The backbone of the post-synaptic density originated in a unicellular ancestor of choanoflagellates and metazoans

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Abstract

**Background:** Comparative genomics of the early diverging metazoan lineages and of their unicellular sister-groups opens new window to reconstructing the genetic changes which preceded or accompanied the evolution of multicellular body plans. A recent analysis found that the genome of the nerve-less sponges encodes the homologues of most vertebrate post-synaptic proteins. In vertebrate excitatory synapses, these proteins assemble to form the post-synaptic density, a complex molecular platform linking membrane receptors, components of their signalling pathways, and the cytoskeleton. Newly available genomes from *Monosiga brevicollis* (a member of Choanoflagellata, the closest unicellular relatives of animals) and *Trichoplax adhaerens* (a member of Placozoa: besides sponges, the only nerve-less metazoans) offer an opportunity to refine our understanding of post-synaptic protein evolution.

**Results:** Searches for orthologous proteins and reconstruction of gene gains/losses based on the taxon phylogeny indicate that post-synaptic proteins originated in two main steps. The backbone scaffold proteins (Shank, Homer, DLG) and some of their partners were acquired in a unicellular ancestor of choanoflagellates and metazoans. A substantial additional set appeared in an exclusive ancestor of the Metazoa. The placozoan genome contains most post-synaptic genes but lacks some of them. Notably, the master-scaffold protein Shank might have been lost secondarily in the placozoan lineage.

**Conclusions:** The time of origination of most post-synaptic proteins was not concomitant with the acquisition of synapses or neural-like cells. The backbone of the scaffold emerged in a unicellular context and was probably not involved in cell-cell communication. Based on the reconstructed protein composition and potential interactions, its ancestral function could have been to link calcium signalling and cytoskeleton regulation. The complex later became integrated into the evolving synapse through the addition of novel functionalities.

Background

Many of the complex traits that distinguish the organisation and biology of multicellular animals originated during the early phase of their evolutionary history. Among these, the emergence of the neuro-sensory system represents a particularly fascinating conundrum, because the biological support of animal perception and behaviour is extraordinarily complex at the molecular, cellular and anatomical scales, and without equivalent in the living world. A promising path to understanding the genetic changes that shaped the body plan of multicellular animals has been recently opened by comparative analyses of genomes from the early-diverging animal lineages and the closest unicellular relatives of metazoans (e.g. [1-10]).

The recent characterisation of a nearly complete set of post-synaptic protein orthologues in the genome of the sponge *Amphimedon queenslandica* [2] represents a successful illustration of this approach. Excitatory synapses in the vertebrate central nervous system are characterised by an electron-dense structure called the post-synaptic density (PSD) below the post-synaptic membrane. Proteomic investigations have identified about one thousand proteins in this complex [11-14]. Some of the proteins encoded by the sponge genome were found to be homologous to vertebrate PSD proteins, suggesting a common origin. However, the evolutionary history of these proteins is not well understood.

The recent availability of genome sequences from *Monosiga brevicollis* (a choanoflagellate) and *Trichoplax adhaerens* (a placozoan) offers an opportunity to refine our understanding of post-synaptic protein evolution. These organisms are the closest unicellular relatives of animals and have nerve-less bodies. The genome of the nerve-less sponge *Amphimedon queenslandica* encodes the homologues of most vertebrate post-synaptic proteins, suggesting that these proteins evolved in a unicellular context.

**Results:** Searches for orthologous proteins and reconstruction of gene gains/losses based on the taxon phylogeny indicate that post-synaptic proteins originated in two main steps. The backbone scaffold proteins (Shank, Homer, DLG) and some of their partners were acquired in a unicellular ancestor of choanoflagellates and metazoans. A substantial additional set appeared in an exclusive ancestor of the Metazoa. The placozoan genome contains most post-synaptic genes but lacks some of them. Notably, the master-scaffold protein Shank might have been lost secondarily in the placozoan lineage.

**Conclusions:** The time of origination of most post-synaptic proteins was not concomitant with the acquisition of synapses or neural-like cells. The backbone of the scaffold emerged in a unicellular context and was probably not involved in cell-cell communication. Based on the reconstructed protein composition and potential interactions, its ancestral function could have been to link calcium signalling and cytoskeleton regulation. The complex later became integrated into the evolving synapse through the addition of novel functionalities.
these proteins (e.g. Shank, DLG, Homer, GRIP) have pivotal roles in building a scaffold for the anchorage of the other PSD components, for instance membrane receptors for neurotransmitters, cation channels, members of intracellular signalling pathways, and cytoskeleton or cytoskeleton regulation proteins [15-18]. This complex sub-membrane plateform thereby connects cell surface receptors with components of signal transduction pathways and with the cytoskeleton machinery. The PSD is also involved in synaptogenesis, dendritic spine formation and synaptic plasticity [17,19,20]. The three-dimensional architecture of the PSD is determined by multiple protein-protein interactions through binding domains (e.g. [17,18,21]). For example, the vertebrate Shank protein, the master scaffold protein of the PSD [18], contains five interacting domains (from N-ter to C-ter: ankyrin repeats, SH3 domain, PDZ domain, proline-rich region, SAM domain) involved in a huge variety of homomeric and heteromeric interactions. Among PSD scaffold proteins, PDZ (for Post-synaptic density) (for Post-synaptic density protein 95/Drosophila disc large tumor suppressor/ZO-1) is the most common type of binding domain, with many proteins containing multiple PDZ domains (e.g. three in the DLG protein; seven in GRIP).

Not only were most PSD proteins identified in the sponge genome (whereas most are absent in fungal or plant genomes), but moreover these proteins essentially contain the same interacting domains arranged into the same architectures as in their vertebrate counterparts [2]. Intriguingly, several of the sponge PSD-like genes (DLG, GKAP, GRIP, HOMER, CRIP) were found co-expressed in a particular non-ciliated epidermal cell type of the Amphimedon larva, the globular cell (erroneously called "flask cells" in [2], see [22]). These observations are consistent with the existence in sponges of an assembled structure similar to the vertebrate PSD, despite absence of recognisable nerve cells and synapses.

The PSD protein complex would therefore have originated before the acquisition of synapses in a nerve-less ancestor, at least under the conventional view that sponges branched off earlier than animals with a nervous system in the phylogeny (e.g. [23-27]). Moreover, since the sponge globular cells express Notch/Delta homologues and a proneural-like bHLH transcription factor [22] in addition to PSD genes, they might represent putative sensory "proto-neural" cells responding to environmental stimuli [22] and a possible "starting point" for the evolution of neural cells [2]. Some recent molecular phylogenies [28-31] have pointed to more complicated evolutionary scenarios (e.g. secondary loss of nerve cells and synapses in sponges) by challenging the monophyly of animals with a nervous system (Cnidaria, Ctenophora and Bilateria, classically grouped under the Eumetazoa) and the early divergence of sponges (discussed in [32,33]). However, a recent phylogenomic study with increased species sampling of non-bilaterian lineages and sophisticated model of sequence evolution [34] lent support to the conventional phylogeny and thereby to a single acquisition of nerve cells and synapses in a eumetazoan ancestor.

Since the publication of [2], additional annotated complete genomes have become available. The present study aims at improving our understanding of the emergence of the PSD protein complex by focusing on sequence data from Monosiga brevicollis and Trichoplax adhaerens, two organisms of particular interest with respect to early metazoan evolution. Monosiga is a member of choanoflagellates, the unicellular sister-group of the Metazoa [1,8]. Previous studies have suggested (without rigorous analyses) the existence of some PSD protein orthologues in the Monosiga genome [2], but these have been considered "very few" [10], supporting the idea that the PSD scaffold is essentially an innovation of the metazoan lineage. However, this remains to be tested rigorously through systematic analyses of choanoflagellate orthologues of the post-synaptic genes. Trichoplax (phylum Placozoa) is a nerve-less animal whose flat and asymmetric body comprises two epithelial cell layers enclosing a loose formation of interconnected fibre cells sandwiched in between [35]. The phylogenetic placement of placozoans remains highly contentious (e.g. [28,34,36]). Analyses of the Trichoplax genome have revealed a degree of genetic complexity that was totally unexpected given its morpho-anatomical simplicity [36]. As the only non-sponge animal lacking a nervous system, Trichoplax is clearly of pivotal importance for understanding the origin of neural cells and synapses.

The gene set investigated in the present study comprises the main scaffold proteins of the PSD, and their more significant interacting partners as recognised in the recent literature (e.g. [15-18]). It is essentially the gene set analysed in [2]. Similarity searches for PSD proteins were conducted against the genomes of M. brevicollis, T. adhaerens, several non-metazoan eukaryotes (as outgroups) and a representative sampling of metazoan species. Our analyses provide evidence for the presence of a substantial subset of functionally important scaffold PSD proteins in the common ancestor of choanoflagellates and metazoans, and for secondary loss of the master scaffold protein Shank in placozoans.

Methods
Data collection
The protein set examined in this study is listed in Fig. 1 and Additional file 1. It is the same as in ref [2], except that we discarded GABA-B receptor, Tamalin and LIM kinase (because we were unable to find evidence from the literature for physical interaction of these proteins
with the PSD scaffold), as well as Kir and Mint (the latter is a presynaptic adaptor protein [37]; the former is a large family of inwardly rectifying potassium channels which is not particularly associated with the PSD, see [38]). In addition to the full protein alignments, we built a separate alignment for PDZ domains (as in ref [2] but with an extended gene sampling) and constructed specific alignments for the SAM and SH3 domains of Shank. The starting data set [2] comprised sequences from Saccharomyces cerevisiae (Ascomycota, Fungi), plants (Arabidopsis thaliana and Oryza sativa), Dictyostelium discoideum (Amebozoa), Amphimedon queenslandica (Demospongiae, Porifera), Neotomastella vectensis (Anthozoa, Cnidaria), Drosophila melanogaster (Hexapoda, Arthropoda) and Homo sapiens. These files were updated by TBLASTN searches (using proteins from Drosophila melanogaster as input sequences) on the most up-to-date collections of predicted transcripts from plants (Arabidopsis thaliana and Oryza sativa), Dictyostelium discoideum (Amebozoa), fungi (Batrachocephalum dendrobatidis, Saccharomyces cerevisiae, Neurospora crassa, Ustilago maydis, Rhizopus oryzae), Monosiga brevicollis (Choanoflagellata), Trichoplax adhaerens (Placozoa), cnidarians (Hydra magnipapillata, Hydrozoa, and Neotomastella vectensis, Anthozoa), and Capitella sp. (Lophotrochozoa, Annelida). An E-value threshold was determined empirically for each protein family by using the E-value associated with the closest paralogue retrieved from Blast of the fly protein against the fly genome (see Additional file 2: E-values used for each family, with names and accession numbers of the closest fly paralogues).

Alignments and phylogenetic analyses

Sequences retrieved from Blast searches were automatically integrated into the corresponding alignments and aligned using CLUSTALW, implemented in our in-house automated Blast and alignment pipeline. Alignments were slightly corrected manually for elimination of major mistakes. Ambiguous regions were identified by visual inspection and removed manually and positions containing more than 80% of missing data were deleted. When several non-overlapping partial sequences were present in the same alignment, the alignment was split into several partitions to avoid artificial grouping in the trees of non-overlapping sequences (observed in preliminary Maximum Likelihood analyses). These partitions, labelled “A, B, C, ...” (see Additional file 1), were analysed separately. The alignments are given in Additional file 3.

Maximum-Likelihood (ML) analyses were performed using the PhyML program [39], with the WAG model of amino-acid substitution and a BioNJ tree as the input tree. A gamma distribution with four discrete categories was used in these ML analyses. The gamma shape parameter and the proportion of invariant sites were optimised during the searches. Branch support was tested with bootstrapping (100 replicates). Bayesian analyses were done with Mr Bayes under the WAG model (prset=invgamma). For each data set, two searches were run in parallel for 500,000 generations, representing 20 times the number of generations discarded as burn in (first 25,000 trees, determined by stabilisation of the likelihoods). We estimated convergence by checking the “average standard deviation in partition frequency values across independent analyses”, using a threshold value of 0.05. Since for most alignments the number of phylogenetically informative positions is low with respect to the number of sequences, trees contain a high number of unstable nodes and therefore it was not possible to constrain searches for convergence with a lower threshold value. In some cases, this threshold of 0.05 was not reached after 500,000 generations (and concomitantly stationarity of the likelihoods was not reached after 25,000 generations). Then, burn-in was expanded and search was prolonged until the index had dropped below 0.05. Posterior probabilities were estimated by constructing a majority-rule consensus of trees sampled every 100 generations. Trees were rooted as in ref [2].

After preliminary (not shown) phylogenetic analyses, all (protein/species) absences were further tested by additional Blast searches on the full (when possible, assembled) nucleotide genome sequences of the corresponding species, with a very relaxed E-value (10) to ensure that these genes have not been missed due to limitations of the initial Blast searches (i.e. the use of predicted transcript data sets, or too strict E-values). In several cases, these genomic searches detected protein orthologues that had been missed during initial searches on predicted transcripts (e.g. IP3R in Hydra magnipapillata; PMCA in Capitella sp.). In these cases, protein alignments were updated using the procedure described above, prior to final phylogenetic analyses (trees shown in Additional file 4).

Orthology assignments

An orthology group is here defined as the clade containing all sequences that are more closely related to the fly and/or human sequence(s) of reference than to any paralogue of ancient origin (i.e. resulting from duplication before the last common ancestor of bilaterians). Our orthology assessments therefore rely primarily on gene tree topologies, and the yellow fields in Fig. 1 indicate the presence of an orthologue, based on the corresponding gene tree. Three additional criteria were taken into account to estimate confidence for these orthology assignments: (i) support values (ML bootstrap support >= 70% and/or Bayesian posterior probability >= 0.95) (“b” in Fig. 1), (ii) congruence between separate analyses of partitions of the same protein alignment (labelled A,
B, ..., see above) or between different domains of the same protein ("c" in Fig. 1), (iii) comparison of domain architectures reconstructed using Smart http://smart.embl-heidelberg.de/ and Pfam http://pfam.sanger.ac.uk/search. Domain architecture was considered as conserved ("d" in Fig. 1) if the main functional domain(s) existing in the human and fly proteins were present and showed similar arrangement. These domain reconstructions are shown in Additional file 5. They were systematically done for the Monosiga and Trichoplax putative orthologues of human PSD proteins. In addition, we compared the five C-ter residues of PDZ-binding proteins in search of PDZ ligand conservation (Additional file 6), but the extent of sequence variability observed among PDZ ligands across taxa and proteins is so high that in practice it is generally impossible to conclude either for or against presence of a PDZ ligand. In addition, intron positions and frames were determined for the Shank proteins using NCBI sequence viewer (human and fly proteins) or JGI browse tool (proteins from Capitella, Nematostella and Monosiga) and the alignment procedure described above.

Reconstruction of protein acquisitions and losses onto the taxon phylogeny

Presence/absence of the PSD proteins across our species sampling was formalised into a taxon/character matrix. These characters were mapped onto the taxon phylogeny under parsimony and a Dollo model of gene gain/loss (each protein has been acquired only once - no convergence - and proteins can be lost but not re-acquired - no reversion). We used the taxon phylogeny of [34], as it is the most comprehensive phylogenomic study of basal metazoan relationships to date. This means in particular that our interpretation of the data relies upon a hypothesis of eumetazoan monophyly (here represented by Cnidaria + Bilateria). However, basal metazoan relationships remain controversial (see alternative views in [28,29,31]), and the reconstruction may differ under alternative topologies. Since the relationships between sponges, Trichoplax and eumetazoans (here represented by cnidarians and bilaterians) were unsupported in ref [34], the reconstruction was done under the three possible tree topologies.

Results and Discussion

Reconstructing evolutionary acquisitions and losses of the major PSD components

Starting from the PSD protein and domain alignments of ref [2] we have extended the taxonomic coverage by adding genes retrieved from the complete genomes of several additional fungal species, the choanoflagellate Monosiga brevicolis, the placozoan Trichoplax adhaerens, the cnidarian Hydra magnipapillata and the annelid Capitella sp. Protein/domain orthology was
determined based on gene trees (Additional file 4) and comparison of domain structures (Additional file 5). Results are summarised in Fig. 1, with a more detailed overview in Additional file 1. The most parsimonious reconstruction of protein acquisitions and losses is shown in Fig. 2 (see less parsimonious reconstructions obtained for alternative placements of Trichoplax, in Additional file 7). The inferred order of emergence of the genes is indicated by a colour code in Fig. 2 and Fig. 3. In Fig. 2, gene gains/losses are labelled in bold when orthology is supported by at least two criteria of confidence (in addition to gene tree topology; see Fig. 1 and Methods); otherwise they are labelled in normal font.

**PSD components originated in two main steps**

Protein acquisitions are concentrated in the two branches preceding and following the divergence of choanoflagellates in the opisthokont phylogeny (Fig. 2). A significant protein set originated in an exclusive ancestor of choanoflagellates and metazoans. This includes the scaffold proteins Shank, Homer and DLG, and the other PSD components CamKII, Citron, Cortactin, Ephrin R, IP3R, PKCaβγ and Shaker K⁺ channels. Another set, including the scaffold proteins CASK, GKAP and S-SCAM (= Magi), and the other PSD components α and δ-catenin, classical Cadherins, NOS, LIN-7, PICK1 and SPAR, is synapomorphic for the Metazoa. Very few proteins are of more ancient origin: PMCA (present in all investigated eukaryotes), β-catenin (present in plants and fungi) and CRIPT (present in plants).

A significant implication is that several core components of the mammalian PSD (red colour in Fig. 3) originated not only before the acquisition of synapses and nerve cells (absent in sponges and placozoan), but even before the acquisition of multicellularity, given that the choanoflagellate-metazoan ancestor was unequivocally unicellular [1,8]. Notably, the scaffold proteins Shank, Homer and DLG/PSD95, known in mammalians to structure the post-synaptic density by linking a huge variety of target proteins (e.g. membrane receptors, components of signal transduction pathways, cytoskeleton-interacting proteins, etc.), originated before the divergence of the choanoflagellates. In our analyses, the orthology of Homer and DLG between choanoflagellates

![Figure 2 Gains (coloured dashes) and losses (ellipses) of post-synaptic proteins reconstructed onto the taxon phylogeny under the parsimony criterion.](image-url)
and metazoans is well supported by the corresponding gene trees, the conservation of their domain structure, high bootstrap support (for Homer) and the congruence between the evolutionary histories of the domains (for DLG) (see Fig. 1).

**Shank emerged before the divergence of choanoflagellates**

The Shank protein is considered as the “master-scaffolding” molecule of the PSD [17,18]. Our PDZ tree identifies a putative choanoflagellate orthologue of metazoan Shank PDZs (PDZ tree in Additional file 4). This orthology assignment is not supported by statistical indices, but several additional lines of evidence argue in favour of orthology. The *Monosiga* protein that contains the putative Shank PDZ shows the combination of protein-interacting domains typical of vertebrate Shanks (N-ter ankyrin repeats, a SH3 domain, a PDZ domain, and a SAM C-ter domain), a domain structure that is unique among vertebrate proteins. Reciprocal Blast searches support orthology between the ankyrin repeats present in this *Monosiga* protein and those of metazoan Shanks (not shown - note that phylogenetic analyses of ankyrin repeats are not feasible). In addition, the putative *Monosiga* Shank protein shares with metazoan Shanks the presence of a conserved N-terminal 126-amino-acids region (upstream of the Ankyrin repeats) with no similarity to any other protein sequence (not shown - result of Blast search against NCBI referenced proteins) (see alignment of this N-terminal region in Additional file 8). Within this region, there are two introns with position conserved down to the phase and present in metazoan Shanks as well as in the *Monosiga* protein (red boxes in the alignment, Additional file 8). Finally, we analysed SH3 domains and found that the SH3 domain of the *Monosiga* putative Shank falls in a clade that contains all metazoan Shank SH3s (and two non-Shank SH3s, from *Drosophila* and *Hydra*) (see SH3 tree in Additional file 4).

These observations indicate that *Monosiga* has a *bona fide* orthologue of Shank. Although there is no experimental evidence that in the choanoflagellate cell Shank acts like its mammalian orthologues, the conserved domain combination suggests comparable scaffold properties. In addition, some of the proteins known in mammals to interact directly with Shank (including Homer, the IP3 receptor, and Cortactin) are also present in *Monosiga*.

The history of Shank has been slightly more complex in the details. Our phylogenetic analyses suggest that the SAM C-ter domain of *Monosiga* Shank might not be orthologous to the SAM domains present in some (but not all) metazoan Shanks (see SAM tree in Additional
file 4). A SAM domain might thus have been added independently in choanoflagellate and metazoan Shanks. This convergence event could have occurred by chance, or it may have been favoured by similar functional constraints acting on the Shank protein. A second peculiarity of choanoflagellate Shank is that its SH3 domain is located between PDZ and SAM, whereas in metazoans it is placed between the Ank repeats and the PDZ domain, indicating that a domain inversion took place in one of these two lineages. We compared intron positions along the Shank proteins of metazoans and Mono-
siga without finding evidence for or against this hypothesis of domain inversion (Additional file 8).

Evolution of the PSD protein repertoire within the Metazoa

Compared to the gene complement present in eumeta-
zooan genomes, different gene subsets happen to be absent in the two nerve-less species Amphimedon queens-
slandica (sponge) and Trichoplax adhaerens (placozoan). This makes the reconstruction of acquisition/losses in this region of the tree highly sensitive to the phylogenetic placement of placozoans with respect to sponges and eumetazoans. Trichoplax lacks orthologues of Shank (present in Monosiga, sponges and eumetazoans), GRIP, ErbB R, mGluR and SynGAP (present in sponges and at least some eumetazoans). If Trichoplax is the sister-
group to other metazoans (Fig. 2), then GRIP, ErbB R, mGluR, and SynGAP were absent in the metazoan ancestor and are synapomorphic for sponges + eumetazoans (brown colour in Fig. 2 and 3). The same taxon phylo-
geny implies secondary loss of α-catenin and CASK (pre-
sent in Trichoplax and in eumetazoans) in the sponge lineage. Alternative taxon phylogenies (e.g. Trichoplax sister-group to eumetazoans as in [34]) imply secondary losses of GRIP, ErbB R, mGluR and SynGAP in placzo-
avs (Additional file 7). Whatever the position of placzo-
avs, the parsimony optimisation suggests secondary loss of Shank in Trichoplax adhaerens.

The apparent loss of Shank in placzoans is particu-
larly puzzling as Shank is the main protein responsible for the higher-order structure of the PSD in mammals [18]. However, alternative scenarios cannot be ruled out at this stage, e.g., Shank could have been over-
looked due to incomplete coverage of the T. adhaerens genome (currently 8×), or the gene might have diverged in the placozoan lineage to the extent that it is no longer recognisable, or it might have been lost in Trichoplax adhaerens but not in other placzoan spe-
cies. If confirmed by future studies, loss of Shank in placzoans would imply disruption or at least consid-
erable alteration of the macromolecular architecture of the PSD-like protein complex. In addition, the Tricho-
plax situation is somewhat difficult to understand when it is realised that Homer and Cortactin, known as direct Shank partners in mammals and present in the Monosiga genome, are still present in Trichoplax despite apparent absence of Shank. As an alternative possibility, Shank could have been replaced by another protein. Functional work is certainly needed in Tricho-
plax to determine the molecular interactions and cellular functions of these proteins. Whether these genes are expressed in the fibre cells, proposed by some [40] to represent a placozan neural-like cell type, will be important to assess. However, placzoans are not likely to teach us much about the ancestral properties and functions of PSD proteins prior to the emergence of a nervous system, because the absence of Shank (if con-
irmed) makes their PSD-like complex totally idiosyncratic.

Whereas the emergence of nerve cells and synapses in a eumetazoan ancestor was seemingly not accompanied by a significant expansion of the PSD complex in terms of pro-
tein number, the few proteins acquired in this branch are significant with respect to neural function. They include the ionotropic glutamate receptors NMDAR (involved in synaptic transmission) and the Neuroligins (involved in synaptic adhesion and signalling in synaptogenesis, through interaction with their ligands the Neurexins [41,42]). Within eumetazoans, we inferred a few protein losses: ErbB receptor and GRIP (a scaffold protein) in cnidarians, α-catenin and Neuroligin in Hydra, SPAR in pro-
tostomians, Erbin in Drosophila and ErbB receptor in the annelid Capitella. Curiously enough, there is no concomi-
tant loss of proteins known to interact with each other (e.
"g. losses of Erbin are not associated with losses of the ErbB receptor), suggesting that proteome and interactome evolved to a large extent independently, or that protein interactions in non-model organisms differ substantially from what happens in the mammalian synapse.

What were the ancestral functions of the PSD protein complex?

Comparative genomics contributes to our understanding of hierarchy in biological systems, by sorting ancient components from more recent additions, a basis for making predictions about ancestral interactions and functions. In bilaterian animals, the PSD is tightly linked functionally to intercellular signalling at the synapse. However, the proto-PSD of the choanoflagellate/ metazoan ancestor (red + violet colours in Fig. 3) was probably not involved in intercellular signalling, since all post-synaptic membrane receptors linked to the mammalian PSD (the different kinds of glutamate receptors, Neuroligin, ErbB receptor, receptor domain of the Ephrin receptor) are of more recent origin. In addition, the global structure of the ancestral proto-PSD, which lacks several important scaffold proteins (GRIP, S-
SCAM, CASK), was certainly much simpler than that of the mammalian PSD.

Shank and its conserved direct or indirect partners appear to form the ancestral core of the post-synaptic protein complex (Fig. 3). The associated associations are actin cytoskeleton regulation (Homer links actin to Shank; the actin-regulating protein Cortactin binds to Shank) and Ca$^{2+}$ signalling (Homer links the IP3 receptor to Shank; CaMKII and PKC$\beta_{\gamma}$, also involved in Ca$^{2+}$ signalling, are present in the choanoflagellate genome). It is thereby proposed here that the main function of the ancestral Shank interactome was to provide a link between intracellular Ca$^{2+}$ signalling and actin cytoskeleton regulation.

The DLG scaffold protein might have ancestrally formed the centre of an additional interactome, possibly independent from the Shank module (GKAP, the physical link between DLG and Shank in mammals, is absent in Monosiga). We identified three DLG partners with clear orthologues in the choanoflagellate genome: CRIPT (a microtubule-interacting protein [43]), the K+ channel Shaker, and the Ca$^{2+}$ plasma membrane channel PMCA. The choanoflagellate orthologue of Citron (a Rho target) is not likely to interact with DLG as in mammals because only the most N-ter part of the protein shows homology with mammalian Citron (Fig. 3). These observations suggest ancestral functions of the DLG interactome in concentrating cation channels on particular plasma membrane areas, and in providing a link between these channels and the regulation of microtubule polymerisation.

From a morphological point of view, the choanoflagellate + metazoan clade is characterised by a unique synapomorphy: acquisition of an apical cell complex with a central flagellum surrounded by a collar of microvilli. This is suggested by the close resemblance between apical cell complexes of choanoflagellates and of sponge choanocytes [44]. According to some [45], this flagellum-collar structure would be homologous with the apical complex of eumetazoan sensory cells. Because choanoflagellate collar microvilli are contractile and labile structures (suggesting highly dynamic regulation), and because their cytoskeleton is principally made of F-actin, it is tempting to speculate about a possible functional link between the emergence of the proto-PSD and the acquisition of the flagellum-collar apical complex. This hypothesis can be tested by determining the subcellular localisation and function of the key proteins Shank, Homer and DLG in Monosiga. Another stimulating path for future investigation on the ancestral functions of core PSD components will involve the characterisation of physical interactions between the various choanoflagellate PSD-like proteins.

Conclusion

The backbone of the post-synaptic density, notably featuring the scaffold proteins Shank, Homer and DLG, was acquired in a common ancestor of choanoflagellates and metazoans, in a unicellular context. It was probably not involved in cell-cell communication, but rather in linking transmembrane cation currents, calcium signalling, and cytoskeleton regulation. Future investigations on complete genome sequences of unicellular opisthokonts that are more closely related to the choanoflagellate + metazoan clade than to Fungi (ichthyosporids and Capaspora) [1] might happen to push even earlier the period of origination of these proteins.

After the addition of several novel components (among which GKAP, the link between DLG and Shank, and the CASK/S-SCAM/classical cadherin interactome), metazoan ancestors became endowed with a nearly complete set of PSD proteins (violet + red + blue colours in Fig. 3). Proteins that are specifically involved in synaptic transmission (i.e. AMPAR, NMDAR and mGluR) or adhesion (neuroligin) in eumetazoans were nevertheless absent from this ancestral animal repertoire (except possibly for mGluRs, present in the metazoan ancestor if Trichoplax is not the basal-most lineage, see Additional file 7). This suggests that PSD proteins ancestrally acted in a non-synaptic context, in consensus with the probable monophyly of Eumetazoa [34] and its corollary, that the last common ancestor of all metazoan animals was nerve-less. In this ancestor, the PSD-like complex was probably a sub-membrane platform for bringing together molecules involved in cation currents, Ca$^{2+}$ and other signalling pathways, and cytoskeleton regulation, possibly (but not necessarily) in relation to cell sensitivity to external stimuli. The complex later became integrated into the evolving synapse, in a eumetazan ancestor, through the addition of novel proteins and functionalities.
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Authors’ contributions

AA and MW conceived and designed the analyses. AA performed the data collection and the analyses. MM drew the figures. AA and MW wrote the paper. Both authors read and approved the final manuscript.

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References

1. Ruiz-Trillo I, Roger AJ, Burger G, Gray MW, Lang BF: A phylogenomic investigation into the origin of metazoa. Mol Biol Evol 2008, 25:646-72.
2. Sakaya C, Armstrong KA, Adamska M, et al: A post-synaptic scaffold at the origin of the animal kingdom. PLoS ONE 2007, 2:e506.
3. Putnam NH, Srivastava M, Helften U, et al: Sea anemone genome reveals ancestral eumetazoan gene repertoire and genomic organization. Science 2007, 317:86-94.
4. Grimson A, Srivastava M, Fahey B, et al: Early origins and evolution of microRNAs and Piwi-interacting RNAs in animals. Nature 2008, 455:1193-1197.
5. Exposito J, Larroux C, Cluzel C, et al: Demosponge and sea anemone fibrillar collagen diversity reveals the early emergence of A/C clades and the maintenance of the modular structure of type V/IXI collagens from sponge to human. J Biol Chem 2008, 283:28226-28235.
6. Manning G, Young SL, Miller WT, Zhai Y: From the Cover: The protist, Monosiga brevicollis, has a tyrosine kinase signaling network more elaborate and diverse than found in any known metazoan. Proc Natl Acad Sci USA 2008, 105:9674-9679.
7. Larroux C, Luke GN, Koopman P, et al: Genesis and expansion of metazoan transcription factor gene classes. Mol Biol Evol 2008, 25:980-96.
8. King N, Westbrook MJ, Young SL, et al: The genome of the choanoflagellate Monosiga brevicollis and the origin of metazoa. Nature 2008, 451:783-8.
9. Rokas A: The origins of multicellularity and the early history of the genetic toolkit for animal development. Annu Rev Genet 2008, 42:235-251.
10. Koski K: Exploring the early origins of the synapse by comparative genomics. Biol Lett 2009, 5:108-11.
11. Collins MO, Hui H, Yu L, et al: Molecular characterization and comparison of the components and multiprotein complexes in the postsynaptic proteome. J Neurochem 2006, 97(1):16-23.
12. Grant SGN: The synapse proteome and phosphoproteome: a new paradigm for synaptic biology. Biochem Soc Trans 2006, 34:59-63.
13. Paddockston AJ, Armstrong JD, Grant SGN: Organization of brain complexity-synapse proteome form and function. Brief Funct Genomic Proteomic 2006, 5:66-73.
14. Klemper P, Smit AB, Li KW: Proteomics analysis of immuno-precipitated synaptic protein complexes. J Proteomics 2009, 72:83-90.
15. Yamauchi T: Molecular constituents and phosphorylation-dependent regulation of the post-synaptic density. Mass Spectrom Rev 2002, 21:266-286.
16. Nishimura W, Yao I, Iida J, Tanaka N, Hata Y: Interaction of synaptic scaffolding molecule and Beta-catenin. J Neurosci 2002, 22:757-765.
17. Boeckers TM: The postsynaptic density. Cell Tissue Res 2006, 326:409-422.
18. Krenkamp H: Scaffolding proteins at the postsynaptic density: shank as the architectural framework. Handb Exp Pharmacol 2008, 162:305-380.
19. Ehlers MD: Molecular morphogens for dendritic spines. Trends Neurosci 2002, 25:64-67.
20. Vessey JP, Kana D: More than just synaptic building blocks: scaffolding proteins of the postsynaptic density regulate dendritic patterning. J Neurochem 2007, 102:324-332.
21. Shiwa-Yamaguchi Y, Furuchi T: The Homer family proteins. Genome Biol 2007, 8:206.
22. Richards GS, Simionato E, Perron M, et al: Sponge genes provide new insight into the evolutionary origin of the nervous circuit. Curr Biol 2008, 18:1156-61.
23. Nielsen C: Six major steps in animal evolution: are we derived sponge larvae? Evol Dev 2008, 10:241-257.
24. Medina M, Collins AG, Silberman JD, Sogin ML: Evaluating hypotheses of basal animal phylogeny using complete sequences of large and small subunit rRNA. Proc Natl Acad Sci USA 2001, 98:9707-9712.
25. Manuel M, Borchartelli C, Alvan E, et al: Phylology and evolution of calcareous sponges: monophyly of calcinea and calcarea, high level of morphological homoplasy, and the primitive nature of axial symmetry. Syst Biol 2003, 52:111-133.
26. Nielsen C: Animal evolution: interrelationships of the living phyla. New York: Oxford University Press, 2001.
27. Brusca RC, Brusca GJ: Invertebrates Sunderland, MA: Sinauer Associates, 2003.
28. Schierwater B, Etel M, Jakob JW, et al: Concatenated analysis sheds light on early metazoan evolution and fuels a modern "urnetazon" hypothesis. PLoS Biol 2009, 7:e20.
29. Dunn CW, Hejnol A, Matus DQ, et al: Broad phylogenomic sampling improves resolution of the animal tree of life. Nature 2008, 452:745-9.
30. Dellaporta SL, Xu A, Sagasser S, et al: Mitochondrial genome of Trichoplax adhaerens supports placozoa as the basal lower metazoan phylum. Proc Natl Acad Sci USA 2006, 103:8751-8756.
31. Hejnol A, Obst M, Stamatakis A, et al: Assessing the root of bilaterian animals with scalable phylogenomic methods. Proc Biol Sci 2009, 276(1677):4261-70.
32. Miller DJ, Ball EE: Animal evolution: Trichoplax, trees, and taxonomic turmoil. Curr Biol 2008, 18(R1003-1005).
33. Telford MJ: Animal evolution: once upon a time. Curr Biol 2009, 19:R339-341.
34. Philippe H, Derelle R, Lopez P, et al: Phylogenomics revives traditional views on deep animal relationships. Curr Biol 2009, 19:706-712.
35. Schierwater B, de Jong D, Desalle R: Placozoa and the evolution of Metazoa and intrasomatic cell differentiation. Int J Biochem Cell Biol 2009, 41:370-379.
36. Srivastava M, Begovic E, Chapman J, et al: The Trichoplax genome and the nature of placozoans. Nature 2008, 454:955-60.
37. Zamponi GW: Regulation of presynaptic calcium channels by synaptic proteins. J Pharmacol Sci 2003, 92:79-83.
38. Neusch C, Weishaupt JH, Bähr M: Kir channels in the CNS: emerging new roles and implications for neurological diseases. Cell and Tissue Research 2003, 311:131-138.
39. Guindon S, Gascuel O: A simple, fast, and accurate algorithm to estimate large phylogenies by maximum likelihood. Syst Biol 2003, 52:696-704.
40. Hadrys T, DeSalle R, Sagasser S, Fischer N, Schierwater B: The Trichoplax Pax6 gene: a putative Proto-PaxA/B/C gene predating the origin of nerve and sensory cells. Mol Biol Evol 2005, 22:1569-1578.
41. Gerrow K, El-Husseini A: Cell adhesion molecules at the synapse. Front Biosci 2006, 11:2400-2419.
42. Dalva MB, McClelland AC, Kayser MS: Cell adhesion molecules: signalling functions at the synapse. Nat Rev Neurosci 2007, 8:206-220.
43. Passafaro M, Sala C, Niethammer M, Sheng M: Microtubule binding by CRIP1 and its potential role in the synaptic clustering of PSD-95. Nat Neurosci 1999, 2:1063-1069.
44. Maldonado M: Choanoflagellates, choanocytes, and animal multicellularity. Invert Biol 2004, 123:1-22.
45. Jacobs DK, Nakanishi N, Yuan D, Camara A, Nichols SA, Hartenstein V: Evolution of sensory structures in basal metazoan. Integr Comp Biol 2007, 47:712-723.

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