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# **OPEN** Experimentally induced metamorphosis in highly regenerative axolotl (ambystoma mexicanum) under constant diet restructures microbiota

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Axolotl (Ambystoma mexicanum) is a critically endangered salamander species and a model organism for regenerative and developmental biology. Despite life-long neoteny in nature and in captive-bred colonies, metamorphosis of these animals can be experimentally induced by administering Thyroid hormones (THs). However, microbiological consequences of this experimental procedure, such as host microbiota response, remain largely unknown. Here, we systematically compared host bacterial microbiota associated with skin, stomach, gut tissues and fecal samples, between neotenic and metamorphic axolotls based on 16S rRNA gene sequences. Our results show that distinct bacterial communities inhabit individual organs of axolotl and undergo substantial restructuring through metamorphosis. Skin microbiota among others, shifted sharply, as highlighted by a major transition from Firmicutes-enriched to Proteobacteria-enriched relative abundance and precipitously decreased diversity. Fecal microbiota of neotenic and metamorphic axolotl shared relatively higher similarity, suggesting that diet continues to shape microbiota despite fundamental transformations in the host digestive organs. We also reproduced the previous finding on reduction in regenerative capacity in limbs of axolotl following metamorphosis, highlighting the need to investigate whether shifts in microbiota is causally linked to regenerative capacity of axolotl. The initial results on axolotl microbiota provide novel insights into microbiological aspects of axolotl metamorphosis and will establish a baseline for future in-depth studies.

Metazoan genomes have diversified and evolved in the presence of associated host microbiota. The evolution of morphology and function of animal organ systems may have been influenced by interactions with their microbial partners<sup>1</sup>. From the host perspective, symbioses between metazoans and microbes provide a synergetic impact to operate essential functions for normal growth, development and behavior<sup>2-4</sup>. Studies on host-microbiome interactions in health and disease conditions indicate that perturbation of the crosstalk between the host and microorganisms may lead to deleterious consequences such as developmental defects<sup>3,5</sup>, increased susceptibility to infectious diseases<sup>6,7</sup> and ultimately fitness costs<sup>8</sup>. Even though host genotypes<sup>9</sup> and environmental factors, such as diet and habitat 10,11, were shown to strongly impact the composition and structure of animal microbiota, ecological forces shaping assembly of the host associated microbial communities have still been poorly understood.

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Amphibians, which undergo dramatic morphological changes through metamorphosis, exhibit explicitly altered biphasic life stages to tackle developmental challenges. Remarkably, metamorphosis in several marine animal species is mediated by bacterial community<sup>1,12</sup>. Thyroid hormones (THs) are key players in initiation and completion of metamorphosis<sup>13,14</sup>. Natural accumulation (as in anurans) or administration (as in axolotl) of THs leads to critical reorganization of organs in order to adapt terrestrial life conditions. This adaptive process includes reconstruction or loss of some existing organs and extremities, and formation of new ones<sup>15,16</sup>. A prominent example of reconstruction is observed in digestive tract of tadpole. In adult frogs acidic stomach and shorter intestine originate from non-acidic stomach and long intestine of tadpole digestive tract<sup>17,18</sup>. Growing evidence supports the notion that reshaping of organs and composition of microorganisms reciprocally influence each other as functions of bacterial communities are increasingly being linked to host metabolic activities<sup>19–21</sup>. Hence, unraveling microbiome compositions in various life stages may offer new insights into the life stage-specific microbial patterns. This notion led to gut microbiota profiling in tadpoles and frogs of the Northern leopard frog (*Lithobates pipiens*)<sup>22</sup> and later in amphibian larvae<sup>23</sup>. Despite this growing body of research, the implication of microbiota in diverse biological processes of amphibian hosts remains largely unknown.

Axolotl (*Ambystoma mexicanum*), a salamander species of amphibians, possess experimentally validated features, such as high regenerative capacity<sup>24</sup>, low cancer incidence<sup>25</sup>, scarless wound healing<sup>26</sup>, life-long neoteny with the ability to undergo induced metamorphosis<sup>27</sup>. These characteristics contributed to the recent establishment of axolotl as a promising vertebrate model organism for regenerative and developmental biology (reviewed in<sup>28</sup>). Reference resources for this model such as transcriptome<sup>29,30</sup>, proteome<sup>31,32</sup>, a draft genome assembly<sup>33</sup> and recently reported sequencing and assembly of axolotl genome data<sup>34</sup> are all publicly available. Current studies on regeneration in axolotl have focused on identification of genes, gene networks and pathways activated during limb and tail regeneration by utilizing transcriptome and proteome profiling tools<sup>29–32,35</sup>. However, there is very limited data on microbial diversity of salamanders<sup>36</sup> and the impact of salamander associated microbiota on biological processes such as immunity<sup>37</sup>. Furthermore, systematic investigation of microbial diversity in multiple organs of salamander species, including axolotl, has been missing in the literature.

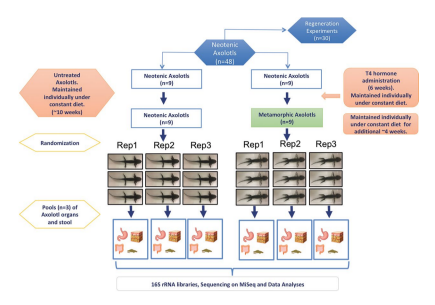
In this study, we hypothesized that induction of metamorphosis in axolotl leads to restructuring of bacterial microbiota due to reorganization of tissues via metamorphosis. To test the hypothesis, we explored the variation in neotenic and metamorphic axolotl's microbiota associated with skin, stomach, gut tissues, and fecal matter using 16S rRNA gene sequencing. Interestingly, limb regenerative capacity was previously shown to be visibly reduced in metamorphic axolotls compared to neotenic counterparts<sup>38</sup>. We therefore attempted to reproduce this finding by comparing limb regenerative capacity between two developmental stages and tested to find correlations with limb microbiota. Overall, the results provide novel insights on restructuring of the microbial communities *via* metamorphosis in this model organism.

#### Results

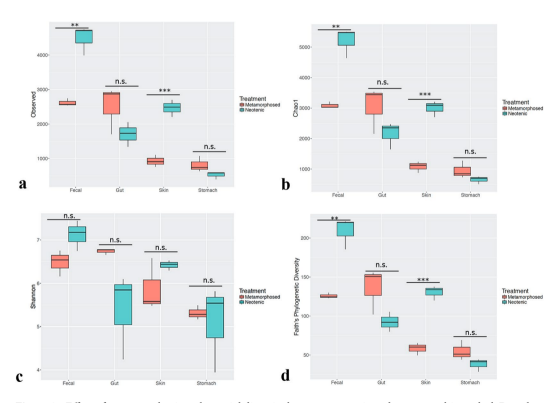
Induction of metamorphosis and comparative approach in axolotl. Details of the experimental design were described in the methods section (Fig. 1). Metamorphosis of axolotls was experimentally induced by administering Thyroid hormones (THs). Within 2–3 weeks of hormonal treatment of the animals, we observed weight loss, progressive disappearance of the fin and decrease in the gills size; and in approximately two months all animals showed characteristics of accomplished metamorphosis (Supplementary Fig. S1). In order to gain novel insights into microbiological aspects of metamorphosis, we first performed comparative analysis of microbiota between neotenic and metamorphic axolotl organs. We also collected water samples from the aquarium of axolotls to identify OTUs present in the water. Furthermore, to obtain additional insights into biological consequences of metamorphosis, we assessed gut tissue histology and limb regenerative capacity at both developmental changes.

Bacterial community structure and membership differ between neotenic and metamorphic axolotl. Sequencing of the V3-V4 region of the 16S rRNA gene produced approximately 3.7 M reads generated from 27 samples (24 samples from axolotls and 3 pools of samples from aquarium water, hereinafter "aqua"). The sequences were clustered into 14451 high quality, singleton-, chloroplast-, and mitochondria-removed and chimera-checked Operational Taxonomic Units (OTUs). Average number of amplicon sequences per sample was found as  $139059 \pm 49159$  sequences. Our data included 12224 (85% of total) *de novo* OTUs (OTU IDs that begin with "New.Reference" or "New.Cleanup.Reference"). We then classified a representative sequence of these OTUs using RDP classifier (v. 2.2) at 70% bootstrap cutoff. We identified 621 OTUs that did not find hits in the RDP database even at the phylum level ("Unclassified Bacteria"). We thus used MOLE-BLAST to determine their identity. Except a few high abundance OTUs (~5% abundance) enriched in the stomach samples hitting plant mitochondria (discarded), most of these OTUs had abundance below 0.1%, which can be considered a rare OTU<sup>39</sup>. The majority of these OTUs were phylogenetically related to the phylum *Proteobacteria* or *Verrucobacteria* (Supplementary Fig. S2; NCBI Accession numbers: MG518658 - MG519278).

Species richness and diversity were analyzed using a variety of alpha-diversity metrics across neotenic and metamorphic samples (Fig. 2a–d). Metamorphosis significantly reduced diversity in fecal and skin samples as follows; Chao1 and Observed OTUs were significantly lower in fecal and skin samples of the metamorphic samples as compared to neotenic samples (Unpaired Student t test, df = 4, p = 0.0018, p = 0.005, respectively). However, Inverse Simpson and Shannon indices were not significantly different (p = 0.34, p = 0.07) for these two particular samples although Faith's phylogenetic tree (PD) did indicate that microbiota diversity of these samples were significantly different from each other (p = 0.0022, p = 0.0005, respectively). Both Simpson and Shannon indices take into account richness and evenness in computing the metrics. Therefore, taxa with high relative abundance being heavily weighted in calculations while the indices are less sensitive to rare taxa when compared to richness only metrics<sup>40</sup>. Interestingly, evenness as calculated by Simpson-E index was not significantly different between

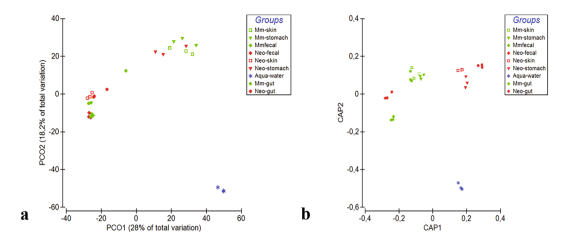


**Figure 1.** Experimental design for the comparison of neotenic and metamorphic axolotl. Of 48 siblings, a subset of 24 axolotls (9 axolotls for metamorphosis and 15 axolotls for regeneration experiments) were randomly selected and induced metamorphosis by T4 hormone administration while the rest kept untreated in neoteny. 30 animals (15 neotenic and 15 metamorphosed) were used in regeneration experiments and 18 animals (9 neotenic and 9 metamorphic) were housed individually for microbiome analysis. Each sample groups for skin, gut, stomach and fecal samples consisted of 3 replicates following randomization and pooling.

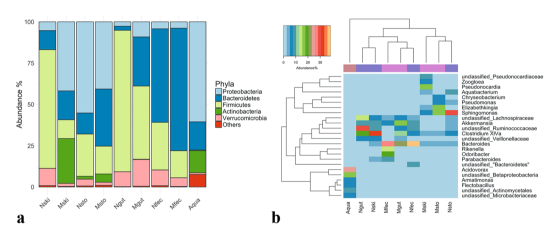


**Figure 2.** Effect of metamorphosis on bacterial diversity between neotenic and metamorphic axolotl. Box plots illustrate the comparison of diversity indices; Observed (a), Chao1 (b), Shannon (c) and Faith's Phylogenetic Diversity (PD) measures.

any sample pairs (p > 0.05 for all comparisons). Correspondingly, we inferred that low abundant taxa drive differences in diversity in fecal and skin samples. Both stomach and gut samples had richness and diversity that were not significantly different on all metrics between neotenic and metamorphic animals (p > 0.05; Fig. 2a-d).



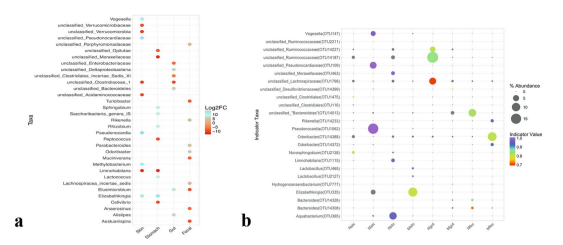
**Figure 3.** Beta diversity analysis based on Bray-Curtis distance matrix showing separation of neotenic and metamorphic bacterial communities. Samples were compared using PCO (a) and Canonical Analysis of Principal Coordinates (CAP) (b) methods.



**Figure 4.** Mean relative abundances of 16S rRNA sequences. Phylum level relative abundance as bar chart (a), genus level relative abundances shown as heatmap (Individual taxa displayed if the its abundance in any sample ≥5%). Samples and bacterial taxa were clustered using average linkage hierarchical clustering of a distance matrix based on Bray−Curtis distance and taxa abundances, respectively. Samples from each group were color coded on the column side bar as follows: aqua samples (brown); samples from neotenic axolotl (light slate blue), metamorphic axolotl (magenta) (b).

Beta diversity of bacterial communities largely differed between neotenic and metamorphic samples based on Bray-Curtis and Jaccard distance metrics (Fig. 3a and Supplementary Fig. S3, respectively). Within and between group differences (Main-effect) using both distance metric were statistically significant as per PERMANOVA test (Pseudo-F (8, 18) = 7.82,  $p(Monte\ Carlo) = 0.0001$ ). Permutational test for homogeneity of dispersions (PERMDISP) was at the border of significance; (F (8, 18) = 8.357, p(perm) = 0.0507, 9999 permutations of residuals), indicating that the average within group dispersions were marginally equivalent among groups but dispersion effect, to some degree, may be confounded in the location effect. We next employed Canonical analysis of principal coordinates<sup>41</sup> based on Bray-Curtis distance matrix, a constrained ordination that maximizes the differences among a priori groups and reveals subtle patterns, which otherwise remain elusive to unconstrained ordinations. CAP analysis clearly separated neotenic and metamorphic samples (Fig. 3b) except for the fecal samples (Correct classification rate 96.3%; trace statistics (tr(Q\_m'HQ\_m): 3,92174 p = 0.001 with 9999 permutations, supporting rejection of the null hypothesis of no difference among the sample groups).

Firmucutes, Proteobacteria, and Bacteriodetes constituted  $86.7\% \pm 8.8$  total average abundance across all axolotl samples (Fig. 4a). Of these phyla, Proteobacteria abundance considerably increased in the skin (5.2% to 41.8%) and in the gut samples (2.7% to 9.1%). In contrast, the abundance of this phylum did not significantly change in the fecal samples yet decreased in the stomach samples (52.0% to 38.9%). Firmicutes abundance seemed to follow the opposite trend. We thus employed Pearson's correlation to identify phylum level taxa (Supplementary Fig. S4), whose abundances negatively correlate as a result of metamorphosis. Abundances of Proteobacteria and Firmicutes along with Verrucomicrobia showed the strongest negative correlation (correlation coefficients were r = -0.64 and r = -0.54, respectively), which was also significant (False Discovery Rate (FDR) adjusted r = 0.0027



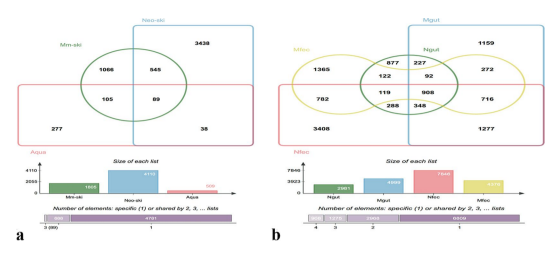
**Figure 5.** Differentially enriched genus level taxa and indicator species in samples from neotenic and metamorphic axolotl. The color scale bar indicates  $\log 2$  fold changes for absolute OTU abundances (DESeq2 analysis (q < 0.01) (**a**), Bubble plot showing indicator species. Only highly significant indicator values (IndVal >0.7, q < 0.01) are displayed. Size of bubble symbol is proportional to the mean relative abundance of indicator OTUs and the color scale bar shows indicator value for each OTU (**b**).

and 0.026, respectively). Interestingly, the phylum *Bacteriodetes* abundance in all samples substantially increased after metamorphosis (56.7% vs. 74.2% in fecal samples; 2.5% vs. 29.7% in gut samples; 11.6% vs. 17.5% in skin samples) and negatively correlated with both *Firmicutes* and *Proteobacteria* (although this result was not significant (q = 0.4). Overall, The axolotl skin microbiota, among others, showed most dramatic shifts between neotenic and metamorphic stages considering these negatively correlated taxa.

Percent average abundances of the genus level taxa are shown in a heatmap (Fig. 4b; for simplicity only taxa with abundance ≥5% in any sample in the same row were shown). Samples were grouped based on Bray-Curtis similarities (top dendogram) and abundances were clustered using hierarchical clustering (average linkage). The observed differences between group similarities were driven by differences in the relative abundance of multiple bacterial taxa. For example, metamorphic skin samples clustered with stomach samples of neotenic and metamorphic axolotl; Clostridium IV, Bacteriodes, and Sphingomonas were abundant in these samples. Notably, the genus Elizabethkingia were observed in high abundance in the skin and the stomach samples of the metamorphic axolotl. Metamorphic gut sample clustered with the fecal samples; Akkermansia and Bacteriodes being in greater abundance in these samples. Finally, neotenic gut and skin samples were grouped together. This pattern was chiefly due to taxa that could not be classified at genus level (unclassified Veillonellaceae, unclassified\_Ruminococcaceae, unclassified\_Lachnospiraceae) and Clostridium XIVa. Remarkably, the aqua samples did not have any taxa with high abundance shared with any samples from axolotl, and separated from other samples.

Indicator and shared species of neotenic and metamorphic axolotl. We next used DESeq2 analysis, a negative Binomial Wald Test $^{42,43}$  to identify differentially abundant taxa (q < 0.01) in neotenic and metamorphic axolotl organs at the genus level taxonomy. We also performed indicator species analysis to delineate high fidelity differentially abundant patterns (IndVal values  $\geq 0.7$ , p < 0.01). We largely observed concordance between both analyses. Skin samples showed the highest number of differentially enriched genera (49 taxa in metamorphic skin samples and 36 taxa enriched in neotenic skin samples); Pseudonocardia, unclassified\_Pseudonocardiaceae, Methylobacterium, Elizabethkingia, Vogesella, Chryseobacterium, and Zoogloea were the top scoring genera in the metamorphic skin while Limnohabitans and several taxa that were classified at higher taxonomic ranking were differentially abundant in the neotenic skin samples (Fig. 5a). These genera were also detected in the indicator species analysis in several OTUs. For example, OTU89, OTU141, OTU1052, OTU1301, OTU1450, OTU2081, were classified as *Pseudonocardia* and OTU89 had the highest abundance of 15.3%. Similarly, the genus Limnohabitans, overrepresented in neotenic samples, was assigned to OTU1542 and OTU2587, although both OTUs had relative abundance (0.01%) that can be considered a rare OTU<sup>39</sup>. The following top scoring taxa were differentially abundant in metamorphic and neotenic samples, respectively; stomach samples: Elizabethkingia (OTU335) and Limnohabitans (OTU46); gut samples: Elusimicrobium, which was not detected by IndVal but OTU7777 (*Hydrogenoanaerobacterium*) was the top scoring taxa (IndVal = 0.85, p = 0.002); *unclassi*fied Ruminococcaceae, unclassified Lachnospiraceae were the indicator species of neotenic gut samples and finally Odoribacter (OTU14388), Rikenella (OTU14233) were both differentially abundant and indicator species of metamorphic fecal samples (Fig. 5b).

Venn diagrams showed number of OTUs shared or unique among the samples isolated from axolotl (Supplementary Fig. S5a and b). The number of unique OTUs in all samples of neotenic (7422 OTUs) and metamorphic axolotl (6237 OTUs) was far grater than shared OTUs, indicating assembly of microbiota is tissue specific as in most other animals<sup>44</sup>. In terms of shared OTUs, 368 and 329 OTUs were present in all neotenic and metamorphic samples, respectively. We also collected water samples ("aqua") from the Aquarium of axolotl to identify OTUs present in the water that may have colonized axolotl skin. We found 89 OTUs shared by the neotenic and metamorphic skin and aqua samples; 38 OTUs were present only in the neotenic and aqua



**Figure 6.** Venn diagrams showing the number of unique and shared OTUs. Skin and aqua samples (**a**), and gut and fecal samples (**b**), collected from neotenic and metamorphic axolotls as indicated in the diagram.

samples while 105 OTUs were shared by metamorphic and aqua samples (Fig. 6a). However, relative abundances of all these OTUs were mostly less than 1%. We also compared unique and shared OTUs between gut tissue and fecal samples (Fig. 6b). Exceptionally, the number of OTUs in the metamorphic gut was 4999, representing substantial increase from 2961 OTUs found in the neotenic gut. Moreover, the two gut tissue samples shared only 227 OTUs. In stark contrast, the number of unique OTUs decreased to 1365 OTUs whereas neotenic fecal sample had 3408 unique OTUs. Notably, the shared number of fecal and gut OTUs, neotenic or metamorphic, were 288 OTUs and 272 OTUs, suggesting the fecal and gut microbiota are compositionally distinct. Finally, we identified the following genera as core taxa, i.e. shared by 90% of all samples (Core90) (Supplementary Fig. S6): Bacteroides, Clostridium XIVa, Clostridium XIVb, Akkermansia, Odoribacter, unclassified\_Veillonellaceae, unclassified\_Lachnospiraceae, Parabacteroides, unclassified\_Rhodospirillaceae, unclassified\_Ruminococcaceae, Rikenella, unclassified\_Clostridiale, and Desulfovibrio.

**Detection of human taxa in axolotl samples and predicted functions.** To answer the question if the captive axolotl may have acquired human bacterial taxa we compared axolotl microbiota with Human Microbiome Project (HMP) $^{45}$  stool and skin samples (HMP reference data: https://portal.hmpdacc.org) using weighted and unweighted Unifrac distances (Supplementary Fig. S7(a–d)). In particular, both neotenic and metamorphic stool and skin samples showed greater similarity to HMP samples compared to other axolotl samples. These results encouraged us to look at predicted functions of the microbiota using Phylogenetic Investigations of Communities by Reconstruction of Unobserved States (PICRUSt). Due to the specific requirement by PICRUSt, a separate OTU table was generated accordingly using a closed reference analysis based on the GreenGenes 99 database version. The CAP analysis of the predicted functions revealed significant differences between the Bray-Curtis distances based on putative pathway abundances (tr(Q\_m'HQ\_m): 5,78539 p = 0.001). However, Bray-Curtis similarities among all distinct groups were around 90%, pointing to the shared predicted functions among the bacterial microbiota of axolotl organs (Supplementary Fig. S8).

Histological comparison of gut tissue and limb regeneration capacity between neotenic and **metamorphic axolotl.** We performed histological comparison of neotenic and metamorphic gut tissue. Results demonstrate structural similarities at both stages rather than differences. Mucus producing goblet cells were more abundant in metamorphic samples, which was also accompanied by a thicker mucosa layer (Supplementary Fig. S9). We then attempted to answer the question whether shifts in microbiota might be accompanied by a change in host regenerative capacity since previous studies provided evidence that limb regenerative capacity is reduced in metamorphic axolotl<sup>38</sup>. We reproduced these observations in this study that regeneration is indeed impeded in the limbs of metamorphic axolotl (Supplementary Fig. S10). We observed that all neotenic animals (n = 15) regenerated (day 64) the limb in a miniaturized form with four digits. In contrast, slower blastema formation and regeneration process were apparent in metamorphic animals. We continued to observe limb generation in metamorphic axolotls up to day 150 (Supplementary Fig. S11(a-c)). At day 150, 2 of 15 metamorphic axolotls (13%) restored the limb with four digits, as exemplified in Fig. S11a. Also, 4 of 15 metamorphic animals (27%) were capable of regenerating amputated limb with 3 digits only (Fig. S11b,c). In addition, another 4 animals (27%) restored limb with only two digits were observed. Rest of the animals (5 out of 15) failed to regenerate a limb to any extent, indicating that the limb generation capacity in metamorphic axolotls is severely impeded.

# Discussion

The main purpose of this study was to comparatively characterize bacterial microbiota of axolotl in neotenic and metamorphic life stages since microbiota of this important biological model has not been reported before. Our results show that substantial shifts occurs in the structure and composition of microbiota, particularly in the skin but also in digestive organs.

Our results in terms of the composition of axolotl microbiota broadly parallel the previous reports on microbiota profile of amphibians, and salamanders in particular. Five major dominating phyla, Firmicutes, Bacteroidetes, Proteobacteria, Verrucomicrobia and Actinobacteria were abundant among all studied samples, which is consistent with previous amphibian studies<sup>37,46–49</sup>. Salamander microbiota was previously studied but often distinct species of salamander skin microbiota was profiled<sup>46,48</sup> and systematic investigation of microbiota diversity in multiple organs of salamander species, particularly that of axolotl, has been missing in the literature with few exception<sup>46</sup>. Characterized both gut and skin microbiota of fire salamanders within the natural habitat. Surprisingly, like captive axolotl in this study, the wild salamanders' gut and skin are associated with the above mentioned five major phyla. The genera Chryseobacterium, Pseudomonas, Flavobacterium, Sphingobacterium, Novosphingobium, were reported to be dominant taxa in skin of fire salamander living in nature. Interestingly, we detected these genera in the neotenic skin samples in this study albeit in low abundance but substantially increased in abundance in the metamorphic skin. These bacterial genera belong to a large, ecologically diverse group, and their members include known pathogens; some could be opportunistic while some others can outcompete emerging fungal pathogens<sup>50</sup>. Interestingly, our analysis revealed that metamorphic axolotl skin was dominated by *Pseudonocardia*  $(15.7\% \pm 10.4)$ , and an unclassified taxa from *Pseudonocardiaceae* family  $(9.7\% \pm 5.7)$ , both are indicator species of the metamorphic skin (IndVal = 0.99, p = 0.006). *Pseudonocardia spp.* is a well known antifungal commensal microorganism<sup>51</sup> and colonize on the integument of fungus-gardening ant species. Recruitment of the members of this genus in high abundance by metamorphic axolotl might reflect host-symobiont synergy against pathogenic fungi.

We sequenced pools of water samples from the axolotl aquarium ("aqua") to identify water-borne bacterial taxa acquired by axolotls. Surprisingly, most abundant genera of aqua samples (e.g. *Acidovorax, Armatimonas, Flectobacillus*) were not associated with axolotl organs but only a subset of low-abundance bacteria were detected. For example, *Aquabacterium* abundance in metamorphic skin and neotenic stomach were 3.9% and 7.3%, respectively while its average abundance in aqua samples was 0.1%. Our results are consistent with previous work reporting amphibian skin may select rare taxa from the environment 50,52-55. In this study, we found that 89 out of 509 OTUs in aqua samples were present in neotenic skin samples while 105 OTUs shared between aqua and metamorphic skin samples (see Fig. 6), and 150 shared OTUs with aqua samples, the highest among others, with metamorphic stomach samples. Consequently, our results support the notion that both host and external factors shape the host microbiota but host genetics applies selective filter.

Conversely, diet is another crucial factor strongly influencing structure of gut microbiota of animal host and even dominate host genotype<sup>52</sup>. Although host genotype and diet are constant in this study, metamorphosis is likely to cause remodeling of epigenetic landscape in the host genome, which in turn is expected to reshape microbiota. Notably, fecal samples from the neotenic and metamorphic axolotls clustered together, albeit richness in metamorphic fecal samples significantly decreased. We observed that feeding behavior of the metamorphic axolotl change during metamorphosis, the animals tend to eat less often (low appetite), which might account for the decreased fecal diversity. Although no major restructuring of intestine via metamorphosis was apparent as described before 16, we observed a higher number of goblet cells and thicker mucus layer in the metamorphic gut tissue compared to neotenic gut (Supplementary Fig. S9). Taken together, relative influence of diet and host epigenetics seems to be compartmentalized; diet appears to influence strongly the bacterial diversity in the fecal microbiota in the gut lumen whereas the host epigenetics (and the resulting changes in transcriptome due to metamorphosis) seems to play a greater and selective role in gut tissues (crypts). Some genera, often associated with symbiosis such as Alistipes and Elusimicrobium, were differentially enriched in the metamorphic gut tissues whereas neotenic gut tissues were represented by two genera with unclassified Clostridiaceae and unclassified Enterobacteriaceae. Surprisingly, the abundance of Akkermansia considerably increased in the metamorphic gut tissues (16%) relative to neotenic gut (9%). A. muciniphila within this genus is known to be a mucin degrading and symbiotic bacterium and the increase in abundance of this genus is meaningful with increasingly thicker mucus layer we observed in the metamorphic gut tissue staining.

Our analysis revealed that axolotl microbiota to certain extent was "humanized" in captivity as manifested from similarity of gut and skin microbiota with the human microbiota, which prompted us to compare predicted functions using PICRUSt<sup>53</sup>, which is optimized for human microbiota. We found that all samples included in microbiota analysis were highly similar based on the abundance of predicted microbial genes (40%). Though further study using shotgun metagenomics technology is warranted, our findings raise the possibility that deficiency in the microbial community function in a given organ can be possibly compensated without taxonomic coherence. However, caution should be exercised in interpreting these results considering the accuracy of the predicted functions is predicated on the closed reference database, whereas the majority of OTUs in this study were clustered using open reference.

Finally, recent studies on structure of microbiota in regeneration and wound healing \$1,54,56,57, point toward potential functional link between microbiota and regeneration. Considering the cross talk between microbes and the host immune system \$36, transformation of skin immunity (thickening mucus layer \$16 and increased secretion of antimicrobial peptides) may partly account for the reconstruction of microbiota after metamorphosis. But whether the restructured microbiota might influence host physiology is unknown. In this regard, we tested whether we can reproduce previous finding on reduced metamorphic limb regenerative capacity \$38 and validated that this trait is indeed steeply diminished in metamorphic axolotl while neotenic animals maintain normal regenerative capacity under identical conditions (Supplementary Fig. S10). Importantly, we detected sharp expansion of *Proteobacteria* in metamorphic axolotl skin relative to the neotenic skin and noted steep decrease of bacterial diversity in the fecal and skin samples. In a previous study \$55 some members of *Proteobacteria* were demonstrated to be causally linked to reduced regenerative capacity in planaria, another biological model for regeneration studies. Even though our observations do not provide a causal link, we hypothesize that some potentially opportunistic members of *Proteobacteria* (e.g. *Vogesella, Sphingomonas*) and others could play a role in

reduced limb regenerative capacity in axolotl. Further research is warranted to test this hypothesis in carefully designed experiments allowing for causal relationship.

#### Conclusions

Our study provides multi-organ profile of axolotl microbiota. To our knowledge this is the first comprehensive report on axolotl microbiota. We demonstrated that microbiota inhabiting axolotl organs considerably restructure upon metamorphosis leading to expansion of opportunistic bacteria within *Proteobacteria*. Lack of systematic studies on axolotl microbiota has hindered its potential as a fruitful model in host-microbe interaction studies. Therefore, the data presented here make a significant contribution for further characterization of a valuable biological model for regenerative and development biology. The paper concludes by arguing that physiological implications of considerable shifts in microbiota should be taken into consideration when captive axolotl is employed in developmental and regeneration studies.

# Methods

Ethical statement and experimental design. The local ethics committee of the Istanbul Medipol University (IMU) authorized experimental protocols and animal care conditions (the authorization number: 38828770-E.7856). All experiments were performed in accordance with relevant guidelines and regulations. The experimental design followed in this study is depicted in Fig. 1. Briefly, a total of 48 adult axolotls (12-15 cm in length, 1 year old) were obtained from the animal care facility of the IMU. Axolotls were chosen among the siblings and they were maintained one animal per aquarium after hatchling till the end of experimental procedures. Out of 48 axolotls 18 were used for metamorphosis experiments; and 30 were reserved for amputation/limb regeneration experiments. Half of both groups (15 for regeneration experiments and 9 for metamorphosis; 24 total) were then induced individually to undergo metamorphosis by L-thyroxine (Sigma-Aldrich, St Louis, MO, USA, Cat. No. T2376) as described elsewhere<sup>27</sup>. Briefly, T4 solution was prepared by dissolving L-thyroxine in Holtfreter's solution (final concentration 50 nM) and was administrated to animals that were maintained individually throughout the experimental period. The medium was replaced with freshly prepared T4 containing solution every third day and animals were monitored for morphological changes such as weight loss, disappearance of the fin and decrease in the gills size which was achieved within 3 weeks following the T4 administration. Hormone treatment continued for another 3 weeks until fully metamorphic axolotls were obtained. Before amputation and sample collection steps, metamorphosed animals were kept for an additional 1-month without T4 administration to allow for metamorphosing axolotls to adapt the terrestrial life conditions. Both neotenic and metamorphic animals were maintained in individual aquaria water at 18 ± 2 °C in Holtfreter's solution. Axolotls were maintained in the university animal facility within a batch-flow aquarium system where the batch water was treated with UV light and filtered to prevent infections. All animals were kept in the same aquatic solution and fed with same diet. Animals were fed once a day using a staple food (JBL Novo LotlM, Neuhofen, Germany). Axolotls did not receive any antibiotic treatment throughout the experiments.

Sample collection. Animals were sacrificed using 0.2% MS222 (Sigma-Aldrich, St Louis, MO, USA. Cat. No. E10521) approximately in two months upon visual observation of metamorphosis. Neotenic and metamorphic animals formed two major experimental groups. To help account for interindividual variation randomly selected three animals were grouped, albeit still maintained in their separate aquarium, to have three biological replicates (R1, R2 and R3) for neotenic and metamorphic experimental groups (Fig. 1). For each replicate skin, stomach, intestine and fecal samples were harvested from three animals within R1, R2, and R3 and pooled together (technical replicates) under sterile conditions in a bio safety cabin. To collect the skin samples, animals were rinsed in sterile water to get rid of the transient bacteria. As the next step, skin samples from the mid styloped level of the right forelimb were isolated for each replicates with punch biopsy using disposable and sterile 6 mm diameter punches (Miltex, York, PA, USA). Stomach and intestine samples were collected after dissection of animals. Intestine contents were first removed, split into approximately equal 4-5 pieces, and rinsed with sterile serum physiologic solution five times. Fecal samples were harvested from the rectum. Isolated and pooled samples for each replicate were frozen in liquid nitrogen immediately for cryopreservation. All samples were stored at  $-80\,^{\circ}$ C till DNA isolation. To compare harvested samples with the microbial structure of Holtfreter's solution ("aqua" samples), water samples (100 ml in total) from randomly chosen aquariums were collected and 3 samples were pooled together to have three replicates (R1, R2 and R3).

**DNA extraction.** DNA isolation from the skin, intestine, and stomach samples was carried out with DNeasy Blood & Tissue kit (Qiagen) according to the manufacturer's protocol. QIAamp DNA Stool Mini Kit (Qiagen) was used to extract the DNA of the fecal samples by following the manufacturer's recommendations. To extract DNA from the water samples,  $100 \, \text{ml}$  of water was filtered through filters with  $0.2 \, \mu \text{m}$  pore size and the filter papers subsequently were used in DNA extractions using metagenomic DNA Isolation Kit for Water (Epicentre, Cat. No. MGD08420) by following the producer's protocol. Spectramax i3 (Molecular Devices, Sunnyvale, CA) was used to measure the concentrations of isolated DNA. Quality of DNA samples was checked by electrophoresis in 1.0% agarose gels.

**PCR and sequencing of 16S rRNA amplicons.** To amplify the variable V3-V4 regions of the 16S rRNA gene, the primers 341 F (5'-CCTACGGGNGGCWGCAG-3') and 805 R (5'-GACTACHVGGGTATCTAATCC-3') were used  $^{58}$ . MiSeq sequencing adaptor sequences were added to the 5' ends of forward and reverse primers. Approximately 12.5 ng of purified DNA from each sample was used as a template for PCR amplification in  $25\,\mu l$  reaction mixture by using 2 × KAPA HiFi HotStart ReadyMix (Kapa Biosystems, MA, USA). For PCR amplification, the following conditions were followed: denaturation at 95 °C for 3 min., followed by 25 cycles of denaturation at 95 °C for 30 sec., annealing at 55 °C for 30 sec. and extension at 72 °C for 5 min. No template negative control samples were included to check PCR contamination and none

of the negative controls yielded detectable level of amplification on agarose gels. Amplified PCR products were purified with Agencourt AMPure XP purification system (Beckman Coulter) and Nextera PCR was performed by using sample-specific barcodes. Constructed Nextera library was then sequenced by Illumina MiSeq platform using MiSeq Reagent Kit v3.

Sequence processing, clustering, and taxonomic assignment. To analyze the paired-end sequencing data Quantitative Insights Into Microbial Ecology (QIIME, v1.9.1)<sup>59</sup> software was used at the Nephele platform (v.1.6, http://nephele.niaid.nih.gov). Nephele platform was also used for the Phylogenetic investigation of communities by reconstruction of unobserved states (PICRUSt) analysis<sup>53</sup> and for comparing the data with the Human Microbiome Project (HMP). Before submitting the raw reads into the Nephele pipeline, primers were removed using cutadapt program<sup>60</sup> and the pipeline was stringently configured to perform the following steps; reads below average quality scores (q < 30) and read-length >450 bp were eliminated. After joining the pair-end reads, the reads were clustered into operational taxonomic units (OTUs) using open reference OTU-picking strategy<sup>58</sup>. The open-reference approach initially runs a closed-reference step. Sequences that fail closed-reference assignment are then clustered as de novo OTU based on pairwise similarity among all sequences in the data set. Open reference clustering was performed based on the 97% clustered SILVA reference (SILVA 123 release;) database<sup>61</sup> and SortMeRNA combined with SUMACLUST algorithms<sup>62</sup>. Non-matching reads to closed reference were subsequently clustered de novo. After obtaining the OTU table from the pipeline, UCHIME (v.4.2) program (http://drive5.com/uchime) integrated into the Mothur (v.1.39.5) tool<sup>63</sup> was separately run to remove chimeric reads. Additionally, The RDP classifier<sup>64</sup> (v. 2.2) was locally run to assign taxonomy for each OTU at a confidence greater than 70% cutoff. Reads that could not be classified at the genus level were sequentially assigned to higher taxonomic hierarchy up to the kingdom level. Unclassified reads at the kingdom level ("Unclassified Bacteria") were extracted from the OTU-representative sequences and searched for nearest neighbor method using MOLE-BLAST<sup>65</sup>. This tool computes a multiple sequence alignment (MSA) between the query sequences along with their top BLAST database hits, and generates a phylogenetic tree. Species richness and diversity were estimated by QIIME with the following alpha diversity metrics: OTU richness, Chao1, Shannon, Simpson E, Inverse Simpson, and Faith's phylogenetic diversity (PD). We assessed normality of alpha diversity data using Shapiro-Wilk tests and compared the metrics between neotenic and metamorphic microbiota using unpaired two-tailed t-test. Venn diagrams were constructed using jvenn, a web based tool (http://jvenn.toulouse.inra.fr)<sup>66</sup>.

Multivariate analysis of community structures and diversity. Bray-Curtis similarity index<sup>67</sup> and Jaccard index of similarity<sup>68</sup> were used to obtain distance matrix after standardizing by the column sums and transforming (square-root) the read abundance data. Similarities in microbial community structures among samples were first displayed using Principal Coordinate Analysis (PCO) (unconstrained). Differences in community structure related to metamorphosis were displayed using a constrained ordination technique, Canonical Analysis of Principal coordinates (CAP). Tests of the multivariate null hypotheses of no differences among a priori defined groups were examined using PERMANOVA and the CAP classification success rate. CAP uses PCO followed by canonical discriminant analysis to provide a constrained ordination that maximizes the differences among a priori groups and reveals patterns that cannot be unraveled using unconstrained ordinations<sup>41</sup>. CAP classification success rates and CAP trace<sub>Q\_m0HQ\_m</sub> statistics were examined in combination to draw conclusions about separation of a priori groups. Permutational analysis of multivariate dispersions (PERMDISP)<sup>69</sup> was used to test for heterogeneity of community structure in a priori groups. PERMANOVA, CAP and PERMDISP were performed with 9999 permutations and run as routines in PRIMER6<sup>70</sup>.

To delineate bacterial taxa responsible for the multivariate patterns and differentially enriched taxa between the neotenic and metamorphic axolotl organs, we used DESeq. 2, a negative Binomial Wald Test<sup>42,43</sup> and indicator species analysis<sup>71</sup> DESeq. 2 analysis results, along with core OTU heatmap, phylum correlation heatmap, and read count figures were obtained using MicrobiomeAnalyst<sup>72</sup>. To perform indicator species analysis, R package labdsv was used. Boxplots, barchart, bubbleplots, and heatmaps were generated using R packages including vegan, ggplot2, heatmap2, Heatplus, reshape, colorramps, and RcolorBrewer (R Core Team 2018, http://www.r-project.org).

**Histology.** Removal of fecal from the isolated intestine samples was followed by fixation in 10% neutral buffered formalin (NBF) for 48 h. The samples were then processed by immersion of materials in ascending alcohol series, toluene and embedding in paraffin.  $4\mu m$  thick tissue sections were obtained by using microtome. Sections were deparaffinized and stained with Hematoxylin and Eosin (Bio-Optica Mayer's Hematoxylin and Eosin Y Plus), Masson's Trichrome (KIT, Masson Trichrome with aniline blue, Bio Optica, 04-010802), Alcian Blue (KIT, Alcian Blue Acid Mucopolusaccharides staining, Bio-Optica, 04-160802) according to manufacturer's instructions. Images were taken by using the NIKON DS-Fi2-U3 Digital Camera. The detailed protocol was described in a previous work<sup>16</sup>.

**Amputation.** A total of 30 Axolot were used for regeneration experiments. Half of the animals were induced to undergo metamorphosis by T4 administration as described above. After complete metamorphosis was achieved, right forelimb of both neotenic and metamorphic animals were amputated from the mid-stylopod site Both macroscopic and microscopic pictures were taken by using Nikon D3200 camera and Zeiss Axio zoom V16 microscope, respectively. Animals were anesthetized in 0.1% MS222 (Sigma-Aldrich, St Louis, MO, USA. Cat. No. E10521) for all animal procedures.

**Availability of data and materials.** The datasets generated and/or analyzed in this study, the OTU table, and the pipeline generated log file, the run parameters and metadata template are available in the figshare repository: https://figshare.com/s/b1f232d8d054e0e20f06. Raw sequencing data have been deposited in the NCBI SRA database (BioProject ID PRJNA432888, www.ncbi.nlm.nih.gov/sra).

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# **Author Contributions**

T.D. and S.Y. conceived the study. T.D., B.Y., İ.K., A.E.İ. and E.C.F. performed animal experiments. T.D., G.Ö. and S.Y. acquired sequencing data; T.D., G.O., and S.Y. analyzed and interpreted the data. T.D., G.O., G.Ö., and S.Y. drafted and critically reviewed/revised the manuscript. All authors read and approved the final manuscript.

# **Additional Information**

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