

1	An Introduction to Phylosymbiosis
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ABSTRACT

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Phylosymbiosis was recently formulated to support a hypothesis-driven framework for the characterization of a new, cross-system trend in host-associated microbiomes. Defining phylosymbiosis as "microbial community relationships that recapitulate the phylogeny of their host", we review the relevant literature and data in the last decade, emphasizing frequently used methods and regular patterns observed in analyses. Quantitative support for phylosymbiosis is provided by statistical methods evaluating higher microbiome variation between host species than within host species, topological similarities between the host phylogeny and microbiome dendrogram, and a positive association between host genetic relationships and microbiome beta diversity. Significant degrees of phylosymbiosis are prevalent, but not universal, in microbiomes of plants and animals from terrestrial and aquatic habitats. Consistent with natural selection shaping phylosymbiosis, microbiome transplant experiments demonstrate reduced host performance and/or fitness upon host-microbiome mismatches. Hybridization can also disrupt phylosymbiotic microbiomes and cause hybrid pathologies. The pervasiveness of phylosymbiosis carries several important implications for advancing knowledge of ecoevolutionary processes that impact host-microbiome interactions and future applications of precision microbiology. Important future steps will be to examine phylosymbiosis beyond bacterial communities, apply evolutionary modeling for an increasingly sophisticated understanding of phylosymbiosis, and unravel the host and microbial mechanisms that contribute to the pattern. This review serves as a gateway to experimental, conceptual, and quantitative themes of phylosymbiosis and outlines opportunities ripe for investigations from a diversity of disciplines.

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Keywords: symbiosis; phylosymbiosis; microbiome; host-microbe interactions



1. INTRODUCTION

The last decade has brought renewed interest in the complexity of microorganisms living in association with hosts, yielding a number of new empirical results, philosophical concepts, and research opportunities (1,2). Any discussion on the study of host-microbiome interactions must begin with clear definitions. Here, we use the term symbiosis (*sym* – "together", *bios* – "life" in Greek) to encompass associations between two or more organisms of different species and without restriction to the length of time of the association or phenotypes produced by the interacting species. Since temporal and functional variation in symbiosis is context-dependent, symbiotic interactions can include a range of obligatory, facultative, transient, and permanent associations with varying degrees of specificity and functional costs and benefits.

The last two decades of research and technological advances have placed microbial symbiosis as a nexus of many subdisciplines within and beyond biology. Scholars now have a suite of tools and increased awareness of the major questions to be answered. These include holistic approaches for the identification of ecological (3) and host (4-7) drivers of microbial taxonomic and functional diversity, as well as reductionist approaches that provide evolutionary and mechanistic insights into transmission processes (8) and phenotypic outcomes of symbiosis (1). The abundance of empirical and theoretical investigations on the ecology and evolution of simple symbioses also comprise fertile ground to build a foundation for the microbiome field that studies frequently complex associations between hosts and their multiple microbial associates. One rapidly growing research area across diverse systems is the recently defined pattern of phylosymbiosis (9). This review aims to synthesize the topic to provide: (a) a long-lasting definition of the term; (b) a practical guide to test phylosymbiosis; (c) an overview of the prevalence of phylosymbiosis; (d) a



72 discourse on the biological significance of phylosymbiosis; and (e) future directions in 73 phylosymbiosis research. 74 75 2. WHAT IS PHYLOSYMBIOSIS? 76 We use the following quote to describe our initial and basic definition of phylosymbiosis, namely 77 "microbial community relationships that recapitulate the phylogeny of their host" (9). 78 Phylosymbiosis is first and foremost a significant association between host phylogenetic 79 relationships and host-associated microbial community relationships wherein "phylo" refers to 80 host clade and "symbiosis" refers to the microbial community in or on the host. 81 82 Prior to the introduction of the term phylosymbiosis in a study of *Nasonia* parasitoid wasp species 83 (9), early investigations specified relationships between host phylogenies or genetic distances with 84 microbial beta diversity in maize (10), insects (5.11), and mammals (4.12). These studies utilized 85 bacterial 16S rRNA gene sequencing across multiple host species to demonstrate that closely-86 related species harbor more similar microbiomes than distantly-related species. For example, the 87 sister species N. giraulti and N. longicornis diverged ~0.4 million years ago and harbor more similar 2nd instar larval, pupal, and adult microbiomes compared to the microbiome in their 88 89 outgroup species N. vitripennis (9,11), which diverged ~1.0 million years ago from the two sister 90 species (13). 91 92 Phylosymbiosis may arise from stochastic and/or deterministic evolutionary and ecological forces. 93 For example, stochastic effects include dispersal fluctuations in microbial communities (ecological 94 drift) or shifts in host geographic ranges (14). Phylosymbiosis can also be shaped by ecological 95 (15-17) and dietary (4) niche variation across host lineages. Deterministic effects include microbial



colonization preferences for certain host backgrounds or host regulation in which microbial community composition is influenced by host trait(s) (18). The first study linking phylosymbiotic patterns to the function of specific host genes found that knockdown of the *Hydra* armenin antimicrobial peptide disrupted phylosymbiosis (6) commonly observed in several freshwater and laboratory *Hydra* species (19). Although phylosymbiosis can potentially arise from long-term, intimate host-microbe associations over evolutionary time, such as through host-microbe coevolution, co-diversification (20), and co-speciation (21), importantly it may also be driven by relatively short-term changes in microbiome composition. Indeed, a recent *Drosophila melanogaster* study revealed the effects of gut microbiome changes on host genomic divergence in as little as five generations (22).

While phylosymbiosis distinguishes itself from non-phylosymbiosis by a significant degree of association between host phylogenetic and microbiome community relationships, it is not universal (Section 5) and therefore provides a testable hypothesis. Determining the presence of phylosymbiosis is a first step preceding further investigations into eco-evolutionary mechanisms, such as the nature of species-species associations, selective or neutral forces driving phylosymbiosis, and the (in)consequences of the pattern on host and microbial phenotypes. If phylosymbiosis results from an evolutionary selective pressure, then decreases in host or microbial fitness are expected upon host exposure to microbiomes from different host lineages in an evolutionary-informed manner. Evolutionary selective pressures for phylosymbiosis could drive the spread of host traits that regulate microbiome composition or microbial traits that enhance host colonization. In this general light, we refer to "functional phylosymbiosis" when host and/or microbial phenotypes impact or are impacted by phylosymbiotic associations.



Interspecific microbiome transplant experiments are useful in elucidating functional phylosymbiosis. A large-scale phylosymbiosis investigation spanning 24 species across four laboratory-reared host clades (*Nasonia* wasps, *Drosophila* flies, mosquitoes, and *Peromyscus* deer mice) demonstrated that interspecific transplants of gut microbial communities between *Peromyscus* species decreased dry matter digestibility and increased food intake, while transplants between *Nasonia* species markedly lowered survival to adulthood by nearly half (23). In addition, interspecific microbiomes are more costly to *Nasonia* larval growth and pupation than intraspecific microbiomes (24). Similarly, reciprocal maternal symbiont transplants between two wild, sympatric *Ontophagus* dung beetle species caused developmental delay and elevated mortality in non-native hosts that persisted to the next generation (25). Collectively, phylosymbiotic associations that impact host fitness support the premise that hosts are adapted to their native microbiomes rather than non-native microbiomes, although more studies are needed to confirm these associations and effects in captive and wild host populations-

Hybridization between host species causes host-microbiome mismatches since combining independently-evolved host genotypes in a hybrid may cause a breakdown in either microbial colonization preferences for certain hosts or host control of the microbiome. As demonstrated in *Nasonia* (9), house mice (26), and whitefish (27), hybrids have an altered microbiome relative to the parental microbiome, suggesting a reduced capacity for hosts to regulate their microbiomes and an increased capacity for pathogenic microbes to bloom. These breakdowns in host-microbiome interactions can associate with maladaptive phenotypes in hybrids including immune dysfunction, pathology, inviability, and sterility (9,26) that can reduce interbreeding between



species or populations. In *Nasonia*, lethality of hybrids between the older species pair was rescued by germ-free rearing and restored by feeding an inoculum of select, resident gut bacterial species from parents to germ-free hybrids (9). In contrast, hybrids between a younger *Nasonia* species pair did not have an altered microbiome nor suffer functional costs. Collectively, the results from interspecific microbiome transplant experiments and host hybridization studies illustrate that host-microbiome interactions across host species can have important functional consequences that impact evolutionary events within and between species, including wedging host populations into species.

3. WHAT IS NOT PHYLOSYMBIOSIS

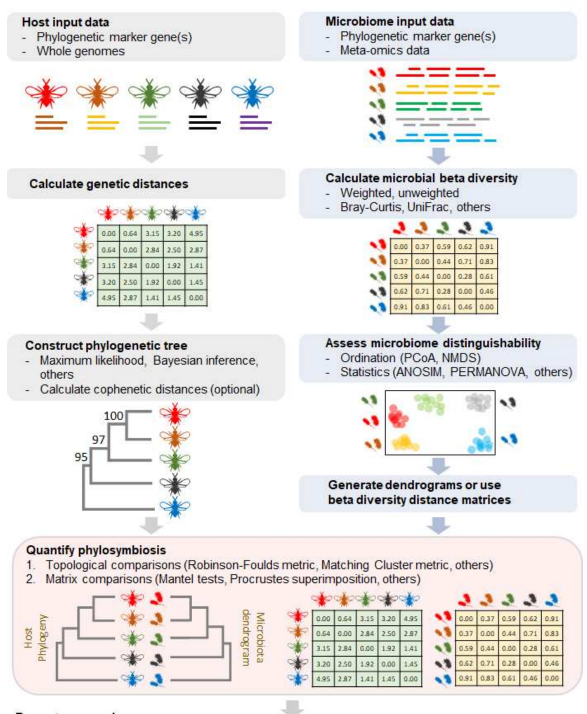
Having now summarized phylosymbiosis, we briefly accentuate what phylosymbiosis is not for clarity. Early misconceptions associated the term with strictly narrow presumptions such as vertical transmission, mutualistic interactions, or evolutionary splitting from a common ancestor via co-evolution, co-speciation, co-diversification, or co-cladogenesis. Although these processes may lead to phylosymbiosis, the pattern may alternatively arise by antagonistic interactions and/or horizontal microbial transmission whereby interactions between hosts and environmental microbes establish phylosymbiosis anew each generation. As such, phylosymbiosis has varied underpinnings subject to empirical investigation, and it may appear at certain points of time and space rather than be stable throughout a host's entire lifespan.

4. A PRACTICAL GUIDE TO STUDYING PHYLOSYMBIOSIS

Investigations of phylosymbiosis vary in approach (qualitative vs quantitative), methodology, and statistical power (18). Thus, a clear, consistent, and robust workflow to detect phylosymbiosis is



desirable for newcomers and experts alike. Here, we suggest a comprehensive workflow for examining phylosymbiosis (**Figure 1**).



Downstream analyses

- Phylogenetic comparative methods, model fitting, functional correlations, experimental validation



Figure 1. Sequential overview of bioinformatic methods commonly used for phylosymbiosis analyses.

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Host taxa and input data. Because phylosymbiosis detection involves the collection of replicated samples across multiple taxa, both optimization of statistical sensitivity (28) and specificity (18) as well as minimization of sequencing batch effects are crucial for differentiating between noise and signal. Although our 2016 study showed that rooted trees with four Nasonia species are sufficient to detect phylosymbiosis within the clade (23), we suggest the use of appropriate power and effect size analyses (reviewed in (29) for microbiome data) to determine sufficient replicates and taxa for the optimization of statistical power (28). Sampling multiple individuals per species will help resolve noise from signal in microbial community relationships, but further study is required on how replicates of inter- and intraspecies samples are best utilized in studying phylosymbiosis across host clades that can vary in divergence times. If available, experimental designs of successful phylosymbiosis studies with similar sample types can also be adapted accordingly (30). Previous studies have successfully detected phylosymbiosis in host taxa spanning ~0.3-100 million years of evolutionary history (21,23), and whether longer times since a last common ancestor impacts phylosymbiosis detection requires further study. Nucleotide or amino acid sequence(s) from host species can be used to generate a phylogenetic or phylogenomic tree that is confidently supported at branching nodes with bootstrap (31) or other measures (32) and across several phylogenetic inference methods (e.g., maximum likelihood (33) and Bayesian inference (34)). Because an accurate host phylogenetic topology is essential for evaluating phylosymbiosis, the tree should be free from systematic artifacts such as long branch attraction; polytomies should be resolved in the host phylogeny when possible. As methods used to





reconstruct a host phylogeny from a sequence alignment have been extensively reviewed (35), we will not discuss them further here. With a host evolutionary tree, pairwise host distances can also be represented as cophenetic distances, computed as the sum of branch lengths connecting a pair of terminal nodes on a phylogenetic tree (36).

Microbiome input data. Phylosymbiosis analysis requires microbial diversity data from each host lineage. Short-read sequencing of microbial phylogenetic marker genes (e.g 16S rRNA gene) is common and economical for microbial profiling. Processed sequenced reads can be analyzed by one of two current methods. First, they can be clustered into operational taxonomic units (OTUs) at different sequence cutoffs (e.g., 97% and 99%) with and/or without reference sequence database (37,38). OTU clustering cutoffs reflect genetic distances between taxa over evolutionary time and may affect phylosymbiosis detection (39); such variability has also been observed in practice (reviewed in (18)). Second, reads can be resolved into amplicon sequence variants (ASVs) without clustering, which may offer single-nucleotide resolution, though sequencing error rates should be accounted for (40). For the greatest sensitivity in phylosymbiosis assessment, meta-omics datasets are advantageous because finer-scale taxonomic and functional profiling can be achieved (41). Metagenomic sequence data were used to demonstrate viral phylosymbiosis in Nasonia (42) as well as the varying effects of host phylogeny and ecology on the composition and functions of non-human, primate gut microbiomes (43,44).

Microbial beta diversity measures. Microbial beta diversity, which measures dissimilarities in microbial composition and structure across host samples, is conventionally used to measure phylosymbiosis. Binary measures, such as Jaccard distance and Sørensen-Dice distance (45,46),



are calculated with OTU presence/absence data. Quantitative descriptors of OTU abundances can also compute beta diversity, including the Bray-Curtis dissimilarity (47) derived from Motyka *et al.*'s coefficient (48). It simplifies as 1-[2w/(a+b)], in which w is the sum of the minimum abundances of common species across two host samples, a is the sum of the abundance of all OTUs/species in one sample, and b is the sum of the abundance of all OTUs/species in the other. Phylogeny-based metrics, such as weighted and unweighted unique fraction (UniFrac), use phylogenetic distances between communities (samples) to calculate microbial community differences, necessitating the use of a phylogenetic tree as input (49).

Because beta diversity metrics reflect different aspects of dissimilarity, the choice of metric is study specific and depends partly on the microbial composition and evolutionary history of the lineages studied. Binary metrics based on presence/absence are more sensitive to variations in rare taxa and were implemented to study host specificity of sponge microbiomes, where rare taxa comprised more than 90% of distinct OTUs (50). Binary metrics may also be sensitive to recent microbial diversification because recently diverged OTUs/ASVs will exert the same effect as OTUs/ASVs with a longer divergence history (39). In contrast, quantitative metrics are more sensitive to variations in abundant taxa. Besides taxonomy-based phylosymbiosis studies (23,51-53), quantitative metrics have also been applied to metagenomics data (42,43). Metrics that consider phylogenetic relationships between OTUs, such as UniFrac distances, (54) are applied in many other phylosymbiosis studies, including bats (55), corals (20), and mammals (4,43).

Microbiome distinguishability, representative of microbial beta diversity differences between host lineages under evaluation, is a prerequisite for phylosymbiosis (20,23,51-53). Microbiome



distinguishability can be visualized from beta diversity data and categorical sample grouping data using ordination plots, such as principle coordinate analysis (PCoA) and non-metric multidimensional scaling (NMDS) plots (56). In addition, microbiome distinguishability can be further evaluated using typically non-parametric multivariable analyses, such as analysis of similarities (ANOSIM) (57) and variants of permutational multivariate analysis of variance (PERMANOVA) (58). Specific pairwise comparisons of intra- and interspecific microbial beta diversity distances can also be performed with an appropriate non-parametric two-sample test (23).

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245 Quantifying phylosymbiosis. The determination of phylosymbiosis relies on evaluating a 246 significant association between host phylogenetic relationships and host-associated microbial 247 community distances. To this end, topological congruency tests directly compare topologies of a 248 host phylogenetic tree and a microbiome dendrogram (23,42,51-53,59). To generate a hierarchical 249 dendrogram, several agglomerative hierarchical clustering methods (reviewed in (56)) can cluster 250 microbial beta diversity distances. The most commonly used method, unweighted pair group 251 method with arithmetic mean (UPGMA), performs pairwise sample clustering from their average 252 dissimilarity values and gives all samples equal weights (60). Compared to linkage clustering 253 approaches, UPGMA prioritizes relationships among groups over individual samples (56). By 254 assigning equal weights to all samples, UPGMA assumes that samples in each group are 255 representative of groups in the larger reference population (56). As such, it may be sensitive to 256 sample sizes and may generate unstable topologies with imbalanced data where some groups are 257 oversampled while some are undersampled. Newer clustering methods, such as the 258 phylogenetically-aware squash clustering method, directly compute distances between samples 259 (rather than differences between beta diversity distances) based on their positions on a



phylogenetic tree (61). In general, the effects of clustering methods on phylosymbiosis detection require further study.

Topological comparison metrics, such as the Robinson-Foulds metric and the more robust and sensitive Matching Cluster metric, are frequently used to detect phylosymbiosis (23,42,51,52,59,62). Robinson-Foulds analyzes the distance between two trees as the smallest number of operations required to convert one topology to the other (63), while Matching Cluster considers congruency at the subtree level and is therefore a more refined evaluation of small topological changes that affect incongruence (64). We strongly recommend the use of both metrics. Statistical significance (p-values) is typically evaluated by determining the probability of 100,000 randomized bifurcating dendrogram topologies yielding equivalent or more congruent phylosymbiotic patterns than the microbiome dendrogram (23); normalized Robinson–Foulds and Matching Cluster scores can be calculated as the number of differences between the two topologies divided by the total possible congruency scores for the two trees, with normalized distances ranging from 0 (complete congruence) to 1 (complete incongruence) (23).

Matrix correlation methods identify phylosymbiosis by comparing the similarities between host-derived and microbial-derived distance matrices. Methods implemented in phylosymbiosis studies (20,21,39,50,65-71) include the Mantel test, which statistically evaluates the linear correlation between all corresponding elements from two independent matrices by permutation (72), and the more powerful Procrustean superimposition approach, which rotates and fits two matrices to minimize their differences association. Partial Mantel tests (73) measuring correlations between two matrices while controlling for the effects of a third variable described in another matrix are



also used to evaluate associations between microbial communities and multiple aspects of host characteristics, such as phylogeny, identity, genetic distances, and geographic distances (39,66,67,69).

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Although both topology-based and matrix-based tests are specific and sensitive enough to detect phylosymbiosis in a variety of empirical cases, there are several differences between them. Topological comparison metrics do not use branch length information as there is no a priori reason to assume rates of host evolution in each lineage should equal rates of ecological community change in the microbiome. Indeed, rates of microbiome change may be expected to be far more rapid than gradual evolution of host genetic changes. As such, tests of topology without relative branch lengths are conservative relative to matrix correlation methods that directly rely on comparisons of host genetic divergence with microbial community dissimilarity. A simulation analysis suggested that the Mantel test has higher sensitivity and power than the Robinson-Foulds metric when phylosymbiosis is based on the assumption of microbial preferences for a host trait (19). The practical relevance of this conclusion is not clear because phylosymbiosis will arise from reasons other than microbial colonization preferences, such as host preferences, neutral processes, and microbe-microbe interactions. Moreover, the performance between Mantel test and the more sensitive topology-based Matching Cluster distance was not evaluated in this simulation, and such comparisons are likely to yield different insights. Systematic benchmarking of type I and II error rates of phylosymbiosis measurement methods across various possible scenarios will aid experimental design and result interpretation. As such, research opportunities for the development and implementation of improved phylosymbiosis detection methods are ample.

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Parameter selection. Phylosymbiosis detection involves selection of various parameters, such as OTU identity cutoff, beta diversity metric, clustering method, and congruency test, each with their strengths and limitations that will vary with study design and questions. Although various parameter combinations can be tested and compared simultaneously (39), in the case when only a few of all possible parameter combinations detect phylosymbiosis, we recommend cautious interpretation of results with respect to the chosen parameters. If available, results should also be compared to those from previous phylosymbiosis studies with similar sample types using the same parameter combinations. Experimental replication is also necessary to confirm phylosymbiosis, especially when it is not consistently detected.

Phylogenetic comparative methods. The effects of phylogenetic signal, defined as "a tendency for related species to resemble each other more than they resemble species drawn at random from the tree" (74), on univariate traits (e.g., microbial alpha diversity) have been examined in parallel with phylosymbiosis studies (66,67). Phylogenetic signal indices like Pagel's λ (75), and Blomberg's K (76) are based on a random Brownian model of trait evolution (77), but can also be used with and compared to more complex models that take into account natural selection. Although these methods are less commonly used on multivariable data and have not yet been applied to evaluate phylosymbiosis explicitly, they are promising alternatives for not only examining host phylogenetic signal on microbial beta diversity, but also testing evolutionary models relevant to phylosymbiosis.

Phylogenetic comparative methods, such as phylogenetic independent contrasts (77) and phylogenetic generalized linear mixed models (pGLMMs) (78), predict the evolutionary



correlation between two or more discrete or continuous traits given a known phylogeny and an evolutionary model. These can also be integrated into phylosymbiosis studies. pGLMMs were recently implemented in coral microbiome (20) and passerine feather microbiome studies (71) to examine the effects of latitude and colony size on coral alpha diversity, cophylogenetic coral-bacteria relationships, and relationships between alpha diversity and relative abundances of bacteriocin-producing bacteria and keratinolytic feather damaging bacteria. These methods can be useful in predicting ecological interactions, such as predator-prey relationships, mutualism, competition, and habitat filtering, as well as environmental interactions, all of which can affect microbial community structure and therefore phylosymbiosis.

Overall, as meta-omics and trait evolution analyses become more widely applicable to phylosymbiosis, one compelling direction of future phylosymbiosis investigations *in silico* is to venture beyond host phylogenetic effects on microbial diversity to resolve linkages between host phylogeny, host functions, microbial diversity, microbial functions, selective forces, and environmental factors.

5. THE PREVALENCE OF PHYLOSYMBIOSIS

A major goal of microbiome science is to find general paradigms and rules, if any, that are comparable across varied systems. In this light, phylosymbiosis is emerging as a bona fide trend because of its frequent recurrence across insect, animal, and plant systems. (**Figure 2**). Phylosymbiosis in insects include viromes of *Nasonia* parasitoid jewel wasps (42) and gut microbiomes of cockroaches, termites (79), lab-reared (23) and wild mosquitoes (59), *Cephalotes* turtle ants (39), and *Apis* social corbiculate bees (69). In *Drosophila* flies, phylosymbiosis patterns are either weakly supported (23) or not detected (80) in lab strains and wild populations.

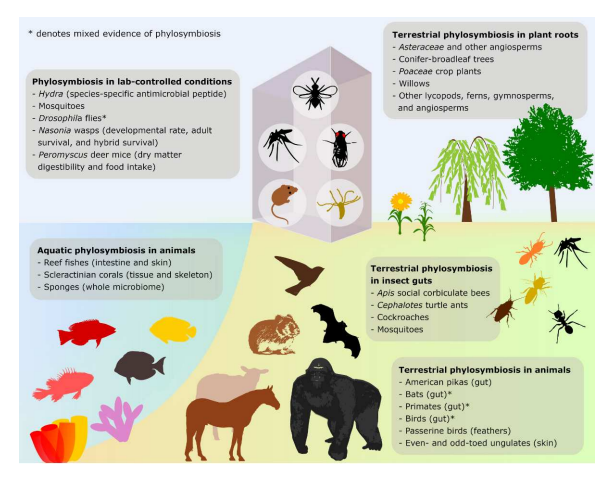


Figure 2. Representative diversity of phylosymbiosis across host species, tissues, habitats, and functions. The * symbol denotes taxa with mixed evidence for phylosymbiosis.

The first phylosymbiosis study on mammalian gut microbiomes (4) demonstrated effects of animal phylogeny and diet on gut microbial community dissimilarity (12,21,23,39,70,81). Studies focusing on gut microbiomes of specific animal groups detected phylosymbiosis in American pikas (51) and *Peromyscus* deer mice (23,52), no phylosymbiosis in western chipmunks (82), and mixed evidence of phylosymbiosis in primates (17,43,44,70), bats (55,83), and birds (62,68,84,85). Besides gut or fecal microbiomes, animal surface microbiomes have also been analyzed for phylosymbiotic associations (86), which for example occur on mammalian skin (53) and passerine feathers (71), but not on amphibian skin (3). A meta-analysis of phylosymbiosis



literature highlighted an increased prevalence of the trend in microbiomes inhabiting internal host compartments in relation to those inhabiting external host compartments (18). However, the finding may be inherently biased due to the larger number of studies investigating phylosymbiosis in the gut in relation to other external host compartments.

Beyond terrestrial and associated habitats, research interest in phylosymbiotic associations in aquatic habitats is steadily growing (**Figure 2**), spanning global sponge microbiome surveys (67,87,88) and taxon-specific sponge surveys (50,65,66) with mixed results. Two previous studies in sponges showed that the host phylogenetic signal on microbial beta diversity was reduced but still significant when host phylogeny is examined given host identity (66,67). In Australian scleractinian corals, phylosymbiosis was generally observed in tissue and skeleton compartments, but not mucus specimens that are predominantly influenced by the environment (20), suggesting different anatomical impacts on the pattern. Phylosymbiosis and host dietary impacts also occur on the skin microbiomes of 44 fish species from the Western Indian Ocean (89), but do not exist on the surface microbiomes of sympatric kelp species (90).

Phylosymbiosis has been assessed in plants, mainly to distinguish the effects of host phylogeny and soil determinants on microbial beta diversity. A comparative analysis of lycopods, ferns, gymnosperms, and angiosperms across a coastal tropical soil chronosequence indicated host phylogeny is a secondary but statistically significant factor shaping root-associated bacterial community structure, after soil age (15). More taxonomically- and/or spatially-restricted surveys have also revealed phylosymbiosis between rhizobacterial communities and *Poaceae* crop plants (91), endosphere bacterial communities and 30 plant species (92), rhizosphere-associated fungal



communities and willows from hydrocarbon-contaminated soils (93), root-associated eumycotan fungal communities and *Asteraceae* flowering plants in a dry grassland (94), ectomycorrhizal fungal communities and conifer-broadleaf forest trees (95), and ectomycorrhizal fungal communities and Estonian Salicaceae willows (96). Contrarily, qualitative incongruency between Brassicaceae host phylogeny and their root microbiomes has been observed (97), whereas non-statistically significant phylosymbiotic correlations have been reported in other plant microbiome studies (16,98).

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6. SIGNIFICANCE AND FUTURE DIRECTIONS OF PHYLOSYMBIOSIS

Microbiome research will continue to be revolutionized by the multi-omics era, where a deluge of data has enabled unprecedented insights into the extensive taxonomic, genetic, and functional composition of microbial communities and their associated hosts. Such large-scale accumulation of empirical and theoretical findings can potentiate the development of new hypotheses, unifying concepts, and frameworks across diverse host-microbiome systems. Indeed, the recurrence of phylosymbiosis across host systems lends itself to large comparative surveys across kingdoms of life that may uncover taxonomic range restrictions of phylosymbiosis as well as the environmental parameters (e.g., soil and water properties) and ecological interactions (e.g., diet and predator-prey relationships) that determine the boundaries of where and when phylosymbiosis occurs. If the microbiome field will have general trends to test in new systems, phylosymbiosis is well-poised for this circumstance. Phylosymbiosis distinguishes itself from non-phylosymbiosis by characterizing a significant degree of association between host phylogenetic and microbiome community relationships. It is not universal, and thus provides a testable hypothesis, reflects the variation likely to be seen in nature, and is amenable to explanation by mechanisms that require further investigation. The



determination of whether phylosymbiosis is present or not is a first step preceding further investigations into mechanistic details, such as the nature of species-species associations and the type(s) of ecological and evolutionary genetic processes underpinning phylosymbiosis. Given the growing evidence for the pattern and increasingly sophisticated tools available to detect phylosymbiosis, phylosymbiosis is relatively clearer and more specific than other terms such as dysbiosis.

Phylosymbiosis also engenders a holistic view of ecology and evolution in which hosts are communities or holobionts whose microbial members can contribute to genetic and phenotypic variation subject to natural selection. Several questions that have been conventionally overlooked include what are the microbial effects on host allele frequencies? Does host gene flow in natural populations impact microbiome variation and phylosymbiosis? Does phylosymbiosis accelerate or decelerate host speciation? What are the genetic and mechanistic factors that regulate phylosymbiosis, and how do these factors vary across populations or species? Collectively, studies determining the magnitude of ecological, evolutionary, and genetic forces in structuring phylosymbiosis is an important area of future research.

CONCLUSIONS

Phylosymbiosis defines a link between host evolutionary relationships and microbial diversity that is quantifiable and applicable across living systems. As research in this area proliferates, a definition, conceptual framework, and workflow for assessing phylosymbiosis will facilitate identification of phylosymbiotic host-microbe interactions. Future cause-and-effect studies of phylosymbiosis will bring a mechanistic understanding of the evolutionary, genetic, and molecular bases. Just as no mature theory of evolutionary genetics was possible until we understood the mode



436	of inheritance, no mature principle of evolutionary ecology for host-associated microbiomes seems
437	possible until we understand the general mechanisms establishing host-microbiome associations.
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445 REFERENCES

- 446 (1) McFall-Ngai M, Hadfield MG, Bosch TC, Carey HV, Domazet-Loso T, Douglas AE, et al.
- 447 Animals in a bacterial world, a new imperative for the life sciences. Proc Natl Acad Sci U S A
- 448 2013 Feb 26;110(9):3229-3236.
- 449 (2) Theis KR, Dheilly NM, Klassen JL, Brucker RM, Baines JF, Bosch TCG, et al. Getting the
- 450 hologenome concept right: an eco-evolutionary framework for hosts and their microbiomes.
- 451 mSystems 2016;1(2):e00028-16.
- 452 (3) Bletz MC, Archer H, Harris RN, McKenzie VJ, Rabemananjara FCE, Rakotoarison A, et al.
- Host ecology rather than host phylogeny drives amphibian skin microbial community structure in
- 454 the biodiversity hotspot of Madagascar. Front Microbiol 2017 Aug 17;8:1530.
- 455 (4) Ley RE, Hamady M, Lozupone C, Turnbaugh PJ, Ramey RR, Bircher JS, et al. Evolution of
- 456 mammals and their gut microbes. Science 2008 Jun 20;320(5883):1647-1651.
- 457 (5) Colman DR, Toolson EC, Takacs-Vesbach C. Do diet and taxonomy influence insect gut
- 458 bacterial communities? Mol Ecol 2012 10/01; 2019/04;21(20):5124-5137.
- 459 (6) Franzenburg S, Walter J, Kunzel S, Wang J, Baines JF, Bosch TCG, et al. Distinct
- antimicrobial peptide expression determines host species-specific bacterial associations. Proc
- 461 Natl Acad Sci USA 2013 09/24;110(39):E3730.
- 462 (7) Bost A, Martinson VG, Franzenburg S, Adair KL, Albasi A, Wells MT, et al. Functional
- variation in the gut microbiome of wild *Drosophila* populations. Mol Ecol 2018 Jul;27(13):2834-
- 464 2845.
- 465 (8) Funkhouser LJ, Bordenstein SR. Mom knows best: the universality of maternal microbial
- 466 transmission. PLoS Biol 2013;11(8):e1001631.
- 467 (9) Brucker RM, Bordenstein SR. The hologenomic basis of speciation: gut bacteria cause hybrid
- 468 lethality in the genus *Nasonia*. Science 2013 Aug 9;341(6146):667-669.
- 469 (10) Bouffaud M, Kyselkova M, Gouesnard B, Grundmann G, Muller D, Moenne-Loccoz Y. Is
- diversification history of maize influencing selection of soil bacteria by roots? Mol Ecol
- 471 2012;21(1):195-206.
- 472 (11) Brucker RM, Bordenstein SR. The roles of host evolutionary relationships (genus: *Nasonia*)
- and development in structuring microbial communities. Evolution 2012;66(2):349-362.
- 474 (12) Ochman H, Worobey M, Kuo CH, Ndjango JB, Peeters M, Hahn BH, et al. Evolutionary
- 475 relationships of wild hominids recapitulated by gut microbial communities. PLoS Biol 2010 Nov
- 476 16;8(11):e1000546.



- 477 (13) Werren JH, Richards S, Desjardins CA, Niehuis O, Gadau J, Colbourne JK, et al. Functional
- and evolutionary insights from the genomes of three parasitoid *Nasonia* species. Science 2010
- 479 Jan 15;327(5963):343-348.
- 480 (14) Moeller AH, Suzuki TA, Lin D, Lacey EA, Wasser SK, Nachman MW. Dispersal limitation
- promotes the diversification of the mammalian gut microbiota. Proc Natl Acad Sci U S A 2017
- 482 Dec 26;114(52):13768-13773.
- 483 (15) Yeoh YK, Dennis PG, Paungfoo-Lonhienne C, Weber L, Brackin R, Ragan MA, et al.
- Evolutionary conservation of a core root microbiome across plant phyla along a tropical soil
- 485 chronosequence. Nat Commun 2017 Aug 9;8(1):215-017-00262-8.
- 486 (16) Erlandson S, Wei X, Savage J, Cavender-Bares J, Peay K. Soil abiotic variables are more
- 487 important than Salicaceae phylogeny or habitat specialization in determining soil microbial
- 488 community structure. Mol Ecol 2018 Apr;27(8):2007-2024.
- 489 (17) Grieneisen LE, Charpentier MJ, Alberts SC, Ran B, Gideon B, Jenny T, et al. Genes,
- 490 geology and germs: gut microbiota across a primate hybrid zone are explained by site soil
- 491 properties, not host species. Proc R Soc B 2019;286.
- 492 (18) Mazel F, Davis KM, Loudon A, Kwong WK, Groussin M, Parfrey LW. Is host filtering the
- main driver of phylosymbiosis across the tree of life? mSystems 2018 Oct 23;3(5):e00097-18.
- 494 (19) Fraune S, Bosch TCG. Long-term maintenance of species-specific bacterial microbiota in
- 495 the basal metazoan *Hydra*. Proc Natl Acad Sci USA 2007 08/07;104(32):13146-13151.
- 496 (20) Pollock FJ, McMinds R, Smith S, Bourne DG, Willis BL, Medina M, et al. Coral-associated
- bacteria demonstrate phylosymbiosis and cophylogeny. Nat Commun 2018 11/22;9(1):4921.
- 498 (21) Groussin M, Mazel F, Sanders JG, Smillie CS, Lavergne S, Thuiller W, et al. Unraveling
- 499 the processes shaping mammalian gut microbiomes over evolutionary time. Nat Commun 2017
- 500 Feb 23;8:14319.
- 501 (22) Rudman SM, Greenblum S, Hughes RC, Rajpurohit S, Kiratli O, Lowder DB, et al.
- Microbiome composition shapes rapid genomic adaptation of *Drosophila melanogaster*. Proc
- Natl Acad Sci U S A 2019.
- 504 (23) Brooks AW, Kohl KD, Brucker RM, van Opstal EJ, Bordenstein SR. Phylosymbiosis:
- relationships and functional effects of microbial communities across host evolutionary history.
- 506 PLoS Biol 2016 Nov 18;14(11):e2000225.
- 507 (24) van Opstal EJ, Bordenstein SR. Phylosymbiosis impacts adaptive traits in *Nasonia* wasps.
- 508 MBio 2019 Jul 16(4):10:e00887-19.



- 509 (25) Parker ES, Dury GJ, Moczek AP. Transgenerational developmental effects of species-
- specific, maternally transmitted microbiota in *Onthophagus* dung beetles. Ecol Entomol
- 511 2019;44(2):274-282.
- 512 (26) Wang J, Kalyan S, Steck N, Turner LM, Harr B, Kunzel S, et al. Analysis of intestinal
- 513 microbiota in hybrid house mice reveals evolutionary divergence in a vertebrate hologenome.
- 514 Nat Commun 2015 Mar 4;6:6440.
- 515 (27) Sevellec M, Laporte M, Bernatchez A, Derome N, Bernatchez L. Evidence for host effect
- on the intestinal microbiota of whitefish (*Coregonus* sp.) species pairs and their hybrids. Ecol
- 517 Evol 2019;0(0):1-13.
- 518 (28) Cohen J. Statistical power analysis for the behavioral sciences. 2nd ed. Hillsdale, NJ:
- 519 Lawrence Erlbaum Associates; 1988.
- 520 (29) Debelius J, Song SJ, Vazquez-Baeza Y, Xu ZZ, Gonzalez A, Knight R. Tiny microbes,
- enormous impacts: what matters in gut microbiome studies? Genome Biol 2016 Oct
- 522 19;17(1):217.
- 523 (30) Conesa A, Madrigal P, Tarazona S, Gomez-Cabrero D, Cervera A, McPherson A, et al. A
- survey of best practices for RNA-seq data analysis. Genome Biol 2016 Jan 26;17:13.
- 525 (31) Felsenstein J. Confidence limits on phylogenies: an approach using the bootstrap. Evolution
- 526 1985;39(4):783-791.
- 527 (32) Anisimova M, Gascuel O. Approximate likelihood-ratio test for branches: a fast, accurate,
- and powerful alternative. Syst Biol 2006 Aug;55(4):539-552.
- 529 (33) Guindon S, Dufayard JF, Lefort V, Anisimova M, Hordijk W, Gascuel O. New algorithms
- and methods to estimate maximum-likelihood phylogenies: assessing the performance of PhyML
- 531 3.0. Syst Biol 2010 May;59(3):307-321.
- 532 (34) Mau B, Newton MA. Phylogenetic inference for binary data on dendograms using Markov
- 533 Chain Monte Carlo. J Comput Graph Stat 1997 03/01;6(1):122-131.
- 534 (35) Wiley EO, Lieberman BS. Phylogenetics: the theory and practice of phylogenetics, second
- edition. Hoboken, New Jersey: John Wiley & Sons, Inc.; 2012.
- 536 (36) Sokal RR, Rohlf FJ. The comparison of dendrograms by objective methods. Taxon
- 537 1962;11(2):33-40.
- 538 (37) Rideout JR, He Y, Navas-Molina JA, Walters WA, Ursell LK, Gibbons SM, et al.
- 539 Subsampled open-reference clustering creates consistent, comprehensive OTU definitions and
- scales to billions of sequences. PeerJ 2014 Aug 21;2:e545.



- 541 (38) Kopylova E, Navas-Molina JA, Mercier C, Xu ZZ, Mahé F, He Y, et al. Open-source
- sequence clustering methods improve the state of the art. mSystems 2016 American Society for
- 543 Microbiology;1(1):e00003-15.
- 544 (39) Sanders JG, Powell S, Kronauer DJC, Vasconcelos HL, Frederickson ME, Pierce NE.
- 545 Stability and phylogenetic correlation in gut microbiota: lessons from ants and apes. Mol Ecol
- 546 2014 03/01; 2019/04;23(6):1268-1283.
- 547 (40) Callahan BJ, McMurdie PJ, Holmes SP. Exact sequence variants should replace operational
- taxonomic units in marker-gene data analysis. ISME J 2017 Dec;11(12):2639-2643.
- 549 (41) Medina M, Sachs JL. Symbiont genomics, our new tangled bank. Genomics 2010 March
- 550 2010;95(3):129-137.
- 551 (42) Leigh BA, Bordenstein SR, Brooks AW, Mikaelyan A, Bordenstein SR. Finer-scale
- phylosymbiosis: insights from insect viromes. mSystems 2018 Dec 18;3(6):e00131-18.
- 553 (43) Amato KR, G Sanders J, Song SJ, Nute M, Metcalf JL, Thompson LR, et al. Evolutionary
- trends in host physiology outweigh dietary niche in structuring primate gut microbiomes. ISME J
- 555 2019 Mar;13(3):576-587.
- 556 (44) Amato KR, Mallott EK, McDonald D, Dominy NJ, Goldberg T, Lambert JE, et al.
- 557 Convergence of human and Old World monkey gut microbiomes demonstrates the importance of
- human ecology over phylogeny. Genome Biol 2019 Oct 8;20(1):201.
- 559 (45) Dice LR. Measures of the amount of ecologic association between species. Ecology 1945
- 560 07/01; 2019/04;26(3):297-302.
- 561 (46) Sørensen TJ. A method of establishing groups of equal amplitude in plant sociology based
- on similarity of species content and its application to analyses of the vegetation on Danish
- commons. Kongelige Danske Videnskabernes Selskab 1948;5:1-34.
- 564 (47) Bray JR, Curtis JT. An ordination of the upland forest communities of southern Wisconsin.
- 565 Ecol Monogr 1957;27(4):326-349.
- 566 (48) Motyka J, Dobrzanski B, Zawadzki S. Wstepne badania nad lagami potudniowowschodniej
- Lubelszezyzny (Preliminary studies on meadows in the southeast of the province Lublin). Univ
- 568 Mariae Curie-SktodowskaA lln Seet E 1950;5(13):367-447.
- 569 (49) Lozupone C, Knight R. UniFrac: a new phylogenetic method for comparing microbial
- communities. Appl Environ Microbiol 2005 Dec;71(12):8228-8235.
- 571 (50) Reveillaud J, Maignien L, Murat Eren A, Huber JA, Apprill A, Sogin ML, et al. Host-
- specificity among abundant and rare taxa in the sponge microbiome. ISME J 2014
- 573 Jun;8(6):1198-1209.



- 574 (51) Kohl KD, Varner J, Wilkening JL, Dearing MD. Gut microbial communities of American
- 575 pikas (*Ochotona princeps*): Evidence for phylosymbiosis and adaptations to novel diets. J Anim
- 576 Ecol 2018 Mar;87(2):323-330.
- 577 (52) Kohl KD, Dearing MD, Bordenstein SR. Microbial communities exhibit host species
- 578 distinguishability and phylosymbiosis along the length of the gastrointestinal tract. Mol Ecol
- 579 2018 Apr;27(8):1874-1883.
- 580 (53) Ross AA, Muller KM, Weese JS, Neufeld JD. Comprehensive skin microbiome analysis
- reveals the uniqueness of human skin and evidence for phylosymbiosis within the class
- 582 Mammalia. Proc Natl Acad Sci U S A 2018 Jun 19;115(25):E5786-E5795.
- 583 (54) Goodrich JK, Di Rienzi SC, Poole AC, Koren O, Walters WA, Caporaso JG, et al.
- 584 Conducting a microbiome study. Cell 2014 Jul 17;158(2):250-262.
- 585 (55) Phillips CD, Phelan G, Dowd SE, McDonough MM, Ferguson AW, Delton Hanson J, et al.
- Microbiome analysis among bats describes influences of host phylogeny, life history, physiology
- and geography. Mol Ecol 2012 Jun;21(11):2617-2627.
- 588 (56) Legendre P, Legendre L. Numerical Ecology. Amsterdam: Elsevier Science B.V.; 1998.
- 589 (57) Clarke KR. Non-parametric multivariate analyses of changes in community structure. Aust J
- 590 Ecol 1993;18(1):117-143.
- 591 (58) McArdle BH, Anderson MJ. Fitting multivariate models to community data: a comment on
- 592 distance-based redundancy analysis. Ecology 2001 01/01; 2019/04;82(1):290-297.
- 593 (59) Novakova E, Woodhams DC, Rodriguez-Ruano SM, Brucker RM, Leff JW, Maharaj A, et
- al. Mosquito microbiome dynamics, a background for prevalence and seasonality of West Nile
- 595 Virus. Front Microbiol 2017 Apr 4;8:526.
- 596 (60) Michener CD, Sokal RR. A quantitative approach to a problem in classification. Evolution
- 597 1957;11(2):130-162.
- 598 (61) Matsen IV FA, Evans SN. Edge principal components and squash clustering: using the
- 599 special structure of phylogenetic placement data for sample comparison. PLoS One
- 600 2013;8(3):e56859.
- 601 (62) Laviad-Shitrit S, Izhaki I, Lalzar M, Halpern M. Comparative analysis of intestine
- microbiota of four wild waterbird species. Front Microbiol 2019 Aug 20;10:1911.
- 603 (63) Robinson DF, Foulds LR. Comparison of phylogenetic trees. Math Biosci 1981 February
- 604 1981;53(1):131-147.
- 605 (64) Bogdanowicz D, Giaro K. On a matching distance between rooted phylogenetic trees. Int J
- 606 Appl Math Comput Sci 2013;23(3):669-684.



- 607 (65) Schottner S, Hoffmann F, Cardenas P, Rapp HT, Boetius A, Ramette A. Relationships
- between host phylogeny, host type and bacterial community diversity in cold-water coral reef
- 609 sponges. PLoS One 2013;8(2):e55505.
- 610 (66) Easson CG, Thacker RW. Phylogenetic signal in the community structure of host-specific
- microbiomes of tropical marine sponges. Front Microbiol 2014 Oct 17;5:532.
- 612 (67) Thomas T, Moitinho-Silva L, Lurgi M, Bjork JR, Easson C, Astudillo-Garcia C, et al.
- Diversity, structure and convergent evolution of the global sponge microbiome. Nat Commun
- 614 2016 Jun 16;7:11870.
- 615 (68) Kropáčková L, Těšický M, Albrecht T, Kubovčiak J, Čížková D, Tomášek O, et al.
- 616 Codiversification of gastrointestinal microbiota and phylogeny in passerines is not explained by
- 617 ecological divergence. Mol Ecol 2017;26(19):5292-5304.
- 618 (69) Kwong WK, Medina LA, Koch H, Sing KW, Soh EJY, Ascher JS, et al. Dynamic
- microbiome evolution in social bees. Sci Adv 2017 Mar 29;3(3):e1600513.
- 620 (70) Gaulke CA, Arnold HK, Humphreys IR, Kembel SW, O'Dwyer JP, Sharpton TJ.
- 621 Ecophylogenetics clarifies the evolutionary association between mammals and their gut
- 622 microbiota. MBio 2018 Sep 11;9(5):e01348-18.
- 623 (71) Javurkova VG, Kreisinger J, Prochazka P, Pozgayova M, Sevcikova K, Brlik V, et al.
- Unveiled feather microcosm: feather microbiota of passerine birds is closely associated with host
- 625 species identity and bacteriocin-producing bacteria. ISME J 2019 May 24.
- 626 (72) Mantel N. The detection of disease clustering and a generalized regression approach. Cancer
- 627 Res 1967 Feb;27(2):209-220.
- 628 (73) Smouse PE, Long JC, Sokal RR. Multiple regression and correlation extensions of the
- 629 Mantel test of matrix correspondence. sysbio 1986 12/01; 6/11;35(4):627-632.
- 630 (74) Blomberg SP, Garland Jr T. Tempo and mode in evolution: phylogenetic inertia, adaptation
- and comparative methods. J Evol Biol 2002;15(6):899-910.
- 632 (75) Pagel M. Inferring the historical patterns of biological evolution. Nature 1999
- 633 10/01;401(6756):877-884.
- 634 (76) Blomberg SP, Garland JR. T, Ives AR. Testing for phylogenetic signal in comparative date:
- behavioral traits are more labile. Evolution 2003;57(4):717-745.
- 636 (77) Felsenstein J. Phylogenies and the comparative method. Am Nat 1985;125(1):1-15.
- 637 (78) Ives AR, Helmus MR. Generalized linear mixed models for phylogenetic analyses of
- 638 community structure. Ecol Monogr 2011;81(3):511-525.



- 639 (79) Dietrich C, Kohler T, Brune A. The cockroach origin of the termite gut microbiota: patterns
- in bacterial community structure reflect major evolutionary events. Appl Environ Microbiol 2014
- 641 Apr;80(7):2261-2269.
- 642 (80) Wong AC, Chaston JM, Douglas AE. The inconstant gut microbiota of *Drosophila* species
- 643 revealed by 16S rRNA gene analysis. ISME J 2013 Oct;7(10):1922-1932.
- 644 (81) Moeller AH, Caro-Quintero A, Mjungu D, Georgiev AV, Lonsdorf EV, Muller MN, et al.
- Cospeciation of gut microbiota with hominids. Science 2016 Jul 22;353(6297):380-382.
- 646 (82) Grond K, Bell KC, Demboski JR, Santos M, Sullivan JM, Hird SM. No evidence for
- phylosymbiosis in western chipmunk species. FEMS Microbiol Ecol 2019 Nov 15:fiz182.
- 648 (83) Lutz HL, Jackson EW, Webala PW, Babyesiza WS, Kerbis Peterhans JC, Demos TC, et al.
- 649 Ecology and host identity outweigh evolutionary history in shaping the bat microbiome.
- 650 mSystems 2019 Nov 12;4(6):e00511-19.
- 651 (84) Hird SM, Sanchez C, Carstens BC, Brumfield RT. Comparative gut microbiota of 59
- neotropical bird species. Front Microbiol 2015;6:1403.
- 653 (85) Liu H, Chen Z, Gao G, Sun C, Li Y, Zhu Y. Characterization and comparison of gut
- microbiomes in nine species of parrots in captivity. Symbiosis 2019;78:241.
- 655 (86) Ross AA, Rodrigues Hoffmann A, Neufeld JD. The skin microbiome of vertebrates.
- 656 Microbiome 2019 May 23;7(1):79-019-0694-6.
- 657 (87) Schmitt S, Tsai P, Bell J, Fromont J, Ilan M, Lindquist N, et al. Assessing the complex
- sponge microbiota: core, variable and species-specific bacterial communities in marine sponges.
- 659 ISME J 2012 Mar;6(3):564-576.
- 660 (88) Lurgi M, Thomas T, Wemheuer B, Webster NS, Montoya JM. Modularity and predicted
- functions of the global sponge-microbiome network. Nat Commun 2019 Mar 1;10(1):992-019-
- 662 08925-4.
- 663 (89) Chiarello M, Auguet JC, Bettarel Y, Bouvier C, Claverie T, Graham NAJ, et al. Skin
- microbiome of coral reef fish is highly variable and driven by host phylogeny and diet.
- 665 Microbiome 2018 Aug 24;6(1):147-018-0530-4.
- 666 (90) Lemay MA, Martone PT, Keeling PJ, Burt JM, Krumhansl KA, Sanders RD, et al.
- Sympatric kelp species share a large portion of their surface bacterial communities. Environ
- 668 Microbiol 2018;20(2):658-670.
- 669 (91) Bouffaud M, Poirier M, Muller D, Moënne-Loccoz Y. Root microbiome relates to plant host
- evolution in maize and other Poaceae. Environ Microbiol 2014;16(9):2804-2814.



- 671 (92) Fitzpatrick CR, Copeland J, Wang PW, Guttman DS, Kotanen PM, Johnson MTJ. Assembly
- and ecological function of the root microbiome across angiosperm plant species. Proc Natl Acad
- 673 Sci U S A 2018 Feb 6;115(6):E1157-E1165.
- 674 (93) Bell TH, El-Din Hassan S, Lauron-Moreau A, Al-Otaibi F, Hijri M, Yergeau E, et al.
- 675 Linkage between bacterial and fungal rhizosphere communities in hydrocarbon-contaminated
- soils is related to plant phylogeny. ISME J 2014 Feb;8(2):331-343.
- 677 (94) Wehner J, Powell JR, Muller LAH, Caruso T, Veresoglou SD, Hempel S, et al.
- Determinants of root-associated fungal communities within Asteraceae in a semi-arid grassland.
- 679 J Ecol 2014 03/01; 2019/05;102(2):425-436.
- 680 (95) Ishida TA, Nara K, Hogetsu T. Host effects on ectomycorrhizal fungal communities: insight
- from eight host species in mixed conifer-broadleaf forests. New Phytol 2007;174(2):430-440.
- 682 (96) Tedersoo L, Mett M, Ishida TA, Bahram M. Phylogenetic relationships among host plants
- explain differences in fungal species richness and community composition in ectomycorrhizal
- 684 symbiosis. New Phytol 2013;199(3):822-831.
- 685 (97) Schlaeppi K, Dombrowski N, Oter RG, Ver Loren van Themaat E, Schulze-Lefert P.
- Ouantitative divergence of the bacterial root microbiota in *Arabidopsis thaliana* relatives. Proc
- 687 Natl Acad Sci U S A 2014 Jan 14;111(2):585-592.
- 688 (98) Vincent JB, Weiblen GD, May G. Host associations and beta diversity of fungal endophyte
- communities in New Guinea rainforest trees. Mol Ecol 2016 Feb;25(3):825-841.

690