Spatial phylogenetics of Japanese ferns: Patterns, processes, and implications for conservation

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Abstract
Premise: Biodiversity is often only measured with species richness; however, this metric ignores evolutionary history and is not sufficient for making conservation decisions. Here, we characterize multiple facets and drivers of biodiversity to understand how these relate to bioregions and conservation status in the ferns of Japan.

Methods: We compiled a community data set of 1239 grid cells (20 × 20 km each) including 672 taxa based on >300,000 specimen records. We combined the community data with a phylogeny and functional traits to analyze taxonomic, phylogenetic, and functional diversity and modeled biodiversity metrics in response to environmental factors and reproductive mode. Hierarchical clustering was used to delimit bioregions. Conservation status and threats were assessed by comparing the overlap of significantly diverse grid cells with conservation zones and range maps of native Japanese deer.

Results: Taxonomic richness was highest at mid-latitudes. Phylogenetic and functional diversity and phylogenetic endemism were highest in small southern islands. Relative phylogenetic and functional diversity were high at high and low latitudes, and low at mid-latitudes. Grid cells were grouped into three (phylogenetic) or four (taxonomic) major bioregions. Temperature and apomixis were identified as drivers of biodiversity patterns. Conservation status was generally high for grid cells with significantly high biodiversity, but the threat due to herbivory by deer was greater for taxonomic richness than other metrics.

Conclusions: Our integrative approach reveals previously undetected patterns and drivers of biodiversity in the ferns of Japan. Future conservation efforts should recognize that threats can vary by biodiversity metric and consider multiple metrics when establishing conservation priorities.

KEYWORDS
apomixis, biodiversity, biogeography, CANAPE, conservation, ferns, Japan, phylogenetic diversity

Characterizing the spatial distribution of biodiversity is a major goal of evolutionary biology with two equally important aspects: to understand the processes generating it and to conserve it. Until recently, the vast majority of studies seeking to understand the spatial distribution of biodiversity have focused on species richness. Hotspots, or areas of exceptionally high species richness or endemism, have received particular attention, resulting in a widely recognized set of 36 terrestrial hotspots (Myers et al., 2000; Noss, 2016) and motivating conservation strategies to preserve them (Margules and Pressey, 2000; Brooks et al., 2006). However, species richness alone cannot provide a complete picture of biodiversity (Miller et al., 2018). All organisms are related to a greater or lesser degree by descent from a common ancestor, and these evolutionary relationships must be taken into account to obtain a full understanding of the biodiversity present in an area.

Presumably, phylogeny has not been taken into account more prominently in biodiversity studies mainly because...
the necessary data (DNA sequences and georeferenced occurrence records) and analytic tools have only become available relatively recently (Soltis and Soltis, 2016; Folk and Siniscalchi, 2021). These data sets and tools now make it possible to analyze other dimensions of biodiversity such as phylogenetic diversity (Faith, 1992) and phylogenetic endemism (Rosauer et al., 2009). Better ways to characterize biological regions (“bioregions”; i.e., areas defined by their taxonomic composition or evolutionary history) are also available (Laffan et al., 2016; White et al., 2019; Daru et al., 2020b), rather than relying on ad hoc characterizations. Furthermore, incorporating these two frameworks—the categorization of areas into phyloregions with analysis of over/under dispersion of biodiversity—can provide powerful insights into the processes structuring biodiversity and suggest conservation priorities.

A comprehensive understanding of the relationships between richness and other metrics requires densely sampled data, but such data sets are rare on the regional (country) scale. The ferns of Japan are excellent model system because they have been the target of intense botanical interest for several decades and are densely sampled (reviewed by Ebihara and Nitta, 2019a). The ferns of Japan include 676 native, nonhybrid taxa (including species and varieties) and hundreds of hybrids (nothotaxa; Ebihara and Nitta, 2019a). The availability of detailed distribution data (distribution maps at a scale of ca. 10 km for nearly all species; Ebihara, 2016, 2017), trait data (multiple quantitative and qualitative traits compiled for identification of nearly all species; Ebihara and Nitta, 2019a), and DNA sequences for >97% of species (Ebihara et al., 2010; Ebihara and Nitta, 2019a; all coverage statistics exclude nothotaxa) make the ferns of Japan an ideal system for investigating the relationships between, and drivers of, multiple dimensions of biodiversity.

One particularly valuable characteristic of the Japanese fern flora is the availability of data on reproductive mode, which is known for 492 native fern taxa excluding hybrids (72.8%; Ebihara and Nitta, 2019a). Reproductive mode is likely to affect population-level genetic diversity (Bengtsson, 2003) and thereby higher-level biodiversity (Krueger-Hadfield et al., 2019). In ferns, clonally reproducing apomictic species are often polyploid hybrids that share identical plastid genotypes among taxa and within populations (Grusz et al., 2009; Chao et al., 2012; Hori et al., 2014; Hori and Murakami, 2019). Therefore, we expect high prevalence of apomictic species within a community to decrease phylogenetic diversity.

Japan is situated along a latitudinal gradient spanning temperate, seasonal areas in the north to subtropical, mostly aseasonal islands in the south. Furthermore, it is a mountainous country with great variation in elevation on the larger islands. Plant distributions often reflect physiological adaptations to climate (e.g., precipitation and temperature; Woodward, 1987). Therefore, we expect the spatial distribution of biodiversity to be determined by climatic variation in addition to reproductive mode.

Here, we leveraged this exceptionally rich data set to analyze the geographic distribution of biodiversity in detail. We asked the following questions in our study system, the ferns of Japan: (1) How is biodiversity distributed? (2) How is biodiversity structured? (3) What environmental and biological factors influence biodiversity? (4) How well is biodiversity protected?

MATERIALS AND METHODS

All computational analyses were carried out in R v4.1.2 (R Core Team, 2021) unless otherwise stated. The R package targets v.0.9.0 was used to control analysis workflow (Landau, 2021).

Study site

Japan (20°N to 45°N, 122°E to 153°E) consists of four main islands and thousands of smaller ones (Figure 1). Most of the main islands have been in contact with the continent at various points in the past during periods of lower sea level, but the Ogasawara archipelago consists of oceanic islands that have never been connected to the mainland. The climate is subtropical in the southern islands and temperate in other parts of the country. Map data for Japan were downloaded from the Geospatial Information Authority of Japan under the Creative Commons Attribution License v4.0 (https://www.gsi.go.jp/kankyochiri/gm_japan_e.html).
Occurrence data

We used a list of native, nonhybrid fern specimens deposited at herbaria in Japan to quantify occurrences (Ebihara, 2016, 2017; Ebihara and Nitta, 2019). We chose to use this over other sources such as the Global Biodiversity Information Facility (GBIF) because of its high quality, which obviates many cleaning steps otherwise needed when working with large, publicly available data sets. Furthermore, the overall sampling completeness of this data set is also high (Figure S1 in Appendix S1). The original list comprises 337,000 specimens representing 673 terminal taxa at the rank of species, subspecies, form, or variety, excluding non-native taxa and nothotaxa. We treated infraspecific taxa as distinct species during analysis and hereafter use “taxon” and “species” interchangeably to refer to entities at this rank, unless otherwise indicated. All names are standardized to a common taxonomy, the Japan Green List v. 1.01 (Ebihara and Nitta, 2019). Occurrence records were georeferenced and thoroughly vetted as follows (Ebihara, 2016, 2017; Ebihara and Nitta, 2019). For specimens that lacked latitude and longitude vetted as follows (Ebihara, 2016, 2017; Ebihara and Nitta, 2019). We chose to use this over other sources such as the Global Biodiversity Information Facility (GBIF) because of its high quality, which obviates many cleaning steps otherwise needed when working with large, publicly available data sets. Furthermore, the overall sampling completeness of this data set is also high (Figure S1 in Appendix S1). The original list comprises 337,000 specimens representing 673 terminal taxa at the rank of species, subspecies, form, or variety, excluding non-native taxa and nothotaxa. We treated infraspecific taxa as distinct species during analysis and hereafter use “taxon” and “species” interchangeably to refer to entities at this rank, unless otherwise indicated. All names are standardized to a common taxonomy, the Japan Green List v. 1.01 (Ebihara and Nitta, 2019).

Occurrence records were georeferenced and thoroughly vetted as follows (Ebihara, 2016, 2017; Ebihara and Nitta, 2019). For specimens that lacked latitude and longitude, georeferencing was done by mapping collection site names to a set of standard grid squares of ca. 10 × 10 km defined by the Statistics Bureau of Japan (the “second-degree mesh”; http://www.stat.go.jp/english/data/mesh/05.html), and the centroid of the second-degree mesh cell used as the specimen location. In the case that the collection site could not be mapped to a single second-degree mesh cell, it was excluded. Next, occurrence maps were generated for each taxon showing presence or absence in each second-degree mesh cell. In the case that a given taxon appeared insufficiently sampled (i.e., not present on the map where it would typically be expected to occur), AE and members of the Nippon Fernist Club (local botanists familiar with the ferns of Japan) searched for additional specimens, which were then added to the list. The occurrence maps were iteratively refined until the vast majority of known Japanese ferns had been observed across their expected ranges; the resulting set of occurrence records can be considered accurate to ca. 10 km (the grain size of the second-degree mesh map).

We further cleaned the list before analysis by filtering out any occurrences not within the second-degree mesh and removing duplicate collections (300,685 specimens, 673 taxa after filtering). Given the high quality of our occurrence data and the fact that automated occurrence cleaning algorithms (e.g., CoordinateCleaner; Zizka et al., 2019) have the potential to erroneously exclude true occurrence points (i.e., false positives; Zizka et al., 2020), we chose not to apply additional automated cleaning steps to our data as is often done with occurrence records obtained from GBIF (e.g., Rice et al., 2019; Suissa et al., 2021).

A necessary step in any analysis of biodiversity is to set the grain size used to accurately define communities (co-occurring species). Here, one must consider that smaller grain size is needed to detect environmental effects, while larger grain size is needed to ensure adequate species sampling. We created sets of 10-km, 20-km, 30-km, and 40-km grid cells covering Japan using a Mollweide equal-area projection (size refers to the length of the side of each square grid cell). At each grain size, species occurrences were converted to a presence–absence community matrix (a species was considered present if at least one specimen was recorded in that grid cell). We calculated sampling redundancy (1 – Richness/Number of specimens; Garcillan et al., 2003) to quantify adequacy of sampling. Preliminary analysis indicated that 20-km grid cells are optimal for our data set; redundancy values greatly increased from 10 to 20 km, but much less as grain size is increased beyond 20 km (Appendix S1; Figure S2). Although the grid cells are defined with equal area, actual land area of each grid cell varies due to coastlines. Strictly filtering out all grid cells with less than complete land area would result in a large loss of data, as Japan has many coastlines and small islands. Therefore, we instead filtered the grid cells by sampling completeness, removing any grid cells with redundancy <0.1; this data set was used for all subsequent analyses. The R package sf v.1.0.5 was used for all GIS analyses (Pebesma, 2018).

Morphological trait data

We used traits originally compiled for identification of ferns and lycophytes of Japan (Ebihara, 2016, 2017), which were formatted to be used with Lucid software (https://www.lucidcentral.org/). Continuous traits were measured on 10 randomly selected specimens per species, then four values per species were obtained from these measurements following the Lucid format: outside (i.e., outside the typical range) minimum, typical minimum, typical maximum, and outside maximum. For species with dimorphic fertile (i.e., spore-bearing) and sterile fronds, fertile and sterile fronds were measured separately. We took the mean of the typical minimum and maximum values to use as the species mean value. Qualitative traits were scored by observing voucher specimens. All qualitative traits were scored in binary format; e.g., a single trait with three states was formatted as three binary traits. From the original trait list, we selected only traits with putative ecological function (Table 1) and excluded any traits that had Pearson correlation coefficient >0.6 or fewer than three observations of a given trait state.

Reproductive mode data

We used data compiled by Ebihara and Nitta (2019a), which classifies each fern taxon as sexual, apomictic, mixed (both sexual and apomictic modes known from the same taxon), or unknown. We calculated the percentage of apomictic taxa in each grid cell as the sum of apomictic and mixed taxa divided by total number of taxa (Appendix S1: Figure S3B).
TABLE 1  Fern traits used in this study. Letters in parentheses indicate trait type: b, binary (trait present or absent); c, continuous; q, qualitative. Qualitative traits were coded as binary for analysis (see Materials and Methods). For complete list of trait states for qualitative traits, see Appendix S1: Table S3.

| Trait                | Significance                                                                 | Reference                      |
|----------------------|------------------------------------------------------------------------------|--------------------------------|
| Indusium (b)         | Protects developing spores                                                  | Poppinga et al. (2015)         |
| Frond width (c)      | Scales with overall plant size                                              | Creese et al. (2011)           |
| No. pinna pairs (c)  | Less-divided fronds have lower rates of evapotranspiration                  | Kluge and Kessler (2007)       |
| Stipe length (c)     | Shorter lengths minimize resistance                                         | Watkins et al. (2010)          |
| Frond margin (q)     | Thermoregulation                                                            | Nicotra et al. (2011)          |
| Frond shape (q)      | Less-divided fronds have lower rates of evapotranspiration                  | Kluge and Kessler (2007)       |
| Frond texture (q)    | Thicker leaves have lower rates of evapotranspiration                       | Givnish (1987)                 |

Environmental data

We downloaded climate data at the 2.5 minute-scale from the WorldClim database using the getData function in the R package raster v.3.5.11 (Hijmans, 2021) and extracted four variables related to our research questions: mean annual temperature (bio1; hereafter “temperature”), temperature seasonality (bio4), annual precipitation (bio12; hereafter “precipitation”), and precipitation seasonality (bio15) (Appendix S1: Figure S3). We identified the intersection of the climate data at the 2.5-minute scale with the 20-km grid cells, then calculated the mean of each climatic variable for each grid cell. We calculated area as rolling mean area (km²) across 100-km latitudinal bands. This approach reflects the idea that the amount of area with a given climate regime enclosing each grid cell may influence species richness of that cell (Fahrig, 2013) and that using raw island areas would not be appropriate in this context because there are just four very large islands with many much smaller ones close by (Figure 1).

Phylogenetic analysis

Sequences for the plastid, coding rbcl gene are available for ca. 97% of the Japanese fern flora (Ebihara and Nitta, 2019a); these sequences originate from samples that were identified using the same taxonomic system as the occurrence data (Ebihara et al., 2010; Ebihara and Nitta, 2019a), so we used these preferentially over other data available on GenBank that might include misidentifications or different taxon concepts. However, using this data set alone for phylogenetic analysis suffers two drawbacks: it is not densely sampled enough to resolve some internal nodes, and it lacks many lineages that have fossils available for molecular dating. Furthermore, use of community phylogenies generated by sampling only the species present in the local community has been shown to produce spurious results in simulation studies (Park et al., 2018). Because one goal of the study is to understand the distribution of paleo- vs. neoendemic areas (defined in units of time), we required an ultrametric tree. Therefore, to obtain a robust, ultrametric tree, we combined the rbcl data of Ebihara and Nitta (2019a) (706 taxa) with a globally sampled plastid data set including all fern sequences on GenBank for four widely sequenced plastid genes (atpA, atpB, rbcl, and rps4; 4492 taxa) and 58 other coding, single-copy plastid genes extracted from 123 complete fern plastomes (FTOL v0.0.1; Nitta et al., 2022) (4787 operational taxonomic units [OTUs] total, including outgroup taxa). This gene sampling is comparable to a recent global fern phylogeny that resolved relationships across ca. 4000 taxa using six plastid markers (Testo and Sundue, 2016), and the addition of plastome data can be expected to increase support along the backbone.

We conducted maximum-likelihood phylogenetic analysis with IQ-TREE v1.6.12 (Nguyen et al., 2015) under the GTR+I+G model (all sites concatenated into a single data matrix), and assessed node support with 1000 ultrafast bootstrap replicates (Minh et al., 2013). Molecular dating was conducted with the program treePL v1.0 (Smith and O’Meara, 2012) using 26 fossil calibration points after Testo and Sundue (2016), with the exception of treating fossil Kuylisporites as belonging to crown Cyathea, not Cyathea + Alsophila (Loiseau et al., 2020). The root of the tree (most recent common ancestor of all land plants) was fixed at 475 million years ago (Ma; Testo and Sundue, 2016). We trimmed the resulting ultrametric tree to only Japanese taxa. treePL requires branch lengths to be >0, and it sets extremely small branches to an arbitrary minimum positive length. This behavior may artificially add branch length between taxa with identical DNA sequences, which occur multiple times in the Japanese fern rbcl data set. We converted any clades in the dated tree consisting of identical sequences to polytomies with the node at 0 Ma (i.e., the present). This tree was used for all subsequent analysis unless indicated otherwise.
Phylogenetic signal

We tested for phylogenetic signal in continuous traits using two alternative metrics, Blomberg’s K (Blomberg et al., 2003) and Pagel’s λ (Pagel, 1999). K describes the ratio between the observed variance in a trait vs. the amount of variance expected under Brownian motion (BM); λ is a scaling parameter between 0 and 1 that transforms the phylogeny such that trait values fit those most closely expected under BM. Both λ and K = 1 under BM; relatively lower values indicate less signal (overdispersion) and higher values indicate more signal (clustering). We measured K and λ using the phylosig function in the R package phytools v.0.7.90 (Revell, 2012), which also calculates significance based on a randomization test for K and a likelihood ratio test for λ. We tested for phylogenetic signal in qualitative (coded as binary) traits using Fritz and Purvis’ D (Fritz and Purvis, 2010) with the phylo.d function in the R package caper v.1.0.1 (Orme et al., 2018), which also assesses significance by comparing the observed value to a null distribution of random values and a simulated distribution of values expected under BM. Values of D range from 0 under BM to 1 under random evolution, but can exceed this range in cases of extreme clumping or over-dispersion, respectively. These methods for phylogenetic signal cannot use a tree with zero dispersion, respectively. These methods for phylogenetic signal exceed this range in cases of extreme clumping or over-dispersion.

Analysis of biodiversity

We measured taxonomic diversity using taxonomic richness (i.e., number of taxa in each community). We measured phylogenetic diversity (PD) using Faith’s PD (the total branch length connecting all terminal taxa in a community, including the root of the phylogeny; Faith, 1992). To detect areas with unusually long or short branches, we calculated relative phylogenetic diversity (RPD), which is the ratio of observed PD to PD measured on a tree where all branch lengths have been transformed to equal length (Mishler et al., 2014).

We measured functional diversity (FD) using the method of Petchey and Gaston (2002). We first calculated functional distances using Gower’s method (Gower, 1971) on the trait data with the gowdis function in the R package FD v.1.0.12 (Laliberté and Legendre, 2010). We weighted continuous and qualitative traits equally, weighted each binary component trait of each qualitative trait equally, and log-transformed and scaled continuous traits prior to calculating distances. We also tested alternative weighting schemes, but these produced very similar results (data not shown). We then used the distances to build a trait dendrogram using the hclust function in the R package stats v.4.1.2 (R Core Team, 2021) and used the dendrogram in place of a phylogenetic tree to calculate PD and RPD as described above. These functional analogs of PD and RPD are hereafter referred to as FD (functional diversity) and RFD (relative functional diversity).

We measured phylogenetic endemism (PE) as the sum of all branches at a site inversely weighted by their range size (Rosauer et al., 2009).

Observed values of PD, RPD, FD, RFD, and PE are known to be related to species richness; as more taxa are drawn at random without replacement from a given set, it becomes increasingly unlikely to draw a taxon that is distantly related (or functionally dissimilar) to the others. Therefore, we determined whether the observed metrics are more or less diverse than random for a given number of taxa by conducting a randomization test. We generated 999 random communities using the curveball algorithm of Strona et al. (2014), which conserves richness per site and species occurrence frequencies while randomizing species identities, then compared the observed value to the distribution of random values. Statistical significance was assessed using a one-tailed test for PE or a two-tailed test for other metrics with $\alpha = 0.5$; observed values in the extreme 5% of the null distribution were considered significant. We also calculated the standard effect size (SES) of each biodiversity metric. The standard effect size measures how extreme the observed value (obs) is relative to the null distribution (null):

$$\text{SES} = \frac{\text{obs} - \text{mean}(\text{null})}{\text{sd}(\text{null})}$$

We tested whether centers of endemism have an over-representation of old (paleoendemic) or new (neoendemic) lineages using categorical analysis of neo- and paleoendemism (CANAPE), which involves measuring PE within an alternate tree where all branch lengths are equal, then comparing the ratio between raw PE and alternative PE (RPE) (Mishler et al., 2014). Low RPE indicates short range-restricted branches (neoendemism), and high RPE indicates long range-restricted branches (paleoendemism). Since all of these measures are affected by species richness, categorization of endemism is done using the rank of each observed variable relative to those calculated for the null distribution. All measures of biodiversity and CANAPE were carried out using the R package canaper v.0.0.2 (Nitta, 2022).

It is possible that endemism may be affected by species that are actually wide-ranging, but only have a small range in Japan (a “border effect”). The border effect is especially expected for southern, subtropical islands, which harbor many tropical species with small ranges in Japan but wider ranges outside of the country. To partially account for this border effect, we also calculated all endemism-related metrics using a data set including species found only in Japan.

Spatial models

Two grid cells lacking environmental data, one grid cell occupied by a single species lacking reproductive mode data, and an outlier with 100% apomictic taxa due to
We tested for spatial autocorrelation in biodiversity metrics (richness, SES of PD, SES of FD, SES of RPD, SES of RFD) and independent variables (environmental variables and percentage apomictic taxa [hereafter, % apomictic taxa]) using Moran’s I with the moran.mc function in the R package spdep v.1.1.13 (Bivand et al., 2013). In all cases, significant spatial autocorrelation was detected (Moran’s I > 0; P value of permutation test < 0.05) (Appendix S1: Table S1). Therefore, we used spatially explicit methods for modeling.

We checked for correlation between independent variables while taking into account spatial autocorrelation using a modified t-test with the modified.ttest function in the R package SpatialPack v.0.3.8196 (Vallejos et al., 2020). This test indicated that temperature seasonality and % apomictic taxa were correlated with mean annual temperature (Appendix S1: Table S2), so for the main modeling analysis, we only included the following environmental variables: mean annual temperature, precipitation, and precipitation seasonality.

We constructed a linear mixed model for each biodiversity metric dependent on the environmental variables (“environmental models”). Initial inspection of the relationship between each biodiversity metric and temperature showed a hump-shaped pattern (Appendix S2), so we also included a quadratic term for temperature only. Richness was fit with the negative binomial response family; others (SES of PD, SES of FD, SES of RPD, SES of RFD) were Gaussian. We fit models including a Matérn correlation matrix based on the geographic centroids of each grid cell as a random effect with the R package spaMM v.3.9.25 (Rousset and Ferdy, 2014). We verified the significance of each term of the model by conducting likelihood ratio tests (LRTs) between the full model and models each with a single term of interest removed, in sequence. We also verified the significance of the full model relative to the null model (only including the Matérn correlation matrix) using LRTs.

We postulated that % apomictic taxa might explain the distribution of phylogenetic diversity better than temperature, so we also constructed a limited set of models to test this hypothesis (“reproductive models”). These models included biodiversity metrics with a phylogenetic component (SES of PD and SES of RPD) as dependent on % apomictic taxa, precipitation, and precipitation seasonality, with random spatial effects as described above. We compared the fit between environmental models and reproductive models for SES of PD and SES of RPD using conditional Akaike information criterion (cAIC) (Vaida and Blanchard, 2005; Courtiol and Rousset, 2017).

Analysis of bioregions

We analyzed bioregions using the framework of Daru et al. (2020a) and their R package phylregion v.1.0.6. Briefly, the analysis involves first calculating beta diversity (change in composition between communities) using either species names (taxonomic bioregions) or phylogenetic distances (phylogenetic bioregions). For taxonomic bioregions, we calculated the turnover in species between sites due to species replacement (Baselga, 2012) with the Sorensen index. For phylogenetic bioregions, we calculated phylogenetic dissimilarities based on the PhyloSor index (Bryant et al., 2008). The distances are used to construct a dendrogram, which is then split into k bioregions. The optimal value of k cannot be known a priori. Previous vegetation zone schemes in Japan have typically included five or six zones (Shimizu, 2014); to enable approximate comparison with these, we tested values of k from 1 to 10, and selected the optimal value using the optimal_phylregion function in the R package phylregion v.1.0.6.

Assessment of conservation status and threats

We downloaded SHP files of conservation zones in Japan from the Japan Ministry of the Environment (https://www.biodic.go.jp/biodiversity/activity/policy/map/map17/index.html) and Ministry of Land, Infrastructure, Transport and Tourism (https://nlftp.mlit.go.jp/ksj/gml/datalist/KsjTmplt-A45.html). We excluded marine zones and those that do not protect plants. We categorized protected areas as either high (no human activities allowed at all) or medium status (some economic activities allowed by permit; Kusumoto et al., 2017); areas not afforded at least medium level of protection were not considered. Conservation status in Japan is administered by multiple laws, and protected areas frequently overlap (Natori et al., 2012). To prevent double-counting, all protected areas within a protection status were merged, and protected areas that overlapped between medium and high status were considered only high status.

Besides habitat loss due to human activity, one major threat to ferns in Japan is herbivory by Japanese deer (Cervus nippon). Although native, Japanese deer have caused extensive damage to native plant communities due to rapid population growth and range expansion since the 1970s (Takatsuki, 2009). Decrease or extirpation of fern populations due to deer herbivory has been observed in multiple sites and species across the country (Minamitani, 2005; Yahara, 2006; Hattori et al., 2010). Furthermore, 33 of 212 (15.5%) fern taxa listed as endangered (Red List status CR, EN, or VU) in Japan include herbivory by deer as one of the reasons for their endangered status (Japan Ministry of the Environment, 2015, 2022). Although not all species of ferns are equally susceptible to deer herbivory (some are unpalatable to deer; Minamitani, 2005), these studies show it is an existential threat to many species. Therefore, analysis of the threat posed to fern biodiversity by deer herbivory is needed to inform future conservation policies.

We downloaded distribution maps (SHP and TIF) of Cervus nippon from the Japan Ministry of the Environment.
To assess change in threat due to herbivory by deer over time, we used three maps available in this data set: range of Japanese deer surveyed in 1978, range surveyed in 2003, and estimated range inferred from a model including snow cover and forest type based on the 2003 survey data (Japan Ministry of the Environment, 2021b). We limited estimated range to areas considered “highly likely” to be occupied by deer (movement cost < 0.10; Japan Ministry of the Environment, 2021b).

We cropped grid cells to only include land area, then calculated the percentage of protected area of each status type or percentage of area occupied by deer for each survey period within grid cells with high levels of biodiversity as measured with PD, FD, PE, or taxonomic richness. These values were compared with the overall percentage of protected area or area occupied by deer across Japan (baseline rate). For PD, FD, and PE, significance was assessed with a one-tailed test at $\alpha = 0.05$ against the 999 null communities described above. For taxonomic richness, grid cells ranked in the top >5% were considered highly diverse.

**RESULTS**

**Data sets**

The phylogeny included 663 taxa (98.08% of the total number of native, nonhybrid ferns [676 taxa]) representing 98 genera and 34 families. The topology generally agreed with earlier, large plastid fern phylogenies (Schuettpelz and Pryer, 2007; Lehtonen, 2011; Testo and Sundue, 2016; Appendix S1: Figure S4). Ferns were assessed with a one-tailed test at $\alpha = 0.05$ against the 999 null communities described above. For taxonomic richness, grid cells ranked in the top >5% were considered highly diverse.

**Phylogenetic signal**

All continuous traits showed significant phylogenetic signal, but estimates of signal strength differed between $K$ and $\lambda$: all $\lambda$ values were >0.95, indicating evolution by BM; however, values of $K$ were <0.1, indicating less phylogenetic signal than expected under BM (Appendix S1: Table S4). Thirty-nine of 74 of binary traits (including qualitative traits coded as binary) showed evidence of phylogenetic signal ($D \leq 0.5$, significantly different from random; Appendix S1: Table S3).

**Observed diversity**

Taxonomic richness was lowest on the northernmost island of Hokkaido, then increased with decreasing latitude until reaching a maximum in southern Kyushu, then decreased again in smaller islands further south (Figure 2A; Appendix S1: Figure S6A). Observed values of PD and FD were highly correlated with richness as expected (Figure 2B, C), showing an initial steep slope that gradually started to level off at higher richness (Appendix S1: Figure S7A, C). Observed values of RPD and RDF were not linearly related to richness; however, they showed high variance at low species richness and low variance at high species richness (Appendix S1: Figure S7B, D).

**Randomization tests for phylogenetic and functional diversity**

Grid cells with significantly low PD were almost all located on the main islands of Japan (Honshu, Kyushu, and Shikoku; Figure 3A). A smaller number of grid cells had significantly high PD, and these were located mostly on small southern islands (Ryukyu, Izu, Ogasawara) or coastal areas (e.g., Kii Peninsula). Grid cells with significantly low RPD were mostly observed in the southern main islands, particularly in and around Kyushu (Figure 3B). Grid cells with significantly high RPD were located both in the small southern islands and northern Honshu and Hokkaido. Results of the randomization test for functional diversity were broadly similar to phylogenetic diversity (Figure 3C, D).

**Relationships between biodiversity and climate or reproductive mode**

All models were significant relative to null models without any fixed effects (Appendix S1: Table S5). The effect of temperature dominated (had the greatest absolute $t$-value) in environmental models (Figure 4A–E), and % apomictic taxa dominated in reproductive models (Figure 4F–G); temperature and % apomictic taxa were significant in each model where they were included, whereas other predictors
(area, precipitation, precipitation seasonality) were not significant in some models. Richness showed a hump-shaped relationship with temperature, reaching maximum values at intermediate temperatures (Figure 5A). Phylogenetic and functional diversity showed either moderately (SES of PD, SES of FD) to strongly (SES of RPD, SES of RFD) hump-shaped relationships with temperature (Figure 5B–E). Standard effect size of RPD showed a clear, linear decline with % apomictic taxa (Figure 5G); a similar pattern was detected in SES of PD, but with wider confidence intervals (Figure 5F). Conditional AIC was lower (indicating better model fit) in the reproductive model compared to the environmental model for SES of RPD, but not SES of PD (Appendix S1: Table S6).

Categorical analysis of neo- and paleoenendism

Nearly all grid cells in islands south of Kyushu (Ryukyu and Ogasawara) have significant levels of phylogenetic endemism, with low (nonsignificant) endemism observed elsewhere (Figure 6). The vast majority of grid cells with significant endemism are mixed or super-endemic. Concentrations of paleoenendism were observed in the Ryukyu archipelago, primarily on Okinawa, Miyako, and Iriomote Islands. Small clusters of grid cells containing significant endemism were also observed northwest of Tokyo (Nagano Prefecture) and on Hokkaido (Figure 6). When we ran CANAPE with a reduced data set including only taxa absolutely restricted to Japan to account for the border effect, there were fewer significant grid cells, but a similar pattern of significant endemism mostly occurring in the southern islands was observed (Appendix S1: Figure S8).

Bioregions

Clustering analysis grouped grid cells into four or five bioregions (Appendix S1: Figure S9) on the basis of taxonomic or phylogenetic distances, respectively. The vast majority of grid cells fell into a subset of these bioregions: based on taxonomic distances, they include bioregion 1 (Hokkaido and high elevation areas of northern Honshu), bioregion 2 (low elevation areas of northern Honshu, southern Honshu, Kyushu, and Shikoku), bioregion 3 (Ryukyu Islands), and bioregion 4 (Ogasawara Islands) (Figure 7A; Appendix S1: Figure S10A). Phylogenetic distances produced similar results, but the Ryukyu and Ogasawara Islands merged into a single bioregion (3), and bioregion 1 covered most of northern Honshu, while extending further south (Figure 7B; Appendix S1: Figure S10B). The other much smaller bioregions (including only one or two grid cells each) are likely artifacts of insufficiently low taxon richness for robust clustering and are not discussed further.

Taxonomic bioregions 3 and 4 had higher mean SES values of PD, RPD, FD, and RFD than bioregions 1 and 2 (Figure 8). Bioregions 3 and 4 also had consistently high PE p-scores, whereas bioregions 1 and 2 had much wider variation (Figure 8E). Bioregion 2 had lower RPD and RFD than bioregion 1. Comparison of biodiversity metrics across phylogenetic bioregions was similar, except that phylogenetic bioregion 3 mostly corresponds to taxonomic bioregions 3 and 4 combined (Appendix S1: Figure S11).
Conservation status and threats

The percentage of protected area in grid cells with significantly high biodiversity ranges from 8.6% to 23.8% total, including 2.8% to 7.0% with high status and 5.8% to 16.8% with medium status (Figure 9A). Cells with significantly high biodiversity had a similar or greater percentage of area protected compared to the baseline rate for Japan at the medium protection status (5.9%) across all biodiversity metrics. In contrast, grid cells with significantly high biodiversity had a similar or lower percentage of area protected than the baseline rate for Japan at the high protection status (3.8%) for most biodiversity metrics. Only grid cells with significantly high PE had a higher percentage area protected (7.0% high, 16.8% medium, 23.8% total) than the baseline rate at both levels of protection.

The range of Japanese deer in Japan increased from 98,916 km² (26.1%) in 1978 to 184,671 km² (48.8%) in 2003 and may be as high as 202,819 km² (53.6%) according to model estimates. The percentage of area occupied by deer in grid cells with significantly high biodiversity increased from 28.2% in 1978 to 42.7% in 2003 when averaged across biodiversity types. Grid cells with high PD, FD, or PE tended to have fewer deer compared to the baseline rate.
However, grid cells with high species richness consistently exceeded the baseline rate for deer presence, with up to 87.4% occupied by deer in the estimated range data set.

DISCUSSION

Distribution and drivers of biodiversity

The latitudinal hump-shaped pattern of species richness observed in ferns here (Figure 2A; Appendix S1: Figure S6A) is well documented (Kusumoto et al., 2017; Ebihara and Nitta, 2019a). Similar patterns have been observed in deciduous broadleaved trees and conifers (Shiono et al., 2015) and niche-modeled distributions of woody plants in Japan (Fukaya et al., 2020). Our spatial modeling analysis indicated that this hump-shaped pattern is strongly linked to temperature, with maximum richness occurring at intermediate temperatures (Figure 5A).

Previous studies have identified temperature, water availability (Bickford and Laffan, 2006; Qian et al., 2012), or a combination of these (Nagalingum et al., 2015; Link-Pérez and Laffan, 2018; Qian et al., 2021) to be important drivers of spatial fern diversity. A hump-shaped pattern of richness along elevation (maximum richness at intermediate elevations) has been frequently observed in ferns on mountains in tropical or subtropical areas (Kessler et al., 2011; Suissa et al., 2021). Hypotheses to explain such a pattern include purely geometrical constraints as well as biological hypotheses positing maximally favorable environmental conditions at mid-elevations (Colwell et al., 2004). Another reason for the greater richness on the main islands compared to southern islands in Japan is that the main islands harbor higher elevational variation that may support a wider range of taxa than is present in the southern islands.

The overwhelming pattern characterizing phylogenetic and functional diversity (PD and FD) is the high diversity of the southern subtropical islands (Ryukyu and Ogasawara Islands) (Figure 3A, C). This pattern is almost certainly due
to the presence of primarily tropical lineages that do not occur in other parts of the country, which lack subtropical climate; 16 genera and four families only occur south of 30.1° latitude (just south of Yakushima). The similarity in patterns of PD and FD is likely due to the moderate degree of phylogenetic signal present in the traits used in our analysis (Appendix S1: Tables S3, S4) and suggests that, at least in the ferns of Japan, phylogeny can be used as a reasonable stand-in for functional diversity. Given their high PD, it is perhaps not surprising that the southern islands also host high amounts of phylogenetic endemism (PE); indeed, the vast majority of grid cells with significant levels of PE, as well as paleoendemic grid cells, are located in the southern islands (Figure 6; Appendix S1: Figure S8). The high PE observed on the southern islands is clearly due to their small land area: small area combined with high PD is expected to lead to high PE.

Another striking pattern revealed here is the predominance of long branches in the northern half of Honshu and the southern islands and the predominance of short branches in southern Honshu, Shikoku, and Kyushu (Figure 3B). The long branches in the southern islands are likely driven by the tropical lineages restricted to this area; those in the north may be due to refugial lineages that are distantly related to others and become more species rich at high latitudes (e.g., Equisetum; Appendix S1: Figure S12A). The preponderance of short branches in southern Honshu, Shikoku, and Kyushu may be due to the radiation of certain species-rich genera such as Deparia (Appendix S1: Figure S12B) and Dryopteris (Appendix S1: Figure S12D).

Furthermore, we identified % apomictic taxa as a strong driver of SES of RPD (Figures 4G, 5G), and to a slightly lesser extent SES of PD (Figures 4F, 5F). Indeed, Dryopteris is one such genus with a high rate of apomictic taxa (Appendix S1: Figure S13), and other genera with high rates of apomixis (Cyrtomium, Pteris, Appendix S1: Figure S13) also reach greatest richness in the southern main islands (Appendix S1: Figure S12). The observation that the % apomictic taxa drives SES of RPD is consistent with the hypothesis that communities with high proportions of asexual taxa have lower PD, and is the first time to our knowledge that this process has been demonstrated in ferns.

FIGURE 5 Relationship between biodiversity metrics and selected predictor variables in the ferns of Japan. (A–E) Environmental models (models including temperature). (F, G) Reproductive models (models including % apomictic taxa). Ribbon shows 95% confidence interval of model. Line fit with the focal predictor variable while averaging over other predictors. For phylogenetic diversity (PD), relative PD (RPD), functional diversity (FD), and relative FD (RFD), the standard effect size (SES) was calculated by comparing raw values to a null distribution of 999 random values, and significance (p-value) determined using a two-tailed test (see Materials and Methods).
We verified that this relationship is also observed when analyzed with a phylogeny with branch lengths in units of genetic change (i.e., the output of the maximum likelihood phylogenetic analysis, before dating), a more direct measure of genetic diversity (Appendix S1: Figure S14). While temperature is also correlated with % apomictic taxa and therefore cannot be discounted, conditional AIC indicates a better fit of the reproductive model than the environmental model for SES of RPD (Appendix S1: Table S6). Ultimately, temperature, in combination with other environmental conditions, is likely driving the predominance of apomictic taxa; Tanaka et al. (2014) also found that the proportion of apomictic taxa reaches a maximum at ca. 34.8°N latitude and increases with temperature in Japan (their study was limited to the main islands and did not include the smaller southern islands). Tanaka et al. (2014) asserted that one limiting factor affecting establishment of apomictic ferns at high latitudes or cold climates is the large cell size of apomicts, which are typically polyploid. As fern gametophytes are only a single cell-layer in thickness with no cuticle or stomata, it is expected they would be particularly susceptible to freezing damage induced by larger cells with higher water content; however, no experimental studies have tested these hypotheses to our knowledge.

Water limitation is thought to play an important role in promoting apomixis in ferns especially in deserts (Grusz et al., 2021), since ferns depend on water for transfer of sperm to egg, and apomictic plants would be able to reproduce in the absence of water. A similar argument has been made to assert that areas with seasonal monsoons are linked to increased rates of apomixis in ferns (Liu et al., 2012;
Tanaka et al., 2014; Picard et al., 2021), but in Japan, although the areas that include high percentages of apomictic taxa are monsoonal, the degree of seasonal water limitation does not approach that of a desert. The monsoon hypothesis was not supported by our results, which showed neither a particularly strong effect of precipitation seasonality in most models (Figure 4), nor a correlation of precipitation seasonality with % apomictic taxa (Appendix S1: Table S2).

The variation in distribution of different biodiversity metrics is reflected by the bioregions analysis, which tended to group grid cells with similar biodiversity values (Figure 8). Interestingly, the bioregions identified here correspond roughly to major vegetation zones previously categorized ad hoc; bioregion 1 corresponds to the cool or subartic deciduous type, bioregion 2 to the warm evergreen type, and bioregions 3 and 4 to the subtropical rainy type (Shimizu, 2014). Furthermore, the border between bioregions 1 and 2 approximately corresponds to the “Gleichenia japonica Line” proposed by Nakaike (1983), which he based on the distribution of Diplopterygium glaucum (Houtt.) Nakai (= Gleichenia japonica Spreng.) as an indicator of evergreen broad-leaf forests in Japan. The agreement between qualitative analysis of bioregions here with previous ad hoc schemes suggests that ferns are useful as bioindicators in Japan. Although some studies have emphasized the importance of deep-sea straits (the Tsugaru Strait separating Hokkaido from Honshu and the Tokara Strait separating the southern islands from Kyushu) in structuring biological communities in Japan (Millien-Parra and Jaeger, 1999; Kubota et al., 2014), they did not play a major role in structuring fern bioregions. The Tsugaru Strait had no relation to any of the bioregions, and although the Tokara Strait somewhat splits phylogenetic bioregions 2 and 3, these regions overlap near Yakushima (Figure 7B).
The weak effect of these straits as barriers makes sense given the presumably high dispersal abilities of ferns, which disperse via tiny spores carried on the wind (Tryon, 1970). Rather, fern bioregions in Japan broadly separate along gradients of temperature and precipitation; in particular, precipitation splits taxonomic bioregion 3 (mostly the Ryukyu Islands; wetter) from bioregion 4 (mostly the Ogasawara Islands; drier) (Appendix S1: Figure S15).

**Conservation status and threats**

We found that grid cells with significantly high amounts of biodiversity are similarly or better protected compared to the baseline protection rate at the medium protection level across all biodiversity metrics, and better than the baseline rate at the high protection level for PE (Figure 9A). The high protection status of grid cells with high PE may be due to an emphasis on conserving endemic plant and animal species in Japanese southern islands such as Irionote and Amami (Japan Ministry of the Environment, 2021a), which include many grid cells with high PE (Appendix S1: Figure S16D). One possible reason for the generally high level of protection overall is that in the past, some ferns have been included in lists of rare and endemic plants for consideration in designation of conservation areas (A. Ebihara, personal communication; Japan Ministry of the Environment, 2021a). Also, (as suggested previously) ferns may serve as bioindicators reflecting overall levels of biodiversity, particularly for other plant groups.

Despite the good news that ferns enjoy relatively strong protection status in Japan, we must bear in mind that there are other threats that conservation zones cannot ameliorate, the most obvious being climate change. One other such threat is herbivory by the native Japanese deer, *Cervus nippon* (Minamitani, 2005; Yahara, 2006; Hattorie et al., 2010; Japan Ministry of the Environment, 2022). The rapid increase in population size observed in surveys from 1978 to 2003 is likely due to a combination of climate change and a decrease in human hunters and natural predators (Takatsuki, 2009). At dense populations, the deer can prevent forest regeneration and result in plant communities dominated by a limited number of species unpalatable to deer (Takatsuki, 2009), although lower population sizes may promote plant diversity as per the intermediate disturbance hypothesis (Nishizawa et al., 2016). They have caused extensive damage even within national parks (Tokita, 2006), which have the highest protection status. Although various efforts to prevent deer herbivory have been deployed such as culling and exclusion fences (Takatsuki, 2009), much work remains to be done to save vulnerable plant populations.

Our analysis shows that the threat to fern biodiversity posed by Japanese deer differs strongly by biodiversity type; although grid cells with significantly high FD, PD, and PE tend to have lower rates of deer occupancy compared to the baseline rate in recent surveys or modeled data, grid cells with high richness experience much greater co-occurrence with deer...
than the baseline rate (Figure 9B). This pattern is due to the fact that Japanese deer do not occur in the southern islands, which harbor high amounts of PD and PE; rather, areas with high species richness are primarily located in southern Honshu, Shikoku, and Kyushu, where deer expansion has been intense (Appendix S1: Figure S17A). This result highlights the importance of using multiple metrics of biodiversity for establishing conservation priorities.

CONCLUSIONS

Our study integrates diverse, thoroughly collected data sets to characterize the spatial biodiversity of Japanese ferns and gain insight into the processes generating these patterns. We conclude by summarizing some caveats that must be kept in mind when interpreting our results, while also suggesting avenues for future research.

First, the data include specimens collected over a wide time period, so they do not necessarily represent the current distribution of ferns in Japan. Furthermore, the collections are not evenly collected throughout the country, but rather some areas have many more specimens than others (Appendix S1: Figure S5A). Indeed, one promising avenue of research may be to take advantage of the most densely sampled areas to investigate changes in abundance over time.

The extreme evolutionary distinctiveness of the southern islands is both a major feature of the Japanese fern flora and a challenge for analysis. The tropical lineages occurring only in these islands are likely responsible for the comparatively low (i.e., clustered) values of PD observed on the main islands (Figure 3A). One way to account for this bimodal distribution pattern would be to use a restricted null model that does not allow all taxa to disperse anywhere within the study area (Mishler et al., 2020). However, ferns are known to have high dispersal abilities (Tryon, 1970; Barrington, 1993; Moran and Smith, 2001). In our data set, the taxon with the greatest latitudinal distribution [Pteridium aquilinum (L.) Kuhn subsp. japonicum (Nakai) A. et S. Löve] spans nearly the entire region (24.3°N to 45.46°N), and 16 taxa have latitudinal distributions spanning the Tokara strait in the south (29°N) to the Togawara strait in the north (41.5°N). Therefore, it seems reasonable to assume that ferns are capable of dispersing across the entire study area. Furthermore, it should be noted that the southern islands are all low altitude (not exceeding 1000 m a.s.l.). If high-elevation habitats were available, the southern islands would be expected to share more species in common with areas farther north. For example, Taiwan, which is just east of the southernmost Japanese islands and harbors high mountains (>3000 m), shares many fern species with the main Japanese islands that are not found in southern Japanese islands. This disjunct pattern supports that the sharp taxonomic turnover between the southern and main islands is more likely due to ecological factors than to dispersal limitation.

One element of biodiversity that we did not consider here is hybrids. The Japanese fern flora includes 374 nothotaxa (Ebihara and Nitta, 2019a). Although hybrids are often thought of as evolutionary “dead ends”, in some cases, they are able to interbreed with diploids and thus contribute to diversification (Barrington et al., 1989). Future studies should consider the distribution of hybrids in Japan and their possible effects on biodiversity. Methods for calculating PD would need to be modified to take into account multiple lineages contributing to one terminal taxon.

Finally, a long under-appreciated aspect of fern ecology is the role of the gametophyte. Unlike seed plants, ferns have gametophytes that are capable of growing independently from the sporophyte. Furthermore, the two stages of the life cycle have vastly different morphology and ecophysiology and may even occur over partially to completely disjunct ranges (reviewed by Pinson et al., 2017). Here, as in the vast majority of ecological fern studies, we considered only the sporophytic stage. However, recent studies in Japan (Ebihara et al., 2013) and elsewhere (Nitta et al., 2017) have revealed different patterns in the community structure of gametophytes and sporophytes. The comprehensive molecular sampling of sporophytes in Japan makes this area ideal for conducting high-throughput DNA sequencing analyses to compare patterns of biodiversity between life stages in ferns across large spatial scales.

AUTHOR CONTRIBUTIONS

J.H.N., B.M., and A.E. conceived the ideas; A.E. provided the data; J.H.N. analyzed the data; W.I. provided resources; J.H.N. wrote the manuscript with input from all coauthors.

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OPEN DATA BADGE

This article has earned an Open Data Badge for making publicly available the digitally shareable data necessary to reproduce the reported results. The data are available at https://doi.org/10.6084/m9.figshare.16655263. Learn more about the Open Practices badges from the Center for Open Science: https://osf.io/tvyxz/wiki.

DATA AVAILABILITY STATEMENT

Code to replicate all analyses, figures, and this manuscript are available at https://github.com/Joelnitta/japan_fernsSpatial_phy. A Docker image to run the code is available at https://hub.docker.com/r/joelnitta/japan_fernsSpatial_phy. Data files needed to run the analysis and selected...
results files are available from Figshare at https://doi.org/10.6084/m9.figshare.16655263.

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SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article. This article includes only online–only Supplemental Data.

Appendix S1. Supplementary tables and figures, including Tables S1–S6 and Figures S1–S17.

Table S1. Spatial autocorrelation in biodiversity metrics (richness, SES of PD, SES of RPD, SES of FD, SES of RFD) and independent variables (environmental variables and % apomorphic taxa) as measured with Moran’s I.

Table S2. Results of modified t-test for correlation between variables while taking into account spatial position.

Table S3. Phylogenetic signal in quantitative (binary) functional traits of the ferns of Japan.

Table S4. Phylogenetic signal in continuous functional traits of the ferns of Japan.

Table S5. Likelihood ratio test (LRT) between full model and null model (model only including the spatial Matérn correlation matrix).

Table S6. Model fit as measured with conditional Akaika information criterion (cAIC).

Figure S1. Species collection curve for the ferns of Japan.

Figure S2. Effect of grain size on sampling redundancy.

Figure S3. Reproductive mode and environmental data in 20-km grid cells across Japan.

Figure S4. Time tree of native Japanese ferns and lycophytes, excluding nothotaxa.

Figure S5. Observed number of specimens and sampling redundancy per 20 km grid cell in the ferns of Japan.

Figure S6. Raw biodiversity metrics of the ferns of Japan plotted by latitude.

Figure S7. Relationships between raw functional and phylogenetic diversity and taxonomic richness in the ferns of Japan.

Figure S8. Phylogenetic endemism of the ferns of Japan measured using CANAPE (categorical analysis of neo and paleoendemism), restricted data set including only taxa endemic to Japan.

Figure S9. Selection of k for bioregions.

Figure S10. Bioregion dendrograms.

Figure S11. Phylogenetic and morphological diversity of the ferns of Japan by phylogenetic bioregion.
Figure S12. Map of species richness for selected genera of ferns of Japan.
Figure S13. Rate of apomixis in genera of Japanese ferns with >1 apomictic taxon.
Figure S14. Relationship between phylogenetic diversity (PD) and relative PD and % apomictic taxa, models inferred using phylogeny with branch lengths in units of expected genetic change (untransformed ML tree).
Figure S15. Scatterplots of grid cell membership in taxonomic or phylogenetic bioregions arranged by mean annual temperature and annual precipitation.
Figure S16. Maps showing overlap of grid cells with significantly high biodiversity for the ferns of Japan and protected areas.

Figure S17. Maps showing overlap of grid cells with significantly high biodiversity for the ferns of Japan and distribution of Japanese deer (*Cervus nippon*).

Appendix S2. Exploratory data analysis for spatial modeling.

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