Temporal Soundscape Patterns in a Panamanian Tree Diversity Experiment: Polycultures Show an Increase in High Frequency Cover

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In this ecoacoustic study we used the setting of a tropical tree diversity planted forest to analyze temporal patterns in the composition of soundscapes and to test the effects of tree species richness on associated biodiversity measured as acoustic diversity. The analysis of soundscapes offers easy, rapid and sustainable methods when assessing biodiversity. During the last years the quantification of regional or global acoustic variability in sounds and the analysis of different soundscapes has been evolving into an important tool for biodiversity conservation, especially since case studies confirmed a relationship between land-use management, forest structure and acoustic diversity. Here we analyzed soundscapes from two seasons (dry and rainy season) and aurally inspected a subset of audio recordings to describe temporal patterns in soundscape composition. Several acoustic indices were calculated and we performed a correlation analysis and a non-metric multidimensional scaling analysis to identify acoustic indices that: (i) were complementary to each other and such represented different aspects of the local soundscapes and (ii) related most strongly to differences in acoustic composition among tree species richness, season and day phase. Thus, we chose “High Frequency Cover,” “Bioacoustic Index,” and “Events Per Second” to test the hypothesis that acoustic diversity increases with increasing tree species richness. Monocultures differed significantly from polycultures during night recordings, with respect to High Frequency Cover. This index covers sounds above 8 kHz and thus represents part of the orthopteran community. We conclude that increasing tree species richness in a young tropical forest plantation had positive effects on the vocalizing communities. The strongest effects were found for acoustic activity of the orthopteran community. In contrast to birds, orthopterans have smaller home ranges, and are therefore important indicator species for small scale environmental conditions.

Keywords: ecoacoustics, tree diversity experiment, high frequency cover, acoustic diversity, soundscape, temporal patterns
INTRODUCTION

Habitat loss, agricultural activities, global warming and the introduction of exotic species lead to massive decreases of biodiversity all over the world (Sala et al., 2000; Cardinale et al., 2012; IPBES, 2019b). With habitat loss being a major driver of species loss and deforestation still increasing dramatically in the tropics (Brooks et al., 2002; IPBES, 2019a), afforestation is an important and widely used tool for restoring ecosystems and mitigating climate change. Although evidence shows that tree species mixtures can improve multi-functionality in forest (van der Plas et al., 2016, 2017), particularly forest productivity and Carbon storage (Potvin and Gotelli, 2008; Potvin et al., 2011; Ammer, 2019; Guillemot et al., 2020), most areas pledged for afforestation are currently set to become monocultures (Lewis et al., 2019). Monocultures are likely less suited than natural secondary forests or mixed-species plantations for promoting and restoring associated biodiversity (Perfecto et al., 1997; Ampoorter et al., 2020). Increasing tree species richness in plantations might have multiple consequences for the associated fauna, by providing a broader range of food sources for both generalists and specialists herbivores, which could scale up to higher trophic levels (Brose, 2003; Potts et al., 2003; Castagneyrol and Jactel, 2012; Pekin et al., 2012; Ebeling et al., 2018). Additionally, changes in tree species richness has consequences for microclimatic conditions that could affect both diversity and abundance of the animal communities as well as temporal activity patterns (Walker, 1975a,b; Akutsu et al., 2007; Checa et al., 2014; Kunz et al., 2019; Campos-Cerqueira et al., 2020; Burivalova et al., 2021).

Biodiversity assessments in species rich tropical regions are time and resource consuming. This is particularly true for arthropods, as species identification for arthropods can take years and demand expert knowledge in taxonomy. But also for well-known taxonomic groups such as birds sampling is time, money and labor intensive. During the last decade passive acoustic monitoring and ecoacoustic methods have been proven a useful and rapidly expanding tool for fast detection of biodiversity trends, especially in response to land-use change (Burivalova et al., 2018, 2019, 2021). Acoustic surveys do have the advantages of being rapid, sensitive to multiple taxa, non-invasive and easy to apply over large areas and long term periods simultaneously at multiple locations (Sueur et al., 2008). Soundscapes are driven by the composition of different vocalizing communities and thus follow their respective vocal activity patterns resulting in specific diurnal and seasonal soundscapes patterns. However, for tropical habitats, little is known about such acoustic temporal patterns, as most ecoacoustic studies so far have focused on temperate regions (Scarpelli et al., 2020). The majority of tropical ecoacoustic studies focused on diurnal patterns indicate that insects dominate most of the diurnal soundscape, leaving a rather small “empty” acoustic niche for birds at dawn and dusk (Aide et al., 2017). In mature tropical forests these dawn and dusk periods result in an overall soundscape saturation peak, with soundscape saturation being defined as the proportion of active acoustic frequency bins within a recording (Burivalova et al., 2018). Much less is known on seasonal acoustic trends in the tropics, which likely relate to changes in precipitation and humidity, often driven by the alternation of dry and rainy seasons (Rankin and Axel, 2017; Opaev et al., 2021). Additionally, some studies indicate that disturbances and changes in microclimate might interact with the diurnal and seasonal (acoustic) activity patterns of the tropical fauna (Akutsu et al., 2007; Checa et al., 2014; Campos-Cerqueira et al., 2020; Burivalova et al., 2021; Fontürbel et al., 2021).

Phenological acoustic activity patterns can be studied by the application of ecoacoustic methods and its large variety of acoustic indices that have been developed so far (Sueur et al., 2014; Eldridge et al., 2018; Oliveira et al., 2021). In the past, these indices have been successfully used as proxies for various aspects of ecosystem biodiversity and as tools for conservation (Sueur et al., 2008; Pekin et al., 2012; Fuller et al., 2015; Buxton et al., 2016, 2018; Mammides et al., 2017; Burivalova et al., 2018; Gasc et al., 2018; Turner et al., 2018; Shaw et al., 2021). These findings suggest that habitat heterogeneity positively affects acoustic diversity and thus also species richness, at least for vocalizing taxonomic groups. Habitat gradients that were tested in previous ecoacoustic studies were sometimes substantial. Dröge et al. (2021) for example confirmed good relationships between acoustic indices and land-use types, testing a gradient from rice paddies to old growth forests. In the present study we wanted to test how changing a single habitat parameter, that is tree species richness, effects the acoustic composition and temporal dynamic of the acoustic community during the diurnal cycle in both the dry and wet seasons. Experimental plantations are an ideal platform for testing such a question as all other environmental variables can be kept rather constant. This research took advantage of the Sardinilla planted forest, set up in Panama in 2001 to study the effects of tree species richness on ecosystem functions (Scherer-Lorenzen et al., 2005). Here we performed acoustic monitoring in plots with different tree species richness ranging from monocultures to five-species polycultures to test our hypothesis that acoustic diversity increases with increasing tree richness. Based on previous studies highlighting the dominance of insects in tropical soundscapes and given the small size of the research plots we further hypothesized that particularly orthopteran vocalizations should increase with increasing tree species richness (Aide et al., 2017; Campos-Cerqueira et al., 2020).

Above and beyond testing the effect of tree species richness on acoustic diversity, our objectives were to examine how tree species richness effected acoustic patterns through different seasons and day phases and which acoustic indices can be used to capture complementary acoustic patterns.

MATERIALS AND METHODS

Study Site

The Sardinilla planted forest is part of the global research network TreeDivNet, which aims to quantify the relation between tree species richness and ecosystem functioning (Paquette et al., 2018). The experimental site of Sardinilla is located in the central north of Panama (9°19′30″N, 79°38′00″W) at an altitude
of 70 m. The area’s original forest was logged in the early 1950s. After 2 years of agriculture, land-use turned into pasture. The total experimental area covers approximately nine hectares (Scherer-Lorenzen et al., 2007).

The tree plantation included six native tree species that form a gradient from fast-growing pioneers to slow long-lived tree species. Planted fast-growing species were *Cordia alliodora* [(Ruiz and Pav.) Oken] and *Luehea seemannii* (Triana and Planch.), whereas intermediate-growing species were *Anacardium excelsum* [(Bertero and Balb. ex Kunth) Skeels] and *Hura crepitans* (L.). *Cedrela odorata* (L.) and *Tabebuia rosea* [(Bertol.) Bertero ex A.DC.] were chosen as slow-growing tree species. In total, there were 24 plots of 45 m by 45 m consisting of twelve monoculture plots (two per species), six three-species mixtures (one per growth rate category) and six identical six-species plots (Potvin and Gotelli, 2008). One tree species, *Cordia alliodora*, failed to establish. We thus considered the realized species richness (5, 3, 2, and 1) rather than the original richness (6-3-1) (Figure 1). In 2017, after 16 years of growth, a data collection campaign took place during which we set up automatic recording devices to monitor acoustic diversity. The recording period ranged from 30 March 2017 to 11 June 2017, covering parts of the dry and the rainy season. The dry season in Sardinilla lasts from end of December to April followed by 8 months of rainy season, which is reflected in monthly precipitation sums of Sardinilla ranging from 48 to 70 mm in March over 115–142 mm in April up to 306–344 mm in May and 290–339 mm in June 2017 (Hydrometeorology Management of ETESA, 2017). Days were differentiated into four phases: dawn (05:30–09:00), day (09:10–15:50), dusk (16:00–19:30) and night (19:40–05:20).

### Recording Methods

Recordings were made in all five-species plots, all three- and two-species plots and five monoculture plots (one for each species) using prototypes of the SET-Recorders (Soundscape Explanatory Terrestrial, Lunilettronik, Fivizzano)\(^1\). One monoculture (*Hura crepitans*) and two five-species plots were excluded from further analysis as recordings stopped after just a few recording minutes. Thus, in total we analyzed data from 14 plots (four 4 monocultures, 3 two-species, 3 three-species, and 4 five-species). The autonomous recorders were equipped with omnidirectional microphone capsule [EMY-63M/P, sensitivity (0 dB = 1 V/Pa. 1 kHz): dB $-38 \pm 3$, signal to noise ratio: $>60$ dB, input voltage of the ADC: $0.75$ Vrms (personal communication with Lunilettronik Coop.)]. The microphone gain was manually set to +25 dB. The signals were sampled at 48 kHz with a 16 bits digitization, recording for 1-min every tenth minute during 24 h a day. This recording schedule allowed us to cover the whole diurnal cycle, while at the same time covering an extended time period from the dry season into the rainy season. The devices were installed on a tree in the center of the plots, positioned at 1.7 m height always facing south, away from the main wind direction. To assess the quality of the audio data the first and last recordings per recording period were listened to.

### Aural Classification of Audio Files

Adapting the protocol developed by Gasc et al. (2018), we randomly selected a subsample of our recordings – 10 audio files per day phase (4), season (2) and richness level (4), resulting in 320 audio files in total. First author SM listened to each

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\(^{1}\)http://www.lunilettronik.it/prodotto/set-soundscape-explorer-terrestrial/
of these audio files and visualized the spectrogram classifying them according to the three soundscape categories biophony, geophony, and anthropophony (Pijanowski et al., 2011). It was possible that the same audio file contained several of these categories. Furthermore, SM noted for all recordings containing biophony the presence and absence of different vocalizing animal groups, that were anurans, birds, cicada, insects vocalizing in mid-frequency ranges (4–8 kHz) and insects vocalizing in high-frequency ranges (8–24 kHz); again it was possible to assign several subcategories to the same audio file. The presence of other soundscape components such as wind, rain, planes, cars, barking dogs, human voices, and other identifiable sounds was also noted, but we did not use this information for further analysis. In accordance with Gasc et al. (2018) the duration of each category and subcategory as well as the range of song types were recorded. As only birds differed very slightly in that respect among tree species richness levels we refrained from presenting that data.

**Calculation and Selection of Acoustic Indices**

In total we calculated different acoustic features using “Analysis Program” (Towsey et al., 2018) and the Bioacoustic Index (Boelman et al., 2007). Data preprocessing involved the package “stringr” (Wickham, 2019). We used “High Amplitude” and “Clipping Index” to identify recordings with very loud wind and rain noises and excluded all recordings with values > 0 from further analysis. False-Color-Spectrograms (Towsey et al., 2014, 2018) were produced to identify periods with strong rain and wind, which were additionally excluded from further analysis. We assumed that strong rain and strong wind events were identical for all plots. This method allowed us to identify the exact start of the rainy season in that year as being the 15 April 2017.

After calculating acoustic indices, a correlation analysis and a non-metric multidimensional scaling (NMDS) analysis to select acoustic indices for further analysis (see Supplementary for further details) were performed. This served to identify indices that were most complementary to each other and thus were most likely to capture different soundscape components during the different day phases and seasons (Towsey et al., 2014; Phillips et al., 2018). The selection was based on the following criteria (i) acoustic indices should not be significantly correlated among each other (Supplementary Figures 1–11); (ii) selected acoustic indices should relate to the NMDS axis that separate the different day phases, seasons and mixture levels (Supplementary Figures 12–14).

Based on these criteria, “High Frequency Cover”, “Bioacoustic Index,” and “Events Per Second” were selected for analysis. “High Frequency Cover” describes the “fraction of noise-reduced spectrogram cells that exceed 3 dB in the high-frequency band” (8–24 kHz) (Towsey, 2018). The “Bioacoustic Index” is calculated as the “area under each curve included all frequency bands associated with the dB value that was greater than the minimum dB value for each curve. The area values are thus a function of both the sound level and the number of frequency bands used by the” fauna (Boelman et al., 2007). “Events Per Second” represents a “measure of the number of acoustic events per second, averaged over the same noise-reduced 1-min segment. An event is counted each time the decibel value in a bin crosses the 3-dB threshold from lower to higher values” (Towsey, 2018). Insect sounds that cover a frequency band for the whole duration of the audio file (e.g., monotonous stridulations) are not counted as an acoustic event.

**Statistical Methods**

Statistical analyses were performed using R 4.0.2 (R Core Team, 2020). For the selected acoustic indices, we model diurnal patterns per season using generalized additive models (GAMs) and tested if parameter estimates for tree species richness level were significant. Tree species richness level (Mix) and season were included as factors, time of the day as smoother and “PlotID” as a random effect. Tropical acoustic communities show diurnal and seasonal activity patterns, to test this we added interaction terms for tree species richness level and season, as well as an additional smoother interaction term for time of day and season. GAMs were calculated using the “mgcv” package.

### TABLE 1 Model specifications and output for High Frequency Cover.

| High frequency cover | Estimate | Std. error | z Value | Pr(>|z|) |
|----------------------|----------|------------|---------|----------|
| SR1: Monoculture dry season (intercept) | −2.98 | 0.09 | −34.73 | <0.001*** |
| SR2 – two tree species | 0.61 | 0.13 | 4.67 | <0.001*** |
| SR3 – three tree species | 0.47 | 0.13 | 3.62 | <0.001*** |
| SR5 – five tree species | 0.30 | 0.12 | 2.49 | <0.05* |
| Season rain | −0.21 | 0.01 | −20.00 | <0.001*** |
| SR2:rain | 0.04 | 0.01 | 2.83 | <0.01** |
| SR3:rain | 0.02 | 0.01 | 1.13 | >0.2 n.s. |
| SR5:rain | −0.003 | 0.01 | −0.22 | >0.8 n.s. |

| Approximate significance of smooth terms | edf | Ref.df | Chi.sq | p-Value |
|----------------------------------------|-----|--------|--------|---------|
| s(ToD):SR1 | 12.26 | 48 | 77.75 | <0.001*** |
| s(ToD):SR2 | 7.26 | 48 | 18.23 | <0.001*** |
| s(ToD):SR3 | 11.16 | 48 | 49.23 | <0.001*** |
| s(ToD):SR5 | 2.33 | 48 | 2.66 | <0.001*** |
| s(ToD):seasondry | 23.44 | 48 | 685.762 | <0.001*** |
| s(ToD):seasonrain | 34.06 | 48 | 619.562 | <0.001*** |
| s(PlotID) | 9.98 | 10 | 4.041 | <0.001*** |

Mix: tree species richness level (SR1, SR2, SR3, and SR5 for 1, 2, 3, and 5 tree species), season: dry and rainy season, ToD, Time of day (hour).

Significant levels are given as: **p < 0.001, *p < 0.01, p < 0.05, p > 0.1.”
### TABLE 2 | Model specifications and output for Bioacoustic Index.

**Bioacoustic index**

Family: Scaled (df=3.383,0.364)

Link function: identity

Formula:

\[
\text{BioAcousticlog} = \alpha + \beta \cdot \text{Species} + \gamma \cdot \text{Season} + \delta \cdot \text{ToD} + \epsilon + \eta \cdot \text{PlotID} + \zeta \cdot \text{Species} \times \text{Season} + \eta \cdot \text{Species} \times \text{ToD} + \eta \cdot \text{Species} \times \text{PlotID} + \varepsilon
\]

Approximate significance of smooth terms

| Estimate | Std. error | z Value | Pr(>|z|) |
|----------|------------|---------|----------|
| SR1: Monoculture dry season (intercept) | 4.62 | 0.073 | 63.05 | <0.001*** |
| SR2 – two tree species | 0.30 | 0.112 | 2.70 | <0.01** |
| SR3 – three tree species | 0.001 | 0.011 | 0.01 | >0.900 n.s. |
| SR5 – five tree species | −0.01 | 0.104 | −0.07 | >0.900 n.s. |
| Season rain | 0.211 | 0.006 | 33.78 | <0.001*** |
| SR2:rain | −0.20 | 0.010 | −21.53 | <0.001*** |
| SR3:rain | 0.12 | 0.010 | 13.42 | <0.001*** |
| SR5:rain | −0.03 | 0.010 | −3.78 | <0.001*** |

### TABLE 3 | Model specifications and output for Events Per Second.

**Events Per Second**

Family: negative binomial (1.58034.093)

Link function: log

Formula:

\[
\text{EventsPerSeconds} = \alpha + \beta \cdot \text{Species} + \gamma \cdot \text{Season} + \delta \cdot \text{ToD} + \epsilon + \eta \cdot \text{PlotID} + \zeta \cdot \text{Species} \times \text{Season} + \eta \cdot \text{Species} \times \text{ToD} + \eta \cdot \text{Species} \times \text{PlotID} + \varepsilon
\]

Approximate significance of smooth terms

| Estimate | Std. error | z Value | Pr(>|z|) |
|----------|------------|---------|----------|
| SR1: Monoculture dry season (intercept) | −1.15 | 0.178 | −6.450 | <0.001*** |
| SR2 – two tree species | 0.79 | 0.272 | 2.904 | <0.01** |
| SR3 – three tree species | 0.33 | 0.272 | 1.220 | >0.200 n.s. |
| SR5 – five tree species | 0.14 | 0.252 | 0.239 | >0.500 n.s. |
| Season rain | −0.35 | 0.024 | −14.377 | <0.001*** |
| SR2:rain | −0.002 | 0.031 | −0.070 | >0.900 n.s. |
| SR3:rain | −0.01 | 0.033 | −0.279 | >0.700 n.s. |
| SR5:rain | −0.16 | 0.032 | −4.861 | <0.001*** |

(Wood, 2011). Model diagnostics were assessed graphically and a specific distribution family was necessary for each acoustic index. Respective model specifications can be found in Tables 1–3.

For graphical outputs we use the packages “ggplot2” (Wickham, 2016) and “tidyverse” (Coretta, 2021).

### RESULTS

**Temporal Patterns and Soundscape Composition**

**Aural Inspection**

The number of recordings containing biophony did not change substantially across day phases and season, as biophony was recorded in almost all recordings that were aurally inspected (Figure 2). Both geophony and anthropophony appeared mostly during the day and rarely at night, but were overall less frequent than biophony. The most common geophony sounds were light wind and rain, as recordings with heavy rain and strong wind were excluded from the recordings. Anthropophony consisted of sounds from planes, undefinable machinery, nearby constructions, vehicles, and chainsaws.

Distinct patterns were found between day – when birds and non-biophony sounds were dominating the soundscape – and night – with orthopterans occupying a large frequency band (Figure 3). Other sound sources in the Sardinilla planted forest included insects vocalizing in mid-frequency ranges (4–8 kHz, mainly crickets), anuran (dominating the range just below the crickets) and cicada (rare events, overlapping with crickets in the frequency domain). Bird vocalizations were recorded from dawn until dusk, and dominated the day recordings. This dominance resulted from a reduced number of recordings with vocalizations from anurans and insects rather than an increase in bird activity. Thus, the aural inspection did not indicate a clear peak of bird vocal activity at dawn and dusk, as can be observed in temperate or boreal biomes. There was no observable seasonal trend for birds. Anurans were least active during the day but this activity increased in the rainy season. Insects singing in mid-frequency ranges were also least active during the day, with a slight increase...
in the rainy season. Insects singing at high frequency ranges showed clear activity peaks at dusk and night with no strong seasonal patterns. Cicadas were not common to observe, and showed an irregular pattern with a slight higher frequency of observations during day and dusk in the dry season.

Comparison of Acoustic Indices

The correlation analysis revealed that acoustic indices could be grouped into four clusters (Supplementary Figure 1). Cluster 1 consisted of Entropy of Average Spectrum and Entropy of Variance Spectrum. These indices showed higher values at night compared to day, with only slight differences among rainy and dry season (Supplementary Figures 2, 3). These indices were negatively correlated with indices from cluster 2 and 4 and were therefore not included in further analysis. Cluster 2 consisted of Acoustic Complexity Index, Mid Frequency Cover and Bioacoustic Index. All indices showed distinct dawn peaks and different patterns between dry and rainy season (Supplementary Figures 4–6). From this cluster we chose the Bioacoustic Index for further hypothesis testing, also based on results from an NMDS analysis (Supplementary Figure 12). The Bioacoustic Index showed peaks at dawn and dusk, slightly lower values during the day and lowest values at night (Figure 4). During the rainy season, the patterns for these indices changed: index values increased throughout the day, but particularly at midday, resulting in less pronounced peaks at dawn and dusk. Thus, the Bioacoustic Index corresponded to anuran and bird acoustic activity patterns identified in the aural inspection. Cluster 3 consisted of High Frequency Cover and Spectral Density. These indices also showed higher values at night compared to day (as cluster 1, Supplementary Figures 7, 8). From this cluster we chose High Frequency Cover for further analysis. High Frequency Cover was designed to represent the orthopteran community that predominantly communicates in high frequencies, while few other vocalizing animal groups occupy this frequency range. Peaks at night for High Frequency Cover, as well as a slight reduced High Frequency Cover in the rainy season (Figure 5) could be verified by the aural inspection (section “Aural Inspection” and Figures 3, 5). Cluster 4 consisted of Low Frequency Cover, Events per Second and Temporal Entropy. These indices showed strong peaks during the day, without peaks at dawn and dusk, and very low values at night (Supplementary Figures 9–11). Dry season had higher peak values than the rainy season. From this cluster Events Per Second was chosen for further analysis (Figure 6).

Effects of Tree Species Richness on Temporal Patterns of Acoustic Indices

There was a significant lower High Frequency Cover in monocultures at night, compared to polycultures (Figure 5), and parametric coefficients for polycultures were significantly different from the monocultures (Table 1). Polycultures did
FIGURE 3 | Detailed biophony composition analysis, from the 320 recordings that were aurally classified, per season, day phase, and tree species richness level. Recordings could contain more than one biophony component. All recordings containing biophony were classified according to presence of birds, anurans, insects vocalizing in mid-frequency ranges (4–8 kHz, MF Insects), insects vocalizing in high-frequency ranges (8–24 kHz, HF Insects) and cicadas.

FIGURE 4 | Fitted values for bioacoustic index over a diurnal cycle (time of day) per season (dry and rainy season) and tree SR level. Dashed vertical line indicate sunrise (ca. 06:00) and sunset (around 18:30).
not differ from each other, but two-species mixtures showed a slight but significant lower decrease of High Frequency cover in the rainy season than the other plots (Table 1). In the dry season the parametric coefficients for two-species mixtures were significantly higher than for the other mixture levels in terms of the Bioacoustic Index, and differences were most pronounced at night (Figure 4 and Table 2). In the rainy season the three-species mixtures showed higher values than the other plots at midday. For Events Per Second the two-species mixtures showed parametric coefficients significantly different from the monocultures while the other polycultures did not (Table 3), and had a significantly higher day peak during the dry season.
Differences among two-species mixtures and the other tree species richness levels were less pronounced but still significant during the rainy season (Figure 6).

**DISCUSSION**

**Indices to Capture Temporal Patterns of Acoustic Diversity**

Different acoustic indices capture different aspects of the soundscape. By choosing non-redundant indices, it is possible to capture complementary patterns in the soundscape, and attribute them to specific groups in the acoustic community (Buxton et al., 2018; Eldridge et al., 2018; Phillips et al., 2018). Since acoustic community composition and dynamics vary among different biomes in interaction with local geophony and anthropophony, it may be necessary for each ecoacoustic study to identify its own unique set of indices to describe soundscape composition. Ross et al. (2021) tested several acoustic indices within different sonic conditions for their performance in capturing biological meaningful patterns and sensitivity in respect to noise. Most commonly used indices showed sensitivity to various confounding sound sources (Ross et al., 2021). Even though they came up with very useful practical recommendation for the suitability of different indices in different sonic conditions, studies in other biomes identified different sets of acoustic indicators to best describe their specific soundscape (Fuller et al., 2015; Ng et al., 2018; Turner et al., 2018; Carruthers-Jones et al., 2019; Oliveira et al., 2021).

If we understand acoustic indices as an integrative measure of biodiversity that reflects the soundscape composition and particularly the acoustic communities as a whole in response to human activity and ecological gradients (Pijanowski et al., 2011; Farina, 2014; Farina and Gage, 2017) we also have to understand that a single acoustic index will rarely correspond only to a single vocalizing animal group (Retamosa Izaguirre et al., 2021). This is probably especially true in the tropics as different vocalizing animal groups often vocalize at the same time (Eldridge et al., 2018) and could explain why in our study the temporal patterns of the chosen acoustic indices did not match the patterns of a single vocalizing animal group perfectly well. This is in line with other studies, mostly on birds, that have shown an often significant but weak correlation of acoustic indices with bird richness (Moreno-Gómez et al., 2019; Dröge et al., 2021). Even though recent studies indicate that such correlations might be improved if species richness is based on bioacoustic identification from acoustic recordings rather than point-count assessments (Shaw et al., 2021). In contrast, studies focusing on the relationship of acoustic indices with parameters that describe ecological condition such as vegetation structure, vegetation complexity, habitat type, land-use intensity and/or distance to the nearest road often report acoustic indices to be good indicators (Tucker et al., 2014; Fuller et al., 2015; Bürüvalova et al., 2018; Ng et al., 2018).

By the selection process we applied and by analyzing different temporal acoustic niches, we could identify three complementary indices that allowed us to test effects of tree species richness on different vocalizing communities. As Metcalf et al. (2020) argued, the interpretability of acoustic indices can be greatly improved by calculating them in ecological meaningful time and frequency bins. But even then, given the integrative properties of acoustic indices in capturing soundscape components, interpretability of acoustic index patterns might not be very straightforward. Aural inspection of a subset of recordings supported the interpretability in our study. Nevertheless, aural inspection only indicated the presence or absence of certain vocalizing groups across day phases and season, while acoustic indices respond also to diversity, distance and abundances of vocalizing animal groups.

High Frequency Cover in our study matched the temporal acoustic activity patterns identified for orthopteran vocalizing in high frequencies, and was therefore a useful tool to identify effects of tree species richness on this insect community. The Bioacoustic Index is frequently used in ecoacoustic studies and it was designed to specifically pick up bird vocalization patterns (Boelman et al., 2007). In our study it seemed to be driven both by bird and anuran vocalizations. While aural inspection of recordings did not suggest a change in activity patterns of birds in our study, the increase in Bioacoustic Index during the rainy season might be mainly driven by an increase in anuran activity. Also Boullhesen et al. (2021) found a very good relationship between the Bioacoustic Index and frog vocal activity patterns and frog species richness. Opav et al. (2021) found a similar increase of Bioacoustic Index with increasing humidity just before the monsoon season, which they associated with the onset of the breeding season for both birds and anurans. In a study of different forest types in the Valdivian temperate rainforests of southern Chile, Fontúrbel et al. (2021) observed a clear peak of bird vocalizations at dawn only in old- growth forest, while it peaked at noon in plantations, and in the afternoon in logged forests and showed a variable pattern in secondary forest, suggesting that daily activity peaks of birds could depend on forest understorey and forest structure. Our data showing that birds were active throughout the day from dawn until dusk and dominated the soundscape at noon is coherent with the observations made in Chilean plantations.

Events Per Second was correlated with Low Frequency Cover and Temporal Entropy and showed a clear peak during the day, but no peaks at dawn and dusk. This might have related to an increase in geophony and anthropophony during the day recordings (Figure 2), which should both be related to Low Frequency Cover (Gage and Axel, 2014; Shaw et al., 2021). The two-species mixtures were a result of establishment failure of *Cordia alliodora*. Which could have resulted in higher wind noises in these plots due to lower stand density and might explain partly that these plots showed significantly higher values for Events Per Second throughout the day, in comparison to the other plots. Additionally, Events Per Second might relate to a dominance of birds while insects showed a reduced activity. This would relate also to the fact that Events Per Second is reduced in the rainy season when the dominance of birds is also reduced due to an increase in anuran activity.

Since the high frequency ranges - in most environments we can think of - are occupied exclusively by orthopterans (and sometimes bats), the usefulness of High Frequency Cover can probably be generalized to other ecosystems and habitats. The
situation is more complex in mid-frequency ranges where more than one vocalizing animal group is active. Other studies have addressed the advantages and disadvantages of different acoustic indices under different environmental conditions, and useful overviews can be found, for example, in Bradfer-Lawrence et al. (2019), Zhao et al. (2019), Ross et al. (2021), Sánchez-Giraldo et al. (2021), and Shaw et al. (2021).

**Effects of Tree Species Richness**

High frequency cover, revealed a positive effect of polycultures on insect acoustic activity that might relate to abundance of this vocalizing insect group. This confirmed our hypothesis that planting polycultures has positive effects on associated biodiversity, specifically manifested within the orthopteran community due to their small home ranges. Birds in contrast are likely to be more mobile and their home ranges might not be restricted to a single plot. A recent meta-analysis revealed that bird diversity in plantations is lower than in natural forests but that overall faunal diversity, including birds, benefits from planting a mix of native tree species (Bohada-Murillo et al., 2020; Wang et al., 2021).

An increase in tree species richness could increase orthopteran abundance and diversity via an improved variety of food sources, especially for specialist plant consumers. A mechanism that is modulated by trophic interactions, as was shown for grassland ecosystems (Siemann, 1998; Ebeling et al., 2018). In forests these relationships were often studied under the objective of testing tree species richness effects on herbivore damage on trees (associational resistance hypothesis, Jactel et al., 2021), and not how it would affect orthopteran diversity. Still, there is evidence that herbivore arthropod diversity and abundance is related to tree diversity and that this relationship between plant diversity and consumer diversity might be particularly strong in the tropics (Becerra, 2015). Given that many orthopteran are highly host tree specific, the effect of tree species identity has to be considered as well (Novotny and Basset, 2005).

Another mechanism could be the indirect effect of tree species richness on microclimate and microhabitats. In a global meta-analysis, canopy cover, which affects microclimate, was the most important driver of faunal biodiversity (Wang et al., 2021). How microclimate can change the acoustic communities was addressed by Campos-Cerqueira et al. (2020) and Burivalova et al. (2021). In their study selective logging resulted in microclimatic changes that decreased humidity in the logging gaps. This had impacts on amphibians and understory specialist birds, and certain insect groups. Anurans are driven by the presence of microhabitats and waterbodies while tree species richness itself is likely less important (Duellman, 1988; Oda et al., 2016). In our study we found tree species richness effects on Bioacoustic Index, suggesting that anuran and maybe also bird activity was positively related to two-species mixtures in the dry season and three-species mixtures in the rainy season.

The sensitivity of insects to land-use induced habitat and microclimatic changes could make them very suitable indicators for disturbance regimes in tropical forests (Akutsu et al., 2007; Campos-Cerqueira et al., 2020). These studies highlight the urgency for including insects into biodiversity assessments. While acoustic identification of insects in the tropics poses a significant challenge, acoustic indices can give valuable insights in temporal activity changes, and indicate changes in abundance and diversity (Oliveira et al., 2021). Thus, acoustic monitoring has great potential to become an important additional monitoring tool which is crucial particularly for tropical ecosystems (Lamarre et al., 2020). In our study, differences in microclimate among plots might be a result of the different tree species richness levels, that result from differences in above ground space use efficiency (Sapijanskas et al., 2014). Additionally, the failure of one tree species resulted in larger canopy openings of some plots, mainly the two-species mixture. If insects profit from canopy gaps this could explain why two-species mixture often showed the highest acoustic activity.

**CONCLUSION**

In this study we showed that planting polycultures increased orthopteran acoustic activity at night, and this may be related to an increase in abundance and maybe also richness of this taxonomic group. Given the small scale of the plots and the young age of the planation this would indicate that this animal group is most sensitive to these small scale habitat differences in our experimental forest plantation, making them important indicator species for monitoring ecosystem changes. Additional to direct effects of tree species richness and tree species identity, variations in microclimate most likely drive the observed differences in acoustic patterns across season and day phases. We have shown that ecoacoustics can provide valuable insights in studying the interaction between forest features, associated biodiversity and acoustic activity patterns in the tropics where these interactions are far less studied than in temperate regions.

**DATA AVAILABILITY STATEMENT**

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

**AUTHOR CONTRIBUTIONS**

SM, MS-L, and CP designed the study. SM analyzed the data with support from OM and LO and wrote the manuscript. All authors read, commented on, and approved the final version of the manuscript.

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SUPPLEMENTARY MATERIAL
The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fevo.2022.808859/full#supplementary-material

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