RESEARCH ARTICLE

Development of spatial models and maps for tree species diversity and biomass in a miombo ecosystem, western Tanzania

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Abstract

Aim: Miombo, a prominent dry forest formation, holds ecological importance for both humans and wildlife. Trees are a driving force behind miombo dynamics, thus, spatially explicit metrics of tree cover are essential for evaluating habitat characteristics, resource availability, and environmental change. We developed predictive models and maps of tree species diversity and biomass within a previously undescribed landscape. **Location:** Mahale Mountains National Park (MMNP), Greater Mahale Ecosystem (GME), Tanzania.

Methods: We created models of tree density, basal area, tree species richness, and tree diversity according to the Shannon Diversity Index. We created a predictive model using an ensemble modeling approach using plot-based data from MMNP and predictor variables derived from satellite data associated with climate, habitat structure, plant productivity, and topography. We assessed predictor importance across models and produced maps based on model predictions and compared them to land cover type and protective status.

Results: Results revealed strong positive correlations between tree metrics ($r \ge 0.70$) and substantial overlap in the selection and relative importance of predictors. Canopy height was the most important predictor across models, followed by climate and topography predictors associated with energy. Predictors derived from the soil-adjusted vegetation index were also valuable. Model performances ranged from R^2 values of 0.45 to 0.55, with tree density performing best. Maps show high tree species diversity and biomass in protected areas.

Conclusions: This study and the maps it produced provide a baseline for land management and future modeling efforts in the GME. Our results highlight the contribution of a wide variety of environmental predictors and the importance of a select few. We confirmed the importance of the current protected area network where conservation efforts align, and help sustain, an abundance and diversity of trees. Current and

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historical disturbance-related predictors should be considered to address remaining unexplained variance.

KEYWORDS

basal area, biomass, ensemble model, Mahale, miombo, satellite data, species diversity, species richness, tree cover, tree density

1 | INTRODUCTION

Miombo is one of Africa's most valuable, threatened, and extensive dry forest formations (Timberlake & Chidumayo, 2011; Ribeiro, 2015). Stretching from Tanzania to Zimbabwe, the miombo ecoregion boundary is primarily determined by an interaction between topography and climate and occurs within a zone that experiences extreme fluctuations in rainfall. Its distinct biodiversity and vast distribution have garnered international recognition, and it is included in the world's top five list of most valuable wilderness areas (Mittermeier et al., 2003). Miombo provides a myriad of services (e.g., nutrient cycling, erosion control) and resources (e.g., food, wood fuels), supporting the approximately 150 million people who live on these lands and in nearby urban settlements (Ryan et al., 2016). This ecosystem is highly threatened by anthropogenic land use activities and climate change, dramatically affecting species distributions and ecosystem processes (Verhegghen et al., 2022). Dry forests like miombo now surpass humid forests (Sunderland et al., 2015) (that have been the subject of far more scientific inquiry and resource allocation) (Schröder et al., 2021) in deforestation rate.

Miombo ecosystems are dominated by tree species from the family Fabaceae in the genera Brachystegia, Julbernardia, and Isoberlinia (Frost, 1996). While they are primarily composed of woodland, these ecosystems are a mosaic of vegetation types, e.g., grasslands, shrublands, dense forests, that vary in size and composition according to environmental factors like soil, water, and topography and anthropogenic factors like land use (Timberlake & Chidumayo, 2011). The dynamics of miombo are driven by the dynamics of trees, which strongly influence vegetation structure, water and nutrient cycling, and fire regimes (Frost, 1996). Data detailing tree cover characteristics such as tree distribution, density, and diversity can help researchers describe land cover types and identify resource-rich areas, as well as assess the health and productivity of the ecosystem to the benefit of both wildlife and humans (Torres & Lovett, 2013; Ribeiro, 2015; Ryan et al., 2016). The importance and vulnerability of miombo and its resources has garnered attention (Syampungani et al., 2009), but inter-site variability warrants expanding research into understudied regions and localities (FAO, 2019; Schröder et al., 2021).

One such understudied region is the Greater Mahale Ecosystem (GME) in western Tanzania. The Tanzania miombo represents the most northern and eastern boundary of the miombo ecoregion, yet it is largely separated from most of the ecoregion due to the expansion of cropland (Timberlake & Chidumayo, 2011). The GME (20,000 km²) is part of this ecoregion and the Albertine Rift, and

encompasses crucial habitat for threatened species and plays a vital role in regional biodiversity and ecosystem services (Plumptre et al., 2007; Bietsch et al., 2016). The land is managed under a variety of designations that include protected and unprotected areas. Mahale Mountains National Park (MMNP), the only national park within the GME, exhibits low human activity and protects key habitat for the biodiversity of the region (Plumptre et al., 2007; Chitayat et al., 2021). Outside MMNP the landscape faces threats from accelerated human population growth and expanding land use practices (e.g., agriculture, charcoal production) that are driving habitat loss and degradation and threatening biodiversity in the region (Pintea, 2012; Thomsen et al., 2023). Tree cover within the GME has yet to be comprehensively evaluated and described, and because of the importance of tree cover in miombo dynamics, this is a significant gap in our understanding of this landscape and the broader miombo ecosystem. One reason this has not yet been accomplished is due to the physical and logistical challenges involved in surveying the vast miombo woodlands of the GME, compounded by the fact that historical research efforts in western Tanzania have primarily concentrated on chimpanzees at established long-term field sites (Nakamura, 2012; Wilson, 2012; Piel et al., 2019).

Satellite remote sensing technology has emerged as an invaluable tool for alleviating the limitations of ground surveys, providing spatial-temporal data that can be used to identify and describe biophysical landscape attributes like trees (Kerr & Ostrovsky, 2003). Moreover, as satellite data is often collected continuously and is freely available to the public, it provides accessible, up-to-date information that can be easily integrated into monitoring frameworks. Spatial predictive models can be used to connect satellite data with field data to make broad-scale predictions that expand the utility of in-situ observations (McNellie et al., 2021; Barreras et al., 2023). These predictions can be used to generate continuous maps that serve as valuable communication tools and can also inform land management decisions, guide conservation efforts, and facilitate infrastructure development projects. Spatial models can also aid in the identification of current and historic landscape conditions, uncovering patterns that aid our understanding of ecosystem dynamics and predicting changes that may arise (Jinga & Palagi, 2020).

The GME is vital for biodiversity, especially threatened species, in the miombo ecoregion. Here, we aim to fill a crucial gap in our understanding of tree cover in the GME with the development of important tools that can describe tree cover characteristics related to species diversity and biomass in a miombo. We conducted field surveys in MMNP where the minimal human impact allows us to

2 | METHODS

2.1 | Study area

The GME is an ecologically rich area that boasts a variety of distinct species and those of conservation concern, including eastern chimpanzees (Pan troglodytes schweinfurthii), African elephants (Loxodonta africana), and mninga trees (Pterocarpus angolensis) (Plumptre et al., 2007; Thomsen et al., 2023). Situated alongside Lake Tanganyika, the health of this ecosystem is inherently connected to the lake, which is significant for its ecosystem services and its diverse aquatic life, hosting 300 endemic species of fish (Sweke et al., 2013). The GME exhibits topography consisting of broad valleys interspersed by steep mountains and flat plateaus, ranging in elevation from 780 to 2460 m above sea level (Carvalho et al., 2022). MMNP protects 1517km² of land and encompasses the GME's highest peaks. The region is highly seasonal, experiencing a rainy season from October to mid-May and a dry season from mid-May to September. Annual precipitation ranges from 900 to 2100mm, and temperatures vary from 11 to 38°C.

2.2 | Response variables – Tree cover characteristics

We collected tree data in 463 plots across MMNP from March 2018 to January 2019 (Figure 1) as part of an extensive study on chimpanzees (Chitayat et al., 2021). Plots were placed along transects at 12 randomly selected sites where we established eight transects, each measuring 1 km in length. These transects featured plots measuring 5 m × 100 m, spaced at 100 m intervals. Within each plot, all trees with a diameter at breast height (1.3 m) \geq 10 cm were identified and measured.

We calculated tree cover characteristics at the plot level that were used as response variables in the development of four separate predictive models: (1) tree density, (2) basal area of trees, (3) tree species richness, and (4) tree species diversity according to the Shannon Diversity Index. Tree density and basal area are both useful biomass proxies (Torres & Lovett, 2013): density provides information about the number of trees per unit area, while basal area quantifies the volume of trees per unit area according to the diameter at breast height. Similarly, species richness and the Shannon index offer different measures of species diversity: species richness counts the number of species per unit area, while the Shannon index considers both the number of species and their relative abundances in its calculation (Shannon, 1948; Whittaker, 1972). We selected these characteristics because they provide valuable and complementary information about forest structure and composition. Non-parametric Spearman rank correlation coefficient tests were conducted to establish the relationship between all pairs of tree cover characteristics. We tested for spatial autocorrelation using Moran's *I* test and found no statistically significant spatial autocorrelations (p > 0.05) for any response variables (Zuur et al., 2010).

2.3 | Predictor variables – Satellite remote-sensing data

We utilized environmental predictor variables derived from remotesensing satellite data to build models of tree cover characteristics in the GME. Tropical dry forests can be challenging to model using satellite data owing to the pronounced temporal and spatial fluctuations of these ecosystems (Bastin et al., 2017; Verhegghen et al., 2022); however, incorporating a wide array of biotic and abiotic predictors can help improve the quality of model outputs (Slik et al., 2010; Zellweger et al., 2015). The predictors were categorized into four groups: (1) climate, (2) habitat structure, (3) plant productivity, and (4) topography (Table 1).

We tested several predictors derived from climatic data, including mean annual temperature and precipitation from the period of 1979-2013 (Karger et al., 2017) and mean annual actual and potential evapotranspiration from the period of 1950-2000 (Zomer et al., 2006). These variables influence the availability of crucial resources like energy and water (White, 1983; Ribeiro et al., 2020) and are often used for modeling vegetation in miombo (Pearson et al., 2006; Slik et al., 2010; Zellweger et al., 2015).

Canopy heights (m) were used to represent habitat structure (Potapov et al., 2021). This variable has demonstrated previous success for modeling tree cover characteristics (Wolf et al., 2012; Knapp et al., 2020).

To represent plant productivity, we included a variety of vegetation indexes (VI) that were derived from Sentinel-2. We used L1C data for its higher spatial resolution and a cloud-masking technique was applied to remove pixels affected by clouds and cirrus using the QA60 band (bits 10 and 11). While surface reflectance data (L2A) are commonly used, atmospheric correction is not always required for classification applications, and vegetation indices such as the normalized difference vegetation index (NDVI) and soiladjusted vegetation index (SAVI) reduce atmospheric effects (Song et al., 2001; Verhegghen et al., 2022). Researchers commonly utilize VIs to model trees because their capacity to quantify greenness is effective for assessing plant abundance and variability (Pettorelli et al., 2005; Timberlake et al., 2010; Barati et al., 2011; Mutowo & Murwira, 2012; Zellweger et al., 2015; Cavada et al., 2017; Mayes et al., 2017). For each VI, we calculated summary statistics using all of the 2018 values, the study period, which served as our predictors



FIGURE 1 Locator map of the study area in western Tanzania. Maps show (a) the location of the Greater Mahale Ecosystem (GME) in Tanzania, (b) a digital elevation model of the GME, (c) land cover types in the GME, and (d) the location of vegetation plots sampled across Mahale Mountains National Park (MMNP).

(Cabacinha & de Castro, 2009). For example, maximum (max) value was the highest observed value from 2018 and the mean value was the mean of all 2018 values.

Topographic predictors were used to describe topographic position and heterogeneity (Amatulli et al., 2018; Mattivi et al., 2019) and obtained from the Shuttle Radar Topography Mission (SRTM). These features can influence tree cover characteristics through associations with local microclimates, soil properties, and water accumulation (de Castilho et al., 2006; Thuiller et al., 2006; Engelbrecht et al., 2007; Homeier et al., 2010; Slik et al., 2010; Fricker et al., 2015).

Satellite data were collected at resolutions ranging from 10 to 1000m (see Table 1). Data were resampled, scaling either up or down, to a uniform resolution of 100m using conventional bilinear interpolation (Divíšek & Chytrý, 2018). This was done to mitigate potential challenges from spatial resolution disparities like data compatibility and model complexity (Moudrý et al., 2019).

2.4 | Predictor variable selection process

To create the most parsimonious predictive models, we applied a variable selection process to remove predictors that provided little

predictive capacity or were highly correlated based on a correlation coefficient of $r \ge |0.7|$ (Jantz et al., 2016). First, we utilized recursive feature elimination to remove predictors of low importance (David et al., 2022). Then, using Pearson product-moment and Spearman rank correlation coefficient tests, we identified pairs of predictors that were highly correlated (Zuur et al., 2010) and retained the predictor with higher relative importance according to recursive feature elimination rankings. Finally, we tested for multicollinearity using a variance inflation factor (VIF) set to 10 or more. This variable selection process was used for each response variable and subsequent model. We only included the selected predictors in future analyses and the building of ensemble models. A spatial evaluation of the calibration range of selected predictors showed that our training data were able to capture the majority of conditions across the GME but that coverage varies between predictors (Appendix S1).

2.5 | Ensemble model development

To build and verify the models, each data set was divided into training (80%) and testing (20%) subsets. We employed an ensemble modeling approach that combines multiple individual base

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Predictor variable	Description	Mean value (range)	Model	Source
Climate	•			
Rain	Mean annual precipitation (mm) ^a	1369 (1042–1828)	ВА	CHELSA
Temperature	Mean monthly temperature (°C) ^a	21.5 (15.7–23.7)	R. SDI	CHELSA
AET	Mean annual actual evapotranspiration ^b	982 (392–1334)	D, BA, R, SDI	CGIAR-CSI
PET	Mean annual potential evapotranspiration ^b	1709 (1351-1862)	BA	CGIAR-CSI
Habitat structure				
Canopy height	Canopy height (m) ^c	9.3 (0-25.9)	D, BA, R, SDI	GLAD
Productivity				
NDVImax	Maximum (max) normalized difference vegetation index (NDVI) ^d	0.78 (0.69-0.84)	ВА	S2
NDVImean	Mean NDVI	0.34 (0.26-0.45)	BA	S2
RGImean	Mean red-green index (RGI) ^e	0.11 (0.01-0.25)	D, BA, R, SDI	S2
RGIstdv	Standard deviation (stdv) of RGI	0.09 (0.04-0.14)	D, BA, R, SDI	S2
RRmean	Mean reflectance ratio (RR) ^e	1.29 (0.74–1.59)	D, BA, R, SDI	S2
RRstdv	Stdv of RR	0.20 (0.07-0.99)	BA	S2
SAVImax	Max soil-adjusted vegetation index (SAVI) ^f	1.17 (1.03–1.26)	D, R, SDI	S2
SAVImean	Mean SAVI	0.51 (0.38-0.78)	D, R, SDI	S2
SAVIstdv	Stdv SAVI	0.10 (0.03-0.17)	D, BA, R, SDI	S2
SLAVImax	Max specific leaf area vegetation index (SLAVI) ^g	1.69 (1.31–2.28)	D, BA, R, SDI	S2
Topography				
Elevation	Elevation (m a.s.l.) ^h	1204 (797–2212)	D	SRTM
Ruggedness	Terrain ruggedness index ^h	2.0 (0.7-6.5)	D	SRTM
Roughness	Topographic roughness index ^h	15.6 (2.6–53.7)	BA, R, SDI	SRTM
ТРІ	Topographic position index ^h	0.04 (-2.34-3.55)	BA	SRTM
TWI	Topographic wetness index ⁱ	7.2 (4.7–12.7)	BA	SRTM

Note: Individual models were developed for tree density (D), stand basal area (BA), tree species richness (R), and tree diversity according to the Shannon Diversity Index (SDI). Data sources include the CGIAR-CSI Global-Aridity and Global-PET Database (CGIAR-CSI) (1000m), Climatologies at high resolution for the Earth land surface areas (CHELSA) (1000 m), Copernicus SENTINEL-2 (S2) (10 m), Global Land Analysis and Discovery (GLAD) (30 m), and Nasa's Shuttle Radar Topography Mission (SRTM) (30 m).

^aKarger et al. (2017). ^bZomer et al. (2006). ^cPotapov et al. (2021). ^dPettorelli et al. (2005). ^eCavada et al. (<mark>2017</mark>). ^fMutowo and Murwira (2012). ^gBarati et al. (2011). ^hAmatulli et al. (2018). ⁱMattivi et al. (2019).

models into a single predictive ensemble model that capitalizes on the collective strength of its components to improve model robustness and reduce uncertainty (Van Der Laan et al., 2007). Ensemble models were developed separately for each tree cover characteristic using the package SuperLearner and a 10-fold cross-validation risk. Three base modeling methods were used to develop ensembles: a regression method (multivariate adaptive regression splines, MARS) and two machine-learning methods (gradient boosted model, GBM, and random forest, RF). All methods account for non-linear relationships and automatically

consider interactions between variables while avoiding overfitting (Leathwick et al., 2006; Zellweger et al., 2016).

For our ensemble models, we included the default model and several tuned models of each base modeling method, which can increase ensemble model performance by leveraging the strengths of different and diverse model configurations (Polley & Van Der Laan, 2010). We tuned base models according to critical hyperparameters that can enhance prediction accuracy and reliability (Appendix S2) (Yates et al., 2023). Tuning was performed using a 10fold cross-validation to ensure robust model performance and avoid



FIGURE 2 Importance of predictor variables used to model individual tree cover characteristics in the Greater Mahale Ecosystem. Plots show a 95% confidence interval around the mean squared error (MSE).

overfitting. The optimal values were determined through a grid search over specified ranges, allowing for systematic exploration of parameter combinations.

We present three statistical metrics to evaluate the performance of our ensemble models: the correlation coefficient (r) depicting the relationship between predicted and observed values, the percentage of explained variance (R^2), and the percentage of root mean squared error derived from a 10-fold cross-validation resampling procedure with data shuffling. Higher r and R^2 values and lower root mean squared error values signify superior model performance. We assessed the importance of predictor variables by measuring their contributions to reducing cross-validated risk through mean squared error.

2.6 | Mapping spatial model predictions

We used ensemble models to generate spatially explicit predictions and produce individual maps of tree cover characteristics. Maps showing the uncertainty of model predictions were also generated for each tree cover characteristic by quantifying the standard deviation of prediction results from all individual models within the ensemble. We created a composite map aggregating all tree cover characteristics by summing each raster cell's normalized predicted values, scaled from 0 to 1. Then, we evaluated the distribution of composite values against land cover designations as determined by the Climate Change Initiative and European Space Agency (CCI/ ESA) land cover map for 2020 (Zanaga et al., 2021). All analyses were performed in R (R Core Team, 2022) and QGIS 3.6 (QGIS Development Team, 2020).

3 | RESULTS

3.1 | Ensemble models of tree cover characteristics

Tree cover characteristics varied between plots, ranging from 0 to 37 for tree density (mean = 9 \pm 7), 0 to 5.1 for basal area (mean = 0.5 \pm 0.5), 0 to 16 for tree species richness (mean = 5 \pm 3), and 0 to 2.6 for Shannon index (mean = 1.2 \pm 0.7). Pairwise correlation tests revealed all tree cover characteristics to be highly correlated with each other (r=0.70-0.97), with the strongest relationship between tree species richness and the Shannon index.

Each model included at least one predictor variable from each category (e.g., climate). Several predictors were selected for basal area that were absent from other models (e.g., precipitation, NDVImean). Other than these predictors for basal area, the selection and relative importance of predictor variables showed substantial overlap among response variables. This was especially true between species richness and the Shannon index models that included the

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same predictors and displayed the same top three most important (Figure 2). Canopy height consistently demonstrated the highest mean importance across ensemble models. Temperature, potential evapotranspiration, and elevation emerged as the second most important predictors across models. These predictors also exhibited strong positive correlations with each other that prevented their inclusion within the same model.

Performance scores varied between ensemble models of tree cover characteristics, with R^2 scores ranging from 0.45 to 0.55 and our model for tree species density performing best across metrics (Table 2). Additionally, predicted values were all significantly correlated with their observed values (p < 0.001).

3.2 Maps of tree cover characteristics in the GME

We expanded model predictions to produce maps of tree biomass and diversity in the GME (Figure 3), which exhibited generally low levels of uncertainty (Figure 4). Higher uncertainty values are predominantly observed in areas with known human settlements along the coast of Lake Tanganyika and the north-central region of the GME around the Mishamo refugee camp, as well as along the highest elevation peaks in MMNP. Similarities and divergences between tree cover characteristics can be readily observed in our composite map (Figure 5), such as high-value areas along the western side of MMNP. Nearly half of the GME is under some protective status (49%), where the majority of the highest and lowest composite values occur, 64% and 66%, respectively. Composite values were disproportionately distributed across land cover types, with results showing that sites dominated by stretches of forest represent only 1% of the landscape but hold 10% of the highest 25% of composite values (Table 3). Similarly, results showed that the lowest 25% of composite values occur in primarily non-tree-dominated, natural or human-modified land cover types, such as grasslands and croplands. Additionally, the land cover classification system was sometimes inaccurate within the study area. For example, some misclassifications occurred in areas that are known to be occupied by human settlements and croplands but were classified as land cover types such as grasslands or shrublands.

DISCUSSION 4

We modeled and made predictions of tree species diversity and biomass for the GME by using an ensemble modeling framework,

TABLE 2 Performance metrics of individual ensemble models for tree cover characteristics related to tree species diversity and biomass.

available remotely sensed satellite data, and plot-based survey data of trees in MMNP. Individual ensemble models achieved moderate performance scores, with our models of tree density and tree species richness performing best. High correlations between observed and predicted values for all response variables suggest that models could generalize well on unseen data. Maps that show the extension of model-based predictions allow for visualizing tree cover characteristics across the GME landscape.

Relative importance of predictor variables 4.1

The diverse environmental predictors we tested showed substantial overlap during the predictor selection process and in relative importance among models. This likely stems from the strong positive correlations between tree cover characteristics similarly affected by environmental factors like soil conditions (Homeier et al., 2010). Canopy height was the most important predictor for all models, which aligns with previous research showing that canopy height can account for a significant portion of the observed variability in tree species diversity and biomass metrics in tropical forests on local scales (Wolf et al., 2012; Knapp et al., 2020). The importance of canopy height may also correspond to structural contrasts between GME vegetation types. Some areas, like grasslands, are not treedominated and exhibit low tree biomass; conversely, areas where trees dominate the landscape display discernible canopy structures. Ogawa et al. (2007) observed differences in canopy height between vegetation types in the GME, showing that forested sites, predominantly located in valley bottoms, have a greater density of tall trees than woodlands. Similar findings have been documented for other sites, showing tree height, basal area, and richness decrease from valleys to plateaus (Homeier et al., 2010; Rodrigues et al., 2020). This pattern may stem from local differences in soil water and nutrient availability that alter plant growth and can impact species coexistence (Ryan & Yoder, 1997; Homeier et al., 2010). Canopy height can vary across different vegetation types under the characteristics of the species found therein, which display minimal overlap in MMNP (Chitayat et al., 2021).

The strong positive correlations between temperature, potential evapotranspiration, and elevation and their concurrent high importance across models suggest that a common underlying factor like atmospheric energy availability may indirectly drive their value (Hawkins et al., 2003; Fisher et al., 2020). Energy availability is known to influence plant growth, reproduction (Dong et al., 2012),

Response variable	Correlation coefficient (r)	Root mean squared error (RMSE)	Explained variance (R ²)
Tree density	0.7	16%	0.55
Tree stand basal area	0.7	19%	0.47
Tree species richness	0.7	19%	0.54
Tree species diversity (Shannon Diversity Index)	0.7	23%	0.45



FIGURE 3 Predictions of tree cover characteristics across the Greater Mahale Ecosystem. Predictions were generated from individual ensemble models developed using plot-based vegetation data from Mahale Mountains National Park (outlined in black) and satellitederived predictors. The color gradient legend applies to all maps.

and species diversity (Fisher et al., 2010). Conversely, water availability is often shown to drive plant dynamics in the tropics (Hawkins et al., 2003; Siyum, 2020); however, water-related predictors were mostly excluded from model development (i.e., topographic wetness index [TWI], precipitation). Precipitation was included in our model of basal area but demonstrated only modest importance toward predictions. This aligns with the results of (Barreras et al., 2023), which showed that mean precipitation held relatively low importance in ensemble models for tree density and height, especially in dry ecosystem types. Actual evapotranspiration was the only water-related predictor chosen across response variables, but it consistently displayed only modest importance. Actual evapotranspiration's broad connection to water and energy dynamics might render it a somewhat simplistic index of the environmental factors affecting plant growth (Fisher et al., 2010), allowing it to be widely applicable but restricting its explanatory power. However, the spatial resolution of our climatic data scale may have also influenced the effectiveness of these predictors in describing tree cover dynamics. These data were upscaled from a coarser resolution of 1000m to 100m, which redistributes the data across a finer grid but does not add any new information. Coarse data have clear benefits for model development, reducing model size and complexity, but course climatic data can also introduce errors when there is a discrepancy between the climate used in analysis and that experienced by organisms and the habitat on the ground (Bütikofer et al., 2020). Particularly in heterogeneous and mountainous regions where local microclimate conditions can differ dramatically, fine-scale data sets may be more

useful. Future modeling efforts may benefit from exploring the use of higher-resolution climatic data, diminishing the risk of missing critical small-scale variations.

Multiple SAVI-based predictors were selected and showed importance among models. SAVI provides advantages over other VIs given the GME's high seasonality and predominantly open vegetation, leading to significant spatial and temporal soil exposure and reduced vegetative coverage (Huete, 1988). Moreover, SAVI's ability to mitigate the influence of soil brightness is valuable in fireimpacted landscapes, like the GME, where exposed and charred soil leads to distinct soil-vegetation contrasts (White & Swint, 2014; Meng et al., 2017). In Tanzania, fire events occur annually even in PAs, and are caused mainly by anthropogenic ignitions for activities such as farm preparation, cattle grazing, hunting, honey harvesting, and charcoal production (Kikula, 1986). PAs like MMNP are also subjected to prescribed burns managed by authorities, which are generally employed to protect the ecosystem under a fire management plan. In miombo, fire plays a pivotal role in shaping tree species composition and is often linked to a decline in tree biomass (Frost, 1996; Chidumayo, 2013).

The importance of VI-based predictors differed between models of tree species diversity and models of tree biomass. RGI (redgreen index) mean was found to be more important in models of tree biomass, likely due to its association with tree crown size which is related to tree size (Cavada et al., 2017). In addition, when predicting the biomass of long-lived species, the mean may be a more relevant metric as it smooths out the short-term fluctuations and FIGURE 4 Maps showing the level of uncertainty of ensemble model predictions for tree cover characteristics across the Greater Mahale Ecosystem, with Mahale Mountains National Park outlined in black. The color gradient applies to all maps and provides the relative level of prediction uncertainty (standard deviation) from low to high.



extremes inherent in metrics such as standard deviation or maximum. Conversely, SAVI standard deviation encapsulates temporal variation in vegetation greenness and demonstrates greater importance in models of tree species diversity. Temporal variations reflect seasonal changes influenced by tree species compositions, distributions, and phenological characteristics (e.g., deciduous). This predictor gains significance by capturing these differences and the occurrence of disturbance events like fires, which often result in dramatic changes to greenness, and connecting them to spatial differences in tree diversity.

4.2 | Spatial predictions across the GME

Model predictions demonstrated spatial similarities between tree cover characteristics, revealing overlapping areas of high or low values. This may reflect stages of ecological succession as the GME experiences annual fires (Naftal et al., 2022) and successional dynamics play a crucial role in shaping vegetation structure and composition (Ouyang et al., 2016), particularly in areas recovering from or experiencing disturbance events (Kalaba et al., 2013; Saito et al., 2014). The observed pattern between tree characteristics may also indicate a potential functional relationship between diversity and biomass, whereby diverse tree communities encourage tree growth through more efficient resource utilization (Grossman et al., 2018). Similar patterns can be expanded to other taxa, with tree diversity and abundance identified as key drivers of taxon-level (e.g., bats, birds, insects) and forest-associated biodiversity in previous studies (Harvey et al., 2006; Ampoorter et al., 2020). Further investigation on this topic is required as the relationship between biomass and biodiversity is often complex and non-linear, at times exhibiting a hump-shaped (unimodal) pattern that reveals a decline in biodiversity at the highest biomass levels (Graham & Duda, 2011). In a woodland-forest mosaic in eastern Tanzania, Shirima et al. (2015) observed a monotonically increasing trend between tree species richness and biomass in miombo woodland, but a unimodal pattern in montane forest. This suggests that there could be an optimal level of biomass that maximizes species diversity in montane forests but not miombo woodland (Shirima et al., 2015).

Forests, particularly non-riverine forests that stretch out over large areas, are rare in the GME. We observe that high tree species diversity and biomass occur disproportionately more in forest and mixed woodland-forest land covers. The highest concentration of high-value predictions can be found along the lakeshore of MMNP, which encompasses large swaths of miombo woodland and the largest forest block in the region. This finding supports the well-established importance of northwestern MMNP, which is known to support relatively unique species assemblages and high densities of species like chimpanzees (Plumptre et al., 2007; Chitayat et al., 2021). PAs in East Africa were historically established with large, charismatic mammal species in mind, yet these efforts can lead to broad advances that extend beyond the original intention, such as biodiversity conservation or carbon sequestration (Banda et al., 2006; Dickson et al., 2020). Outside MMNP, we see high values clustered within the Ntakata



TABLE 3 Percentage of composite values of tree cover characteristics occurring in different land cover types.

	% GME landscape	Tree cover composite values			
Land cover type		Low quartile 1	Med-low quartile 2	Med-high quartile 3	High quartile 4
Cropland, human settlement	5%	10%	14%	2%	1%
Grassland, wetland	5%	32%	7%	3%	4%
Natural mosaic	10%	32%	16%	6%	3%
Woodland (15%–40% trees)	40%	14%	41%	42%	26%
Woodland, forest mix	39%	12%	22%	46%	56%
Forests (≥40% trees)	1%	0%	0%	1%	10%

Note: Land cover types may encompass multiple vegetation types, for example, natural mosaics can include bamboo thicket, shrubland, and miombo woodland. Composite values are divided into equal interval quartiles, with quartile 1 containing the lowest 25% of values and quartile 4 the highest 25% of values calculated across the GME.

forest reserve. This area also possesses significant ecological importance, providing sanctuary for endangered and endemic species and corridor habitat for elephants and chimpanzees beyond the boundaries of national parks (Plumptre et al., 2007; Lindsey et al., 2020). This reserve is part of the Ntakata Mountains REDD project, spearheaded by Carbon Tanzania (Dickson et al., 2020).

The composite map highlights the importance of the PA network in the GME for conserving tree species diversity and biomass. However, there is clear variation both within and between PAs likely stemming from multiple factors. This includes the natural heterogeneity of the GME and occurrence of vegetation types that display low tree cover, such as grasslands, and thus low composite values. Additionally, there are differences in when PAs were established, the history of land use across the GME, and PA designation (e.g., national park, village forest reserve), which dictates restrictions on human activity and often the availability

of resources to enforce those restrictions. This results in a spectrum of conservation outcomes, showcasing the value of longestablished and strictly regulated PAs like MMNP, as well as those occurring in areas with a history of sustainable land use practices like Ntakata forest reserve. Comparatively, those with less restrictive protections or lacking resources may fail to prevent habitat loss and degradation (Itoh et al., 2012; Thomsen et al., 2023), or may simply require more time to benefit from their recent changes to their status and resources, such as the Tongwe West Forest reserve.

4.3 | Model improvements

The performance scores of our models demonstrate unexplained variance that may result from the scale of the predictors used or a lack of predictors related to current and historic environmental disturbances. After MMNP was gazetted in 1985, authorities prohibited human activities except those related to tourism and research. However, illegal activities (e.g., wood harvesting, medicinal plant collection) and human encroachment into the park can be challenging to manage and vary over time. Miombo demonstrates the ability to recover from human activities, but the process can take decades, and recurrent disturbances can significantly impede the recovery rate (Williams et al., 2008; Chidumayo, 2013; Kalaba et al., 2013). Because of these dynamics, Ribeiro (2015) argue that land use data and land cover recovery rates are valuable for modeling vegetation patterns in miombo.

Future models may also benefit from the inclusion of fire-related predictors that go beyond the scope of the VI predictors we included. Fire has a well-documented influence on tree cover in miombo woodlands (Frost, 1996) and studies indicate that fire characteristics like frequency, intensity, and timing can have varied effects on trees and tree communities (Ryan & Williams, 2011; Chidumayo, 2013; Mapaure, 2013; Saito et al., 2014; Tarimo et al., 2015; Buramuge et al., 2023). While fire has the potential to open up areas for colonization and promote diversity through the establishment of new species, frequent and intense fires can reduce diversity and biomass by favoring fire-tolerant species and suppressing tree growth because of mortality or damage to the stems, bark, and roots (Ribeiro, 2015; Buramuge et al., 2023). Moreover, disturbance impacts can be synergistically influenced by additional factors like large herbivores (Mapaure, 2013; Ivory & Russell, 2016). Improved model accuracy may require the integration of novel variables that, independently or in synergy with other variables, can help explain the spatial variability of tree cover characteristics and enhance the reliability of models.

Lastly, to capture a greater diversity of environmental conditions across the GME and enhance the reliability of predictions, future efforts should incorporate data from areas outside MMNP and protected zones. By using data solely from MMNP in the current study, we lacked data from sites where human activities like agriculture or cattle herding occur—practices prohibited in MMNP but - Applied Vegetation Science 🛸

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widespread outside the GME – that are known to impact tree cover characteristics (Chidumayo, 2013; Macave et al., 2022; Buramuge et al., 2023).

5 | CONCLUSIONS

This study provides a crucial baseline for the development of spatial models and tree maps in the heterogeneous miombo landscape of the GME, addressing a previously lacking area of research. The challenges encountered underscore the complexity of modeling trees in this landscape. Our results emphasize the significance of the existing PA network where high composite values align with unique species assemblages and conservation efforts. The strong positive correlations found between tree cover characteristics suggest a potential functional relationship between tree diversity and biomass, which may have implications for broader biodiversity conservation efforts. This finding warrants further investigation as it may yield insights into biodiversity and biomass patterns across taxa that can inform broad monitoring and conservation strategies. We are cautious in interpreting some results due to methodological limitations and encourage future research to incorporate additional explanatory factors and data from areas outside protected zones to enhance the reliability of estimates. Given the environmental impacts we can expect to occur as a result of climate change and human population growth (Warren et al., 2018; Elisa et al., 2024), there is an urgent need to expand and refine spatial products and tools that can help monitor any changes and guide the effective management of the GME.

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DATA AVAILABILITY STATEMENT

The data and code that support the findings of this study are openly available in figshare (https://figshare.com/) at https://doi.org/10. 6084/m9.figshare.c.6874501.

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SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.

Appendix S1. Maps showing the calibration range of predictor variables used in ensemble models of tree cover characteristics.

Appendix S2. Grid search space for hyperparameter tuning of ensemble models of tree cover characteristics.

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