Ecosystem size and complexity dictate riverine biodiversity

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Abstract: Larger ecosystems support more species; this ubiquitous pattern is the foundation of current conservation schemes. However, many ecosystems possess a complex spatial structure that cannot be represented by area, and the role of such complexity in controlling biodiversity is largely unknown. Here, we use theory and extensive fish community data from two distinct geographic regions (Japan and United States) to show that ecosystem size and complexity dictate riverine biodiversity. We found that larger and more branched 'complex' river networks harbored greater species richness due to increased space and environmental heterogeneity. The complexity effect was comparable to the size effect, and this pattern has emerged regardless of ecological contexts. The dual control of biodiversity may be a pervasive feature that has far-reaching implications for biodiversity conservation.

7 Introduction

Ecologists have long sought to understand the general drivers of biodiversity. One of the most robust 18 empirical generalizations in ecology is the positive relationship between species richness and area, i.e., the 19 species-area relationship (the SAR) (1). In 1921, Arrhenius (2) formulated the SAR as a power-law $S = cA^z$, 20 an equation currently known as the Arrhenius species-area relationship (S) is the number of species observed in a given geographic area A, c the constant, and z the scaling exponent). Since then, the spatial scaling 22 of species richness has been observed in many taxonomic groups (3). The SAR is ubiquitous because multiple mechanisms produce an apparently similar pattern. Larger ecosystems typically support more 24 diverse metacommunities due to increased habitat diversity (4), larger metacommunity size (5), and/or enhanced colonization dynamics (6). Importantly, the SAR provides the foundation for global conservation 26 efforts (7-9). For example, conservation ecologists have used SAR estimates to design marine and terrestrial 27 protected areas (7, 8), which currently encompass more than 30 million km² globally (10).

Many ecosystems, however, possess a complex spatial structure that cannot be represented by area - a dimension referred to as scale-invariant complexity (11, 12). Such complexity is evident in branching ecosystems, including rivers, trees, and mountain ranges, to name just a few (12). Geomorphic or biological processes 31 generate a pronounced self-similarity in complex branching patterns such that the part and the whole look alike (12). Even though the branching structure is independent of spatial scale, it forms a physical template 33 that dictates habitat diversity and dispersal corridors for living organisms (13). Limited, but accumulating evidence suggests that classical metapopulation and metacommunity theories cannot predict ecological 35 dynamics driven by branching structure (14-16), and this recognition has led to recent developments of spatial theories devoted to complex branching ecosystems (17). For example, these studies have highlighted 37 key roles of branching structure in driving local biodiversity patterns, such as increased species richness at merging points of branches (18). However, most empirical research has explored the consequences of 39 branching complexity for local community structure (19) or has relied solely on theoretical arguments with limited replications of branching architecture (20). At present, we lack a comprehensive evaluation of how 41 branching complexity controls biodiversity patterns at the metacommunity level. Filling this knowledge gap may provide common ground for achieving successful conservation in spatially complex ecosystems, where accelerated species loss threatens the delivery of ecosystem services (21).

Here, we show that ecosystem size and complexity play comparable roles in controlling biodiversity patterns

in rivers - a prime example of complex branching ecosystems. Individual streams and rivers flow through different landscapes with distinct geological and climatic backgrounds, serving as a spatial unit of unique 47 in-stream environments (16, 22-25). The recurrent merging of diverse tributaries ultimately forms a fluvial 48 network with fractal branching patterns (12). As such, the complexity of branching structure, which we define here as the probability of branching per unit river distance (24, 26), may represent a macro-scale control of the 50 ecosystem's habitat heterogeneity (habitat diversity per unit area) (13, 23, 24). Meanwhile, ecosystem size (watershed area) should determine the metacommunity size and total habitat diversity (area x heterogeneity), 52 two factors that regulate biodiversity at the metacommunity level (4, 5). We predict that ecosystem size 53 and branching complexity increase watershed-scale species richness (γ diversity) by enhancing local species 54 richness (α diversity) and/or spatial difference of species composition (β diversity) under different ecological 55 scenarios. The present study combines theory and empirical analysis of extensive community data from two different regions of the globe to provide crucial insights into how ecological communities are structured in 57 complex branching networks.

59 Results and Discussion

Theoretical synthesis of ecosystem size and complexity influences

First, we developed a theoretical framework synthesizing the influences of ecosystem size and complexity on biodiversity patterns in branching ecosystems. We depicted branching ecosystems as a spatial network 62 of connected habitat patches which local communities inhabit (24, 26) (Figure 1A and B). Two factors 63 determine mean environmental conditions at each habitat patch: (1) headwater environments (the most 64 upstream habitat patch) and (2) local environmental noise. Environmental values at headwaters are drawn 65 randomly from a normal distribution and propagate downstream with local environmental noise (i.e., a spatial autoregressive process with white noise). These environmental values recurrently 'mix' at confluences 67 considering the relative size of joining tributaries (see Materials and Methods). Therefore, our networkgeneration procedure resembles natural processes of how branching river networks create diverse habitats 69 in a metacommunity. We then simulated metacommunity dynamics in the theoretical branching networks 70 using a general metacommunity model (27). In this model, 50 species with different abiotic niches (optimum 71 and width) disperse along a network and compete for resources with varied strengths in temporally dynamic 72 environments (see Materials and Methods for model details). 73

Given the results of extensive sensitivity analysis (Supplementary Text), we considered 32 simulation 74 scenarios comprising a combination of four landscape and eight ecological scenarios. We distinguished 75 landscape scenarios by setting four combinations of environmental variation at headwaters ($\sigma_h = 0.01, 1$) 76 and the degree of local environmental noise ($\sigma_l = 0.01, 1$), both of which are defined as standard deviations of 77 normal distributions. When $\sigma_h > \sigma_l$, branching produces greater habitat heterogeneity because headwaters 78 are the primary source of environmental variation (Figure 1A). This landscape scenario should reproduce 79 natural patterns of habitat heterogeneity, in which environmental conditions differ greatly among tributaries 80 but are highly correlated within a tributary (16, 22, 23). In the meantime, when $\sigma_h \leq \sigma_l$, local environmental 81 noise masks environmental variation among tributaries, leading to limited influences of branching on habitat 82 heterogeneity (Figure 1B). This scenario may reflect human-modified landscapes where the physical or biological distinctiveness of tributaries has been compromised due to human activities (24, 28). Thus, the 84 inequality between σ_h and σ_l creates contrasting patterns of habitat heterogeneity within a network. We also considered eight ecological scenarios that differ in dispersal distance (controlled by the rate parameter of an 86 exponential dispersal kernel θ), dispersal probability p_d , and the maximum value of interspecific competition 87 strength α_{max} (see Materials and Methods). These simulation scenarios were capable of reproducing 88 common spatial patterns of local species richness in rivers, corroborating the appropriateness of our choice 89 in parameter combinations (Supplementary Text). Under each simulation scenario, we simulated 1400 time steps of metacommunity dynamics (including 400 time steps of initialization and burn-in periods) in 91 1000 branching networks with the gradients of ecosystem size (the number of habitat patches: 10 to 150) and complexity (branching probability: 0.01 to 0.99). 93

Our theoretical analysis yielded results consistent with our prediction. Ecosystem size and complexity both increased γ diversity under a realistic landscape scenario (**Figures 2 and 3**), where the environmental variation at headwaters σ_h is greater than the degree of local environmental noise σ_l (**Figure 1A**). The

relationships had a characteristic of power-law (i.e., linear in a log-log scale) and were consistent under various ecological scenarios (compare panels in **Figures 2 and 3**). Importantly, the impact of ecosystem complexity was comparable to that of ecosystem size. Hence, regardless of ecological scenarios, ecosystem size and complexity are likely to be equally important in regulating γ diversity.

However, dispersal processes affected mechanisms that underlie the positive effects of ecosystem size and complexity on γ diversity (compare left and right columns in Figures 2 and 3). We observed a greater contribution of β diversity (defined as $\frac{\gamma}{\alpha}$) to increased γ diversity when dispersal limitation existed (i.e., species travel short distances). This pattern reflected significant spatial variation in species composition over the branching network and was likely driven by the local association of species' niche and abiotic environments (i.e., species sorting) (27, 29). In contrast, when the dispersal limitation was relaxed (species travel long distances), a clear increase in α diversity underpinned the positive relationships between γ diversity and ecosystem properties. The results agree with previous predictions that increased dispersal homogenizes community composition while enhancing local diversity through increased immigrants from suitable habitat patches (i.e., mass effects) (27, 29, 30). Therefore, ecosystem size and complexity can increase γ diversity via enhanced species sorting or mass effects, and which takes primacy is dependent on the degree of dispersal limitation. These patterns were consistent across different levels of dispersal probabilities (Figures S5 and S12). The strength of competition decreased the maximum of α diversity but did not change the scaling relationships with ecosystem properties (Figures 2 and 3). In summary, our theory highlights how apparently similar patterns in γ diversity emerge through different ecological pathways.

Influences of ecosystem size and complexity differed significantly in their dependence on landscape scenarios. Ecosystem size had positive effects on γ diversity regardless of landscape scenarios, although the slopes were steeper with greater environmental variation (higher σ_h and/or σ_l) (Figures 2 and S5-S11). This result is attributable to the fact that larger ecosystems can hold more individuals in a metacommunity (5). In contrast, we observed limited or no influences of branching complexity when local environmental noise was equal to or exceeded environmental variation at headwaters ($\sigma_l \geq \sigma_h$; Figures S13-S18). Under this scenario, branching has a minor influence on the ecosystem's habitat heterogeneity because patch-level environmental variation is equivalent to or greater than environmental differences between tributaries (Figure 1B). Therefore, this landscape scenario eliminates the positive effect of branching complexity on γ diversity. This theoretical prediction may not apply to pristine or semi-natural river networks where individual streams show distinct and spatially-correlated environmental conditions, including water temperature, water chemistry, and flow/sediment regimes (22, 23). Instead, it may be most relevant to severely altered landscapes where localized human disturbance disrupts the environmental distinctiveness of branches through, for example, flow regulations by dams (28). Hence, our theory has important implications for riverine biodiversity conservation by pointing to the crucial role of habitat diversity produced by branching structure.

Empirical evidence from distinct geographic regions

The proposed theory provided important insights into how ecological communities are structured in branching networks; however, empirically testing the predictions is extremely difficult because it requires metacommunity-level replications. To confront this logistical challenge, we compiled fish community data across two geographic regions: Hokkaido Island in Japan and the Midwestern US (Midwest). These regions are located in comparable latitude ranges (Figure 1D) but support distinct fish communities (Tables S5 and S6). Therefore, this data set provides an excellent opportunity to examine the generality of our theoretical predictions. After careful data selection (see Materials and Methods and Supplementary **Text**), we estimated α , β , and γ diversity (asymptotic species richness; **Materials and Methods**) at 180 watersheds (59 in Hokkaido and 121 in the Midwest), each of which comprised > 10 sites of presence-absence fish community data (a total of 6605 sites). These watersheds are small enough (40 to 4939 km²) to assume that fishes can disperse therein at a multi-generation time scale (31), while posing challenges to traverse across watersheds (the ocean or lentic habitats; see Materials and Methods for watershed definition). We combined this data set with geospatial information, including watershed characteristics (watershed area and branching probability), climates (annual mean temperature and annual cumulative precipitation), and land use patterns (the fraction of agricultural land use and dam density). Using this data set, we assessed whether observed effects of the macro-scale factors are consistent across the two geographic regions by developing global and region-specific robust regression models. In the global model, we assumed that effects of ecosystem size (watershed area) and complexity (branching probability) are constant across the two regions (i.e., fixed slopes). Meanwhile, the region-specific model assumes region-specific slopes of ecosystem size and complexity (Materials and Methods). We compared the support of these competing models using the Bayes factor, a measure of the strength of evidence in favor of one model over the alternative. In our definition (Materials and Methods), the Bayes factor of > 1 indicates the support for the global model over the region-specific model.

We found decisive support for the global model in explaining γ diversity (Bayes factor = 153.8) despite the substantial difference in fish fauna between the study regions. The estimated γ diversity increased with increasing watershed area (ecosystem size) and branching probability (ecosystem complexity), and these patterns were consistent across the study regions (Table 1 and Figure 4). In particular, the effect of ecosystem complexity was striking in its magnitude. Average predictive comparisons (32) revealed that an expected increase of γ diversity per 0.1 branching probability was 10.98 \pm 4.49 species (Materials and Methods and Supplementary Text); this level of increase in γ diversity requires 1911 km² in the watershed area (see Table S7 for the estimated average predictive comparisons). This result is reasonable because numerous studies have shown that tributaries are the primary source of environmental heterogeneity in river networks (16, 22-25) - the sole theoretical assumption required for the dual control of γ diversity to emerge. Further, the consistent patterns across geographic regions with distinct fish fauna suggest that ecological traits of species in metacommunities may have a minor effect on the scaling relationships, as predicted by our theory.

Similarly, there were weak to moderate supports for the global models of α and β diversity (Bayes factor = 1.5 and 8.9, respectively). In both regions, β diversity responded significantly to ecosystem size and complexity while α diversity showing a weaker or a vague response to these ecosystem properties (Table 1 and Figure 4). In our simulations, this pattern has emerged under the scenarios with dispersal limitation, which elegantly matches the previous observations of stream fish movement. Field studies (e.g., markrecapture) recurrently revealed the restricted movement of stream fish, typically limited to several hundreds of meters at an annual time scale (33). The reciprocal agreement of theoretical and empirical patterns provides indirect but convincing evidence that dispersal limitation, which results in the increasing importance of species sorting process (Figures 2 and 3), plays a key role in driving the associations between γ diversity and ecosystem properties in rivers.

The consistent effect of branching probability on γ diversity across the study regions is noteworthy because many watersheds in the Midwest have been altered by agricultural land use (mean % agricultural land use: 179 55% in the Midwest and 6% in Hokkaido). If the intensive land use by humans impairs biological or physical 180 distinctiveness among tributaries, theory predicts a weakened effect of branching probability on γ diversity. However, γ diversity increased significantly with increasing branching probability in this highly modified landscape, suggesting that tributaries still sustain unique environmental conditions to support high spatial 183 variation in species composition. Indeed, β diversity increased with increasing branching probability in both regions (Figure 4). It is conceivable that local geological and geomorphological differences, such as slope, aspect, and soil porosity, persist in human-dominated landscapes to maintain the diversity of in-stream processes among tributaries. The lack of land use effects (the fraction of agricultural land use and dam density) further corroborates our interpretation (**Table 1**). Although our analysis is correlative, the finding is encouraging because the branching complexity of river networks may serve as a natural defense system to human-induced environmental changes.

A potential criticism is that our results are driven by spurious correlations; however, this is very unlikely. We 191 have statistically controlled the potential influences of climates and land use patterns (**Table 1**). Moreover, 192 our empirical results were fairly consistent with theoretical predictions, which are free from any confounding 193 factors. Thus, we are confident in our statistical inference.

Implications 195

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Using theory and extensive community data, our study provides the strong evidence that ecosystem size 196 and complexity play equally important roles in regulating riverine biodiversity. In particular, the emerging complexity-diversity relationship points to several important avenues for riverine biodiversity conservation. First and foremost, there is now a clear need to explicitly consider the dimension of ecosystem complexity to achieve successful conservation. Human-induced habitat alterations, including flow regulation (28), habitat fragmentation (34), and stream burial (35), may compromise or restrict access to the diverse habitats that complex branching networks may support. Hence, it is imperative to recognize the role of branching complexity and minimize the homogenizing effects of human activities. Second, the complexity perspective may provide insights into the spatial planning of conservation efforts. Riverine reserves and local restoration are increasingly recognized as an effective management tool, and designing a spatial network of conservation sites in rivers is an area of active research (36). While large reserves or restoration sites are undoubtedly important, our results indicate the potential of coordinated networks of local conservation sites in protecting biodiversity. For example, synergies of multiple small reserves may emerge at the watershed scale when ecologically-distinct tributaries are involved in the design, as evidenced by the recent successful conservation of tropical fishes in Thailand's Salween basin (36).

While the prevailing evidence supports the importance of ecosystem size in scaling biodiversity patterns (3, 211 5, 37, 38), ecosystem complexity has not received the attention it deserves. However, the ubiquity of scale-212 invariant complexity across terrestrial (11) and aquatic ecosystems (12) calls for more research embracing the 213 two orthogonal dimensions of the ecosystem's geometric structure. Our fundamental findings should apply 214 beyond branching ecosystems because ecosystem size and complexity both represent a physical 'template' 215 providing a wide spectrum of niche opportunities for living organisms. Further, the proposed framework 216 may be extended to other aspects of biodiversity; for example, a logical next step is to explore how these 217 ecosystem properties regulate functional diversity (39, 40) and shifts in community assembly processes (41). 218 Recognizing the dual control of biodiversity broadens viable options for spatial planning of protected areas 219 or restoration sites, thereby helping conserve biodiversity from societal demands that threaten it. The 220 generalization of our findings in spatially complex ecosystems represents a frontier for future research. 221

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231 Author contributions

AT conceived of the project. AT performed theoretical and statistical analysis with inputs from TK. AT, SK, CLD, and YM compiled and managed the fish community data set. AT wrote the first draft while SK, CLD, TK, and YM contributed significantly to the final version.

Data and code availability

Third parties provided data for empirical data analysis; references are provided where appropriate. R functions for the generation of branching networks and the metacommunity simulation are provided as the R package 'mcbrnet' (available at https://github.com/aterui/mcbrnet). Codes for simulations, statistical analysis, figures, and tables are available at https://github.com/aterui/public-proj stream-diversity.

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