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Neuroglobins: pivotal proteins associated with emerging neural systems and precursors of metazoan globin diversity

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Key words: Neuroglobin, structure, nervous system evolution, globin evolution, acoel, cnidarian

Background: Neuroglobins are expressed in vertebrate neurons.

Results: Neuroglobins are located in acoel and medusa neural systems (two basal animals) but also ubiquitous in metazoan transcriptomes.

Conclusion: Neuroglobin has been recruited early in neural cell prototypes and later co-opted in hemoglobin-based blood systems.

Significance: Universality of neuroglobin sheds new light on the origin and evolution of globins.

SUMMARY

Neuroglobins, previously thought to be restricted to vertebrate neurons, were detected in the brain of a photosymbiotic acoel, *Symsagittifera roscoffensis*, and in neuro-sensory cells of the jellyfish *Clytia hemisphaerica*. For *S. roscoffensis*, a member of a lineage that originated either at the base of the Bilateria or of the deuterostome clade, we report the ligand-binding properties, crystal structure at 2.3 Å and specific brain immuno-cytochemical pattern. Furthermore, we describe *in situ* hybridizations of two neuroglobins specifically expressed in differentiating nematocytes (neuro-sensory cells) and in statocytes (ciliated mechanosensory cells) of the nervous system of *C. hemisphaerica*, a member of the early-branching animal phylum Cnidaria. *In silico* searches using these neuroglobins as queries revealed the presence of previously unidentified neuroglobin-like sequences in most metazoan lineages. Since neural systems are ubiquitous in Metazoa (except Porifera and Placozoa), the constitutive expression of neuroglobin-like proteins in an acoel and a cnidarian, two metazoans with simple body plans, strongly supports the notion of an intimate association of neuroglobins with the evolution of animal neural systems and hints at the preservation of a vitally important function. Neuroglobins were probably recruited in the first proto-neurons in early metazoans, from globin precursors we identified in choanoflagellates, sponges or placozoans, and were strongly

conserved concomitantly with nervous system evolution. Since the origin of neuroglobins predates the origins of other metazoan globins, it is likely that neuroglobin gene duplication followed by co-option and subfunctionalization led to the emergence of polyphyletic families of globins in protostomes and deuterostomes (i.e. convergent evolution).

Interest in the structure, function and evolutionary relationships of circulating hemoglobins (Hbs) and intracellular myoglobins (Mbs) of animals dates back to the first 3D structural determination of these proteins in the 1960s (1) (2) (3). The large range of animal globins and the extensive occurrence of globins in prokaryotes (4) is now recognized. Prominent among the recently described metazoan globins is vertebrate neuroglobin (Ngb) (5) which is expressed in neurons of the central and peripheral nervous systems. The *in vivo* function of Ngb remains undefined despite a major effort over the last decade. Suggested functions include oxygen (O₂) supply in hypoxia and ischemia (6), scavenging of reactive oxygen free radicals (7), protection from apoptosis (8), redox-regulated nitrite reductase activity (9) and involvement in respiratory chain function (10). In murine models of human neuropathology, Ngb is also expressed in reactive astrocytes, a subtype of glia cells in the nervous system (11).

Recently, Blank and collaborators (12) demonstrated that the functional hexacoordinated Globin X (GbX) protein of the

cypriniform adult Zebrafish is located in nervous central system and retina, suggesting a neural-based function but contradicting a previous result obtained from the other cypriniform *Carassius auratus* GbX showing that mRNA GbX was not detected in brain and eye but in other tissues (muscle, heart, gut, liver) (13). Thought to be restricted to vertebrate, GbX-like sequences have been recently *in silico* identified in other deuterostomes and in protostomes supporting an early emergence of this gene family in metazoan evolution (14). However further cellular investigations must be performed for assigning a non ambiguous neural function of the GbX-like sequences so far identified in metazoans.

In protostomes, globins have been observed in the nerve tissue of certain annelids, molluscs and a nematode (15), but have not been phylogenetically linked to vertebrate Ngbs or other deuterostome globins. Their O₂ binding affinities resemble those of vertebrate Mbs and their function is considered to be O₂ storage and thus protection against hypoxia (16), (17).

Recent phylogenomic analyses of vertebrate globins have demonstrated that they can be separated into two groups, one derived from vertebrate-specific duplications (Cytoglobins, Globin E, Globin Y, the Hb chains and Mb), and another resulting from duplications preceding the emergence of chordates (Ngb, HbX) (18), (19), (20). The most recent molecular phylogenetic analysis of globin sequences from the five major groups comprising the deuterostomes, i.e.

cephalochordates, echinoderms, hemichordates, urochordates and vertebrates, suggests that all deuterostome globins occur in four clades (21). Despite the fact that a molecular analysis of metazoan globins (including echinoderm and cnidarian globins) suggested an ancestral connection to the nervous system (22), Ngbs have not been reported in deep branching metazoan lineages, and evolutionary patterns of emergence of metazoan globin lineages are still unresolved.

We have employed the discovery of the hitherto unknown Ngbs in an acoel and a cnidarian, that exhibit simple morphological organizations characteristic of ancestral Bilateria / bilaterian Deuterostomes and Radiata, to further clarify the origin of globins in metazoan lineages.

Symsagittifera roscoffensis is a photosymbiotic acoel (Fig. 1A), thus occupying a phylogenetic position either preceding the deuterostome-protostome split or branching at the base of deuterostomes (23) (24). This hermaphroditic marine flatworm has a simple body plan with a digestive syncytium (no epithelially-lined gut), a ventral mouth, a muscle system, a nervous system with a simple central brain, but no excretory or blood circulatory systems (25).

We report the discovery of Ngb-like sequences in EST libraries from *S. roscoffensis*, the cloning and purification of a Ngb, its immunocytochemical localization within neural cells, its ligand binding properties and crystal structure. We examined the sites of expression of putative Ngbs in the jellyfish *Clytia hemispherica* (Cnidaria, Anthozoa), which, like

the “higher” animals (the Bilateria), exhibits a complex body organization, including striated musculature, reproductive organs and a specialized nervous system (26). *In situ* hybridization experiments using two specific Ngb-like probes highlight differentiating neural cell type called nematocytes or stinging cells (mechanoreceptors, i.e. neuro-sensory cells) and statocysts (gravito-sensors) in the jellyfish *Clytia hemispherica*.

A broad *in silico* transcriptome survey revealed expressed Ngb-like proteins in most of the metazoan phyla, ranging from animals with no symmetrical body plan (sponges, placozoans) to complex bilaterians, through the symmetrically radial cnidarians. Based on Ngb conservation throughout metazoans and recent biomedical studies underlying the irreversible detrimental effects of Ngb dysfunction in neurons, we assume that Ngb played a crucial role early in the subsequent evolution of metazoan nervous systems and brains in metazoan exhibiting more complex body-plan. Indeed, Ngb appears as a key and central partner in neurone physiology as a neuroprotectant preventing the oxidative damages and neurodegenerescence as illustrated in Alzheimer’s disease transgenic mice models (27).

Our data and results suggest that an ancestral globin-like gene was recruited in emerging proto-neural cells and system in the first diploblastic animals (ancestors of extant cnidarians such as sea anemones, corals or medusa) and specifically evolved as a neural globin. A natural corollary is a novel scenario for metazoan globin evolution, namely, the

independent emergence of globins such as extracellular annelid Hbs, mollusc and arthropods Hbs, and vertebrate Hbs, via functional shifts from Ngb copies early during metazoan radiation and concomitantly with increasing body plan complexity and the development of blood circulatory systems.

EXPERIMENTAL PROCEDURES

Expression, purification and characterization of S. roscoffensis Ngb - The coding sequence of *S. roscoffensis* Ngb i.e. (SrNgb1), (ID number European Nucleotide Archive HE972520) was amplified by PCR and subsequently cloned into a pET-3a cloning vector (Invitrogen). The construct was transformed into *E. coli* BL21DE3 for protein expression in auto-inducible medium (28). The protein was purified with an Akta purifier system (GE Healthcare), due to the low pI of the *S. roscoffensis* globin the samples were loaded on a 5 ml HiTrap DEAE FF column (GE Healthcare) equilibrated with Tris HCl 50mM pH 8.5, and eluted at a concentration of 25mM NaCl. The obtained samples were loaded on a desalting Sephadex G-25 column (GE Healthcare) suspended in PBS, pH 7.4 and the material was finally purified on a Superose 12 HR 16/50 (Amersham Biosciences) column equilibrated with PBS, pH 7.4. Finally, ferric and ferrous spectra UV/visible spectra (O₂ and CO) were measured with a Cary model 400 spectrophotometer.

Autoxidation kinetics and ligand rebinding of SrNgb1 - Full spectra were measured versus time on HP 8453 diode-array spectrophotometer. The sample was first thoroughly deoxygenated in a sealed optical cuvette under a stream of N₂. Then a slight excess of sodium dithionite was added to reduce the globin heme moiety. Finally the cuvette was equilibrated under air to obtain the oxy reduced species and to allow the depletion of the residual unreacted dithionite. Ligand recombination kinetics were measured at a single wavelength after photodissociation by 10 ns pulses at 532 nm, as previously described (29). Samples in sealed cuvettes were equilibrated under various fractions of CO or oxygen. A mixed atmosphere of both CO and O₂ was used to study the oxygen to CO replacement reaction after photolysis of CO.

Immuno-cyto-localization with SrNgb1 and RF-amide antibodies - Acoel flatworms were collected in Roscoff (Brittany, France) and anesthetized with 7% MgCl₂ and fixed during 45 min in 4% PFA at 4°C. Animals were then washed with phosphate buffer pH 7.4, permeabilized with 0.1% Triton X-100 in PBS 3 times for 15 min at room temperature. They were then incubated with 5% BSA, 0.1% Triton and 0.05% Tween 20 in PBS for 2-3 hours at room temperature and then incubated overnight at 4°C alternatively with 1/700 polyclonal *S. roscoffensis* anti-Ngb, produced against whole recombinant protein by Eurogentec (Speedy 28-day polyclonal packages) or with anti-RF-amide (courtesy of Thomas Leitz,

Kaiserslautern). The next day, acoels were washed three times for 15 min in PBS and incubated with the appropriate secondary antibodies. They were then incubated for 10 min in a DAPI solution (2µg/ml in PBS), washed 3 times in PBS and mounted on a glass slide for microscope observation. Image acquisition of fluorescence labeling was monitored with a confocal microscope (Leica sp5) equipped with a 20x objective and using Leica LAS-AF software.

Animal collection and in situ hybridization - Medusae were obtained in Paris by culture of *C. hemisphaerica* colonies in artificial seawater (Reef Crystals®) established from polyps provide by Evelyn Houliston (Villefranche-sur-Mer) as previously described (30). Medusae were left unfed for one day before fixation. They were fixed for 40 min at 4°C in 3.7% PFA, 0.2% glutaraldehyde, PBT 1X (10mM Na₂HPO₄, 150 mMNaCl, pH7.5, 0.1% Tween 20). DIG-labelled antisense RNA probes synthesis and whole-mount *in situ* hybridizations were carried out as previously described (31) The only modification to the *in situ* protocol was an acetic anhydride treatment before hybridization. Alkaline phosphatase activity was revealed using NBT/BCIP (NitroBlueTetrazolium/Bromochlorylindolophosphate, blue staining) or fast red TR-naphthol reagent ® (Sigma, red staining). After postfixation and DAPI staining (32), samples were mounted in Citifluor®. Double *in situ* hybridizations were performed as described in (33). DIC images were obtained with an

Olympus BX61 microscope using Q-imaging Camera with Image Pro plus® software (Media Cybernetics).

Protein crystallization - All crystallization experiments were carried out at 292 K. Initial crystallization trials were performed with the PACT, JCSG+, PEG I and PEG II Suites (Qiagen) that is a total of 384 conditions in four 96-well plates from Corning. The trials were set up using a Cartesian crystallization robot, and the sitting drops were made by mixing 300 nl of protein (13 mg/mL in 30 mM PBS buffer pH 7.5, 100 mM NaCl) with 150 nl of reservoir solution. A single hit was identified in the PEG II screen, containing 1M LiCl, 0.1 M Na acetate and 30% (w/v) PEG 6000. Subsequently, this crystallization condition was optimized in 24-well Linbro plates by the hanging-drop vapour-diffusion method, screening ranges from 0.6 to 1.0 M LiCl and 30% to 39% PEG 6000. These drops were prepared on siliconized cover slips by mixing 2 ml of protein with 1 ml of well solution. The drops were equilibrated against reservoir solutions of 0.75 ml volume. Best crystals were obtained for 32% PEG 6000, 1.0 M LiCl and 0.1 M Na acetate. For cryo-protection, 5% glycerol was added to the crystal drop solution before flash-freezing the crystals in the gaseous N₂-stream at 100 K.

Data collection and X-ray diffraction analysis - X-ray diffraction data were first collected from globin crystals at 100 K on beamline ID23-I at the ESRF (Grenoble, France) using an ADSC

Quantum 4R CCD detector. All crystals were flash-cooled in a liquid nitrogen stream. The crystals were rotated through 120° with a 0.5° oscillation range per frame at a wavelength of 0.933 Å. All raw data were processed using the program XDS and the resultant data were merged and scaled using the program XSCALE (34). Models for structure solution by molecular replacement were selected by a sequence search using BLAST against the PDB sequence database. However, all attempts, to solve the structure of this globin by molecular replacement performed with the program AMORE (35), using various neuroglobin or myoglobin models stayed unsuccessful. A second data set was therefore collected at the Fe absorption edge at a wavelength of 1.7387 Å on beamline BM30A, covering an angular section of 90° with an oscillation range of 1.0°. The data treatment was performed with XDS in the same way as for the native data set. All further data collection statistics are given in Table S1.

Crystal structure determination and refinement

- The iron atom substructure solution was calculated with SHELXD (36) followed by phasing and density modification performed with SHELXE, using the graphical interface HKL2MAP (37) and the resulting electron density map was displayed with Coot (38). Both possible enantiomorph space groups were tried and the phasing procedure allowed a selection of a clear and contrasted structure solution in P6₂22. These starting phases were used to build the initial model using ARP/wARP and REFMAC as part of the CCP4

suite (39), and switching to the higher resolution data at 2.3 Å. Roughly 70% of the helices were constructed by the automatic procedure. The subsequent manual adjustment and model building was carried out with Coot and alternated with refinement cycles using REFMAC. Water molecules were added automatically with the REFMAC-ARP/wARP option and visually verified, one by one, using Coot. The final model contained residues ranging from 6 to 154, the prosthetic heme group, 98 water molecules and an oxygen ligand bound to the iron atom. The asymmetric unit contains one globin molecule leading to a Matthews coefficient of 4.9 and a solvent content of 74.9%. The phasing and final refinement statistics are given in Table S1. (*Symsagittifera roscoffensis* neuroglobin PDB accession number: 4B4Y)

Phylogenomics and molecular phylogeny - The identification of Ngb-like / putative neural globin sequences was performed using *S. roscoffensis* Ngb1 and vertebrate Ngb sequences queries in blastp searches of the non-redundant nucleotide database maintained by NCBI (<http://www.ncbi.nlm.nih.gov/nucest/>) and of non-annotated ESTs databases from various metazoans, deposited and archived at (<ftp://ftp.ncbi.nlm.nih.gov/pub/TraceDB/>).

A multiple alignment of a representative subset of Ngb-like sequences has been automatically generated with HMMER v3.0 package (40) using the hmmalign program and the Globin (PF0042) raw HMM as a guide. Molecular

phylogenetic analysis was carried out using the Maximum likelihood approach with PhyML software (41) with LG option as model of amino acid substitution, NNI moves option for the tree topology search operation and SH-like support option for the default branch support. The tree topology (Newick format) was edited with MEGA5.1 (42).

In addition to be accessible in Mat&Met / Results paragraphs and supplementary data, the molecular phylogeny analysis has been performed and deposited using respectively the BioSide software and his dedicated website at <http://www.bioside.org>. Indeed, in order to be easily traceable and reproducible by anyone that would like to replay the molecular phylogeny procedure, a file including the original multiple alignment of sequences and PhyML setups are available and detailed following the permalink http://www.bioside.org/workflow/BS12111400_01 (id number is BS121114001) or at <http://www.bioside.org/community>. Prediction of N-terminal myristoylation of Ngb-like sequences was performed with the program The MYR Predictor, a web-service available at <http://mendel.imp.ac.at/myristate/SUPLpredictor.htm>. This program calculates whether a protein is predicted as myristoylated with reliable/twilight zone confidence or not.

RESULTS and DISCUSSION

The Ngb-like protein 1 of S. roscoffensis is a functional neuroglobin - SrNgb1 is expressed in the brain and nervous system of *S. roscoffensis* (Fig. 1B; 1C). The acael brain is formed by a

layer of neuronal cell bodies surrounding a central neuropile, embedding the statocyst, a gravity sensor (25). The SrNgb1 signal mainly occurs in the anterior tip (“head”) where photoreceptors and frontal sensory organs collect environmental information. The signal surrounds the statocyst and the photoreceptors and is superimposable with the anti-RFamide antibody pattern (Fig. 1B) and the serotonergic nervous system (43). Constitutive expression of SrNgb1 during embryogenesis and in juvenile and adult stages indicates its implication throughout nervous system development and in maintenance of brain activity.

The spectroscopic properties of purified SrNgb1 (UV and visible absorption spectra of the ferrous and ferric forms) indicate that in the absence of external ligands it is pentacoordinated, in contrast to vertebrate Ngbs in which a sixth coordination bond is formed with a distal histidine (Fig. 2A). The rate constants of O₂ and CO binding and of O₂ dissociation are similar to those of vertebrate Mbs, and consequently so is its O₂ binding affinity (Table 1). The rate of heme autoxidation under pure O₂ at 25 °C is slow (first order rate 0.053 h⁻¹; Fig. 2B), which is not surprising in view of the fact that there is a well established inverse relationship between O₂ affinity and autoxidation rate for pentacoordinated globins. This reaction is much slower than those observed for vertebrate Ngbs, probably due to a higher capacity of the hexacoordinated form for transferring an electron to molecular O₂ (44). Overall, these observations are consistent with an *in vivo* function involving reversible binding of the

diatomic ligand rather than a redox reaction with O₂ as a terminal electron acceptor.

The structure of S. roscoffensis neuroglobin -

The structural model consists of 149 residues (including Ala6 to Glu154) that bind a heme b prosthetic group, with a bond between the heme iron and the proximal histidine (H103), the distal ligand being an O₂ molecule (Fig. 3A). The tertiary structure corresponds to the classical globin fold, consisting of eight helices (A to H, Fig. 3A), the heme binding cleft formed by helices E and F. Despite being deoxy-pentacoordinated, SrNgb1 shares certain structural features with vertebrate Ngbs that are quite different from classical Hb and Mb structures. Although the identity of SrNgb1 with mouse Ngb is only 19% (Fig. 3B), all of the conserved globin-fold residues (45) are present, including the heme ligand residues E7His and F8His. The C and D helix regions most closely resemble those described in murine Ngb (46). The Trp residue at position 52 in SrNgb1 (Fig. 3A) may present a barrier to ligand exit and entry by forming a stable hydrogen bond to one of the heme propionates (distance 2.8Å; Fig. 3A). This interaction is reinforced by a water molecule located near by (heme-propionate-O2D/HOH, distance 3.8Å; HOH117), which is further hydrogen bonded to the distal histidine (ND1, 2.7Å) and the second propionate group of the heme (HOH/heme-O2D, 3.0Å). In murine Ngb, residues Lys67 and Tyr44 form a similar hydrogen bonding network involving a water molecule also binding to the distal His(46). Structural equivalence is provided by superimposition of

HOH117 with its murine counterpart, and by superimposition of the Tyr44 OH-group in murine Ngb with the Trp52 NH-group in SrNgb1. Moreover, Tyr44 in murine Ngb and Trp52 in SrNgb1 are at equivalent positions in the sequence alignment (Fig. 3B). SrNgb1 also shares with murine Ngb the high flexibility of the connection between helices E and F (data not shown). SrNgb1 displays a unique feature in that helix F is bent by the presence of a proline (Pro94) (Fig. 3C). This could be analogous to the transition of human Ngb structures that is triggered by a disulfide bond in the CD region (47). The closest match to SrNgb1 in the PDB database was ferrous CO-bound murine Ngb (1W92). Overall, the SrNgb1 structural sequence matches Ngbs and plant Hbs, with a slightly better Z-score (48) than to Mbs (data not shown).

In the cnidarian medusa Clytia hemisphaerica, two globins (CheNgb1 and CheNgb2) are expressed in differentiating neuro-sensory cells (nematocytes) - Nematocytes exhibit many characteristics of neuro-sensory cells, including mechano-sensitive cilia, neurite-like outgrowths and synapses. They contain a single-use dart specialized for killing prey. Nematogenesis (the generation of nematocytes) in Cnidaria is used as a model for non-bilaterian neurogenesis (26), (49), as these neural cells are continuously generated throughout larval and adult life.

The *CheNgb1* and *CheNgb2* genes are mainly expressed in the nematogenic ectoderm of tentacle bulbs and manubrium (compare Fig.

4A, 4B and 4C). In the tentacle bulbs, their expression patterns are crescent-shaped and interrupted on the external side of the bulb (Fig. 4D-4F, blue staining), thus exactly matching the expression of minicollagen 3-4a (*mcol3-4a*, Fig. 4H, red staining). The latter belongs to a family of small collagen-like proteins known in hydrozoans to be a major component of the nematocyst wall (33). Double *in situ* hybridizations revealed extensive co-expression of *mcol3-4a* with both *CheNgb1* (purple color in Fig. 4E) and *CheNgb2* (purple color in Fig. 4G), indicating that both genes are expressed in differentiating nematoblasts over a large time window.

CheNgb2 mRNA was also detected in the statocysts (Fig. 4F arrowhead, 4F' and 4F''), the equilibration organs arranged regularly around the rim of the bell of the animal. *CheNgb2*-expressing cells are located in the basal epithelium of the statocyst, near the bell margin and interpreted as ciliated mechano-sensory cells (figure 4F' and 4F'').

CheNgb1 and *CheNgb2* transcripts were also abundant in the proximal part of the manubrium ectoderm and mimicked the expression pattern of minicollagen, with which they are co-expressed as demonstrated by double *in situ* hybridization. *CheNgb1* and *CheNgb2* were also localized in the female gonad in an unidentified cell type (not germ line cells) (fig 4A and 4B).

Neuroglobins are ubiquitously expressed in Metazoa - Using SrNgb1 as an in silico probe

for blasting genomic resources, we identified 50 or so transcripts never described so far from different phyla (Table S2A-B) mostly related to other neuroglobins / neuroglobin-like sequences according to classical blastp searches against the NCBI non-redundant nucleotide database. After a cross verification systematically conducted with the Panther predictive tool, all the new globins fall into the Panther Leghemoglobin-related family that encompass 14 subfamilies including Neuroglobin, Globin X, Non-Symbiotic Hemoglobin and Leghemoglobin. None of the new sequences we found are related to the Panther Hemoglobin-family that encompasses vertebrate Hemoglobin, Cytoglobin or Myoglobin.

The taxonomic distribution of the neuroglobin-related sequences suggests broad conservation throughout metazoan evolution (Fig. 5A, Table S2A-B). They were detected in non-symmetrical body plan basal metazoans with neither nervous system nor circulatory system, i.e. in the metazoan lineages Porifera (the sponges *Amphimedon queenslandica* and *Carterospongia foliascens*) and Placozoa (*Trichoplax adherens*). In the radially symmetrical cnidarians which have a simple nervous system but no circulatory blood system, Ngb-like sequences were present in Anthozoa (the coral *Montastraea faveolata* and the sea anemones *Anemona viridis* and *Nematostella vectensis*) and Hydrozoa (*Clytia hemisphaerica* and *Hydra magnipapillata*). No other types of globin (neither homologs of circulating Hbs nor Mb-like globins) were detected in these basal metazoans. In

protostomes, expressed Ngb-like sequences were found in (1) cephalopod mollusks such as the cuttlefish *Sepia officinalis* and *Euprimna scolopes* and the squid *Dorytuthis paeleii*, (2) many arthropods such as the hymenopter *Apis mellifera* (bee), the crustacean *Carcinus maenas* (green shore crab) and *Daphnia pulex* (a common species of water flea) or the social insects *Harpegnathos saltator* (ant), (3) the sipunculid *Themiste sp.* (the peanut worm), (4) the brachiopod *Terebratalia transversa* (the common lampshell), (5) various annelids such as the polychaetes *Alvinella pompejana* (Pompeii worm from deep-sea hydrothermal vents) or the hirudinea *Hellobdella robusta* (leech). Expressed Ngb-like sequences were also identified in so called “minor phyla” such as platyhelminthes, tardigrads, kinorhynchs, and nemertodermatids (a sister group of acoels) (Table S2A-B). In deuterostomes, Ngb-like sequences were identified in all phyla preceding the emergence of vertebrates: in the echinoderms *Strongilocentrotus purpuratus* and *Paracentrotus lividus* (sea urchins), the hemichordates *Saccoglossus kowalevskii* (acorn worm) and *Balanoglossus clavigerus*; the cephalochordate *Branchiostoma lanceolatum* (amphioxus, also known as the lancelets), and the urochordates *Molgula tectiformis* and *Botryllus schlosseri* (tunicates).

Vertebrate species have a single Ngb gene copy while many of the other metazoans have several copies, indicating gene duplication events correlated with subfunctionalization. The existence of a second *S. roscoffensis* or cnidarian Ngb sequence (Table S2A-B)

illustrates classical cases of diversification by a gene duplication event. The molecular unrooted phylogenetic tree (Fig. 5B) clearly shows that Vertebrate Hbs, Mbs, and Cybs clearly form a distinct monophyletic group (Fig. 5B), in agreement with earlier results (50) (21). Vertebrate Ngb and GbXs are included into a group of functional neuroglobins and neuroglobin-related sequences that harbors the neuroglobins characterized in this study i.e. the Ngb duplicates of *S. roscoffensis* and of *C. haemispherica*. The presence of Vertebrate GbXs sequences in this group supports a likely connection of these proteins with neural systems. The cluster that contains the choanoflagellates leghemoglobin-related sequences (the closest living unicellular relative to metazoan (38)), the poriferan and the vertebrate Ngbs sequences likely represents the ancestral Ngb lineage with plesiomorphic characteristics. Indeed we noticed in Blast results that choanoflagellates and poriferans, cnidarians and *S. roscoffensis* neuroglobins produced significant alignments with protists, especially with the unicellular green algae *Micromonas* and the diatom (unicellular brown algae) *Thalassiosira* globin that both exhibit Leghemoglobin-related signature according Panther prediction system. These findings are in agreement that metazoan globins were likely inherited from a unicellular eukaryote globin. The second cluster with SrNgb2, CheNgb1 and Vertebrates GbX represents another cluster of neuroglobin-related sequences. The other sequences diagnosed as putative neuroglobin-related proteins (with a leghemoglobin-related signature) that do not cluster specifically within

the Ngb group (including reflect primary sequence divergence and likely species-specific functional diversification. Further exploratory approaches such as gene or protein expression localization will be required for formally establishing the involvement of these proteins (including the so-called GbX) in the nervous system.

We also noticed, when the coding sequences we recovered were complete, that some neuroglobin-related sequences exhibited a meristoylation site and some not, with no clear pattern in the phylogenetic tree (Fig. 5B). Vertebrates Hbs, Mbs, and Cybs clearly form a distinct monophyletic group (Fig. 5B).

It is clear that our molecular phylogeny of Ngb-like sequences is inevitably based on a heteroclitic subset of paralogous and orthologous Ngb-like sequences, but as transcriptomes do not reveal 100% of transcripts and especially cryptically expressed genes (those with a low number of corresponding transcripts), the number of Ngb-like proteins is likely to be significantly underestimated. In other words, more sequences with more functional data from more taxa will refine the phylogenetic relationship among Ngb-related sequences.

Neuroglobin is likely an early constitutive actor in nervous systems and brain evolution - It is clear that Ngb-like proteins are ubiquitous in metazoans (Fig. 5A). The emergence of neural structures in metazoans represented an innovation resulting in functions such as

interneuronal and neuro-muscular transmission, allowing feeding, reproduction, vision and complex behaviours like predation (51). Although the origin of nerve cells remains unknown, the Cnidaria, whose name derived from cnidocytes (i.e. nematocytes), occupy a key position in Metazoan with respect to early nervous system evolution (52). Together with the ctenophores, the Cnidaria form the Coelenterata, the sister group of eumetazoans (Bilateria) (53). It is assumed that transduction of chemical and mechanical stimuli in nematocytes are hallmarks of primitive nerve cells and that nematocytes are thus representative of ancestral sensory cells that preceded the differentiation of neuronal cell types in animal evolution (54). The unequivocal expression of Ngbs in nematocytes of the jellyfish *Clytia hemispherica* appears to be a robust indication of the essential role of these proteins in early evolution of the nervous system. The fact that acoel and jellyfish statocysts (the sensory organs measuring pressure) are respectively and specifically targeted by Ngb antibody and Ngb probes illustrates the intimate connection of Ngb with nerve nets and transmission of information. We assume that an original exaptation, i.e. the recruitment of a globin by proto-nervous cells and proto-nervous circuitry, laid the foundations for elaborated nervous systems and brains in the first metazoans displaying anatomical polarity (radial then bilateral symmetry) and differentiated nervous systems. Neuroglobin precursors are likely homologous to those identified in unicellular eukaryotes (choanoflagellates) and simple metazoans

(sponges and placozoans) devoid of neural cells, but possessing the basic genetic toolkit encoding proteins homologous to those involved in nervous system development in higher animals (55,56) (Fig. 5A).

The deleterious effects on nerve cells of Ngb silencing (57) (10) and the conservation of this protein throughout metazoan evolution underline the pivotal function of Ngbs in development and physiology of neurons. Sub-cellular expression of Ngb in mitochondria of neuronal cells in regions of the brain with high metabolic activity (58) (10) is an indicator of the implication of Ngb in cellular homeostasis in extant organisms and, by extension, in early emerging metazoan neuronal cells. The Ngb-like sequences of certain cnidarians, protostomes and deuterostomes exhibit a predicted N-terminal myristoylation site indicating a possible interaction with membranes, putatively including those of the mitochondria (Fig.5B). The presence of such a site has already been described for the globin expressed in the gills of the crab *Carcinus maenas* mentioned above, a Ngb-like protein (Leghemoglobin-related family) according to Panther prediction (59).

In the core of the globin-fold, hexacoordination of the heme iron atom leads to a high autoxidation rate, suggesting that hexacoordinated vertebrate Ngbs are involved in redox metabolism connected to oxidative phosphorylation either with electron carriers or with reactive oxygen species produced by the mitochondria (60). Our results show that some Ngbs, such as SrNgb1, can be functionally

pentacoordinated. SrNgb1, whose O₂ binding affinity is similar to that of Mb, is likely to be involved in O₂ storage, and thus provision of O₂ during periods of hypoxia. This proposal is in agreement with the most likely roles of nerve Hbs in the annelid (*Aphrodite aculeata*), the clams (*Spisula solidissima* and *Tellina alternata*) and the nemertean (*Cerebratulus lacteus*), which have been established to be the provision of O₂ to the metabolically highly active neural cells and thus protection under hypoxic conditions (15), (61), (62), (16).

It remains to be determined which form of coordination (penta- or hexa-) of metazoan Ngbs was associated with neofunctionalization and which was the ancestral state. It is pertinent to note that human Ngb exists as an equilibrium between the two forms, with the hexacoordinated form being dominant (~99:1) (9).

Neuroglobins could also be precursors of the metazoan globin repertoire - The results of our survey highlight the presence of putative Ngbs proteins in radial and bilateral animals irrespective of the presence or absence of a blood circulatory system and of the respiratory protein employed (hemocyanin in mollusks and arthropods, hemerythrin in sipunculids and brachiopods, hemoglobin in other metazoans). The presence of Ngb in ice fish, where circulating Hb has disappeared from the blood circulatory system, is not paradoxical as claimed by Cheng (63), but illustrates the separate evolutionary pathways of Ngbs and O₂

binding Hbs, the mandatory constitutive expression of Ngb in the nervous system, and a clear case of disadaptation, i.e. loss of the circulating oxygen carrier.

Assuming that the ancestral bilaterian body plan was very simple with a nervous system but no blood circulatory system, it is obvious that the presence of Ngb predates the emergence of circulatory Hb. Given that Ngbs are ancestral and constitutively expressed in all metazoans (Fig. 5A), the sporadic presence of O₂ binding Hb in individual metazoan lineages strongly suggests that they are polyphyletic. In other words, the emergence of circulating Hb in metazoans is likely due to convergent evolution. The globin lineages other than Ngb found in many metazoan groups have probably emerged as the result of functionalization (64) and cooption of a Ngb-like globin in early metazoans. Indeed, most of the metazoan transcriptomes checked in this study exhibit multiple Ngb-like paralogs, likely originating from gene duplication events.

CONCLUSION

We demonstrate the presence of a functional Ngb in neural cells of the acoel *S. roscoffensis* and expression of homologous Ngbs specific to neuro-sensory cells (differentiating nematoblasts) in the cnidarian jellyfish *Clytia hemispherica*. These results suggest that the first globins expressed in early bilaterians and symmetrically radial cnidarians were specifically linked to the metazoan nervous system. The pentacoordination of SrNgb1 vis a

vis the hexacoordination of the vertebrate Ngbs may be due to differences in function, with the acoel Ngb playing an O₂ storage role providing neuroprotection during hypoxic periods. This interpretation is supported by reports of the functions of “nerve globin” in several protostomes.

Extensive *in silico* mining of genomic data using SrNgb1 as a probe revealed the occurrence of expressed Ngb-like sequences in most metazoan phyla, including sponges and Placozoa, basal metazoans lacking neural and circulatory systems. Our results clearly demonstrate that the emergence of Ngb in metazoans chronologically preceded the emergence of other globin families. Consequently, we propose a novel scenario for metazoan globin evolution, based on two broad and complementary statements. On the one hand, our experimental and *in silico* results suggest that an ancestral globin-like gene was recruited in the emerging proto-neural system in the ancestor of Bilateria and diploblastic animals (ancestors of extant cnidarians such as sea anemones, corals or medusa) to become a functional Ngb. On the other hand, metazoan globins other than Ngbs, such as annelid, mollusc, arthropod, and vertebrate Hbs, likely originated independently from early Ngbs, via co-option of duplicated Ngb genes and functionalization during metazoan radiation, concomitant with increasing body plan complexity and the emergence of blood circulatory systems.

Access to multiple ontogenetic stages of emerging marine models, for which genomic

resources and molecular tools are increasingly available (65), will be of a prime importance for functional genomic exploration using Ngbs as key developmental markers in animal lineages exhibiting complex nervous tissues (cephalopods), elaborated social behavior (ants), or subject to anthropogenically-induced stresses or diseases (corals, mussels, oysters).

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FOOTNOTES

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FIGURE LEGENDS

FIGURE 1. A: photograph of a colony of the symbiotic acoel *Symsagittifera roscoffensis* (4 to 5 mm long) at low tide. The green color is due to the presence of about 50000 photosymbionts (the

unicellular green algae *Tetraselmis convolutae*) harbored within each adult acoel. **B:** (1) Light micrograph of the anterior tip of a juvenile with the statocyst (S) flanked by two photoreceptors (P). Cilia are visible on the periphery of the head; (2) Composite confocal image showing the red SrNgb1-antibody signal surrounding the statocyst with peripheral extensions, DAPI stained nuclei appear in blue; (3) Confocal image illustrating RF-amide stained *Symsagittifera roscoffensis* nervous system. **C:** Magnification of the extremity of the anterior tip (“the head”). Arrows indicate fiber-like structures labelled with SrNgb1 anti-body occurring at the same place of the frontal glands (frontal sensors). These fiber-like structures are superimposable with serotonergic nervous system and especially neurites.

FIGURE 2. A: UV and visible spectra for SrNgb1. In the inset is shown the partially oxygenated spectrum measured under an oxygen tension of 10 Torr. The dashed lines refer to the maximum absorption for the fully oxy and deoxy spectra. **B:** Autoxidation of SrNgb1 in 50 mM Tris-HCl 0.2 mM EDTA 10 U SOD and catalase at pH 8.0 under 1 atm O₂ at 25 °C. In the inset is shown the variation of absorption occurring during the redox kinetics. (right panel) Autoxidation of SrNgb1 in 50 mM Tris-HCl 0.2 mM EDTA 10 U SOD and catalase at pH 8.0 under 1 atm O₂ at 25 °C. The inset shows the variation of absorption occurring during the redox kinetic.

FIGURE 3. A: Ribbon representation of SrNgb1 crystal structure (4B4Y): (left panel) close up view of the heme binding pocket of SrNgb1 highlighting the hydrogen bonding network involving the distal heme binding position and a tightly bound water molecule (HOH117); (right panel) 3D structural super imposition with murine Ngb (1W92). **B:** Multiple sequence alignment based on the structural superimposition with murine Ngb (1W92), bovine hemoglobin (1JEB) and sperm whale myoglobin (107M), as obtained with the program ESPRIPT (<http://espript.ibcp.fr/ESPrIPT/cgi-bin/ESPrIPT.cgi>). The conserved histidines (axial heme ligands) are marked by black triangles. The red triangle marks a Trp residue involved in the tight binding of a water molecule in the distal heme pocket. The eight helices that form the classical globin fold are numbered from A to H and color coded from blue (N-terminus) to red (C-terminus) in the same manner as in the ribbon representation of SrNgb1 in Figure 1b. **C:** Extract of the SrNgb1 crystal structure highlighting the relative orientations of the heme-ligand containing helices E and F. A proline at position 94 in helix F leads to a discontinuous and bent helix F in SrNgb1 (4B4Y). The same structural extract showing the relative orientations of helices E and F in murine Ngb (1W92), where Helix F is continuous and straight.

FIGURE 4. Expression patterns of two *Clytia globin* genes *CheNg1* and *CheCyto* in several territories of the medusa. **A-C:** Whole-mount *in situ* hybridizations for *CheNg1*, *CheCyto* (Blue= NBT/BCIP development) and *Chemcol3-4a* (red=Fast red development). **D-H:** All bulbs have the same orientation, proximal area on the top and tentacle on the bottom. Crescent shaped expression patterns in the ectodermal layer of a tentacle bulb. All of them are interrupted on the external side of the bulb but sometimes the continuity of the staining on the inner face is visible (**E**, **G**). **E** and **G:** purple staining indicates expression of two genes: in each case, minicollagen 3-4a staining was revealed first with fast red and then the other probe was revealed using NBT/BCIP. **E'** and **G':** details of the staining in nematoblasts (black arrowhead). **F'** and **F'':** Higher magnification of a statocyst delimited by the dotted circle: *In situ* hybridization (in black and white) and DAPI counterstaining (in red) merged after conversion of the *in situ* staining in grey scale. **I-L:** Gene expression in the manubrium views (mouth on the bottom). The signal is concentrated in the ectoderm layer at the base of the manubrium. **J'** and **L':** detailed views of double-stained cells corresponding to nematoblasts (black arrowhead), note that there is no signal in mature nematocytes (white arrowhead). go: gonad, ma: manubrium, tb: tentacle bulb. Scale bars: A-C: 100 μm ; D-H: 25 μm ; I-M: 50 μm ; E', G', J', L': 5 μm ; F', F'': 10 μm .

FIGURE 5. A: Schematic and consensual representation of metazoan phylogeny illustrating the presence /absence of Ngbs, other globins, the two other respiratory proteins (hemocyanin and hemerythrin) and blood circulatory systems. The sporadic presence of globins in certain metazoan lineages can be explained by independent functional shifts from Ngb-like proteins (i.e convergent evolution). Acoelomorphs and Xenoturbella are represented in two alternative phylogenetic positions, reflecting the ongoing debate as to their affiliations. **B:** Unrooted molecular phylogeny based on multiple alignments of a subset of 84 sequences that comprise 138 amino acids of Ngbs, LGB-related (Ngb-like), Hb, Mb and Cyb sequences from diverse phyla. Dots indicate a possible miristoylated Ngb-like (green) or not (red). Vertebrates globins are in the yellow clusters: Hemoglobin, myoglobin and cytoglobins appear clearly as an invention of Vertebrates while vertebrate Ngbs and GbXs are imbedded within the large green group where functional neuroglobin of *Syngasteria roscoffensis* and *Clytia haemispherica* occur (respective names are in bold red). The blue cluster that includes Ngb-like sequences from Choanoflagellates, Porifera (sponges), Placozoa, some Cnidaria, some protostome and deuterostomes including vertebrate Ngbs (yellow cluster) likely represents the plesiomorphic members of Ngbs.

TABLE 1. O₂ and CO binding data. Experimental conditions: 50 mM Tris-HCl 100 mM NaCl, 5 mM DTT, pH 8.0. Human Ngb experimental conditions: 100 mM potassium phosphate, 2 mM DTT, pH 7.0. O₂ solubility coefficient was 1.82×10^{-6} moles/liter at 25 °C and for CO solubility coefficient was 1.36×10^{-6} moles/liter at 25 °C. ° O₂ affinity was estimated equal to 1.8 Torr.

FIGURE 1

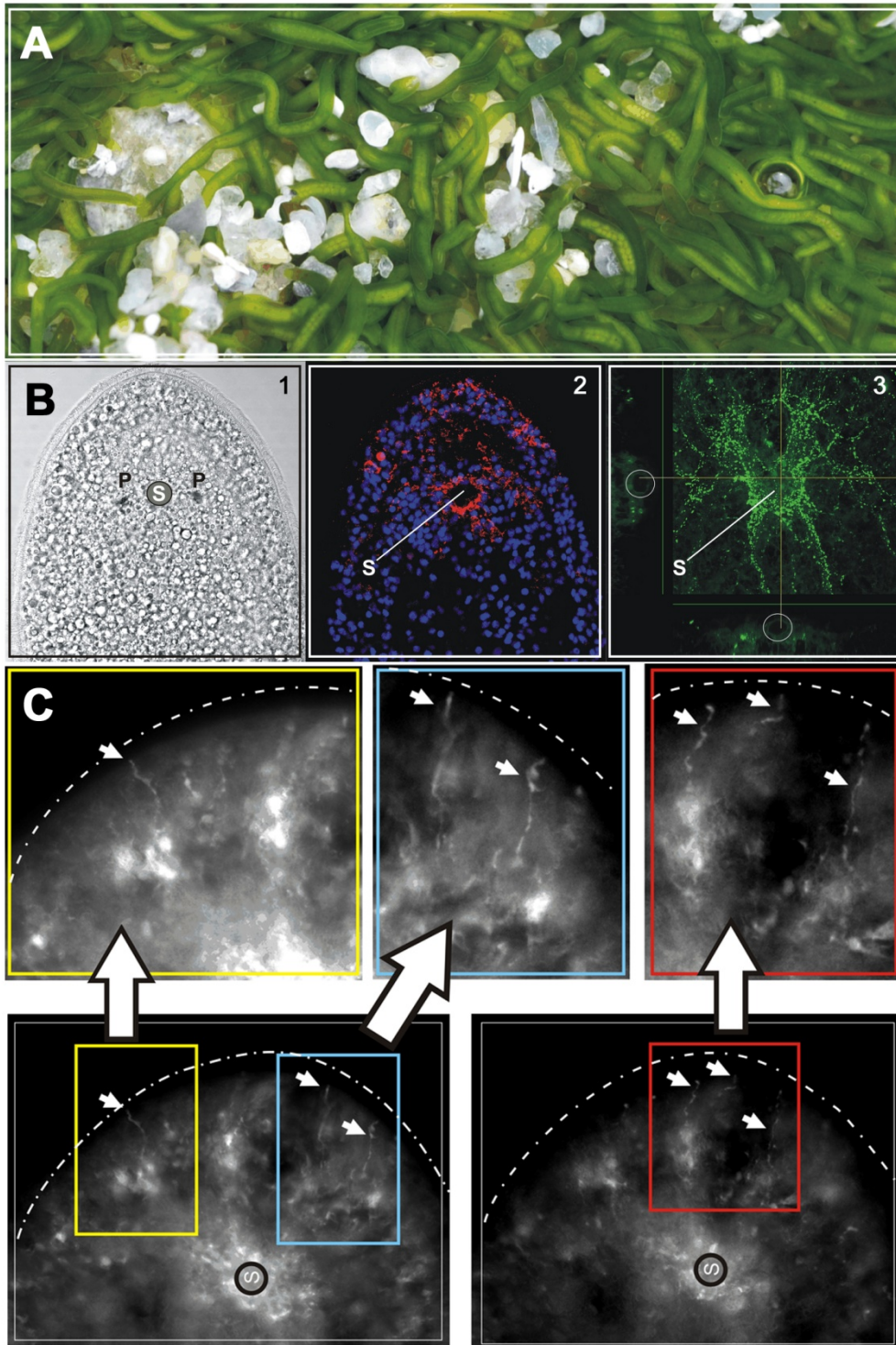


FIGURE 2

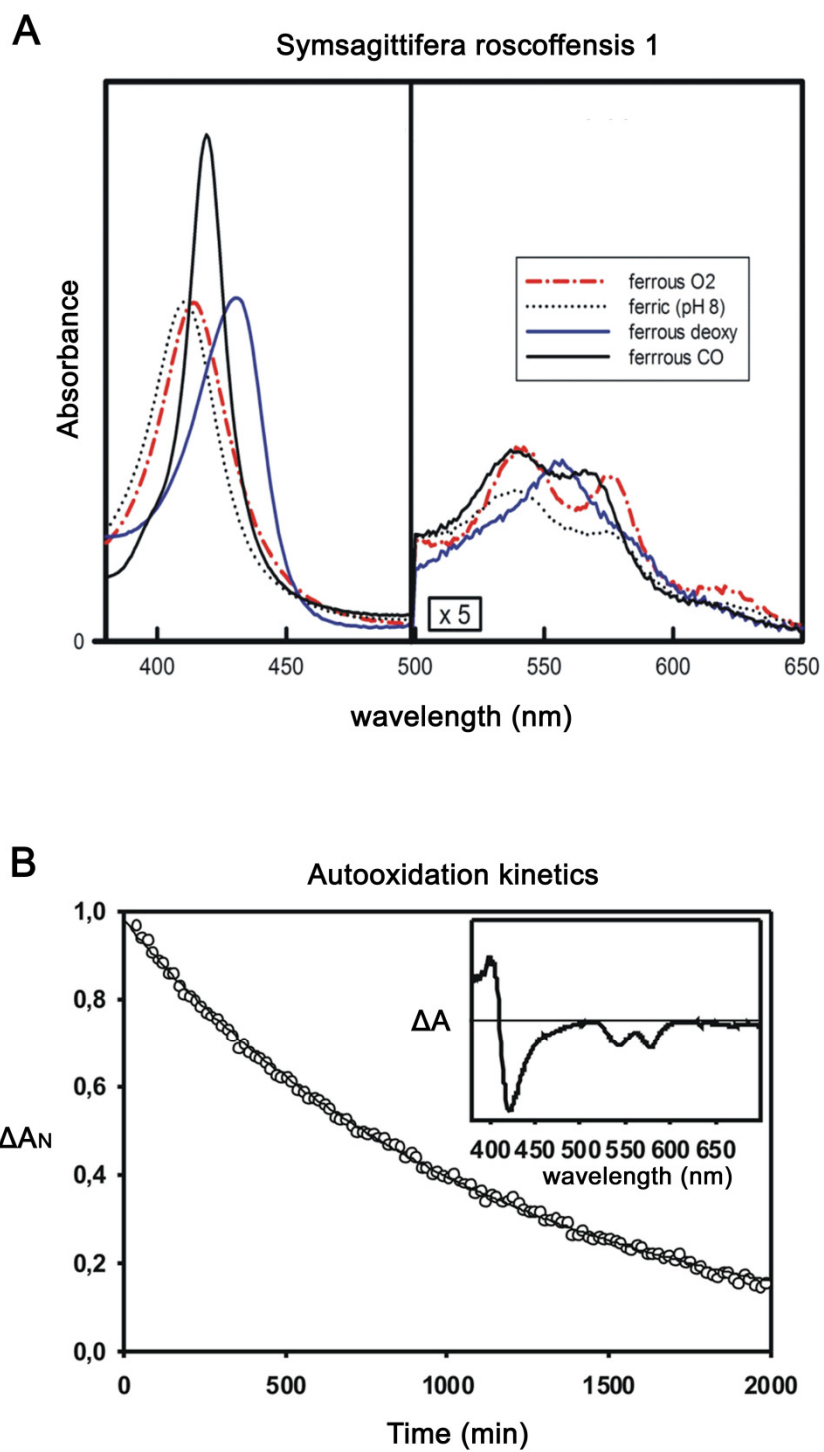


FIGURE 4

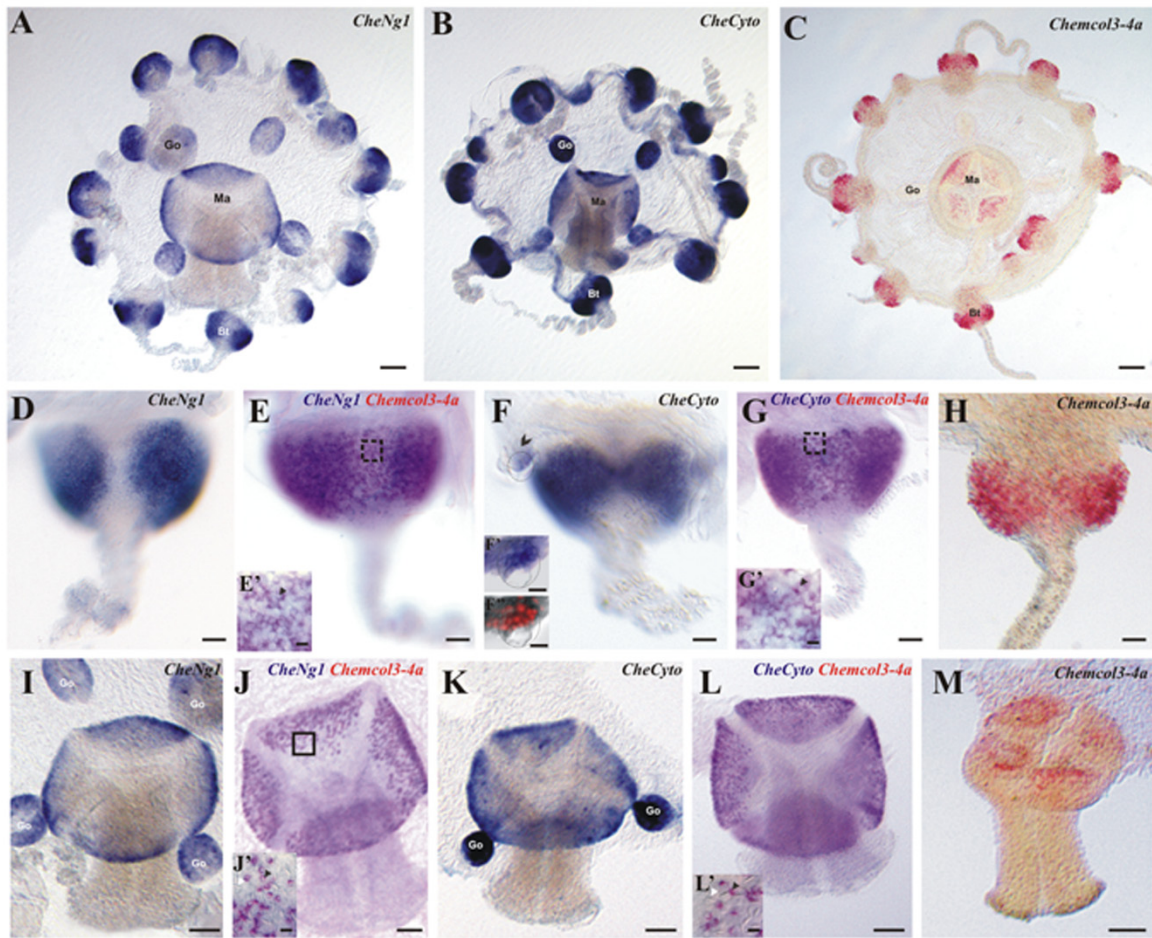
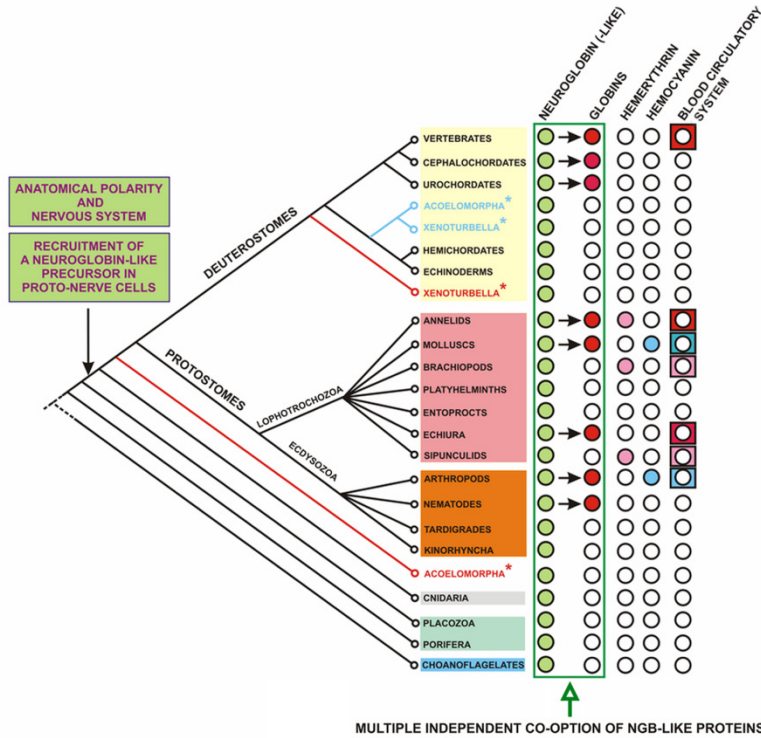


FIGURE 5

A



B

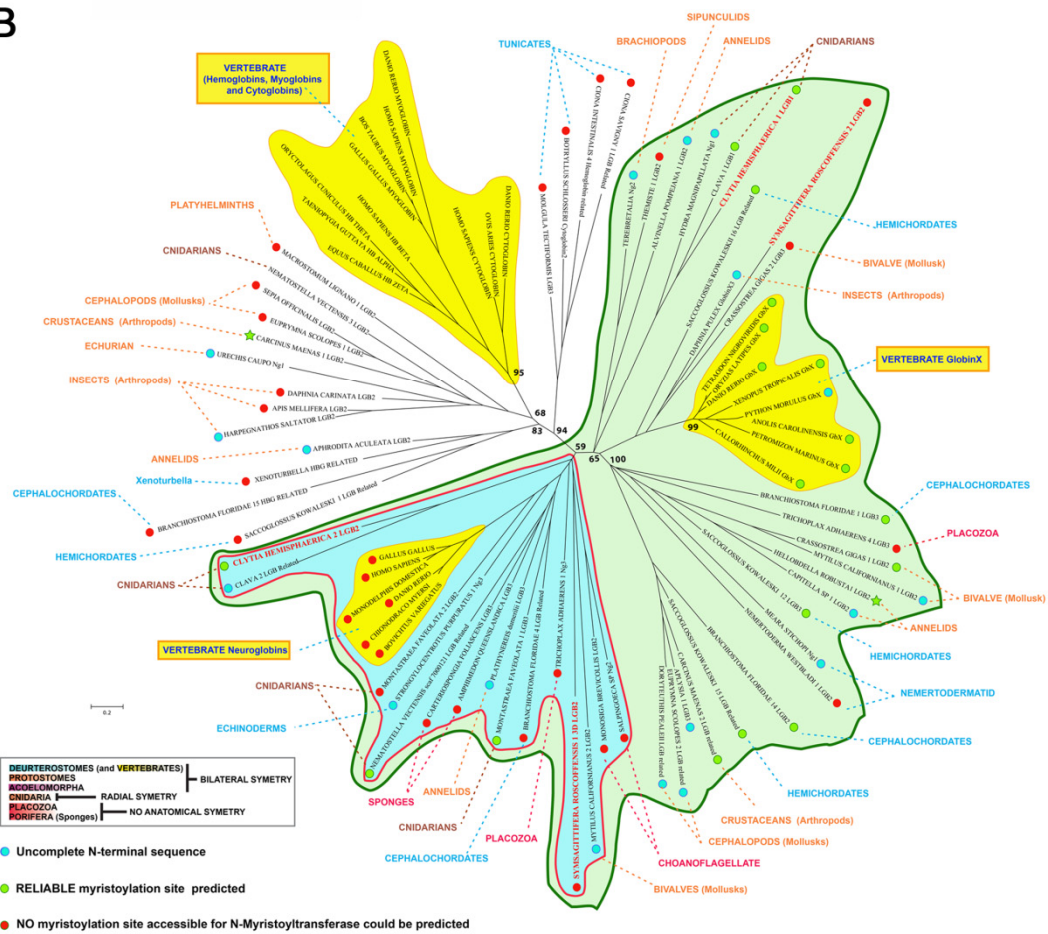


TABLE 1

	SrNgb1	Human Mb	human Ngb (25°C)
k_{onCO} (/μM/s)	0.35	0.65	40
k_{onO_2} (/μM/s)	7	15	170
k_{offO_2} (/s)	35+/-5	27	0.7
K_{O_2} (μM)	5.0	1.8	0.004
$P_{50 \text{ O}_2}$ (Torr)	2.8	1	6.8
k_{onHis} (/s)			1800
k_{offHis} (/s)			0.6
K_{His}			3000

TABLE S1 Data collection, phasing and refinement statistics on globin crystals, space group P6₂22

Beamline at ESRF	ID23-I	BM30A
Wavelength (Å)	0.93	1.7389
Unit cell parameters in Å and degrees (°)	a=b=97.07, c=140.10, $\alpha=\beta=90, \gamma=120$	a=b=97.07, c=140.10, $\alpha=\beta=90, \gamma=120$
Resolution range (Å)	30.06–2.3 (2.36-2.30) ^a	50.06–3.2 (3.29-3.20) ^a
No. of observations	57993 (3605)	60276 (3599)
No. of unique reflections	12137 (688)	11480 (675)
Completeness (%)	91.7 (77.4)	93.3 (77.4)
$\langle I/\sigma(I) \rangle$	16.6 (2.2)	14.9 (2.0)
Redundancy	4.8 (5.2)	5.2 (5.3)
R_{sym} (%) ^b	4.8 (43.3)	7.6 (56.7)
Phasing statistics		
Anomalous difference (CC in %, given by ShelxE)		35.21
Figure of merit		0.335
Refinement statistics		
R_{cryst} (%)	21.4	
R_{free} (%) ^c	25.1	
Esu based on Free R value	0.17	
Overall B factor (Å ²)	38.1	
Protein	38.0	
Heme	33.4	
Solvent	41.2	
Rms deviation in bond lengths (Å)	0.029	
Rms deviation in bond angles (°)	2.46	

^a Values for the highest resolution shell are given in parenthesis.

^b $R_{\text{sym}} = \Sigma |I - I_{\text{av}}| / \Sigma |I|$, where the summation is over all symmetry-equivalent reflections.

^c R_{free} values were calculated on 5% of the data (904 reflections) that were set aside in the minimization steps.

TABLE S2A

Below are the primary sequences (in fasta format) used for molecular phylogeny.

In order to see the original alignment used to perform the molecular phylogeny this file has to be opened with any sequence editor.

New LegHemoglobin related (including Ngb-like) sequences found in various phyla are highlighted in yellow (with supplemental information concerning their origin).

For each sequences except for vertebrate globins, Panther prediction are mentioned in the single line description such as Ng for neuroglobin-like, LGB for LegHemoglobin-related, Cytoglobin for cytoglobin-like, HGB for Hemoglobin related, and GlobinX for globinX-like.

Note that for each of these sequences the score of Panther prediction hit is specified: 1 (the score of the Panther hit is better than E-3, but worse than E-11 (protein is evolutionarily related but function may have diverged), 2 (the score of the Panther hit is better than E-11, but worse than E-23 (molecular function likely to be the correct but biological process/pathway less certain), 3 (the score of the Panther hit is better than E-23 (very likely to be a correct functional assignment)).

>VERTEBRATE GlobinX

>ANOLIS CAROLINENSIS GbX (from Droge et al 2011)
 -----MGCALSGAQDPPVSEECSPDDG
 LDLNRETTLGSNGRTTEPFPLSGAQKELIRGSWEILHK---DIARVGIIVFIRLFETHPE
 -CKDVFFLFR-D--ID--DFQQLKMSKELQAHGLRVMSFIEKSVARM---DQ-EPKLHHL
 AFELGRSHCRYK----APPKYEYIGIQFIQAAQPILKEAWT-PETEKAWEGLFQYLAAT
 MRRGFYKEQKATGKN-----

>CALLORHINCHUS MILII GbX (from Droge et al 2011)
 -----MGCAIS
 GPGQYPASGREDDVAVASLSLSDRQTQLVKETWRLVQE---DIAKVGIIIMFVRLFETHPE
 -CKDAFFLFR-D--ID--DLQQLRKSGLRAHGLRVMSFIEKTVARL---DQ-EDRLQQL
 XLELGKSHFRYS----AAPKYYPYVGNFICAVQPILKEKWT-AEVEEAWKGLFHYLTSV
 MKKGYQDEERGSCPREKPKHGPNV-----

>DANIO RERIO GbX (from Droge et al 2011)
 -----MGCAISG
 SGLTAGAPEIRPGEETPAGLTTNHIRLIKESWRLIQE---DIAKVGIIIMFVRLFETHPE
 -CKDVFFLFR-D--VE--DLERLRTSRELRAHGLRVMSFIEKSVARL---DQ-LERLETL
 ALELGKSHYRYN----APPKYGYVGAEFICAVRPILKDRWT-PELEEAWKTLFQYVTSI
 MREGFLEEERNKRSNTQTSSRERPDKRSTAI-----

>ORYZIAS LATIPES GbX (from Droge et al 2011)
 -----MGCAIS
 GLAAKTDLAERSREDAAVEHPNEEQIQMIKDSWKVIRD---DIAKVGIIIMFVRLFETHPE
 -CKDVFFLFR-D--VE--DLERLRTNRELRAHGLRVMSFIEKSVARL---DQ-PERLEAL
 AVELGKSHYHYN----APPKYNYVGAEFICAVQPILKEQWT-TELEKAWQTLFQFVTAL
 MKQGYQEE-SARQRQLA-TSPKDRDLKRNAL-----

>PETROMIZON MARINUS GbX (from Droge et al 2011)
 -----MGCTVSTDERTGAQSSSQGQSASRKQQ
 QPEQQRAAGEGHQPPGPPQAPSESQRRLVRDLSWLALQC---DIARVGVIMFVRLFETHPE
 -CKDVFYQFR-D--CE--DLQKLKMNKQLQAHGLRVMSFIEKSVARL---EQ-ECVLEQL
 IVEMGRKHYKYN----ASPKYYSFVGIEFIATVQPFLQEKWT-NEVEDAWQCLFRYIAAV
 MKRGYLEEEAASNGVNTANYDRGQGNHGATAM-----

>PYTHON MORULUS GbX (from Droge et al 2011)

 -----QKELIRESWKILHK---NITRVGIIVFIRLFETHPE
 -CKDVFFLFR-D--ID--DLQQLKMNKELQAHGLRVMSFIEKSVARL---DQ-EGKLEVL
 AFELGRSHFRYK----APPNYEYIGIQFIQAVQPILKEDWT-LEVEKAWK-----

```

>TETRAODON NIGROVIRIDIS GbX (from Droge et al 2011)
-----MGCALSSLGAKA
EFGDRSAEEEDAAAAA AVVYPREDQIQMIKDSWKVIRD---DIAKVGIIIMFVRLFETHPE
-CKDVFFLFR-D--VE--DLERLRSSRELRAHGLRVMSFIEKSVARL---DQ-QDRLEAL
AVELGKSHYHYN----APPKYYSYVGAEFICAVQPILKERFT--SELEEAWKTLFQYVTGL
MRKGHQEEGSRQRHLALPPKDGPEKRTSAL-----
-----
>XENOPUS TROPICALIS GbX (from Droge et al 2011)
-----MGCILSSLG
WQWRDSL DHTETSPLLPTLNLSEQQOQLLVESWRLIQH---DIAKVGIVILFVRLFETHPE
-CKDVFFLFR-D--VD--DLQALRANKDLRAHGLRVLSFVEKSVARI---AD-CARLEEL
ALELGRSXRYRN----APPRYYQYVGTEFISAVCPMLHDKWT--AEVEEA WKGLFAYICTV
MERGYQEEE--RRHSDGRSLIDGLQGNKGLI-----
-----
>VERTEBRATE MYOGLOBINS
-----
>HOMO_SAPIENS_MYOGLOBIN_gi|4885477|ref|NP_005359.1|
-----
-----MGLSDGEWQLVNLVWGKVEA---DIPGHGQEV LIRLFKGHPE
-TLEKFDKFK-H--LK--SEDEMKASEDLKKHGATVLTALGGILKK-----KGHHEAE
IKPLAQSHATKHK---IPVKYLEFISECIIQVLQSKHPGDFG-ADAQGAMNKALELFRKD
MASNYKELGFQG-----
-----
>DANIO_RERIO_MYOGLOBIN_gi|41053652|ref|NP_956880.1|
-----
-----MADHDLVLKCGAVEA---DYAANGGEVLNRLFK EYPD
-TLKLFPKFS-G--IS--QGDLAGSPA VAAHGATV LKKGEL LKA-----KGDHAAL
LKPLANTHANIHK---VALNNFRLITEVLVKVMAEKAGLDAA-GQ GALRRVMDA VIGDID
GYKKEIGFAG-----
-----
>BOS_TAURUS_MYOGLOBIN_gi|27806939|ref|NP_776306.1|_
-----
-----MGLSDGEWQLVNLAWGKVEA---DVAGHGQEV LIRLFTGHPE
-TLEKFDKFK-H--LK--TEAEMKASEDLKKHGNTVLTALGGILKK-----KGHHEAE
VKHLAESHANKHK---IPVKYLEFISDAI IHV LHAKHPSDFG-ADAQAAMSKALELFRND
MAAQYKVLGFHG-----
-----
>GALLUS_GALLUS_MYOGLOBIN_gi|268607704|ref|NP_001161224.1|_
-----
-----MGLSDQEWQQLTIWGKVEA---DIAGHGHEVLMR LFDHPE
-TLDRFDKFK-G--LK--TPDQMKGSEDLKKHGATVLTQLGKILKQ-----KGNHESE
LKPLAQTHATKHK---IPVKYLEFISEV I I KVIAEKHAADFG-ADSQAAMK KALELFRND
MASKYKEFGFQG-----
-----
>VERTEBRATE CYTOGLOBINS
-----
>DANIO_RERIO_CYTOGLOBIN_gi|23308615|ref|NP_694484.1|_
-----
-----MEGDGGVQLTQSPDSLTEEDVCVIQDTWKPVYA---ERDNAGVAVLVRFFTNFPS
-AKQYFEHFR-E--LQ--DPAEMQONAQLKKHGQRVLNALNTLVENL---RD-ADKLN TI
FNQMGKSHALRHK---VDPVYFKILAGVILEVLVEAFPQCFS PAEVQSSWSKLMGILYWQ
MNRVYAEVGWEN-----
-----
>OVIS_ARIES_CYTOGLOBIN_gi|261244962|ref|NP_001159664.1|_
-----
--MEKVPGEMEIERRERSEELSEAERKAVQATWARLYA---NCEDVGVA I LVRFFVNFPS
-AKQYFSQFK-H--ME--EPLMERSPQLRKHACRVMGALNTVVENL---HD-PEKVSSV
LALVGKAHALKHK---VEPVYFKILSGVILEVIAEEFASDFP--PETQRAWAKLRGLIYSH
VTAAYKEVGWVQ-----
-----
>HOMO_SAPIENS_CYTOGLOBIN_gi|20987458|gb|AAH29798.1|_

```



```

-----
--MEKVPGEMEIERRERSEELSEAERKAVQAMWARLYA---SCEDVGVAILVRRFFVNFPS
-AKQYFSQFK-H--ME--DPLEMERSPQLRKHACRVMGALNTVVENL---HD-PDKVSSV
LALVGKAHALKHK---VEPVYFKILSGVILEVVAEEFASDFP-PETQRAWAKLRGLIYSH
VTAAYKEVGVWVQ-----

```

>VERTEBRATE HEMOGLOBINS

>EQUUS_CABALLUS_HB_ZETA_gi|167621441|ref|NP_001108014.1|_hemoglobin_subunit_zeta_

```

-----
-----MSLTKAERTMVVSIWVKISM---QADAVGTEALQRLFSSYPQ
-TKTYFPHF-----DLHEGSPQLRAHGSKVAAAVGDAVKSI-----DNVAGA
LAKLSELHAYILR---VDPVNFKFLSHCLLVTLASRLPADFT-ADAHAAWDKFLSIVSSV
LTEKYR-----

```

>HOMO_SAPIENS_HB_BETA_gi|256028940|gb|ACU56984.1|_beta-globin_

```

-----
-----MVHLTPEEKSAVTALWGKV-----NVDEVGGEALGRLLVVYPW
-TQRFESFG-D--LS--TPDAVMGNPKVKAHGKKVLGAFSDGLAHL-----DNLKGT
FATLSELHCDKLH---VDPENFRLLGNVLVCLVAHHFGKEFT-PPVQAAAYQKVAVGANA
LAHKYH-----

```

>TAENIOPYGIA_GUTTATA_HB_ALPHA_gi|323668297|ref|NP_001191172.1|_hemoglobin_subunit_alpha-A_

```

-----
-----MVLSAGDKSNVKAVFGKIGG---QADEYGADALERMFFATYPQ
-TKTYFPHF-----DLGKGSQVKGHGKVAALVEAANNV-----DDLAGA
LSKLSDLHAQKLR---VDPVNFKLLGQCFLVVVATRNPSLLT-PEVHASLKDFLCAVGTV
LTAKYR-----

```

>ORYCTOLAGUS_CUNICULUS_HB_THETA_gi|284005393|ref|NP_001164887.1|_theta_1_globin_

```

-----
-----MALSAERALLRALWKKLGS---NVGVYATEALERTLEAFPR
-TKIYFSHM-----DLSPGSAQVRAHGRKVADALTLAADHL-----DDLPGA
LSALSDDLHVRTL---VDPHFHGLLGHCLLVTLARHYPGDFG-PAMHASVDKFLHHVISA
LTSKYR-----

```

>VERTEBRATE NEUROGLOBINS

>HOMO_SAPIENS_gi|10864065|ref|NP_067080.1|_neuroglobin_[Homo_sapiens]

```

-----
-----P--ELIRQSWRAVSR---SPLEHGTVLFARLFALEPD
-LLPLFQYNC-R--QFS-SPEDCLSSPEFLDHIRKVMLVIDAAVTNV---ED-LSSLEEY
LASLGRKHRAVG---VKLSSFSTVGESLLYMLEKCLGPAFT-PATRAAWSQLYGAVVQA
MSRGWDG-----

```

>DANIO_RERIO_gi_18859087_ref_NP_571928.1__neuroglobin_[Danio_rerio]

```

-----
-----KLSEKDKGLIRDSWESLKG---NKVPHGIVLFTLRFELDPA
-LLTLFSYST-N--CG--DAPECLSSPEFLEHVTKVMLVIDAAVSHL---DD-LHTLEDF
LLNLGRKHQAVG---VNTQSFALVGESLLYMLQSSLGPAFT-TSLRQAWLTMYSIVVSA
MTRG-----

```

>GALLUS_GALLUS_gi_154816290_gb_ABS87379.1__neuroglobin_[Gallus_gallus]

```

-----
-----MLSRTQQUALIRESWRRVSG---SPVQHGVVLFSLFDLDPD
-LLPLFQYNC-K--RFA-SQOECLAAPEFLDHIRKVMLVIDAAVSHL---ED-LPCLEEY
LCNLGKKHQAVG---VKVESFSTVGESLLYMLENCLGAAFS-PDVREAWIELYGAVVKA
MQR-----

```

 >MONODELPHIS_DOMESTICA_gi_78486584_ref_NP_001030592.1__neuroglobin_[Monodelphis_domestica]

-----RRLSGPEQELIRESWQKVNS---NPLQHGMI L FTRLFDLEPD
 -LLPLFQYNC-R--QFS-SPQDCLSSPEFLDHIRKVMLVIDAAVTHV---EN-LSSLEEY
 LTNLGKKHKAVG----VKLSSFSTVGESLLYMLQCLGSTFT-VTMKEAWTQLYGAVVQA
 MSRGW-----

 >CHIONODRACO_MYERSI_gi|270341071|emb|CAR57914.1|_neuroglobin_[Chionodraco_myersi]

-----KLSGKDKELIRGSWESLKG---NKVPHGVV MFSR L FELDPE
 -LLTLFHYYT-N--CG--STQDCLSSPEFLEHVTKVMLVIDAAVSHL---DD-LPSLEDF
 LLNLGRKHQAVG----VNTQSFAEVGESLLYMLQCSLGQAYT-APLRQAWLNLYSIVVAA
 MM-----

 >BOVICTUS_VARIEGATUS_gi_270341067_emb_CAR57912.1__neuroglobin_[Bovichtus_variegatus]

-----KLSGKDKELIRGSWESLKG---NKVPHGVV MFSR L FELDPE
 -LLTLFHYYT-N--CG--STQDCLSSPEFLEHVTKVMLVIDAAVSNL---DD-LPSLEDF
 LLNLGGKHQAVG----VNTQSFAEVGESLLYMLQCSLGQAYT-APLRQAWLNLYSIVVAA
 MSR-----

 >SIPUNCULA

>THEMISTE_1_LGB2_gnl|ti|2001595631_T1_MM2_10H06_From_transcriptome_available_at_ftp://ftp.ncbi.nlm.nih.gov/pub/TraceDB/

-----MNQAPEDQTKFTPDELMLRLSWNAWVA--GDLAGXGFD MFV KMFEQRKE
 -TKEVFPFAR-G--N---DAXQMHQIINLVLFHVTKFMKNIDEVVKNA---DR-LEDVVSM
 LRRVEGRHGHQGHN--VPSAYFPFLGAAMH-TLIKANYKSYD-SKLDDCWVPLWNFMNNE
 MTTGQEVYRGGKI-----

>THEMISTE_2_LGB2_gnl|ti|2001595657_T1_MM2_11B08_From_transcriptome_available_at_ftp://ftp.ncbi.nlm.nih.gov/pub/TraceDB/

-----LRLSWNAWVA--GDLVGRGFDSFV KMFEQKKE
 -TKEVFSFAR-G--N---DAQMQQSSNVLFHVTRVMKNIDEVVKHA---DR-LEDVVSM
 LRQVGGRHGHQGHN--VPSAYFPYLGEAMR-TLIKANYKAYD-SKLDDCWVRLWDFINKQ
 MTTGQEYTEEEKS-----

>THEMISTE_3_LGB2_gnl|ti|2001594772_T1_MM2_01H11_From_transcriptome_available_at_ftp://ftp.ncbi.nlm.nih.gov/pub/TraceDB/

-----MNL DLEDQCRFDPEELINLRLSWNAWVA--GDL PGRGFDMFTKMFQRKE
 -TKEVFSFVR-G--S---DAHMQQSSKFLFHVTRVMKNIDEVVKHA---DR-LEDVVSM
 LRQVGGRHGS SHN--VPSAYFPFLGIALR-NLMSQSYKGYD-SKLDDAWTRLWGFMTSQ
 MMYG-----

 >BRACHIOPODA

>TEREBRETALIA_Ng2_gnl|ti|1987128689_Tt_MM1_02C05_From_transcriptome_available_at_ftp://ftp.ncbi.nlm.nih.gov/pub/TraceDB/

-----FTTQEK NLRSSWRAFVA--EDPAKRGYQMF KMFEENRPD
 -TKKFFK-FV-Q--KD--ALGTMKSSTSLVFHTGRVAKYIQLVCDCL---DD-PTEAVPL
 LRQLGGKHGCGPNNFNVDATYFPELGKAMRQLMESSGLQGYD-QELDALWEKLYTWITKR
 MEEGM-----

>ANNELIDA

>ALVINELLA_POMPEJANA_1_LGB2_gi|301587212|gb|FP491105.1|FP491105_FP491105_Alvinella_pompejana_whole_body_library_Alvinella_pompejana_cDNA_clone_MH0AAB59YJ14_5', mRNA sequence From ESTs stored at <http://www.ncbi.nlm.nih.gov/nucleotide/>

-----YRLS---DISERGMDFVRLFELHPV
 -YKSYFQKLR-D--V---DIEDLRQSGKLRVHSTSVMSKSIDLVETL---DH-PPDLRDM
 AIKIAHPHFDRG----VRPSQYRELFALILEYLLKDKAKVVFN-DEAEAAWQKLFDYVLDI
 TAAVMDLQIEKMG-----

>ALVINELLA_POMPEJANA_2_LGB2_gi|223839268|gb|GO218263.1|GO218263_CAGB27239.rev_CAGB_Alvinella_pompejana_Normalized_library_RN05_posterior_end_Alvinella_pompejana_cDNA_clone_CAGB27239_3', mRNA sequence From ESTs stored at <http://www.ncbi.nlm.nih.gov/nucleotide/>

-----MASYKPDPRCPLTERQLYSITKSWKAINR---EMASTAVNMFIRLLEHDGI
 --RSFFTQKFK-D--HK--TVAELRASKVFESHALMVISVIDDVITNL---DD-MDYVMSL
 LQATGESHSIKFKN--FNPDFLWNVEGAFLLWAVKETLGDRYT-ISIENIYTITIRYILQS
 LHDAFTKHRERQNSTNNDCEKTNLLNQELSTADRKT-----

>APHRODITA_ACULEATA_LGB2_gi|1491803|gb|AAC47259.1|nerve myoglobin [Aphrodita aculeata]

-----glsqaDIAVIRSTWAKVQG-sgSATDIGRSIFIKFFELDPA
 -AQNEFPCKG-E-----SLAALKTNVLLGQHGAKFMEYITTAVNGL---dDYAGKAHGP
 LTELGSRHKTRG----TTPANFGKAGEALLailasvvggdf-t-paakdawtkvyntisst
 mqa-----

>CAPITELLA_SP_1_LGB2_From_transcriptome_available_at_ftp://ftp.ncbi.nlm.nih.gov/pub/TraceDB/

-----DQLLTPEEIVLVRVTWEQLKT-nlTLANLGKKVFLRIFNLKPD
 -IKKLFPFSD-V-----WGDDLIRHPKFVLSERFMLVVDCCVQNL---ECIKSEhGEM
 LANLGRAHVNYkG---FSRENFEVFMKAIWVYVYHQLKDSMD-SEVECAWKKLLlfiivq
 qragydaekeappnglsfllqt

>CAPITELLA_SP_2_LGB2_From_transcriptome_available_at_ftp://ftp.ncbi.nlm.nih.gov/pub/TraceDB/

-----magnaqpfv
 scaggqppdqkpsnipqHEFLTQNKLGAKSTWEFLCH-tsTPTERGMRVFLRIFEIAPV
 -TKTLFPFKD-M-----PNEDLHRNSLFGKHATRFMKSVEFTMQNL---DALDVIvNPT
 LVSIGNKHVHIkG---FHPDYLDTFQTALMDIWDDELGKKFS-KETKEAWIKIFALITRK
 VFEGfqeettrfrpplpyegkq

>CAPITELLA_SP_3_LGB3_gnl|ti|1068987169_BGYZ80398.g1_From_transcriptome_available_at_ftp://ftp.ncbi.nlm.nih.gov/pub/TraceDB/

-----LFTDRQKAIITKTWRHMGN---DLTGRGSKVFLKIFNLHPE
 -VKQLFPSLK-N--D---NEDQLLKNPCFRGHASRFMQSVGAVVENL---DS-PGDLSPL
 LIDLGRKHVLFGG---FTPEYFAAFTEGMMCIWSEELGKGFT-DEVSVAWKTVFDFIMSQ
 LQDGYA-----

>PLATHYNEREIS_dumerilii_LGB3_454_sequencing_of_a_normalized_cDNA_library_from_Plathynereis_heads_and_mixed_larval_stages)_ (normalization_and_sequencing_was_done_by_the_company_Agowa)_ Kindely_provided_by_Kristin_Tessmar_Raible_from_Max_F._Perutz_Laboratories/_University_of_Vienna,_Austria.

-----PLNDNQKELIKKLWEIVRE--GDIESTGVSMYMHMFTLQPE
 -VMKMFsfvp-K--NVT-NPEDLKNsarflRHARNLMTTVTTAVDML---GD-MEGLSEV
 LVDLGRRHKKYR---AKTEHFPIVGRSLTHAISAATGDAFT-PETAFAFTAFYGVVTH-

>HELLOBDELLA_ROBUSTA1_LGB2_Helobdella_robusta_leech_jgi|Helro1|171404|fgene
sh4_pg.C_scaffold_20000312_From_transcriptome_available_at_ftp://ftp.ncbi.n
lm.nih.gov/pub/TraceDB/

```
-----MGANGF
KKVKSEPLLNLLNYLNNDVLTIREKQLVRESWTLLSI---KLKSLGKQVFLRIFELRPS
-TKNLFPFKT-V-----WGDKLIKHPFLFLTHSKRFVKVIGCVVDRL---DYLQEECAQP
LIELGKKHVSIEG---FLPDYYDVYIRAIISIWKQELKDVYT-NELSEAWHKVLVYIVSK
LKEGYETEWKVATYFNPO-----
-----
```

>ECHIUARA

>URECHIS_CAUPO_Ng1_gn1|ti|1987440364_Uc_MM1_01H01_From_transcriptome_availa
ble_at_ftp://ftp.ncbi.nlm.nih.gov/pub/TraceDB/

```
-----VSLQNIALLVLSWAILKK---DLLSNGAALFLALFEAYPD
-YKDLFKQFP-G--L---RLEDLTRMPPLRALGATFMHSLGSMVDNL---GD-LECVVEL
LRERTHTHWERG----IRFEHFQNVFDLLPAFLKSKLGYNFD-DATGAACAAASVMITVL
QAEKTL-----
-----
```

>PLATYHELMINTHES

>MACROSTOMUM_LIGNANO_1_LGB2_M1_aW_013_G17_c_From_transcriptome_available_at
ftp://ftp.ncbi.nlm.nih.gov/pub/TraceDB/

```
-----MLTADETOQLILSGWNQAMK---DAKGLGLDIFLTLFEMFPQ
-HQELFrDFK-G--K---SRAELEKMPKLRAGHLRVVNTLDGAIQSL---DD-MEVCVSS
LELIGASHKGRN----MNAGHFEDLNKALDVVFTRRLLGAAAYT--DNKAVWMKLegvipv
Iqrgm-----
-----
```

>MACROSTOMUM_LIGNANO_2_LGB2_M1_aW_008_G17_c_From_transcriptome_available_a
t_ftp://ftp.ncbi.nlm.nih.gov/pub/TraceDB/

```
-----MLNEVEKKIILSGWQQAIK---DKKALGMDVFMVTLFEMFPQ
-HQELFrDFK-G--K---SRAELEKMPKMRALRVVNTLDGAIQSL---DD-MEVCASS
LELIGASHKSHH----LSAKHFEDLNAALAVVFERRLLGKAFV--DNKAVVWVWVLLQGIpv
Iqrgl-----
-----
```

>CHOANOFLAGELLATE

>MONOSIGA_BREVICOLLIS_LGB2_gi|167520949|ref|XP_001744813.1|_hypothetical_pr
otein_[Monosiga_brevicollis_MX1]

```
-----MS
DRRRSSGSSEDEAHAPDYPSFDPVAVKLARKQWKRVVQ---LVPNWHEVFFSYLFFERAPY
-ARTLFPFD-----VDRLQGNSSLAEHAKRVGQALETALQGL---FE-YYSLVEV
LEKLGRRHFYKYG----VEPEHIDLFEETFYKTLAIGLGKKWN-PEARRAWEIVCGLILSP
IRTGILQARTKANHLRAKEAERKRQLEMAAARLEGRVASSGVQFSSDTERSRRSTAASAT
ATPHCGLKSNRFSSNSLRKTIL
```

>SALPINGOECA_SP_Ng2_gi|326430027|gb|EGD75597.1|hypothetical protein
PTSG_06664 [Salpingoeca sp. ATCC 50818]

```
-----MRLDMEQLKIALGSWTAVVE---LVPTWHEVFFAELFQAHPE
-TERLLYSS-----DKSKSWNERHMARVGVKSGDVIKSL---SN-YDDVIEH
LTALGTRHARYG----LHVDQLDLFINAFLWTLGAGLGDSWD-HSVKKAWMHVLPFILSP
LKSGLVVARTLRNDYNTSGCLRCRLLIPLHGRRLRPITVSLV-----
-----
```

>PLACOZOA

>TRICHOPLAX_ADHAERENS_1_Ng3_fgeneshTA2gi|196007506|ref|XP_002113619.1|hypot
hetical protein TRIADDRAFT_57230 [Trichoplax adhaerens]

```
-----MDQAQTDVQTPPQPSLTEEQKAIIRENQDVVEE---NMSEVGLYLFSLFTIAPE
```

-YREVFPFPE-----TTTNDVRLRVHATGVMKTVGKAVQNL---DQ-FSELQSA
LSTLQGQFHRKA----IKFENFQAVGQALIQTLSDKLQENFT-PEVHEAWSKTFDMITAA
MKSGMN-----

>TRICHOPLAX_ADHAERENS_2_LGB3_gi|196012120|ref|XP_002115923.1|hypothetical
protein TRIADDRAFT_59832 [Trichoplax adhaerens]

-----MAPTAQDLQTIRETWALVAP---DLKKHGTVLFRLFEQHPD
-VQRLFEEKIK-D--V---PHDQLATNENFVFHTTRVMETIDHAVKGI---DN-LPALTVL
LKQLGSSHAQYN---VKKEYFKIGLRIFE-----

>TRICHOPLAX_ADHAERENS_3_Ng1_gi|196001583|ref|XP_002110659.1|hypothetical
protein TRIADDRAFT_54901 [Trichoplax adhaerens]

-----MVLVNNYSLIKLSPA
-TKIYFHGVD-F--EK--RDSYLAKNFTFLRNHAARFMEAINVIIGQD---MD-IFSVESY
FRVVGSKHHSYN----LKLEHVQDISDAFLEMARNALKKKFT-KSTEAAWSFFQMVTD
IKNGIMKAQNRN-----

>TRICHOPLAX_ADHAERENS_4_LGB3_gi|196016934|ref|XP_002118316.1|_hypothetical
protein TRIADDRAFT_62364 [Trichoplax adhaerens]

-----MTKI
DSENVKSKNVVTGNDIKSYLNYQERQAIIDSWNAIST--EKQKYGTILFLKLFLEPR
-VKSLFTIFD-F--NE--PLEDIIQSPHFRSHAMRFMQSLETGVLMG---FD-KESCDFL
FKSLGSRHHFYD----LKSEFLDVIPECILHTIKKGCNNWS-NETADAWKIATKVLCEL
FREGLETKPKK-----

>PORIFERA

>CARTERIOSPONGIA_FOLIASCENS_LGB3_gi|241971149|gb|GO083496.1|GO083496_DMP57
15864_Carteriospongia_foliascens_DMP_cDNA_Library_Carteriospongia_foliascens
_cDNA, mRNA sequence From ESTs stored at <http://www.ncbi.nlm.nih.gov/nucest/>

-----MSHPLSSEELKLVEKSWALIQE--FGLQEAGMVMFKRLFEMSPG
-LQNLFPFA-----GDSLSDNEGMKHSFIVMTSLDDCLKIV---HD-VPKLDKDE
LISLGAVHHIQG---VTSEHFAPVGSALIWALEAALKEKFT-PEVKAAWLALYKVVQSL
MEQGMEEGI-----

>AMPHIMEDON_QUEENSLANDICA_LGB3_gi|340378768|ref|XP_003387899.1|_PREDICTED:
neuroglobin-like [Amphimedon queenslandica]

-----SLTSAQVALIESTWVKVVK---DLQGAGNIMFLKLFQIDVS
-VRDKFPFRD-V-----PYEELEDSESLKHSQVMETIDLAITLLLG-GE-MEKLVEA
LVDLGMAMHMQG----LKPEDFDHVGEALVHALGVALGKEFN-DEAKKAWTLLYSVVTAK
MKEG-----

>CNIDARIA

>NEMATOSTELLA_VECTENSIS_1_LGB3_gi|156408000|ref|XP_001641645.1|_predicted_p
rotein [Nematostella vectensis]

-----TENTLTSVPLSTRKLVRESWELIEP---VKITIGKRLFTRFLFDVNP
-MQDTFPNFK-G--K---ELKDILNSRSLYLHAKRVMVAVENAVTVL---DD-AETFESY
LINLGGRHLPWG---VTKDHFVVGGEAFIWAQDVLGEGCT-SDVAEAWIDLYGYIVQA
MLEG-----

>NEMATOSTELLA_VECTENSIS_2_LGB2_gi|162098184|gb|FC288990.1|FC288990_CAGN4759
.fwd_CAGN_Nematostella_vectensis_Nemve_mixed_stages_unfert_eggs_to_primary
polyps_Nematostella_vectensis_cDNA_clone_CAGN4759_5', mRNA sequence

```

-----
MHGVIEEGLLQLERINPITGLSAREVAVVKQTNLTKP---DLMGVGMRIFKSLFEAFPA
-YQAVFPKFS-D--V---PLDKLEDTPAVGKHAI SVTTKLDELIQTL---DE-PANLALL
ARQLGEDHIVLK----VNKPMFKSFGKVLVRLLENDLGQRFS-SFASRSWHKAYDVIVEY
IEEGLQQSYKQDPVTGITDAEKVLVQESWDLKPDLLGLG-----
-----

```

```

>NEMATOSTELLA_VECTENSIS_3_LGB2_gi|162098183|gb|FC288989.1|FC288989_CAGN4759
.rev_CAGN_Nematostella_vectensis_Nemve_mixed_stages_unfert_eggs_to_primary
polyps_Nematostella_vectensis_cDNA_clone_CAGN4759_3', mRNA sequence
-----

```

```

-----NPQNAFSAADIQAIQGTWALAK---PDLMGKGAMVFKQLFTEHGY
--QPLFSNLA-Q--Y---EITGLEGSPELNTHARNVMAQLDTLVGSL---QN-SIELGQS
LAQLGKDHVPRK----VNRVHFKDFAEHFIPLMKADLGDEFT-PLAESAWKKAQFDVMIAT
IEQGQRARRSVATFLTNPVA-----
-----

```

```

>NEMATOSTELLA_VECTENSIS_4_LGB2_gi|162112725|gb|FC303536.1|FC303536_CAIC1367
1.fwd_CAIC_Nematostella_vectensis_Nemve_whole_embryos_normalized_Nematostel
la_vectensis_cDNA_clone_CAIC13671_5', mRNA sequence
-----

```

```

-----CFKAFN-K--V---SLEDLEKSPFLKAHATSVMSAINEVVCNL---DE-VEILGIL
LEKIGFSHARRE----IRRIHFENLAKVVVAYLRQALGSHLT-EEGADAWRKALCVMIDI
IEKGSTSERW-----
-----

```

```

>NEMATOSTELLA_VECTENSIS_5_LGB3_gi|156405932|ref|XP_001640985.1|predicted
protein [Nematostella vectensis]
-----

```

```

--MGCSSSTFKPPREPVKIPLSVAQKYLVRWETWETIEQ---HSKAVGKKTFLRFFEMNPD
-YQKLFPEFA-T--L---DQVELEQANALHGAERVMKAVENAVSAM---DD-AESFAAY
LENLGRHKKARA----LKPAYLDAMQVAYTDTIQDLLKQWT-DGTAEAWNKLFRFIADT
MKHGLSS-----
-----

```

```

>NEMATOSTELLA_VECTENSIS_scaffold_7000121_LGB3_from_Bailly_and_Vinogradov_20
08
-----

```

```

-----MCGGASKTLTTPHGTEEH
LTKKSQSENGNQSFVGNRPLRTERQIKLVQDTRWLLIP---SQKKTAMIFYLKLFTLDPI
-FKEVFS-FH-T--E---NEGQLEQDERFLFQSRKFMEMINSAVDRL---ND-ISLLVMI
LKSLGCVHWTKF---KIKPEYYEPVKGALIIYSISKGLGSLFN-DEIGEAWQAMYDLMSG
MISGTKAVQARSQNSL-----
-----

```

```

>NEMATOSTELLA_VECTENSIS_scaffold_3000224_LGB3_from_Bailly_and_Vinogradov_20
08
-----

```

```

-----MCGGASSTV
RPFIFIRQPASDTENTLTSVPLSTRRKKLVRESWELIEP---VKITIGKRLFTRLFDVNP
-MQDTFPNFK-G--K---ELKDILNSRSLYLHAKRVMVAVENAVTVL---DD-AETFESY
LINLGRHLPWG----VTKDHFVVGAEFIWALQDVLGEGCT-SDVAEAWIDLYGYIVQA
MLEGLQQAQKGR-----
-----

```

```

>NEMATOSTELLA_VECTENSIS_scaffold_76000030_LGB2_from_Bailly_and_Vinogradov_2
008
-----

```

```

-----MGCASSLATQTKL
LKGHLPTCETQYLTQVDQLPLTETQKYYIKQSWMGLES---NKGELGIEIFLRLFSENPT
-LQLMFPEFR-E--YS--TLEELKESRSLQGHGTRVMKVVENAVNSL---ED-GHALMEY
LQELGRRHKTRQ--IKPTVSNLQEI SQAINETFEENLGIKWT-VEIAESWKLKLLDYVMAM
IIRGLRSP-----
-----

```

```

>NEMATOSTELLA_VECTENSIS_scaffold_50000067_LGB2_from_Bailly_and_Vinogradov_2
008
-----

```

```

-----MGCVVSKNPST
VAKIVPGGGEELKFETSRIPLDAKETQLVRKTWAILGD---RQVEVGKSLFLRFFEEHPT
-SKDLFPFR-N--I---SNEKIAESPALYGHARRVMKSVDNAVASI---EN-VQVYSAY
-----

```

LYELGTRHQTRQ----LSEEQLKFMGGAFLLFAMRLHLRKEWS-RATSKAWEKIFSFMADA
MMRGCKG-----

>NEMATOSTELLA_VECTENSIS_scaffold_4200019_LGB2_from_Bailly_and_Vinogradov_2
008

-----MGCG
SSVVTQGNMPLHCLGLKLECDMTYEQKYLIRETWKFLEV--SKKEIGVSVYKRFLNMHPG
-LQTYFSEFK-----HIKIDNINGSHGHPRLMLAIDNAVTAL---GD-SDSFSAY
LVELGRRHHGMNFRPGPTHFNDLRKCFLSVIEEILATASLWD-FQVEEAWNRLFDSITAM
ILRGIQLAKV-----

>NEMATOSTELLA_VECTENSIS_scaffold_5000153_LGB3_from_Bailly_and_Vinogradov_2
08

--MGCGSSTFKPPREPVKIPLSVAQKYLVRTWETIEQ--HSKAVGKKTFLRFFEMNPD
-YQKLFPEFA-T--L---DQVELEQANALHGHAQRVMKAVENAVSAM---DD-AESFAAY
LENLGRHAKARA----LKPAYLDAMQVAYTDTIQDLLKTQWT-DGTAEAWNKLFRFIADT
MKHGLSS-----

>NEMATOSTELLA_VECTENSIS_scaffold_141000032_LGB3_from_Bailly_and_Vinogradov_2
008

---MGCSSLSQANLPRTMPLSEAQKYLVRTWETIEP---QKQTVGKKAFLRFFDMNPD
-YQNLFPPEFK-S--L---SYEELQKANALHGHAQRVMKAVENAVMSI---DD-VMSFSAY
LEELGRRHAKTRA----LKPSTYLEAMHGALMDTLRNLQSQWT-EETAEAWNKLFSFISTT
MVRGLQSRD-----

>CLYTIA_HEMISPHAERICA_1_LGB1_gi|294376172|gb|FP931337.1|FP931337_FP931337_C
lytia_hemisphaerica_library_Clytia_hemisphaerica_cDNA_clone_SA0AAB120YK22_5
, mRNA sequence

-----MGAICSHKTL
VSKSFVYIKSIQKSSNIKVPLSAKEVNILKKTWPPVKN--NWLKICLVAFERWFTLYPQ
-LRYMFRSLP-E--DI--QFEDLFKTDALKMHVDKVRDVLLELLIKKI---DN-VEELVNT
LVDFGRQHMLG----AEQRYATALAASFQYIGICMIMDVD---SSVENAWDSLRLRFVMDL
LKLGMROMEKAEQEKESLNKGYNTEELLEKAQDGGALDENENSAPMPLALINEDDSTSSV
RTFSCR-----

>CLYTIA_HEMISPHAERICA_2_LGB2_gi|294400477|gb|FP945496.1|FP945496_FP945496_C
lytia_hemisphaerica_library_Clytia_hemisphaerica_cDNA_clone_SA0AAB133YH06_5
, mRNA sequence

-----MGSS
GSCLKFTMKPNGKVDAAPADYTEAEINIVQKQWSVAMR--NLDSVGYKLFKALFQYKDI
MAKFDFAVAN-N--L---DFQKSMNDARLSFHIRRVFHTINTVVVSL---ND-GDFVASQ
LEHVGAIHAEYG----LQATHLARFKDVMLETLEEFKQEGFQ-EDSKTAWSKIVDAIAKY
MLKGDIDKAKEQTKTSELSSDDGDGKMEQ-----

>CLAVA_1_LGB1_Locus_11167_Transcript_3/8_Confidence_0.360_Kindly_provided_b
y_Stefano_Piraino_from_Universita_del_Salento_Italy

-----MGSKLCLAIHK
SVARSTQPSIHTKYGNICIPLTKKDIKLRKSWGIMKM--NWKICLVVFDWRWFSMYPE
-LRNMFKSFS-S--VL--TLQELLASRTMKSHIQQLEELVEDLLYKV---DD-PTDFIET
VIQFGEHHLG-----AKQLYATSLAAAFQYAICISLNLD-RDTENAWDSLFRFLMDC
LKFGMRNEMHKETTSPDKLEVQDG-----

>CLAVA_2_Ng_Locus_28538_Transcript_4/6_Confidence_0.667_Kindly_provided_by
Stefano_Piraino_from_Universita_del_Salento_Italy

-----EVGILRKQWSIMMS---DADANGIMLFTALFAINGH
-LRTKFAFGD-V--YGF-DCKTTFGDERLLGHLRGVFSVEMLINSLVQ-NT-FYESLGK
IKELGKVHSLAG----IHLSDLQAFKKAMIQTLKKEKSEFD-DKALKAWSKFIDIVIEA
FDVNDTDDDEKTQPIGK-----

>HYDRA_MAGNIPAPILLATA_Ng1_gi|221117935|ref|XP_002162062.1|_PREDICTED:_similar_to_neuroglobin_[Hydra_magnipapillata]

-----LSGKEIETLKKSWTTAKQ---FWNEICTCAFSRWFSTYPE
 -IQSKFGVYG-D--NL--TMNEVLASESLCIHIRKSVELIEIIKKV---DE-RHELSEY
 LIELGKLHHKFG----AEQKYATALGSSSFVFAISQICPNIDM--ITEGAWDSLFKYIVT-

>MONTASTRAEA_FAVEOLATA_1_LGB3_gi|282539129|gb|GW263294.1|GW263294_CCHW10651.b1_CCHW_Montastraea_faveolata_heat/dark/disease-stressed_adult+_6_day_old_larvae_Montastraea_faveolata_cDNA_clone_CCHW10651_5', mRNA sequence

MGCGGSKAIKNRTAPAPVAEQTQSPGLKETSPAKQHRQQPEGKRTEETVEETAPADGYGR
 QGTEGDKQATEEQSENVEGPITQEQISLVQDTWKLNVNG---DLEQVGVVEFYTRLFKENPE
 -LLQMFSFRD-L--ANS-TEDAMRTDDRFRKROGLVTMQHVDLAVASL---SD-LGSIVPA
 LKDLGARHSMYK----VEEHFGPVG-----

>MONTASTRAEA_FAVEOLATA_2_LGB2_gi|282548677|gb|GW271367.1|GW271367_CCHW15436.b1_CCHW_Montastraea_faveolata_heat/dark/disease-stressed_adult+_6_day_old_larvae_Montastraea_faveolata_cDNA_clone_CCHW15436_5', mRNA sequence

-----MERSKESDGLTDLQIEMIRSSWEKVTP--NKKHHGQLLFHKLFEIAPE
 -MTDLFPFG-----DDFTKQFTTHALNIMNALDHAIQNL---DN-PDVLPK
 LRELQMHAGFE----LTIKEFQHVGEALIWVLTGLGDDFT-----

>ACOELA

>SYMSAGITTIFERA_ROSCOFFENSIS_2_LGB2_From_the_present_study:_illumina_library_made_by_Genoscope,_French_Sequencing_Center

-----MQLNNIQKCLPFL
 FPKRNSHMSDYFAFEGLPPVPNSSESEEIRKSWKKIEL---DAAKLGIAVVFVGLFERYPE
 -IQASFSKIA-N--V---NKSSLNSNVMLHAHSLHIMHMIGKLIQLL---NE-PENLLSK
 VVELGERHFDRK----ANDELLQYFCPAY-VEAMAKKGQWK--KKTIIAWEKFFDFIRAA
 MVHGLKKRKGHSSISNTTSAANTAAEKNHNSPSSQ-----

>SYMSAGITTIFERA_ROSCOFFENSIS_1_3D_LGB2_From_the_present_study:_illumina_library_made_by_Genoscope,_French_Sequencing_Center

-----MATLESMQVSEEQOSLIMEDVQVLLP---NYDDFVEDVLQQFMEENPE
 -TFQIFPWAD-A--SK--TAKEMRSHPRFKSHAKSIGKVISDCLVDL---NG-VKKHEPK
 LSSLGAMHTKKK----VPTELFQKLGGCILTQVVKRVSEAKWSEEKKEAWLKAYGIITVM
 VTE-----

>NEMERTODERMATIDA

>MEARA_STICHOPI_Ng1_transcript_1_Kindly_provided_by_Andreas_Hejnol_from_SARS_International_Center_for_Marine_Molecular_Biology

-----SRKILVCRLFESHPO
 -VRDIFEHSE-----DKDLVPADIIKAHGMRIMGLLGRFVSNLESNVD-DNLLGQP
 IHDLGRKHVDFK----APPYLFDFVALQIQHIIKSKLEQVWT-DEIGDAWKVMFDIIVFN
 LKSGQNQEMQDRGITV-----

>NEMERTODERMA_WESTBLADI_1_LGB2_gnl|ti|2247831830_Nw_MM1_39E05_From_transcriptome_available_at_ftp://ftp.ncbi.nlm.nih.gov/pub/TraceDB/

-----MGLTETQRVLIKQSWKKVIS--GGRVEAGWVLFKFFTSPE
 -AQNYFKAFK-G--K---PLSELQNNTQMKGHVLRVINYITDIVDTL---EV-DEMREEM

SINIGRTHGRRRA----IPAEMFQCLKPAVFGTVIDLNGGSLS-DEAAAAWGLLWEALVLC
VLEGMKKPGQPGMGQTHSHADQIIIGELLVTHPAHVVALYITNYLLLFIKYLHHEPK----

>NEMERTODERMA WESTBLADI_2_LGB2_gnl|ti|2247830945_Nw_MM1_30C08_From_transcriptome_available_at_ftp://ftp.ncbi.nlm.nih.gov/pub/TraceDB/

-----MG
SYLSGLLWNSVDPIDELTKLTPSEKNALVDSWMLVID--DGLQQGGIKLFLKFFTLDP
-ARPYFTKFL-K--L---SDDELRESKMLRAHVIRVMNTLNSLVDGL---DD-PELFVEL
SKFIGRTHYRHN----IDTEHLVNLGEAVIWLVEQSNNGRLP-PAVEDSWQRLWERITHF
ILAGEEEEEKTDGGETGVSVEQNSHVIMSTGVQSNLNFHHIFFNNPCYCIPLT---

>NEMERTODERMA WESTBLADI_3_LGB1_gnl|ti|2247830237_Nw_MM1_22H08_From_transcriptome_available_at_ftp://ftp.ncbi.nlm.nih.gov/pub/TraceDB/

-----MALTDSQQFLVKKVWDKVVVA--DGYHEPGLIFMNKFFTDTP
-AQNYFWFKV-D--R---PLYELRENIHMEAHLTRGLSHITCTINKV---AE-PTVCADM
WRNLGRTHTAHT----VPFKMFDLMPAFVNMVIQCCGMVD-QDTLVALSLYWFIMRC
VKEGMEEAAAKVWSDSTTLNGSLYLKIINTPLLYCPCHGRTVSDIITRCMLRVAII----

>XENOTURBELLIDA

>XENOTURBELLA HEMOGLOBIN_RELATED_S02970_XB_MM1_31H06_From_transcriptome_available_at_ftp://ftp.ncbi.nlm.nih.gov/pub/TraceDB/

-----MVFAGLSGGEVAAVKSSFGKVTD----LAGTGAELFLKYFQARPS
-AQNYFKWKG-Q-----SLDAVKADAYLQTHGNRMVSTLKTIVIGSI---ED-AGKCEAL
ITQLGNLHTGRN----VAAADFEALFPIIAGVLSEKCGGALA-GDASSGWQKLYNEIVRL
IDAGQKTPAAGGARAFLAAIGAN-----

>MOLLUSCA

>SEPIA OFFICINALIS_LGB2_Sequence has been provided by L.Bonnaud and Y.Bassaglia from an ESTs library built from Sepia officinalis embryos by the Genoscope/CEA (project AP07/08_n?07-http://www.genoscope.cns.fr/spip/Collection-d-ESTs-d-embryons-de.html)ADY0AAA118YM16CM1

-----MGALFSVIDAL
IRKIGIMSPLDDEEVDSTGLKVKKEKRAMTQSWKYISP---NLKAEGIHFFNMLFTEHPD
-YIDYFPSFR-G--K---KLEEFNTKASFMAHAKNVFYAVTLIVDTL---DD-ADELVEI
LLKTGRNHRRS----VPLSPFQGRAIVFEKIMTERLGKAIP-PLGKGLVDKSIDCS---

>EUPRYMNA SCOLOPES_1_LGB2_gi|84449782|gb|DW284378.1|DW284378_UI-S-HH0-aea-p-02-0-UI.s1_UI-S-HH0_Euprymna_scolopes_cDNA_clone_UI-S-HH0-aea-p-02-0-UI_3',_mRNA_sequence
From ESTs stored at http://www.ncbi.nlm.nih.gov/nucest/

-----MGALLSVIDAV
IRKIGIMSPRDDEEVDSTGLKVKKEKRAITQSWKHVSQ---NLKQESMNFNLLFTEHPA
-YIEYFPAFC-G--K---KLEEFNTKPKFIAHAKNVFYALILVVDL---DE-PDELVEI
LLKTGRDHNRG----VPMAAFHNLAIVFDKFLTIRLGNNYT-PLAKESWTKALTVVNAV
IEKGIEDGIDCSECRYRERNRST-----

>DORYTEUTHIS PEALEII_LGB_related_gi|342663023|gb|JK329464.1|JK329464_oy57h01.y1_Woods_Hole_Squid_Stellate_Ganglia_cDNA_Library_Doryteuthis_pealeii_cDNA,_mRNA_sequence_From_ESTs_stored_at_http://www.ncbi.nlm.nih.gov/nucest/

-----VRADGLPYPPPPPTDPRPLPLSPLQVFKLKKSWKGIKR---SIELTGVMFVRMFRTQPG
-LKNLFKDFR-E--LE--TDDEMRENEALEKHATLVMNTLDDAITNI---EN-VDLVLDL
LHRIGKSHLRFQG---FNVEYFWLAEQPLLDAIKITLGDrys-DNMDIYKLVIRFLLTE
VTKGARVDVSS-----

>EUPRYMNA_SCOLOPES_2_LGB_related_gi|84436671|gb|DW271268.1|DW271268_UI-S-GS1-acj-k-06-0-UI.s1_UI-S-GS1_Euprymna_scolopes_cDNA_clone_UI-S-GS1-acj-k-06-0-UI_3', mRNA sequence From ESTs stored at <http://www.ncbi.nlm.nih.gov/nucest/>

-----I--E---TDDQMRNEALEKHATLVMNTIDEVIANI---EN-VDLVLDL
LHRIGKSHCRFQG---FNVEYFWLAEQPLIDAIAIKITLGDRYS-DNMNIYKLVIRFLLTE
VTKGARANVSST-----

>MYTILUS_CALIFORNIANUS_1_LGB2_gi|145897418|gb|ES403000.1|ES403000_MUT03-C03.xld-t_SHGC-
MUT_Mytilus_californianus_cDNA_5', mRNA sequence From ESTs stored at <http://www.ncbi.nlm.nih.gov/nucest/>

-----GELATDTIRSTWPLLSLSD---DIERTGIKVFLRIFYEYEPK
-IRNVFKRFG-E--M---DEIELRRSPIFKEHAYRFMRVVDLVDNM---DPPKAHIQQN
LMMLGAKHATFEG---FRIEYFEAYSESLIDVWEYTGEEFI-PEVRESWTEFFDYLVKY
MCQGYNVFTNET-----

>MYTILUS_CALIFORNIANUS_2_LGB2_gi|223024721|gb|FL490027.1|FL490027_Mg_Nor01_51M10_Nor01_Mytilus_galloprovincialis_cDNA_3', mRNA sequence From ESTs stored at <http://www.ncbi.nlm.nih.gov/nucest/>

-----QTMESSSTISAIKNTAAALEE---HSSQLGLKTYELLYEEHPE
-VKATFNKS-----EYSTEADAVPKDLAVKVGKAIATFAKHC---DD-LDSQKEL
FGGIAIKHVKLGG---ILEEQYSAMEGSFLKAVKEVLGDAAT-DEVLDHWKKGFFHFLAEH
IRGLEAEK-----

>SPISULA_SOLIDISSIMA_LGB2_gi|76058055|emb|CAJ31107.1|_nerve_hemoglobin_[Spisula_solidissima]

-----DPCPVTKLTKAEKDAVANSWAALKQ---DWKTIGADFFVKLFETYPN
-IKAYFKSFD-N--M---DMSEIKQSPKLRHSINFCHGLNSFIQSL---DE-PDVLVIL
VQKLTVNHFRK----IAVDRFQEAFFALYVSYAQDHAKFD---DFTAAAWTKTLKVVADV
IG-----

>APLYSIA_1_LGB3_CNSN01-C-002803-
501 From transcriptome available at <ftp://ftp.ncbi.nlm.nih.gov/pub/TraceDB/>

-----IFSLKQSWKGIKR---KMEDTGVDMFVRLFKTSSH
-LQTMFQGF-E--D--IR--SDELRSNEALEYHATLVMNTLDDAITHI---DN-YDFVKQL
LSKTGASHVKFAG---FKSTNFLAIKGPFLAVRVTLGDRYT-ENMQNIYTIATIFILET
LQQGMEEALEKAGSSEVAQGHIRV-----

>APLISIA_2_LGB2_CNSN01-F-074342-
501 From transcriptome available at <ftp://ftp.ncbi.nlm.nih.gov/pub/TraceDB/>

-----MGAEQSLS
CFPCTSVRRRCRRRHQCLSKYLNPRQVQLVQCTWSILKM---DLTTLGVTVFLNFFETEPD
-LKMMFPKMI-RM-NESNQLEWDVDKMDLQKHAVTVMEGLGAAVETL---ND-SQFLNTV
LIALGQTHEKRN----IKPNMLKRMWPSMHVGLGAVLGEQY-KEVSEAWCKLYSYICLQ
MQIGMENPN-----

>LIOLOPHURA_JAPONICA_Ng1_gi|47115693|sp|Q7M416.1|GLB1_LIOJA RecName:
Full=Globin-1; AltName: Full=Myoglobin I

-----gISADQAKALKDDIAVVAQ---NPNGCGKALFIKMFEMNPG
-WVEKFPKAWK-G--K---SldeIKASDKITNHGGKVINELANWINNI-----NSASGI
LKSQGTAKHGRS----IGIEYFENVLPVIDATFAQQMGGAYT-AAMKDALKAAWTgvi

```

gmkagy-----
-----
>CRASSOSTREA_GIGAS_2_LGB3_gi|313365341|gb|HS221690.1|HS221690_CCTS16119.b1
CCTS_Crassostrea_gigas_mixed_adult_tissues_library_4_normalized_Crassostrea
gigas_cDNA_clone_CCTS16119_5', mRNA sequence From ESTs stored at http://ww
w.ncbi.nlm.nih.gov/nucest/
-----
-----MTSTEFTDEQKQLVKTTWNIVRE---DISKVGVITFLRLFEEKFPD
-VQDLFVFPFR-G--L---NSEELRQNVGLREHGMRVMGTIEKCITRI---DQ-PDKLTSM
LESLGAKHVVD---TKIEYFDLLSPQLIQAITTRHWSVD--SQREQAWNNFYCTL---
-----
-----
>CRASSOSTREA_GIGAS_1_LGB2_gi|318048551|gb|FQ663445.1|FQ663445_FQ663445_Cras
sostrea_gigas_library_(Genoscope_-
CEA)_Crassostrea_gigas_cDNA_clone_WY0AAA49YL07FM1, mRNA sequence From ESTs
stored at http://www.ncbi.nlm.nih.gov/nucest/
-----MGC
DASKSTDVIENEEIESDTDFDFTDTQIDTIRSTWPILAC---DMVDIGSKVFLKIFIDEFK
-LKYAFPSFS-D--M---EENELLRHPPFIDHVTRFMQIIDYLVENL---DQQNSDFHQA
LLMLGAKHATYPG---FQVSHFTVFNKALLEAWESAIGEEFI-PEVQNCWTQLFAYIMRY
IVQGYELYFSEC-----
-----
>ARTHROPODA

>APIS_MELLIFERA_LGB2_gi|118150510|ref|NP_001071291.1|globin 1 [Apis
mellifera]
-----MGT
FLRFLGISSDDNRIDQATGLTERQKKLQNTWAVVRK---DEVASGIAVMTAFFKKYPE
-YQRYFTAFM-D--T---PLNELPANKRFQAHACAGVITALNNVIDFL---HD-PGLMEAS
LIGLVERHKKRG---QTKEEFQNLKEVMLEVLRLQALGQYT-PEVAEAWNKTLDMMF GK
IYQ-----
-----
>CARCINUS_MAENAS_1_LGB2_gi|299757081|emb|CBN88274.1|_hemoglobin_[Carcinus_m
aenas]
-----MGAVLSVVGWLSPGTQVGAV
TFPEEGSLGPEADVDPKTTGLTLRHRTAIYRTWDLVRP---NPKLHGINLFLTMFQEEPV
-LQTRFKGFA-G--K---SIEELKNSKRLAAHGTTVVMATAMVDNL---ED-VSVLVEL
LKNTGANHRDRG---VPKGFELLAPVLRFLKDNLGSAWS-PVAEEAWTQAMKVINAV
IFTSYDACSILS-----
-----
>CARCINUS_MAENAS_2_LGB_related_gi|84413153|gb|DW250579.1|DW250579_Cm_mx1_36
e01_SP6_Green_Shore_Crab_Multiple_Tissue, Normalized_Carcinus_maenas_cDNA_c
lone_Cm_mx1_36e01_5' similar to gb|EAL38715.1|ENSANGP00000028536_Anophele
s_gambiae_str. PEST. Score = 156_bits (395), Expect = 4e37, mRNA sequence_F
rom ESTs stored at http://www.ncbi.nlm.nih.gov/nucest/
-----MGCHITKNKKNKETE
VKVIDLPEPPEPPPPDPRPLPTARQRFNIIKSWKGIAR---AIEPTGVNMFVKLFENHSE
-LITFFTKFR-Q--LR--TRDEQAESLELAEHATIVMNSIDEGIKAM---DN-VDFFFDL
LHQIGASHRKIPG---FKKEYFWKIEHPFLEAVRLTLGDRYT-DNMDNIYRITIKLLIET
VVRGYELAELEKE-----
-----
>DAPHNIA_PULEX_GlobinX3_gi|321478927|gb|EFX89883.1|hypothetical protein
DAPPUDRAFT_231989 [Daphnia pulex]
-----RLEK---DIAQVGIIVFINLFFETHPD
-MQSVFLPFT-----GVVLDLKKSKLLSEHALRVMGAVQRAVHRLQEPKLFHAF
LSELGRKHEKNG---AKLEYIDYIGPQFLCAIRPILGDDRT-LETEKAWTLLLDYMTAA
MKESLVEARNASAAESSKPLTLPPSSSSSSSAATDD-----
-----

```

>DAPHNIA_CARINATA_LGB2_gi|290774776|gb|GR506052.1|GR506052_LAaa_0006_F05_Ga
mogenetic_water_flea_(Daphnia_carinata)_cDNA_library_Daphnia_carinata_cDNA
5',_mRNA_sequence_From_ESTs_stored_at_http://www.ncbi.nlm.nih.gov/nucest/

-----MDTLKTVNVAAVQNTWAIIVKK---DLNTHAPQFYVALLTAHPE
-YQPMFPTIA-N--V---PAGELLNPNALKTLSVNVLTKLSELIDCMG----NPDALQGQ
LVDLANQHKQRG----TTRAHFNDLSKVLIDFLAAKLGGEFT-PEARQAWTATMQGINTV
VEASS-----

>HARPEGNATHOS_SALTATOR_LGB2_gi|307192580|gb|EFN75768.1|_Globin_[Harpegnathos
s_saltator]

-----EKTGMSEKQKKLVQNTWAIIVRK---DDVSSGLAIMNAFFTRYPE
-YQDQFKSFK-G--I---PFEELSKNKKFQAHCVSVIAGLSNVIDHI---HN-PELMEAS
LINLAERHKNRG----QTREHFQNLRYVLEDLIPSVFGKQYT-QEVQEAWKKMFDYLF--

>TUNICATA

>Ciona_intestinalis_Gb4_HBG_related_NP_001027701.1

-----MQSMSTPNPNQCVNGPNYFCSCCCGNSVCNSNWLTC
VTAGSVAPTIVSTTVPPADEGLKRSDIINIQDSWNTLKGKGF--GYETVGMVLVHRLFNDAPO
-TRYLFSQLS-LSSNESFTLEQMRNNSRVVYHANRVARAVGRLVDLI---ELP-TNFTDH
LVWLGQRHAYHG----VAPVNFDMGVPVLETIKVNLELPSD-SPTLSAWAKAYGVIKNG
IKDAIIATYAEG-----

>Ciona_savigny_Gb1_LGB3_(from_Hoffmann_et_al_2012)

-----MEMNAQEIQDVRDSWKRLCAD--GEKTVGLMLMQKLFNTYPE
-SIKVSRLG-ITNKAIITIDDLSTNSAASRHAESLTSRIGTLVDLM---HNT-HEFKEC
STEVGEIHIKYG----VTAEHVDILGNVLLSVICDSQGLSKS-SDLYLCWTKTWEGIAKY
VKIGLQQ-----

>MOLGULA_TECTIFORMIS_LGB3_gi|68107250|gb|CJ414980.1|CJ414980_CJ414980_Molgu
la_tectiformis_unpublished_cDNA_library,_larva_Molgula_tectiformis_cDNA_clo
ne_mtlv020d18_5',_mRNA_sequence_From_ESTs_stored_at_http://www.ncbi.nlm.nih
.gov/nucest/

-----MGGLTQEEIQAVRESWAAVIKK-VGVTETGLAVLHRLFADVPE
-TKTLFYRNIG-LDSYDTITMDQLKDNKKIRNHALRVVTSISNVLKNI---KN-GEKLAEL
FKKLGEIHKKNK----VPPEYFDKMGNVLLAVLTSVLEVDLC-HPTSLAWGKVYGILRDQ

>BOTRYLLUS_SCHLOSSERI_Cytoglobin2_gi|322520564|gb|JG298067.1|JG298067_CCAO1
145.b1_CCAO_Botryllus_schlosseri_total_asexual_and_embryonic_development_Bo
tryllus_schlosseri_cDNA_clone_CCAO1145_5',_mRNA_sequence_From_ESTs_stored_a
t_http://www.ncbi.nlm.nih.gov/nucest/

-----MEEKSLFTAEESDAIKSSWSKMTE--NGVSTAGRLILCRLFQDVPE
-VTTLFYRLG-ISGDVSYTLESLSQKSQFNHAKRLSLALDAIVKKL---DD-TAFITQK
CTDLGKAHEKHN----VKPHYFDLLGQVLVKVICQALNLTES-HPTVKAWIKAYGAISTG
LKNGLTSELYLS-----

>CEPHALOCHORDA

>Branchiostoma_floridae_GB1_LGB3_XP_002608549.1_hypothetical_protein
BRAFLDRAFT_98913

-----MGAFLTKPFSLVGRLL
WKVLFSSWVVKQIETPSDVTGLTPTQSRLVKESWKMFLS---KKRENGFVIFRVLFTDYPV
-TRKLFKGVQ--LDLDAPGQLESSITLRAHVTRFMHSFDTYMESESL---DDP-EDLKQL
LYDTGKSHLIHD----IKPEYFDVLETVLMKSLRIVFGSKLT-PQLEEAWQTAYSHLKV

```

IKQGLEDAIQKRDQADTSVVVTVE-----
-----
>Branchiostoma floridae Gb14 LGB2 XP_002610160.1 hypothetical protein
BRAFLDRAFT_77082
-----MGANMGCSNSKKMSHESESAN
SGDSTPPKSSTPSALDERLPLTQKQKFLLLKSWKGVAR---QISQCGKTM LIRLFKDDPQ
-LMAVFNQKF-R-HLRERDADVLYQDAILDAAHAATVMEALHEAITHL---DDS-VFVMKV
LHDVVGKMHQRYN----VDPSVFLKVEKPFILTAVSEVLGDRYT-KNMEEIYTITIKFILAT
LSEGATMELTEDEQKNLGRLLWRPPGRVHKFVRPEKVAAIVDAQSEENGVH-----
-----
>Branchiostoma floridae Gb15 HGB related GenBank: CBL51564.1
-----
-----MGLTSEDKSAVLDSWAKMSGP--TFQDAGEKVFLLLLKTDS-
-TKALFPKFR-D-----IPYDQLAGHPDVRDHGGKVMQVLDDFIKGL---DNG---GDGA
VQKVGLLHKGVG----VSHDNINLMKPVLMTLLGELGCSSA-----AGAWENLWARFMDV
HRTCY-----
-----
>Branchiostoma floridae Gb4 LGB3 XP_002589215.1 hypothetical protein
BRAFLDRAFT_74626 XP_002589215.1
-----MGTIADGE
GTELNGYGGEKEPPGGGHGGPLTQEQVHGIKETWAILAQ---DPVERGVDLFMKIFEEEDPD
-LKKLFYFAD-D-----GRELSREDQRMRSHERVMEAVGGAVDSL---GDL-TAVVPV
LTELGALHHKYG----VQPSYFDTVGAALIYILETNLGDKLT-PNIRQGWVLVYGIVGAT
MKKGMQQAMDHQNMAKTRP-----
-----
>HEMICHORDATA

>Saccoglossus kowalevskii Gb1 NGB2 neuroglobin-like protein NP_001161601.1
-----
-----MADPVTTLTSDEVAAIKSSWSAVYD---KKKESGVTLFVKLF TENPS
-FKSQFGYMS-G-V----ADGDMKTLPALENHGVKVM DRIN EWGNL----TNGAELVKQ
LKHLGTTHIALK----VTEDNFNAMDSVLMYTLQEQGGS AFT-PAAKAAWQKAWGMKSV
IVGALKG-----
-----
>Saccoglossus kowalevskii Gb12 LGB3 Saccoglossus kowalevskii scaffold 38407
in Acorn worm assembly (see Hoffmann et al 2012)
-----M
GCTPSINERDFQTPVDDKHLDDRQKRIVRKTWRPLAN---DMTENGQKIFINIFESHPE
-IKYMFPTRD-I-E----GRDNL SANPHFRMHSSRFMQSVGAAIDNL---NDLDNALRPL
LVKLA KTHVRFK---GFKPDYFDAFEEAMLSVWQEELGQRFT-TEVEESWKLLFFYIKDC
LKEGYDIAMNEKTS GELNNSDFINQ-----
-----
>Saccoglossus kowalevskii GB15 LGB related scaffold 38908 in Acorn worm
assembly (see Hoffmann et al 2012)
-----MGCTVSTKPDLYSQ
NGDADPYKKKKVAPVDSRLPLTARQKFQITKSWKGIAR---NMENTGKSMFMRLFQSNIE
-LKNMFTGFE-E-FD---DLED MRESQQL ENHASLVMYTIDEAIASI---DDI-DFVV EL
LGKIGRTHTRTD----FNPQLFWRIEQPFLSAVKETLED RYT-KNIEE IYKITFRFIVDA
LIDGVVAGVNERKAEAEAEAEAKAEAEDEQDNEMMKKTMHNGDKEDNCIESSKQKDITVI
ASTKQR-----
-----
>Saccoglossus kowalevskii Gb16 LGB 3 scaffold 38908 in Acorn worm assembly
(see Hoffmann et al 2012)
-----
-----MGCLISSNGEHPMITKEQTKILTSTWHSIHG---DLEKIGLLMFMGMFDNYPE
-TRQFFGLSG-G-SI---VLEDPAVIQKIREHGLRFMTTARKLVMNL---DDK-DKFDRI
LLDLGRRHHGYK----ADVDLIEVFGQQFIASIQPTLKD NWN-PAVGEAWEQLFKCVSSR
MKDGF LQAQSSPSNTELLK-----
-----
>BALANOGLLOSSUS_CLAVIGERUS_Cytoglobin2_gi|311137357|gb|FN985678.1|FN985678_F
N985678_dmp027_Balanoglossus_clavigerus_cDNA_clone_dmp027P0033D09_mRNA_seq
uence_From_ESTs_stored_at_http://www.ncbi.nlm.nih.gov/nucest/

```

-----MG
 SIIGYFWASSDGNVDPGTTLTPNERRAIRDTWNAIYE---NKGENGVAFFMKLFEAHPV
 -YKKLFKNLE-G-ID---DMAKLSKHPRMKAHGLRVMASFNSEIENL---DD-AEVLVQL
 LVDIGISHSKHK----VTEEQFNVSNHFLSYFLRSLFLFAFI-YLAIPFPNFVVVLIYLF
 VRLSLLSVYLSV-----

>ECHINODERMATA

>STRONGYLOCENTROTUS_PURPURATUS_1_Ng3_Strpur_D1_(147-305)

-----LDERQKTLVENTWKTLEK---NTELYGSIMFAKLTTDHPD
 -IGKLFPPGG-K--NL--TYGELLVDPDVRVHGKRVIETLGSVVEDL---DD-MELVIQI
 LEDLGQRHNAYN----AKKTHIIAVGGALLFTIEEALGAGFT-PEVKAAWAAVYNIVSDT
 MSTKL-----

>KINORHYNCHA

>ECHINODERES_HORNI_Globin_NeuroBlast_KINORHYNCHA_GNL|TI|1987172666_EH_MM1_21H06 FROM TRANSCRIPTOME AVAILABLE AT FTP://FTP.NCBI.NLM.NIH.GOV/PUB/TRACE DB/

-----MGCG
 QSLESQGDGGAADIDSNKCPDRRQIFELTKSWRMISK---NMESAGVMMFVKLFTTTNE
 -INHMFALI-T--EDG-KVRNSEAQKQLEHHAAMVMTLDESIQAF---TD-LDVLVFL
 TTRIGQQHRKMDG---FSANYFWKIEDPFLTAVKQTLDDKYA-AELGHIYRRPIRFILKL
 LYNGYMKKD-----

>ENTOPROCTA

>PEDICELLINA_SP_LGB1_ENTOPROCT_GNL|TI|1987415660_PC_MM1_28H03 FROM TRANSCRIPTOME AVAILABLE AT FTP://FTP.NCBI.NLM.NIH.GOV/PUB/TRACEDB/

-----IRDRLSLDSNEMMAMQLMWNDWVG--GDAARGFDSFILMFKQHPE
 -TQTLDFDAR-G--C---SAAQMNSSRLLFHVTRVIHNITDVMDDL---HE-LEEVVVK
 LRQLGGRHGTPGTCYNVPGAYFPWLGDMKELMQGKGGWG---DSHNKAWAFVWSFICNS
 LTSGQAIYGGGSCPAKKA-----

>TARDIGRADA

>RICHTERSIUS_CORONIFER_HEMOGLOBIN_RELATED_TARDIGRADA_GNL|TI|1987149271_RC_MM1_21B09 FROM TRANSCRIPTOME AVAILABLE AT FTP://FTP.NCBI.NLM.NIH.GOV/PUB/TRACEDB/

-----MG
 GRASRILSSGSANVPDKVTGLSDVEKRAIQENWRIVYR---DLKNGVELFVRYFVSRYPE
 -YKDAFESLR-D--I---RMQDIGKSHKLAHSVQVMQYGTKPRKAR---YA-AGTFYGE
 PLNL---HTTTVDF-----LAFKLGHNFGHVHKTAWNKMLELVMV
 IQQGMTAEADDSKYKENQTA AVRQSEHEELTLTNLTETT-----

TABLE S2B

Multiple alignments with emphasis on informative sites selected for the molecular phylogeny analysis (underlined in yellow)

	10	20	30	40	50	60	70
VERTEBRATE GlobinX
ANOLIS CAROLINENSIS GMGC	ALSGAQDPFV	SEECSPPLDDG	LDLNRETTLG
CALLORHINCHUS MILII G	----MGCAIS	GPGQYPASGR
DANIO RERIO GbX (from	----MGCAISG	SGLTAGAPEI
ORYZIAS LATIPES GbX (.....	----MGCAIS	GLAAKTDLAE
PETROMIZON MARINUS Gb	----MGCTVSTD	ERTGAQSSSQ
PYTHON MORULUS GbX (f
TETRAODON NIGROVIRIDI	----MG	CAISSLGAKA
XENOPUS TROPICALIS Gb	----MGCILSSLG	WQWRDSL DHT
VERTEBRATE MYOGLOBINS
HOMO_SAPIENS_MYOGLOBI
DANIO_RERIO_MYOGLOBIN
BOS_TAUROS_MYOGLOBIN
GALLUS_GALLUS_MYOGLOB
VERTEBRATE CYTOGLOBIN
DANIO_RERIO_CYTOGLOBI
OVIS_ARIES_CYTOGLOBIN
HOMO_SAPIENS_CYTOGLOB
VERTEBRATE HEMOGLOBIN
EQUUS_CABALLUS_HB_ZET
HOMO_SAPIENS_HB_BETA
TAENIOPYGIA_GUTTATA_H
ORYCTOLAGUS_CUNICULUS
VERTEBRATE NEUROGLOBI
HOMO_SAPIENS_gi 10864
DANIO_RERIO_gi_188590
GALLUS_GALLUS_gi_1548
MONODELPHIS_DOMESTICA
CHIONODRACO_MYERSI_gi
BOVICTUS_VARIEGATUS_
SIPUNCULA
THEMISTE_1_LGB2_gn1 t
THEMISTE_2_LGB2_gn1 t
THEMISTE_3_LGB2_gn1 t
BRACHIOPODA
TEREBRETTALIA_Ng2_gn1
ANNELIDA
ALVINELLA_POMPEJANA_1
ALVINELLA_POMPEJANA_2
APHRODITA_ACULEATA_LG
CAPITELLA_SP_1_LGB2_F
CAPITELLA_SP_2_LGB2_F
CAPITELLA_SP_3_LGB3_g
PLATHYNEREIS_dumerili
HELLOBDELLA_ROBUSTA1_
ECHIURA
URECHIS_CAUPO_Ng1_gn1
PLATYHELMINTHES
MACROSTOMUM_LIGNANO_1
MACROSTOMUM_LIGNANO_2
CHOANOFLLAGELLATE
MONOSIGA_BREVICOLLIS
SALPINGOECA_SP_Ng2_gi
PLACOOZA
TRICHOPLAX_ADHAERENS_
TRICHOPLAX_ADHAERENS_
TRICHOPLAX_ADHAERENS_
TRICHOPLAX_ADHAERENS_
PORIFERA
CARTERIOSPONGIA_FOLIA
AMPHIMEDON_QUEENSLAND
CNIDARIA
NEMATOSTELLA_VECTENSI
NEMATOSTELLA_VECTENSI
NEMATOSTELLA_VECTENSI
NEMATOSTELLA_VECTENSI

DANIO RERIO GbX (from	RPGEETPAG	LTTNHIRLIK	ESWRLIQE	--	DIAKVGIIM	FVRLFETHPE	-CKDVFFLFR	-D--VE--DL
ORYZIAS LATIPES GbX (RSREDAAVEH	PNEEQIQMIK	DSWKVIRD	--	DIAKVGIIM	FVRLFETHPE	-CKDVFFLFR	-D--VE--DL
PETROMIZON MARINUS Gb	GHQPPGPPQA	PSESQRRLVR	DSWLALQC	--	DIARVGVIM	FVRLFETHPE	-CKDVFFYQFR	-D--CE--DL
PYTHON MORULUS GbX (f	-----	---QKELIR	ESWKILHK	--	NITRVGIIV	FIRLFETHPE	-CKDVFFLFR	-D--ID--DL
TETRAODON NIGROVIRIDI	DAAAAAVVY	PREDQIQMIK	DSWKVIRD	--	DIAKVGIIM	FVRLFETHPE	-CKDVFFLFR	-D--VE--DL
XENOPUS TROPICALIS Gb	ETSPLLPTLN	LSEQQQQLLV	ESWRLIQH	--	DIAKVGVIL	FVRLFETHPE	-CKDVFFLFR	-D--VD--DL
VERTEBRATE MYOGLOBINS
HOMO SAPIENS MYOGLOBI	-----MG	LSDGEWQLVL	NVWGKVEA	--	DIPGHGQEV	LIRLFKGHPE	-TLEKFDKFK	-H--LK--SE
DANIO RERIO MYOGLOBIN	-----	--MADHDLVL	KCWGAVEA	--	DYAANGGEV	LNRLFKEYPD	-TLKLFPKFS	-G--IS--Q
BOS TAURUS MYOGLOBIN	-----MG	LSDGEWQLVL	NAWGKVEA	--	DVAGHGQEV	LIRLFTGHPE	-TLEKFDKFK	-H--LK--TE
GALLUS GALLUS MYOGLOB	-----MG	LSDQEWQQVL	TIWGKVEA	--	DIAGHGHEV	LMRLFHDHPE	-TLDRFDKFK	-G--LK--TP
VERTEBRATE CYTOGLOBIN
DANIO RERIO CYTOGLOBI	GVQLTQSPDS	LTEEDVCVIQ	DTWKPVYA	--	ERDNAGVAV	LVRFFTNFPS	-AKQYFEHFR	-E--LQ--DP
OVIS ARIES CYTOGLOBIN	EIERERSEE	LSEAERKAVQ	ATWARLYA	--	NCEDVGVAI	LVRFFVNFPS	-AKQYFSQFK	-H--ME--EP
HOMO SAPIENS CYTOGLOB	EIERERSEE	LSEAERKAVQ	AMWARLYA	--	SCEDVGVAI	LVRFFVNFPS	-AKQYFSQFK	-H--ME--DP
VERTEBRATE HEMOGLOBIN
EQUUS CABALLUS HB ZET	-----MS	LTKAERTMVV	SIWGKISM	--	QADAVGTEA	LQRLFSSYPQ	-TKYFFPHF-	-----
HOMO SAPIENS HB BETA	-----MVH	LTPEEKSAVT	ALWGKVV	--	NVDEVGGEA	LGRLLVVYPW	-TQRFESFG	-D--LS--TP
TAENIOPTYGIA GUTTATA H	-----MV	LSAGDKSNVK	AVFGKIGG	--	QADEYGADA	LERMFATYPQ	-TKYFFPHF-	-----
ORYCTOLAGUS CUNICULUS	-----MA	LSAAERALLR	ALWKKLGS	--	NVGVYATEA	LERTLEAFPR	-TKIYFSHM-	-----
VERTEBRATE NEUROGLOBI
HOMO SAPIENS gi11864	---P--ELIR	QSWRAVSR	---	SPLEHGTVL	FARLFALEPD	-LLPLFQYNC	-R--QFS-SP
DANIO RERIO gi188590	-----K	LSEKDKGLIR	DSWESLKG	--	NKVPHGIVL	FTRLFELDPA	-LLTLFYSYST	-N--CG--DA
GALLUS GALLUS gi1548	-----M	LSRTQQALIR	ESWRRVSG	--	SPVQHGIVL	FSRLFELDPD	-LLPLFQYNC	-K--RFA-SP
MONODELPHIS DOMESTICA	-----RR	LSGPEQELIR	ESWQKVNS	--	NPLQHGMLL	FTRLFLEDPD	-LLPLFQYNC	-R--QFS-SP
CHIONODRACO MYERSI gi	-----K	LSGKDKELIR	GSWESLKG	--	NKVPHGVVM	FSRLFELDPE	-LLTLFHYTT	-N--CG--ST
BOVICHOTUS VARIEGATUS	-----K	LSGKDKELIR	GSWESLKG	--	NKVPHGVVM	FSRLFELDPE	-LLTLFHYTT	-N--CG--ST
SIPUNCULA
THEMISTE_1_LGB2_gnl t	MNQAPEDQTK	FTPDELMLAL	LSWNAWVA	--	GDLAGXGFD	FVKMFEQRKE	-TKEVFPFAR	-G--N--DA
THEMISTE_2_LGB2_gnl t	-----	---LR	LSWNAWVA	--	GDLVGRGFD	FVKMFEQRKE	-TKEVFPFAR	-G--N--DA
THEMISTE_3_LGB2_gnl t	MNLDLEDQCR	FDPEELINLR	LSWNAWVA	--	GDLPRGFD	FTKMFEQRKE	-TKEVFSFVR	-G--S--DA
BRACHIOPODA
TEREBRETALIA Ng2_gnl	-----	FTTQEKRLR	SSWRAFVA	--	EDPAKRGYQ	FMKMFENRPD	-TKKFFK-FV	-Q--KD--AL
ANNELIDA
ALVINELLA POMPEJANA_1	-----	-----	---YRLS	---	DISERGMID	FVRLFELHPV	-YKSYFQKLR	-D--V--DI
ALVINELLA POMPEJANA_2	ASYKPDPRCP	LTERQLYSIT	KSWKAINR	--	EMASTAVNM	FIRLLEHDGI	--RSFFTKFK	-D--HK--TV
APHRODITA ACULEATA LG	-----g	lsgaDIAVIR	STWAKVQG	-s	gSATDIGRSI	FIKFFELDPA	-AQNEFPCKG	-E-----SL
CAPITELLA SP_1_LGB2_F	-----DQL	LTPEEIVLVR	VTWEQLKT	-n	lTLANLGKV	FLRIFNLKPD	-IKKLFPPSD	-V-----WG
CAPITELLA SP_2_LGB2_F	kpsnipqHEF	LTQNKQLGAK	STWEFLHC	-t	sTPTERGMRV	FLRIFEIAPV	-TKTLFPFKD	-M-----PN
CAPITELLA SP_3_LGB3_g	-----L	FTDRQKAIIT	KTRWHMGN	--	DLTGRGSKV	FLKIFNLHPE	-VKQYFPSLK	-N--D--NE
PLATHYNEREIS dumerili	-----P	LNDNQKELIK	KLWEIVRE	--	GDIESTGVSM	YMHMFTLQPE	-VMKMFSEVP	-K--NVT-NP
HELLOBDELLA ROBUSTA1	NLNYLNNDV	LTIREKQLVR	ESWTLLSI	--	KLKSLGKQV	FLRIFELRPS	-TKNLFPFKT	-V-----WG
ECHIURA
URECHIS CAUPO Ng1_gnl	-----	VSLQNIALLV	KSWAILKK	--	DLLSNGAAL	FLALFEAYPD	-YKDLFKQFP	-G--L--RL
PLATYHELMINTHES
MACROSTOMUM LIGNANO_1	-----M	LTADETQLIL	SGWNQAMK	--	DAKGLGLDI	FLTLFEMFPQ	-HQELFRDFK	-G--K--SR
MACROSTOMUM LIGNANO_2	-----M	LNEVEKKIIL	SGWQQAIK	--	DKKALGMDV	FMTLFEMFPQ	-HQELFRDFK	-G--K--SR
CHOANOFLAGELLATE
MONOSIGA BREVICOLLIS	DEAHAPDYP	FDPVAVKLAR	KQWKRVOQ	--	LVPNWHEVF	FSYLFERAPY	-ARTLFPFD-	-----V
SALPINGOECA SP_Ng2_gi	-----MR	LDMEQKIAL	GSWTAVVE	--	LVPTWHEVF	FAELFOAHPE	-TERLLYSS-	-----
PLACOZOA
TRICHOPLAX ADHAERENS	DSVQTPPQPS	LTEEQKAIIR	ENWQDVEE	--	NMSEVGLYL	FSKLFETIAP	-YREVFPEE-	-----
TRICHOPLAX ADHAERENS	-----MA	PTAQDLQTR	ETWALVAP	--	DLKHHGTVL	FLRLFQHPD	-VQRLFEEKI	-D--V--PH
TRICHOPLAX ADHAERENS	-----	-----	-----	---	---MVLVN	NYSLIKLSPA	-TKIYFHGVD	-F--EK--RD
TRICHOPLAX ADHAERENS	VVTGNDIKSY	LNQYERQAI	DSWNAIST	--	EKQKYGTIL	FLKLFELRPS	-VKSLFTTFD	-F--NE--PL
PORIFERA
CARTERIOSPONGIA FOLIA	-----MSHP	LSSEELKIVE	KSWALIQE	--	FGLQEAGVM	FKRLFEMSPG	-LQNLFFPA-	-----GD
AMPHIMEDON QUEENSLAND	-----S	LTSAQVALIE	STWKVVKK	--	DLQGAGNIM	FLKLFQIDVS	-VRDKFPFRD	-V-----PY
CNIDARIA
NEMATOSTELLA VECTENSI	-TENTILTSVP	LSTRRKKLVR	ESWELIEP	--	VKITIGKRL	FTRLFDVNP	-MQDTFPNFK	-G--K--EL
NEMATOSTELLA VECTENSI	QLERINPTG	LSAREVAVVK	QTWNLVVK	--	DLMGVMRI	FKSLFEAFPA	-YQAVFPKFS	-D--V--PL
NEMATOSTELLA VECTENSI	----NPQNA	FSAADIQAIQ	GTWALAK	--	PDLMGKAMV	FKQLFTEHGY	--QPLFSNLA	-Q--Y--EI
NEMATOSTELLA VECTENSI	-----	-----	-----	---	-----	-----	---CFKAFN	-K--V--SL
NEMATOSTELLA VECTENSI	KPPREPVKIP	LSVAQKYLVR	ETWETIEQ	--	HSKAVGKKT	FLRFFEMNPD	-YQKLFPEFA	-T--L--DQ
NEMATOSTELLA VECTENSI	NQSFVGNRPR	LTERQIKLVQ	DTWRLLIP	--	SQKKTAMIF	YKLFETLDP	-FKEVFS-FH	-T--E--NE
NEMATOSTELLA VECTENSI	DTENTILTSVP	LSTRRKKLVR	ESWELIEP	--	VKITIGKRL	FTRLFDVNP	-MQDTFPNFK	-G--K--EL
NEMATOSTELLA VECTENSI	QYLTQVDQLP	LTETQKYIYK	QSMWGLS	--	NKGELGIEI	FLRLFENPT	-LQLMPEFR	-E--YS--TL
NEMATOSTELLA VECTENSI	EKLFETSRIE	LDAKETQLVR	KTWAILGD	--	RQVEVGKSL	FLRFFEEHPT	-SKDLFPEFR	-N--I--SN
NEMATOSTELLA VECTENSI	HLCGLKLECD	MTYEQKYLVR	ETWKFLEV	--	SKKEIGVSV	YKRLFNHHPG	-LQYFSEFK	-----H
NEMATOSTELLA VECTENSI	KPPREPVKIP	LSVAQKYLVR	ETWETIEQ	--	HSKAVGKKT	FLRFFEMNPD	-YQKLFPEFA	-T--L--DQ
NEMATOSTELLA VECTENSI	SQANLPRTMP	LSEAQKYLVR	ETWETIEP	--	QKQTVGKKA	FLRFFDMNPD	-YQNLPEFK	-S--L--SY
CLYTIA HEMISPHERICA	IQKSSNIKVP	LSAKEVNILK	KTWPPVKN	--	NWLKICLVA	FERWFTLYPQ	-LRYMFRSLP	-E--DI--QF
CLYTIA HEMISPHERICA	NGKVDAAPAD	YTEAEINIVQ	KQWSVAMR	--	NLDSVGYKL	FAKLFQYKDI	MAKPDFAVAN	-N--L--DF
CLAVA_1_LGB1_Locus_11	HTKYGNICIP	LTKKDIKKLR	KSWGIMKM	--	NWYKICLVV	FDRWFSMYPE	-LRNMFKFS	-S--VL--TL
CLAVA_2_Ng_Locus_2853	-----	---EVGILLR	KQWSIMMS	--	DADANGIML	FTALFAINGH	-LRTKFAFGD	-V--YGF-DC
HYDRA MAGNIPAPILLATA	-----	LSGKEIETLK	KSWTTAKQ	--	FWNEICTCA	FSRWFSSTYPE	-IQSKFGVYG	-D--NL--TM

OVIS_ARIES_CYTOGLOBIN	LEMERSPQLR KHACRVMGAL NTVVENI	HD-PEKVSSV LALVGKAHAL KHK	VEPV YFKILSGVIL
HOMO_SAPIENS_CYTOGLOBIN	LEMERSPQLR KHACRVMGAL NTVVENI	HD-PDKVSSV LALVGKAHAL KHK	VEPV YFKILSGVIL
VERTEBRATE_HEMOGLOBIN
EQUUS_CABALLUS_HB_ZET	DLHEGSPQLR AHGSKVAAAV GDAVXSI	--DNVAGA LAKLSELHAY ILR	VDPV NFKFLSHCLL
HOMO_SAPIENS_HB_BETA	DAVMGNPKVK AHGKVKLGAF SDGLAHL	--DNLKGK FATLSELHCD KLH	VDPE NFRLLGNVIV
TAENIOPIYGIA_GUTTATA_H	DLGKGSQAQVK GHGKVKAAAL VEAANNV	--DDLAGA LSKLSDLHAQ KLR	VDPV NFKLLGQCFL
ORYCTOLAGUS_CUNICULUS	DLSPGSAQVR AHGRKVDAL TLAADHL	--DDLPGA LSALSDLHVR TLR	VDPH HFGLLGHCLL
VERTEBRATE_NEUROGLOBIN
HOMO_SAPIENS_gi 10864	EDCLSSPEFL DHIRKVMLVI DAAVTNV	ED-LSSLEEY LASLGRKHRA VG	VKLS SFSTVGSLL
DANIO_RERIO_gi_188590	PECLSSPEFL EHVTKVMLVI DAAVSHI	DD-LHTLEDL LNLGRKHQA VG	VNTQ SFALVGSLL
GALLUS_GALLUS_gi_1548	QECLAAPEFL DHIRKVMLVI DAAVSHI	ED-LPCLEEY LCNLGKHHQA VG	VKVE SFSTVGSLL
MONODELPHIS_DOMESTICA	QDCLSSPEFL DHIRKVMLVI DAAVTHV	EN-LSSLEEY LTNLGGKHHQA VG	VKLS SFSTVGSLL
CHIONODRACO_MYERSI_gi	QDCLSSPEFL EHVTKVMLVI DAAVSHI	DD-LPSLEDF LNLGRKHQA VG	VNTQ SFAEVGSLL
BOVICHTUS_VARIEGATUS_SIPUNCULA	QDCLSSPEFL EHVTKVMLVI DAAVSNL	DD-LPSLEDF LNLGGKHHQA VG	VNTQ SFAEVGSLL
THEMISTE_1_LGB2_gnl t	XQMHQIINVL FHVTKFMKNI DEVVKNA	DR-LEDVVSM LRRVEGRHGH QGHN	VPSA YFPFLGAAMH
THEMISTE_2_LGB2_gnl t	KQMQQSSNVL FHVTRVMKNI DEVVKHA	DR-LEDVVSM LRQVGRHGH QGHN	VPSA YFPYLGAMR
THEMISTE_3_LGB2_gnl t	HQMQQSSKFL FHVTRVMKNI DEVVKHA	DR-LEDVVSM LRQVGRHGH SHHN	VPSA YFPFLGIALR
BRACHIOPODA
TEREBRENTALIA_Ng2_gnl	GTMKSSTSLV FHTGRVAKYI QLVCDCL	DD-PTEAVPL LRQLGGKHHG GPNNFN	VDAT YFPELGKAMR
ANNELIDA
ALVINELLA_POMPEJANA_1	EDLRQSGKLR VHSTSVMKSI TDIVETI	DH-PPDLRDM AIKIAHPHFD RG	VRPS QYRELFAILL
ALVINELLA_POMPEJANA_2	AELLRASKVEF SHALMVISVI DDVITNL	DD-MDYVMSL LQATGESHSI KEKN	VNPE FLWNVEGAFI
APHRODITA_ACULEATA_LG	AALKTNVLLG QHGAKFMEYI TTAVNGI	DD-YAGRAHP LTELGRHKT RG	TTPA NFGKAGEALI
CAPITELLA_SP_1_LGB2_F	DDLIRHPKEV LHSERFMLVV DCCVQNL	ECIKSEHGEM LANLGRAHVN YK	FSRE NFEVFMKAIW
CAPITELLA_SP_2_LGB2_F	EDLHRNSLFK GHATRFMKSV EFTMQNL	DALDVIIVNPT LVSIGNKHVH IK	FHPD YLDTFQALM
CAPITELLA_SP_3_LGB3_g	DQLLKNPCFR GHASRFMQSV GAVVENL	DS-PGDLSP LIDLGRKHVL FGG	FATPE YFAAFTEGMM
PLATHYNEREIS_dumerili	EDLKN SARFL RHARNLMTTV TTAVDML	GD-MEGLSEV LVDLGRHKK YR	AKTE HFPYVGRSLT
HELLOBDELLA_ROBUSTA1_ECHIURA	DKLIKHPFL THSKRFVKVI GCVVDRI	DYLOECAAQF LIELGKHHVS IEG	FLPD YDVIYIRAI
URECHIS_CAUPO_Ng1_gnl	EDLTRMPPLR ALGATFMHSL GSMVDNL	GD-LECVVEL LRERTHTHWE RG	IRFE HFQNVFDLLP
PLATYHELMINTHES
MACROSTOMUM_LIGNANO_1	AELKMPKLR AHGLRVVNTL DGAIQSI	DD-MEVCVSS LELIGASHKG RN	MNAG HFEDLNKALD
MACROSTOMUM_LIGNANO_2	AELKMPKMR AHALRVVNTL DGAIQSI	DD-MEVCASS LELIGASHKS HH	LSAK HFEDLNAALA
CHOANOFLAGELLATE
MONOSIGA_BREVICOLLIS	DRLOGNSSLA EHAKRVGQAL ETALQGL	FE-YYSLVEV LEKLGRRHFK YG	VEPE HIDLFEEFTY
SALPINGOECA_SP_Ng2_gi	--DKSKSWNE RHMARVGKSV GDVIKSI	SN-YDDVIEH LTALGTRHAR YG	LHVD QLDLFINAFL
PLACOZOA
TRICHOPLAX_ADHAERENS	-TTDNVRLR VHATGVMTV GKAVQNI	DQ-FSELQSA LSTLQGFHHR KA	IKFE NFOAVGQALI
TRICHOPLAX_ADHAERENS	QDLATNENFV FHTTRVMTI DHAVKGI	DN-LPALTVL LKQLGSSHAQ YN	VKKE YFKIGLRISE
TRICHOPLAX_ADHAERENS	SYLAKNTFLR NHAARFMEAI NVIIGQD	MD-IFSVESY FRVVGSKHHS YN	LKLE HVQDISDAFL
TRICHOPLAX_ADHAERENS	EDIIQSPHFR SHAMRFMQSL ETGVLMG	FD-KESCDPL FKSLGSRHFF YD	LKSE FLDVIPICIL
PORIFERA
CARTERIOSPONGIA_FOLIA	SLSDNEGMK HHSFIVMTSL DDCLKIV	HD-VPKLDE LISLGAHVHI QG	VTSE HFAPVGSALI
AMPHIMEDON_QUEENSLAND	EELEDESFL KHSLQVMTI DLAITLLG	GE-MEKLVEA LVDLGMAMAH QG	LKPE DFDHVGREALV
CNIDARIA
NEMATOSTELLA_VECTENSI	KDILNRSRLY LHAKRVMVAV ENAVTVI	DD-AETFESY LINLGRHLP WG	VTKD HFGVVGGAFI
NEMATOSTELLA_VECTENSI	KDLEDTPAVG KHAI SVTKL DELIQT	DE-PANLALL ARQLGEDHIV IK	VNKP MFKSFGKVLV
NEMATOSTELLA_VECTENSI	TGLEGSPELN THARNVMAQL DTLVGS	QN-SIELGQS LAQLGKDHVP RK	VNVR HFKDFAEHFI
NEMATOSTELLA_VECTENSI	EDLEKSPFL AHATSVMSAI NEVVCM	DE-VEILGIL LEKIGFSHAR RE	IRRI HFENLAKVVV
NEMATOSTELLA_VECTENSI	VELEQANALH GHAERVMAK ENAVSAM	DD-AESFAAY LENLGARHKA RA	LKPA YLDAMQVAYT
NEMATOSTELLA_VECTENSI	GQLEQDERFL FQSRKFMEMI NSAVDRI	ND-ISLVMI LKSLGEVHWT KF	IKPE YFEPVKGALI
NEMATOSTELLA_VECTENSI	KDILNRSRLY LHAKRVMVAV ENAVTVI	DD-AETFESY LINLGRHLP WG	VTKD HFGVVGGAFI
NEMATOSTELLA_VECTENSI	EELKESRSLQ GHTKRVMKV ENAVNSL	ED-GHALMEY LQELGRRHKT RQ	IKPTVS NLQEISQAIN
NEMATOSTELLA_VECTENSI	EKIAESPALY GHARRVMKV DNAVASI	EN-VQVYSAY LYELGTRHQT RQ	LSEE QLKFMGGAFL
NEMATOSTELLA_VECTENSI	IKIDNINGSH GHPRRLMAI DNAVATL	GD-SDSFSAY LVELGRRHGH MFRP	PTHF NDLRKCFLSV
NEMATOSTELLA_VECTENSI	VELEQANALH GHAKRVMKAV ENAVSAM	DD-AESFAAY LENLGARHKA RA	LKPA YLDAMQVAYT
NEMATOSTELLA_VECTENSI	EELQKANALH GHAKRVMKAV ENAVMSI	DD-VMSFSAY LVELGRRHKT RA	LKPS YLEAMHGALM
CLYTIA_HEMISPHERICA	EDLFKTDALK MHVDKVRDVL ELLIKKI	DN-VEELVNT LVDFGRQHMM LG	AEQR YATALAASFQ
CLYTIA_HEMISPHERICA	QKSMNDARLS FHIRRVFHTI NTVVSCI	ND-GDFVASQ LEHVGAIAE YG	LQAT HLARFKDVML
CLAVA_1_LGB1_Locus_11	QELLASRTMK SHIQLEELV EDLLYKV	DD-PTDFIET VIQFGEHKK LG	--AK QLYATSLAAA
CLAVA_2_Ng_Locus_2853	KTFGDERLL GHLRGVFSSV EMLINSLVQ	NT-FYESLGK IKELGKVHSL AG	IHLS DLQAFKKAMI
HYDRA_MAGNIPAPILLATA	NEVLASESLC IHIRKSVELI EIIIKKV	DE-RHELSEY LIELGKLHKK FG	AEQK YATALGSSFV
MONTASTRAEA_FAVEOLATA	DAMRTDDRFK RQGLVTMQHV DLAVASI	SD-LGSIVPA LKDLGARHSM YK	VEEH HFGPVG---
MONTASTRAEA_FAVEOLATA	--DFTKPOFT THALNIMNAL DHAIQNL	DN-PDVLIPK LRELQGMHAG FE	LTIK EFQHVGEALI
ACOELA
SYMSAGITTIFERA_ROSCOF	SSLNSNVMHL AHSLHIMMI GKLIQLL	NE-PENLLSK VVELGERHFD RK	ANDE LLQYFCPAY-
SYMSAGITTIFERA_ROSCOF	KEMRSHPRFK SHAKSIGKVI SDCLVDI	NG-VKKHEPK LSSLGAMHTK KK	VPTL LFGKLGCCIL
NEMERTODERMATIDA
MEARA_STICHOPI_Ng1_tr	KDLVPADIK AHGMRIMGLL GRFVSNLESN	VD-DNLLGQP IHDLGRKHVD FK	APPY LFDVFALQIQ
NEMERTODERMA_WESTBLAD	SELQNTQMK GHVLRVINYI TDIVDTL	EV-DEMREEM SINIGRTHGR RA	IPAE MFQCLKPAVF
NEMERTODERMA_WESTBLAD	DELRESKMLR AHVIRVMTL NSLVDGI	DD-PELFVEL SKFGRTHYR HN	IDTE HVLNLGEAVI
NEMERTODERMA_WESTBLAD	YELRENHME AHLTRGLSHI TCTINKV	AE-PTVCADM WRNLGRTHTA HT	VPFK FMDLMKPAVF
XENOTURBELLIDA
XENOTURBELLA_HEMOGLOBIN	DAVKADAYLQ THGNRVMSTL KTVIGSI	ED-AGKCEAL ITQLGNLHTG RN	VAAA DFEALFPIIA
MOLLUSCA

SEPIA OFFICINALIS_LGB	EEFNTKASFM AHAKNVFYAV TLIVDTL---	DD-ADELVEI LLKTGRNHRH RS---	VPLS PFQGRAIVFE
EUPRYMNA_SCOLOPES_1_L	EEFNTKPKFI AHAKNVFYAL ILVVDLS---	DE-PDELVEI LLKTGRDHHN RG---	VPMA AFHNLAIVFD
DORYTEUTHIS_PEALEII_L	DEMRENEALE KHATLVMTL DDAITNI---	EN-VDLVLDL LHRIGKSHLR FQG---	FNVE YFWLAEQPLI
EUPRYMNA_SCOLOPES_2_L	DQMRREALE KHATLVMTI DEVIANI---	EN-VDLVLDL LHRIGKSHCR FQG---	FNVE YFWLAEQPLI
MYTILUS_CALIFORNIANUS	IELRRSPIFK EHAHYRFMRV DDLVDNM---	DDPKAHIQQN LMLLGAKHAT FEG---	FRIE YFEAYSBSLI
MYTILUS_CALIFORNIANUS	YSTEADAVPK DLAVKVGKAI ATFAKHC---	DD-LDSQKEL FGGIAIKHVK LGG---	ILEE QYSAMEGSFL
SPISULA_SOLIDISSIMA_L	SEIKQSPKLR AHSINFCHGL NSFIQSI---	DE-PDVLVIL VQKLTVNHFR RK---	IAVD RFQELAFALYV
APLYSIA_1_LGB3_CNSN01	DELRNSNEALE YHATLVMTL DDAITHI---	DN-YDFVKQL LSKTGASHVK FAG---	FKST NFLAIKGPFL
APLYSIA_2_LGB2_CNSN01	EDVDVDDKMLQ KHAVTVMEGL GAAVETL---	ND-SQFLNTV LIALGQTHEK RN---	IKPN MLKRMWPSMH
LIOLOPHURA_JAPONICA_N	DEIKASDKIT NHGGKVINEL ANWINNI---	---NSASGI LKSQGTAAHK RS---	IGIE YFENVLPVID
CRASSOSTREA_GIGAS_2_L	EELRQNVGLR EHGMRVMGTI EKCITRI---	DQ-PDKLTSM LESLGEKHVV FD---	TKIE YFDLLSPOLI
CRASSOSTREA_GIGAS_1_L	NELLRHPPFI DHVTRFMQII DYLVENI---	DQONSDFHQA LMLLGAKHAT YPG---	FQVS HFTVFNKALL
ARTHROPODA
APIS_MELLIFERA_LGB2_g	NELPANKRFQ AHCAGVITAL NNVIDFT---	HD-PGLMEAS LIGLVERHKK RG---	QTKE EFQNLKEVML
CARCINUS_MAENAS_1_LGB	EELKNSKRLA AHGTVVMMAI TAMVDNI---	ED-VSVLVEL LKNTGANHRD RP---	VPKG DFELLAERVLV
CARCINUS_MAENAS_2_LGB	DEQAESLELA EHATIVMNSI DEGIKAM---	DN-VDFFFDL LHQIGASHRK IPG---	FKKE YFWKIEHPFL
DAPHNIA_PULEX_GlobinX	GVVLDLKKK KLLSEHALRV MGAVQRAVHR	LQPEKLHAF LSELGRKHEK NG---	AKLE YIDYIGQOFL
DAPHNIA_CARINATA_LGB2	GELLLNPALK TFSVNVLTKL SELIDCMG---	---NPDALQGQ LVDLANQHKQ RG---	TTRA HFDNLSKVLID
HARPEGNATHOS_SALTATOR	EELSKNKKFO AHCVSVIAGL SNVIDHI---	HN-PELMEAS LINLAERHKN RG---	QTRF HFQNLRYVLE
TUNICATA
Ciona_intestinalis_Gb	EQMRNNSRVV YHANRVARAV GRLVDLI---	EELP-TNFTDH LVWLQORHAY HG---	VAPV NFDYMGPVLL
Ciona_savigny_Gb1_LGB	DDLSTNSAAS RHAESLTSRI GTLVLDL---	HNT-HEFKEC STEVGEIHIK YG---	VTAE HVDILGNVLL
MOLGULA_TECTIFORMIS_L	DQLKDNKKIR NHALRVVTSI SNVLKNI---	KN-GEKLAEL FKKLGEIHKK NK---	YVKG YFDKMGNVLL
BOTRYLLUS_SCHLOSSERI	EELSKNSQFN SHAKRLSLAL DAVVKKI---	DD-TAFITQK CTDLGKAHEK HN---	VKPH YFDLLGQVLV
CEPHALOCHORDA
Branchiostoma florida	GQLESSITLR AHVTRFMHSF DTYMESI---	DDP-EDLKQL LYDTGKSHLI HD---	IKPE YFVLETVML
Branchiostoma florida	DVLYQDAILD AHAATVMEAL HEATHIL---	DDS-VFVMKV LHDVGKMHQR YN---	VPKG YFLKVEKPFPL
Branchiostoma florida	DQLAGHPDVR DHGGKVMQVL DDPITKGI---	DNG---GDGA VQKVGLLHKG VG---	VSHD NINLMKPVLM
Branchiostoma florida	ELSREDQMR SHGERVMEAV GGAVDSL---	GDL-TAVVPV LTELGAHHK YG---	VQPS YFDTVGAALI
HEMICHORDATA
Saccoglossus kowalevs	DMKTLPALE NHGKVMMDRI NEWMGNI---	---TNGAELVKQ LKHLGTTTHIA LK---	VTED NFNAMDSVLM
Saccoglossus kowalevs	NDLSANPHFR MHSSRFMQSV GAAIDNI---	NDDLNALRPL LVKLAKTHVR FK---	GFKPD YFDFAFEAML
Saccoglossus kowalevs	EDMRESQLE NHASLVMTI DEIASI---	DDI-DFVVEL LGKIGRTHTR TD---	FNPQ LFWRIEQPFL
Saccoglossus kowalevs	EDPAVIQKIR EHGLRFMTTA RKLVMNI---	DDK-DKFDRI LLDLGRRHG YK---	ADVD LIEVFGQQFI
BALANOGLOSSUS_CLAVIGE	AKLSKHPRMK AHGLRVMASF NSFIENT---	DD-AEVLVQL LVDIGISHSK HK---	VTEE QFNVSNHFLS
ECHINODERMATA
STRONGYLOCENTROTUS_PU	GELLVDPDVR VHGRVIETL GSVVEDI---	DD-MELVIQI LEDLGQRHNA YN---	AKKT HIIAVGGALL
KINORHYNCHA
ECHINODERES_HORNI_Glo	RNSEAQKQLE HHAAMVMYTL DESIQAF---	TD-LDVLPEL TTRIGQQHRK MDG---	FSAN YFWKIEDPFL
ENTOPECTA
PEDICELLINA_SP_LGB1_E	AQMNSRRL FHVTRVIHNI TDVMDHI---	HE-LEEVPK LRLGGRHGT NPGCYNVPGA	YFPWLGDMAM
TARDIGRADA
RICHTERSIUS_CORONIFER	QDIGKSHKLR AHSVQVMQYG TKPKRAR---	YA-AGTFYGE PLNL---HTT TVDF---	-----

VERTEBRATE_GlobinX
ANOLIS_CAROLINENSIS_G	QAAQPILKEA WT-PETEKAW EGLFQYLAAT MRRGFYKEQK ATGKN---	220	230	240	250	260	270
CALLORHINCHUS_MILII_G	CAVQPILKEK WT-AEVEEAW KGLFHYLTSV MKKGYQDEER GSCPREKPKH GPNSV---						
DANIO_RERIO_GbX (from	CAVRPILKDR WT-PELEEAW KTLFQYVTSI MREGFLEEBER NKRSNTQTS RERPDKRSTA I-----						
ORYZIAS_LATIPES_GbX (CAVQPILKEQ WT-TELEKAW QTLFQFVTAL MKQGYQEE-S ARQRQLA-TS PKDRLDKNT AL-----						
PETROMIZON_MARINUS_Gb	ATVQPFLQEK WT-NEVEDAW QCLFRYIAAV MKRGYLEEEA ASNGVNTANY DRGQGNHGAT AM-----						
PYTHON_MORULUS_GbX (f	QAVQPILKED WT-LEVEKAW K-----						
TETRAODON_NIGROVIRIDI	CAVQPILKER FT-SELEEAW KTLFQYVTGL MRKGHQEEGS RQRHLALPPK DGPEKRTSAL -----						
XENOPUS_TROPICALIS_Gb	SAVCPMLHDK WT-AEVEEAW KGLFAYICTV MERGYQEEB- RRHSDGRSLI DGLQGNKGLI -----						
VERTEBRATE_MYOLOBINS						
HOMO_SAPIENS_MYOGLABI	QVLSQKHPGD FG-ADAQAM NKALELFRKD MASNYKELGF QG-----						
DANIO_RERIO_MYOGLABIN	KVMAEKAGLD AA-GQALRR VMDAVIGDID GYKYEIGFAG -----						
BOS_TAURUS_MYOGLABIN	HVLHAKHPSD FG-ADAQAM SKALELFRND MAAQYKVLGF HG-----						
GALLUS_GALLUS_MYOGLAB	KVIAEKHAAD FG-ADSQAAM KKALELFRND MASKYKEFGF QG-----						
VERTEBRATE_CYTOGLOBIN						
DANIO_RERIO_CYTOGLOBI	EVLVEAFPQC FSPAQVQSSW SKLMGILYQW MNRVYAEVGV EN-----						
OVIS_ARIES_CYTOGLOBIN	EVIAEEFASD FP-PETQRAW AKLRGLIYSH VTAAYKEVGV VQ-----						
HOMO_SAPIENS_CYTOGLOB	EVVAAEFASD FP-PETQRAW AKLRGLIYSH VTAAYKEVGV VQ-----						
VERTEBRATE_HEMOGLOBIN						
EQUUS_CABALLUS_HB_ZET	VTLASRLPAD FT-ADAHAAM DKFLSIVSSV LTEKYR---						
HOMO_SAPIENS_HB_BETA	CVLAAHFGKE FT-PPVQAAQ QKVAVGANA LAHKYH---						
TAENIOPYGIA_GUTTATA_H	VVAVTRNPSL LT-PEVHASL DKFLCAVGTV LTAKYR---						
ORYZCTOLAGUS_CUNICULUS	VTLARHYPGD FG-PAMHASV DKFLHHVISA LTSKYR---						
VERTEBRATE_NEUROGLOBI						
HOMO_SAPIENS_gi110864	YMLEKCLGPA FT-PATRAAW SQLYGAVVQA MSRGWDG---						
DANIO_RERIO_gi_188590	YMLQSSLGPA YT-TSLRQAW LTMYSIVVSA MTRG-----						
GALLUS_GALLUS_gi_1548	YMLENCLGAA FS-PDVREAW IELYGAVVKA MQR-----						
MONODELPHIS_DOMESTICA	YMLEQCLGST FT-VMKEAW TQLYGAVVQA MSRGW-----						
CHIONODRACO_MYERSI_gi	YMLQCSLGQA YT-APLRQAW LNLYSIVVAA MM-----						

BOVICHTUS VARIEGATUS_	YMLQCSLGQA YT	APLRQAW	LNLYSIVVAA	MSR	-----	-----	-----	-----
SIPUNCULA
THEMISTE_1_LGB2_gn1 t	-TLIKANYKS YD	SKLDDCW	VPLWNFMNNE	MTTGQEVYRG	GKI	-----	-----	-----
THEMISTE_2_LGB2_gn1 t	-TLIKANYKA YD	SKLDDCW	VRLWDFINKQ	MTTGQEYTEE	EKS	-----	-----	-----
THEMISTE_3_LGB2_gn1 t	-NLMSQSYKG YD	SKLDDAW	TRLWGFMTSQ	MMYG	-----	-----	-----	-----
BRACHIOPODA
TEREBRETRALIA_Ng2_gn1	QLMSSGLQG YD	QELDALW	EKLYTWTITKR	MEEGM	-----	-----	-----	-----
ANNELIDA
ALVINELLA POMPEJANA_1	EYLKDKAKVV FN	DEAEAAM	OKLFDYVLDI	TAAVMDLQIE	KMG	-----	-----	-----
ALVINELLA POMPEJANA_2	WAVKETLGDR YT	ISIENIY	TITIRYILQS	LHDAFTKHRE	RQNSTNNDCE	KTNLLNQELS	TADRKT	---
APHRODITA ACULEATA LG	ailasvvggd ft	paakdaw	tkvyntisst	mqa	-----	-----	-----	-----
CAPITELLA_SP_1_LGB2_F	YVWYHQLKDS MD	SEVECAW	KKLLlfiivq	qragydaeke	appnglsfll	qt
CAPITELLA_SP_2_LGB2_F	DIWDEDLGGK FS	KETKEAW	IKIFALITRK	VFEGfgeett	rfrpplpyeg	kg
CAPITELLA_SP_3_LGB3_g	CIWSEELGKG FT	DEVSVAV	KTVFDFIMSQ	LQDGYA	---	---	---	---
PLATHYNEREIS_dumerili	HAISSAATGDA FT	PETAFAF	TAFYGVVTH	---	---	---	---	---
HELLOBDELLA_ROBUSTA1_	SIWKQELKDV YT	NELSEAW	HKVLVYIVSK	LKEGYETEWK	VATYFNPQ	---	---	---
ECHIURA
URECHIS CAUPO_Ng1_gn1	AFLKSKLGYN FD	DATGAAC	AAASVMITVL	QAEKLT	---	---	---	---
PLATYHELMINTHES
MACROSTOMUM LIGNANO_1	VVFTRRLGAA YT	DNKAVV	MKLLegvipv	Iqrgm	-----	-----	-----	-----
MACROSTOMUM LIGNANO_2	VVFERRLGKA FV	DNKAVV	VKLLQGIipv	Iqrgl	-----	-----	-----	-----
CHOANOFMAGELLATE
MONOSIGA BREVICOLLIS	KTLLAIGLGGK WN	PEARRAW	EIVCGLLISP	IRTGILQART	KANHLRAKEA	ERKRQLEMAA	ARLEGRVASS	---
SALPINGOECA_SP_Ng2_gi	WTLGAGLGDS WD	HSVKKAW	MHVLFFILSP	LKSGLVVART	LRNDYNTSGC	LRCRRLLIPL	HGRRLRPITV	---
PLACAZOA
TRICHOPLAX ADHAERENS_	QTLSDKLQEN FT	PEVHEAW	SKTFDMITAA	MKSGMN	---	---	---	---
TRICHOPLAX ADHAERENS_	F	---	---	---	---	---	---	---
TRICHOPLAX ADHAERENS_	EMARNALKKK FT	KSTEAAM	RSFFQMVTD	IKNGIMKAQN	RN	---	---	---
TRICHOPLAX ADHAERENS_	HTIKKCGGNN WS	NETADAW	KIATKVLCEL	FREGLETGPK	K	---	---	---
PORIFERA
CARTERIOSPONGIA_FOLIA	WALEAALKEK FT	PEVKAAM	LALYKVQSL	MEQGMEEGI	---	---	---	---
AMPHIMEDON_QUEENSLAND	HALGVALGKE FN	DEAKKAW	TLLYSVVTA	MKEG	-----	-----	-----	-----
CNIDARIA
NEMATOSTELLA VECTENSI	WALQDVLGEG CT	SDVAEAM	IDLYGIYVQA	MLEG	-----	-----	-----	-----
NEMATOSTELLA VECTENSI	RLENDLQGR FS	SFASRSW	HKAYDVIVEY	IEEGLQSSYK	QDPVTGITDA	EKVLVQESWD	LLKPDLLGLG	---
NEMATOSTELLA VECTENSI	PLMKADLQDE FT	PLAESAW	KKAFDVMIA	IEQQRRARS	VATFLTNPVA	---	---	---
NEMATOSTELLA VECTENSI	AYLRQALGSH LT	EEGADAW	RKALCVMIDI	IEKGSTSERW	---	---	---	---
NEMATOSTELLA VECTENSI	DTIQDLLKTQ WT	DGTAEAM	NKLFRFIADT	MKHGLSS	---	---	---	---
NEMATOSTELLA VECTENSI	YSISKGLGSL FN	DEIGEAM	QAMYDLMSGA	MISGTKAVQA	RSQNSL	---	---	---
NEMATOSTELLA VECTENSI	WALQDVLGEG CT	SDVAEAM	IDLYGIYVQA	MLEGLQQA	GR	-----	-----	-----
NEMATOSTELLA VECTENSI	ETFEENLGK WT	VEIAESW	KLLLDYVMAM	IIRGLRSP	---	---	---	---
NEMATOSTELLA VECTENSI	FAMRLHLRKE WS	RATSKAW	EKIFSFMADA	MMRGCKG	---	---	---	---
NEMATOSTELLA VECTENSI	IEEILATASL WD	FQVEEAM	NRLFDSITAM	ILRGIQLAKV	---	---	---	---
NEMATOSTELLA VECTENSI	DTIQDLLKTQ WT	DGTAEAM	NKLFRFIADT	MKHGLSS	---	---	---	---
NEMATOSTELLA VECTENSI	DTLRNLLQSQ WT	EETAEM	NKLFSPFIST	MVRGLQSRD	---	---	---	---
CLYTIA HEMISPHERICA	YGICMIMD VD	SSVENAM	DSLRFVMDL	LKLGMRMEKE	AQEKESLNKG	YNTBELLEKA	QDGGALDEN	---
CLYTIA HEMISPHERICA	ETLEEAFKEG FQ	EDSKTAW	SKIVDAIAKY	MLKGIDKAKE	QTKTSELSSD	DGDGKMEQ	---	---
CLAVA_1_LGB1_Locus_11	FQYAIICISLN LD	RDTENAM	DSLFRFLMDC	LKFGMRNEMH	KETTSPLDKL	EVQDG	---	---
CLAVA_2_Ng_Locus_2853	QTLKKEKSE FD	DKALKAW	SKFIDIVIEA	FDVNDTDE	KTQPIGK	---	---	---
HYDRA MAGNIPAPILLATA	FAISQICPNI DM	ITEGAW	DSLFKYIVT	---	---	---	---	---
MONTASTRAEA FAVEOLATA
MONTASTRAEA FAVEOLATA	WVLTATGLGDD FT	-----	-----	-----	-----	-----	-----	-----
ACOELA
SYMSAGITTIFERA_ROSCOF	VEAMAKKGQW K	KKTIIAM	EKFFDFIRAA	MVHGLKRRKG	HSSISNTTSA	ANTAACKNHN	SPSSSQ	----
SYMSAGITTIFERA_ROSCOF	TQVVKRVSEA	KWSEKKEAM	LKAYGIITVM	VTE	-----	-----	-----	-----
NEMERTODERMATIDA
MEARA STICHOPI_Ng1_tr	HIKSKLEQV WT	DEIGDAM	KVMFDIIVFN	LKSGQNQEMQ	DRGITV	---	---	---
NEMERTODERMA WESTBLAD	GTVIDLNGGS LS	DEAAAAM	GLLWEALVLC	VLEGMKKPGQ	PGMGQTHSHA	DQIIGELLVT	HPAHVVALYI	---
NEMERTODERMA WESTBLAD	WLVEQSNNGR LP	PAVEDSW	QRLWERITHF	ILAGEEESK	TDSGGTGV	VEQNSHVIMS	TGVQNSLNF	---
NEMERTODERMA WESTBLAD	NMVIQCCGGM VD	QDTLVAL	SLYWDFIMRC	VKEGMEEAAA	KVWSDSTTLN	GSLYLKIINT	PLLYCPCCHR	---
XENOTURBELLIDA
XENOTURBELLA_HEMOGLOB	GVLSEKCGGA LA	GDASSGW	OKLYNEIVRL	IDAGQKTPAA	GGARAEFLAA	IGAN	-----	-----
MOLLUSCA
SEPIA OFFICINALIS_LGB	KIMTERLGKA IP	PLGKGLV	DKSIDCS	---	---	---	---	---
EUPRYMNA SCOLOPES_1_L	KFLTIRLGN YT	PLAKESW	TKALTVVNAV	IEKGIEDGID	CSECRYRERN	RRST	-----	-----
DORYTEUTHIS PEALEII_L	DAIKITLQDR YS	DNMDIY	KLIVRFLTE	VTGARVDVS	ST	-----	-----	-----
EUPRYMNA SCOLOPES_2_L	DAIKITLQDR YS	DNMNIY	KLIVRFLTE	VTGARANVS	ST	-----	-----	-----
MYTILUS CALIFORNIANUS	DVWEYTI GEE FI	PEVRESW	TEFFDYLVKY	MCQGYNVFTN	ET	-----	-----	-----
MYTILUS CALIFORNIANUS	KAVKEVLGDA AT	DEVLDHW	KKGFHFLAEH	IRGLEAEK	---	---	---	---
SPISULA SOLIDISSIMA_L	SYAQDHAKFD --	DFTAAAM	TKTLKVVADV	IG	---	---	---	---
APLYSIA_1_LGB3_CNSN01	EAVRVTLGDR YT	ENMQNIY	TIAIFILET	LQOQMEEALE	KAGSSEVAQG	HIRV	-----	-----
APLYSIA_2_LGB2_CNSN01	VGLGAVLGGG YT	KEVSEAM	CKLYSYICLQ	MQIGMENPN	---	---	---	---
LIOLOPHURA JAPONICA_N	ATFAQQMGGA YT	AAMKDAL	KAATGvivp	gmkagy	---	---	---	---
CRASSOSTREA GIGAS_2_L	QAITTRHWSV D	SQREQAW	NNFYCTL	---	---	---	---	---
CRASSOSTREA GIGAS_1_L	EAWESAIGEE FI	PEVQNCW	TQLFAYIMRY	IVQGYELYFS	EC	-----	-----	-----
ARTHROPODA

CAPITELLA_SP_3_LGB3_g
PLATHYNEREIS_dumerili	-----	-----	-----	-----	-----
HELLOBDELLA_ROBUSTA1	-----	-----	-----	-----	-----
ECHIURA
URECHIS_CAUPO_Ng1_gn1	-----	-----	-----	-----	-----
PLATYHELMINTHES
MACROSTOMUM_LIGNANO_1	-----	-----	-----	-----	-----
MACROSTOMUM_LIGNANO_2	-----	-----	-----	-----	-----
CHOANOFLAGELLATE
MONOSIGA_BREVICOLLIS	GVQFSSDTER	SRRSTAASAT	ATPHCGLKSN	RFSSNSLRKT	IL
SALPINGOECA_SP_Ng2_gi	SLV-----	-----	-----	-----	-----
PLACOOZA
TRICHOPLAX_ADHAERENS	-----	-----	-----	-----	-----
TRICHOPLAX_ADHAERENS	-----	-----	-----	-----	-----
TRICHOPLAX_ADHAERENS	-----	-----	-----	-----	-----
TRICHOPLAX_ADHAERENS	-----	-----	-----	-----	-----
PORIFERA
CARTERIOSPONGIA_FOLIA	-----	-----	-----	-----	-----
AMPHIMEDON_QUEENSLAND	-----	-----	-----	-----	-----
CNIDARIA
NEMATOSTELLA_VECTENSI	-----	-----	-----	-----	-----
NEMATOSTELLA_VECTENSI	-----	-----	-----	-----	-----
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NEMATOSTELLA_VECTENSI	-----	-----	-----	-----	-----
NEMATOSTELLA_VECTENSI	-----	-----	-----	-----	-----
CLYTIA_HEMISPHAERICA	ENSAPMPLAL	INEDDSTSSV	RTFSCR----	-----	-----
CLYTIA_HEMISPHAERICA	-----	-----	-----	-----	-----
CLAVA_1_LGB1_Locus_11	-----	-----	-----	-----	-----
CLAVA_2_Ng_Locus_2853	-----	-----	-----	-----	-----
HYDRA_MAGNIPAPILLATA	-----	-----	-----	-----	-----
MONTASTRAEA_FAVEOLATA	-----	-----	-----	-----	-----
MONTASTRAEA_FAVEOLATA	-----	-----	-----	-----	-----
ACOELA
SYMSAGITTIFERA_ROSCOF	-----	-----	-----	-----	-----
SYMSAGITTIFERA_ROSCOF	-----	-----	-----	-----	-----
NEMERTODERMATIDA
MEARA_STICHOPI_Ng1_tr	-----	-----	-----	-----	-----
NEMERTODERMA_WESTBLAD	TNYLLLFIKY	LHHEPK----	-----	-----	-----
NEMERTODERMA_WESTBLAD	HHIFFNNNCP	YCIPQLT---	-----	-----	-----
NEMERTODERMA_WESTBLAD	TVSDIITRCM	LRVAII----	-----	-----	-----
XENOTURBELLIDA
XENOTURBELLA_HEMOGLOB	-----	-----	-----	-----	-----
MOLLUSCA
SEPIA_OFFICINALIS_LGB	-----	-----	-----	-----	-----
EUPRYMNA_SCOLOPES_1_L	-----	-----	-----	-----	-----
DORYTEUTHIS_PEALEII_L	-----	-----	-----	-----	-----
EUPRYMNA_SCOLOPES_2_L	-----	-----	-----	-----	-----
MYTILUS_CALIFORNIANUS	-----	-----	-----	-----	-----
MYTILUS_CALIFORNIANUS	-----	-----	-----	-----	-----
SPISULA_SOLIDISSIMA_L	-----	-----	-----	-----	-----
APLYSIA_1_LGB3_CNSN01	-----	-----	-----	-----	-----
APLYSIA_2_LGB2_CNSN01	-----	-----	-----	-----	-----
LILOPHURA_JAPONICA_N	-----	-----	-----	-----	-----
CRASSOSTREA_GIGAS_2_L	-----	-----	-----	-----	-----
CRASSOSTREA_GIGAS_1_L	-----	-----	-----	-----	-----
ARTHROPODA
APIS_MELLIFERA_LGB2_g	-----	-----	-----	-----	-----
CARCINUS_MAENAS_1_LGB	-----	-----	-----	-----	-----
CARCINUS_MAENAS_2_LGB	-----	-----	-----	-----	-----
DAPHNIA_PULEX_GlobinX	-----	-----	-----	-----	-----
DAPHNIA_CARINATA_LGB2	-----	-----	-----	-----	-----
HARPEGNATHOS_SALTATOR	-----	-----	-----	-----	-----
TUNICATA
Ciona_intestinalis_Gb	-----	-----	-----	-----	-----
Ciona_savigny_Gb1_LGB	-----	-----	-----	-----	-----
MOLGULA_TECTIFORMIS_L	-----	-----	-----	-----	-----
BOTRYLLUS_SCHLOSSERI	-----	-----	-----	-----	-----
CEPHALOCHORDA
Branchiostoma_florida	-----	-----	-----	-----	-----

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Branchiostoma florida DAQSEENGVH -----
Branchiostoma florida -----
Branchiostoma florida -----
HEMICHORDATA .....
Saccoglossus kowalevs -----
Saccoglossus kowalevs -----
Saccoglossus kowalevs NGDKEDNCIE SSKQKDITVI ASTKQR----
Saccoglossus kowalevs -----
BALANOGLOSSUS_CLAVIGE -----
ECHINODERMATA .....
STRONGYLOCENTROTUS_PU -----
KINORHYNCHA .....
ECHINODERES_HORNI_Glo -----
ENTOPROCTA .....
PEDICELLINA_SP_LGB1_E -----
TARDIGRADA .....
RICHTERSIUS_CORONIFER -----

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