Cross-phyla protein annotation beyond sequence similarity through structural prediction and alignment

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\section*{Abstract}

\textbf{Background:} Annotating protein function is a major goal in molecular biology, yet experimentally determined knowledge is often limited to a few model organisms. In non-model species, the sequence-based prediction of gene orthology can be used to infer function, yet this approach loses its predictive power with longer evolutionary distances. Here we propose a pipeline for the functional annotation of proteins using structural similarity, exploiting the fact that protein structures are directly linked to function and can be more conserved than protein sequence.

\textbf{Results:} We propose a pipeline for the functional annotation of proteins via structural similarity (CoFFE) and use this to annotate the complete proteome of a sponge, an early-branching animal uniquely positioned for inferring the early history of animal cell types. CoFFE accurately identifies proteins with known homology in >90\% cases, and annotates an additional 50\% of the proteome beyond sequence-based methods. Using this, we uncover new functions for sponge cell types, including extensive FGF, TGF and Ephrin signalling in sponge epithelia and redox metabolism and control in myopeptidocytes. Notably, we also annotate...
genes that arose via horizontal gene transfer and likely participate in digesting algal and plant
 cell walls, which are specific to the enigmatic sponge mesocytes.

**Conclusions:** Our work demonstrates that structural similarity is more powerful to
generate functional prediction and annotation than sequence alone, bridging longer evolutionary
distances. We anticipate that with a growing number of experimental and predicted structures
the approach outlined here will boost annotation efficiency especially for phyletically distant
non-model species, and will facilitate discovery in numerous -omics datasets.

**Keywords**

Functional annotation, proteins, *Spongilla*, scRNA-seq, structural similarity, structure
conservation, ortholog conjecture
Background

For many types of high-throughput experiments in molecular biology, knowing the protein function of the assayed genes is crucial to interpret the data. Since protein functional studies are limited to a few model species, amino acid sequence similarity has been used to infer the function of protein homologs. This approach exploits the observation that proteins with similar structures often have similar sequences, and that proteins with similar functions often have similar structures [1,2]. However, homology detection over longer evolutionary distances remains challenging owing to the decay of protein similarity that abolishes evidence of historical continuity. This presents a severe bottleneck for inferring protein function across a wide expanse of the tree of life, particularly in distant organisms where many proteins fall in the “twilight zone”, sharing a sequence identity only between 10-20% with proteins in characterised models [3,4].

A way to venture more deeply into the twilight zone is to use structural similarity for functional annotation transfer, as structures are more conserved in evolution [5]. Until recently, this was not feasible since predicting protein structure from amino acid sequence required prior inference of sequence homology [6]. This has changed with the advent of AlphaFold [7], a deep learning AI system that can predict de novo protein structures with atomic resolution, together with novel approaches for identifying similar structures in large databases [8]. Entire proteomes can now be aligned to known structures, either experimentally solved or predicted by AlphaFold, and aligned to structures with characterised functions from model systems.

In this work, we predicted structures for the entire proteome of the freshwater sponge *Spongilla lacustris*, an early-branching animal, and aligned them against available structural databases. We show a 50% expansion of previous functional annotation, and revisit recent single-cell RNA-sequencing data [9] to discover additional aspects of sponge cell biology, such as extended cell signalling in pinacocytes, redox metabolism and control in myopeptidocytes, and polysaccharide digestion as a key function of the previously uncharacterized mesocytes.
Results

A protein structure-based pipeline enriches functional annotation for *Spongilla lacustris*

We created a structure-based pipeline for functional annotation transfer, which we refer to as **CoFFE** (ColabFold-FoldSeek-EggNOG). Instead of using amino acid sequences directly, we predict protein structures, align them against structural databases, and transfer the functional annotation of the best hits to the queries (for an overview, see Suppl. Fig. 1; details in Methods). As a test case, we chose to annotate proteins in the freshwater demosponge *Spongilla lacustris*, an early-branching animal with only about 20 cell types, making it a unique model for understanding the origins of specialised animal cells [9].

The *Spongilla lacustris* proteome contains 41943 predicted proteins, including protein isoforms. As a first step, we used the ColabFold [10] pipeline to predict three-dimensional structures of all 41943 predicted *Spongilla* proteins (all structures and metadata deposited to ModelArchive, see Methods). This represents the entirety of the *Spongilla* proteome except for 11 proteins, which were too long (> 2900 amino acids) to be predicted by the available hardware. Confidence of predicted protein structures was assessed by calculating average pLDDT (predicted local distance difference) values. Notably, average pLDDT values for *Spongilla* protein structures were similar to those of more well-characterised animal models (Fig. 1A).

Next, we used the predicted protein structures as queries to search with FoldSeek [8] against AlphaFoldDB [11], SwissProt [12], and PDB [13]. After removing lower-quality matches we transferred the functional annotation of the best hits to the sponge proteins, producing annotations for slightly more than 60% of the proteome (25232 proteins). This represents an approximately 50% increase in annotated proteins compared to the state-of-the-art orthology pipeline EggNOG-mapper [14]. This moves Spongilla into a comparable level of functional annotation as *C. elegans* (70% at least described), one of the most intensively studied model species (Fig. 1B).

Exploration of the CoFFE results (Supplemental Note A) in terms of the distributions and correlation of different measures (pLDDT, structure/sequence bit score, perc. identity, etc.), as well as various protein characteristics (hydrophobicity, gravy index, polarity, etc.), did not reveal specific cause for concern and suggested little correlation between protein structure
prediction quality and chemical properties of the proteins.

Fig 1. Structural prediction and alignment of the *Spongilla* proteome A) Distribution of average pLDDT per protein structure of predicted proteomes from common model species in comparison to *Spongilla lacustris*. B) Percentage of genes that receive functional annotation from EggNOG, and the enrichment of the *Spongilla lacustris* functional annotation by the CoFFE pipeline. Highlighted organisms appear in A). C) Overlap between EggNOG and CoFFE annotations. **Ortholog**: proteins identified as belonging to the same orthology group in the most recent common ancestor in the EggNOG database. **Protein fam.**: proteins identified as belonging to the same eukaryote orthology group in the EggNOG database, indicating annotations represent homologs in the same gene family. 50% PFAM: half of the sequence-based PFAM domains are shared. No agreement: the CoFFE and EggNOG annotations differ. Subcategories with “same name” denote the fractions where EggNOG and CoFFE returned the same preferred name for a protein. D) Percent enrichment of *Spongilla* annotation at different levels with EggNOG-mapper (+ BLASTp) as the baseline.

**CoFFE annotations agree with sequence-based annotation transfer**

As a first step towards validating our pipeline, we compared CoFFE annotations to sequence-based annotations generated by EggNOG-mapper. This method annotates proteins by searching precomputed gene trees in the eggNOG orthology database and assigns query proteins to fine-grained orthology groups. A total of 16589 proteins were annotated by both
pipelines. Overall, 90.6% of proteins were annotated to homologs (Fig. 1C), including 56.7% orthologs and an additional 33.9% representing proteins in the same gene family. In the remaining cases, most (5% overall) shared PFAM domains in common \cite{15}, bringing the percentage of proteins with at least minimal agreement between CoFFE and EggNOG-mapper to \approx 95%. These results indicate that CoFFE can generally be trusted to find true homologs or proteins related by shared domains.

We also compared our results to “legacy” annotations from the recently published Spongilla cell type atlas, which used BLASTp to supplement EggNOG-mapper annotations \cite{9}. Compared to this combined sequence-based approach, CoFFE annotates more proteins organism-wide (60% to 40%). More importantly, CoFFE markedly improved the proportion of annotated cell type-specific markers (70%) compared with sequenced-based approaches (56%; Fig. 1D and Suppl. Note XX). This indicates CoFFE will be valuable for exploring cell type-specific functions in non-model organisms.

**CoFFE expands annotation of signalling pathways in Sponge pinacocytes**

Pinacocytes are contractile epithelial cells that line the sponge canal system, playing important roles in morphogenesis, barrier formation, and sponge whole-body contractions \cite{16,17}. Musser et al. showed pinacocytes upregulate genes involved in TGF/BMP and Wnt signalling, key pathways regulating development and morphogenesis in multicellular animals \cite{18,19}. Supporting this, CoFFE identified additional pinacocyte marker genes in each of these pathways.

In the FGF pathway, sequence-based annotations indicated expression of Fgf receptors in incumbent pinacocytes and apendopinacocytes, along with FGF mediator Frs and regulator Grb2, which are expressed in the entire family of pinacocyte cell types (Figure 2A). Extending this, CoFFE identified GAB1 and GAB2 (GBR2-associated-binding protein 1/2) as well as PTPN11/SHP2, which are necessary for signal transmission into the cell \cite{20,21}. Strikingly, CoFFE also detected the FGF ligand in *Spongilla*, which was not found using sequence-based approaches, revealing it is most highly expressed in incumbent pinacocytes, lophocytes, and sponge neuroid cells. Structural superposition of the protein with its best FoldSeek hit (UniProtID: P48804, *Gallus gallus* FGF4) revealed an extensive alignment of large parts of the protein, with a RMSD of 0.89 over 543 atoms despite a sequence similarity of only 11.8% (Fig. 2B).
Musser et al. also described multiple genes involved in TGF-β signalling in the pinacocyte family, including Tgfbr1, Acvr, Smad and Smurf [22]. CoFFE extends the list of known actors in pinacocyte TGF signalling, adding important ligands such as INHBE, CHRDL1 or KCP.

Lastly, Ephrin and the ephrin receptor (Eph) are membrane-anchored signalling molecules that mediate communication between adjacent cells [23, 24]. Whereas multiple ephrin receptors were detected via sequence similarity, ephrin itself was only found in the Spongilla proteome using highly sensitive HMM profile searches [25, 26]. CoFFE annotates a Spongilla gene with differential expression in various pinacocytes as a homolog of Caenorhabditis elegans Efn-3 (UniProtID: Q19475). Although these proteins share only 22% sequence identity, the aligned structures achieve an RMSD of 1.59 over 580 atoms (Fig. 2C). A separate Hmmer search [27] using the ephrin Pfam profile (PF00812) picked up the same gene, suggesting CoFFE is at least as sensitive as curated HMM profile searches.

FGF, TGF-beta as well as Ephrin interestingly exhibit converging downstream signalling pathways, including PI3K/Akt, ERK/MAPK1, JNK or RhoA/ROCK, responsible for cell growth, differentiation, migration and cytoskeletal organisation [23, 28, 29]. Together, CoFFE and sequence-based methods identified the principal proteins involved in the downstream pathways, showing they are broadly expressed across most Spongilla cell types, consistent with their diverse functional roles. This example highlights that CoFFE is able to vastly extend the information about already hypothesised signalling pathways in cell types by annotating proteins showing extensively divergent sequences.

Redox metabolism and control in myopeptidocytes

The mesohyl of sponges is a collagenous, dynamic tissue forming large parts of the body between pinacocytes and the feeding choanocyte chambers [30]. Musser et al. identified five novel mesenchymal cell types [9] in Spongilla. Among them, myopeptidocytes are an abundant, largely uncharacterised cell type, forming long projections that contact other cells. Sequence-based annotations suggested the generation and degradation of hydrogen peroxide in myopeptidocytes, taking into account the expression of dual oxidase (Duox1), its maturation factor (DuoxA), and Catalase (Cat) [31]. Reduced states of metal ions, such as Cu\(^{+}\), imported by copper transporters such as S1C31a2, are able to react with H\(_2\)O\(_2\) (Fenton reaction) [32].
leading to the generation of hydroxyl radicals, prominent reactive oxygen species (ROS) in cells (Fig. 3), supported by the presence of Cyba [33]. However, further roles of ROS metabolism and function had so far remained unclear.

CoFFE identifies potential iron-based generation of $H_2O_2$ in myopeptidocytes by detecting ferric-chelate reductase (FRRS1 and FRRS1L), which recycles Fe$^{3+}$ to its reduced state. At the same time, CoFFE proposes downstream metabolism and function of the ROSs: Disulfide oxidoreductase (DsBB) as well as Flavin carrier protein (FLC) both play a role in oxidative protein folding [34,35]. NmrA-like acts as a redox sensor in the cell [?]. Consistent with a possible redox regulation role, myopeptidocytes express a range of additional ROS metabolising and responsive proteins: Sulfiredoxin 1 (SRXN1) promotes resistance against oxidative stress damage [36] while AP endonuclease 1 (APEX1) protects against ROS induced DNA damage [37,38]. We also identified a glutathione S-transferase member (HPGDS) as well as an additional member of methionine sulfoxide reductases (msrA), which are important enzymes involved in the repair of proteins damaged by oxidative stress [39,40]. Finally, myopeptidocytes also express the LDL-receptor LRP2, which is known to bind apolipoprotein M (ApoM) [41]. Notably, CoFFE also detected ApoM in myopeptidocytes, suggesting a role in lipid metabolism. Consistent with this, Musser et al. documented round inclusions in myopeptidocytes that may represent lipid droplets. Lipid metabolism and redox control is tightly coupled in peroxisomes which are responsible for beta-oxidation of long-chain fatty acid [42].

Finding digestive roles for the uncharacterised mesocytes

Electron microscopy studies of the Spongilla feeding behaviour have demonstrated that prey endocytosis is initially carried out by pinacocytes and choanocytes. Food particles are passed on to digestive and phagocytic cells in the mesohyl, which are then thought to release undigested pellets into the excurrent canals [43]. These food particles often represent bacteria and algae, which are protected by polysaccharide and glycoprotein cell walls [44]. Breaking down cell walls would allow the sponge to extract additional energy and boost filter-feeding efficiency. Mesocytes are newly discovered sponge cell types and functionally uncharacterised medium-sized cells, whose name refers to their sparse location in the mesenchymal mesohyl [9].

Our new CoFFE annotations of previously unidentified mesocyte markers include genes such as expansin (yoaJ), glucan endo-1,3-beta-glucosidase (BG3), and spore cortex-lytic enzyme (sleB),
Table 1. Hydrolytic enzymes in *Spongilla* mesocytes.

| Annotation               | Annotation origin | Taxonomy     | Function                                                                 |
|--------------------------|-------------------|--------------|--------------------------------------------------------------------------|
| Aminohydrolase           | legacy EggNOG     | Bacteria     | hydrolase acting on aminogroups                                           |
| GMHA                     | legacy EggNOG     | Bacteria     | xanthan biosynthesis                                                     |
| Metallopeptidase M20     | legacy EggNOG     | Bacteria     | metallopeptidase                                                         |
| yoaJ                     | refined by CoFFE  | Bacteria     | cell wall degradation                                                    |
| sleB                     | refined by CoFFE  | Bacteria     | hydrolase activity, cell wall organization                               |
| T5orf172                 | new annotation    | Bacteria     | hydrolase activity                                                       |
| BG3                      | new annotation    | Streptophyta | cellulase activity                                                       |
| cellulase A family member | new annotation  | Streptophyta | cellulase activity                                                       |
| Endochitinase            | new annotation    | Streptophyta | endochitinase                                                            |
| Chitinase class I        | new annotation    | Streptophyta | endochitinase                                                            |

all hydrolases that specifically degrade cell walls, cellulase, chitin, other polysaccharides [45–48] and proteins (Table 1). It is thus tempting to speculate that mesocytes represent cells specialised to digest polysaccharides that are otherwise difficult to hydrolyse.

Interestingly, enzymes such as cellulases are famously absent from the metazoan digestive repertoire [45]. CoFFE annotations of these mesocyte marker genes reveal many have most structural similarity to bacteria and plant/algal proteins, suggesting these may have arisen via horizontal gene transfer (HGT). In line with this, recent evidence from nematodes showed they expanded their dietary options via the HGT of a cellulase gene of eukaryotic origin [49]. Furthermore, a previous study analysing origins of genes in the marine demosponge *Amphimedon queenslandica* identified a high proportion of polysaccharide hydrolases and metallopeptidases originating from HGTs [50].

Analysis of the GC-content and codon usage of genes shown in Table 1 compared to the entire *Spongilla* transcriptome revealed no significant deviation for putative HGT genes (Figure 3A), consistent with the hypothesis that they are present in sponge genomes. To confirm this and explore the relative timing of HGT events, we searched for homologs of all proteins shown in Table 1 in openly available genomes of sponge species (Figure 3B). We detected every protein...
in at least one other sponge. Corroborating previous studies, we identified hydrolases present in nearly all examined sponges. The only protein present in all analysed sponges is a metallopeptidase, similar to that detected in ctenophore comb jellies [51], indicating this transfer may have occurred prior to the last common ancestor of all animals. Strikingly, we found that three hydrolases were shared by only freshwater sponges, suggesting mesocyte specific functions may have expanded via HGT events after colonisation of new freshwater environments. Finally, the presence of predicted chitinases is intriguing, as chitin has been shown to be an important structural component of *Spongilla lacustris* [52]. This finding suggests another possible role for mesocytes as structural remodelers of the sponge skeleton made of polysaccharides.

**Identifying well-folded, unannotated proteins in *Spongilla lacustris***

Using the CoFFE pipeline we annotated a total of 26344 out of 41943 predicted proteins. The remaining 15599 unannotated proteins were relatively short, and may represent incomplete fragments, untranslated sequences, or de novo lineage-specific genes. Notably, we found XX unannotated proteins with a pLDDT score greater than 70, indicating well-folded structures. Although many of these had poor FoldSeek alignments falling below our accepted bit score threshold, 316 unannotated and well-folded predicted proteins had no hit using FoldSeek. Manual inspection revealed that the overwhelming majority of these are predicted to be long helices, except for 35 non-helical structures. To ensure these hadn’t somehow eluded sequence similarity searches, we used NCBI BLASTp to identify potential homologs in the NR database, even very remote ones (Suppl. Tabel 1). Seven of the sequences find no annotation nor get significant PFAM domain hits, but are broadly expressed in sponge cell types, presenting prime candidates for truly novel structures.
Fig 2. Signalling pathways in *Spongilla* pinacocytes A) Dotplot of pinacocyte signalling and effectors. Cell types of the pinacocyte family are highlighted by a red square. Genes on blue background are annotated by sequence-based methods (‘legacy’). Genes on yellow background are annotated by CoFFE. B) Structural alignment of *Spongilla* FGF (blue, 61 - 230 aa) with *Gallus gallus* FGF4 (UniProtID: P48804) (yellow, 54 - 194 aa) (RMSD = 0.89 over 543 atoms). C) Structural alignment of *Spongilla* ephrin (blue, 1 - 153 aa) with *C. elegans* efn-3 (UniProtID: Q19475) (yellow, 29 - 179 aa) (RMSD = 1.59 over 580 atoms). In both cases the structural similarity is apparent despite their low sequence identity of 11.8% and 22% respectively. Superpositions were created using the super command in PyMOL (V2.3.5)[]. Cell type abbreviations: incPin - incurrent pinacocytes; apmPin - apendopinacocytes; lph - lophocytes; basPin - basopinacocytes; scp - sclerophorocytes; met - metabolocytes; chb - choanoblasts; choanocytes; apo - apopylar cells; myp - myopeptidocytes; amb - amoebocytes; grl - granulocytes; nrd - neuroid cells; mes - mesocytes; arc - archaeocytes; scl - sclerocytes.
**Fig 3.** **ROS metabolism and redox-control in Myopeptidocytes**

Myopeptidocytes differentially express multiple genes involved in redox control and ROS defence. Genes in black and green have been annotated using sequence based methods. Green proteins are known to interact. Genes in red have been identified using CoFFE.

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**Fig 4.** **Horizontal gene transfer of mesocyte marker genes**

A) Comparison of GC-content and codon usage of mesocyte marker genes of non-metazoan origin with whole *Spongilla* transcriptome. B) Homology search of mesocyte marker genes in various sponge species. The green dot denotes the last common freshwater sponge ancestor.
Conclusions-Discussion

By predicting the protein structures of the entire proteome of the freshwater sponge *Spongilla lacustris* and aligning them against all currently available structures, we were able to increase the annotation of the proteome by approx. 50% (60% of the proteome in total, up from 40% when only using sequence information). For proteins with annotation from both sources, we found that in more than 90% of cases, sequence-based and structure-based annotation agreed at least to the protein superfamily level.

To explore the usefulness of structural similarity for functional annotation we revisited the annotation of marker genes previously identified from *Spongilla* scRNA-seq data. In the epithelial pinacocyte family, this significantly improved sequence-based annotation via the detection of key players in FGF, TGF-beta, as well as ephrin signalling. For the digestive peptidocyte family, we were able to infer complex modes of redox regulation as an important function of the myopeptidocytes. Finally, we detected polysaccharide and protein hydrolyzing enzymes of bacterial and plant/algal origin in the enigmatic mesocytes which strongly suggests that these digest cell walls of endocytosed prey. The mesocyte example also reveals cell types whose functional role is largely defined by genes acquired via HGTs from other pro- and eukaryotic organisms. This is remarkable both from a cell type evolutionary perspective and because such innovation had previously escaped our notice due to extensive sequence divergence following the HGT event.

The lack of reliable functional annotation has so far been a major bottleneck in the analysis of non-model species, in particular those separated from traditional models by large phylogenetic distances. Here, we demonstrate that by exploiting the evolutionary conservation of protein structure it is possible to dramatically improve protein functional annotations in non-model species. The pipeline described here can thus be used to query highly informative candidate genes from proteomics or single-cell -omics experiments. Although protein structural predictions for an entire proteome might be outside the technical capabilities of many labs due to the cost of procuring the necessary hardware, plans to expand the AlphaFold database [1] have been announced. Given the power of AlphaFold and the huge potential of expanded annotations, it is reasonable to expect that in the future protein sequences deposited in public databases will automatically receive predicted structures, paving the way for unique insights.
into biological functions across the tree of life.
Materials and methods

In all boxplots, the box extends from the lower to upper quartile values of the data, with a line at the median. The whiskers represent the 5 – 95% percentiles.

Sequence-based annotation of the *Spongilla lacustris* proteome

Juvenile freshwater sponges (*Spongilla lacustris*), grown from gemmules, were used for bulk RNA isolation and sequencing. *De novo* transcriptome assembly with Trinity, returned 62180 putative isoforms, covering 95.2% of Metazoan BUSCOs [53]. To identify putative proteins, Transdecoder [54] (version 3.0.1) was used with a minimum open reading frame length of 70 amino acids, resulting in 41945 putative proteins. The longest putative protein per gene ID was kept. The resulting predicted proteome was annotated by EggNOG mapper [14,55] (v2.1.7, default settings) via the website.

Legacy annotation

Musser *et al.* [9] used the putative proteins to create a phylome by constructing gene/protein trees for each protein (available online). The phylome information was used to refine the assignment of transcripts to genes. In some cases, 3’ and 5’ fragments of a gene were assigned to two different transcripts. These fragments were merged into the same merged gene name using the gene tree information. Functional annotations were supplemented by EggNOG mapper (v1) and blastp searches against human RefSeq (default parameters). This annotation was used in the original *Spongilla* scRNA sequencing publication and is present in the single-cell data. The legacy annotation was used for the single-cell data analysis, but not for the comparison between the CoFFE pipeline and the sequence-based annotation transfer.

Structure-based annotation of the *Spongilla lacustris* proteome

CoFFE and constituent tools

We designed a simple pipeline that goes from sequence to predicted structure to functional annotation (symbolically, ColabFold-FoldSeek-EggNOG, or CoFFE), and used it to annotate the *Spongilla lacustris* predicted proteome. We used ColabFold to predict structures and subsequently aligned the predicted structures against all currently available (solved and
predicted) protein structures using FoldSeek. Finally, we used structural similarity to transfer annotations from the best structural targets to their corresponding *Spongilla* protein queries. In the following we show an overview of the tools in use and a more detailed description of the CoFFE pipeline.

**MMseqs2** [56] is a software suite for sequence-sequence, sequence-profile, and profile-profile search and clustering. It is orders of magnitude faster than BLAST at the same sensitivity and is widely adopted [14].

**AlphaFold2** [7] is a neural network-based model that predicts protein three-dimensional structures from sequence, regularly achieving atomic accuracy even in cases where no similar structure is known. AlphaFold is widely considered to have revolutionised the field of structural bioinformatics, greatly outperforming the state of the art in the most recent iteration of the CASP challenge [57].

AlphaFold quantifies prediction confidence by pLDDT, the predicted local distance difference test on the $\text{C}_\alpha$ atoms. Regions with pLDDT > 90 are modelled to high accuracy; regions with 70 < pLDDT < 90 should have a generally good backbone prediction; regions with 50 < pLDDT < 70 are low confidence and should be treated with caution; regions with pLDDT < 50 should not be interpreted and probably are disordered.

**ColabFold** [10] is a pipeline that combines fast homology searches via MMseqs2 [56] with AlphaFold2 [7] to predict protein structures 40 to 60 times faster than the original AlphaFold2.

**FoldSeek** [8] enables fast and sensitive comparison of large structure databases. FoldSeek’s key innovation lies in the appropriate translation of structure states to a small alphabet, thus gaining access to all the heuristics of sequence search algorithms.

## The CoFFE pipeline

A visual representation of the CoFFE pipeline can be found in supplement figure XX. To predict structures, we adapted the ColabFold pipeline as outlined in [https://colabfold.mmseqs.com/](https://colabfold.mmseqs.com/)

**Multiple sequence alignment generation:** We downloaded reference sequence databases (UniRef30, ColabFold DB) ([databases.sh](https://colabfold.mmseqs.com/databases.sh)) and calculated indices locally ([create_index.sh](https://colabfold.mmseqs.com/create_index.sh)). We were interested in homology detection at the limit of the twilight zone, so UniRef30, a 30% sequence identity clustered database based on UniRef100 [58].
was the adequate choice. Similarly, we indexed the *Spongilla* predicted proteome (spongilla_create_index.sh). We calculated MSAs for each *Spongilla* predicted protein using MMseqs2 [56] (align.sh) adapted from colabfold_search.sh).

**Structure prediction:** We predicted structures for all *Spongilla* predicted proteins using ColabFold [10] as a wrapper around AlphaFold2.

We split the MSAs in 32 batches and submitted each one to the EMBL cluster system (managed by slurm [59]); we used default arguments but added --stop-at-score 85. (milot.sh) M. Mirdita, personal communication). The calculations were done on NVIDIA A100 GPUs, on computers running CentOS Linux 7. We used GCC [60] version 10.2.0 and CUDA version 11.1.1-GCC-10.2.0 [61]. We processed the resulting PDB-formatted model files with Biopython’s PDB module [62].

**Structure search and annotation transfer:** Structural search was conducted using FoldSeek. FoldSeek allows fast comparison of large structural databases. We downloaded PDB, SwissProt, and AlphaFold DB (fs_pdb.sh fs_sp.sh and fs_afdb.sh, respectively). For each *Spongilla* protein we kept the best-scoring AlphaFold2 model, and used them to construct a FoldSeek database fs_query.sh. These models were then used to search against the three structural databases. For each search we kept the FoldSeek hit with the highest corrected bit score in each database and aggregated the three result tables (AlphaFoldDB, PDB, SwissProt) into one. We imposed a bit score cutoff (1xe-05) on FoldSeek hits based on their bimodal distribution. Annotations of the best hits were gathered from either UniProt via its API [63] or through EggNOG mapper (v2.1.7, default settings) [14] by using the sequences of the best structural hits (pulled by UPIMAPI [64]). To facilitate downstream analysis, we extracted summary tables from each resource type. This procedure can be found in the corresponding notebook. A total of 1401 proteins received sequence annotation but their FoldSeek best hits were below the bit score cutoff.

**Differential gene expression in single-cell transcriptomics data**

We obtained the processed Seurat file from [9], and downloaded the lists of differentially expressed genes of clusters, cell types, and cell type clades from the supplemental material of the same publication (Supplemental Data S1 to S3; file science.abj2949.data_s1.xlsx; tabs “Diff. exp. 42 clusters”, “Diff. Exp. cell types”, “Cell type clade genes (OU tests”). The single-cell
data operates on the level of genes, so we transformed the sequence-derived and CoFFE annotations by merging isoform entries and keeping the entry with the best bit score.

We used the legacy annotation included in the file (phylome, EggNOG-mapper, and BLASTp-based), the sequence-derived annotation, and the CoFFE output to propose names for *Spongilla* proteins. We prioritized sequence-derived annotations (legacy annotation, EggNOG preferred name, EggNOG description) and fell back to CoFFE (CoFFE preferred name, CoFFE description) when there were none. We produced dotplots for the top 200 differentially expressed genes in each cell type and manually inspected them. We focused on the terminal (named) cell types; owing to the presence of continuously differentiating stem cells, the single-cell data contains many clusters of maturing or differentiating cells whose expression patterns do not distinguish them from their mature counterparts. Code and detailed explanations are available in the corresponding notebook.

**Protein structure visualisation and superposition**

In order to visualize predicted structures from the *Spongilla* proteome, we used PyMOL Version 2.3.5 [65]. Superposition with their respective best Foldseek hit was carried out using the **super** command. **super** creates sequence-independent superpositions and is more reliable for protein pairs with low sequence similarity.

**Detection of ephrin orthologs in *Spognilla* using hmmer**

To validate the ephrin ortholog detected by CoFFE in *Spongilla*, we recapitulated a previous effort to detect ephrins in different species using extensive hmmer protein profile search [26]. For this, we performed a hmmer search (v3.3.2) using the ephrin Pfam profile (PF00812) against the *Spongilla* proteome using default settings.

**0.1 Detecting HGT in Spongilla**

To detect HGT we followed the proposal by Degnan [66] and sought to get multiple indications for HGT. In particular: phylogenetic evidence that the candidate gene is more closely related to foreign than to animal genes; genome data showing the candidate gene assembles into a contiguous stretch of DNA with neighboring genes unambiguously of animal origin (this requires, of course, the availability of a sequenced and assembled animal genome; the more
complete the assembly, the more confident the HGT identification); and gene sequence revealing metazoan-like compositional traits, including presence of introns, GC content and codon usage. Where possible, gene expression data showing active transcription of candidate genes in animal cell nuclei can enormously strengthen a case, and also addresses the issue of whether or not the HGT-acquired gene is active in its new genomic context.

We used https://spongebase.net to obtain transcriptomes and genomes for 13 sponge species, and obtained a 14th one directly from its repository (https://bitbucket.org/molpalmuc/tethya_wilhelma-genome/). We built sequence databases with MMseqs2 version 12-113e3 and searched against them (mmseqs easy-search) with the protein sequences of all isoforms of the HGT candidates. The resulting alignments were filtered to keep the best-scoring alignment per species per gene.

The table of best hits per genome was visualised in Python, and a sponge phylogeny was added manually based on 67 (also A. Riesgo, personal communication).
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A Extracting quality indicators from the MAF pipeline

To get a better understanding of the relationship between the various metrics, we assessed the correlation between them (Fig. 5). We hoped to find a combination of measures that together would be a robust indicator of prediction quality for MAF.

Most obviously, query length, (uncorrected) structural bit score, and (uncorrected) sequence bit score correlate very strongly. Longer proteins will have more or longer aligned regions with their targets, and as the bit score is additive, will accumulate higher bit scores, regardless of whether the alignment happens in structure or sequence space. It seems these measures can be used interchangeably; out of principle, we would prefer to use query length, as it is the least derived.

A way to overcome the dependence on length is to normalize alignment scores by the length of the aligned region \( \text{(corrected bit score - (FS))} \). This measure correlates strongly with the percent structure state identity. This is equivalent to percent sequence identity correlating strongly with the bit score in a sequence similarity search, and is not surprising at all. It seems that these measures can also be used interchangeably, but we would prefer to use percent structure identity, since it correlates less strongly with the other measures in the plot.

The rest of the measures (pLDDT, EggNOG bit score of FoldSeek best hit, percent aligned query, and input MSA size) don’t correlate too strongly to each other. We decided to use them as-is.

Three things become apparent: first, higher pLDDT scores generally correlate with higher FoldSeek bit scores. This means that FoldSeek does a better job of identifying structural similarity for well-folded proteins. This seems intuitive: the atomic coordinates of well-folded proteins will be more deterministic than disordered proteins, so well-folded orthologs will overlap structurally over their entire length, while heavily disordered orthologs might only overlap structurally over very short patches.

Second, agreement of structural and sequence annotation correlates with higher bit score. This is also ok: the peptides that share the most specific orthogroup are the ones that are recognisably the same, regardless if we look at sequence or structure; peptides that at least share the root orthogroup are the ones with high sequence conservation but lower

The more confident the overlap between structure and sequence annotation, the higher the
average bit score is. This happens in all pLDDT buckets, but has a stronger effect the higher the average pLDDT is.

FoldSeek is more confident in identifying structural similarity for peptides whose structure annotation agrees with the sequence annotation to the most specific orthogroup. Peptides with root orthogroup overlap come in second, and peptides with PFAM domain overlap third. These categories correspond to structures being in the same protein family or superfamily, or at least sharing a couple of domains, indicating three broad levels of structural similarity.

Three, peptides without sequence annotation and peptides where structure and sequence annotation disagree don’t have meaningful differences in their FoldSeek bit scores. This happens regardless of average pLDDT.
| isoform   | q len | #MSA | %seq. id. | bit score | plddt | BLASTp  |
|-----------|-------|------|-----------|-----------|-------|---------|
| c88419_g1_i1 | 154   | 462  | 0.337     | 580.0     | 85.01 | CRN5    |
| c104256_g1_i1 | 375   | 163  | 0.272     | 785.0     | 82.42 | CRN5    |
| c104256_g1_i2 | 299   | 184  | 0.267     | 781.0     | 90.35 | CRN5    |
| c104256_g1_i3 | 336   | 167  | 0.262     | 777.0     | 88.39 | CRN5    |
| c104256_g1_i4 | 332   | 158  | 0.263     | 796.0     | 90.26 | CRN5    |
| c104256_g1_i6 | 253   | 196  | 0.275     | 774.0     | 92.81 | CRN5    |
| c94352_g2_i1 | 127   | 1725 | 0.500     | 299.0     | 86.06 | DD3-3   |
| c89711_g1_i1 | 144   | 2424 | 0.441     | 380.0     | 86.73 | DD3-3   |
| c112476_g1_i1 | 578   | 3328 | 0.355     | 2091.0    | 81.55 | DD3-3   |
| c103626_g1_i2 | 187   | 1994 | 0.416     | 522.0     | 89.04 | DD3-3   |
| c101972_g1_i1 | 778   | 3481 | 0.378     | 2536.0    | 85.58 | DD3-3   |
| c101972_g1_i2 | 778   | 3481 | 0.378     | 2534.0    | 85.58 | DD3-3   |
| c101972_g1_i3 | 778   | 3481 | 0.378     | 2537.0    | 85.54 | DD3-3   |
| c104453_g2_i1 | 359   | 368  | 0.177     | 662.0     | 75.80 | Foll. epith. yolk prot. subunit |
| c104453_g2_i2 | 280   | 356  | 0.172     | 658.0     | 89.90 | Foll. epith. yolk prot. subunit |
| c104453_g2_i3 | 340   | 378  | 0.185     | 634.0     | 78.61 | Foll. epith. yolk prot. subunit |
| c104453_g2_i4 | 236   | 374  | 0.183     | 594.0     | 83.73 | Foll. epith. yolk prot. subunit |
| c98985_g1_i2 | 73    | 1852 | 0.138     | 45.0      | 70.12 | XPO5 (Exportin-5) |
| c88005_g3_i1 | 130   | 3213 | 0.274     | 363.0     | 94.66 | Hemopexin repeat-containing protein |
| c70965_g1_i1 | 169   | 299  | 0.209     | 354.0     | 77.62 | Pinocchio |
| c103612_g1_i1 | 89    | 1253 | 0.152     | 46.0      | 73.88 | Kinesin-like protein KIF23 |
| c100331_g1_i3 | 312   | 257  | 0.179     | 649.0     | 86.28 | BamB |
| c100031_g2_i1 | 281   | 43509| 0.219     | 821.0     | 82.62 | Hydralysin-2-like |
| c93555_g2_i2 | 77    | 439  | 0.109     | 80.0      | 87.77 | ? |
| c86044_g1_i1 | 87    | 7052 | 0.077     | 90.0      | 88.87 | ? |
| c83472_g1_i1 | 98    | 553  | 0.177     | 90.0      | 84.77 | ? |
| c108675_g1_i1 | 73    | 3174 | 0.224     | 51.0      | 78.17 | ? |
| c10071_g1_i2 | 69    | 1    | 0.123     | 61.0      | 81.77 | ? |
| c102929_g1_i1 | 79    | 43996| 0.152     | 85.0      | 71.74 | ? |

**Table 2.** High confidence structures with non-helical appearance. The last column contains a summary of the significant BLASTp results in the nr database. Question marks denote peptides who did not find any named sequence homologs or significant PFAM domain hits.
Fig 5. **Pairwise correlation of various metrics.** *Spongilla MSA size (AF):* size of the input MSA used for AlphaFold. *pLDDT (AF):* average pLDDT of the predicted structure. *best match seq. id. (FS):* structure state identity in the alignment between the query and the best FoldSeek target. *corrected bit score (FS):* the FoldSeek bit score divided by alignment length. *Spongilla query length (AF):* the query length of the input peptide. *bit score (FS):* bit score of the FoldSeek alignment between the query *Spongilla* structure and the structure of the best FoldSeek target. *bit score (EggNOG-seq):* bit score of the EggNOG alignment between the query *Spongilla* peptide sequence and the best target sequence. *rel. ali. length (FS):* percentage of the query (*Spongilla*) structure aligned with the best target structure. *bit score (EggNOG-FS):* bit score of the sequence alignment between the sequence of the best FoldSeek target and the EggNOG database.
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