



The Utility of Macroecological Rules for Microbial Biogeography

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Dickey JR, Swenie RA, Turner SC, Winfrey CC, Yaffar D, Padukone A, Beals KK, Sheldon KS and Kivlin SN (2021) The Utility of Macroecological Rules for Microbial Biogeography. Front. Ecol. Evol. 9:633155. doi: 10.3389/fevo.2021.633155 Macroecological rules have been developed for plants and animals that describe large-scale distributional patterns and attempt to explain the underlying physiological and ecological processes behind them. Similarly, microorganisms exhibit patterns in relative abundance, distribution, diversity, and traits across space and time, yet it remains unclear the extent to which microorganisms follow macroecological rules initially developed for macroorganisms. Additionally, the usefulness of these rules as a null hypothesis when surveying microorganisms has yet to be fully evaluated. With rapid advancements in sequencing technology, we have seen a recent increase in microbial studies that utilize macroecological frameworks. Here, we review and synthesize these macroecological microbial studies with two main objectives: (1) to determine to what extent macroecological rules explain the distribution of host-associated and free-living microorganisms, and (2) to understand which environmental factors and stochastic processes may explain these patterns among microbial clades (archaea, bacteria, fungi, and protists) and habitats (host-associated and free living; terrestrial and aquatic). Overall, 78% of microbial macroecology studies focused on free living, aquatic organisms. In addition, most studies examined macroecological rules at the community level with only 35% of studies surveying organismal patterns across space. At the community level microorganisms often tracked patterns of macroorganisms for island biogeography (74% confirm) but rarely followed Latitudinal Diversity Gradients (LDGs) of macroorganisms (only 32% confirm). However, when microorganisms and macroorganisms shared the same macroecological patterns, underlying environmental drivers (e.g., temperature) were the same. Because we found a lack of studies for many microbial groups and habitats, we conclude our review by outlining several outstanding questions and creating recommendations for future studies in microbial ecology.

Keywords: microorganisms, Rapoport's rule, Abundance/Occupancy, Bergmann's rule, Theory of Island Biogeography, Species-Area Relationship, Latitudinal Diversity Gradient, macroecology

INTRODUCTION

Understanding the factors that determine the abundance, distribution, and diversity of organisms across spatial and temporal scales is a fundamental challenge in ecology. For plants and animals, many macroecological rules have been theorized to explain the physiological, ecological, and some evolutionary processes that underlie these patterns. Microorganisms also exhibit patterns in

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abundance, distribution, and diversity over space and time (Martiny et al., 2006, 2011; Hanson et al., 2012; Talbot et al., 2014; Kivlin, 2020). However, it is unclear if macroecological rules developed for plants and animals apply to microorganisms and if they could be used to improve predictions for abundance, distribution, and diversity of microorganisms (Prosser et al., 2007; Soininen, 2012; Shade et al., 2018; Kivlin et al., 2020). Recent review papers have highlighted how macroecological frameworks can be leveraged to investigate biogeographic patterns in microorganisms (Shoemaker et al., 2017; Xu et al., 2020). Yet, to our knowledge, no comprehensive literature search has defined when microbial biogeographic patterns conform to those expected by various macroecological rules or the mechanisms driving microbial biogeographic patterns. Defining which macroecological rules are universal across the tree of life will allow for ecosystem-wide development of theory of common mechanisms affecting community assembly, response to global change, and recovery from human-induced disturbance.

Some macroecological rules should apply across the spectrum of life. For example, those that describe patterns of abundance or biodiversity of plants and animals (e.g., latitudinal/elevational clines in biodiversity, species-area relationships, speciesabundance distributions, and island biogeographic patterns) should easily translate to microorganisms since these rules share a common theoretical underpinning of applying the drivers of local, regional and continental community assembly to larger scales. Community assembly of all organisms, including microscopic ones, is driven by deterministic and stochastic processes (Vellend, 2010; Nemergut et al., 2013). Deterministic processes include environmental selection or filtering, and biotic interactions such as competition and facilitation (Funk et al., 2008; Goldford et al., 2018). Stochastic processes that affect community assembly include dispersal limitation, neutral processes of ecological and evolutionary drift (Hubbell, 2001; Martiny et al., 2011), priority effects of colonization (Fukami et al., 2010), legacy effects of previous environmental conditions (Hawkes et al., 2017), and historical vicariance of geographic position of land masses (Matheny et al., 2009) or suitable habitat (Takacs-Vesbach et al., 2008). Finally, selection regimes over long time periods can influence community assembly through evolution (Kraft et al., 2007; Starnawski et al., 2017). Because these explanatory drivers of community assembly apply regardless of the focal organism, community-level macroecological patterns driven by these processes should hold across the tree of life (Horner-Devine et al., 2004; Shade et al., 2018).

In contrast, other organismal and functional macroecological rules, such as the Metabolic Theory of Ecology (MTE; Brown et al., 2004), Species-Energy Theory (Wright, 1983), Bergmann's Rule (Bergmann, 1847), Foster's Rule (Foster, 1964), and Rapoport's Rule (Stevens, 1989) may lack direct analogs to microbial consortia because the mechanisms posited to underlie these rules relate to traits (e.g., body size, thermoregulation, or homeostasis) or energy sources (e.g., thermodynamics) that may not exist or apply to microorganisms or that are difficult to measure. This is particularly true of macroecological rules that assume sexual dimorphism of two sexes (Rensch's Rule; Rensch, 1950), which apply well to many animals but break down in organisms with simpler (e.g., asexual bacteria) or much more complex mating systems (e.g., fungal species with hundreds of mating types). Moreover, macroecological rules based on body size lack direct analogs for most microorganisms where body size could refer to cell size, colony size, or sporocarp size and therefore difficult to systematically investigate. Even if a consensus could be reached, DNA-based inference of microbial abundance is unlikely to represent organismal size when ribosomal copy number varies 100-fold among microbial taxa and lacks clear phylogenetic signal (Lofgren et al., 2019).

In addition to whether or not macroecological rules apply to microorganisms, microbial groups may also vary in their ability to conform to macroecological rules of plants and animals due to differences in habitat or due to difference in microbial groups (e.g., prokaryotes vs. multicellular eukaryotes). For example, dispersal limitation of soil-borne and plant-associated microorganisms, in general, should be much higher compared to air-borne, water-borne, or animal-associated microorganisms that are actively transported as animals explore surrounding landscapes. Dispersal limitation should also be greater for microbes with a multicellular life stage than those microbes that remain unicellular throughout their whole life. This could limit range sizes of terrestrial and plant-associated microorganisms relative to aquatic or animal-associated microorganisms, with the exception of isolated, island-like aquatic systems such as lakes or other small bodies of water. In addition, climate fluctuations of terrestrial ecosystems may be more pronounced than those in aquatic ecosystems, which could affect organism range sizes (Sorte et al., 2013). Finally, biotic interactions may be more coupled in confined terrestrial spaces (e.g., soil pores) compared to large or more well mixed aquatic and marine ecosystems.

Perhaps one of the most crucial distinctions among microbial lifestyles is that of free-living versus host-associated microorganisms (Figure 1). Some microorganisms are freeliving in soil or aquatic habitats. Distributions of these taxa may be mostly affected by dispersal limitation and subsequent environmental filtering, where traits such as dormancy (Lennon and Jones, 2011) or reduced metabolic rates (Wisnoski et al., 2020) allow some microorganisms to cope with unfavorable environmental conditions. However, a large portion of microbial diversity lives in symbiosis inside (endosymbiotic) or on (episymbiotic) animal and plant tissues. For many vertically transmitted host-associated microorganisms in animals and some plants, dispersal of the microorganism often depends on dispersal of the host (Salerno et al., 2016; Shade et al., 2017), eventually leading to co-cladogenesis in each group (Takiya et al., 2006; Schardl et al., 2008) and a strong signal of biotic filtering on the microbiome among host taxa. This is especially true when host-associated microbes are more buffered from environmental conditions compared to freeliving microorganisms. Multiple lines of evidence suggest that host-associated microorganisms (especially vertically transmitted microorganisms) often have narrower thermal tolerances (Dunbar et al., 2007; Kikuchi et al., 2016) and reduced genome sizes and functions (Moran et al., 2008). However, for other plant-associated microorganisms (e.g.,



mycorrhizal fungi), independent dispersal of plants and microorganisms allows dispersal limitation and environmental filtering (Hazard et al., 2013) to influence host-associated microbial communities.

In this review, we first introduce seven macroecological rules and discuss cases where they may apply to microorganisms. We then conduct a literature meta-analysis to synthesize information available on macroecological trends of free-living and host-associated microorganisms from terrestrial and aquatic ecosystems to understand the extent to which microorganisms follow the same largescale macroecological patterns as animals and plants. The objectives of this review are to (1) determine to what extent macroecological rules explain the distribution of host-associated and free-living microorganisms, and (2) understand which environmental factors most affect the distributions of host-associated versus free-living microorganisms among microbial clades and habitats. We focus on four macroecological rules at the organismal level (Bergmann's Rule, Rapoport's Rule, Abundance/Occupancy Relationships, and Gloger's Rule), three rules that describe/explain community assembly and biodiversity [Latitudinal Diversity Gradient (LDG), Theory of Island Biogeography, and Species-Area Relationships], and one functional rule (MTE).

For each rule we collected studies by querying Web of Science or Google Scholar with the search terms "rule name" AND "micro*" or "fung*" or "bacteri*" or "protist*" or "archaea*" (query completed 06.15.2020). These queries returned studies on fungi, bacteria, archaea, and protists (including diatoms). Because many studies report trends consistent with these rules without studying them directly, we also surveyed all papers that were cited by or cited the papers in our query. We did not include the term "macroecology" because many papers including this term do not include primary data or did not survey a given macroecological rule of interest. We also did not include papers that collected appropriate data but did not conduct statistics to analyze a macroecological pattern as raw data availability varies and therefore was beyond the purview of this synthesis. We did not include human-associated microorganisms or green algae in our survey. Human travel, diet, and medicinal applications can affect microbiomes in complicated ways and therefore human microbiomes were not deemed as natural communities for the purposes of this review. Overall, this search resulted in 175 studies with 218 records (macroecological patterns for a given microbial group within a study) across all continents. For each study, we classified which macroecological rule(s) were surveyed and each study organism (archaea, bacteria, fungi, or protist), habitat (terrestrial or aquatic/marine), and niche (free-living or host-associated). We considered speciesarea relationships only for contiguous habitats and included species-area relationships among island-like habitats (e.g., lakes, tree holes, plant rhizosphere, and animals) with other island biogeography studies. When available, we classified which deterministic (e.g., resources and climate) and stochastic (e.g., dispersal limitation, priority effects) drivers were correlated with each observed macroecological pattern. Despite collecting 218 records for this synthesis, data limitation precluded a quantitative meta-analytical approach to addressing both adherence of microorganisms to macroecological rules and the environmental drivers influencing these distributions. Instead, we use vote counting (Hedges and Olkin, 1980) to define which macroecological rules were supported among microbial guilds and lifestyles and which environmental factors may have influenced these distributional patterns. Each time a study record confirmed a macroecological rule, it received one positive vote, and each time a study record provided contradictory evidence for a given macroecological rule, it received one negative vote. While vote counting techniques are sensitive to sample size within studies, this analysis clearly demonstrates both when microbial distributions conform to macroecological rules and more importantly when gaps in scientific inquiry do not allow us to make this conclusion.

REVIEW OF MACROECOLOGICAL RULES

Organismal Macroecological Rules Bergmann's Rule

Bergmann's rule, which states that body size decreases among species with decreasing temperature, has primarily been studied in mammals and birds. Not all birds and mammals follow the rule, and even fewer smaller organisms, such as insects, conform to it (Blackburn et al., 1999a; Meiri and Dayan, 2003). A possible clue for ectotherms may be that Bergmann's rule partially agrees with the temperature-size rule, which states that ectotherms develop more quickly with smaller body sizes under higher temperatures (Zuo et al., 2012); a species' body size is a result of temperature dependent energy consumption and reallocation which can vary over elevation and across latitudes.

There is some evidence supporting Bergmann's rule for microorganisms, such as bacterioplankton (Meador et al., 2009), with smaller cells at higher temperatures (Daufresne et al., 2009). The temperature-size relationship has also been examined as a possible mechanism behind a Bergmann's rule pattern in microorganisms, stating that as temperature increases in aquatic systems and nucleic acid content is reduced, and thus microbes have smaller cells (Huete-Stauffer et al., 2016). Temperature-size relationships have also been studied with ubiquitous microbes, such as diatoms in geothermal streams, whereby there is no observed relationship of this relationship (Adams et al., 2013). While some evidence of Bergmann's rule exists, significant challenges in defining the size of many microorganisms, such as multicellular fungal hyphae, must be standardized before rigorous comparisons can be drawn.

Rapoport's Rule

Rapoport's rule states that the latitudinal ranges of plants and animals are greater at higher compared to lower latitudes (Stevens, 1989). Stevens (1989) originally proposed a mechanism underlying this rule that relates to the climatic conditions under which different organisms have evolved. Specifically, species at higher latitudes must be able to cope with greater temporal variability in climate compared to species at lower latitudes. As a result, species that have evolved at higher latitudes should be able to occupy larger spatial extents than species at lower latitudes. The rule was later extended to elevational gradients with the expectation that at higher elevations, climate is more variable and thus high elevation species should have broader elevational ranges (Stevens, 1992). Other possible mechanisms for the rule have since been proposed, such as decreasing land area available to species at lower latitudes, reduced competition at higher latitudes, or extinction of species with narrow ranges at high latitudes due to glaciation (Gaston et al., 1998). Rapoport's rule tends to be weaker in marine compared to terrestrial systems and in the Southern compared to Northern Hemisphere (Ruggiero and Werenkraut, 2007). The rule has received such mixed support that some researchers have questioned whether it is "time for an epitaph" for Rapoport's rule (Gaston et al., 1998; Gaston and Chown, 1999). Despite this, ecologists are still intrigued by this macroecological rule and continue to examine whether it holds in a host of taxa, including microorganisms.

Thus far, support for Rapoport's rule across latitude and elevation in microorganisms is equivocal at best. In terrestrial systems, the latitudinal ranges of fungal taxa (Tedersoo et al., 2014; Cox et al., 2016) and ciliates appear to increase toward the poles [W. Foissner, pers. comm. to Azovsky and Mazei (2013)]. By contrast, several studies have observed smaller range sizes of bacterial taxa at higher elevations and latitudes (Okie et al., 2015; Lear et al., 2017; Wu et al., 2018). In marine systems, a global study of bacteria found strong support for Rapoport's rule, with bacteria in the tropics having smaller ranges than bacteria in temperate regions (Amend et al., 2013). The relationship was so strong that the most abundant bacteria in the tropics were completely absent from higher latitudes, but even narrowly distributed bacteria in temperate regions were found at lower latitudes (Amend et al., 2013). A second study of marine bacteria also found that bacteria showed narrower ranges at lower compared to higher latitudes, but this was only true for bacteria in the phyla Bacteroidetes and Cyanobacteria and the classes α -, β -, and γ - of the phylum Proteobacteria (Sul et al., 2013). Researchers have also found a reverse Rapoport's rule in marine benthic ciliates. One study found that ciliates had narrower ranges at higher latitudes (Azovsky and Mazei, 2013). Support is also mixed for an elevational Rapoport's rule across microbial taxa. In a study across fungal phyla (Basidiomycota, Ascomycota, Zygomycota, Chytridiomycota, and Glomeromycota), fungi at higher elevations had greater distributional ranges (Ogwu et al., 2019). In contrast, bacteria and diatoms in streams in Asia and Europe did not follow an elevational Rapoport's rule (Teittinen et al., 2016; Wang and Soininen, 2017).

Researchers have identified several possible mechanisms leading to the distributions of microorganisms across latitudinal and elevational gradients. Range sizes of some taxa have been associated with variation in temperature and precipitation (Lear et al., 2017), suggesting that environmental selection determines range extents. However, there is some evidence that suggests dispersal limitation may be the primary driver of range sizes for some groups of microorganisms at larger spatial scales (Sul et al., 2013), while environmental selection, competition, and niche differentiation becoming more important factors at smaller spatial scales (Kivlin et al., 2014).

Abundance/Occupancy Relationships

The abundance–occupancy (A/O) relationship refers to the prevalent pattern among macroorganisms, whereby more abundant species occupy more locations within a given range (Gaston et al., 2000), both within and among species. Most A/O relationships report positive correlations but fail to investigate the potential reasons why this relationship forms. There are a handful of empirical studies that show weakly positive, or even negative trends amongst plants (Boeken and Shachak, 1998), fish (Marshall and Frank, 1994), and birds (Blackburn et al., 1999b). In addition, evidence of positive A/O relationships have been found for protists (Warren and Gaston, 1997), bacteria (Holt et al., 2004), and freshwater diatoms (Heino and Soininen, 2006; Spatharis et al., 2009; Soininen and Teittinen, 2019).

In general, the mechanisms resulting in the A/O pattern are not well understood. The A/O relationship can occur when extinction and colonization are density-dependent (Warren and Gaston, 1997). In experimental microcosms with constant resources, when dispersal was allowed, the relationship tracked the carrying capacity hypothesis (i.e., locally abundant species have lower extinction and higher colonization rates than species with low abundance; Holt et al., 2002). However, this study found that the relationship was not contingent on dispersal; without dispersal, the A/O relationship arose through extinction (Holt et al., 2002). Moreover, the A/O relationship does not always depend on environmental resource consistency. For example, the relationship held in heterogeneous environments, though to a lesser degree than in homogeneous environments (Holt et al., 2004). In field-based observations, the A/O relationship has been observed with soil fungal assemblages where dispersal had less of a role than environmental filtering on species distributions (Kivlin et al., 2014). Mechanisms behind this pattern were attributed to soil nutrients, though the relationship was weak (Kivlin et al., 2014).

Niche-based processes can also create A/O relationships among organisms. In microcosm experiments with protists, biotic interactions strengthened A/O relationships (Holt et al., 2002). Likewise, in natural systems Rocha et al. (2018) found that niche-based variables were the primary predictors for the positive A/O relationship observed in freshwater diatoms and that the effect of niche breadth on the relationship was less important than local environmental niche position. In animal host-associated intestinal bacteria, Green et al. (2016) found support for the A/O relationship that they attributed to multiple factors, including dispersal limitation, host selection, historical contingency, and stochastic processes.

Gloger's Rule

Gloger (1833) observed a relationship between plumage coloration and climate variation in birds, predicting that darker coloration due to pigmentation occurred in warmer, more humid tropical regions while lighter colors occurred in cooler areas toward the poles. There is limited evidence for Gloger's rule in organisms other than animals. However, pigments such as melanin are also an important protective mechanism for many microorganisms, since darker pigments can provide protection from ionizing radiation and desiccation (Dadachova and Casadevall, 2008; Fernandez and Koide, 2013). Pigmentation may also play a role in thermoregulation and energy exchange by increasing or decreasing light absorption or reflectance (Cordero et al., 2018). Consequently, pigmentation could potentially provide an important adaptive function for microbes.

However, rather than following Gloger's rule, evidence from microbial distributions thus far supports the theory of thermal melanism, whereby greater melanism increases solar radiation absorption (e.g., Kettlewell, 1973; Brakefield, 1984). This has been shown both observationally and experimentally in yeast where pigmentation increases heat capture. For example, in contrast to Gloger's rule, free-living, dark-pigmented yeasts were found more often outside of the tropics (Cordero et al., 2018). This suggests pigmentation provides an adaptive advantage for these microorganisms at higher latitudes by increasing heat capture. Similarly, a survey of ~3,000 European macrofungal assemblages showed mushrooms were more darkly colored in colder climates (Krah et al., 2019). Mean temperature and in some cases, seasonality were identified as drivers of mushroom coloration, and increased reproductive success was suggested as a potential advantage of darker pigmentation in colder climates. While suggestive that Gloger's rule may not apply to microorganisms, evidence thus far is mainly derived

from free-living fungi and has not been examined in other microbial groups.

Community Level Macroecological Rules Latitudinal Diversity Gradient

The LDG, whereby biodiversity increases from the poles to the equator, is a common large-scale pattern in ecology (Jablonski et al., 2006). Many invertebrate and vertebrate species exhibit this gradient, as do vascular plants and some fungi and marine bacteria (Hillebrand, 2004). However, many macroorganisms do not exhibit a LDG (Gillman et al., 2015), owing to legacies of environmental disturbance, dependence on environmental factors that do not trend with latitude, or dependence on biotic interactions with other organisms whose diversity also does not scale linearly with latitude. However there is more observed variation amongst studies that investigate species richness and latitude patterns for microorganisms than there is for macroorganisms (Thompson et al., 2017).

Early reports on microbial diversity suggested that some fungi and marine bacteria may follow the classic LDG pattern much as plants and animals do (Hillebrand, 2004). Further studies confirmed this pattern in stream diatoms, marine bacterioplankton, marine tintinnids, Streptomyces bacteria, freshwater fungi, and most fungal taxonomic groups (Fuhrman et al., 2008; Tedersoo et al., 2014; Andam et al., 2016; Hyde et al., 2016). The main drivers of this pattern were mean annual precipitation, temperature, carbon availability, and substrate pH, all of which can affect microbial activity, energy acquisition and longevity. However, microbes also tend to follow a hump-shaped diversity pattern with latitude within a hemisphere such that the greatest diversity is found at intermediate latitudes. This trend was demonstrated in planktonic bacteria (Milici et al., 2016), ectomycorrhizal fungi (Tedersoo et al., 2014), aquatic hyphomycete fungi (Jabiol et al., 2013; Duarte et al., 2016), saprotrophic stream fungi (Seena et al., 2019), and other fungal guilds (Větrovský et al., 2019). Potential drivers of this type of distribution include sea water temperature, thermoclines, day length, and phosphorus levels for aquatic species, dispersal limitation in terrestrial microbes, and plant host-related diversity and distribution in ectomycorrhizal fungi.

Other patterns of diversity have also been found across latitudes. A study of stream diatoms found that richness showed a U-shaped pattern. However, the study ranged only from subtropical to temperate latitudes in the northern hemisphere (Passy, 2010). Thus, a different pattern might have been observed had a larger latitudinal gradient been examined. In addition, for both the hump-shaped and U-shaped diversity distributions, sampling bias toward temperate areas may have influenced these results. Overall, a majority of studies found no pattern in microbial diversity across latitude. These include microbes in the Pacific Ocean (Baldwin et al., 2005), marine free-living ciliates (Azovsky and Mazei, 2013), freshwater diatoms (Soininen and Teittinen, 2019), heterotrophic bacterioplankton (Schiaffino et al., 2013), autotrophic eukaryotes (Schiaffino et al., 2016), soil bacteria (Fierer and Jackson, 2006; Hendershot et al., 2017), and soil fungi (Hendershot et al., 2017). Instead, soil pH was an important driver of distributions for terrestrial microorganisms in these studies, but localized climatic factors, nutrient availability, dispersal limitations, and scale of study may also play a role. Species richness of some ecto- and endoparasites also does not vary with latitude, since they are insulated from environmental conditions by their hosts (Rohde, 1978, 1999).

Theory of Island Biogeography

The dynamic equilibrium Theory of island Biogeography is an influential framework for understanding species diversity patterns. Its original proponents (MacArthur and Wilson, 1967) predicted two general patterns: (1) larger islands have more species at equilibrium, since extinction rates decline and colonization rates increase with increased area and (2) islands at a greater distance from the "mainland" or source population have fewer species at equilibrium, as colonization rates decrease with distance. Traditionally applied to groups of oceanic islands, the theory has recently been used to understand microbial biogeography across many scales on patchy, islandlike systems. These include roots (Peay et al., 2007), leaves (Andrews et al., 1987; Kinkel et al., 1987), lakes or pools (Reche et al., 2005), and non-human animal organs (Loudon et al., 2016; Moeller et al., 2017). The theory has also been applied to other discrete habitats that are separated by sharp differences in temperature (Whitaker et al., 2003; Darcy et al., 2018), substrate textural or chemical qualities such as oxygen availability or host-secreted metabolites (Loudon et al., 2016), or other abiotic factors that have been shown to structure microbial distributions.

Results from several studies support the general relationship between microbial diversity and island size, although there is no consensus on how habitat size is measured amongst varying "island-like" systems (Itescu, 2019). For example, Darcy et al. (2018) observed that bacterial diversity within cryoconite holes in Antarctic glaciers is positively correlated with hole area, whereas Bell et al. (2005) found bacterial genetic diversity in water-filled tree holes increased with water volume. Studies have also found fungal species richness increases with tree host population size in arctic and alpine environments (Chlebicki and Olejniczak, 2007) and as a function of tree photosynthetic tissue volume for ectomycorrhizal fungi (Peay et al., 2007; Glassman et al., 2017a). Yet, some studies fail to detect a significant relationship between species diversity and island area, such as AM fungal communities on true islands, bacterial richness in oak litter patches and filamentous fungi on apple leaves (Kinkel et al., 1987; Davison et al., 2018; Spiesman et al., 2018). In some cases, the lack of an observed relationship is likely because environmental heterogeneity relevant for focal taxa does not scale with island size (Teittinen and Soininen, 2015). The application of the Theory of Island Biogeography across many scales (e.g., tree holes to oceanic islands) for microorganisms reflects the multitude of scales at which habitat heterogeneity and dispersal affect microbial distributions. Yet, how microbial distributions on island-like habitats differ across these scales warrants further attention.

The mechanisms that produce microbial diversity patterns across islands of various sizes are still debated. Recent studies have found these patterns hold even after explicitly accounting for environmental variables. For example, forest fragment size predicted diversity of root-associated fungi in a dominant tree even after accounting for the effects of soil carbon (Vannette et al., 2016). Dinnage et al. (2019) determined that tree size explained more variation in rhizobia diversity than other edaphic factors. Various host-associated microbial clades have differing responses to ecological drift, host size, and other environmental factors (Li et al., 2020). For example, bacterial alpha diversity increased with island area, which may be explained by differences in overall immigration and extinction rates (Dinnage et al., 2019). However, fungal beta diversity increased with area, which instead suggests an effect of greater microenvironmental variation in larger patch sizes (Chlebicki and Olejniczak, 2007). Further research into mechanisms underlying diversity-area patterns may clarify such differences across taxa, although studying processes such as local extinction in microbial communities may not be straightforward, particularly in natural systems (but see Andrews et al., 1987).

Empirical evidence that microbial diversity and community similarity decline as a function of distance from the source population is mixed across the literature, reflecting taxon-specific patterns as well as the need for more research. A significant challenge in applying island biogeographic theory to microbial communities in island-like systems is that the microorganism source, or "mainland" population, is often difficult to define, and thus most studies instead consider only distances among islands. Interestingly, most of the studies that assess the effects of distance from the mainland have used the roots of isolated or planted pine trees as "islands" and a denser, nearby pine forest as the "mainland" (Peay et al., 2010, 2012; Bahram et al., 2013; Glassman et al., 2017a,b). For example, Bahram et al. (2013) and Glassman et al. (2017a) both found that ectomycorrhizal fungal community similarity and diversity declined with increasing distance from the natural forest's edge. The majority of studies examining how distance among "islands" shapes community similarity and diversity find spatial autocorrelation. For instance, communities of hot spring-dwelling archaea (Whitaker et al., 2003) and protists in lakes (Lepère et al., 2013) become more dissimilar as a function of distance, rather than variation in environmental factors. This suggests that for these organisms, the difficulties of dispersing among patches of suitable habitat are more important for community structure than environmental filtering.

Exceptions to the predictions of island biogeographic theory reveal that the relative importance of environmental filtering and dispersal limitation is taxon- and scale-specific. For example, Vannette et al. (2016) determined that the degree of forest fragment isolation was important in structuring the communities of some classes of root-associated fungi but not others. In other cases, lack of support may instead be due to the spatial scale considered. Darcy et al. (2018) observed positive spatial autocorrelation of cryoconite glacier hole bacterial communities among glaciers, but very little within glaciers, suggesting that had they sampled at a smaller scale, they may not have found evidence in line with island biogeographic theory. The importance of scale is also taxon-specific because taxa differ in their dispersing and colonizing abilities (Peay et al., 2012), likely explaining why Teittinen and Soininen (2015) did not find spatial structuring of diatom communities in springs. Overall, more manipulative experiments to assess why and how taxa differ in their dispersal capacities would help assess the relevance of an island-distance relationship for microorganisms.

Species-Area Relationships

The positive relationship between species richness (S) and the surveyed area (A) is one of the clearest, least disputed macroecological patterns and is fundamental to understanding the inherent heterogeneity of global biodiversity (Arrhenius, 1921). The power-law $S = cA^z$ has been used to compare the relationship between species number and area, where c is the intercept and the exponent z measures the rate of community turnover across a standardized spatial scale. This phenomenon has been explored across myriad taxa, including birds (Jetz and Rahbek, 2001), mammals (Lomolino, 1982), plants (Honnay et al., 1999), terrestrial microbial eukaryotes (Green et al., 2004), soil bacteria (Horner-Devine et al., 2004), aquatic bacteria (Reche et al., 2005; Martiny et al., 2011), ectomycorrhizal fungi (Peay et al., 2007), and vertebrate gut bacteria (Godon et al., 2016). Yet, the power-law often fails to predict species richness for many organisms (McCoy and Connor, 1976; Connor and McCoy, 1979) indicating that this relationship may only apply to a subset of spatial scales (i.e., local or regional), habitat types, or taxonomic groups.

Among microbial studies that investigate species-area or taxaarea relationships, microbial diversity often increases with spatial scale but to a much lower extent (slope z = 0.02-0.1) than observed for macroorganisms [slope z = 0.3-0.6, see Green and Bohannan (2006) and references therein]. The observed lower slopes in microorganisms in comparison to macroorganisms may be due to inconsistency in taxonomic comparisons made with direct observation versus DNA sequencing. For instance, 97% clustering is a fairly coarse clustering strategy for microorganisms that may obscure patterns of species turnover (Storch and Šizling, 2008). Spatial scaling of microbial diversity may vary from that of macroorganisms for many reasons including the higher diversity of microorganisms versus macroorganisms, loose species definitions for microorganisms, dormancy of microbial taxa (Jones and Lennon, 2010), or extracellular DNA confounding species richness estimates (Carini et al., 2016).

In addition to methodological concerns, many variables other than spatial extent affect the distribution of free-living and host-associated microorganisms. For example, microbes associated with marshland habitat respond to both geographic distance and sediment moisture (Horner-Devine et al., 2004). While species-area relationships are beginning to be explored across microbial systems, many confounding factors exist when comparing studies, such as differences in spatial extent or environmental heterogeneity. Illustrating this point is the finding that the rate of community turnover depends on whether microbial species richness is measured within or across habitats (Martiny et al., 2011). In addition, determining patterns of species richness over space is even more intractable for host-associated microbial taxa, such as ectomycorrhizal fungi, where host effects and dispersal limitation might conflate the relative importance of other environmental factors in prediction of species richness (Tanesaka, 2012). Nevertheless, despite inconsistencies in the strength of the relationship of species richness and area, taxa within most microbial guilds and lifestyles increase in diversity as more space is sampled.

Functional Macroecological Rules Metabolic Theory of Ecology

The MTE predicts that the metabolism of individual organisms increases with environmental temperature, often with a 3/4th scaling to body size (Brown et al., 2004). Corollary predictions suggest that metabolic rates of all organisms are a combination of the allometric scaling of their body mass and biochemical kinetics (West et al., 1997) which leads to a negative association between population carrying capacity and temperature and therefore a positive association between temperature and species richness (Brown et al., 2004). While supported in some plant and animal systems, considerable controversy surrounds the universality of metabolic theory. For example, many underlying assumptions of metabolic theory, such as the actual scaling of surface area to volume ratio across organisms and enzymatic activation energies, remain untested or unproven across the tree of life (Duncan et al., 2007; Price et al., 2012).

In microbial systems, this theory has been extended to extracellular enzyme activities (Elias et al., 2014), which are assumed to reflect intracellular metabolic rates. Many studies across environmental temperature gradients find positive relationships between temperature and soil enzyme activities and respiration (Xu et al., 2017) as well as between temperature and microbial diversity (Okie et al., 2015; Zhou et al., 2016; Wu et al., 2018). Similarly, meta-analyses confirm that metabolic efficiency (i.e., the quantity of resources incorporated into biomass relative to uptake) scales with body size among microorganisms in terrestrial and aquatic ecosystems (Sinsabaugh et al., 2015). However, metabolic scaling in microorganisms rarely follows the 3/4th power to body size (DeLong et al., 2010) and studies are often confounded by other abiotic and biotic factors that also shift with temperature, such as variation in water, nutrient availability, plant diversity, or productivity (Zhou et al., 2016). Furthermore, thermal acclimation (Bradford et al., 2008) and adaptation (Alster et al., 2020) can alter microbial temperature responses in less than a decade. Thus, while evidence in support of metabolic scaling of microbial processes seems confirmatory, the rule should be examined more carefully across gradients that decouple temperature from other environmental drivers over evolutionary time. In addition, cell size is not linearly related to metabolic rates in marine phytoplankton communities (Marañón, 2015), suggesting that the relationship between organismal size and metabolism may apply to aggregates of microorganisms but break down at the smallest, cellular scales.

LITERATURE SYNTHESIS RESULTS

Across all macroecological rules the majority of macroecological microbial studies (78%) have been conducted on free-living microorganisms, mainly in aquatic ecosystems (**Table 1** and **Figure 2**). Citations corresponding with **Table 1** can be found in Supplemental Material 1. In contrast, few studies have examined similar patterns in host-associated microorganisms with the notable exceptions of studies of latitudinal diversity and island biogeographic patterns focused on plant-associated fungi (e.g., mycorrhizal fungi and endophytes; **Figure 2**). Furthermore, the bulk of these studies examined community-level trends in microbial macroecology (71%), with less than 35% of studies occurring at the organismal level (**Figure 2**). Within community-based macroecological rules, LDGs were the more commonly studied, followed by the Theory of Island Biogeography and then species-area relationships.

Conformance to Macroecological Rules

Sixty-three percent of the studies we examined in this synthesis confirmed similar patterns of microbial and macro-organismal distributions. In particular, organismal-level microorganism patterns supported macroecological rules the most (Bergmann's Rule - 75% confirm; Rapoport's Rule - 85% confirm; and Area-Occupancy - 100% confirm). Notably, Gloger's Rule of increasing pigmentation at lower latitudes was reversed for fungi in the only two published studies, which suggested that melanization was associated with thermal adaptation to colder, more polar latitudes. However, community-level patterns were not as consistent among taxa. For example, patterns of microbial diversity did not consistently decrease from the equator to the poles as expected from macroorganisms (32% confirm). Instead, microorganisms exhibited reverse, humpshaped, and indistinct diversity patterns with latitude (Figure 3). In contrast, rules predicted by the dynamic equilibrium Theory of Island Biogeography, were largely upheld (74% confirm) across many systems (e.g., terrestrial host-associated mycorrhizal fungi, aquatic bacteria and protists) with increasing diversity on larger islands (but see notable exceptions, e.g., leaf endophytic fungi; Figure 3). Similarly, diversity was often negatively correlated with island distance from the source population, especially for planthost associated microorganisms. All descriptions of microbial species-area relationships noted increased species diversity as more habitat area was sampled. Finally, for functionalbased macroecological rules, evidence for metabolic scaling of microbial function confirmed expected positive temperaturedependence of microbial activity in 93% of studies but was examined in only 14 studies and never in host-associated microbial communities.

Drivers of Macroecological Rules

When macroecological rules were upheld in microorganisms, factors driving these patterns were mostly similar to those hypothesized for the macroorganisms studied to date (**Table 1**). For example, patterns consistent with the rules predicted by the dynamic equilibrium Theory of Island Biogeography were often correlated with island patch size and dispersal, whereas

LDGs and metabolic scaling followed temperature gradients (but see Hawkins and Diniz-filho, 2004 for a discussion of confounding effects of temperature and latitude). However, there were notable exceptions. For example, we expected substrate pH to be a strong determining factor for microbial richness patterns due to its pivotal role in explaining beta diversity at large spatial scales. Instead, pH was only related to speciesarea relationships and island size in island biogeography (Fierer and Jackson, 2006). Other environmental factors related to climate and resources (carbon, nutrients) often were correlated with many microbial macroecological patterns. In addition, dispersal limitation and patch size were among the main correlates of microbial distributions, regardless of habitat or host-associated versus free-living status. While data were scarce, the distributions of host-associated microorganisms tended to be structured more heavily by biotic factors (e.g., host identity, microbial interactions) than were free-living taxa, as expected in Figure 1. Similarly, when comparisons were possible (e.g., LDG, Species Area Relationship, and Theory of Island Biogeography) there were no clear differences in drivers of terrestrial and aquatic microorganisms or among microbial kingdoms (Table 1). Clearly more investigation into several macroecological rules such as Gloger's Rule and Bergmann's Rule is warranted, as is more in-depth inquiry of the processes driving macroecological patterns, since over 30 studies (14%) found support for microbial congruence with macroecological rules but did not test or describe a process underlying these patterns.

DISCUSSION

Many factors should be considered when one compares these microorganismal trends to those of macroorganisms. Many microbial studies did not explicitly test a macroecological rule and thus were not searchable in Web of Science or Google Scholar. In addition, null results are less likely to be published. These factors could have biased our findings toward rule confirmation rather than exception. Biologically, patterns of microbial distributions obtained from sequencing may sample a different subset of organisms than those derived from observational studies of plants and animals. DNA-based sampling encompasses both dormant and active states, which may increase estimates of microbial diversity as compared to plants and animals. An analog would be to sample all plant and animal gametes, which clearly is not feasible. Microorganisms that undergo horizontal gene transfer or have higher DNA substitution rates may also evolve faster than macroorganisms, allowing greater ability to respond to microclimates and shifts in environments than plants and animals. This could be why microbial distributions seem to track local nutrients or niches more than would be predicted by theories mentioned above for macroorganisms.

Understudied Macroecological Rules in Microorganisms

Many macroecological rules are still not addressable for microorganisms. For example, validation of the species-energy

TABLE 1 | Evidence for each group of organisms is represented by kingdom for protists (P), fungi (F), bacteria (B), and archaea (A).

	Organismal (physiological)				Community			Functional
Factor	Bergmann's Rule	Rapoport's Rule	A/O	Gloger's Rule	Latitudinal Diversity Gradient	Theory of Island Biogeography	Species-Area Relationship	Metabolic Theory of Ecology
Substrate Properties								
atch Size						 B^T (Godon et al., 2016; Dinnage et al., 2019), F^T (Peay et al., 2007; Tanesaka, 2012; Vannette et al., 2016; Glassma et al., 2017a,b), P^T (Ishtiaq et al., 2010; Spurgin et al., 2012), B^A (Bell et al., 2005; Reche et al., 2005; Darcy et al., 2018), B^T (Li et al., 2020), F^T (Tanesaka, 2012; Li et al., 2020), P^A (Patrick, 1967; Lepère et al., 2018) 	F ^T (van der Gast et al., 2011), F ^A (Duarte et al., 2017)	
litrogen	P ^A (Svensson et al., 2014)		F ^T (Kivlin et al., 2014)		Ρ^τ (Lara et al., 2016)	F ^T (Vannette et al., 2016; Glassman et al., 2017a; Boeraeve et al., 2019)	B ^A (Horner-Devine et al., 2004; Martiny et al., 2011; Santos et al., 2016)	B ^A (Sinsabaugh et al., 2015; Arandia-Gorostidi et al., 2017), B ^T (Okie et al., 2015; Sinsabaugh et al., 2015), F ^A (Sinsabaugh et al., 2015) F ^T (Sinsabaugh et al., 2015)
Carbon					F ^T (Davison et al., 2015), F ^A (Hyde et al., 2016), P ^T (Lara et al., 2016)	F ^T (Vannette et al., 2016; Glassman et al., 2017a), B ^T (Delgado-Baquerizo et al., 2018)	B ^A (Horner-Devine et al., 2004)	 B^A (Sinsabaugh et al., 2015), B^T (Sinsabaugh et al., 2015), F^A (Sinsabaugh et al., 2015), F^T (Sinsabaugh et al., 2015)
'hosphorous	P ^A (Svensson et al., 2014)					F ^T (Glassman et al., 2017a), B ^T (Delgado-Baquerizo et al., 2018)	B ^A (Santos et al., 2016), P ^A (Goldenberg Vilar et al., 2014)	 B^A (Sinsabaugh et al., 2015), B^T (Sinsabaugh et al., 2015), F^A (Sinsabaugh et al., 2015), F^T (Sinsabaugh et al., 2015)
Vicronutrients			F^T (Jiao and Lu, 2020)		F ^T (Tedersoo et al., 2014)	B ^T (Dinnage et al., 2019), F ^T (Glassman et al., 2017a)	B^T (Meyer et al., 2018), P^A (Goldenberg Vilar et al., 2014)	2010)
Salinity	P ^A (Svensson et al., 2014)		B^A (Logares et al., 2013)					
Нс						B ^T (Dinnage et al., 2019), F ^T (Glassman et al., 2017a; Boeraeve et al., 2019), B ^T (Delgado-Baquerizo et al., 2018), P ^A (Soininen et al., 2007)	 B^T (Meyer et al., 2018), B^T (Fierer and Jackson, 2006; Tu et al., 2016) 	Β^T (Okie et al., 2015)
Conductivity Biotic						F^T (Glassman et al., 2017a), P ^A (Aarnio et al., 2019)		
nteractions ⊣ost					F ^T (Arnold and Lutzoni, 2007)	B ^T (Godon et al., 2016; Loudon et al., 2016), F ^T (Chlebicki and Olejniczak, 2007; Belisle et al., 2012)	 F^T (Van Der Linde et al., 2018), F^T (Feinstein and Blackwood, 2012) 	

(Continued)

TABLE 1 | Continued

Factor	Organismal (physiological)					Functional		
	Bergmann's Rule	Rapoport's Rule	A/O	Gloger's Rule	Latitudinal Diversity Gradient	Theory of Island Biogeography	Species-Area Relationship	Metabolic Theory of Ecology
Niche Position and Breadth			P ^A (Heino and Soininen, 2006; Heino and Tolonen, 2018; Rocha et al., 2018; Vilmi et al., 2019)			B ^T (Godon et al., 2016; Moeller et al., 2017)		
Microbe-Microbe Competition			B^T (Holt et al., 2004), P^T (Holt et al., 2004)		FT (Liu Setal 2020)	B^T (Loudon et al., 2016)		
Climate					- (Ed 01 of dil, 2020)			
Temperature	P ^A (Svensson et al., 2014)	B^A (Lear et al., 2017)			 F^T (Davison et al., 2015; Lu et al., 2018), B^A (Fuhrman et al., 2008; Stomp et al., 2011; Kruk et al., 2012; Sul et al., 2013; Vogt et al., 2019), B^T (Tu et al., 2016), F^A (Hyde et al., 2016), F^T (Liu S. et al., 2020), P^A (Dolan et al., 2016), P^T (Lara et al., 2016) 	F ^T (Botnen et al., 2019)	 B^A (Martiny et al., 2011), B^T (Fierer and Jackson, 2006; Tu et al., 2016; Deng et al., 2018) 	 A^T (Wu et al., 2018), B^A (Arandia-Gorostidi et al., 2017; Jiang et al., 2019), B^T (Okie et al., 2015; Zhou et al., 2016; Xu et al., 2017; Wu et al., 2018), F^T (Zhou et al., 2016; Xu et al., 2017)
Precipitation		B^A (Lear et al., 2017)			F ^T (Tedersoo et al., 2014)	B ^T (Dinnage et al., 2019; Li et al., 2020), F ^T (Botnen et al., 2019)	 B^A (Martiny et al., 2011), B^T (Fierer and Jackson, 2006; Tu et al., 2016) 	
Stochastic Processes								
Dispersal Limitation		B^A (Sul et al., 2013)	B ^A (Mo et al., 2018), B ^T (Holt et al., 2004), F ^T (Kivlin et al., 2014), P ^T (Holt et al., 2004)		B ^A (Sul et al., 2013), B ^T (Andam et al., 2016)	 B^T (Loudon et al., 2016; Moeller et al., 2017), F^T (Peay et al., 2007, 2010, 2012; Bahram et al., 2013; Vannette et al., 2016; Glassman et al., 2017a,b; Boeraeve et al., 2018), P^T (Ishtiaq et al., 2010; Spurgin et al., 2012)A^A (Whitaker et al., 2003), B^A (Darcy et al., 2018), F^T (Li et al., 2020), P^A (Soininen et al., 2007; Lepère et al., 2013; Santos et al., 2016) 	F ^T (Green et al., 2004)	
Unknown Mechanism	B ^A (Meador et al., 2009), P ^A (Soininen and Kokocinski, 2006)	 A^A (Sintes et al., 2015; Raes et al., 2018b), B^A (Amend et al., 2013; Sul et al., 2013; Wang and Soininen, 2017; Raes et al., 2018b), F^A (Raes et al., 2018b), F^T (Tedersoo et al., 2014; Cox et al., 2016; Ogwu et al., 2019) 	 B^T (Nemergut et al., 2011; Green et al., 2016; Stopnisek and Shade, 2020), A^A (Zhou et al., 2019; Liu J. et al., 2020), B^A (Pommier et al., 2007; Humbert et al., 2009; Östman et al., 2010; Nemergut et al., 2011; Liu et al., 2015; Bienhold et al., 2016; Liao et al., 2017; Mo et al., 2018; B^T (Nemergut et al., 2011; Valverde et al., 2014; Ji et al., 2020), F^T (Zhang et al., 2018; Gange et al., 2019; Dickie et al., 2020), P^T (Warren and Gaston, 1997) 		 B^A (Shu and Jiao, 2008; Thompson et al., 2011), B^T (Weiser et al., 2018), F^T (Weiser et al., 2018) 	 F^T (Helander et al., 2007; Botnen et al., 2019), B^A (Barberán and Casamayor, 2014), P^A (Macingo et al., 2019) 	F ^T (Kranabetter et al., 2018), P ^T (Ishtiaq et al., 2010), A ^A (Barreto et al., 2014; Li and Ma, 2019), A ^T (Zhou et al., 2008), B ^A (Smith et al., 2005; Barretc et al., 2014; Zinger et al., 2014), B ^T (Noguez et al., 2005; Zhou et al., 2008), F ^I (Zhou et al., 2018), F ^I (Zhou et al., 2008), P ^A (Finlay et al., 1998; Azovsky, 2002), P ^T (Wetzel et al., 2012))

(Continued)

Macroecological Rules and Microbes

TABLE 1 | Continued

Factor		Organismal (phys	siological)			Functional		
	Bergmann's Rule	Rapoport's Rule	A/O	Gloger's Rule	Latitudinal Diversity Gradient	Theory of Island Biogeography	Species-Area Relationship	Metabolic Theory of Ecology
Contradictory Evidence	P ^A (Adams et al., 2013)	P ^A (Azovsky and Mazei, 2013; Teittinen et al., 2016)		F ^T (Krah et al., 2019), F ^T (Cordero et al., 2018; Krah et al., 2019)	F ^T (Tedersoo et al., 2010, 2012, 2014; Timling et al., 2012; Holt et al., 2015; Glynou et al., 2016; Pärtel et al., 2017; de Menezes et al., 2018; Nave et al., 2017; de Menezes et al., 2018; Nave et al., 2019), P ^A (Rohde, 1978), A ^A (Sintes et al., 2019), P ^A (Rohde, 1978), A ^A (Sintes et al., 2019), A ^T (Ma et al., 2017; Weiser et al., 2018), B ^A (Baldwin et al., 2009; Ghiglione et al., 2012; Schiaffino et al., 2013, 2016; Sul et al., 2009; Ghiglione et al., 2012; Schiaffino et al., 2013, 2016; Sul et al., 2013; Milici et al., 2016; Raes et al., 2013; Milici et al., 2016; Raes et al., 2018a; Ibarbalz et al., 2016; Humbert et al., 2019; Johnson et al., 2017; Ma et al., 2016; Hendershot et al., 2017; Ma et al., 2016; Hendershot et al., 2017; Ma et al., 2018; Bickel et al., 2019; Delgado-Baquerizo and Eldridge, 2019; Hedĕnec et al., 2019; Delgado-Baquerizo and Eldridge, 2019; Hedĕnec et al., 2019; Delgado-Baquerizo and Eldridge, 2019; Hedĕnec et al., 2019; Diarbalz et al., 2016; Schiaffino et al., 2016; Raes et al., 2018; Bickel et al., 2019; Diarbalz et al., 2019; Seena et al., 2019; Hassett et al., 2019; Ma et al., 2017; Ma et al., 2017; Ma et al., 2017; Ma et al., 2017; Ma et al., 2019; Pino et al., 2019; Pino et al., 2019; Pino et al., 2019; Pino et al., 2019; Colline et al., 2016; Co	 F^T (Andrews et al., 1987; Kinkel et al., 1987; Grilli et al., 2015; Da Silva et al., 2017; Davison et al., 2018), B^A (Sommaruga and Casamayor, 2009; Logue et al., 2012; Várbíró et al., 2017), B^T (Spiesman et al., 2018), P^A (Hessen et al., 2006; Teittinen and Soininen, 2015; Bolgovics et al., 2016, 2019) 		B ^A (Marañón, 2015)

Letters are color-coded green for host-associated microbiomes and purple for free-living microbiomes. Superscript letters designate aquatic (A) or terrestrial (T) habitat. Rules examined are Bergmann's Rule, Rapoport's Rule, Abundance/Occupancy Relationships (A/O), Gloger's Rule, Latitudinal Diversity Gradient, Theory of Island Biogeography, Species-Area Relationships, and Metabolic Theory of Ecology. Rows are ecological factors that are grouped by Substrate Properties (both physical and chemical properties), Biotic Interactions, Climate, Stochastic Processes, Unknown Mechanisms, and Contradictory Evidence. A study was counted and placed under Unknown Mechanism, if the rule was upheld for microorganism but did not investigate or model any ecological factor in regard to this observed pattern. The row Contradictory Evidence includes studies where no support, or another pattern, was observed for each rule. This row helps elucidate inconsistencies or disagreements in the applicability of macroecological rules and patterns to the field of microbial ecology.



theory (Wright, 1983) is largely lacking for microorganisms that gain energy from multiple sources (e.g., photoautotrophic and chemolithotrophs). Examinations of center-periphery dynamics in which organisms are expected to have the highest abundance at the center of their range (Hengeveld and Haeck, 1982) are also lacking. Accurate range maps are not available for the majority of microorganisms (but see Kivlin, 2020), and when they are, do not contain information on microbial abundance. Other macroecological rules related to organismal size (e.g., Foster's Rule; Foster, 1964) or sexual selection (e.g., Rensch's Rule; Rensch, 1950) require detailed morphological measurements across space that are not generated from DNA-based microbial observations. As data on microbial distributions and microbial traits (e.g., Zanne et al., 2020) accumulates, many of these understudied rules may be addressable in the near future.

Outstanding Questions

Several outstanding questions and subsequent discussion points remain after our synthesis. First, if a given macroecological

rule is supported, are there important environmental factors or interactions among environmental drivers specific to microbial taxa (e.g., soil pore size) that were not captured in previous studies? If so, can these explain some of the cases where microorganisms follow macroecological patterns but where the proposed causal mechanism for macroorganisms seems inapplicable? Second, how should we conceptualize traits of microorganisms, such as body size, to compare with macroorganisms? So far, microbial body size has been measured as cell size, but extensions to sporocarp or colony size could better reflect ideas of resource allocation and use underlying some theories of macroecology (Young, 2006). Third, how do the evolutionary rates of microbial speciation and extinction affect our interpretation of macroecological rules, such as those associated with the MTE or the dynamic equilibrium Theory of Island Biogeography? For example, are biologically diverse islands hotspots for speciation or cold spots for extinction, or are patterns relating to island biogeography theory solely driven by ecological dispersal



(purple) and host-associated (green) microorganisms grouped at organismal (A), community (B), and functional (C) levels. Trends here are generated from the predominant pattern or all patterns from our synthesis when there was no consensus on vote counting. Microbial patterns match those of plants and animals for organismal (but see Gloger's Rule) and functional levels, but community-level patterns are variable, reflecting different ecological drivers of macro- and microorganism distributions.

trajectories? Finally, how does temporal turnover of microbial communities influence our conceptualization of macroecological patterns that are inferred from single time point studies? Are macroecological patterns of microorganisms consistent intraand inter-annually? Data from multi-year studies may address most of these questions and can allow assessment of ecological and evolutionary turnover in microbial communities in a variety of habitats.

Best Practices in Microbial Macroecology

To spur future studies, we have developed a list of best practices for the future of microbial macroecology. First, we recommend hypothesis-driven inquiry where researchers use macroecological rules as null hypotheses instead of surveying DNA for *de novo* patterns. To illustrate this further, one could test the null hypothesis that rhizosphere microbes have no difference in mean range size at higher elevations than lower elevations (Rapoport's Rule) given their association with their host. Second, macroecological rules should be used as a framework to connect multiple factors influencing biogeographic patterns (e.g., patch size and dispersal limitation among islands). Third, we should consider unique limitations and prospects of microbial ecology (microbial morphology and physiology, DNAbased studies, destructive sampling, horizontal gene transfer, the effect of spatial scale relative to the system, pooling across replicates within sites for DNA sequencing) and interpret findings in this light. For example, Dickey et al. (2020) shows that rhizosphere bacterial communities vary within the rhizosphere habitat and that DNA sequencing resolution affects estimates of microbial richness. When comparing macroecological rules across the tree of life, collection methods and statistics should be standardized as much as possible to avoid over-estimating microbial diversity. This could be accomplished by standardizing sampling effort among taxa through rarefaction or creating universal species concepts that encompass variable selection regimes instead of universal cutoff of 97% to define microbial taxa (e.g., Beaulieu et al., 2012). Furthermore, sampling efforts in tropical regions should be increased to counterbalance the sampling bias in temperate regions. Fourth, researchers should examine more than DNA (e.g., organismal traits, metabolomes) when surveying microorganisms to understand organismal trait-based and functional macroecological patterns. Finally, we should conduct longitudinal studies across years to understand microbial fluctuations due to ecological and evolutionary processes. The future is open for many lines of inquiry into the patterns and processes underlying microbial macroecology. The promising analogs to established macroecological rules for macroorganisms and notable exceptions we found here provide a roadmap to guide forecasts of microbial distributions under current future environmental conditions.

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AUTHOR CONTRIBUTIONS

All authors made substantial contributions to the concept and design of this study, contributed to the data acquisition for this study, and contributed in writing and revising the manuscript. JD and SK created figures and finalized revisions prior to submission.

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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GLOSSARY

Endosymbiotic: Microorganisms that colonize inter- or intracellular space of plants, animals, and fungi.

Episymbiotic: Microorganisms that live on the surface of plants, animals, and fungi.

Historical vicariance: Contemporaneously observed geographic disjunction in species ranges caused by historical severance of a contiguous species range due to separation of land masses or geological events (e.g., river gorges).

Island: An island is both a true island of land surrounded by water and island-like habitats that are spatially and temporally isolated from habitats of the same type (*sensu* Itescu, 2019).

Legacy effects: Effects of prior environmental conditions on subsequent organismal physiology, diversity, composition, and distributions after environmental conditions have changed.

Macroecological rules: Generalizable patterns of organismal distributions, abundances, and diversity supported by known processes of organismal evolution, physiology, community assembly, and/or dispersal.

Microorganisms: Here, we define microorganisms as microscopic fungi, bacteria, archaea, and protists.

Priority effects: When the order of arrival at a given habitat gives primary colonizers an advantage in using resources to be able to outcompete subsequent colonizers.