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Mapping single-cell atlases throughout Metazoa unravels cell type evolution — Source link

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18 Abstract

19 Comparing single-cell transcriptomic atlases from diverse organisms can elucidate the 20 origins of cellular diversity and assist the annotation of new cell atlases. Yet, 21 comparison between distant relatives is hindered by complex gene histories and 22 diversifications in expression programs. Previously, we introduced the self-assembling 23 manifold (SAM) algorithm to robustly reconstruct manifolds from single-cell data 24 (Tarashansky et al., 2019). Here, we build on SAM to map cell atlas manifolds across 25 species. This new method, SAMap, identifies homologous cell types with shared 26 expression programs across distant species within phyla, even in complex examples 27 where homologous tissues emerge from distinct germ layers. SAMap also finds many 28 genes with more similar expression to their paralogs than their orthologs, suggesting 29 paralog substitution may be more common in evolution than previously appreciated. 30 Lastly, comparing species across animal phyla, spanning mouse to sponge, reveals 31 ancient contractile and stem cell families, which may have arisen early in animal 32 evolution.

33 Introduction

34 There is much ongoing success in producing single-cell transcriptomic atlases to 35 investigate the cell type diversity within individual organisms (Regev et al., 2017). With 36 the growing diversity of cell atlases across the tree of life (Briggs et al., 2018; Cao et al., 37 2019; Fincher et al., 2018; Hu et al., 2020; Musser et al., 2019; Plass et al., 2018; Siebert 38 et al., 2019; Wagner et al., 2018), a new frontier is emerging: the use of cross-species 39 cell type comparisons to unravel the origins of cellular diversity and uncover speciesspecific cellular innovations (Arendt et al., 2019; Shafer, 2019). Further, these 40 41 comparisons promise to accelerate cell type annotation and discovery by transferring 42 knowledge from well-studied model organisms to under-characterized animals.

43

44 However, recent comparative single-cell analyses are mostly limited to species within the 45 same phylum (Baron et al., 2016; Geirsdottir et al., 2019; Sebé-Pedrós et al., 2018; 46 Tosches et al., 2018). Comparisons across longer evolutionary distances and across 47 phyla are challenging for two major reasons. First, gene regulatory programs diversify 48 during evolution, diminishing the similarities in cell type specific gene expression patterns. 49 Second, complex gene evolutionary history causes distantly related organisms to share 50 few one-to-one gene orthologs (Nehrt et al., 2011), which are often relied upon for 51 comparative studies (Briggs et al., 2018; Shafer, 2019). This effect is compounded by the 52 growing evidence suggesting that paralogs may be more functionally similar than 53 orthologs across species, due to differential gain (neo-functionalization), loss (non-54 functionalization), or partitioning (sub-functionalization) events among paralogs (Nehrt et 55 al., 2011; Prince & Pickett, 2002; Stamboulian et al., 2020; Studer & Robinson-Rechavi, 2009). 56

57

Here, we present the Self-Assembling Manifold mapping (SAMap) algorithm to enable 58 59 mapping single-cell transcriptomes between phylogenetically remote species. SAMap 60 relaxes the constraints imposed by sequence orthology, using expression similarity 61 between mapped cells to infer the relative contributions of homologous genes, which in 62 turn refines the cell type mapping. In addition, SAMap uses a graph-based data 63 integration technique to identify reciprocally connected cell types across species with 64 greater robustness than previous single-cell data integration methods (Haghverdi et al., 65 2018; Hie et al., 2019; Polański et al., 2019; Stuart et al., 2019).

66

67 Using SAMap, we compared seven whole-body cell atlases from species spanning animal 68 phylogeny, which have divergent transcriptomes and complex molecular homologies (Figure 1A-B and Supplementary Table 1). We began with well-characterized cell types 69 70 in developing frog and fish embryos. We found broad concordance between 71 transcriptomic signatures and ontogenetic relationships, which validated our mapping 72 results, yet also detected striking examples of homologous cell types emerging from 73 different germ layers. We next extended the comparison to animals from the same phylum 74 but with highly divergent body plans, using a planarian flatworm and a parasitic blood 75 fluke, and found one-to-one homologies even between cell subtypes. Comparing all 76 seven species from sponge to mouse, we identified densely interconnected cell type 77 families broadly shared across animals, including contractile and stem cells, along with 78 their respective gene expression programs. Lastly, we noticed that homologous cell types 79 often exhibit differential expression of orthologs and similar expression of paralogs,

suggesting that the substitution and swapping of paralogs in cell types may be more common in evolution than previously appreciated. Overall, our study represents an important step towards analyzing the evolutionary origins of specialized cell types and their associated gene expression programs in animals.

84 Results

85 The SAMap algorithm

SAMap iterates between two modules. The first module constructs a gene-gene bipartite 86 87 graph with cross-species edges connecting homologous gene pairs, initially weighted by 88 protein sequence similarity (Figure 1C). In the second module, SAMap uses the gene-89 gene graph to project the two single-cell transcriptomic datasets into a joint, lower-90 dimensional manifold representation, from which each cell's mutual cross-species 91 neighbors are linked to stitch the cell atlases together (**Figure 1D**). Then, using the joint 92 manifold, the expression correlations between homologous genes are computed and 93 used to reweight the edges in the gene-gene homology graph in order to relax SAMap's 94 initial dependence on sequence similarity. The new homology graph is used as input to 95 the subsequent iteration of SAMap, and the algorithm continues until convergence, 96 defined as when the cross-species mapping does not significantly change between 97 iterations (Figure 1E).

98

99 algorithm overcomes several challenges inherent to mapping This sinale-cell 100 transcriptomes between distantly related species. First, complex gene evolutionary 101 history often results in many-to-many homologies with convoluted functional relationships 102 (Briggs et al., 2018; Nehrt et al., 2011). SAMap accounts for this by using the full 103 homology graph to project each dataset into both its own and its partner's respective principal component (PC) spaces, constructed by the SAM algorithm, which we previously 104 105 developed to robustly and sensitively identify cell types (Tarashansky et al., 2019). The 106 resulting within- and cross-species projections are concatenated to form the joint space.

For the cross-species projections, we translate each species' features into those of its partner, with the expression for individual genes imputed as the weighted average of their homologs specified in the gene-gene bipartite graph. Iteratively refining the homology graph to only include positively correlated gene pairs prunes the many-to-many homologies to only include genes that are expressed in the same mapped cell types.

112

Second, frequent gene losses and the acquisitions of new genes result in many cell type gene expression signatures being species-specific, limiting the amount of information that is comparable across species. Restricting the analysis of each dataset to only include genes that are shared across species would result in a decreased ability to resolve cell types and subtypes with many species-specific gene signatures. SAMap solves this problem by constructing the joint space through the concatenation of within- and crossspecies projections, thus encoding all genes from both species.

120

121 Lastly, the evolution of expression programs gradually diminishes the similarity between 122 homologous cell types. To account for this effect, SAMap links cell types across species 123 while tolerating their differences. Cells are mapped by calculating each of their k mutual 124 nearest cross-species neighbors in the combined projection. To establish more robust 125 mutual connectivity, we integrate information from each cell's local, within-species 126 neighborhood (Figure 1D), overcoming the inherent stochasticity of cross-species 127 correlations. Two cells are thus defined as mutual nearest cross-species neighbors when 128 their respective neighborhoods have mutual connectivity. It is important to note that the 129 magnitude of connections is not directly calculated from their expression similarity,

allowing cell types with diverged expression profiles to be tightly linked if they are amongeach other's closest cross-species neighbors.

132

133 Paralog substitutions are prevalent between homologous cell types in frog and fish 134 We first applied SAMap to the *Xenopus* and zebrafish atlases, which both encompass 135 embryogenesis until early organogenesis (Briggs et al., 2018; Wagner et al., 2018). 136 Previous analysis had linked cell types between these two organisms by matching 137 ontogeny, thereby providing a reference for comparison. SAMap produced a combined 138 manifold with a high degree of cross-species alignment while maintaining high resolution 139 for distinguishing cell types in each species (Figure 2A). We measured the mapping 140 strength between cell types by calculating an alignment score (edge width in **Figure 2B** 141 and color map in **Figure 2C**), defined as the average number of mutual nearest cross-142 species neighbors of each cell relative to the maximum possible number of neighbors.

143

144 SAMap revealed broad agreement between transcriptomic similarity and developmental 145 ontogeny, linking 26 out of 27 expected pairs based on previous annotations (Figure 2B 146 and **Supplementary Table 2**) (Briggs et al., 2018). The only exception is the embryonic 147 kidney (pronephric duct/mesenchyme), potentially indicating that their gene expression 148 programs have significantly diverged. In addition, SAMap succeeded in drawing parallels 149 between the development of homologous cell types and matched time points along 150 several cell lineages (Figure 2C). While the concordance was consistent across cell 151 types, we noticed that the exact progression of developmental timing can vary, suggesting 152 that SAMap can quantify heterochrony with cell type resolution.

153

154 SAMap also weakly linked several closely related cell types with different ontogeny. For 155 example, optic cells from both species are also connected to eye primordium, frog skeletal 156 muscles to fish presomitic mesoderm, and frog hindbrain to fish forebrain/midbrain. 157 Notable exceptions also included mapped secretory cell types that differ in their 158 developmental origin and even arise from different germ layers (black edges in Figure 159 **2B**). They are linked through a large set of genes including conserved transcription factors 160 (e.g., foxa1 (Dubaissi et al., 2014), grhl (Miles et al., 2017)) and proteins involved in 161 vesicular protein trafficking (Figure 2 – figure supplement 1). This observation supports 162 the notion that cell types may be transcriptionally and evolutionarily related despite having 163 different developmental origins (Arendt et al., 2016).

164

165 To benchmark SAMap performance, we used Eggnog (Huerta-Cepas et al., 2019) to define one-to-one vertebrate orthologs between fish and frog and fed these gene pairs 166 167 as input to several broadly used single-cell data integration methods, Seurat (Stuart et 168 al., 2019), Liger (Welch et al., 2019), Harmony (Korsunsky et al., 2019), Scanorama (Hie 169 et al., 2019), and BBKNN (Polański et al., 2019). We found that they failed to map the two 170 atlases, yielding minimal alignment between them (Figure 2D and Figure 2 - figure 171 **supplement 2**). We also compared the results when restricting SAMap to using the one-172 to-one orthologs instead of the full homology graph. Even when removing the many-to-173 many gene homologies and the iterative refinement of the homology graph, we identified 174 similar, albeit weaker, cell type mappings. This suggests that, at least for the frog and fish

175 comparison, SAMap's performance is owed in large part to its robust, atlas stitching176 approach.

177

178 The key benefit of using the full homology graph is to enable the systematic identification 179 of gene paralogs that exhibit greater similarity in expression across species than their 180 corresponding orthologs. These events are expected to arise as the result of gene 181 duplications followed by diversification of the resulting in-paralogs (Studer & Robinson-182 Rechavi, 2009). In addition, genetic compensation by transcriptional adaptation, where 183 loss-of-function mutations are balanced by upregulation of related genes with similar 184 sequences (EI-Brolosy et al., 2019), could also result in this signature. In total, SAMap 185 selected 8,286 vertebrate orthologs and 7,093 eukaryotic paralogs, as enumerated by 186 Eggnog, for manifold alignment. Among these, 565 genes have markedly higher 187 expression correlations (correlation difference > 0.3) with their paralogs than their 188 orthologs (Figure 2E and Figure 2 – figure supplement 3), and 209 genes have 189 orthologs that are either completely absent or lowly-expressed with no cell-type specificity 190 (Supplementary Table 3), suggesting that they may have lost their functional roles at 191 some point and were compensated for by their paralogs. We term these events as 192 "paralog substitutions". SAMap linked an additional 297 homologous pairs not previously 193 annotated by orthology or paralogy, but which exhibited sequence similarity and high 194 expression correlations (>0.5 Pearson correlation). These likely represent unannotated 195 orthologs/paralogs or isofunctional, distantly related homologs (Gabaldón & Koonin, 196 2013). These results illustrate the potential of SAMap in leveraging single-cell gene 197 expression data for pruning the networks of homologous genes to identify evolutionary

substitution of paralogs and, more generally, identify non-orthologous gene pairs that may
 perform similar functions in the cell types within which they are expressed.

200

201 Homologous cell types between two flatworm species with divergent body plans

202 To test if we can identify homologous cell types in animals with radically different body 203 plans, we mapped the cell atlases of two flatworms, the planarian Schmidtea 204 mediterranea (Fincher et al., 2018), and the trematode Schistosoma mansoni, which we 205 collected recently (Li et al., 2020). They represent two distant lineages within the same 206 phylum but have remarkably distinct body plans and autecology (Laumer et al., 2015; 207 Littlewood & Waeschenbach, 2015). While planarians live in freshwater and are known 208 for their ability to regenerate (Reddien, 2018), schistosomes live as parasites in humans. 209 The degree to which cell types are conserved between them is unresolved, given the vast 210 phenotypic differences caused by the transition from free-living to parasitic habits 211 (Laumer et al., 2015).

212

SAMap revealed broad cell type homology between schistosomes and planarians. The schistosome had cells mapped to the planarian stem cells, called neoblasts, as well as most of the differentiated tissues: neural, muscle, intestine, epidermis, parenchymal, protonephridia, and *cathepsin*⁺ cells, the latter of which consists of cryptic cell types that, until now, have only been found in planarians (Fincher et al., 2018) (Figure 3A). These mappings are supported by both known cell type specific marker genes and numerous homologous transcriptional regulators (Figure 3B and Figure 3 – figure supplement 1).

221 We next determined if cell type homologies exist at the subtype level. For this, we 222 compared the neoblasts, as planarian neoblasts are known to comprise populations of 223 pluripotent cells and tissue-specific progenitors (Fincher et al., 2018; Zeng et al., 2018). 224 By mapping the schistosome neoblasts to a planarian neoblast atlas (Zeng et al., 2018), 225 we found that the schistosome has a population of neoblasts (ε -cells (Wang et al., 2018)) 226 that cluster with the planarian's pluripotent neoblasts, both expressing a common set of 227 TFs (e.g., soxp2, unc4, pax6a, gcm1) (Figure 3C-D). The ε-cells are closely associated 228 with juvenile development and lost in adult schistosomes (Wang et al., 2018), indicating 229 pluripotent stem cells may be a transient population restricted to their early developmental 230 stages. This is consistent with the fact that, whereas schistosomes can heal wounds, they 231 have limited regenerative ability (Wendt & Collins, 2016). SAMap also linked other 232 schistosome neoblast populations with planarian progenitors, including two populations 233 of schistosome neoblasts (denoted as μ (Tarashansky et al., 2019) and μ) to planarian 234 muscle progenitors, all of which express myoD, a canonical master regulator of 235 myogenesis (Scimone et al., 2017). These likely represent early and late muscle 236 progenitors, respectively, as µ-cells do not yet express differentiated muscle markers 237 such as *troponin*, whereas μ '-cells do (Figure 3 – figure supplement 2).

238

239 Cell type families spanning the animal tree of life

To compare cell types across broader taxonomic scales, we extended our analysis to include juvenile freshwater sponge (*Spongilla lacustris*) (Musser et al., 2019), adult *Hydra* (*Hydra vulgaris*) (Siebert et al., 2019), and mouse (*Mus musculus*) embryogenesis (Pijuan-Sala et al., 2019) atlases. In total, SAMap linked 1,051 cross-species pairs of cell

types, defined by the annotations used in each respective study. 95% of the cell type
pairs are supported by at least 40 enriched gene pairs, and 87% are supported by more
than 100 gene pairs, indicating that SAMap does not spuriously connect cell types with
limited overlap in expression profiles (Figure 4A).

248

249 We next extended the notion of cell type pairs to cell type trios, as mapped cell types gain 250 additional support if they share transitive relationships to other cell types through 251 independent mappings, forming cell type triangles among species. The transitivity of a 252 cell type pair (edge) or a cell type (node) can be quantified as the fraction of triads to 253 which they belong that form triangles (Figure 4B). The majority (81%) of cell type pairs 254 have non-zero transitivity independent of alignment score and the number of enriched 255 gene pairs (Figure 4 – figure supplement 1-2). Cell type pairs with fewer than 40 256 enriched gene pairs tend to have lower (<0.4) transitivity (Figure 4 – figure supplement 257 2). In addition, 16% of mapped cell type pairs have zero edge transitivity but non-zero 258 node transitivity, meaning that at least one of the cell types connects to only a single 259 member of an interconnected cell type group (Figure 4C). Such edges may be of lower 260 confidence as they should connect to other members of the same group and are thus 261 excluded from downstream analysis.

262

Among the interconnected groups of cell types, we identified families of contractile cells and neural cells (**Figure 4D**). Both cell type families are highly transitive compared to the overall graph transitivity (bootstrap p-value < 1×10^{-5}), meaning that their constituent cell types have more transitive edges within the group than outside the group (**Figure 4E**). In

267 addition, the dense, many-to-many connections within the contractile and neural families 268 are each supported by at least 40 enriched gene pairs (Figure 4F). Consistent with the 269 nerve net hypothesis suggesting a unified origin of neural cell types (Tosches & Arendt, 270 2013), the neural family includes vertebrate brain tissues, both bilaterian and chidarian 271 neurons, cnidarian nematocytes that share the excitatory characteristics of neurons (Weir 272 et al., 2020), and Spongilla choanocytes and apopylar cells, both of which are not 273 considered as neurons but have been shown to express postsynaptic-like scaffolding 274 machinery (Musser et al., 2019; Wong et al., 2019). The contractile family includes 275 myocytes in bilaterian animals, Hydra myoepithelial cells that are known to have 276 contractile myofibrils (Buzgariu et al., 2015), and sponge pinacocytes and 277 myopeptidocytes, both of which have been implicated to play roles in contractility (Musser 278 et al., 2019; Sebé-Pedrós et al., 2018). In contrast to the families encompassing all seven 279 species, we also found a fully interconnected group that contains invertebrate pluripotent 280 stem cells, including planarian and schistosome neoblasts, Hydra interstitial cells, and 281 sponge archeocytes (Alié et al., 2015). The lack of one-to-one connections across phyla 282 is in keeping with recent hypotheses that ancestral cell types diversified into families of 283 cell types after speciation events (Arendt et al., 2016, 2019). Our findings thus suggest 284 that these cell type families diversified early in animal evolution.

- 285
- 286 Transcriptomic signatures of cell type families

The high interconnectedness between cell types across broad taxonomic scales suggests that they should share ancestral transcriptional programs (Arendt et al., 2016; Tosches et al., 2018). SAMap identified broad transcriptomic similarity between bilaterian and non-

290 bilaterian contractile cells that extends beyond the core contractile apparatus. It links a 291 total of 23,601 gene pairs, connecting 5,471 unique genes, which are enriched in at least 292 one contractile cell type pair. Performing functional enrichment analysis on these genes. 293 we found cytoskeleton and signal transduction functions to be enriched (p-value < 10^{-3}) 294 based on the KOG functional classifications (Tatusov et al., 2003) assigned by Eggnog 295 (Figure 5A). These genes include orthology groups spanning diverse functional roles in 296 contractile cells, including actin regulation, cell adhesion and stability, and signaling 297 (Figure 5B and Supplementary Table 4), indicating that contractile cells were likely 298 multifunctional near the beginning of animal evolution.

299

300 We also identified several transcriptional regulators shared among contractile cells 301 (Figure 5B). Previously known core regulators involved in myocyte specification (Brunet 302 et al., 2016) were enriched only in bilaterian (e.g., myod, and tcf4/E12) or vertebrate 303 contractile cells (e.g., *mef2*). In contrast, we found homologs of Muscle Lim Protein (*Csrp*) 304 and Forkhead Box Group 1 (Larroux et al., 2008) enriched in contractile cells from all 305 seven species. The Fox proteins included FoxC, which is known to regulate cardiac 306 muscle identity in vertebrates (Brunet et al., 2016) and is contractile-specific in all species 307 except schistosome and Spongilla. Notably, we also identified FoxG orthologs to be 308 enriched in three of the four invertebrates (Figure 5 – figure supplement 1), suggesting 309 that FoxG may play an underappreciated role in contractile cell specification outside 310 vertebrates.

311

312 For the family of invertebrate stem cells, we identified 3,343 genes that are enriched in at 313 least one cell type pair and observed significant enrichment (p-value < 10^{-3}) of genes involved in translational regulation such as RNA processing, translation, and post-314 315 translational modification (Figure 5C). These genes form 979 orthology groups, 17% of 316 which are enriched in all cell types of this family (**Supplementary Table 4**). Importantly, 317 other stem cell populations in Hydra and planarian lineage-restricted neoblasts have 318 significantly reduced expression of these genes (Figure 5D). These results suggest that 319 SAMap identified a large, deeply conserved gene module specifically associated with 320 multipotency.

321

322 Discussion

323 Cell types evolve as their gene expression programs change either as integrated units or 324 via evolutionary splitting that results in separate derived programs. While this notion of 325 coupled cellular and molecular evolution has gained significant traction in the past years, 326 systematically comparing cell type-specific gene expression programs across species 327 has remained a challenging problem. Here, we map single-cell atlases between 328 evolutionarily distant species in a manner that accounts for the complexity of gene 329 evolution. SAMap aligns cell atlases in two mutually reinforcing directions, mapping both 330 the genes and the cells, with each feeding back into the other. This method allows us to 331 identify one-to-one cell type concordance between animals in the same phylum, whereas between phyla, we observe interconnected cell types forming distinct families. These 332 333 findings support the notion that cell types evolve via hierarchical diversification (Arendt et 334 al., 2019), resulting in cell type families composed of evolutionarily related cell types

sharing a regulatory gene expression program that originated in their common ancestor.
One-to-one cell type homologies should exist only if no further cell type diversification has
occurred since the speciation.

338

339 In parallel, SAMap systematically identifies instances where paralogs exhibit greater 340 expression similarity than orthologs across species. Paralog substitution likely occurs due 341 to differential loss or retention of cell type-specific expression patterns of genes that were 342 duplicated in the common ancestor (Studer & Robinson-Rechavi, 2009). Alternatively, 343 paralog substitutions could arise due to compensating upregulation of paralogs following a loss-of-function mutation acquired by an ortholog (EI-Brolosy et al., 2019). While the 344 345 analysis presented here focuses on comparisons between two species, incorporating 346 multiple species into a single analysis that also accounts for their phylogenetic 347 relatedness could enable determining the specific order of paralog substitutions, 348 associated cell type diversification events, and the mechanism by which they arose. 349 However, this would require cell atlases that consistently sample key branching points 350 along the tree of life. Nevertheless, identifying lineage-specific paralog substitution 351 signatures should be accessible in extensively studied vertebrate single-cell atlases, as 352 the vertebrate clade is where existing data and knowledge are most concentrated.

353

Besides applications in evolutionary biology, we anticipate SAMap can catalyze the annotation of new cell atlases from non-model organisms, which often represents a substantial bottleneck requiring extensive manual curation and prior knowledge. Its ability to use the existing atlases to inform the annotation of cell types in related species will

- 358 keep improving as more datasets become available to better sample the diversity of cell
- 359 types. Moreover, our approach allows leveraging existing and forthcoming single-cell
- 360 gene expression data to predict functionally similar gene homologs, which can serve as
- 361 guideposts for mechanistic molecular studies.
- 362

363 Materials and Methods

364 Data and Code Availability

- 365 The source code for SAMap is publicly available Github at • (https://github.com/atarashansky/SAMap), along with the code to perform the 366 367 analysis and generate the figures.
- The datasets analyzed in this study are detailed in Supplementary Table 1 with
 their accessions, and annotations provided.

370

371 The SAMap Algorithm

372

373 The SAMap algorithm contains three major steps: preprocessing, mutual nearest 374 neighborhood alignment, and gene-gene correlation initialization. The latter two are 375 repeated for three iterations, by default, to balance alignment performance and 376 computational runtime. SAMap runs up to one hour on an average desktop computer for 377 200,000 total cells.

378

379 1. Preprocessing.

380 1.1. Generate gene homology graph via reciprocal BLAST.

We first construct a gene-gene bipartite graph between two species by performing reciprocal BLAST of their respective transcriptomes using *tblastx*, or proteomes using *blastp. tblastn* and *blastx* are used for BLAST between proteome and transcriptome. When a pair of genes share multiple High Scoring Pairs (HSPs), which are local regions

of matching sequences, we use the HSP with the highest bit score to measure homology.
Only pairs with E-value < 10⁻⁶ are included in the graph.

387

Here we define similarity using BLAST, though SAMap is compatible with other protein homology detection methods (e.g. HMMER (Eddy, 2008)) or orthology inference tools (e.g. OrthoClust (Yan et al., 2014) and Eggnog (Huerta-Cepas et al., 2019)). While each of these methods has known strengths and limitations, BLAST is chosen for its broad usage, technical convenience, and compatibility with low-quality transcriptomes.

393

We encode the BLAST results into two triangular adjacency matrices, A and B, each 394 395 containing bit scores in one BLAST direction. We combine A and B to form a gene-gene adjacency matrix G. After symmetrizing G, we remove edges that only appear in one 396 direction: $G = Recip(\frac{1}{2}[(A+B) + (A+B)^T]) \in \Re^{m_1+m_2 \times m_1+m_2}$, where Recip only keeps 397 398 reciprocal edges, and m_1 and m_2 are the number of genes of the two species, respectively. To filter out relatively weak homologies, we also remove edges where G_{ab} < 399 $0.25 max(G_{ab})$. Edge weights are then normalized by the maximum edge weight for each 400 gene and transformed by a hyperbolic tangent function to increase discriminatory power 401 between low and high edge weights, $\hat{G}_{ab} = 0.5 + 0.5 \tanh(10G_{ab}/max_b^{ab}(G_{ab}) - 5)$. 402

403

404 1.2. Construct manifolds for each cell atlas separately using the SAM algorithm.

The scRNAseq datasets are normalized such that each cell has a total number of raw counts equal to the median size of single-cell libraries. Gene expressions are then lognormalized with the addition of a pseudocount of 1. Genes expressed (i.e., $log_2(D + 1) >$ 408 1) in greater than 96% of cells are filtered out. SAM is run using the following parameters: 409 *preprocessing* = '*StandardScaler*', *weight_PCs* = *False*, *k* = 20, and *npcs* = 150. A detailed 410 description of parameters is provided previously (Tarashansky et al., 2019). SAM outputs 411 N_1 and N_2 , which are directed adjacency matrices that encode *k*-nearest neighbor graphs 412 for the two datasets, respectively.

413

SAM only includes the top 3,000 genes ranked by SAM weights and the first 150 principal components (PCs) in the default mode to reduce computational complexity. However, downstream mapping requires PC loadings for all genes. Thus, in the final iteration of SAM, we run PCA on all genes and take the top 300 PCs. This step generates a loading matrix for each species $i, L_i \in \Re^{300 \times m_i}$.

419

420 2. Mutual nearest neighborhood alignment.

421 2.1. Transform feature spaces between species.

For the gene expression matrices $Z_i \in \Re^{n_i \times m_i}$, where *n* and *m* are the number of cells 422 423 and genes respectively, we first zero the expression of genes that do not have an edge 424 in \hat{G} and standardize the expression matrices such that each gene has zero mean and unit variance, yielding \tilde{Z}_i . \hat{G} represents a bipartite graph in the form of \hat{G} = 425 $\begin{bmatrix} 0_{m_1,m_1} & H \in \Re^{m_1 \times m_2} \\ H^T \in \Re^{m_2 \times m_1} & 0_{m_2,m_2} \end{bmatrix}, \text{ where } 0_{m,m} \text{ is } m \times m \text{ zero matrix and } H \text{ is the biadjacency}$ 426 matrix. Letting $H_1 = H$ and $H_2 = H^T$ encoding directed edges from species 1 to 2 and 2 427 to 1, respectively, we normalize the biadjacency matrix H_i such that each row sums to 1: 428 $\widehat{H}_i = SumNorm(H_i) \in \Re^{m_i \times m_j}$, where the *SumNorm* function normalizes the rows to sum 429

430 to 1. The feature spaces can be transformed between the two species via weighted 431 averaging of gene expression, $\tilde{Z}_{ij} = \tilde{Z}_i \hat{H}_i$.

432

433 2.2. Project single-cell gene expressions into a joint PC space.

We project the expression data from two species into a joint PC space (Barkas et al., 2019), $P_i = \tilde{Z}_i L_i^T$ and $P_{ij} = \tilde{Z}_{ij} L_j^T$. We then horizontally concatenate the principal components P_i and P_{ij} to form $\hat{P}_i \in \Re^{n_i \times 600}$.

437

438 2.3. Calculate *k*-nearest cross-species neighbors for all cells.

Using the joint PCs, \hat{P}_i , we identify for each cell the *k*-nearest neighbors in the other dataset using cosine similarity (k = 20 by default). Neighbors are identified using the *hnswlib* library, a fast approximate nearest-neighbor search algorithm (Malkov & Yashunin, 2020). This outputs two directed biadjacency matrices $C_i \in \Re^{n_i \times n_j}$ for (i, j) =(1,2) or (2,1) with edge weights equal to the cosine similarity between the PCs.

444

445 2.4. Apply the graph-coarsening mapping kernel to identify cross-species mutual nearest
446 neighborhoods.

To increase the stringency and confidence of mapping, we only rely on cells that are *mutual* nearest cross-species neighbors, which are typically defined as two cells reciprocally connected to one another (Haghverdi et al., 2018). However, due to the noise in cell-cell correlations and stochasticity in the kNN algorithms, cross-species neighbors are often randomly assigned from a pool of cells that appear equally similar, decreasing the likelihood of mutual connectivity between individual cells even if they have similar

expression profiles. To overcome this limitation, we integrate information from each cell's
local neighborhood to establish more robust mutual connectivity between cells across
species. Two cells are thus defined as mutual nearest cross-species neighbors when their
respective neighborhoods have mutual connectivity.

457

Specifically, the nearest neighbor graphs N_i calculated in step 1.2 are used to calculate 458 the neighbors of cells t_i hops away along outgoing edges: $\bar{N}_i = N_i^{t_i}$, where \bar{N}_i are 459 adjacency matrices that contain the number of paths connecting two cells t_i hops away, 460 461 for i = 1 or 2. t_i determines the length-scale over which we integrate incoming edges for 462 species *i*. Its default value is 2 if the dataset size is less than 20,000 cells and 3 otherwise. 463 However, cells within tight clusters may have spurious edges connecting to other parts of 464 the manifold only a few hops away. To avoid integrating neighborhood information outside 465 this local structure, we use the Leiden algorithm (Traag et al., 2019) to cluster the graph 466 and identify a local neighborhood size for each cell (the resolution parameter is set to 3 by default). If cell *a* belongs to cluster c_a , then its neighborhood size is $l_a = |c_a|$. For each 467 row a in N_i we only keep the l_a geodesically closest cells, letting the pruned graph be 468 denoted as \hat{N}_i . 469

470

Edges outgoing from cell a_i in species *i* are encoded in the corresponding row in the adjacency matrix: C_{i,a_i} . We compute the fraction of the outgoing edges from each cell that target the local neighborhood of a cell in the other species: $\tilde{C}_{i,a_ib_j} = \sum_{c \in X_{j,b_j}} C_{i,a_ic}$, where X_{j,b_i} is the set of cells in the neighborhood of cell b_i in species *j* and \tilde{C}_{i,a_ib_i} is the fraction

of outgoing edges from cell a_i in species *i* targeting the neighborhood of cell b_j in species *j*.

477

To reduce the density of \tilde{C}_i so as to satisfy computational memory constraints, we remove edges with weight less than 0.1. Finally, we apply the mutual nearest neighborhood criterion by taking the element-wise, geometric mean of the two directed bipartite graphs: $\tilde{C} = \sqrt{\tilde{C}_1 \circ \tilde{C}_2}$. This operation ensures that only bidirectional edges are preserved, as small edge weights in either direction results in small geometric means.

483

484 2.5. Assign the k-nearest cross-species neighborhoods for each cell and update edge
485 weights in the gene homology graph.

Given the mutual nearest neighborhoods $\tilde{C} \in \Re^{n_1 \times n_2}$, we select the *k* nearest neighborhoods for each cell in both directions to update the directed biadjacency matrices

488 C_1 and C_2 : $C_1 = KNN(\tilde{C}, k)$ and $C_2 = KNN(\tilde{C}^T, k)$, with k = 20 by default.

489

490 2.6. Stitch the manifolds.

We use C_1 and C_2 to combine the manifolds N_1 and N_2 into a unified graph. We first weight the edges in N_1 and N_2 to account for the number of shared cross-species neighbors by computing the one-mode projections of C_1 and C_2 . In addition, for cells with strong crossspecies alignment, we attenuate the weight of their within-species edges. For cells with little to no cross-species alignment, their within-species are kept the same to ensure that the local topological information around cells with no alignment is preserved.

Specifically, we use N_1 and N_2 to mask the edges in the one-mode projections, $\widetilde{N}_1 =$ 498 $U(N_1) \circ (Norm(C_1)Norm(C_2))$ and $\widetilde{N}_2 = U(N_2) \circ (Norm(C_2)Norm(C_1))$, where U(E) sets 499 500 all edge weights in graph E to 1 and Norm normalizes the outgoing edges from each cell 501 to sum to 1. The minimum edge weight is set to be 0.3 to ensure that neighbors in the original manifolds with no shared cross-species neighbors still retain connectivity: $\tilde{N}_{1,ij}$ = 502 $min(0.3, \tilde{N}_{1,ij})$ and $\tilde{N}_{2,ij} = min(0.3, \tilde{N}_{2,ij})$ for all edges (i, j). We then scale the within-503 species edges from cell *i* by the total weight of its cross-species edges: $\widetilde{N}_{1,i} = (1 - 1)^{-1}$ 504 $\frac{1}{k}\sum_{j=1}^{n_2}C_{1,ij}\widetilde{N}_{1,i}$ and $\widetilde{N}_{2,i} = (1 - \frac{1}{k}\sum_{j=1}^{n_1}C_{2,ij})\widetilde{N}_{2,i}$. Finally, the within- and cross-species 505 graphs are stitched together to form the combined nearest neighbor graph N: $N = [\widetilde{N}_1 \bigoplus$ 506 $C_1] \oplus [C_2 \oplus \widetilde{N}_2]$. The overall alignment score between species 1 and 2 is defined as S =507 $\frac{1}{n_1+n_2} (\sum_{i=1}^{n_1} \sum_{j=1}^{n_2} C_{1,ij} + \sum_{i=1}^{n_2} \sum_{j=1}^{n_1} C_{2,ij}).$ 508

509

510 *3. Gene-gene correlation initialization.*

511 3.1. Update edge weights in the gene-gene bipartite graph with expression correlations. 512 To compute correlations between gene pairs, we first transfer expressions from one species to the other: $\bar{Z}_{i,n_im_j} = C_{i,n_i}Z_{j,m_j}$, where \bar{Z}_{i,n_im_j} is the imputed expressions of gene 513 514 m_j from species j for cell n_i in species i, and C_{i,n_i} is row n_i of the biadjacency matrix 515 encoding the cross-species neighbors of cell n_i in species *i*, all for (i, j) = (1, 2) and (2, 1). 516 We similarly use the manifolds constructed by SAM to smooth the within-species gene expressions using kNN averaging: $\overline{Z}_{j,m_i} = N_{j,m_i}Z_{j,m_i}$, where N_j is the nearest-neighbor 517 518 graph for species *j*. We then concatenate the within- and cross-species gene expressions such that the expression of gene m_j from species *j* in both species is $\bar{Z}_{m_j} = \bar{Z}_{i,m_j} \oplus \bar{Z}_{j,m_j}$. 519

520

For all gene pairs in the unpruned homology graph generated in step 1.1., \hat{G} , we compute their correlations, $\hat{G}_{ab} := \theta(0)Corr(\bar{Z}_a, \bar{Z}_b)$, where $\theta(0)$ is a Heaviside step function centered at 0 to set negative correlations to zero. We then use the expression correlations to update the corresponding edge weights in \hat{G} , which are again normalized through $\hat{G}_{ab} = 0.5 + 0.5 \tanh(10\hat{G}_{ab}/max(\hat{G}_{ab}) - 5)$.

526

527 Annotation of cell atlases

528 To annotate the primary zebrafish and *Xenopus* cell types, the cell subtype annotations 529 provided by the original publications (Briggs et al., 2018; Wagner et al., 2018) are 530 coarsened using a combination of the manual matching and developmental hierarchies. 531 For example, as "heart - mature", "heart - hoxd9a", "heart", and "heart field" in zebrafish 532 are all manually matched to "cardiac mesoderm" in Xenopus, we label these cells as 533 "heart". In cases where the matching is insufficient to coarsen the annotations, we use 534 the provided developmental trees to name a group of terminal cell subtypes by their 535 common ontogenic ancestor. The annotations provided by their respective studies were 536 used to label the cells in the Spongilla, Hydra, and mouse atlases. To annotate the 537 schistosome cells, we used known marker genes to annotate the main schistosome tissue 538 types (Li et al., 2020). Annotations for all single cells in all datasets are provided in 539 Supplementary Table 1.

540

541 Visualization

542 The combined manifold *N* is embedded into 2D projections using UMAP implemented in

543 the scanpy package (Wolf et al., 2018) by scanpy.tl.umap with the parameter min dist = 0.1. The sankeyD3 package (https://rdrr.io/github/fbreitwieser/sankeyD3/man/sankeyD3-544 545 package.html) in R is used to generate the sankey plots. Edge thickness corresponds to 546 the alignment score between mapped cell types. The alignment score between cell types a and b is defined as $s_{ab} = \frac{1}{|c_a| + |c_b|} (\sum_{i \in c_a} \sum_{j \in c_b} C_{1,ij} + \sum_{i \in c_b} \sum_{j \in c_a} C_{2,ij})$, where c_a and c_b 547 are the set of cells in cell types a and b, respectively. Cell type pairs with alignment score 548 549 less than z are filtered out. By default, z is set to be 0.1. Cell types that did not cluster 550 properly in their respective manifolds were omitted from the sankey plot. In the zebrafish-Xenopus comparison, we excluded heart, germline, and olfactory placode cells from both 551 552 species because they did not cluster in the Xenopus atlas. Similarly, the iridoblast, 553 epiphysis, *nanoq*+, apoptotic-like, and forerunner cells were excluded because they did 554 not cluster in the zebrafish atlas.

555

556 The network graphs in **Figure 4D** are generated using the *networkx* package 557 (https://networkx.github.io) in python. To focus on densely connected cell type groups, 558 we filter out cell type pairs with alignment score less than 0.05.

559

560 Identification of gene pairs that drive cell type mappings

561 We define g_1 and g_2 to contain SAMap-linked genes from species 1 and 2, respectively. 562 Note that a gene may appear multiple times as SAMap allows for one-to-many homology. 563 Let $X_{a_1b_2}$ denote the set of all cells with cross species edges between cell types a_1 and 564 b_2 . We calculate the average standardized expression of all cells from species *i* that are 565 in $X_{a_1b_2}$: $Y_{i,g_i} = \frac{1}{|\{x,x\in X_{a_1b_2}\}|} \sum_{x\in X_{a_1b_2}} \tilde{Z}_{i,x,g_i}$, where $\tilde{Z}_{i,x,g_i} \in \Re^{|g_i|}$ is the standardized

expression of genes g_i in cell x. The correlation between Y_{1,g_1} and Y_{2,g_2} can be written as 566 $Corr(Y_{1,g_1}, Y_{2,g_2}) = \sum_{j=1}^{|g_1|} S(Y_{1,g_1})_j \circ S(Y_{2,g_2})_j$, where S(Z) standardizes vector Z to have 567 568 zero mean and unit variance. We use the summand to identify gene pairs that contribute most positively to the correlation. We assign each gene pair a score: $h_g = T(S(Y_{1,g_1})) \circ$ 569 $T(S(Y_{2,q_2}))$, where T(Z) sets negative values in vector Z to zero in order to ignore lowly-570 571 expressed genes. To be inclusive, we begin with the top 1,000 gene pairs according to h_q and filter out gene pairs in which one or both of the genes are not differentially 572 573 expressed in their respective cell types (p-value > 10^{-2}), have less than 0.2 SAM weight, 574 or are expressed in fewer than 5% of the cells in the cluster. The differential expression 575 of each gene in each cell type is calculated using the Wilcoxon rank-sum test 576 implemented in the scanpy function scanpy.tl.rank genes groups.

577

578 Orthology group assignment

We used the Eggnog mapper (v5.0) (Huerta-Cepas et al., 2019) to assign each gene to 579 580 an orthology group with default parameters. For the zebrafish-to-Xenopus mapping, 581 genes are considered paralogs if they map to the same eukaryotic orthology group and 582 orthologs if they map to the same vertebrate orthology group. For the pan-species 583 analysis, we group genes from all species with overlapping orthology assignments. In 584 Figure 5B, each column corresponds to one of these groups. As each group may contain 585 multiple genes from each species, we present the expression of the gene with the highest 586 enrichment score per species. All gene names and corresponding orthology groups are 587 reported in Supplementary Table 4.

588

589 Phylogenetic reconstruction of gene trees

We generate gene trees to validate the identity of genes involved in putative examples of paralog substitution and of *Fox* and *Csrp* transcriptional regulators that are identified as enriched in contractile cells. For this, we first gather protein sequences from potential homologs using the eggnog version 5.0 orthology database (Huerta-Cepas et al., 2019). For the *Fox* and *Csrp* phylogenies, we include all Fox clade I (Larroux et al., 2008) and Csrp/Crip homologs, respectively, from the seven species included in our study.

596

597 Alignment of protein sequences is performed with Clustal Omega version 1.2.4 using 598 default settings as implemented on the EMBL EBI web services platform (Madeira et al., 599 2019). Maximum likelihood tree reconstruction is performed using IQ-TREE version 600 1.6.12 (Nguyen et al., 2015) with the ModelFinder Plus option (Kalyaanamoorthy et al., 601 2017). For the *Csrp* tree, we perform 1,000 nonparametric bootstrap replicates to assess 602 node support. For Fox, we utilize the ultrafast bootstrap support option with 1,000 603 replicates. For each gene tree we choose the model that minimizes the Bayesian 604 Information Criterion (BIC) score in ModelFinder. This results in selection of the following 605 models: DCMut+R4 (Csrp) and VT+F+R5 (Fox). The final consensus trees are visualized 606 and rendered using the ete3 v3.1.1 python toolkit (Huerta-Cepas et al., 2016) and the 607 Interactive Tree of Life v4 (Letunic & Bork, 2019).

608

609 KOG functional annotation and enrichment analysis

610 Using the eggnog mapper, KOG functional annotations are transferred to individual 611 transcripts from their assigned orthology group. For enrichment analysis, all genes

612 enriched in the set of cell type pairs of interest are lumped to form the target set for each 613 species. For example, the target set for Spongilla archaeocytes used in Figure 5C is 614 composed of all genes enriched between Spongilla archaeocytes and other invertebrate 615 stem cells. Note that this set includes genes from other species that are linked by SAMap 616 to the Spongilla archeocyte genes. We include genes from other species in the target set 617 to account for differences in KOG functional annotation coverage between species. As 618 such, the annotated transcripts from all 7 species are combined to form the background 619 set. We used a hypergeometric statistical test (Eden et al., 2009) to measure the 620 enrichment of the KOG terms in the target genes compared to the background genes.

621

622 Mapping zebrafish and xenopus atlases using existing methods

623 For benchmarking, we used vertebrate orthologs as determined by Eggnog as input to 624 Harmony (Korsunsky et al., 2019), Liger (Welch et al., 2019), Seurat (Stuart et al., 2019), 625 Scanorama (Hie et al., 2019), BBKNN (Polański et al., 2019), which are all run with default 626 parameters. One-to-one orthologs were selected from one-to-many and many-to-many 627 orthologs by using the bipartite maximum weight matching algorithm implemented in 628 networkx. When using the one-to-one orthologs as input for SAMap, we ran for only one 629 iteration. The resulting integrated lower-dimensional coordinates (PCs for Seurat, 630 Harmony, and Scanorama and non-negative matrix factorization coordinates for Liger) 631 and stitched graph (BBKNN and SAMap) were all projected into 2D with UMAP (Figure 632 **2 – figure supplement 2A**). The integrated coordinates are used to generate a nearest 633 neighbor graph using the correlation distance metric, which is then used to compute the

alignment scores in Figure 2 – figure supplement 2B. The alignment scores for SAMap
and BBKNN are directly computed from their combined graphs.

636

637 In-situ hybridization in schistosomes

638 S. mansoni (strain: NMRI) juveniles are retrieved from infected female Swiss Webster 639 mice (NR-21963) at ~3 weeks post-infection by hepatic portal vein perfusion using 37°C DMEM supplemented with 5 % heat inactivated FBS. The infected mice are provided by 640 the NIAID Schistosomiasis Resource Center for distribution through BEI Resources, NIH-641 642 NIAID Contract HHSN272201000005I. In adherence to the Animal Welfare Act and the 643 Public Health Service Policy on Humane Care and Use of Laboratory Animals, all 644 experiments with and care of mice are performed in accordance with protocols approved 645 by the Institutional Animal Care and Use Committees (IACUC) of Stanford University 646 (protocol approval number 30366). In situ hybridization experiments are performed as described previously (Tarashansky et al., 2019), using riboprobes synthesized from gene 647 648 fragments cloned with the listed (Smp 170340): primers: collagen 649 GGTGAAGAAGGCTGTTGTGG, ACGATCCCCTTTCACTCCTG; tropomyosin 650 (Smp 031770): AAGCTGAAGTCGCCTCACTA, CATATGCCTCTTCACGCTGG; 651 troponin (Smp 018250): CGTAAACCTGGTCAGAAGCG, 652 ATCCTTTTCCTCCAGAGCGT; myosin regulatory light chain (Smp 132670): 653 GAGACAGCGAGTAGTGGAGG, TGCCTTCTTTGATTGGAGCT; wnt (Smp 156540): TGTGGTGATGAAGATGGCAG, 654 CCACGGCCACAACACATATT; frizzled 655 (Smp 174350): CGAACAGGCGCATGACAATA, TGCTAGTCCTGTTGTCGTGT.

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868 Figures





875 fractions of transcriptomes lacking homology, which limits the amount of information 876 comparable across species. (C) SAMap workflow. Homologous gene pairs initially weighted by protein sequence similarity are used to align the manifolds, low dimensional 877 878 representations of the cell atlases. Gene-gene correlations calculated from the aligned 879 manifolds are used to update the edge weights in the bipartite graph, which are then used 880 to improve manifold alignment. (D) Mutual nearest neighborhoods improve the detection 881 of cross-species mutual nearest neighbors by connecting cells that target one other's 882 within-species neighborhoods. (E) Convergence of SAMap is evaluated by the root mean 883 square error (RMSE) of the alignment scores between mapped clusters in adjacent 884 iterations for all 21 pairwise comparisons of the 7 species.



886

Figure 2: SAMap successfully maps *D. rerio* and *X. tropicalis* atlases and reveals prevalent paralog substitutions. (A) UMAP projection of the combined zebrafish (yellow) and *Xenopus* (blue) manifolds, with example cell types circled. (B) Sankey plot summarizing the cell type mappings. Edges with alignment score < 0.1 are omitted. Edges that connect developmentally distinct secretory cell types are highlighted in black. (C)

892 Heatmaps of alignment scores between developmental time points for ionocyte, 893 forebrain/midbrain, placodal, and neural crest lineages. X-axis: zebrafish. Y-axis: 894 Xenopus. (D) SAMap alignment scores compared to those of benchmarking methods 895 using one-to-one vertebrate orthologs as input. Each dot represents a cell type pair 896 supported by ontogeny annotations. (E) Expression of orthologous (left) and paralogous 897 (right) gene pairs overlaid on the combined UMAP projection. Expressing cells are color-898 coded by species, with those that are connected across species colored cyan. Cells with 899 no expression are shown in gray. More examples are provided in Figure 2 - figure 900 supplement 3.



902

903 Figure 3: SAMap transfers cell type information from a well-annotated organism 904 (planarian S. mediterranea) to its less-studied cousin (schistosome S. mansoni) 905 and identifies parallel stem cell compartments. (A) UMAP projection of the combined 906 manifolds. Tissue type annotations are adopted from the S. mediterranea atlas (Fincher 907 et al., 2018). The schistosome atlas was collected from juvenile worms, which we found 908 to contain neoblasts with an abundance comparable to that of planarian neoblasts (Li et 909 al., 2020). (B) Overlapping expressions of selected tissue-specific TFs with expressing 910 cell types circled. (C) UMAP projection of the aligned manifolds showing planarian and 911 schistosome neoblasts, with homologous subpopulations circled. Planarian neoblast data

is from (Zeng et al., 2018), and cNeoblasts correspond to the Nb2 population, which are
pluripotent cells that can rescue neoblast-depleted planarians in transplantation
experiments. (D) Distributions of conserved TF expressions in each neoblast
subpopulation. Expression values are *k*-nearest-neighbor averaged and standardized,
with negative values set to zero. Blue: planarian; yellow: schistosome.



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Figure 4: Mapping evolutionarily distant species identifies densely connected cell
type groups. (A) Violin plots showing the number of enriched gene pairs in cell type
mappings from all 21 pairwise mappings between the 7 species. 87% of cell type
mappings have greater than 40 enriched gene pairs (dotted line). Species acronyms are

923 the same as in Figure 1A. (B) Schematic illustrating edge (left) and node (right) 924 transitivities, defined as the fraction of triads (set of three connected nodes) in closed 925 triangles. (C) The percentage of cell type pairs that are topologically equivalent to the 926 green edge in each illustrated motif. (D) Network graphs showing highly connected cell 927 type families. Each node represents a cell type, color-coded by species (detailed 928 annotations are provided in **Supplementary Table 5**). Mapped cell types are connected 929 with an edge. (E) Boxplot showing the median and interguartile ranges of node 930 transitivities for highly connected cell type groups. For all box plots, the whiskers denote 931 the maximum and minimum observations. The average node transitivity per group is 932 compared to a bootstrapped null transitivity distribution, generated by repeatedly 933 sampling subsets of nodes in the cell type graph and calculating their transitivities. p < p $5x10^{-3}$, ** p < $5x10^{-5}$, ***p < $5x10^{-7}$. (F) Boxplot showing the median and interguartile 934 935 ranges of the number of enriched gene pairs in highly connected cell type groups. All cell type connections in these groups have at least 40 enriched gene pairs (dotted line). 936



Figure 5: SAMap identifies muscle and stem cell transcriptional signatures 938 939 conserved across species. (A) Enrichment of KOG functional annotations calculated 940 for genes shared in contractile cell types. For each species, genes enriched in individual contractile cell types are combined. (B) Expression and enrichment of conserved muscle 941 942 genes in contractile cell types. Color: mean standardized expression. Symbol size: the 943 fraction of cells each gene is expressed in per cell type. Homologs are grouped based on 944 overlapping eukaryotic Eggnog orthology groups. If multiple genes from a species are 945 contained within an orthology group, the gene with highest standardized expression is 946 shown. Genes in blue: core transcriptional program of bilaterian muscles; red: 947 transcription factors conserved throughout Metazoa. (C) Enrichment of KOG functional 948 annotations for genes shared by stem cell types. (D) Boxplot showing the median and

interquartile ranges of the mean standardized expressions of genes in hydra and
planarian stem cells/progenitors that are conserved across all invertebrate species in this
study. Planarian progenitors: *piwi*+ cells that cluster with differentiated tissues in Fincher
et al. (Fincher et al., 2018). Neoblasts: cluster 0 in Fincher et al. (Fincher et al., 2018) that
does not express any tissue-specific markers.



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Figure 2 – figure supplement 1: Expression of selected genes enriched in *D. rerio*and *X. tropicalis* secretory cell types. Expressions of orthologous gene pairs linked by
SAMap are overlaid on the combined UMAP projection. Expressing cells are color-coded
by species, with those connected across species colored cyan. Cells with no expression
are shown in gray. The mapped secretory cell types are highlighted with circles.



Figure 2 – figure supplement 2: Existing methods failed to map *D. rerio* and *X. tropicalis* atlases. (A) UMAP projections of the integration results from SAMap using the

965 full homology graph, compared to Liger, BBKNN, Scanorama, Seurat, Harmony, and 966 SAMap using 1-1 orthologs. For fair comparisons, all methods were run on the *D. rerio* 967 and *X. torpicalis* atlases subsampled to approximately 15,000 cells to satisfy 968 computational constraints of Seurat and Liger. (B) Distribution of alignment scores 969 between individual cells.



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Figure 2 – figure supplement 3: Representative examples of paralog substitution
events in *D. rerio* and *X. tropicalis* atlases. Expressions of orthologous and paralogous
gene pairs are overlaid on the combined UMAP projection. Expressing cells are colorcoded by species, with those that are connected across species colored in cyan. Cells
with no expression are shown in gray.



978

979 Figure 3 – figure supplement 1: SAMap-linked gene pairs that are enriched in cell 980 type pairs between S. mediterranea and S. mansoni. (A) Rows: linked cell types. 981 Schistosome cell types correspond to leiden clusters. Columns: genes linked by SAMap 982 with overlapping eukaryotic Eggnog orthology groups. We calculate the average 983 standardized expression of each gene in an orthology group for its corresponding cell 984 type in a particular pair and report the highest expression. A selected set of orthology 985 groups corresponding to transcriptional regulators are labeled. (B) Fluorescence in situ 986 hybridization shows the co-expression of wnt (Smp 156540) and a panel of muscle 987 markers (collagen, troponin, myosin and tropomyosin) in S. mansoni juveniles. The body 988 wall muscles are expected to be located close to the parasite surface (dashed outline).

989 The images are maximum intensity projections constructed from ~10 confocal slices with 990 optimal axial spacing recommended by the Zen software collected on a Zeiss LSM 800 991 confocal microscope using a 40× (N.A. = 1.1, working distance = 0.62 mm) water-992 immersion objective (LD C-Apochromat Corr M27). (C) Whole mount in situ hybridization 993 images showing that the expression of wnt and frizzled (Smp 174350) are concentrated 994 in the parasite tail (arrows) with decreasing gradients extending anteriorly. In planarian 995 muscles, Wnt genes provide the positional cues for setting up the body plan during 996 regeneration (Scimone et al., 2017; Reddien, 2018). The presence of an anterior-997 posterior expression gradient of wnt and frizzled in muscles of schistosome juveniles 998 suggests that they may have similar functional roles in patterning during development.



1000

Figure 3 – figure supplement 2: Schistosome neoblasts express canonical muscle markers in muscle progenitors. UMAP projections of schistosome neoblasts with gene expressions overlaid. μ and μ ' cells are circled. Colormap: expression in units of $log_2(D +$ 1). For visualization, expression was smoothed via nearest-neighbor averaging using SAM. Note that *myod1* and *cabp* are expressed in both presumptive muscle progenitor populations, whereas all other markers are enriched in μ ' cells. All genes displayed are also expressed in fully differentiated muscle tissues.





- 1013 correlation coefficient reported at the top. Alignment scores and edge transitivity for
- 1014 individual species pairs are shown in the remaining subplots.



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Figure 4 – figure supplement 2: Number of enriched gene pairs are mostly independent of edge transitivity. Top left: The edge transitivity is plotted against the number of enriched gene pairs for all cell type pairs in the connectivity graph. Dotted line: the linear best fit, with the Pearson correlation coefficient reported at the top. Top right: magnified view of the mapped cell type pairs supported by small numbers of gene pairs

- 1022 (<40) to show those edges have low transitivity scores (<0.4). The sublots below show
- 1023 the number of enriched gene pairs and edge transitivity for individual species pairs.



1026 Figure 5 – figure supplement 1: Phylogenetic reconstruction of animal contractile 1027 cell transcriptional regulators. Trees depict Csrp/Crip (A) and Fox group I (B) gene 1028 families. Genes labelled red are enriched in at least one contractile gene pair identified 1029 via SAMap. Support values indicate bootstrap support from 1,000 nonparametric (Csrp) 1030 or ultrafast (Fox) bootstrap replicates. Besides these two transcriptional regulators, 1031 contractile cells in all seven species were found to be also enriched for transcription 1032 factors from the C2H2 Zinc Finger, Lim Homeobox, and Paired Homeobox families, 1033 although in different cell types we found enrichment of a number of distinct orthologs. 1034 Whether this reflects an ancestral role for these transcription factor families in regulating 1035 contractility or their independent evolution will require additional taxonomic sampling and 1036 broader coverage of muscle cell diversity to resolve.

1037 Supplementary table captions

1038

1039 Supplementary Table 1: Cell atlas metadata and cell annotations. Metadata includes 1040 the number of cells, number of transcripts in the transcriptome, median number of 1041 transcripts detected per cell, the reference transcriptome used in this study, database 1042 through which the transcriptomes are provided, technology used for constructing the cell 1043 atlases, atlas data accessions, processing notes, and references. Leiden clusters and 1044 cell type annotations are reported for cells in each atlas. The Zebrafish and Xenopus 1045 tables include both the original cell type annotations and those used in this study. D. rerio. 1046 X. tropicalis, and mouse annotations include developmental stages.

1047

Supplementary Table 2: Cell type annotations for the zebrafish-*Xenopus* mapping.
Correspondence between the cell type annotations provided in the original study (Briggs
et al., 2018; Wagner et al., 2018) and corresponding annotations used in this study is
provided for both *D. rerio* and *X. tropicalis* atlases.

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Supplementary Table 3: Identified paralogs with greater expression similarity than orthologs in the zebrafish-*Xenopus* mapping. Each row contains a pair of vertebrateorthologous genes and a corresponding pair of eukaryotic paralogs with higher correlation in expression compared to the orthologs, the expression correlations for ortholog and paralog pairs, the difference between their correlations, and whether the paralogs are considered as a paralog substitution (defined as when the substituted ortholog is either

absent or lowly-expressed with no cell-type specificity). Highlighted rows are shown in
Figure 2E and Figure 2 – figure supplement 3.

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Supplementary Table 4: Genes enriched in contractile cell types and invertebrate stem cells highlighted in Figure 4D. The IDs of the genes enriched in the contractile and invertebrate stem cell types are provided along with the IDs of the Eggnog orthology groups to which they belong. In cases where multiple genes from a species belonging to the same orthology group are enriched, the most differentially expressed gene is shown. The descriptions in the stem cell table are orthology annotations associated with the Spongilla genes provided in the original study (Musser et al., 2019).

1069

1070 Supplementary Table 5: Cell types in the cell type families shown in Figure 4D. For 1071 the schistosome cell types, we annotated two neural clusters, both of which express the 1072 neural marker complexin (Li et al., 2020). One of the clusters expresses the antigen 1073 SmKK7, so we label the clusters "Neural" and "Neural KK7", respectively. The "Muscle" 1074 population contains non-neoblast cells expressing troponin. The "Tegument prog" and 1075 "Teaument" populations consist of cells expressing tegument progenitor and 1076 differentiated marker genes, respectively, as reported in a previous study (Wendt et al., 2018). 1077