSAFSFGKRGF FSFGKRGRDAGLSAFSFGKRGF		KRDEEDGAFE-EENYD-EKRSRIGALTGLTYGKR KREGGEEEDPAFE-EENNN-EEKRAGYNGLSQFTFGKR KREGLEEEDAALE-EEDNNDDEKRAGYNGLSQFTFGKR KREGLEEDGAFE-EENQDEEEKRGGYNGLSQFTFGKR KRDDAEEDGAFE-EENQDEEEKRGGYNGISGYTFGKR KRDDAEEDGAFE-DENDD-EKR-GFNGISGYTFGKR KRDDAEEGAFE-DENDD-EKR-GFNGISGYTFGKR KREDLD-EEGAFE-DENDD-EKR-GFNGISGYTFGKR KREDLD-EEGAFE-DENDD-EKR-GFNGISGYTFGKR KREDLD-EDGAFE-EENED-EKR-GFNGISGYTFGKR KREDLD-EGGAFE-EENED-EKR-GFNGISGYTFGKR KREDLEEDGAFE-EENDD-EKR-GFNGISGYTFGKR KREDL-D-EEGAFE-EENDD-EKR-GFNGISGYTFGKR KREDL-D-EEGAFE-EENDD-EKR-FNGISGYTFGKR KREGLDEEGAFE-ENDD-EKR-FNGISGTFGKR KREGL-DEEGAFE-EENDD-EKR-FNGISGTFGKR KREGL-DEEGAFE-EENDD-EKR-FNGISGTFGKR KREGL-DEEGAFE-EENDD-EKR-FNGISGTFGKR KREGL-DEEGAFE-EENDD-EKR-FNGISGTFGKR
Op_filo Mi_grac Am_squa Op_resi Op_abys Op_angu Op_scha Op_lame Op_bisp Op_brev Ba_hero Op_brev Ba_hero Op_vivi Op_vivi Op_wend Op_plic Op_perf Cl_cana Op_exim_1 Op_liod	P-SGYSAFTFCKR-RMEPLSAFSFCKR-GMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SGFSAFSFCKR-RMEPLSAFSFCKR-GMDPSALSA P-SGFSAFTYCKR-R-EPLSAFSFCKR-GMDPSALSA P-SGFSAFTYCKR-R-DPLSAFSFCKR-GMDPSALSA P-SGFSAFTYCKR-R-DPLSAFSFCKR-GMDPSALSA P-AGFSAFNFCKR-R-DPLSAFNFCKR-GMDPSGLSA P-AGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLGAFSFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLGAFSFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLGAFSFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLGAFSFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA	FSFG KRGRDPSALSAFSFG KR FSFG KRGMDPSGLSAFSFG KR FSFG KRGMDPSALSAFSFG KR FSFG KRGMDPSALSAFNFG KR YSFG KRGRDPSALSAFNFG KR YSFG KRGRDPSALSAFNFG KR FSFG KRGRDPSGLSAFSFG KR SF FSFG KRGRDPSGLSAFSFG KR SF FSFG KRGRDASGLSAFSFG KR SF FSFG KRGRDATGLSAFSFG KR SF		KRDEEDGAFE-EENYD-EKRSRIGALTGLTYGKR KREGGEEEDPAFE-EENNN-EEKRÄGYNGLSOFTFGKR KREGLEEEDAALE-EEDNNDDEKRÄGYNGLSOFTFGKR KREGLEEDGAFE-EENQDEEEKRGGYNGLSOFTFGKR KRDDAEEDGAFE-EENQDEEEKRGGYNGISGYTFGKR KRDDAEEDGAFE-EENND-EKR-GFNGISGYTFGKR KRDDAEEGAFE-DEDND-EKR-GFNGISGYTFGKR KREDLD-EEGAFE-DENDDEKR-GFNGISGYTFGKR KREDLD-EGGAFE-DENEDEKR-GFNGISGYTFGKR KREDLD-EDGAFE-EENED-EKR-GFNGISGYTFGKR KREDLD-EDGAFE-EENED-EKR-GFNGISGYTFGKR KREDLD-EGGAFE-EENDD-EKR-GFNGISGYTFGKR KREGL-D-EEGAFE-EENDD-EKR-GFNGISGYTFGKR KREGL-D-EEGAFE-ENDD-EKR-FNGISGYTFGKR KREGL-D-EEGAFE-ENDD-EKR-FNGISGLTFGKR KREGL-D-EEGAFE-ENDD-EKR-FNGISGLTFGKR KREGL-D-EEGAFE-EENDD-EKR-FNGISGYTFGKR KREGL-D-EEGAFE-EENDD-EKR-FNGISGYTFGKR KREGL-D-EEGAFE-EENDD-EKR-FNGISGYTFGKR KREGL-D-EEGAFE-EENDD-EKR-FNGISGYTFGKR KREGL-D-EEGAFE-EENDD-EKR-FNGISGYTFGKR KREGL-D-EEGAFE-DENDD-EKR-FNGISGYTFGKR
Op_filo Mi_grac Am_squa Op_resi Op_abys Op_angu Op_scha Op_bisp Op_brev Ba_hero Op_brev Ba_hero Op_op_vivi Op_vivi Op_vivi Op_vivi Op_perf Cl_cana Op_exim_1 Op_liod Op_prol	P-SGYSAFTFCKR-RMEPLSAFSFCKR-GMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SGFSAFSFCKR-RMEPLSAFSFCKR-GMDPSALSA P-SGFSAFSFCKR-R-EPLSAFSFCKR-GMDPSALSA P-SGFSAFSFCKR-R-DPLSAFSFCKR-GMDPSALSA P-SGFSAFFCKR-R-DPLSAFSFCKR-GMDPSALSA P-AGFSAFNFCKR-R-DPLSAFNFCKR-GMDPSGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLGAFSFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLGAFSFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDATGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDATGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFSFCKR-R-DPLSAFNFCKRGGMDASGLSA	FSFG KRGRD PSALSAFSFG KR FSFG KRGMD PSGLSAFSFG KR FSFG KRGMD PSALSAFSFG KR FSFG KRGMD PSALSAFNFG KR YSFG KRGRD PSALSAFNFG KR FSFG KRGRD PSALSAFNFG KR FSFG KRGRD PSGLSAFSFG KR GF FSFG KRGRD PSGLSAFSFG KR GF FSFG KRGRD AGLSAFSFG KR GF FSFG KR GRD FSG KR GF FSFG KR GF FSFG KR GRD FSG KR GF FSFG KR GRD FSG KR GF FSFG KR		KRDEEDGAFE-EENYDEKR SRIGALTGLTYGKR KREGGEEEDPAFE-EENNN-EEKRAGYNGLSQFTFGKR KREGLEEEDAALE-EEDNNDDEKRAGYNGLSQFTFGKR KREGLEEDGAFE-EENQDEEEKRGGYNGISGYTFGKR KRDDLEEDGAFE-EENQDEEEKRGGYNGISGYTFGKR KRDDAEEDGAFE-EENNDEKR-GFNGISGYTFGKR KRDDEEGAFE-DEDNDEKR-GFNGISGYTFGKR KREDLD-EEGAFE-DENDDEKR-GFNGISGYTFGKR KREDLD-EGGAFE-DENDDEKR-GFNGISGYTFGKR KREDLD-EGGAFE-DENDDEKR-GFNGISGYTFGKR KREDLD-EGGAFE-DENDDEKR-GFNGISGYTFGKR KREDLD-EGGAFE-DENDDEKR-GFNGISGYTFGKR KREGLD-EEGAFE-EENEDEKR-FNGISGYTFGKR KREGLD-EEGAFE-EENDDEKR-FNGISGLTFGKR KREGLD-EEGAFE-EENDDEKR-FNGISGLTFGKR KREGLD-EEGAFE-EENDDEKR-FNGISGLTFGKR KREGL-D-EEGAFE-EENDDEKR-FNGISGTFGKR KREGL-D-EEGAFE-EENDD-EKR-FNGISGTFGKR KREGL-D-EEGAFE-EENDD-EKR-FNGISGTFGKR KREGL-D-EEGAFE-DENDD-EKR-FNGISGTFGKR KREGL-D-EEGAFE-DENDD-EKR-FNGISGTFGKR KREGL-D-EEGAFE-DENDD-EKR-FNGISGTFGKR KREDMD-EEGAFE-DENDD-EKR-AYNGISGLTFGKR
Op_filo Mi_grac Am_squa Op_resi Op_abys Op_angu Op_scha Op_lame Op_bisp Op_brev Ba_hero Op_op_vivi Op_vivi Op_vivi Op_vivi Op_perf Cl_cana Op_liod Op_prol As_tubi	P-SGYSAFTFCKR-RMEPLSAFSFCKR-GMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKKRGMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKKRGMDPSGLSA P-SGFSAFSFCKR-RMDPLSAFSFCKR-GMDPSALSA P-SGFSAFSFCKR-R-EPLSAFSFCKR-GMDPSALSA P-SGFSAFSFCKR-R-DPLSAFSFCKR-GMDPSALSA P-SGFSAFSFCKR-R-DPLSAFSFCKR-GMDPSALSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDPSALSA P-AGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLGAFSFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA	FSFGKRGRDPSALSAFSFGKR- FSFGKRGMDPSGLSAFSFGKR- FSFGKRGMDPSALSAFSFGKR- FSFGKRGRDPSALSAFNFGKR- SFGKRGRDPSALSAFNFGKR- SFGKRGRDPSALSAFNFGKR- FSFGKRGRDPSGLSAFSFGKR- FSFGKRGRDSGLSAFSFGKRGF FSFGKRGRDAGLSAFSFGKRGF FSFGKRGRDAGLSAFSFGKRGF FSFGKRGRDAAGLSAFSFGKRGF FSFGKRGRDAAGLSAFSFGKRGF FSFGKRGRDAAGLSAFSFGKRGF FSFGKRGRDAAGLSAFSFGKRGF FSFGKRGRDAAGLSAFSFGKRGF FSFGKRGRDAAGLSAFSFGKRGF FSFGKRGRDAGLSAFSFGKRGF FSFGKRGRDAGLSAFSFGKRGF FSFGKRGRDAGLSAFSFGKRGF FSFGKRGRDAGLSAFSFGKRGF FSFGKRGRDAGLSAFSFGKRGF FSFGKRGRDAGLSAFSFGKRGF FSFGKRGRDAGLSAFSFGKRGF FSFGKRGRDAGLSAFSFGKRGF FSFGKRGRDSGMGAFSFGKRGF FSFGKRGRDSGMGAFSFGKRGF		KRDEEDGAFE-EENYDEKR SRIGALTGLTYGKR KREGGEEEDPAFE-EENNN-EEKR AGYNGLSQFTFGKR KREGLEEEDAALE-EEDNNDDEKR AGYNGLSQFTFGKR KREGLEEDGAFE-EENQDEEEKR GGYNGIAGYTFGKR KRDDLEEDGAFE-EENQDEEEKR GGYNGISGYTFGKR KRDDAEEDGAFE-EENNDEKR-GFNGISGYTFGKR KRDDAEEGAFE-DEDNDEKR-GFNGISGYTFGKR KREDLD-EEGAFE-DENDDEKR-GFNGISGYTFGKR KREDLD-EGGAFE-DENDDEKR-GFNGISGYTFGKR KREDLD-EGGAFE-DENDDEKR-GFNGISGYTFGKR KREDLD-EGGAFE-DENDNEKR-GFNGISGYTFGKR KREDLD-EGGAFE-DENDNEKR-GFNGISGYTFGKR KREDLD-EGGAFE-DENDNEKR-GFNGISGYTFGKR KREGL-D-EEGAFE-DENDNEKR-GFNGISGLTFGKR KREGL-D-EEGAFE-EENDDEKR-FNGISGLTFGKR KREGL-D-EEGAFE-EENDDEKR-FNGISGTFGKR KREGL-D-EEGAFE-DENDDEKR-FNGISGTFGKR KREGL-D-EEGAFE-DENDDEKR-FNGISGTFGKR KREGL-D-EEGAFE-DENDDEKR-FNGISGTFGKR KREDMD-EEGAFE-DENDDEKR-FNGISGTFGKR KREDMD-EEGAFE-DENDDEKR-FNGISGTFGKR KREDMD-EEGAFE-DENDDEKR-FNGISGTFGKR KREDMD-EEGAFE-DENDDEKR-FNGISGTFGKR
Op_filo Mi_grac Am_squa Op_resi Op_abys Op_angu Op_scha Op_bisp Op_brev Ba_hero Op_brev Ba_hero Op_vivi Op_vivi Op_vivi Op_vivi Op_perf Cl_cana Op_prol As_tubi Op_oedi	P-SGYSAFTFCKR-RMEPLSAFSFCKR-GMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SGFSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SGFSAFSFCKR-R-EPLSAFSFCKR-GMDPSALSA P-SGFSAFSFCKR-R-DPLSAFSFCKR-GMDPSALSA P-SGFSAFSFCKR-R-DPLSAFSFCKR-GMDPSALSA P-AGFSAFNFCKR-R-DPLSAFSFCKR-GMDPSGLSA P-AGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLGAFSFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLGAFSFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGLSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGLSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGLSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA	FSFG KRGRDPSALSAFSFG KR FSFG KRGMDPSGLSAFSFG KR FSFG KRGMDPSALSAFSFG KR FSFG KRGRDPSALSAFNFG KR FSFG KRGRDPSALSAFNFG KR FSFG KRGRDPSALSAFNFG KR FSFG KRGRDPSGLSAFSFG KR GF FSFG KRGRDPSGLSAFSFG KR GF FSFG KRGRDAGLSAFSFG KR GF FSFG KRGRDPSGMGAFSFG KR GF FSFG KRGRDPSGMGAFSFG KR GF FSFG KRGRDPSGMGAFSFG KR GF FSFG KRGRDPSGMGAFSFG KR GF FNFG KRGVDQSGLSAFSFG KR GF		KRDEEDGAFE-EENYDEKRSRIGALTGLTYGKR KREGGEEEDPAFE-EENNN-EEKRAGYNGLSQFTFGKR KREGLEEEDAALE-EEDNNDD-EKRAGYNGLSQFTFGKR KREGLEEDGAFE-EENQEEEKRGGYNGIAGYTFGKR KRDDLEEDGAFE-EENQEEEKRGGYNGISGYTFGKR KRDDAEEDGAFE-EENNDEKR-GFNGISGYTFGKR KRDDAEEGAFE-DENDDEKR-GFNGISGYTFGKR KREDL-D-EEGAFE-DENDDEKR-GFNGISGYTFGKR KREDL-D-EGGAFE-DENDDEKR-GFNGISGYTFGKR KREDLD-EGGAFE-DENDDEKR-GFNGISGYTFGKR KREDLEEDGAFE-DENDN-EKR-GFNGISGYTFGKR KREDLD-EGGAFE-DENDN-EKR-GFNGISGYTFGKR KREGL-D-EEGAFE-DENDN-EKR-GFNGISGYTFGKR KREGL-D-EEGAFE-DENDN-EKR-FNGISGYTFGKR KREGL-D-EEGAFE-EENDD-EKR-FNGISGLTFGKR KREGL-D-EEGAFE-EENDD-EKR-FNGISGLTFGKR KREGL-D-EEGAFE-DENDD-EKR-FNGISGYTFGKR KREGL-D-EEGAFE-DENDD-EKR-FNGISGYTFGKR KREDM-D-EEGAFE-DENDD-EKR-FNGISGYTFGKR KREDM-D-EEGAFE-DENDD-EKR-AYNGISGYTFGKR KREDM-D-EEGAFE-DENDD-EKR-AYNGISGYTFGKR KREDM-D-EEGAFE-DENDD-EKR-AYNGISGYTFGKR KREDM-D-EEGAFE-DENDD-EKR-AYNGISGYTFGKR KREDL-D-EEGAFE-DENND-EKR-AYNGISGYTFGKR
Op_filo Mi_grac Am_squa Op_resi Op_abys Op_angu Op_scha Op_lame Op_bisp Op_brev Ba_hero Op_operv Ba_hero Op_op_vivi Op_wend Op_plic Op_perf Cl_cana Op_exim_1 Op_liod Op_prol As_tubi Op_oedi Go_pust	P-SGYSAFTFCKR-RMEPLSAFSFCKR-GMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SGFSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SGFSAFSFCKR-R-EPLSAFSFCKR-GMDPSALSA P-SGFSAFSFCKR-R-DPLSAFSFCKR-GMDPSALSA P-SGFSAFSFCKR-R-DPLSAFSFCKR-GMDPSALSA P-AGFSAFNFCKR-R-DPLSAFSFCKR-GMDPSGLSA P-AGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGLSAFNFCKR-R-DPLSAFNFCKR-GKDASGLSA P-SGLSAFNFCKR-R-DPLSAFNFCKR-GKDASGLSA P-SGLSAFNFCKR-R-DPLSAFNFCKR-GKDASGLSA P-SGLSAFNFCKR-R-DPLSAFNFCKR-GKDASGLSA P-SGLSAFNFCKR-R-DPLSAFNFCKR-GKR-SGLSASLSA P-SGLSAFNFCKR-R-DPLSAFNFCKR-GKR-SGLSASLSA P-SGLSAFNFCKR-R-DPLSAFNFCKR-GKDASGLSA P-SGLSAFNFCKR-R-DPLSAFNFCKR-GKDASGLSA P-SGLSAFNFCKR-R-DPLSAFNFCKR-GKDASGLSA P-SGLSAFNFCKR-R-DPLSAFNFCKR-GKR-SGLSASLSA P-SGLSAFNFCKR-R-DPLSAFNFCKR-GKR-GKR-SGLSASLSA	FSFG KRGRDPSALSAFSFG KR FSFG KRGMDPSGLSAFSFG KR FSFG KRGMDPSALSAFSFG KR FSFG KRGMDPSALSAFNFG KR FSFG KRGRDPSALSAFNFG KR FSFG KRGRDPSALSAFNFG KR FSFG KRGRDPSGLSAFSFG KR GF FSFG KRGRDPSGLSAFSFG KR GF FSFG KRGRDAGLSAFSFG KR GF FNFG KRGYDQSGLSAFSFG KR GF FNFG KRGYDQSGLSAFSFG KR GF		KRDEEDGAFE-EENYD-EKRSRIGALTGLTYGKR KREGGEEEDPAFE-EENNN-EEKRAGYNGLSQFTFGKR KREGLEEEDAALE-EEDNNDD-EKRAGYNGLSQFTFGKR KREGLEEEDGAFE-EENQDEEEKRGGYNGIAGYTFGKR KRDDLEEDGAFE-EENQEEEKRGGYNGISGYTFGKR KRDDAEEDGAFE-EENND-EKR-GFNGISGYTFGKR KRDDAEEGAFE-DENDD-EKR-GFNGISGYTFGKR KREDL-D-EEGAFE-DENDD-EKR-GFNGISGYTFGKR KREDL-D-EGAFE-DENDD-EKR-GFNGISGYTFGKR KREDL-D-EGGAFE-DENDD-EKR-GFNGISGYTFGKR KREDL-D-EGGAFE-DENDN-EKR-GFNGISGYTFGKR KREDL-D-EGGAFE-DENDD-EKR-GFNGISGYTFGKR KREGL-D-EGGAFE-DENDN-EKR-GFNGISGYTFGKR KREGL-D-EEGAFE-EENED-EKR-FNGISGYTFGKR KREGL-D-EEGAFE-ENDD-EKR-FNGISGLTFGKR KREGL-D-EEGAFE-ENDD-EKR-FNGISGYTFGKR KREGL-D-EEGAFE-EENDD-EKR-FNGISGYTFGKR KREGL-D-EEGAFE-EENDD-EKR-FNGISGYTFGKR KREGL-D-EEGAFE-DENDN-EKR-AYNGISGYTFGKR KREDM-D-EEGAFE-DENDN-EKR-AYNGISGYTFGKR KREDM-D-EEGAFE-GENDD-EKR-FNGISGYTFGKR KREDM-D-EEGAFE-GENDD-EKR-AYNGISGYTFGKR KREDL-D-EEGAFE-GENDD-EKR-AYNGISGYTFGKR KREDL-D-EEGAFE-GENDD-EKR-AYNGISGYTFGKR KREDL-D-EEGAFE-GENDD-EKR-AYNGISGYTFGKR KREDL-D-EEGAFE-GENDD-EKR-AYNGISGYTFGKR KREDL-D-EEGAFE-GENDD-EKR-AYNGISGYTFGKR KREDL-D-EEGAFE-GENDD-EKR-AYNGISGYTFGKR KREDL-D-EEGAFE-GENDD-EKR-AYNGISGYTFGKR KREDL-D-EEGAFE-DENND-EKR-AYNGNSGYTFGKR KREDL-D-EEGAFE-DENND-EKR-AFNGMSGYTFGKR KREDL-D-EEGAFE-DENND-EKR-AFNGMSGYTFGKR
Op_filo Mi_grac Am_squa Op_resi Op_abys Op_angu Op_scha Op_lame Op_bisp Op_brev Ba_hero Op_op_vivi Op_vivi Op_vivi Op_vivi Op_vivi Op_perf Cl_cana Op_liod Op_prol As_tubi Op_oedi	P-SGYSAFTFCKR-RMEPLSAFSFCKR-GMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SGFSAFSFCKR-RMDPLSAFSFCKR-GMDPSALSA P-SGFSAFSFCKR-R-EPLSAFSFCKR-GMDPSALSA P-SGFSAFSFCKR-R-DPLSAFSFCKR-GMDPSALSA P-SGFSAFSFCKR-R-DPLSAFSFCKR-GMDPSALSA P-AGFSAFNFCKR-R-DPLSAFNFCKR-GMDPSGLSA P-AGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLGAFSFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLGAFSFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLGAFSFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGLSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGLSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGLSAFNFCKR-R-DPLSFFCKR-GMESSLSA P-SGLSAFNFCKR-R-DPLSFFCKR-GMESSLSA P-SGLSAFNFCKR-R-DPLSFFCKR-GMESSLSA P-SGLSAFNFCKR-R-DPLSFFCKR-GMESSLSA P-SGLSAFNFCKR-R-DPLSFFCKR-GMESSLSA P-SGLSAFNFCKR-R-DPLSFFCKR-GMESSLSA P-SGLSAFNFCKR-R-DPLSFFCKR-GMESSLSA P-SGLSAFNFCKR-R-DPLSFFCKR-GMESSLSA P-SGLSAFNFCKR-R-DPLSFFCKR-GMESSLSA P-SGLSAFNFCKR-R-DPLSFFCKR-GMESSLSA P-SGLSAFNFCKR-R-DPLSFFCKR-GMESSLSA P-SGLSAFNFCKR-R-DPLSFFCKR-GMESSLSA P-SGLSAFNFCKR-R-DPLSFFCKR-GMESSLSAF	FSFG KRGRDPSALSAFSFG KR FSFG KRGMDPSGLSAFSFG KR FSFG KRGMDPSALSAFSFG KR FSFG KRGMDPSALSAFNFG KR FSFG KRGRDPSALSAFNFG KR FSFG KRGRDPSALSAFNFG KR FSFG KRGRDPSGLSAFSFG KR SF FSFG KRGRDPSGLSAFSFG KR SF FSFG KRGRDASGLSAFSFG KR SF FSFG KRGRDPSGMSAFSFG KR SF FNFG KRGYDQSGLSAFSFG KR SF FNFG KR GYDQSGLSAFSFG KR SF FNFG KR GYDQ F FNFG KR GYD F F FNFG KR GYD F F F F F F F F F F F F F F F F F F F		KRDEEDGAFE-EENYD-EKRSRIGALTGLTYGKR KREGGEEEDPAFE-EENNN-EEKRAGYNGLSQFTFGKR KREGLEEEDAALE-EEDNNDDEKRAGYNGLSQFTFGKR KREGLEEDGAFE-EENQDEEEKRGGYNGISGYTFGKR KRDDEEDGAFE-EENQDEEEKRGGYNGISGYTFGKR KRDDEEDGAFE-EENQEEEKRGGYNGISGYTFGKR KRDDEEGAFE-DENDD-EKR-GFNGISGYTFGKR KRDDEEGAFE-DENDD-EKR-GFNGISGYTFGKR KREDL-D-EEGAFE-DENDD-EKR-GFNGISGYTFGKR KREDL-D-EGGAFE-DENDD-EKR-GFNGISGYTFGKR KREDL-D-EGGAFE-DENDD-EKR-GYQISGYTFGKR KREDL-D-EEGAFE-ENDD-EKR-FNGISGYTFGKR KREGL-D-EEGAFE-ENDD-EKR-FNGISGYTFGKR KREGL-D-EEGAFE-ENDD-EKR-FNGISGYTFGKR KREGL-D-EEGAFE-ENDD-EKR-FNGISGYTFGKR KREGL-D-EEGAFE-ENDD-EKR-FNGISGTFGKR KREGL-D-EEGAFE-ENDD-EKR-FNGISGTFGKR KREGL-D-EEGAFE-ENDD-EKR-FNGISGTFGKR KREGL-D-EEGAFE-EENDD-EKR-FNGISGTFGKR KREGL-D-EEGAFE-DENDD-EKR-AYNGISGTFGKR KREGL-D-EEGAFE-DENDD-EKR-AYNGISGTFGKR KREDM-D-EEGAFE-GENDD-EKR-AYNGISGTFGKR KREDD-D-EEGAFE-GENDD-EKR-AFNGSGTFGKR KREDL-D-EEGAFE-DENND-EKR-AFNGSGTFGKR KREDL-D-EEGAFE-DENND-EKR-AFNGSGTFGKR KREDL-D-EEGAFE-DENND-EKR-AFNGSGTFGKR
Op_filo Mi_grac Am_squa Op_resi Op_abys Op_angu Op_scha Op_lame Op_bisp Op_brev Ba_hero Op_operv Ba_hero Op_op_vivi Op_wend Op_plic Op_perf Cl_cana Op_exim_1 Op_liod Op_prol As_tubi Op_oedi Go_pust	P-SGYSAFTFCKR-RMEPLSAFSFCKR-GMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SGMSAFSFCKR-RMEPLSAFSFCKRGMDPSGLSA P-SGFSAFSFCKR-RMDPLSAFSFCKR-GMDPSALSA P-SGFSAFTYCKR-R-EPLSAFSFCKR-GMDPSALSA P-SGFSAFTYCKR-R-DPLSAFSFCKR-GMDPSALSA P-SGFSAFTYCKR-R-DPLSAFSFCKR-GMDPSALSA P-SAFDAFSFCKR-R-DPLSAFSFCKR-GMDPSALSA P-AGFSAFNFCKR-R-DPLSAFNFCKR-GMDPSGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLGAFSFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLGAFSFCKR-GMDASGLSA P-SGFSAFNFCKR-R-DPLGAFSFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGFSAFNFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGLGAFSFCKR-R-DPLSAFNFCKRGGMDASGLSA P-SGLSAFNFCKR-R-DPLSFFCKRGGMDASGLSA P-SGLSAFNFCKR-R-DPLSFFFCKRGGMDASGLSA P-SGLSAFNFCKR-R-DPLSFFFCKRGGMDASGLSA P-SGLSAFNFCKR-R-DPLSFFFCKRGGMDASGLSA P-SGLSAFNFCKR-R-DPLSFFFCKR-GME-SGLSA P-SGLSAFNFCKR-R-DPLSFFFCKR-GME-SGLSA P-SGLSAFNFCKR-R-DPLSFFFCKR-GME-SGLSA P-SGLSAFNFCKR-R-DPLSFFFCKR-GME-SGLSA P-SGLSAFNFCKR-R-DPLSFFFCKR-GME-SGLSA R-SGFNAFSFCKR-R-DPLSAFSFGKR-GMD-RLNA	FSFG KRGRDPSALSAFSFG KR FSFG KRGMDPSGLSAFSFG KR FSFG KRGMDPSALSAFSFG KR FSFG KRGRDPSALSAFNFG KR FSFG KRGRDPSALSAFNFG KR FSFG KRGRDPSALSAFNFG KR FSFG KRGRDPSGLSAFSFG KR GF FSFG KRGRDSGLSAFSFG KR GF FSFG KRGRDASGLSAFSFG KR GF FNFG KRGYDQSGLSAFSFG KR GF FNFG KR GF FNF		KRDEEDGAFE-EENYD-EKRSRIGALTGLTYGKR KREGGEEEDPAFE-EENNN-EEKRAGYNGLSQFTFGKR KREGLEEEDAALE-EEDNNDD-EKRAGYNGLSQFTFGKR KREGLEEEDGAFE-EENQDEEEKRGGYNGIAGYTFGKR KRDDLEEDGAFE-EENQEEEKRGGYNGISGYTFGKR KRDDAEEDGAFE-EENND-EKR-GFNGISGYTFGKR KRDDAEEGAFE-DENDD-EKR-GFNGISGYTFGKR KREDL-D-EEGAFE-DENDD-EKR-GFNGISGYTFGKR KREDL-D-EGAFE-DENDD-EKR-GFNGISGYTFGKR KREDL-D-EGGAFE-DENDD-EKR-GFNGISGYTFGKR KREDL-D-EGGAFE-DENDN-EKR-GFNGISGYTFGKR KREDL-D-EGGAFE-DENDD-EKR-GFNGISGYTFGKR KREGL-D-EGGAFE-DENDN-EKR-GFNGISGYTFGKR KREGL-D-EEGAFE-EENED-EKR-FNGISGYTFGKR KREGL-D-EEGAFE-ENDD-EKR-FNGISGLTFGKR KREGL-D-EEGAFE-ENDD-EKR-FNGISGYTFGKR KREGL-D-EEGAFE-EENDD-EKR-FNGISGYTFGKR KREGL-D-EEGAFE-EENDD-EKR-FNGISGYTFGKR KREGL-D-EEGAFE-DENDN-EKR-AYNGISGYTFGKR KREDM-D-EEGAFE-DENDN-EKR-AYNGISGYTFGKR KREDM-D-EEGAFE-GENDD-EKR-FNGISGYTFGKR KREDM-D-EEGAFE-GENDD-EKR-AYNGISGYTFGKR KREDL-D-EEGAFE-GENDD-EKR-AYNGISGYTFGKR KREDL-D-EEGAFE-GENDD-EKR-AYNGISGYTFGKR KREDL-D-EEGAFE-GENDD-EKR-AYNGISGYTFGKR KREDL-D-EEGAFE-GENDD-EKR-AYNGISGYTFGKR KREDL-D-EEGAFE-GENDD-EKR-AYNGISGYTFGKR KREDL-D-EEGAFE-GENDD-EKR-AYNGISGYTFGKR KREDL-D-EEGAFE-GENDD-EKR-AYNGISGYTFGKR KREDL-D-EEGAFE-DENND-EKR-AYNGNSGYTFGKR KREDL-D-EEGAFE-DENND-EKR-AFNGMSGYTFGKR KREDL-D-EEGAFE-DENND-EKR-AFNGMSGYTFGKR