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SNAREing the basis of multicellularity: Consequences of protein family expansion during evolution

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Abstract

Vesicle trafficking between intracellular compartments of eukaryotic cells is mediated by conserved protein machineries. In each trafficking step, fusion of the vesicle with the acceptor membrane is driven by a set of distinctive SNARE proteins that assemble into tight four-helix bundle complexes between the fusing membranes. During evolution about twenty primordial SNARE types were modified independently in different eukaryotic lineages by episodes of duplication and diversification. Here we show that two major changes in the SNARE repertoire occurred in the evolution of animals, each reflecting a main overhaul of the endomembrane system. In addition, we found several lineage-specific losses of distinct SNAREs, particularly in nematodes and platyhelminthes. The first major transformation took place during the transition to multicellularity. The primary event that occurred during this transformation was an increase in the numbers of endosomal SNAREs, but the SNARE-related factor Lgl also emerged. Apparently, enhanced endosomal sorting capabilities were an advantage for early multicellular animals. The second major transformation during the rise of vertebrates resulted in a robust expansion of the secretory set of SNAREs, which may have helped develop a more versatile secretory apparatus.

Introduction

A defining feature of eukaryotic cells is their complex system of internal compartments. Directional transport between the compartments is mediated by vesicles that bud from a donor organelle and then fuse with an acceptor organelle. As each of the internal compartments serves a specific and vital cellular function, it must maintain its identity during vesicle trafficking. This is achieved by specific protein machineries that tightly regulate each transport step (Bonifacino and Glick 2004; Behnia and Munro 2005; Cai et al. 2007). It has been recognized that several of the key machineries involved in vesicle trafficking steps appear to be the result of gene duplications, followed by diversifications (Cavalier-Smith 2002; Dacks and Field 2007). For example, in each trafficking step the core of the protein machinery involved in the fusion process is composed of members of the so-called SNARE (Soluble NSF Attachment Protein Receptors) protein family (Hong 2005; Jahn and Scheller 2006). SNARE proteins are small cytoplasmatically orientated membrane-associated proteins with a relatively simple domain architecture. In most SNARE proteins, an extended coiled coil segment, the so-called SNARE motif, is directly adjacent to a single trans-membrane domain at the C-terminal end. As a common mechanism, the SNARE motifs of heterologous sets of SNARE proteins assemble in a zipper-like fashion into tight four-helix bundle units between two membranes, a process that clamps the membranes together and initiates fusion. The SNARE machinery has diversified only modestly in the course of eukaryotic evolution. In fact, we were able to show that a basic set of twenty types of SNARE proteins is conserved in all eukaryotes, suggesting that they constitute the set of SNARE genes of the proto-eukaryotic cell (Kloepper et al. 2007).

In the previous study, we recognized that the basic set of SNARE genes had been independently modified by paralogous expansion in several different eukaryotic lineages (Kloepper et al. 2007). So far, an increase in the number of SNARE genes has been observed mainly in animals and in land plants (Bock et al. 2001; Dacks and Doolittle 2004; Yoshizawa et al. 2006; Dacks and Field 2007; Sanderfoot 2007). A genome-wide inspection of the first four eukaryotic species sequenced came to the conclusion that mammals (e.g. *Homo sapiens*) contain an enlarged set of secretory SNAREs compared to invertebrates (e.g. the fruitfly *Drosophila melanogaster* and the round worm *Caenorhabditis elegans*). Since the number of SNARE genes in the two invertebrates was comparable to that of the yeast *Saccharomyces cerevisiae*, the authors concluded that multicellular organisms do not necessarily require an enlarged repertoire of SNARE proteins (Bock et al. 2001). Rather, they proposed, the enlargement of the secretory SNAREs took place in mammals. In other studies, by contrast, it was suggested that in both, animals and land plants, the increase in the number of SNAREs, in particular the ones involved in secretion, occurred much earlier in evolution, paralleling the leap from single to multicellular organisms (Sanderfoot et al. 2000; Dacks and Doolittle 2004). On that account, the idea has been put forward that the secretory pathway of multicellular organisms requires an additional level of complexity compared to that of their single-cell predecessors (Dacks and Field 2007; Sanderfoot 2007). Only with an increased number of secretory SNARE proteins, according to this idea, an individual cell within a tissue is able to deliver material to different regions of its plasma membrane. This line of reasoning, however, neglects the fact that the capability to transport and secrete material to specific regions of the plasma membrane is an essential part of the life cycle of every eukaryotic cell, multicellular or unicellular (Soldati and Schliwa 2006).

For example, polarized secretion is the basis of the well-studied process of asymmetric enlargement of the budding daughter cell during cytokinesis in the single-cell yeast *Saccharomyces cerevisiae* (Brennwald and Rossi 2007). Furthermore, multicellular species are found throughout the eukaryotic kingdoms, for example in the lineage of fungi, yet all inspected fungi species appear to maintain an almost unchanged basic set of SNARE proteins throughout evolution (Kloepper et al. 2007). Moreover, considerably enlarged SNARE repertoires, including extra secretory SNAREs, have been detected in several unicellular eukaryotes, for example for the ciliate *Paramecium tetraurelia* with its multifaceted endomembrane system (Kloepper et al. 2007). Together, these observations cast doubt on a strict correlation between the increase in the number of secretory SNARE genes and the emergence of multicellularity.

So far, all accounts of SNAREs in animals were based on only few model organisms. In addition, in the previous studies, some more-derived SNAREs have not been included (Bock et al. 2001; Dacks and Doolittle 2004; Yoshizawa et al. 2006). Thus, it still remains unclear exactly which SNARE factors have been added at which point during animal evolution. Clearly, in order to gain deeper insights into adaptations of the metazoan endomembrane system, a much broader database containing the SNARE repertoires of species that represent most major branches of the complex evolutionary tree of animals must be used. In addition, not only the number of SNARE genes but, more importantly, the evolutionary history of the different functional SNARE types needs to be reconstructed more thoroughly. Only this knowledge will allow us to address a series of pressing questions: How did the transition to multicellularity affect the endomembrane system? Are the changes mostly observable in the SNAREs actually involved in the exocytotic step of the secretory pathway? Is it possible to associate the rise or loss of certain SNARE factors with specific lineages within animals? Are the routes of the secretory pathway indeed comparable between various animal species?

In order to answer these questions, we considerably expanded our previous collection (Kloepper et al. 2007) by integrating sequences from a multiplicity of different animal species, making use of the abundance of new genomic data that have been established lately. A point-by-point account of the SNARE repertoires was facilitated by our recent classification of the multigene SNARE family, with its largely improved ability to distinguish between orthologues and paralogues (Kloepper et al. 2007). Moreover, this approach allows us to integrate new SNARE sequence data from genome and EST projects quickly. Only this allowed us to pinpoint the changes in the SNARE sets throughout all metazoan lineages investigated. In addition, we constructed phylogenetic trees for each individual SNARE subgroup. All these factors enabled us to infer a probable sequence of events for the observed changes in the SNARE repertoire during animal evolution.

We discovered that during metazoan evolution, two major changes in the SNARE repertoire have occurred. The first major change took place during the transition from single to multicellular organisms. During this transition, mostly SNARE proteins involved in endosomal trafficking were duplicated and diversified, as well as some SNAREs involved in secretion. In addition, SNARE-related factors like the tumor suppressor lethal giant larvae (Lgl), which plays a key role in establishing epithelial cell polarity, originated. The second major change took place in the lineage of vertebrates, where the secretory SNARE repertoire in particular enlarged drastically.

Materials and Methods

Sequences

In a recent classification of SNARE proteins, we published a set of more than 2000 protein sequences from 145 eukaryotic species, including 811 sequences from 59 metazoan species (Kloepper et al. 2007). These sequences were classified into twenty distinct groups. Here, we used the established HMM models (Kloepper et al. 2007) to classify the SNAREs of newly available metazoan genomes and EST libraries. We gathered 1234 additional sequences from the nr-database at NCBI and various genome projects (DOE Joint Genome Institute (JGI); Baylor College of Medicine, J. Craig Venter Institute and Broad Institute) and various EST databases.

After removing sequences with more than one occurrence, sequences which seemed to be misassembled, or sequences which failed an a visual verification, we obtained a set of 1774 unique SNARE sequences from 177 metazoan species. In addition, we identified sequences of two SNARE-related protein families using an extensive blast search on the nr-database at NCBI, identifying 37 sequences of lethal giant larvae (Lgl) and 41 sequences of Sec22-like. Both proteins exhibit a very high N-terminal sequence similarity to tomosyn and Sec22, respectively, but, interestingly, have lost their SNARE domain. Since we were especially interested in pinpointing the emergence of innovations within the SNARE set of metazoans, we supplemented our collected sequences with a set of 31 sequences from two choanoflagellates and two sequences from two ichthyosporeans (JGI and EST data). All species and SNARE sequences used in this study are listed in Suppl Table 1. These newly classified sequences were integrated into the SNARE database (Kloepper et al. 2007) for public access, which can be accessed via our projects homepage (<http://bioinformatics.mpibpc.mpg.de/snare/>). Any sequence set used in the phylogenetic analysis was aligned using muscle (Edgar 2004) with standard settings. Sites with more than 25 percent gaps were removed from the alignments. After removing those sites, we removed aligned sequences which contained more than 50 percent gaps. The reduced alignments are available via the supplementary section of our projects homepage.

Phylogeny

The phylogenetic reconstruction was separated into two parts and was carried out essentially as described (Kloepper et al. 2007). Firstly, we reconstructed a tree from the conserved alignments using IQPNNI (Important Quartet Puzzling and Nearest Neighbor Interchange) (Vinh le and Von Haeseler 2004) with a gamma distribution as a model for rate heterogeneity. The estimation for the gamma distribution parameter used four rate categories. Additionally, the proportion of invariable sites was estimated from the data, and the Jones, Taylor and Thornton (JTT-) distance matrix was used as a substitution matrix, and the stopping rule of the algorithm was used but the algorithm had to run for at least the suggested number of iterations. All other settings of the application were set to default values. For each edge of the constructed tree, we estimated the confidence using Likelihood-Mapping. Secondly, we used the *phylip* package (Felsenstein 1998) to apply a distance-based bootstrap analysis with 1000 replicates. We used standard settings for seqboot, the JTT distance matrix once again and also a gamma distribution (with parameter approximation from tree-puzzle) for protdist and standard options for neighbour. Whenever necessary, we used a random seed of nine. Because bootstrap values have been shown to be systematically

biased, we used the almost unbiased (AU) test (Shimodaira 2002) to correct for this. The site wise log-likelihoods needed for the AU-test were obtained using a modified version of *phyml* (Guindon and Gascuel 2003) and the test was performed using *consel* (Shimodaira and Hasegawa 2001). We joined the results of both estimations using the tree of IQPNNI as a starting point and labeled the inner edges of the tree with their Likelihood-Mapping and corrected bootstrap support values. All trees can be downloaded via the supplemental section of our projects homepage. The trees are stored in Nexus format and can be interactively explored using, for example, *SplitsTree 4* (Huson and Bryant 2006).

Results

In order to catalog the SNARE inventory of animals, we complemented our previous collection (Kloepper et al. 2007) with newly published genomes and ESTs from different animal species, obtaining an overall set of 1839 SNARE sequences from 177 different metazoan species. This data set was complemented by 31 sequences from two choanoflagellates and 2 sequences from two ichthyosporeans, both of which are believed to be closely related to metazoans. All sequences and species used in this study are listed in Suppl. Table 1 and can also be found in our SNARE data base on our projects homepage. The data set comprises about 55 species with nearly complete sets of SNAREs (Suppl. Table 2). These sequences were classified into their corresponding basic subgroups according to our recently established HMM profiles. The classification scheme served as the framework to reconstruct the history of these changes. It is based on the highly conserved SNARE motif, as the other sequence regions are usually less conserved. However, we observed that within the animal kingdom, other sequence regions are also usually well conserved. Therefore, we constructed alignments of the entire sequences for each subgroup using *muscle* (Edgar 2004). Sites with more than 25 percent gaps were removed from the alignments. In addition, we excluded all sequences which contained more than 50 percent gaps. The length of these conserved alignments varied from 99 sites for the R.IV subgroup to 292 sites for the Qa.I subgroup.

Based on these alignments, we constructed phylogenetic trees for each of the fundamental SNARE types. When we used SNARE protein sequences from several other eukaryotic kingdoms as an outgroup, we observed that the different SNARE sequences from animals usually congregated within the phylogenetic trees and were well separated from the outgroup (data not shown), substantiating the view that animals constitute a monophyletic group.

The secretory SNARE set of Choanoflagellates is metazoan-like

The multicellular progenitor of animals, the urmetazoan, is believed to have evolved from an unicellular flagellated protist, probably closely related to extant heterotrophic, single-cell choanoflagellates (Brooke and Holland 2003; King 2004). In order to investigate whether SNARE genes were modified during this pivotal transition, we collected SNARE sequences from the genome of the choanoflagellate *Monosiga brevicollis* and supplemented these data with EST sequences from another choanoflagellate, *Monosiga ovata*. We detected a relatively simple repertoire of SNARE proteins in *M. brevicollis*, in principal consisting, except for a few duplications, of the basic set of SNARE proteins we had recently deduced for the proto-eukaryotic cell. Comparable basic sets of SNAREs can be found in green algae and fungi, for example. It should be noted, however, that we cannot rule out the possibility that the genome data of *M. brevicollis* is incomplete or that *M. brevicollis* has secondarily lost some SNARE factors.

In general, our phylogenetic analysis places *M. brevicollis* closer to metazoans than to other eukaryotes, supporting the notion that choanoflagellates are indeed closely related to the unicellular ancestor of animals. In particular, the secretory SNAREs of choanoflagellates are closely related to the set of animal SNAREs believed to mediate regulated secretion. For example, the SNAP-25 like sequences (Qbc-SNARE) from choanoflagellates contain a stretch of cysteines at the C-terminal end of their Qb-SNARE motif. These cysteines are known to be palmitoylated and serve as a membrane-anchor for animal SNAP-25 (Veit et al. 1996).

An enlarged set of endosomal SNAREs in multicellular animals

In order to reconstruct the SNARE repertoire of the metazoan ancestor, we compared the set of SNARE genes of lower metazoans with that of choanoflagellates. Remarkably, in comparison to the basal SNARE set of *M. brevicollis*, we detected a considerably enlarged repertoire of SNARE and SNARE-related genes in cnidarians, particularly in *Nematostella vectensis*, and also in the placozoan *Trichoplax adhaerens*, bringing to light a clear expansion of the SNARE repertoire during the transition from single-cell to multicellular organisms. Notably, both phyla exhibit a very similar SNARE repertoire, including several new molecular “inventions” (see Fig. 1a for the current view of the phylogenetic relationships of metazoans). Remarkably, we observed that the SNARE sets of the lower metazoans are very similar to those of various bilaterians, suggesting that the extended SNARE repertoire was already present in the last common ancestor of lower metazoans and bilaterians.

Our rigorous classification analysis was able to delineate which of the original functional SNARE types were duplicated during this first phase of expansion. In detail, we found at least nine evolutionary novel SNARE types in lower metazoans compared to single-cell choanoflagellates (Fig. 1b; for details, see also Supp. Table 2), suggesting the presence of a more multifaceted membrane trafficking system in metazoans in general. Notably, the exact function of most of these new factors and whether they were ubiquitously expressed is not entirely clear yet. Remarkably, most of these changes occurred in SNAREs involved in endosomal trafficking steps.

Besides the canonical R-SNARE Vamp7 (also referred to as Ti-Vamp; R.III-type), two additional endosomal R-SNAREs, Vamp7-like and Vamp4, were detected in lower metazoans (Fig. 1b). In animals, Vamp7 is involved in endosomal trafficking towards lysosomes (Advani et al. 1998; Advani et al. 1999), but it has also been implicated in secretion from lysosomal compartments (Martinez-Arca et al. 2003; Marcet-Palacios et al. 2007; Pocard et al. 2007). The function of the additional, R-SNARE Vamp7-like (not described to date), which is only present in lower metazoans, is unknown. However, its domain structure is almost identical to Vamp7, suggesting a similar function for both factors. Interestingly, in the phylogenetic tree (Fig. 2a), Vamp7-like appears to be closely related to endobrevin/Vamp8, which only exists in vertebrates. Endobrevin does not contain an N-terminal extension. It is therefore possible that Vamp7-like has lost the entire N-terminal profilin-like domain, giving rise to the novel R-SNARE endobrevin in the vertebrate lineage. Endobrevin was originally identified as an endosomal R-SNARE (Advani et al. 1998; Wong et al. 1998b), but later has been shown to be involved in regulated secretion of pancreatic zymogen granules (Wang et al. 2004; Wang et al. 2007). The third metazoan R.III-type SNARE, Vamp4, contains a shorter N-terminal extension comprising a highly conserved double-leucine motif followed by an acidic cluster (Advani et al. 1998). This region is important for its interaction with the cytosolic coat adaptor, AP-1, that targets Vamp4 to the TGN (Peden et al. 2001; Hinners et al. 2003; Zeng et al. 2003; Tran et al. 2007).

In choanoflagellates, we found only one Qa.III.b-SNARE, Syx7, whereas two additional endosomal syntaxins were found in most metazoans, Syx17 and Syx20 (Fig. 1b). Syx17 is the most deviated SNARE of the Qa.III.b-group in animals (Fig. 2b). Originally, Syx17 had been implicated in trafficking towards the smooth-surfaced ER (Steggmaier et al. 2000), but this function is debated (Zhang et al. 2005). Notably, and in contrast to all other known SNARE proteins, all Syx17 possess two adjacent transmembrane regions at the C-terminal end (Steggmaier et al. 2000). Syx20

is a SNARE gene we had first discovered in vertebrates only (Kloepper et al. 2007). A more detailed phylogenetic analysis of all Qa.III.b-type SNAREs (Fig. 2b) in animals now indicates that this factor is present in several other branches of metazoans as well. In insects, this factor originally has been termed Syntaxin 13. This nomenclature is somewhat confusing, since the name Syx13 (also called Syx12) (Advani et al. 1998; Tang et al. 1998d) is usually reserved for a Qa.III.b-SNARE closely related to Syx7 that is found only in vertebrates (Wong et al. 1998a) (see also below). We therefore propose to generally use the name Syx20 for this Qa.III.b-type.

Furthermore, two independent genes encoding for Qb.III.b-SNAREs, Vti1a and Vti1b (Advani et al. 1998; Xu et al. 1998), were found in most metazoans (Fig. 1b). Both are thought to function in early and late endosomal trafficking, respectively. A set of two independent Vti1 genes is present in *M. brevicollis* as well, although our analysis is not able to determine whether these genes correspond to the split of Vti1a and Vti1b found in metazoans (Suppl. Data) or if these two genes represent an independent duplication.

In addition to the enlargement of the endosomal SNARE set, some other (non-endosomal) SNAREs have been duplicated in this first expansion phase. Instead of the single secretory Qbc-SNARE (Qbc.IV) SNAP-25 discovered in *M. brevicollis*, the metazoan ancestor probably possessed three different Qbc-SNARE genes; SNAP-25 (Oyler et al. 1989), SNAP-29 (Steggmaier et al. 1998; Wong et al. 1999) and SNAP-47 (Holt et al. 2006) (Fig. 1b). In contrast to SNAP-25, both SNAP-29 and SNAP-47 do not own a stretch of cysteines in the center of the linker between the two SNARE motifs. In the phylogenetic tree (Fig. 2c), SNAP-29 and SNAP-47 are well separated from SNAP-25. As with several of the other novel SNAREs mentioned above, the exact function of SNAP-29 and SNAP-47 is not entirely clear. Both SNAP-29 and SNAP-47 show a widespread distribution on intracellular organelles. SNAP-29 is thought to be involved in trafficking within the Golgi apparatus, but has also been suggested to mediate constitutive secretion, whereas the function of SNAP-47 is not known so far.

Besides the bona-fide SNARE factors, i.e. proteins that possess a functional SNARE motif, we found two additional factors that are clearly derived from SNARE genes (Fig. 1b and Fig. 3a). One is Sec22-like (also referred to as Sec22a (Hay et al. 1996)), which is derived from the R.I-SNARE Sec22 (Sec22b) that functions in trafficking between the ER and the Golgi apparatus (Hay et al. 1997; Liu and Barlowe 2002). Sec22-like carries a profilin-like N-terminal domain closely related to Sec22, but instead of a C-terminal SNARE motif, it carries three consecutive C-terminal transmembrane domains. Sec22-like is localized to the ER, but its function is unknown (Hay et al. 1996; Tang et al. 1998b).

The second novel factor is Lethal giant larvae (Lgl), which has a function in establishing epithelial cell polarity in multicellular animals (Vasioukhin 2006; Wirtz-Peitz and Knoblich 2006). Lgl is derived from tomosyn (Pobbati et al. 2004; Fasshauer and Jahn 2007), a regulatory R-SNARE (R.Reg) that is thought to be involved in polarized secretion (Fig. 1b). As tomosyn does not contain a transmembrane anchor at the end of its R-SNARE motif, it cannot serve as a fusogenic R-SNARE; instead, it is thought to regulate the accessibility of the SNARE-acceptor complex for the R-SNARE residing on secretory vesicles. Tomosyn carries a large N-terminal domain composed of two consecutive seven-bladed β -propeller domains (Hattendorf et al. 2007). Lgl possesses a very similar N-terminal domain, but it has lost the C-terminal R-SNARE motif.

Furthermore, animals, like several other eukaryotic lineages, for example

fungi and plants, usually possess two different Qc.II-SNAREs, which are thought to participate in consecutive trafficking steps within the Golgi apparatus. Nevertheless, it remains unclear, whether a duplication of an original Qc.II-SNARE occurred only once in the eukaryotic ancestor or whether the pair of Qc.II-SNAREs in animals, Bet1 (Hay et al. 1996) and Gs15 (Xu et al. 1997), arose by an independent duplication.

Losses and gains of individual SNARE genes in different animal-lineages

The increased set of 29 to 30 different SNARE proteins in urmetazoans discovered in this study, has been modified differently in different lineages of animals. We discovered nested duplications, particularly in vertebrates, and probably independent losses of individual factors in different lineages of invertebrates. Typically, gene losses affected the novel metazoan SNAREs rather than the basic repertoire. A schematic overview of the observed lineage-specific changes in the SNARE repertoire is given in Suppl. Table 2.

The genes SNAP-47 and Sec22-like were often lost, for example in nematodes, platyhelminths, insects, and crustaceans, and also in tunicates. Furthermore, we were unable to detect the Qc.II SNARE Gs15 in nematodes and insects. Loss of this gene must have occurred independently in both lineages, which are often grouped together as ecdysozoans, as we found Gs15 to be present in crustaceans and arachnidans, both of which are grouped with insects to the arthropods.

Both nematodes and platyhelminths have lost, probably independently, the endosomal SNAREs Vti1b (Qb.III.b) and Syx8 (Qc.III), of which the latter belongs to the basic SNARE set. Both proteins are thought to cooperate in one SNARE unit during transport from early towards late endosomes. Interestingly, it has been observed that mice deficient in Vti1b were viable and fertile, but had reduced amounts of Syx8, whereas the quantity of the other putative complex components, endobrevin and Syx7, was not changed (Atlashkin et al. 2003). This suggests that Syx8 and Vti1b are functionally tightly linked. Yet upon first inspection, we noted that the fruitfly *Drosophila melanogaster* contained Syx8 but apparently only one Vti1 gene. However, as we observed that several other insects, e.g. *Anopheles gambiae*, possess two different Vti1 genes, we re-inspected the genome of *D. melanogaster* and discovered a second, so far not annotated Vti1b gene on chromosome 3R. In addition, we came across a loss of the R.III-SNARE Vamp4 within holometabolous insects: so far, we have discovered Vamp4 only in Orthoptera (grasshoppers) and Hymenoptera (bees, wasps and ants), but not in Coleoptera (beetles), Lepidoptera (butterflies and moths), or Diptera (flies). In nematodes as well as in platyhelminthes, only one copy of R.III-SNAREs was found, suggesting that they have secondarily lost the novel types. However, the assignment of their R.III-SNAREs to the three metazoan types, Vamp7, Vamp7-like/endobrevin and Vamp4, is challenging, because the sequences are rather deviated, something that is frequently observed for SNARE sequences from these two phyla.

Besides several gene losses in “invertebrates” discussed above, we also came across several nested duplications within these lineages, mostly of secretory SNAREs. For example, the fruitfly possesses a duplicated set of SNAREs involved in regulated secretion, the Qbc-SNAREs SNAP-24 and SNAP-25 (Niemeyer and Schwarz 2000; Vilinsky et al. 2002), and the R-SNAREs syb (Sudhof et al. 1989; Chin et al. 1993) and nsyb (DiAntonio et al. 1993). Furthermore, we noticed that even the simple animal *Trichoplax adhaerens* possesses an enlarged set of secretory SNAREs.

Two rounds of whole-genome duplications gave rise to the enlarged set of secretory SNAREs in vertebrates

A much more pronounced enlargement by nested duplications of the SNARE repertoire occurred in the vertebrate lineage: several SNARE genes can be found in multiple copies in vertebrates, while invertebrates and basal deuterostomes typically possess only one orthologue. This observation is consistent with two consecutive rounds of whole genome duplications that are believed to have occurred early in vertebrate evolution (Ohno 1970; Holland 2003; Panopoulou and Poustka 2005).

Notably, the gene duplications in vertebrates mostly affected the secretory SNAREs. For example, duplication of the SNAP-25 gene gave rise to a second, closely related Qbc-SNARE, SNAP-23 (Ravichandran et al. 1996; Wang et al. 1997). It is possible that the alternative splicing of SNAP-25, caused by duplication of exon 5 (Bark and Wilson 1994), into two different gene products, SNAP-25a and b, also occurred in the vertebrate lineage. Even more extensive were the duplications of the secretory syntaxin (Qa.IV) and synaptobrevin (R.IV) genes, yielding a multiplicity of new gene products (Fig. 1b). The large set of secretory syntaxins (Qa.IV) in vertebrates renders the phylogenetic tree more complex (Fig. 2d). The changes in the secretory apparatus also involved the regulatory R-SNARE tomosyn. Vertebrates usually contain two different tomosyn genes (Groffen et al. 2005) and the related factor amisyn (Scales et al. 2002), which, like tomosyn, does not possess a TMR. In contrast to tomosyn, however, amisyn carries a shorter N-terminal region that is homologous to the N-terminal region of the exocyst component Sec3 (Scales et al. 2002). Moreover, the tomosyn-related gene Lgl was duplicated in vertebrates. In addition, we unearthed a few more duplications outside of the secretory set. For example, a duplication of an ancestral Qc.III gene gave rise to Syx6 (Bock et al. 1996) and Syx10 (Tang et al. 1998c) in vertebrates. In addition, an ancestral Qa.III.b SNARE was duplicated, giving rise to Syx7 and Syx13 in vertebrates. These two Qa.III.b SNAREs are thought to mediate distinct endosomal trafficking steps (Antonin et al. 2000; Collins et al. 2002; Zwillig et al. 2007). Furthermore, the duplication yielded two Sec22-like genes in vertebrates, which had originally been termed Sec22a (Hay et al. 1996) and Sec22c (Tang et al. 1998b).

A third round of whole genome duplication in bony fishes

The second genome duplication in the vertebrate lineage probably occurred before the divergence of chondrichthyes (i.e. cartilaginous fishes). This is supported by EST sequences from *Leucoraja erinacea* and *Squalus acanthias*. The position of more basic vertebrates, like the sea lamprey *Petromyzon marinus*, in relation to the two rounds of genome duplication is not entirely clear. Up to now, only few EST sequences of SNARE proteins from the sea lamprey have been available. For example, we found a fragmentary sequence that is closely related to the secretory syntaxins Syx11 and Syx19. Both factors are closely related and probably arose in a second round of genome duplication. This suggests that at least one whole genome duplication must have taken place in this organism.

Furthermore, our data support the notion that a third whole genome duplication (3R) took place in the lineage of teleost fishes (Meyer and Van de Peer 2005); (Panopoulou and Poustka 2005), since we found several duplications of secretory SNAREs in teleosts. For example, the aforementioned Syx11 gene was duplicated. In addition, several factors involved in intracellular trafficking steps were affected. For example, two independent sequences for Sec20, Sec22, Gos28, endobrevin, and possibly also Syx5 were found (Suppl. Table 2). It should be noted,

however, that this assessment is preliminary, as the genome assemblies of fishes often appear to be partly fragmentary.

Discussion

The transition to multicellularity was a pivotal step during the evolution of animals from their protozoan ancestors. Staying together as a multicellular organism brought about radical changes in lifestyle and cellular organisation, yet the basic cellular functions needed to be preserved. The emergence of multicellularity also launched the development of different cell types. The way for multicellular traits like intercellular cohesion, communication, and differentiation was paved by molecular innovations. Indeed, several domain architectures found in proteins involved in these processes are largely unique to animals (Brooke and Holland 2003; King 2004).

Phylogenetic studies of the animal kingdom often focus on protein families involved in processes related to multicellularity. For example, the evolutionary history of animal-specific homeobox genes, particularly of the collection of Hox and ParaHox genes, which are transcription factors involved cell differentiation in animals, have been intensively studied (Garcia-Fernandez 2005; Lemons and McGinnis 2006). These genes participate in the developmental program that shapes animal morphology. Obviously, the body form is acted upon strongly by natural selection, whereas it is less clear whether vital cellular functions like vesicular trafficking are also affected in multicellular organisms.

In fact, the family of SNARE proteins has diversified only modestly in the course of eukaryotic evolution (Klopper et al. 2007). Probably, the proto-eukaryotic cell contained about twenty different SNARE proteins to catalyze the vesicular trafficking steps between the main compartments found in contemporary cells (Klopper et al. 2007). We have now found this type of basic repertoire of SNARE genes in several eukaryotic organisms including the choanoflagellate *M. brevicollis*, an organism that is thought to be closely related to the unicellular ancestor of animals (Brooke and Holland 2003; King 2004). In fact, it has been shown that the precursors of several of the molecular building blocks that were largely exploited during animal evolution already existed in choanoflagellates (King and Carroll 2001; King et al. 2003; Segawa et al. 2006; Abedin and King 2008; King et al. 2008; Ruiz-Trillo et al. 2008). In line with this notion, many of the SNAREs of *M. brevicollis* are closely related to the ones from metazoans. In particular, the secretory set of SNAREs (type IV) from *M. brevicollis* closely resembles the set of SNAREs from multicellular animals, i.e. Syx1, SNAP-25 and Syb1. This set of SNAREs, which is commonly attributed to regulated secretion, is highly conserved in all animals.

Remarkably, we uncovered a major increase in the set of SNARE proteins that probably occurred upon emergence of multicellularity in animals, i.e. before the split of basal metazoans and bilaterians. This expansion alludes to a major overhaul of the endomembrane system during this critical period in metazoan evolution. The expanded, primordial metazoan set is largely conserved in most animal phyla. In particular, placozoans, cnidarians, annelids, molluscs, and lower deuterostomes usually have only marginally modified SNARE repertoires. Moreover, the SNAREs from these phyla often form relatively short branches in our phylogenetic trees, reflecting their basal character. This can even be seen for the SNAREs from the faster evolving urochordates (Dehal et al. 2002), which, together with cephalochordates and vertebrates, make up the chordates.

The nature of the first expansion of the metazoan SNARE set alludes to a larger genome change, possibly whole genome duplication(s), at the base of animal evolution. Although the function of SNARE proteins does not allow for larger domain re-arrangements, we came across a few of these examples during the first phase of

expansion: Syx17, Lgl, Sec22-like and Vamp4. Some of these re-arrangements, in particular the acquisition of additional transmembrane domains in Syx17 and Sec22-like, are clear evolutionary novelties that corroborate the monophyletic origin of all animals inspected. However, it should be borne in mind that although the species used for our analysis provided a significant coverage of most main extant metazoan lineages, at the time, only limited sequence information for the basal metazoans, the phyla Porifera and Ctenophora, are available. Nevertheless, the few SNARE sequences collected from poriferans suggest that the first phase of expansion of the SNARE repertoire occurred already before the split of the poriferans.

In contrast to previous claims that animals generally possess an enlarged set of secretory SNARE proteins (Dacks and Doolittle 2004; Yoshizawa et al. 2006; Dacks and Field 2007), our more detailed analysis now demonstrates that most novel SNAREs that arose during the first expansion originated from SNAREs involved in endosomal trafficking, whereas the major expansion of secretory SNAREs only occurred later, during the rise of vertebrates. But why in particular did the number of endosomal SNAREs increase during the emergence of multicellularity in animals? Although unicellular eukaryotes already have the ability to segregate different types of secretory proteins and to deliver them to different subdivisions of the plasma membrane (Soldati and Schliwa 2006), our analysis suggests that this capability has been largely amplified in animals, where the majority of cells are polarized, facing two different environments. It is possible that these morphological changes provided a fertile soil for adaptations of the endosomal SNARE repertoire. The differentiation of cells is initiated by genetic programs under the control of secreted morphogenic cues. Indeed, in the last few years, it has become clear that the endocytic pathways play important roles not only in secretion of these cues, but also in modulation of their signal transduction in the receiving cell (Gonzalez-Gaitan 2003; Rodriguez-Boulan et al. 2005; Emery and Knoblich 2006; Leibfried and Bellaiche 2007). In fact, endosomal SNARE proteins have been shown to be important for establishing cell polarity (Lu and Bilder 2005; Balklava et al. 2007). For example, the loss of the early endosomal SNARE protein Syx7 (in *Drosophila*, also referred to as avalanche) strongly resembles the phenotypes of *Drosophila* neoplastic tumor-suppressor mutations such as *scrib*, *dlg*, and *Lgl* (Hariharan and Bilder 2006; Humbert et al. 2006; Vasioukhin 2006; Wirtz-Peitz and Knoblich 2006; Chia et al. 2008). Intriguingly, one of these tumor-suppressor genes involved in establishment of cell polarity, Lgl, also emerged during the transition to multicellularity in animals by duplication of the regulatory SNARE tomosyn. Like tomosyn, Lgl has been shown to interact with the secretory SNARE apparatus, suggesting that both factors still function via homologous molecular pathways (Brennwald and Rossi 2007).

The road map of the endosomal compartments is rather elaborate as they serve as a sorting hub for cargo arriving from the exocytic and the endocytic pathway. It is conceivable that the increased collection of SNAREs resulted in a more complex network of endosomal trafficking routes in animals compared to their unicellular progenitors. However, it should be kept in mind that the exact role of several of the novel metazoan SNAREs is not entirely clear yet. In addition, the endosomal R-SNAREs seem not to be confined to one transport step only. For instance, Vamp7 (Ti-Vamp) not only mediates trafficking towards the lysosomal compartment, but it is also involved in secretion of lysosome-related organelles (Luzio et al. 2007). Comparably, endobrevin mediates trafficking towards late endosomes, but is also involved in granule secretion in exocrine tissues (Wang et al. 2004; Ren et al. 2007; Wang et al. 2007). It needs to be clarified in the future whether the participation of

R.III-type SNAREs in lysosomal secretion represents a new adaptation of these proteins in animals. However, considering that several other eukaryotic lineages do not possess a specific secretory R-SNARE (R.IV-type) but use R.III-SNAREs for secretion, it is likely that this ability of R.III-type of SNAREs rather reflects an ancient function of this group of proteins. Furthermore, the novel Qbc-SNAREs SNAP-29, and SNAP-47, are “exocytotic” by origin. However, their localization on intracellular membranes might also be consistent with functions in intracellular trafficking. Altogether, systematic cell biological investigations are necessary to test our predictions.

The second major expansion of the metazoan SNARE repertoire is restricted to the lineage of vertebrates, leading to their multifaceted set of secretory SNARE proteins. This expansion in vertebrates is accompanied by few domain rearrangements, but no losses of the original metazoan set were observed. Hence, our finding is in line with an earlier notion that the gene content of lower metazoans and vertebrates is highly conserved (Kortschak et al. 2003; Kusserow et al. 2005; Raible et al. 2005; Technau et al. 2005; Putnam et al. 2007; Miller and Ball 2008). This expansion of the SNARE repertoire in vertebrates can probably be attributed to two rounds of whole genome duplications (Ohno 1970; Holland 2003; Panopoulou and Poustka 2005). In line with observations on other factors, our data suggest that many other duplicated SNARE genes in vertebrates were not retained after whole genome duplications. This phenomenon is usually explained by extensive gene loss of duplicates (Holland 2003; Panopoulou and Poustka 2005). The fact that mostly duplicates of the secretory SNARE were retained, suggests that these factors presented a selective advantage during the rise of vertebrates. Often the novel secretory SNAREs are largely confined to different subcellular locations like the apical or basolateral membrane, or their expression is restricted to particular tissues or different developmental stages (for a review, see (Hong 2005; Jahn and Scheller 2006; Stow et al. 2006)). Hence, the extended set of secretory SNAREs appears to have facilitated the development of new secretory cell types in vertebrates.

It is very likely that not only the major expansions but also the losses of individual SNARE factors in various animal lineages affected the itinerary of transport vesicles as the changes probably reflect lineage-specific adaptations of the endomembrane system. Hence, one should be cautious when comparing the vesicular trafficking pathways of different animal species. Losses occurred for example in the invertebrates *D. melanogaster* and *C. elegans*, which are well-studied model organisms. Particularly in the repertoire of nematodes and platyhelminthes, we noticed the absence of some novel metazoan SNARE types involved in endosomal trafficking. As both phyla possess a somewhat differently reduced set of novel SNAREs, these losses seem to reflect secondary gene losses, rather than a more basal status of these phyla. Hence, it seems possible that comparable losses of several endosomal SNARE genes in nematodes and platyhelminthes are a sign of convergent adaptations of their endosomal sorting pathways caused by comparable life-style and feeding requirements. In addition, and in line with observations on other genes, the SNARE sequences from both phyla often form long but well-separated branches, supporting their more derived status. In fact, the relative position of these phyla in the phylogeny of animals is still debated (for discussion see e.g. (Philippe and Telford 2006)).

It is worth mentioning that we also observed a few differences in the SNARE repertoire of different vertebrate lineages. For example, the Qc.III genes Syx6 and Syx10, which are the products of a gene-duplication in vertebrates, participate in

different SNARE units that mediate trafficking of different types of vesicles from endosomes towards the TGN. Remarkably, Syx10 is secondarily absent from mouse and rat genomes, suggesting that a subtle difference in the endosomal trafficking pathways between murine rodents and other vertebrates exists (Ganley et al. 2008). The existence of differences in their endosomal pathways is corroborated by the fact that Syx20 (Qa.III.b), an endosomal syntaxin with unknown function, appears to have been lost in mice and rats as well.

Overall, the animal genomes examined in this work comprise a remarkably consistent and homologous set of SNARE proteins. As outlined above, the two major expansions can probably be attributed to two distinct periods of larger genome remodeling, during which several individual SNARE factors were duplicated. Yet, duplications of SNAREs, in particular of the ones involved in secretion, followed by functional diversification, occurred recurrently in different lineages. For example, we came across duplicated sets of secretory SNAREs in several invertebrates but also in another choanoflagellate, *Monosiga ovata*. The identification of lineage-specific duplications was possible because we have integrated sequence data from a large quantity of species that represent a very broad spectrum of the animal kingdom. In addition, we rigorously removed duplicates from our dataset. Furthermore, we have used a highly accurate classification and a detailed phylogenetic analysis. We noted that these methodological key elements were paid less attention to in earlier studies, explaining why these studies were unable to distinguish the two major expansions of the metazoan SNARE repertoire found by us (Dacks and Doolittle 2004; Yoshizawa et al. 2006; Dacks and Field 2007).

Although it was not the primary goal of this study, we noticed that the trees built from orthologous SNAREs generally recapitulate their presumed evolutionary relationships. In fact, the sequences of species that belong to the same phylum usually congregated. This is seen best in the trees of SNAREs involved in trafficking within the ER and the Golgi, since these factors are present usually as singletons in all animal genomes. These factors apparently underwent slow lineage-specific adaptations (Fig. 3). More drastic changes occurred in SNARE proteins involved in endosomal and secretory trafficking steps, probably, as discussed above, evoked by episodes of larger genome re-arrangements. For example, expansion of the set of secretory syntaxins (Qa.IV) in vertebrates led to a highly diversified set that obscures the underlying information of the phylogenetic relationships (Fig. 2d). Generally, the trees based on single sequences are not ideally suited to reflect the relationships between the larger phylogenetic groups of animals. In fact, the paralogous expansions render it challenging to assemble and align concatenated sequences to increase the resolution of the phylogenetic tree. Also, several animal lineages are still under-represented in the genome sequences that are currently available. Nevertheless, as new genome data can be quickly integrated, our detailed analysis of the evolutionary history of SNARE genes in animals is a promising supplementary tool for future phylogenetic studies.

Supplementary Material

The supplementary data of this study can be downloaded from the SNARE projects homepage (<http://bioinformatics.mpibpc.mpg.de/snare/>). The SNARE projects homepage allows access to the SNARE database (Kloepper et al. 2007), into which all newly classified sequences of this study have been integrated.

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Figure Legends

Fig. 1. Schematic depiction of the major changes in the SNARE repertoire during animal evolution.

a) Phylogenetic relationships between the major animal groups represented in this study. The tree is based on the current view of the phylogeny of metazoans, but several diversifications are still heavily debated (Philippe and Telford 2006). For example, the phylogenetic placement of placozoan *Trichoplax adhaerens* among the lower metazoans is not clear (Miller and Ball 2005; Schierwater 2005), although recent data strongly suggest that their position should be as the most basal extant lower metazoan (Dellaporta et al. 2006; Signorovitch et al. 2007). The numbers indicate the major changes in the SNARE repertoire uncovered in this study. The details of the changes indicated by the numbers 1 and 2 are illustrated in b); the other lineage-specific changes (3 through 7) are displayed in Suppl. Table. 2.

b) Schematic depiction of the major changes in the SNARE repertoire during the rise of multicellular animals and of vertebrates.

To illustrate the changes of the SNARE sets representative species with well-maintained genomes were used: *Monosiga brevicollis* for choanoflagellates, the cnidarian *Nematostella vectensis* for basal metazoans, and *Mus musculus* for vertebrates/tetrapods. The domain architecture of the SNARE proteins is drawn to scale. The highly conserved SNARE motif is depicted by a gray box; the adjacent transmembrane region by a black box. The degenerated SNARE motif of Lgl is depicted as a white box. The two β -propeller domains of tomosyn and Lgl are indicated as white ovals. Note that the C-terminal region of Syx6 from *M. brevicollis* lacks a transmembrane region and is shown as a dashed line to indicate that this region of the sequence most likely is not correctly annotated. The expansion of the SNARE set in vertebrates is probably caused by two rounds of whole genome duplications. Consequently, tetrapods can have up to four orthologous copies – although two or three copies are more often seen. Note that an enlarged set of regulatory R-SNAREs in vertebrates is not shown for space limitations. Notably, one has to assume that at least two Qa.IV-genes existed in the initial repertoire of the vertebrate ancestor to be able to account for the numerous novel Qa.IV genes in vertebrates, e.g. Syx1-4, Syx11 (Tang et al. 1998a; Prekeris et al. 2000), Syx19 (Wang et al. 2006), and Syx21. To account for this, the putative second Qa.IV-SNARE of the vertebrate ancestor is shown in gray. Interestingly, in few animal species (e.g. *Nematostella vectensis*), we found only a single set of secretory Qa- and R-SNAREs. This is comparable to many unicellular eukaryotes, for which it is thought that the single secretory set generally is capable of mediating constitutive and regulated secretion. Possibly, discrimination between both types of secretion can be imposed on this SNARE unit by additional factors. It is thus conceivable that even in multicellular organisms on the level of the SNARE factors, no strict separation between constitutive and regulated secretion exists.

Fig. 2. Outlines of unrooted phylogenetic trees of individual SNARE subgroups that have undergone major expansions in the evolution of animals.

Each tree is shown as a schematic outline. For each tree, a detailed figure is given at the SNARE projects homepage (<http://bioinformatics.mpibpc.mpg.de/snare/>). In addition, all individual phylogenetic trees, and the corresponding sequence alignments are available there. In each tree, the diverged metazoan SNARE types are shown by different colors. In addition, the major groups of animals are indicated. The labels at the major branches represent the likelihood mapping (left) and AU support values (right).

a) The endosomal R-SNAREs (R.III-type) of animals split into three major branches Vamp7, Vamp4 and endobrevin. The gene, Vamp7-like, which is only present in a few basal metazoans, is not especially indicated in the schematic tree, but can be seen in the enlarged version.

b) The endosomal Qa-SNAREs (Qa.III.b-type) of animals split into the three elementary branches, Syx7, Syx20, and Syx17. In the lineage of vertebrates, another type, Syx13, arose.

c) The Qbc-SNAREs (Qbc.IV-type) of animals basically split into three branches, SNAP-25, SNAP-29 and SNAP-47. Note that SNAP-47 contains an extended linker region compared to the other metazoan Qbc-SNAREs. In the lineage of vertebrates, another type, SNAP-23, emerged, but independent duplications of the SNAP-25-type occurred in other lineages as well.

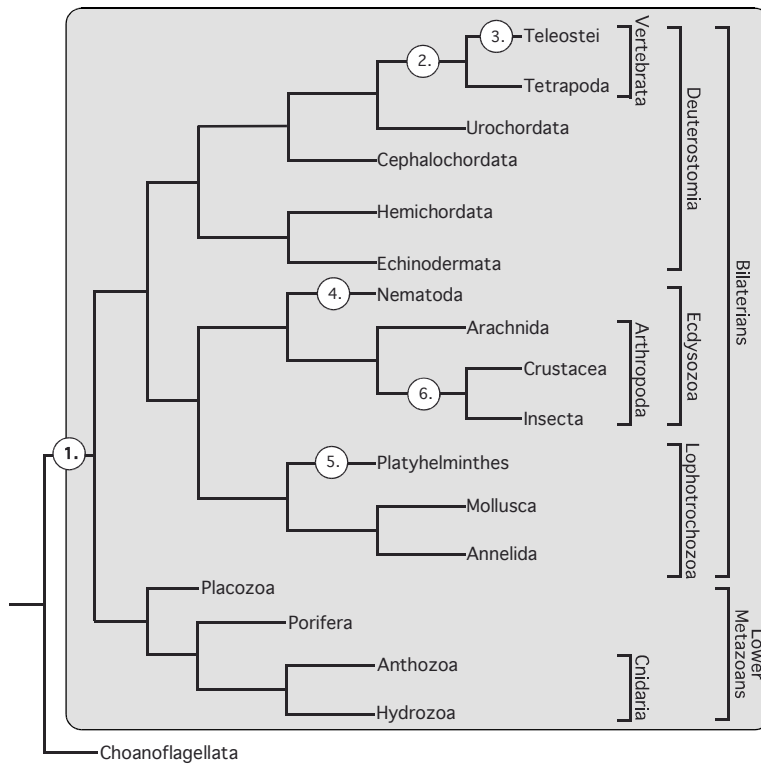
d) The phylogenetic tree of secretory syntaxins (Qa.IV-type) is dominated by the major expansion in the vertebrate lineage that gave rise to various new types. For example, both Syx11 and Syx19 belong to a more deviated vertebrate specific group of secretory syntaxins that, instead of a transmembrane region (TMR), possess a C-terminal cysteine rich sequence, which is probably palmitoylated.

Originally, metazoans appear to have contained only one or two isoforms of secretory syntaxins. For example, besides the highly preserved Syx1, several lineages possess a much more diverged secretory syntaxin (e.g. Syx4 of insects).

Fig. 3. The phylogenetic trees of individual SNAREs reflect the relationship of animal groups.

In metazoans, a new factor, Sec22-like, arose by gene duplication of the R-SNARE Sec22 (R.I-type) followed by domain rearrangements. Sec22-like has lost its R-SNARE motif, but possess, three consecutive TMRs. **(a)** The schematic overview of the unrooted phylogenetic tree of metazoan Sec22 subfamily shows that the two factors are well separated. Sec22-like appears to have been lost again in different animal lineages, whereas Sec22, besides a duplication in teleost fishes, has been retained as a singleton. The phylogenetic tree of Sec22 reflects the evolutionary relationships among the larger animal groups well. For example, Sec22 from fishes lies on the vertebrate branch, but is separated from tetrapod sequences. Comparably, Sec22 sequences from insects are well confined, despite the enormous morphological diversity of this animal group. **(b)**. A close-up view of this branch shows that various insect groups are also well separated.

A



B

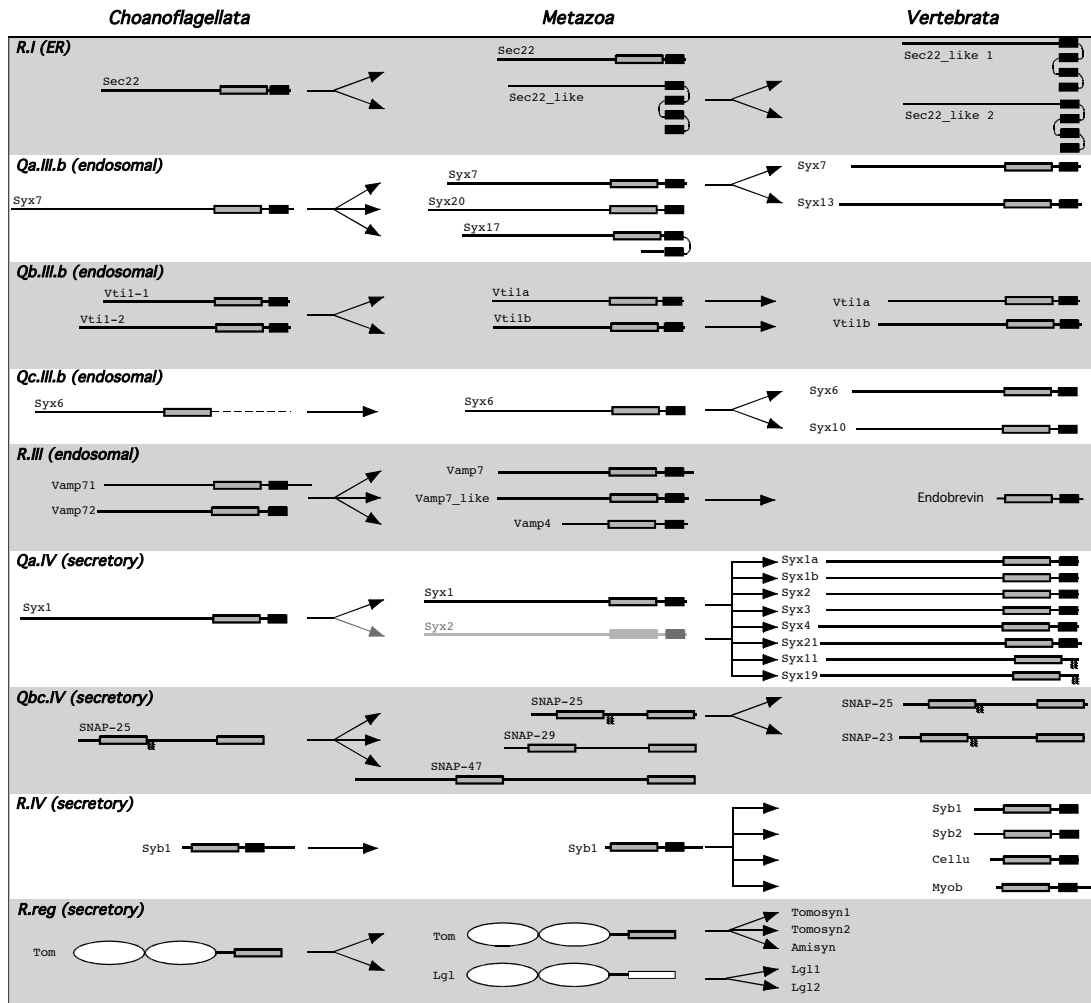


Fig1- Kloepper et al

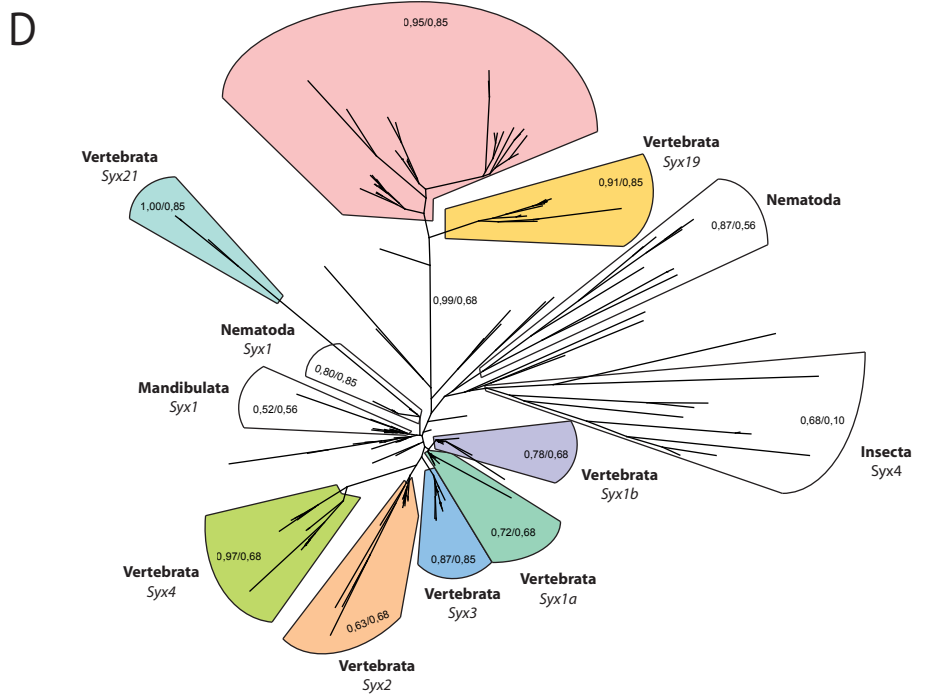
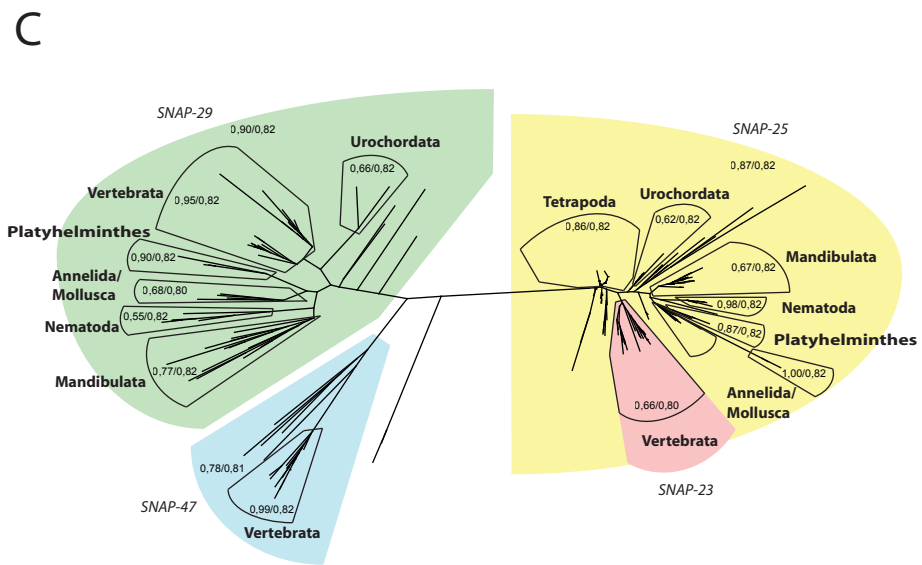
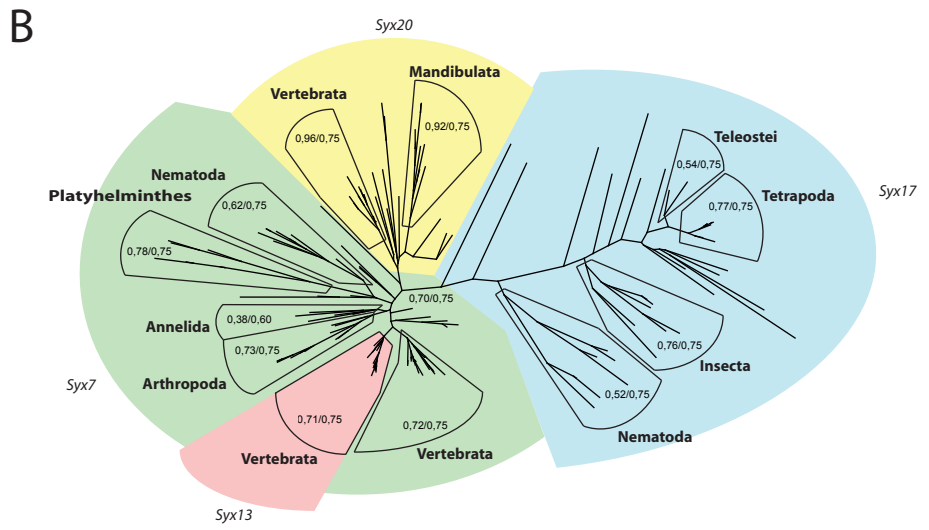
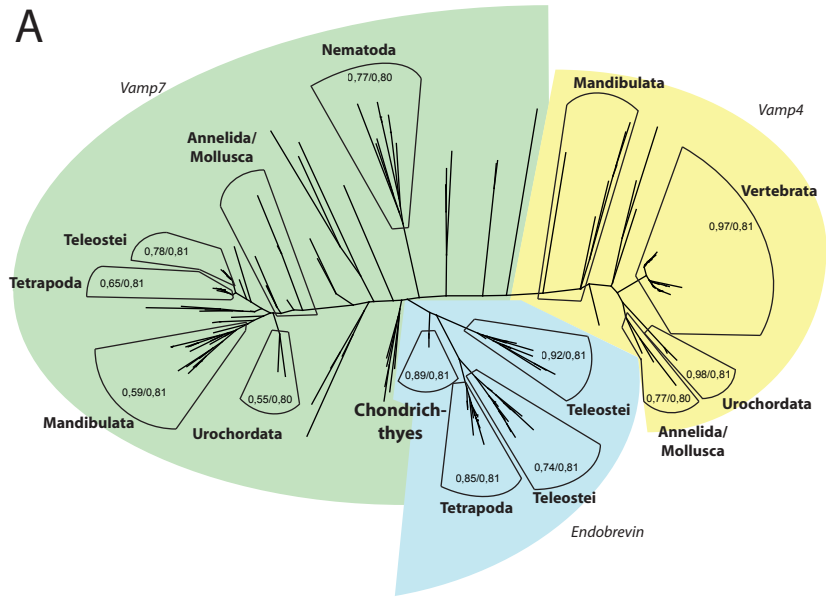
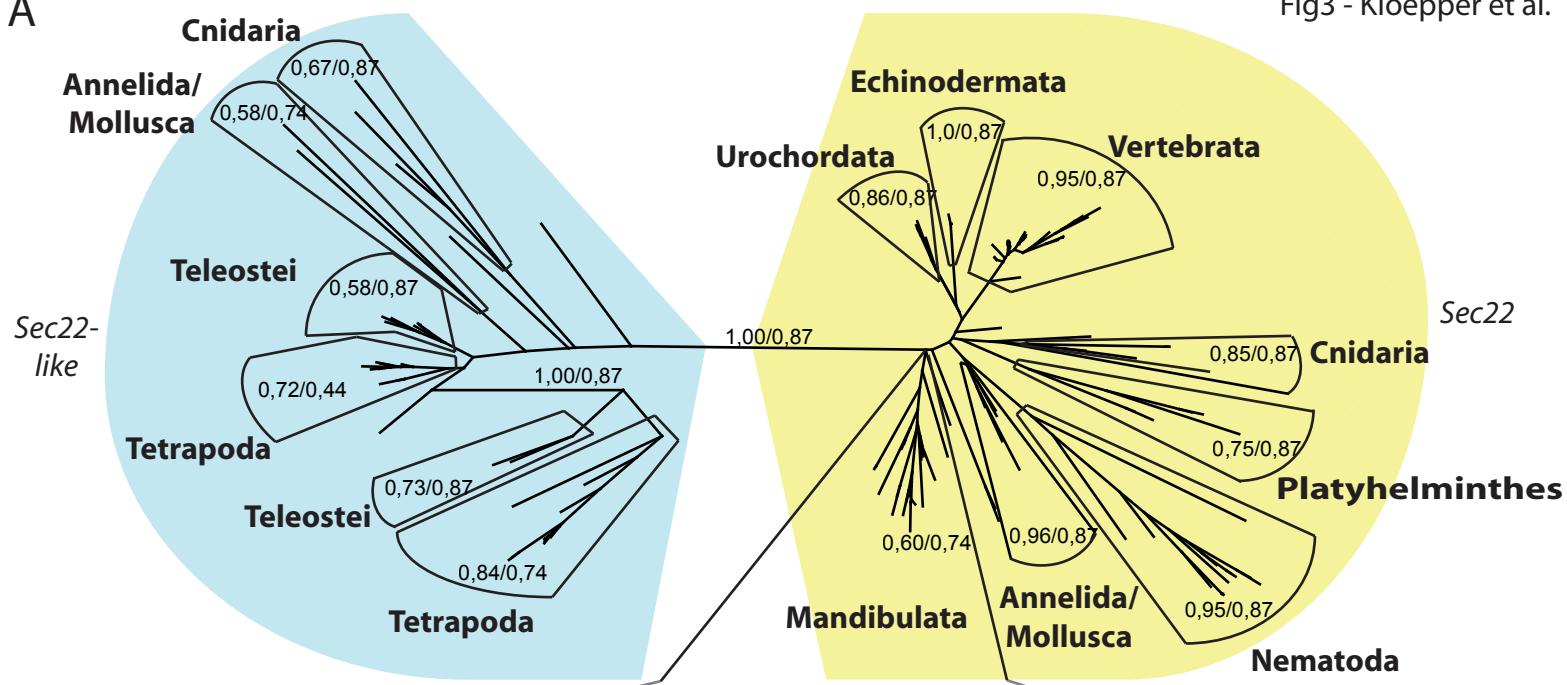
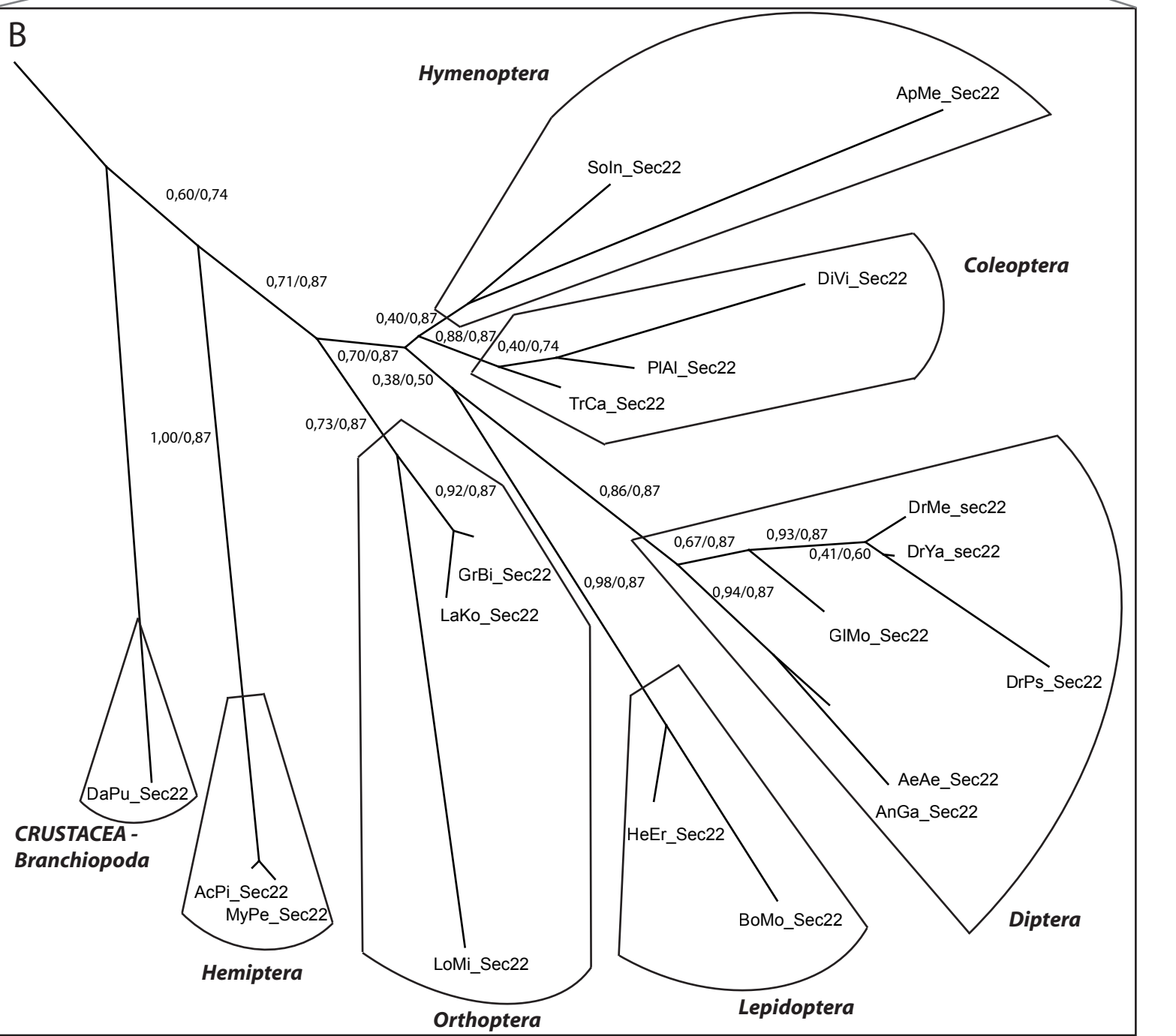


Fig2 - Kloepper et al.

A



B



Supplemental Data

SNAREing the basis of multicellularity: Consequences of protein family expansion during evolution

Tobias H. Klöpper, C. Nickias Kienle, and Dirk Fasshauer

The supplementary data of this study can be downloaded from the SNARE projects homepage (<http://bioinformatics.mpibpc.mpg.de/snare/>). The SNARE projects homepage allows access to the SNARE database (Klopper et al., 2007), into which all newly classified sequences of this study have been integrated.

The supplementary data contain detailed version of the trees depicted in Figures 2a-d and 3a (i), the supplementary tables 1 and 2 (ii), and the generated Nexus files and Alignments of all SNARE types (iii).

i) The following detailed schematic trees of Figures 2a-d and 3a can be downloaded:

Suppl. Fig. 1. Detailed schematic depiction of the unrooted phylogenetic trees of the endosomal R-SNAREs (R.III-type) shown in Fig 2a.

Suppl. Fig. 2. Detailed schematic depiction of the unrooted phylogenetic trees of the endosomal Qa-SNAREs (Qa.III.b-type) shown in Fig 2b.

Suppl. Fig. 3. Detailed schematic depiction of the unrooted phylogenetic trees of the Qbc-SNAREs (Qbc.IV-type) shown in Fig 2c.

Suppl. Fig. 4. Detailed schematic depiction of the unrooted phylogenetic trees of the secretory syntaxins (Qa.IV-type) shown in Fig 2d.

Suppl. Fig. 5. Detailed schematic depiction of the unrooted phylogenetic trees of the secretory syntaxins (R.I-type) shown in Fig 3a.

ii) The following supplementary tables can be downloaded:

Suppl. Table 1. List of all SNARE sequences used.

Suppl. Table 2. List of the SNARE sets of 55 different species, highlighting important changes during the evolution of distinct animal lineages.

The two major expansion, of the SNARE repertoire (i.e. during the transition to multicellularity in animals and during the rise of vertebrates) are indicated in an

additional row, in which the new SNAREs are highlighted in light green. In addition, putative losses of distinct SNARE genes are highlighted in light orange. The additional round of duplication of distinct SNARE genes in bony fishes is highlighted in light yellow.

iii) The following 17 phylogenetic trees in Nexus file format and their corresponding alignments can be downloaded:

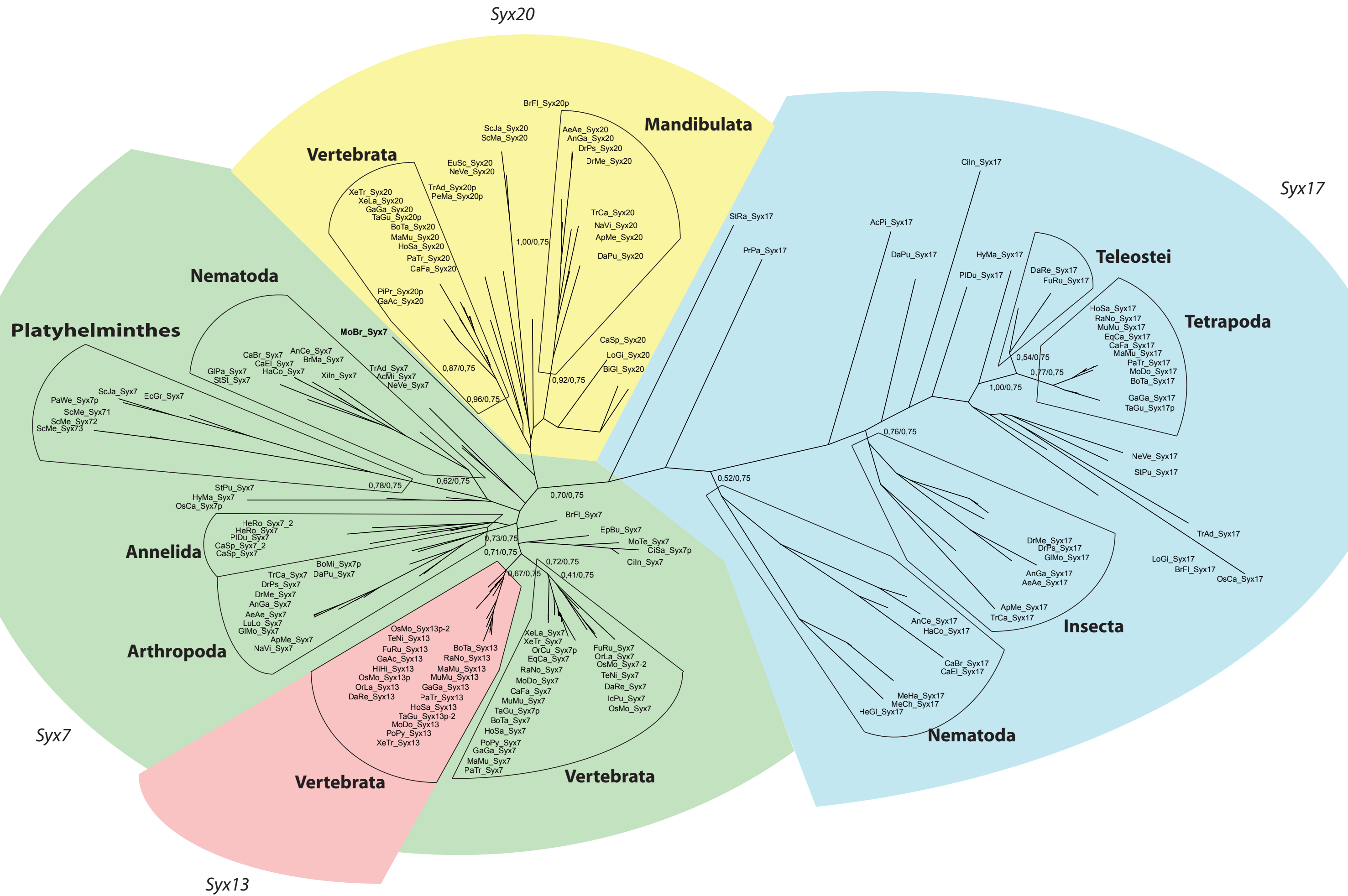
<i>SNARE subgroups</i>	<i>Common names of the basic metazoan SNARE types included in analysis</i>
Qa.I	Syx18
Qa.II	Syx5
Qa.III.a	Syx16
Qa.III.b	Syx7, Syx17, Syx20
Qa.IV	Syx1
Qb.I	Sec20
Qb.II	Bos1, Gos1
Qb.III	Vti1a, Vtila
Qc.I	Use1
Qc.II	Bet1, Gs15
Qc.III	Syx6, Syx8
SNAP (Qbc.IV)	SNAP-25, SNAP-29, SNAP-47
R.I	Sec22, Sec22-like
R.II	Ykt6
R.III	Vamp7, Vamp7-like, Vamp4
R.IV	Syb
R.Reg	Tomosyn, Lgl

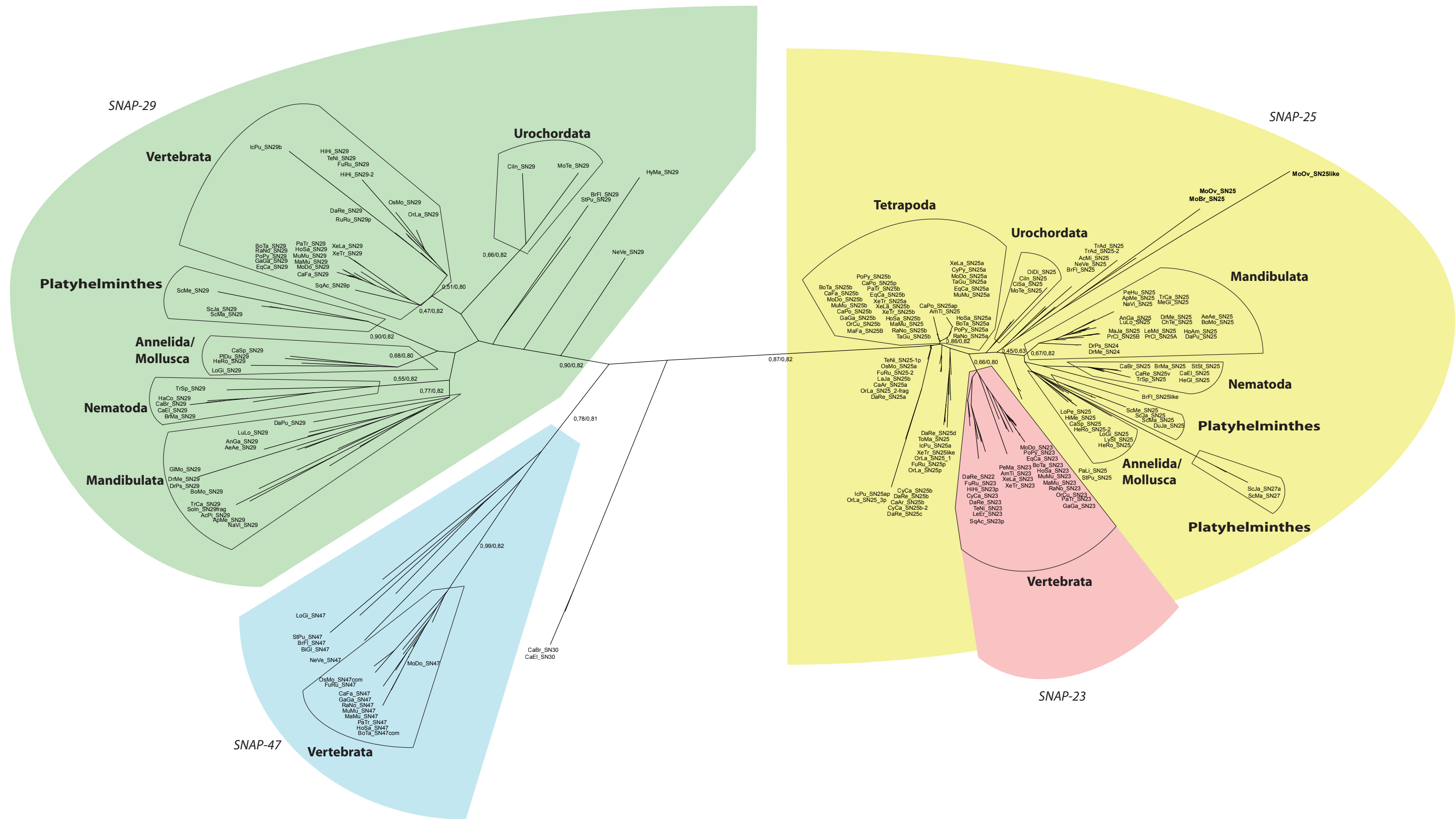
Syx20

Syx17

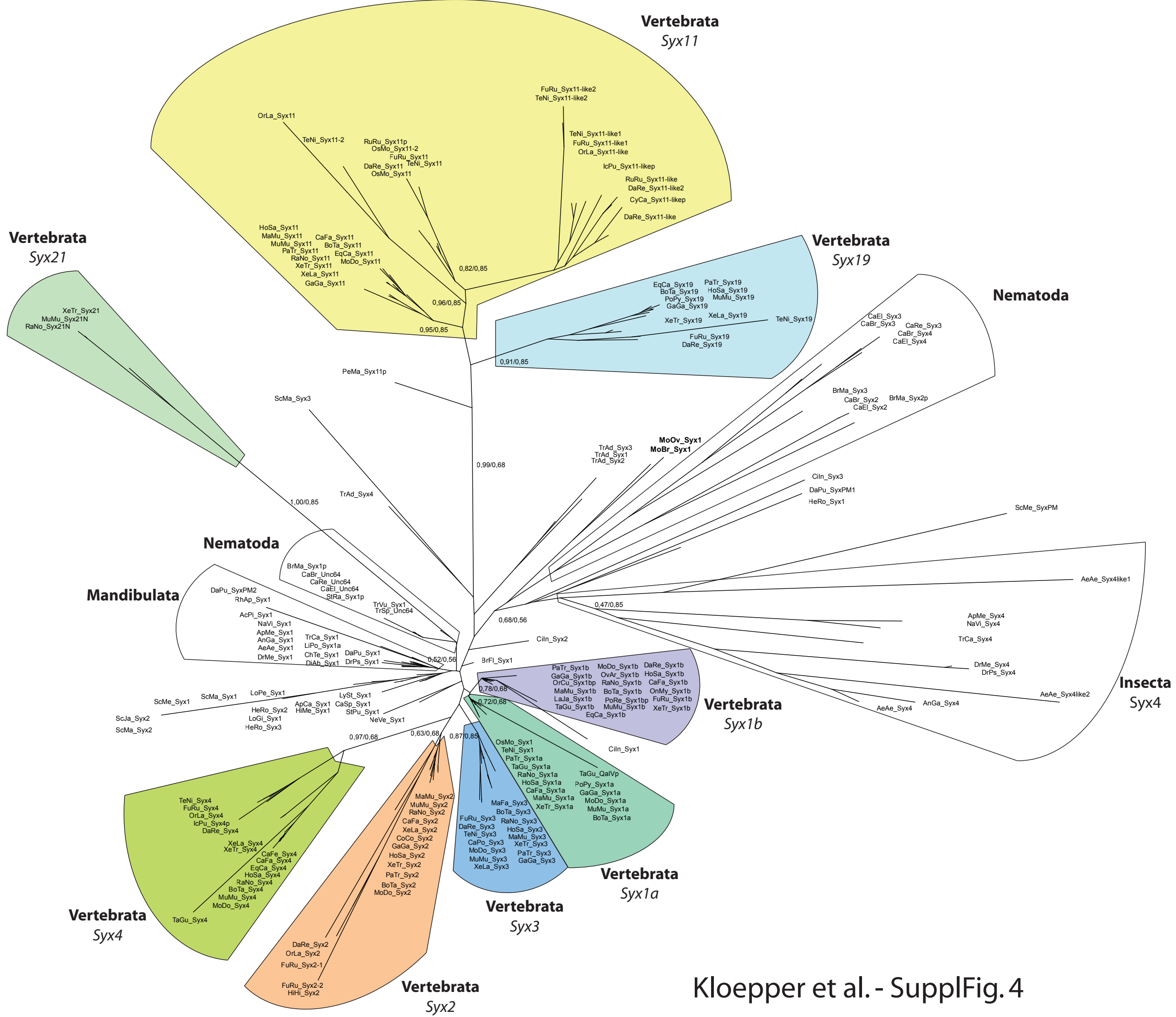
Syx7

Syx13





Kloepper et al. - SupplFig. 3



Kloepper et al. - SupplFig.4

