Molecular, phylogenetic and developmental analyses of Sall proteins in bilaterians

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Abstract

Background: Sall (Spalt-like) proteins are zinc-finger transcription factors involved in a number of biological processes. They have only been studied in a few model organisms, such as Drosophila melanogaster, Caenorhabditis elegans, Schmidtea mediterranea and some vertebrates. Further taxon sampling is critical to understand the evolution and diversification of this protein and its functional roles in animals.

Results: Using genome and transcriptome mining, we confirmed the presence of sall genes in a range of additional animal taxa, for which their presence had not yet been described. We show that sall genes are broadly conserved across the Bilateria, and likely appeared in the bilaterian stem lineage. Our analysis of the protein domains shows that the characteristic arrangement of the multiple zinc-finger domains is conserved in bilaterians and may represent the ancient arrangement of this family of transcription factors. We also show the existence of a previously unknown zinc-finger domain. In situ hybridization was used to describe the gene expression patterns in embryonic and larval stages in two species of snails: Crepidula fornicata and Lottia gigantea. In L. gigantea, sall presents maternal expression, although later on the expression is restricted to the A and B quadrants during gastrulation and larval stage. In C. fornicata, sall has no maternal expression and it is expressed mainly in the A, C and D quadrants during blastula stages and in an asymmetric fashion during the larval stage.

Discussion: Our results suggest that the bilaterian common ancestor had a Sall protein with at least six zinc-finger domains. The evolution of Sall proteins in bilaterians might have occurred mostly as a result of the loss of protein domains and gene duplications leading to diversification. The new evidence complements previous studies highlighting an important role of Sall proteins in bilaterian development. Our results show maternal expression of sall in the snail L. gigantea, but not C. fornicata. The asymmetric expression shown in the ectoderm of the trochophore larva of snails is probably related to shell/mantle development. The observed sall expression in cephalic tissue in snails and some other bilaterians suggests a possible ancestral role of sall in neural development in bilaterians.

Keywords: Lottia gigantea, Crepidula fornicata, Gastropoda, Spiralia, Gene evolution, Protein domains, Spalt, Sall
The spalt (sal) genes were originally described as homeotic genes in Drosophila melanogaster [2], which has two paralogs: spalt major (salm) and spalt-related (salr) [7]. Afterward, spalt orthologs have been also described in the nematode Caenorhabditis elegans [8], the planarian Schmidtea mediterranea [9], as well as in some species of vertebrates [10]. All these studies have shown that spalt is activated in response to several signal transduction pathways in different tissues and developmental processes [11], and the expression patterns and functions of sall genes, together with the analysis of their regulation, indicate they cannot be universally assigned to a specific signaling pathway.

For instance, Sall proteins have been described as transcriptional repressors, mainly through two mechanisms [12]: (1) the interaction between 12 amino acids located at the N-terminal part of Sall proteins and the histone deacetylase complex NuRD [13–15] and (2) the direct binding to an AT-rich region of the heterochromatin of the central region of Sall proteins that includes ZF2 and ZF3 pairs [16]. In addition, SalI proteins can interact with PIN2, an isoform of telomeric repeat-binding factor 1 (TRF1), which might indicate an involvement of Sall proteins in the regulation of higher-order chromatin structures and that the Sall proteins could be components of a distinct heterochromatin-dependent silencing process [17]. In addition, Sall proteins have also been described as transcriptional activators of several genes, such as the cyclin CDK inhibitor p21 [18], Nanog [19], Pou5f1 [20] and Sall itself [21].

The subcellular localization and transcriptional capacity of Sall proteins might be conditioned by post-translational modifications and protein interactions. SUMOylation modifies its localization inside the nucleus [12], while phosphorylation reduces its repression activity [22]. Interaction between Sall proteins can also have functional consequences, such as differences in subcellular localization that might be mediated by the poly-Q region [23].

Sall is therefore involved in a number of different biological processes. The Drosophila sal homeotic genes are implicated in many developmental processes [11, 12] such as the specification of head and tail during embryogenesis [24], organogenesis [25–27] and the determination of neural fate in the peripheral nervous system [28–30]. Similarly, in the nematode C. elegans, the sall gene sem-4 controls the fate of several different cell types including neurons, muscle, hypodermis, sex myoblasts, coelomocytes and multiple neuronal lineages [8, 31]. Studies in the nematode and also in flies and vertebrates indicate that sall genes might regulate this patterning and cellular identity through repression of Hox genes [32]. In addition, sal is involved in the formation of excretory systems in planarians [9]. Finally, vertebrate homologs of spalt (SALL) have been shown to be involved in normal development and tumor suppression and are implicated in several human genetic disorders [10]. They have important roles during neural development [20, 33–35] and organogenesis, especially in kidney [9, 34, 36–38], heart [33, 34, 36] and limb development [23, 39, 40].

Therefore, knowledge about spalt and its expression derives from studies in a few model organisms representing the three major clades of bilaterians: Ecdysozoa D. melanogaster Saln and Salr and C. elegans SEM-4; Spiralia S. mediterranea Sall; and Deuterostomia Vertebrate Sall proteins. Unlike the deuterostomes and ecdysozoans, spiralian have not been so widely used in modern cell and molecular research. Spiralian comprise 14 morphologically diverse phyla (including mollusks, annelids and nemertans) that share some developmental processes like the presence of spiral cleavage, from which the name of the group is derived [41]. The diversity of developmental programs, life histories and body plans makes them an excellent group to undertake comparative studies aimed at understanding the molecular and genetic bases of the evolution of morphological diversity of triploblast bilaterian metazoans.

Advances in sequencing have made available new genomic and transcriptomic data for some non-model Spiralia. The snail species Lottia gigantea was the first representative of Spiralia to have its genome sequenced [42], and the availability of sequence data for other members of this group in public databases like NCBI keeps increasing by the day. In addition, more studies have recently focused on the developmental genetics of Spiralia in the last few years, mostly with a focus on understanding the role of certain genes in the origin

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**Fig. 1** Organization of conserved domains in “Sal-box” containing proteins. Colored ovals represent the zinc-finger motifs. The blue rectangle represents the poly-Q region. The turquoise diamond represents the 12 conserved amino acids at the N-terminal end of the Sall proteins that interact with the HDC NuRD [14]. Protein length range is indicated at the right.
of animal body plan diversity and in molecular evolution of genes and genetic pathways. Although there are no classical model species in developmental genetics of Spiralia, some efforts of several labs including ours have focused on a few species of snails, nemertans and annelids, which have provided valuable information on gene expression patterns and on experimental manipulation of certain genes [43-48]. One conserved feature shared by the spiralian is the distinct pattern of alternating oblique embryonic cell divisions referred to as "spiral cleavage." In spiralian embryos, the first and second orthogonal cleavage planes led to the formation of four cells (macromeres A, B, C and D) and the progeny of these cells will define the future left, ventral, right and dorsal body sides (referred to as A, B, C and D "quadrants," respectively). Subsequent divisions result in the stereotypic formation of successive tiers of generally smaller micromeres, which are generated toward the animal pole. Cleavage is then followed by gastrulation and finally by the formation of a larva [49].

In order to understand the origin and the evolutionary history of Sall proteins in bilaterians, here we have identified Sall proteins in several Spiralia, including mollusks, an annelid and a brachiopod, as well as in some other bilateral organisms. We have identified the main protein domains and the potential protein interaction sites in all these newly determined Sall orthologs. We also report the expression pattern of sall gene in two species of gastropods (L. gigantea and Crepidula fornicata) during their embryonic development. Finally, we discuss the evolution of Sall proteins, the evolutionary conservation of their domains, and the temporal and local activation in a wider comparative phylogenetic analysis.

Methods
Identification of candidate Sall proteins
In order to identify candidate Sall proteins, we performed searches in different databases (Additional file 1: Table S1). Potential snail Sall sequences were derived from Biomphalaria glabrata and C. fornicata from RNA-seq datasets generated in our lab [50] and uploaded to Geneious version 6.1.2 [51]. Potential sequences of the snail L. gigantea were retrieved from the JGI genome portal [42] using blastn and pblast alignment algorithms [52]. Potential sall orthologs for the Xenoturbellid Xenoturbella, the acelis Convolutriloba and Isodiametra, the annelid Dinophilus, the brachiopods Terebratalia and Novocrania, the nemertean Lineus, the priapulid Priapulus, the platyhelminth Prostheceraeus, the nemertodermatid Meara, and the bryozoan Membranipora were searched for in RNA-seq datasets (Additional file 1: Table S1). In order to have more representatives of other bilaterian (arthropods, nematodes, mollusks, tunicates, echinoderms, hemichordates, vertebrates) and non-bilaterian clades (cnidarians, ctenophorans, placozoans, poriferans) for a wider analysis of the phylogeny and structure of Sall proteins, additional searches were performed in the NCBI databases [53] using keyword search (Spalt, Spalt-like, sall, sal-like), tblastn and pblast. In addition, the zinc-finger proteins containing the Sal-box motif Schnurri, PRDII-BF1 and HIVEP1 were also retrieved from the NCBI databases using keyword search (Schnurri) and tblastn and pblast search using the Sal-box motif as template. Translation into protein sequences was carried out using MacVector version 12.7 [54], assuming standard codon usage.

Phylogenetic analyses
Full-length sequences of available Sal-box containing proteins (Schnurri, PRDII-BF1, HIVEP1) (Additional file 1: Table S1) were aligned with Sall potential orthologs (Additional file 2: Fig. S1) using ClustalX version 2.1 [55] followed by refinement by eye and trimmed in MacVector, selecting the homologous sequences and excluding sites of ambiguous alignment and gaps. In order to determine whether the newly determined potential Sall proteins were indeed Sall proteins or other proteins containing a Sal-box, we performed a phylogenetic analysis including the zinc-finger domains three and four (ZF3 and ZF4), the only two zinc-finger domains present in all proteins containing a Sal-box (Fig. 1; Additional file 2: Fig. S1), for all the sequences retrieved in this study. Once orthology was established, a second phylogenetic analysis was performed including the zinc-finger domains two, three and five (ZF2, ZF3 and ZF5) (Additional file 3: Fig. S2) for all the Sall sequences retrieved in this study (Additional file 1: Table S1). These two datasets were subjected to coalescent-based, Bayesian inference (BI) phylogenetic analyses implemented using BEAST 1.8.3 software [56]. The JTT + G model [57] was selected as the best-fit model of protein evolution using ProtTest [58]. We assumed a strict molecular clock and the Yule speciation model as the coalescent prior. Analyses were run for 3,000,000 generations, sampling trees and model parameters every 300 generations. Convergence of results was assessed by visual inspection of the log file using Tracer software [59] and accordingly a burn-in period of 300,000 generations (10%) was established. We used TreeAnnotator software (distributed as part of the BEAST software package) to recover the maximum clade credibility (MCC) consensus tree from the post-burn-in sample of trees. The robustness of the inferred clades was evaluated based on Bayesian posterior probabilities (BPPs). Candidate sequences were identified as orthologs when they grouped in a clade with high statistical support (BPP > 0.95) with sequences of known identity.
Cloning, sequencing and RNA probe generation

Embryos from *L. gigantea* were handled and stored in the freezer as described in Grande and Patel [60]. Embryos from *C. fornicata* were collected and reared as previously described (see, for instance, Henry et al. [49]). High-quality total RNA from embryos of the gastropods *L. gigantea* and *C. fornicata* was extracted using TriZol and purification methods followed the manufacturer’s suggested protocol. The purity and concentration of total RNA was verified with a NanoDrop ND-1000 Spectrophotometer (Thermo Scientific, Wilmington, DE) and approximately 1 µg of total RNA from each developmental stage was used to synthesize cDNA (iScript cDNA Synthesis kit, Bio-Rad, Hercules, CA). Gene-specific primers were designed for each gene (Additional file 4: Table S2). PCR amplification reactions were performed with Phusion HF DNA polymerase, Q5 high-fidelity DNA polymerase and Q5 high GC enhancer buffer (New England Biolabs, Ipswich, MA), according to the manufacturer’s suggested ratios. Amplified PCR products were run on 1% agarose gels, gel-purified (GeneClean Turbo kit, MP Biomedicals, Solon, OH) and cloned into pGem-T Easy vector (Promega, Madison, WI). Digoxigenin (DIG)-labeled and fluorescein (FITC)-labeled antisense and sense RNAs were synthesized from purified pGEM-T easy plasmid DNA that was amplified using SP6 and T7 primers via PCR. Template DNA was used with SP6 and T7 RNA polymerase (Invitrogen, Carlsbad, CA, USA), DIG and fluorescein labeling mix (Roche, Indianapolis, IN, USA) to generate probes. Reactions were purified with RNeasy MinElute Cleanup kit (Qiagen, Valencia, CA), and probe concentrations were verified using a NanoDrop ND-1000 Spectrophotometer.

In situ hybridization and Hoechst staining

The in situ hybridization protocol for single-gene labeling was modified from Finnerty et al. [61] and Perry et al. [46]. The double labeling in situ protocol in *L. gigantea* was performed as described in Grande and Patel [60]. Embryos were counterstained with 1:10,000 dilution of Hoechst 33342 and stored in 80% glycerol/20% 1× PBS for 1 h to visualize nuclei, followed by three washes in PBS and stored in 80% glycerol/20% 1× PBS.

Microscopy and image analysis

Fixed embryos processed for in situ hybridization were mounted on Rain-X coated (ITW Global Brands, Houston, TX) glass slides in 80% glycerol/20% 1X PBS. Coverslips were prepared as described in Lyons et al. [62] for *C. fornicata* and with two additional supporting coverslips at the sides for *L. gigantea*. All images were acquired using an Axioskop 2 plus microscope (Zeiss) in conjugation with a CoolSnap FX color camera (Roper Scientific) and MetaVue 5.07 (Universal Imaging) software from Centro de Biología Molecular Severo Ochoa SMOCS facilities. Additional image processing was done with Helicon Focus (Helicon Soft Ltd. Kharkov, Ukraine) to combine multifocal stacks of images and get focused images and Adobe Photoshop (Adobe Systems Software Ireland Ltd.) for the stacking of bright-field and fluorescent images of *C. fornicata* embryos.

Results

Phylogenetic analyses

One single potential Sal1 ortholog was found in several Spiralia, including mollusks, nemertean, annelids, brachiopods, bryozoans and platyhelminthes (Additional file 1: Table S1). One single potential ortholog was also found in some ecdysozoan groups, such as arthropods, and arthropods (except Diptera, which has two well-known paralogs: Salm and Salr) (Additional file 1: Table S1). In addition, one single ortholog was identified for some other bilaterians, such as acelcs, nemertodermatids and Xenoturbella (Additional file 1: Table S1). Deerterostomes except vertebrates also have one Sall ortholog (Additional file 1: Table 1). Searches in the databases from non-bilaterian metazoans did not retrieve any potential orthologs (Additional file 1: Table S1).

In order to assign orthology, we first performed a phylogenetic analysis including the newly determined Sall sequences along with previously reported Sall sequences and the related zinc-finger proteins Schnurri, PRDII-BF1 and HIVEP1 (Additional file 5: Fig. S3). The resulting tree confirmed with high statistical support that the newly identified Sall sequences were indeed more closely related to the previously reported Sall sequences than to other zinc-finger-related sequences (BPP = 1) (Additional file 5: Fig. S3). Once orthology of the Sall proteins was established, we performed a second phylogenetic analysis including all Sall orthologs (Additional file 6: Fig. S4). This analysis pointed out the existence of three highly divergent sequences: the platyhelminth Sall, the nematode SEM-4 and the bryozoan Sall proteins (Additional file 6: Fig. S4). All vertebrate Sall sequences formed a clade (BPP = 1) with one supported group for each paralog (Sall1, Sall2, Sall3 and Sall4) (Additional file 6: Fig. S4). All ecdysozoan Sall sequences but SEM4 formed a single clade (BPP = 91) (Additional file 6: Fig. S4). Finally, all spiralian Sall sequences, except the platyhelminth, the bryozoan and the annelid Sall, formed a clade (BPP = 1) (Additional file 6: Fig. S4).

Analysis of Sall protein domains

All Sall orthologs were aligned in order to identify the protein domains. Gene sequences were incomplete in the gastropod *C. fornicata*, the nemertodermatid, the
nemertean, the bryozoan and the tunicate (Fig. 2). The analysis of the protein domains showed that the molecular structure of Sall proteins is highly consistent in bilaterians with some exceptions discussed below (Fig. 2). The ZF domain located in the N-terminal part of the protein (ZF1) corresponds to the C2HC class, and we found it in ecdysozoans, spiralians and deuterostomes, as well as in Xenoturbella, a member of the clade Xenacoelomorpha, the sister group of all bilaterians [63] (Fig. 2). The 12 amino acid sequence responsible for the interaction with the histone deacetylase complex NuRD is similarly conserved, but lost in some groups (Fig. 2; Additional file 7: Fig. S5). The poly-Q region is highly conserved, only lost in the nematode SEM-4, the platyhelminth and most mollusks ( gastropods and the cephalopod), and partially conserved in the bryozoan and the tunicate (Fig. 2; Additional file 7: Fig. S5). The other ZF domains (2–5) correspond to the C2H2 class arranged in pairs connected by an H/C link and are highly conserved through the evolution of Bilateria (Fig. 2). The second zinc finger (ZF2) is more conserved and contains the characteristic Sal-box. The ZF2 domain is lost in the nematode, the platyhelminth and the gastropod B. glabrata, and shows modifications in the nemertean and the annelid, which have lost the N-terminal zinc finger of the pair and the bryozoan, which has lost the C-terminal zinc finger (Fig. 2; Additional file 7: Fig. S5). The ZF3 domain is the most conserved in all groups analyzed here, and it has an additional zinc-finger-associated domain, except in Platyhelminthes, which may have lost it (Fig. 2; Additional file 7: Fig. S5). The ZF4 domain is lost in acoels and in the nematode. Sall2 of all vertebrates, as well as Sall3 of the platyhelminth and the gastropod, also lack ZF4, while it is modified in the nematode. 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Diptera (*Drosophila melanogaster*)

Lepidoptera (*Papilio machaon*)

Hymenoptera (*Duforea novaeangliae*)

Crustacea (*Limulus polyphemus*)

Arachnida (*Parasteatoda tepidariorum*)

Nematoda (*Pristionchus pacificus*)

Priapulida (*Priapulus caudatus*)

Cephalopoda (*Octopus bimaculoides*)

Gastropoda (*Biomphalaria glabrata*)

Gastropoda (*Crepidula fornicata*)

Gastropoda (*Lottia gigantea*)

Bivalvia (*Crassostrea gigas*)

Nemertea (*Lineus ruber*)

Annelida (*Dinophylus gyrociliatus*)

Brachiopoda (*Terebratalia transversa*)

Bryozoa (*Membranipora membranacea*)

Platyhelminthes (*Prostheceraeus vittatus*)

Vertebrata (*Homo, Mus, Gallus, Xenopus*)

Tunicata (*Ciona intestinalis*)

Echinodermata (*Strongylocentrotus*).

Hemichordata (*Saccoglossus kowalevskii*)

Fig. 2 Organization of conserved domains in Sall proteins in Bilateria. The light blue diamond represents the 12 conserved amino acids at the N-terminal end that interact with the HD‐NuRD [14]. Colored ovals represent the zinc‐finger motifs from ZF1 to ZF6. The blue rectangle represents the poly‐Q region. The striped lines represent missing sequence. Phylogenetic tree based on Cannon et al. [63] and Peters et al. [64]. The blue box on the tree highlights the deuterostomes; the orange box, the spiralia; the green box, the ecdysozoans; and the red box, the xenacoelomorphs. Bold names highlight the snail species studied in more detail in this work.
Fig. 3  Sall expression in C. fornicata. In situ hybridization of sall mRNA in embryos from 16-cell stage to organogenesis. The distribution of labeled mRNA is shown by the dark blue staining. In all images orientation is indicated at the bottom left corner of each panel. Scale bar at the bottom right corner equals 40 µm. Yellow arrowhead indicates the blastopore/stomodeum. a, c 16- and 24-cell stage embryos, respectively. mRNA is expressed in the macromeres. The lighter colored nuclei are labeled with Hoechst (blue). b, d Schematic representation of 16- and 24-cell embryos, respectively, in animal view (modified from Henry et al., [47]). Capital letters indicate the macromeres and lowercase letters the micromeres. e-j’ Localization of sall mRNA in blastula stage. e-e’, j-j’ Bright-field and fluorescent Hoechst-labeled corresponding images. f-i Lateral view of each quadrant. Green and red arrowheads indicate stained reference cells in quadrants C and A, respectively, in different views. q quadrant, A animal pole, V vegetal pole. k-n’ Localization of sall mRNA in gastrula stage. Each bright-field image has a corresponding fluorescent Hoechst-labeled image on the right. o-q’ Localization of sall mRNA in organogenesis. o, o’ Bright-field and fluorescent Hoechst-labeled corresponding images. p Bright-field right view. The blue arrow indicates the asymmetric patch of expression present in this side. q Bright-field left view. A: anterior; P: posterior; V: ventral; D: dorsal; R: right; L: left
in C and D quadrants after the 32-cell stage during blastula and gastrula stages, but is still restricted to the same cells and their progeny (Fig. 4e, f). The progeny of those cells that expressed \(sall\) in the A and B quadrants at the 32-cell stage maintained the expression of \(sall\) up to the larva stage (Fig. 5). Finally, in the trochophore stage \(sall\) is also expressed in a ring in the cephalic region (Fig. 5c, e) and in a dorsolateral ectodermic strip in the post-trochal region at both sides of the trochophore (Fig. 5c–e).

**Discussion**

Our study is the first to investigate Sall proteins in a broad phylogenetic framework. Previous studies had identified Sall proteins exclusively in nematodes, flies, planarians and vertebrates. Here we showed that \(sall\) is also present in members of all main groups across Bilateria (i.e., Xenacoelomorpha, Ecdysozoa, Spiralia and Deuterostomia) (Fig. 2). In addition, we could not find any Sall protein in the genome of non-bilaterians (the cnidarian *Nematostella vectensis*, the ctenophore *Pleurobrachia bachei*, the placozoan *Trichoplax adhaerens* and the sponge *Amphimedon queenslandica*), which suggests that Sall proteins might be an exclusive feature of bilaterians.

The analysis of the Sall protein domains and their comparison among bilaterians showed that orthologous zinc fingers in different Sall proteins are more similar in sequence among them than to other zinc-finger domains of the same Sall protein. This may indicate that the characteristic arrangement of the multiple double zinc fingers is not the result of independent duplications from a unique double zinc finger in different organisms, but is instead the ancient arrangement of this family of transcription factors. The novel zinc-finger domain described here (ZF6) is present in Ecdysozoa, Spiralia and Xenacoelomorpha, and one can interpret this as being plesiomorphic for all bilaterian Sall proteins and then secondarily lost in some specific groups like deuterostomes, nematodes, priapulids and insects (Fig. 2; Additional file 7: Fig. S5). Similarly, we found the ZF1 domain in most phyla, implying that it may also be an ancestral domain in
bilaterian Sall proteins, and not a domain exclusive of the vertebrate homologs, as previously thought [12] (Fig. 2).

In summary, this new evidence suggests that the bilaterian common ancestor probably had a Sall protein with at least 6 zinc-finger domains.

From this ancestral Sall protein, the evolution of Sall proteins in bilaterians might have occurred mostly as a result of the loss of protein domains and gene duplications leading to diversification of functions in paralogs, as it occurs in vertebrate or Drosophila Sall proteins. For instance, the presence/absence/modification of either ZF4 or ZF5 in paralogs of chordates and Drosophila might be related to the differential repression or activation mechanisms of these Sall proteins [10]. The ZF4 and ZF5 domains are necessary for the localization to heterochromatin in vertebrates [64]. In addition, this family of zinc fingers is capable of binding to DNA at specific AT-rich regions [16], as do other known zinc-finger proteins.

**Fig. 5** Expression of sall during trochophore stages in *L. gigantea*. The distribution of the labeled mRNA is shown by the dark blue staining. Orientation is indicated at the bottom left corner of each panel. Scale bar at the bottom right corner equals 40 µm. A: anterior; P: posterior; V: ventral; D: dorsal; R: right; L: left. a, b Localization of sall mRNA in pre-trochophore stage. Expression starts to extend from the progeny of 2 m micromeres. c–e In situ hybridization of sall mRNA at trochophore stage. Black arrowhead indicates the stained ring in the cephalic region. Green arrowhead indicates the dorsolateral ectodermic strip. Yellow arrowhead indicates the stomodeum.
could be explained by their functions, as the representative Fig. S5), and they can be easily homologized. How-
and cell-specific protein partners, resulting in differ-
ence and absence of both motifs observed in the analy-
protein is responsible for the repression activity through the recruitment of the histone deacetylase complex NuRD [14], but its function is correlated with its localization in the nucleus, which might be dependent on the ZF1 [13]. The almost absolute correlation between the presence and absence of both motifs observed in the analysis of the protein domains (Fig. 2; Additional file 7: Fig. S5) could be explained by their functions, as the repression through NuRD would not be possible without the localization provided by the ZF1. Another region located between ZF2 and ZF3 is a putative repression and heterochromatin localization domain [15] in vertebrate Sall and its functions might be conserved during evolution. These ZFs are encoded in a single large exon [1], show high identity and are especially well conserved, with ZF3 being conserved in all species analyzed (Fig. 2; Additional file 7: Fig. S5), and they can be easily homologized. However, ZF2 is not present in SEM-4 and Biomphalaria Sall (Fig. 2; Additional file 7: Fig. S5). Also, in contrast, in Drosophila Salm and Salr there is an activation domain located between the poly-Q and ZF3 domains [12]. In addition, the repression activity does not appear to be related to ZF3, which is able to bind to DNA. Rather, repression may be related to the region located at the N-terminal side of ZF3 [15], where we observed a highly conserved sequence in all species analyzed (consensus sequence: SETSKLQQLVENID). One more region, the poly-Q domain, could be important for the biological activity of these proteins, making possible the interaction among the paralogs as suggested by Sweetman et al. (2003) [23]. However, our data show that the poly-Q domain is not only restricted to vertebrates, but is widely present across the Bilateria, even in species with only one Sall form. Therefore, further studies are needed to understand the role of the poly-Q domain in Sall proteins.

The best homology among the double zinc fingers was found in the eight amino acids of the “Sal-box,” which are present in all C-terminal finger motifs of the ZF pairs and the H/C link [3]. Interestingly, in all cases where one finger of a pair is lost, it is the N-terminal. This kind of zinc-finger pair has also been observed in other proteins: the human transcription factor PRDII-BF1 [5] and Drosophila Schnurri [4, 5]. Sequence similarity within the pairs of zinc fingers suggests the importance of the Sal-box for the structure of the motif and may define a conserved subfamily of zinc-finger proteins.

Given the implication of sall in the development of Drosophila [11], C. elegans [8], S. mediterranea [9] and vertebrates [12], we studied the expression pattern through the development of two snail species: C. fornicata (Fig. 3) and L. gigantea (Figs. 4, 5) for a better understanding of the relevance of this protein and its functions in animal development. In both species, sall was expressed through most of the development process, suggesting that sall has a role in the development of spiralian, as in other bilaterians.

However, the differences in the expression patterns found in the snails L. gigantea and C. fornicata indicate that the degree of conservation of sall coding sequences does not necessarily imply similarities of expression patterns (Figs. 3, 4, 5). Interestingly, L. gigantea show maternal expression of sall, a feature that was previously described only in murine sall4, which presents maternal mRNA at the 2-cell stage, but is degraded in the next cleavage [67]. While Lottia shows a high level of maternal expression in all cells at the 2-cell and 4-cell stages and in the 2 m micromeres at early cleavage, this does not seem to be the case for Crepidula (Figs. 3, 4). The appearance of sall mRNA at the 16-cell stage in Crepidula embryos implies that sall mRNA is not maternal in this species and that zygotic expression of sall begins later, at the 16-cell stage (Fig. 3). Further work in other bilaterian embryos will help elucidate whether maternal expression is an ancestral feature of Sall proteins.

The transcriptional expression of sall differs greatly between Lottia and Crepidula during gastrulation and early organogenesis stages, although there are some common patterns observed in larvae (Figs. 3, 4, 5). The restriction in the expression pattern to A and B quadrants seen in Lottia was not previously described for any other gene in this species (Fig. 4). By contrast, in Crepidula, the expression is mainly restricted to A, C and D quadrants (Fig. 3). The lateral asymmetries in the expression pattern in the posterior region in C. fornicata during
organogenesis (Fig. 3) might be related to the differential proliferation during shell/mantle growth as previously described for other genes [46]. However, in both species sall is expressed close to the stomodeum in the posterior lateral ectoderm (Figs. 3, 4, 5). In addition, in Lottia sall is expressed in a ring in the cephalic domain (Fig. 5).

Determination of functional conservation of Sall proteins across bilaterians would require further experiments in additional groups. However, several lines of published evidence indicate that at least some functions of sall have been evolutionarily conserved. For instance, the sall genes appear to function as cell fate determinants, regulators of Hox genes and AP patterning, and as transcriptional repressors. In addition, their function in neural development seems to be conserved. In Drosophila, Sall is required in neuronal precursors and differentiated neurons to restrict neuronal fates to the proper cells [11], SEM-4 controls neural development in C. elegans [8] and the sall genes are involved in determining neural fates in chordates [20, 35]. Here we have shown that in L. gigantea sall is expressed in the cephalic region (Fig. 5), which may reflect its role in neural fate specification, although additional experiments are needed to confirm this hypothesis. If new evidence corroborates this neural expression in Spiralia, this may support the potential ancestral role of Sall proteins in neuronal development. An interesting question would be whether Sall is expressed in C. fornicata nephridia during organogenesis, since Sall1 is necessary in mice to develop kidneys [37, 38] and the human SALL1 is mutated in patients with Townes–Brocks syndrome (TBS) [68], which produces among other symptoms abnormal kidney development. Even in planarians, Sall is required for protonephridia regeneration [9].

Conclusions
The results of our analyses provide novel evidence about the evolution of Sall proteins and their functional domains. Specifically, they show that sall is conserved across Bilateria and might be exclusive to this group. The ancestral Sall protein probably presented six zinc-finger domains, including a sixth zinc-finger domain that is reported here for the first time (ZF6, Fig. 2).

We also present the first report of sall gene expression in snails; the results highlight its importance in the development of bilaterians. sall has maternal expression and is expressed in the ectoderm, surrounding the stomod- eum and in a cephalic ring in snails. The expression in the cephalic region in snails suggests a possible ancestral role of sall in neural development in bilaterians.

Additional files

- **Additional file 1: Table S1.** Protein sequences used in this study with their length, accession number and the corresponding species.
- **Additional file 2: Fig. S1.** Sequence alignment of Sal proteins. Alignment of the deduced amino acid sequences of Sall proteins of different taxa. Amino acids shared by more than 50% of the species are shaded in gray.
- **Additional file 3: Fig. S2.** Sequence alignment of Sal proteins. Alignment of the deduced amino acid sequences of Sall proteins of different taxa. Amino acids shared by more than 50% of the species are shaded in gray.
- **Additional file 4: Table S2.** Primer sequences used in this study.
- **Additional file 5: Fig. S3.** Phylogenetic analysis of zinc-finger proteins containing the “Sal-box” (FTTKGNLK). Maximum clade credibility tree from the BEAST analysis including the deduced amino acid sequences of Sall proteins together with Schnurri, PRDII-BF1 and HIVEP1. Numbers indicated above the nodes are Bayesian posterior probabilities (BPP). The yellow box highlights Sall sequences across the Bilateria (BPP = 1) and the red box, the sequences of the other zinc-finger-related proteins (BPP = 1).
- **Additional file 6: Fig. S4.** Phylogenetic analysis of Sall proteins across the Bilateria. Maximum clade credibility tree from the BEAST analysis including the deduced amino acid sequences of Sall proteins. Numbers above nodes are Bayesian posterior probabilities. The dark blue line groups Sall sequences of Vertebrates; the light blue line, the sequences of deuterostomes; the green line, the sequences of ecdysozoans (E); the orange line, the sequences of spiralian (S); and the red line, the sequences of xenacoelomorphs (X).
- **Additional file 7: Fig. S5.** Hypothesized Sall protein gains and losses during Bilateria evolution. The turquoise diamond represents the 12 conserved amino acids at the N-terminal end that interact with the HDC NuRD [14]. Colored ovals represent the zinc-finger motifs from ZF1 to ZF6. The blue rectangle represents the poly-Q region. The striped lines represent missing sequence. Phylogenetic tree based on Cannon et al. [63] and Peters et al. [64]. The blue box on the tree highlights the deuterostomes; the orange box, the spiralian; the green box, the ecdysozoans; and the red box, the xenacoelomorphs. Bold names highlight the snail species studied in more detail in this work. On each branch, proposed gains and losses of Sall protein domains are indicated.
- **Additional file 8: Fig. S6.** Amplification of spalt from cDNA extracted from zygotes and 24-h post-fertilization Crepidula embryos. Amplification of spalt (Cfo-spalt) was only detected using cDNA from 24-h post-fertilization Crepidula embryos. There was no amplification at all when cDNA from zygotes was used. The gene twist from Crepidula (Cfo-twist) was used as a positive control since twist is detected at both stages of development. PCR products were run on 1% agarose gels.
- **Additional file 9: Fig. S7.** Schematic drawings of sall expression at 4-cell, 8-cell, 16-cell stages and trochophore larva in L. gigantea. a: expression in all cells at 4-cell stage; b: Expression in the macromeres at 8-cell stage; c: Expression in the 2 m micromeres at 16-cell stage. Micromere 2a underlined in green and 2b underlined in orange. d: expression in the trochophore. In green and orange, expression of sall in the dorsolateral ectoderm derived from 2a and 2b, respectively. In pink sall, expression in the cephalic ring and next to the stomodeum, not derived from 2a and 2b micromeres. Drawings modified from Dictus and Damen [69].

Authors’ contributions
CG conceived and designed research. JL-S, MT-G, KJP and CG conducted experiments. All authors analyzed data. JL-S wrote the manuscript with input from all authors. All authors read and approved the final manuscript.
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Competing interests
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