

Research Article

Edge-Interior Disparities in Tree Species and Structural Composition of the Kilengwe Forest in Morogoro Region, Tanzania

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A survey to determine the variation in species and structural composition of trees along the edge-interior gradient was done in the Kilengwe forest in Morogoro region, Tanzania. The forest was categorized into three habitats, namely, edge (0–100 m), intermediate (100–200 m), and interior (>200 m) depending on the distance from the forest margin. A total of six plots of 0.04 ha each were randomly placed in each of the habitats whereby all trees with DBH ≥ 10 cm were inventoried. A total of 67 species representing 26 families were recorded. Fabaceae was the most speciose and abundant family. *Brachystegia spiciformis* was the most abundant species. Of the recorded species, 10.45% were common in the three habitats while 8.95%, 13.43%, and 26.86% occurred exclusively to the edge, intermediate, and interior habitats, respectively. The forest interior was significantly rich in terms of species richness, diversity, density, and basal area than the edge and intermediate habitats. The edge had significantly higher number of stumps/ha. In summary, the results suggest that edge/intermediate and interior are contrasting habitats in terms of tree species richness, diversity, and structural composition. Moreover, the forest edge and intermediate habitats were found to be characterized by high anthropogenic activities compared to the forest interior habitat.

1. Introduction

Forest fragmentation and deforestation are among the critical environmental problems with possible implications on a global scale [1, 2] as human activities convert continuous forests into number of patches [3]. The tropical forests that encompass 6% of the world's land area and harbor at least 50% of the world's biodiversity are deforested and fragmented at an alarming rate [4, 5]. For instance, the global annual losses of 5.2×10^6 ha/year and 8.3×10^6 ha/year due to fragmentation and deforestation in the tropical forest area were recorded between 1990–2000 and 2000–2010, respectively [6]. In Tanzania, the annual loss of forest cover increased by 37% from the period of 1990–1995 (322,000 ha/year) to 2000–2010 (403,000 ha/year), which was mainly due to increased demand as the population grows [6, 7]. The forest fragmentation process disrupts structure and spatial continuity as it reduces original area, increases edge formation, and

isolates remaining forest patches [3, 8]. The formation of edges is recognized to be fundamental of ecological change as it involves alteration of microclimatic conditions, enhances invasion by exotic species, and increases the human pressure [9, 10]. Due to this, several organisms, for instance, mammals, amphibians, birds, and tree communities, have been reported to suffer significant changes in their local abundance and distribution in the forest fragments [11–14].

Various studies done in the tropical forests have reported negative impacts of forest fragmentation, mainly associated with the edge effects. These impacts include (1) reduction in recruitment rates of trees due to habitat desiccation and seedling damage by litter and tree falling near forest edges [11], (2) increasing sapling mortality rate by competition with lianas, vines, and ruderal species [15], and (3) increased mature tree mortality due to increased rates of uprooting and breakage near forest edges [2, 15, 16], which results in the decrease in canopy height [17, 18]. Due to the above explained

effects and other edge-related processes, it is reasonable to expect that tree species in the forest edges will differ markedly from the forest interior in terms of species richness, diversity, and structure as well as ecological and taxonomical composition [12, 19, 20].

The Eastern Arc and Coastal forests (Kilengwe inclusive) of Tanzania are known to be centers of endemism, but the forests are facing danger of losing some of these due to increased anthropogenic activities as a result of population growth and fragmentation [21–23]. Thus, determining how tree species diversity and abundance vary within forests is a vital step in plant ecology and biodiversity conservation [9]. In Tanzania, no known study has examined the edge-interior differences in the tree species and structural composition in any forest. Thus, due to the existence of this knowledge gap, this study intends to (1) provide an understanding of the existing knowledge discrepancy by comparing the species richness, diversity, abundance, and basal area of tree communities along edge-interior gradient in the Kilengwe forest reserve and (2) quantify and compare impact of anthropogenic disturbances using number of stamps/ha observed in each established habitat. The findings from this study will contribute to the management of the forest reserve and other similar tropical forests in Tanzania and elsewhere.

2. Materials and Methods

2.1. Description of the Study Site. The Kilengwe Forest Reserve is located at latitude 7° 29' South and longitude 37° 32' East at an elevation of 182 to 228 m above mean sea level covering an area of 995 ha. The forest is in Kisaki Ward, Bwakira Chini Division in the Morogoro Rural District. The forest is surrounded by two villages, namely, Kilengwe and Zongomero, and is owned and managed by the local government. A number of seasonal streams that provide water to the local community for domestic purpose originate from this forest reserve. The climate of the region is oceanic due to nearness (about 200 km) to the Indian Ocean and the rainfall regime is bimodal. The long rains last from March to May, peaking in April while the short rains last from October to December. The mean annual rainfall and temperature in the Morogoro region are about 740 mm and 25.1°C, respectively. Agriculture is the most important socioeconomic activity done by locals living in the two adjacent villages. Other land use practices done by locals include livestock keeping, especially poultry, goats, and cows. Bee keeping and carpentry are done at small scale. Illegal logging activities were observed in the forest, especially close to the edges.

2.2. Data Collection. To characterize tree species and structural composition between edge (0–100 m), intermediate (100–200 m), and interior (>200 m) of the forest, six plots of 20 m × 20 m were randomly placed in each of the three established forest habitats. This design was due to the fact that the edge effects can penetrate to 100 m to the forest interior from the margin [2]. In each plot, all trees with diameter at breast height (DBH) ≥ 10 cm measured at 1.3 m above the

ground were counted and identified and stem diameters were recorded from each plot in the habitats. Trees with multiple stems at 1.3 m height were treated as the single individual whereby the diameters of all stems were taken and averaged. If a tree had buttress or an abnormality at 1.3 m height, the diameter was measured just above the buttress where the stem assumes near cylindrical shape. The identification was done with the help of an expert from the forest department of the Morogoro region. Also, in each plot, tree stumps were counted and recorded. Each plot among the established habitats was considered as an independent sample.

2.3. Data Analysis. Species diversity was calculated using the Shannon-Wiener diversity index (H') and Margalef index (D) while the equitability was determined by Pielou's evenness using the Species Richness and Diversity IV (SDR IV) software [24]. Species richness was expressed by the number of observed species in the forest while the first order Jackknife species richness estimator was used to estimate potential species richness in the three studied habitats [25]. The species accumulation curves were constructed for comparing the increase of the number of species with increasing sample area within the habitats. The forest structure was explained in terms of density (stems/ha) and basal area (m²/ha). A multivariate agglomerative clustering technique using the Ward's group linkage based on the Bray-Curtis distance measure was performed using the Community Analysis Package 4 (CAP IV) software [24] to analyze species compositional similarities between the studied habitats. The significant differences in structural composition and stumps/ha between the three habitats were tested using the One Way Analysis of Variance (ANOVA) followed by the *post hoc* Tukey's Highly Significance Difference (Tukey's HSD) test at the 5% significance level using the QED statistical software [24]. Before undertaking ANOVA, all the data were tested for normality using the Shapiro-Wilk tests.

3. Results

3.1. Overall Floristic Composition and Abundance. A total of 199 stems belonging to 67 tree species and 26 families were recorded from an area of 0.72 ha. Family Fabaceae was the most dominant family with 21 species in the studied area of the forest, followed by Moraceae (5 species), and Sterculiaceae (4 species). The most abundant species were *Brachystegia spiciformis* with a relative abundance of 4.02%, followed by *Julbernardia globiflora*, *Burke Africana*, and *Synsepalum cerasiferum* with a relative abundance of 3.52% each. *B. spiciformis* and *J. globiflora* were the most frequent species in the studied area with a relative frequency of 3.73% each. Of the total observed species, 10.45% (7 species: *Acacia nigrescens*, *Albizia glaberrima*, *B. spiciformis*, *Cussonia spicata*, *S. cerasiferum*, *Vitex doniana*, and *Stereospermum kunthianum*) were common in all the three studied forest habitats while 8.95% (6 species), 13.43% (9 species), and 26.86% (18 species) occurred exclusively to the edge, intermediate, and interior habitats, respectively (see Table 1).

TABLE 1: The tree species distribution and abundance in the edge, intermediate, and interior of the forest.

Scientific names	Family	Edge	Intermediate	Interior	Total	Relative abundance
<i>Brachystegia spiciformis</i> Benth.	Fabaceae	2	3	3	8	4.02
<i>Burkea africana</i> Hook. f.	Fabaceae	1	—	6	7	3.52
<i>Jubernadia globiflora</i> (Benth.) Troupin	Fabaceae	—	2	5	7	3.52
<i>Synsepalum cerasiferum</i> (Welw.) T.D.Penn.	Sapotaceae	2	1	4	7	3.52
<i>Bauhinia petersiana</i> Bolle	Fabaceae	1	—	5	6	3.02
<i>Dombeya natalensis</i> Sond.	Sterculiaceae	—	—	6	6	3.02
<i>Ophrypetalum odoratum</i> Diels.	Annonaceae	—	2	4	6	3.02
<i>Acacia nigrescens</i> Oliv.	Fabaceae	1	2	2	5	2.51
<i>Diospyros squarrosa</i> Klotzsch.	Ebenaceae	—	2	3	5	2.51
<i>Ehretia amoena</i> Klotzsch.	Boraginaceae	1	4	—	5	2.51
<i>Stereospermum kunthianum</i> Cham.	Bignoniaceae	2	2	1	5	2.51
<i>Vitex doniana</i> Sweet	Verbenaceae	1	2	2	5	2.51
<i>Albizia glaberrima</i> (Schum. and Thonn.) Benth.	Fabaceae	2	2	1	5	2.50
<i>Dalbergia boehmii</i> Taub.	Fabaceae	—	1	4	5	2.50
<i>Albizia versicolor</i> Welw. ex Oliver	Fabaceae	—	—	4	4	2.01
<i>Cynometra uluguruensis</i> Harms.	Fabaceae	1	—	3	4	2.01
<i>Deinbollia borbonica</i> Scheff.	Sapindaceae	3	—	1	4	2.01
<i>Markhamia obtusifolia</i> (Baker) Sprague.	Bignoniaceae	—	1	3	4	2.01
<i>Myrianthus holstii</i> Engl.	Moraceae	—	—	4	4	2.01
<i>Oncoba spinosa</i> Forssk.	Salicaceae	—	4	—	4	2.01
<i>Sclerocarya birrea</i> (A. Rich.) Hochst	Anacardiaceae	—	—	4	4	2.01
<i>Strychnos spinosa</i> Lam.	Loganiaceae	—	4	—	4	2.01
<i>Terminalia sambesiaca</i> Engl. and Diels.	Combretaceae	—	—	4	4	2.01
<i>Trema orientalis</i> (L.) Blume	Ulmaceae	2	—	2	4	2.01
<i>Combretum molle</i> R.Br. ex G.Don.	Combretaceae	3	—	—	3	1.51
<i>Commiphora africana</i> (A. Rich.) Endl.	Bursaceae	—	1	2	3	1.51
<i>Cussonia spicata</i> Thunb.	Araliaceae	1	1	1	3	1.51
<i>Dalbergia melanoxylon</i> Guill. and Perr.	Fabaceae	—	2	1	3	1.51
<i>Dombeya rotundifolia</i> (Hochst.) Planch.	Sterculiaceae	1	2	—	3	1.51
<i>Khaya anthotheca</i> (Welw.) C. DC.	Meliaceae	1	2	—	3	1.51
<i>Sterculia quinqueloba</i> (Garcke) K. Schum.	Sterculiaceae	1	—	2	3	1.51
<i>Acacia polyacantha</i> Wild.	Fabaceae	—	1	2	3	1.50
<i>Acacia seyal</i> Del.	Fabaceae	—	1	2	3	1.50
<i>Albizia petersiana</i> (Bolle) Oliv.	Fabaceae	—	—	3	3	1.50
<i>Acacia caffra</i> Thunb. Wild.	Fabaceae	1	—	1	2	1.01
<i>Allanblackia uluguruensis</i> Engl.	Clusiaceae	1	1	—	2	1.01
<i>Anthocleista grandiflora</i> L.	Loganiaceae	1	—	1	2	1.01
<i>Antiaris toxicaria</i> Lesch.	Moraceae	—	—	2	2	1.01
<i>Brachystegia temarindoides</i> Benth.	Fabaceae	1	—	1	2	1.01
<i>Breonadia salicina</i> (Vahl) Happer and J.R.I. Wood	Rubiaceae	1	1	—	2	1.01
<i>Bridelia micrantha</i> (Hochst.) Baill.	Euphorbiaceae	—	2	—	2	1.01
<i>Cassipourea mallosana</i> Alston	Rhizophoraceae	2	—	—	2	1.01
<i>Englerophytum natalense</i> (Sond.) T.D. Penn.	Sapotaceae	—	—	2	2	1.01
<i>Ficus exasperata</i> Vahl.	Moraceae	—	1	1	2	1.01
<i>Ficus lutea</i> Vahl.	Moraceae	—	2	—	2	1.01
<i>Grewia similis</i> K. Schum.	Tiliaceae	1	—	1	2	1.01
<i>Margaritaria discoidea</i> (Baill.) G.L. Webster	Euphorbiaceae	—	1	1	2	1.01
<i>Parkia filicoidea</i> Welw.	Fabaceae	1	—	1	2	1.01
<i>Annona senegalensis</i> Pers.	Annonaceae	—	1	—	1	0.50
<i>Cassia abbreviate</i> Oliv.	Fabaceae	—	—	1	1	0.50
<i>Combretum adenogonium</i> Steud. ex Rich.	Combretaceae	—	1	—	1	0.50

TABLE 1: Continued.

Scientific names	Family	Edge	Intermediate	Interior	Total	Relative abundance
<i>Commiphora eminii</i> Engl.	Burseraceae	—	—	1	1	0.50
<i>Cussonia zimmermannii</i> Harms.	Araliaceae	—	—	1	1	0.50
<i>Diplorhynchus condylocarpon</i> (Muell. Arg.) Pichon	Apocynaceae	—	—	1	1	0.50
<i>Erythrophleum suaveolens</i> (Guill and Perr) Brennan	Fabaceae	—	1	—	1	0.50
<i>Harrisonia abyssinica</i> Oliv.	Simaroubaceae	—	—	1	1	0.50
<i>Harungana madagascariensis</i> Lam. ex Poiret	Clusiaceae	—	—	1	1	0.50
<i>Lonchocarpus bussei</i> Harms.	Fabaceae	1	—	—	1	0.50
<i>Markhamia zanzibarica</i> Bojer ex DC.	Bignoniaceae	1	—	—	1	0.50
<i>Milicia excelsa</i> (Welw.) C.C. Berg	Moraceae	—	1	—	1	0.50
<i>Oxyanthus goetzei</i> K. Schum	Rubiaceae	—	—	1	1	0.50
<i>Pouteria altissima</i> Baehni	Sapotaceae	—	—	1	1	0.50
<i>Pseudolachnostylis maprouneifolia</i> Pax	Phyllanthaceae	—	—	1	1	0.50
<i>Pterocarpus tinctorius</i> Welw.	Fabaceae	—	—	1	1	0.50
<i>Scorodophloeus fischeri</i> (Taub.) J. Leon	Fabaceae	1	—	—	1	0.50
<i>Sterculia appendiculata</i> K. Schum.	Sterculiaceae	1	—	—	1	0.50
<i>Tabernaemontana pachysiphon</i> Stapf	Apocynaceae	—	1	—	1	0.50
Total		39	55	105	199	100

3.2. *Species Diversity, Similarity, and Species Accumulation Curves.* The species richness, Shannon-Wiener diversity index (H'), Margalef index (D), Pielou's Evenness (E), and the first order Jackknife estimates of the studied habitats are shown in Table 2. The forest interior habitat was revealed to be the most diverse community with significantly higher species richness (47 species), Shannon-Wiener index (3.65), and Margalef index (9.86) compared to edge and intermediate habitats, which did not differ appreciably. Though evenness did not differ significantly between habitats, the edge community had relatively higher evenness than other habitats. Figure 1 shows the similarity between the three studied habitats. The distance correlation (Ward linkage using the Bray-Curtis measure) between the edge and intermediate is minimal, and this explains that edge and intermediate had a high similarity index (0.32). The forest interior habitat was less similar to edge and intermediate habitats (see Figure 1). The species accumulation curves (Figure 2) revealed an increasing trend as the number of sample areas increased in each studied habitat. The first order Jackknife species richness estimator projected higher species richness in each studied habitat than the observed ones (Table 2). Moreover, the rate of species increase per unit area (i.e., regression slope) was significantly higher in the interior habitat compared to edge and intermediate habitats, which did not differ substantially (Table 3).

3.3. *Structural Composition.* The densities of stems between the studied forest habitats varied significantly ($F(2, 15) = 15.18$, $P = 0.002$) ranging from 163 stems/ha (at the edge) to 442 stems/ha (at the interior) with an overall average of 278 stems/ha when all habitats are pooled together (Figure 3). *Post hoc* Tukey's HSD test ($P < 0.05$) confirmed the interior to have a significantly higher density than edge and intermediate, which did not differ appreciably ($P > 0.05$) in their tree densities. The basal areas differed significantly

between the studied habitats ($F(2, 15) = 5.98$, $P = 0.01$) ranging from 3.7 m²/ha (at the edge) to 11.1 m²/ha (at the interior) (Figure 4). *Post hoc* Tukey's HSD test ($P < 0.05$) revealed the interior of the forest to have a significantly higher basal area compared to the edge and intermediate, which also did not differ considerably ($P > 0.05$) in their basal areas. The species with the highest basal areas were *Dalbergia melanoxylon* and *J. globiflora* with a relative basal area of 8.10% and 6.00%, respectively. Moreover, number of stumps/ha among the habitats varied extensively ($F(2, 15) = 5.19$, $P = 0.02$) from 2 to 14 stumps/ha (Figure 5). *Post hoc* Tukey's HSD test ($P < 0.05$) revealed the forest edge (2 stumps/ha) to have a significantly lower number of stumps/ha than intermediate (8 stumps/ha) and interior (14 stumps/ha), which also differed significantly in their number of stumps/ha ($P < 0.05$).

4. Discussion

4.1. *Overall Floristic Composition of the Forest.* Tropical forests are recognized to harbor more than half of the global species richness, and they are often subjected to increasing anthropogenic pressure, which poses a great threat to existing biodiversity [26]. The Eastern Arc and Coastal forests of Tanzania (Kilengwe inclusive) are known to be rich in species diversity as they are located within the region of high rainfall and habitat heterogeneity [27]. This study has observed Fabaceae to be the most dominant family (with 21 species), which implies that the Fabaceae could be the most dominant tree family in the region. This finding is in line with the results of other authors [22, 28–30] who reported the same family to be the most dominant tree family in the Coastal and Eastern Arc forests of Tanzania. The family Fabaceae was reported to dominate by 25%–50% [28, 31, 32] and by 33% [22] in their studies. The present study revealed the Fabaceae to dominate by 31%. Moreover, like many other studies done

TABLE 2: Species richness, diversity indices, and evenness values of the three studied habitats.

Habitats	Observed species richness (\pm SE)	Jackknife 1 (\pm SE)	Shannon-Wiener (H')	Margalef index (D)	Pielou's evenness (E)
Edge	29 \pm 4 ^a	51 \pm 4 ^a	3.28 \pm 0.19 ^a	7.64 \pm 1.03 ^a	0.97 \pm 0.06
Intermediate	32 \pm 2 ^a	47 \pm 2 ^a	3.34 \pm 0.05 ^a	7.74 \pm 0.40 ^a	0.96 \pm 0.02
Interior	47 \pm 5 ^b	68 \pm 5 ^b	3.65 \pm 0.11 ^b	9.86 \pm 0.94 ^b	0.95 \pm 0.03

Note: values with different letters indicate significant differences between habitats (Tukey's HSD test, $P = 0.05$).

elsewhere [33–37] this study confirms that the Fabaceae is the speciose family in most of the lowland forests of Africa. Other recorded families in this study have similarly been reported by various authors [29, 30, 32] in their studies in the Eastern Arc and Coastal forests of Tanzania.

4.2. Species and Structural Composition along the Edge-Interior Gradient. The study has registered a clear pattern of changes in the tree community whereby the tree species diversity, richness, density, and basal area increased towards the forest interior (Table 2, Figures 2 and 4). In all cases, the forest interior habitat had significantly higher values than the edge and intermediate habitats, which did not differ appreciably. The results signify that edge effects can penetrate to a distance of 200 m from the forest margin. Also, these findings suggest that the forest interior and the edge/intermediate are contrasting habitats. A forest community is said to be speciose if it has a Shannon-Wiener diversity index value ≥ 3.5 [38]. This confirms the observation of this study, which revealed significantly higher Shannon-Wiener diversity index value of 3.65 compared to edge (3.28) and intermediate (3.34) habitats. The presence of less species diversity, richness, density, and basal area at the edges and intermediate can be linked to the fact that these two habitats are easily accessible by local people while fetching for their basic needs like firewood, charcoal, building poles, and traditional medicines [22]. The observed illegal timber harvest contributes to the observed low values, especially on tree density and basal area on the edge/intermediate habitats as it is easier for the timber thieves to cut and carry logs to the truck during the night [22, 39]. The grazing and cattle trampling can also be the causal factor for the observed low values at edges as livestock were observed to reach the forest edges in the studied forest. The Luguru people prefer to use fire to prepare their farms before starting a new agricultural season and for hunting bush animals like *Hyrax pumilus*, *Sus scrofa*, and *Thryonomys* spp., which are used for nutritional purposes. The penetration of fire to forest edges could also account for the observed low species diversity and density values as it kills the fire-sensitive species and affects their regeneration [10, 22]. The periodic fires reduce canopy cover and drastically change vegetation structure and composition [15, 39]. Another observed important factor was the action of sporadic winds that cause great damage to tree communities and high mortality of canopy trees at the edges, which subsequently affect the composition and abundance. Moreover, the present findings are comparable to other works done in Brazil and other countries, in which the forest interior was found to be rich in species diversity and the penetration of edge effects was reported to vary from 15 m to 200 m [12, 40–43]. Distinct abiotic conditions [18, 40–42, 44],

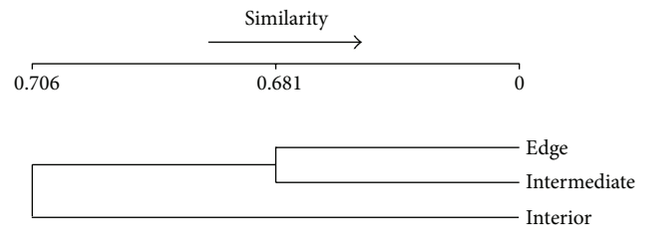


FIGURE 1: Similarity in species composition between the three studied habitats.

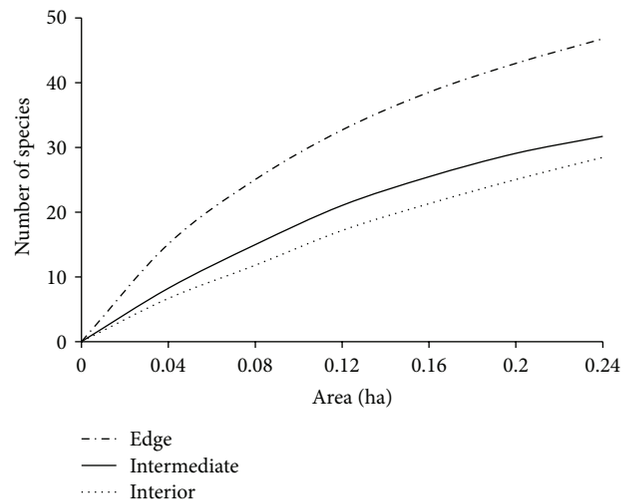


FIGURE 2: Species-area curves based on the cumulative sample area in the studied habitats.

vegetation structure [40], tree mortality rate [2], and high predation, loss of pollinators, and seed dispersers, at edges [11], were reported to be the common cause of differences in species composition between forest edge and interior in various studies done elsewhere.

The presence of significantly less tree density at the edges/intermediate habitats may also be the result of the interplay of factors of two kinds: firstly, those that reduce the possibility of seedling establishment as seedling is the first size class to be affected by edge effects due to its sensitivity to environmental changes and biotic interactions [9, 10], and secondly, those factors that increase seedling, sapling, and adult tree mortality rate [11]. Some processes related to edge effects that could have contributed to low tree density on the edge/intermediate habitats include; reduction in seedling recruitment at edges due to uprooting and breakage due to wind turbulence [2, 45], seedling damage caused by an increasing litter fall near forest edges [11], sapling mortality

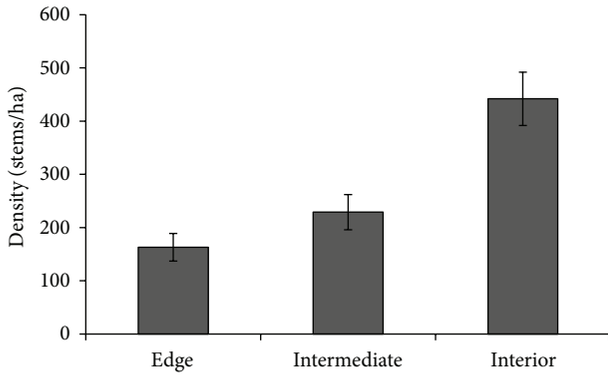


FIGURE 3: Mean tree density (\pm standard error) in the three studied forest habitats.

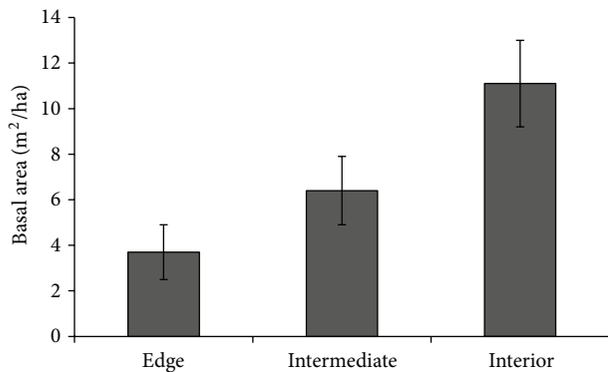


FIGURE 4: Mean basal area (\pm standard error) in the three studied forest habitats.

TABLE 3: Relationship between species richness and forest area.

Location	r^2 -value	Regression equation
Edge	0.99	Species richness = 14.72 ln area - 2.15
Intermediate	0.98	Species richness = 16.77 ln area - 1.78
Interior	0.97	Species richness = 24.32 ln area - 0.88

by competition with lianas, vines, and invasive species [2, 46], and easy accessibility to edges by locals and their livestock [47]. Also, the low basal area at the edge and intermediate habitats could be the result of continuing anthropogenic activities especially illegal logging done by unfaithful locals. Various authors have indicated that the presence of harsh microenvironmental conditions such as air and soil temperatures, high light transmittance, lower relative humidity [17, 23, 48], increased wind forces [2], lower availability of soil carbon, total nitrogen and phosphorus [49], and lower soil and litter moisture [17, 49] near edges contributes to drastic changes in the abundance and distribution of trees in the forest.

4.3. Species Accumulation Curves. The species-area curves displayed an escalating trend, which suggest that increasing the sampling effort could have increased the species richness observed in each habitat. This is due to the fact that the larger

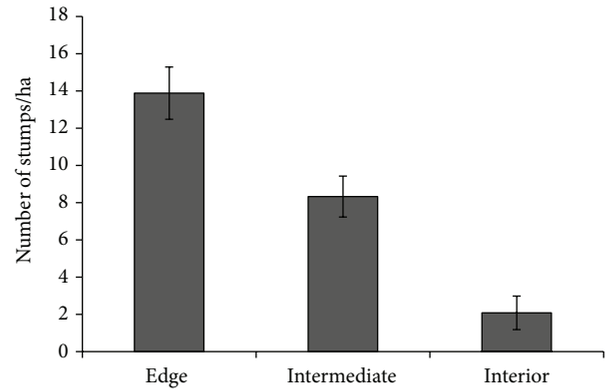


FIGURE 5: Mean stumps/ha (\pm standard error) in the three studied forest habitats.

the forest area sampled is the more environmentally heterogeneous the sampling area becomes and hence the higher the possibility of having many species is. The observed trend in both curves concurs with the first order Jackknife species richness estimator, which provides higher species richness estimates than the observed in each studied habitat. This observation basically informs us that the sample size used was not enough to capture all the species in the forest habitats, which implies that more plots will be required for future inventories in the habitats. Furthermore, the regression slopes have registered an increasing trend from the edge to the forest interior habitat (Table 3). This observation provides a clear indication that the interior habitat is rich and has a high recruitment rate per unit area than edge and intermediate habitats, which did not differ markedly. The observation can possibly be linked with ongoing anthropogenic disturbances observed at the edge and intermediate habitats (Figure 5) whereby the number of stumps/ha was significantly higher in these two habitats compared to the forest interior habitat.

5. Conclusion

In summary, the findings indicate that the edge/intermediate habitats are different from the forest interior in terms of tree species richness, diversity, and structural composition. The forest interior possesses higher species richness, diversity, tree density, and basal area while forest edges and intermediate habitats had impoverished assemblage of tree species, diversity, and density. Also, the edge and intermediate were characterized by high anthropogenic activities by having more stumps/ha than forest interior habitat. If the Kilengwe forest will continue being fragmented, there will be an increase of edge related habitats, which will cause structural and floristic composition changes due to increased edge effects and the forest will face a great threat of losing its original biota especially the rare species. The study recommends long-term research to study microenvironmental factors such as light availability, air and soil temperature, humidity, and nutrients along the edge-interior gradient in the forest in order to determine their influence on tree species richness, composition, and structure.

Conflict of Interests

The author declares that there is no conflict of interests regarding the publication of this paper.

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