Correction: Climate change versus deforestation: Implications for tree species distribution in the dry forests of southern Ecuador

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There is an error in the first sentence of the second paragraph in the Introduction section. The correct sentence is: Ecuador, one of the ten most biodiversity-rich nations in the world [11, 12], has one-sixth of its territory covered by deciduous and semi-deciduous forests [13] with a reported national deforestation rate of approximately 475 km$^2$/year during 2008–2014 [14].

Reference
1. Manchego CE, Hildebrandt P, Cueva J, Espinosa CI, Stimm B, Günter S (2017) Climate change versus deforestation: Implications for tree species distribution in the dry forests of southern Ecuador. PLoS ONE 12(12): e0190092. https://doi.org/10.1371/journal.pone.0190092 PMID: 29267357
Climate change versus deforestation: Implications for tree species distribution in the dry forests of southern Ecuador

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Abstract

Seasonally dry forests in the neotropics are heavily threatened by a combination of human disturbances and climate change; however, the severity of these threats is seldom contrasted. This study aims to quantify and compare the effects of deforestation and climate change on the natural spatial ranges of 17 characteristic tree species of southern Ecuador dry deciduous forests, which are heavily fragmented and support high levels of endemism as part of the Tumbesian ecoregion. We used 660 plant records to generate species distribution models and land-cover data to project species ranges for two time frames: a simulated deforestation scenario from 2008 to 2014 with native forest to anthropogenic land-use conversion, and an extreme climate change scenario (CCSM4.0, RCP 8.5) for 2050, which assumed zero change from human activities. To assess both potential threats, we compared the estimated annual rates of species loss (i.e., range shifts) affecting each species. Deforestation loss for all species averaged approximately 71 km²/year, while potential climate-attributed loss was almost 21 km²/year. Moreover, annual area loss rates due to deforestation were significantly higher than those attributed to climate-change (P < 0.01). However, projections into the future scenario show evidence of diverging displacement patterns, indicating the potential formation of novel ecosystems, which is consistent with other species assemblage predictions as result of climate change. Furthermore, we provide recommendations for management and conservation, prioritizing the most threatened species such as Albizia multiflora, Ceiba trichistandra, and Cochlospermum vitifolium.

Introduction

Seasonally dry tropical ecosystems have harbored humans for thousands of years. The Americas are no exception because these ecosystems have historically been the preferred zones for settlement and agriculture [1–3]. Due to these and other anthropogenic influences, neotropical seasonally dry forests are the most threatened tropical forests in the world [4], which similarly
to other dry areas, could be at risk of degradation due to the effects of climate change [5]. The latest estimates indicate that two-thirds of original neotropical dry forest has been converted to other types of land uses [6]. Some authors argue that the combination of anthropogenic pressure, variability in climatic conditions, and climate change makes tropical dry forests particularly vulnerable regions [4,7]. Of these threats, climate change is perhaps the greatest uncertainty as it might cause species extinctions, range shifts, and biodiversity loss [8–10], particularly in areas where the magnitude of the threats have not been explored yet.

Ecuador, one of the ten most biodiversity-rich nations in the world [11,12], has one-sixth of its territory covered by deciduous and semi-deciduous forests [13] with a reported national deforestation rate of approximately 30 km²/year during 2008–2014 [14]. A high proportion of Ecuador’s seasonally dry forests is located in the southwestern part of the country, situated in the Tumbes–Chocó–Magdalena region and adjacent to the Tropical Andes, two large biodiversity hotspots with great species diversity and high levels of species endemism, but also with high habitat loss caused by land-use change [15]. Moreover, these dry forests are particularly susceptible because they are highly fragmented, less than 2.3% of their areas are represented in natural reserves [6], and almost all major conservation threats are linked to habitat degradation [16]. Moreover, these forests not only provide timber and non-timber forest products, but also key ecosystem services such as water flux balance and erosion prevention.

In addition, deforestation rates for the seasonally dry forests of southwestern Ecuador were approximately 29.2 and 57.2 km²/year from 1976 to 1989 and 1989 to 2008 [17], respectively, where the most prominent native forest conversions were toward pasture or crops [18,19]. On this subject, there is high certainty that land-use change contributes to environmental degradation and exacerbates the negative impacts of climate change [20]. For instance, temperature increases of 0.1°C to 0.2°C per decade and precipitation variations of 4% per decade have already been detected in Ecuador between 1961 and 1990 [21]. In the case of southern Ecuador, Peters et al. [22] found a similar warming pattern of 0.13°C per decade and weak but significant trends in increasing rainfall. In addition, future climate projections for southwestern Ecuador predict a 2°C to 5°C increase in air temperature and a 10% to 40% increase in precipitation by the end of the century [23,24]. Regardless of the future climate scenario, most projections indicate an increase in temperatures and a variation in precipitation values, suggesting precipitation increases in southwestern Ecuador by the end of the century [20].

Altogether, climate change simulations signal an increase in seasonality by 2030 in the areas proximal to the Andes [25], and the effects on native tree species are already being manifested as upslope range shifts [26,27]. However, given that individual species are expected to have different range shifts depending on internal and external traits [28], quantifying the magnitude and direction of these shifts is important in assessing whether the current species composition will remain constant or disaggregate with future changes. Convergent changes could indicate that ecosystem compositions will remain stable (assuming equal displacement ability) while divergent patterns may indicate new ecosystem compositions, with unknown consequences on synecology, ecosystem functions, and thus ecosystem services. In this regard, a meaningful approach using a response-and-effect functional framework was suggested by Suding et al. [29] to minimize these uncertainties.

Furthermore, for efficient planning, implementation of conservation measures, and sustainable land use, prioritizing efforts according to threats and vulnerabilities is important. Therefore, it is critical to differentiate between potential climate change and deforestation threats, identifying patterns at both species and community levels. In this study, we use species distribution models to estimate and compare potential climate change threats with current deforestation patterns for a characteristic plant community of 17 tree species in the seasonally dry forest of Ecuador. We hypothesize that (a) deforestation differ in magnitude and spatial
distribution from potential range shifts due to climate change; (b) both patterns do not exhibit species-specific effects; and (c) individual species responses reveal a convergent pattern, maintaining community structure. In this study, we aim to provide a scientific reference frame to identify the lesser of two evils and provide a basis for effective resource allocation in forest conservation and sustainable land use.

**Materials and methods**

**Study area**

Although the precise geographical extent of the dry deciduous forest region in Ecuador lacks unanimous consensus [30–35], authors agree on the presence of several distinct ecosystems within this region. In this study, we focus on the dry deciduous forest on hillsides of southwestern Ecuador, as proposed and described by Aguirre et al. [36], because this ecological unit is heavily threatened by human intervention [17,37,38]. The locality is characterized by a 5-month dry season, mean annual temperature of 20°C–26°C, precipitation ranging from 300 to 700 mm/year [39], and a high number of endemic species [40,41].

**Species records**

According to the criteria of Aguirre et al. [36], we selected all characteristic tree species of the dry deciduous forest on hillsides of southwestern Ecuador, excluding predominantly shrub life-forms [42,43] (Table 1). All 17 tree species are used as local timber or other wood products [43], and although these species are categorized as distinctive of the area, they do not exclusively occur in this region [44]. Presence records were obtained from the GBIF database [45] and complemented with inventory data from our permanent plots as well as herbarium records at the Universidad Nacional de Loja (S1 Dataset).

| Selected species | Records | Synonyms | Family | Elevation |
|------------------|---------|----------|--------|-----------|
| Albizia multiflora (Kunth) Barneby & J.W. Grimes | 51 | Acacia multiflora; Pithecellobium multiflorum | Mimosaceae | 0–1000 |
| Bursera graveolens (Kunth) Triana & Planch. | 42 | Elaphrium graveolens; Spondias edmonstonei | Burseraceae | 0–2000 |
| Caesalpinia gibrata Kunth | 65 | Caesalpinia paapi; Caesalpinia corymbosa | Caesalpinaceae | 0–500 |
| Cavanillesia plataniolphi (Bonpl.) Kunth | 07 | Pourretia plataniolphia | Malvaceae | 0–500 |
| Ceiba trichistandra (A. Gray) Bakh | 39 | Eriodendron trichistandrand | Malvaceae | 0–500 |
| Chloroleucon mangense (Jack.) Britton & Rose | 53 | Pithecellobium mangense; Mimosa mangensis | Mimosaceae | 0–1000 |
| Cordia macrantha Chodat | 17 | - | Boraginaceae | 0–500 |
| Coccoloba ruiziana Lindau | 33 | - | Polygonaceae | 0–1000 |
| Colicodendron scabridum (Kunth) Seem | 26 | Capparis scabrida | Capparaceae | 0–500; 1000–2000 |
| Cochlospermum vitilolum (Willd.) Spreng. | 40 | Bombax vitilolum | Cochlospermaceae | 0–1000 |
| Erythrina velutina Wild. | 23 | Erythrina splendida | Fabaceae | 0–500 |
| Geoffroea spinosa Jacq. | 51 | Geoffroea striata; Robinia striata | Fabaceae | 0–500 |
| Guazuma ulmifolia Lam. | 52 | - | Sterculiaceae | 0–2500 |
| Handroanthus chrysanthus (Jacq.) S.O. Grose | 52 | Tabebia chrysanthha | Bignoniaceae | 0–1000 |
| Loxopterygium suaveolens Spruce ex Engl. | 24 | - | Anacardiaceae | 0–2000 |
| Piscidia carthagenerica Jacq. | 55 | Piscidia acuminata; Ichthyomethia acuminata | Fabaceae | 0–500 |
| Prosopis juliflora (Sw.) DC. | 30 | Mimosa juliflora | Mimosaceae | 0–500 |

Tree species and number of occurrence records used to produce the species distribution models, along with register data from the Catalogue of the Vascular Plants of Ecuador [42] that were validated through The Plant List [46]. Elevation is given in m a.s.l.

https://doi.org/10.1371/journal.pone.0190092.t001
The geographical accuracy of species records was ensured by validating metadata and verifying individual coordinates through the OpenLayers plugin 1.1.4 for QGIS. Then, we used the R script ElimCellDups [47] to retain a single species occurrence per raster cell.

**Predictor variables**

Present and future bioclimatic layers were obtained from WorldClim.org [48] at 30-arc second resolution, approximately $1 \times 1$ km near the equator. In addition, the following three topographical variables were used: soil classification based on the USDA denominations; absolute depth to bedrock; and soil organic carbon stock. All three variables were obtained from SoilGrids.org [49–51], and their grid resolutions were adjusted to match the bioclimatic layers. The chosen future scenario was the most extreme possible outcome for 2050 and utilized the Representative Concentration Pathway (RCP) 8.5 from the global circulation model CCSM4.0 in accordance with the fifth intergovernmental panel on climate change assessment report [52]. The complete list of variables considered in the models is shown in Table 2. Furthermore, to account for the fundamental role of environmental space during modeling [53,54], we delimited the spatial grid coverage to the Coastal and Andean regions of Ecuador, excluding major islands and Amazon region provinces (Fig 1).

**Modeling potential species distributions**

The environmental niche modeling was produced with Maxent v3.3, a widely used algorithm for assessing species distributions that can rely on presence-only data and retain a strong predictive power compared to other approaches [55]. To obtain biological meaningful outcomes,

| Variable                                      | Code | Data resolution |
|-----------------------------------------------|------|-----------------|
| Annual mean temperature [$\degree$C]          | bio01| $\sim1000$      |
| Mean diurnal range (Mean of monthly (max temp–min temp)) [$\degree$C] | bio02| $\sim1000$      |
| Isothermality ($(bio2/bio7) \times 100)$ [%]  | bio03| $\sim1000$      |
| Temperature seasonality (standard deviation $\times 100$) [$\degree$C] | bio04| $\sim1000$      |
| Max. temperature of warmest month [$\degree$C]| bio05| $\sim1000$      |
| Min. temperature of coldest month [$\degree$C]| bio06| $\sim1000$      |
| Temperature annual range (Bio5-Bio6) [$\degree$C]| bio07| $\sim1000$      |
| Mean temperature of wettest quarter [$\degree$C]| bio08| $\sim1000$      |
| Mean temperature of driest quarter [$\degree$C]| bio09| $\sim1000$      |
| Mean temperature of warmest quarter [$\degree$C]| bio10| $\sim1000$      |
| Mean temperature of coldest quarter [$\degree$C]| bio11| $\sim1000$      |
| Annual precipitation [mm]                    | bio12| $\sim1000$      |
| Precipitation of wettest month [mm]          | bio13| $\sim1000$      |
| Precipitation of driest month [mm]           | bio14| $\sim1000$      |
| Precipitation seasonality (coefficient of variation) [%] | bio15| $\sim1000$      |
| Precipitation of wettest quarter [mm]        | bio16| $\sim1000$      |
| Precipitation of driest quarter [mm]         | bio17| $\sim1000$      |
| Precipitation of warmest quarter [mm]        | bio18| $\sim1000$      |
| Precipitation of coldest quarter [mm]        | bio19| $\sim1000$      |
| Soil classification, TAXOUSDA [predicted most probable class] | Sclass | 250 $^b$       |
| Soil depth (absolute depth to bedrock) [cm]  | Sdepth| 250 $^b$       |
| Soil organic content (fine earth fraction) [g/kg]| Sorgco| 250 $^b$       |

Units indicated inside brackets. Data resolution expressed in m$^2$. Sources of information are indicated below:

- $^a$ Worldclim.org.
- $^b$ Soilgrids.org.

[https://doi.org/10.1371/journal.pone.0190092](https://doi.org/10.1371/journal.pone.0190092)
we followed the recommendations of Merow et al. [56]. To minimize multicollinearity and model overfitting, a principal component analysis (PCA) was performed with all predictor variables and those highly correlated were removed ($r^2 > 0.8$). In addition, to account for occurrence record sampling biases, we built a simple biased raster file [56] based on Ecuadorian access roads, where we assigned a 2.5 km buffer around roadways and assumed that the probability of finding records inside this zone was double that of the surrounding area. We programmed the console to run 5 replicates of each model and left the remaining settings at default values.

The outcomes of the environmental niche modeling were converted to binary-type using the 10% training presence threshold as the absence criteria. Analyses of rasters, vectors, and area calculations were performed using QGIS 2.2.0, including the principal component analysis through the python plugin PCA v0.3. Model evaluation was performed by the area under the receiver-operator (AUC) of the receiver operator characteristic (ROC), a debated, but prevalent rank-based metric to assess predicted distribution model accuracy [56, 57]. This metric is the probability that a random presence locality is ranked higher than a random absence location. An AUC value of 0.5 indicates that the prediction is not better than random; < 0.5 is worse than random; 0.5–0.7 indicates poor performance; 0.7–0.9 represents reasonable or moderate performance; and > 0.9 signifies high performance [58].

Calculation of deforestation and climate change metrics

To calculate the deforestation and climate change metrics affecting each species, we used 2008 and 2014 public land-cover and land-use data from the Ecuadorian Ministry of Environment that is available in digital cartography form [13]. Next, we used forest loss and/or remnant
native forest information to mask the binary outcomes of individual species distribution models, for both present and future models. Thus, we obtained species-specific approximations of the affected area for the periods: 2008–2014 and 2014–2050. Given that estimated area loss attributed to deforestation was calculated for a 6-year period and the area threatened by climate change was determined for a 36-year period, we standardized values by calculating annual rates of loss and compared values for all 17 species using a paired t-test.

Distribution area measurements and landscape metrics for each species were computed by the python plugin LecoS v2.0.7 [59], an alternative to the more comprehensive FRAGSTATS, that has the advantage of working within the QGIS processing framework. For this, we transformed the deforestation and climate change masked outputs to raster format using a $250 \times 250$ m cell size, which set the minimum detectable area for any landscape metric to $0.0625 \text{ km}^2$. In addition, we also superimposed (i.e., stacked) species distributions in native forests that were unthreatened by climate change, to evaluate how the modeled species overlap differed from the deforestation and climate change threats.

To summarize and visually compare the changes in distribution attributed to deforestation and climate change, we calculated the core distributional shifts (i.e., area centroids) for each species according to two time frames: 2008–2014 and 2014–2050. For this, we used SDMtoolbox [60], a python-based GIS toolkit for automating analyses in ecology and species distribution models. This analysis reduces the distribution area to a single point and creates a line that represents the magnitude and direction of change. In this section, climate change vector lengths were corrected to reflect six years of change to match the deforestation time interval.

**Results**

**Species records and predictor variables**

Presence records for all 17 species ranged from 7 to 62 unique points per raster cell, with more than two-thirds of all species having $> 30$ records, generally perceived as an optimal number of locations to generate consistent models [55]. From the pool of 22 predictor variables, eight were selected based on their correlation coefficients and were considered biologically meaningful for dry forest ecosystems (Table 3). The designated variables to produce the models included annual mean temperature, mean diurnal range, precipitation seasonality, wettest quarter precipitation, driest quarter precipitation, soil classification, depth to bedrock, and soil organic content.

Model outcomes consistently indicated that the two most important predictor variables were the precipitation of driest quarter and soil classification, continuous and categorical variables, respectively. In contrast, variables that contributed least to the model were precipitation of wettest quarter and precipitation seasonality (Table A in S1 Appendix).

**Species distribution modeling and evaluation**

We obtained robust evaluation metrics for 14 species ($\text{AUC} \geq 0.90$), while the remaining three species had lower AUC values (between 0.79 and 0.88), which are still considered an indicative of reasonable to moderate performance (Table 4).

In addition, to corroborate the model outcomes, we built a stacked map of all species present in remnant native forests (Fig 2), which revealed a concentration of species distributions in southwestern Ecuador that agreed with ecological descriptions.

**Assessment of deforestation and climate change**

Binary potential distributions were combined with land-cover and land-use data to obtain the following three area estimates for each species: area lost by deforestation, remnant native forest
### Table 3. Correlation matrix of predictor variables.

|       | Temperature | Precipitation | Topography |
|-------|-------------|---------------|------------|
| bio01 | 0.76 0.88   | 0.98 0.97     |            |
| bio02 | 0.98 0.72   | 0.77 0.99     |            |
| bio03 | 0.74 0.92   | 0.87 0.88     |            |
| bio04 | 0.86 0.84   | 0.87 0.85     |            |
| bio05 | 0.96 0.93   | 0.99 0.99     |            |
| bio06 | 0.80 0.98   | 0.98 0.98     |            |
| bio07 | 0.88 0.89   | 0.89 0.88     |            |
| bio08 | 0.99 1.00   | 0.80 0.87     |            |
| bio09 | 0.99 1.00   | 0.80 0.87     |            |
| bio10 | 0.99 0.99   | 0.80 0.87     |            |
| bio11 | 0.80 0.87   | 0.44 0.82     |            |
| bio12 | 0.95 0.75   | 0.95 0.77     |            |
| bio13 | 0.55 0.77   | 0.99 0.99     |            |
| bio14 | 0.16 0.56   | 0.59 0.55     |            |
| bio15*| 0.75 0.17   | 0.70 0.22     |            |
| bio16*| 0.57 0.97   | 0.53 0.35     |            |
| bio17*| 0.56 0.80   | 0.08 0.36     |            |
| bio18 | 0.44 0.32   | 0.82 0.51     |            |
| bio19 |              | 0.09 0.28     |            |
| Sclass*|        | 0.45 0.33     |            |
| Sdepth*|         | 0.58          |            |

Highly correlated variables were ignored (correlation > 0.8).

* Selected variables to produce the models included annual mean temperature (bio1), mean diurnal range (bio2), precipitation seasonality (bio15), precipitation of wettest quarter (bio16), precipitation of driest quarter (bio17), soil classification (Sclass), absolute depth to bedrock (Sdepth), and soil organic content (Sorgco).
area unthreatened by climate change, and area threatened by climate change. An illustration for one species is shown in Fig 3, and maps for all 17 species are listed in Table B in S1 Appendix.

For all 17 tree species except *Cavanillesia platanifolia*, deforestation consistently represents a greater threat in reducing distribution areas compared with climate change (CCSM4.0, RCP 8.5, 2050). Our annual loss estimates from deforestation ranged from approximately 9 to 200 km²/year across species, while estimated annual loss from climate change ranged from 4 to 60 km²/year. Results from the paired t-test identified a significant difference between the distribution-loss associated with deforestation and the loss attributed to climate change (p-value = 0.001, t = 3.93, and df = 16). Similarly, t-tests for landscape variables also showed significance, particularly for landscape cover (p-value < 0.0001), edge length (p-value = 0.01), number of patches (p-value = 0.0002), and mean patch area (p-value < 0.0001) (Table 5). In summary, there is evidence that all tree species are highly affected by deforestation processes, but there are three species, namely *Albizia multiflora*, *Ceiba trichistandra*, and *Cochlospermum vitifolium* that might be susceptible to additional climate change pressure, which strongly affects the area and number of patches of these species.

In addition, seven species displayed an increase in their future distributions after area gain was accounted for in models, with an estimated area gain ranging from 22 to 74 km²/year. These seven species (i.e., *Caesalpinia glabrata*, *Coccoloba ruiziana*, *Colicodendron scabridum*, *Cordia macrantha*, *Guazuma ulmifolia*, *Handroanthus chrysanthus*, and *Prosopis juliflora*) also showed corresponding increases in their total edge lengths and number of patches, assuming recolonization.

Stacked distributions to explore species overlap predictions also suggest that deforestation has a greater effect on annual loss than climate change. For instance, areas with higher species overlap (13 to 17 species) decrease at approximately 34 km²/year owing to deforestation, compared to less than 6 km²/year owing to the future climate scenario (Table C in S1 Appendix).

### Table 4. Average model AUC values.

| Species                        | AUC     | SD      |
|--------------------------------|---------|---------|
| *Albizia multiflora*           | 0.995   | ± 0.014 |
| *Bursera graveolen*            | 0.948   | ± 0.021 |
| *Caesalpinia glabrata*         | 0.936   | ± 0.018 |
| *Cavanillesia platanifolia*    | 0.981   | ± 0.011 |
| *Ceiba trichistandra*          | 0.958   | ± 0.022 |
| *Chloroleucon mangense*        | 0.944   | ± 0.043 |
| *Coccoloba ruiziana*           | 0.926   | ± 0.018 |
| *Cochlospermum vitifolium*     | 0.920   | ± 0.054 |
| *Colicodendron scabridum*      | 0.950   | ± 0.029 |
| *Cordia macrantha*             | 0.900   | ± 0.069 |
| *Erythrina velutina*           | 0.925   | ± 0.062 |
| *Geoffroea spinosa*            | 0.926   | ± 0.041 |
| *Guazuma ulmifolia*            | 0.797   | ± 0.019 |
| *Handroanthus chrysanthus*     | 0.869   | ± 0.029 |
| *Loxopterygium huasango*       | 0.965   | ± 0.031 |
| *Piscidia carthagenensis*      | 0.933   | ± 0.029 |
| *Prosopis juliflora*           | 0.884   | ± 0.038 |

AUC, or area under the receiver-operating characteristic (ROC) curve. Values correspond to the mean of 5 model runs.

https://doi.org/10.1371/journal.pone.0190092.t004
Moreover, in southwestern Ecuador, deforestation seems to reduce the number of species at all elevations, while the future climate scenario appears to shrink species overlaps primarily at higher altitudes and not in the lowlands, where projections show an increase in the number of overlapping species. In addition, the spatial deforestation hotspot does not coincide with the hotspot of species loss attributed to climate change (Figure D in S1 Appendix).

Finally, based on the centroids of distribution, we produced a summary map showing the core distributional shifts for all species considered in this study (Fig 4). From this map, deforestation appears to push the majority of species distributions northward at a mean rate of 0.8 km/year. However, the climate change projections result in a set of species headed southward at a mean rate of 1.4 km/year (i.e., Albizia multiflora, Cochlospermum vitifolium, Erythrina...
velutina, Geoffroea spinosa, Bursera graveolens, Piscidia carthagenensis, Ceiba trichistandra, and Loxopterigyum huasango), while another group shows a northern habitat displacement migration at 0.7 km/year (i.e., Guazuma ulmifolia, Handroanthus chrysanthus, Prosopis juliflora, Caesalpinia glabrata, Cordia macrantha, Colicodendrum scabridum, Cocosolba ruiziana, and Cavanillesia plataniifolia). Overall, the core distributional shift directions indicate that the pathways for habitat displacements are species-specific and display divergent patterns.

**Discussion**

Species distribution models are useful tools for conservation planning, resource management, and policy development [61], although the implementation and interpretation of these models may present challenges [62, 63]. Our study represents the first attempt to use exploratory

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**Fig 3. Potential distribution areas for Albizia multiflora.** The map on the left shows the species model for continental Ecuador, while the enlarged map on the right highlights species presence calculations in southwestern Ecuador. Green, black, and gray colors represent areas of remnant native forest, areas lost to deforestation, and remnant forest areas threatened by climate change, respectively.

https://doi.org/10.1371/journal.pone.0190092.g003
Table 5. Annual landscape change attributed to deforestation and climate change.

| Species                           | Deforest. | Climate | Deforest. | Climate | Deforest. | Climate | Deforest. | Climate | Deforest. | Climate | Deforest. | Climate | Deforest. | Climate | Deforest. | Climate |
|-----------------------------------|-----------|---------|-----------|---------|-----------|---------|-----------|---------|-----------|---------|-----------|---------|-----------|---------|-----------|---------|
| Coccoloba ruiziana                | -0.05     | -0.03   | -0.04     | -0.01   | +0.65     | -0.06   | -0.02     | -0.01   | +2.3      | -1.5    | -97.98    | -21.49  | 0.07      | -0.02   |
| Chloroleucon mangense             | -0.03     | -0.02   | -0.02     | -0.01   | -13.62    | -30.51  | +1.1      | -0.5    | -82.30    | -6.97   | -0.07     | -0.02   | 0.01      | -0.01   |
| Cordia macrantha                  | -0.08     | +0.05   | -0.05     | +0.02   | -40.45    | +74.63  | +1.0      | +0.3    | -102.62   | -1.12   | -0.07     | -0.01   | 0.01      | -0.01   |
| Colicodendron scabridum           | -0.03     | +0.03   | -0.03     | +0.03   | -66.50    | +132.69 | +1.2      | +1.2    | -14.47    | +19.53  | 0.07      | -0.01   | 0.01      | -0.01   |
| Cochlodermum vitifolium           | -0.09     | -0.05   | -0.09     | -0.05   | -83.45    | -201.47 | +2.0      | -0.7    | -101.17   | +3.91   | -0.11     | -0.01   | 0.01      | -0.01   |
| Erythrina velutina                | -0.06     | -0.01   | -0.06     | -0.01   | -80.55    | -24.78  | +2.0      | 0.00    | -105.20   | +0.88   | -0.10     | -0.01   | 0.01      | -0.01   |
| Geoffroea spinosa                 | -0.05     | -0.00   | -0.05     | -0.00   | -94.38    | -7.09   | +1.6      | -0.05   | -101.40   | +4.05   | -0.08     | -0.02   | 0.01      | -0.01   |
| Guazuma ulmifolia                 | -0.11     | +0.04   | -0.11     | +0.04   | -164.00   | +217.36 | +1.0      | +0.6    | -9.22     | +46.94  | -0.07     | -0.03   | 0.01      | -0.01   |
| Handroanthus chrysanthus          | -0.13     | +0.01   | -0.13     | +0.01   | -148.97   | +88.87  | +1.4      | +0.3    | -27.85    | +8.74   | -0.09     | -0.01   | 0.01      | -0.01   |
| Loxostegium huasango              | -0.03     | -0.01   | -0.03     | -0.01   | -7.53     | -27.09  | +2.0      | -0.7    | -99.47    | -14.93  | 0.12      | -0.03   | 0.01      | -0.01   |
| Piscidia carthagensis             | -0.04     | -0.01   | -0.04     | -0.01   | -38.88    | -48.38  | +0.4      | -0.4    | -97.73    | -1.08   | 0.04      | -0.01   | 0.01      | -0.01   |
| Prosopis juliflora                | -0.08     | +0.03   | -0.08     | +0.03   | -57.20    | +174.87 | +1.0      | +1.2    | -26.38    | +10.62  | -0.09     | -0.01   | 0.01      | -0.01   |

Minimum patch area detected for all species was equal to the grid cell size (0.0625 km²). The proportion (%) of total area was calculated based on the environmental space used during modeling (approximately 130,000 km²). The number of patches depicts the percent change relative to all patches. Asterisks indicate statistical difference:

(*) represents significance at p-value = 0.01.

(**) indicates high significance at p-value = 0.001.

(***) very high significance at p-value < 0.0001.

https://doi.org/10.1371/journal.pone.0190092.1005

In our models, the precipitation of the driest quarter and soil class were the two variables that exerted the largest influence on the distribution of dry forest tree species. Precipitation during the dry season has been suggested as the most critical element for determining vegetation patterns [64] because water remains a critical factor driving basic physiological processes of tropical trees [65]. Conversely, soil characteristics differentiate seasonally dry forests from savannas, even under similar climate regimes [66] and have been shown to affect the vegetation structure in Northern Peruvian dry forests [67]. In the context of major environmental processes, our study region represents typical characteristics for tropical dry forests.

With regards to model validation, we focused on the operational capability of the outputs rather than testing their inferences about the real system (e.g., Rykiel [68]). Therefore, validations used AUC values, which also show a degree of correlation with other model performance metrics [69]. In all cases, we obtained high AUC values with considerably low replicate variation (< 0.07 SD), which is half of the recommended value by other studies for “accurate and
stable” models that meet practical needs [70,71]. Furthermore, as additional support to our models, the highest concentration (i.e., stacked distribution) of all 17 species agrees with the southwestern Ecuador habitat description (as depicted by Aguirre et al. [36]), providing a more practical validation of model accuracy.

According to a study based on satellite and aerial photography data, deforestation in southwestern Ecuador dry deciduous forests was approximately 29 and 57 km²/year from 1976 to 1989 and 1989 to 2008, respectively [17]. However, official reports indicate a 15 km²/year mean annual deforestation rate between 2008 and 2014 across the study area (i.e., environmental space used during modeling) [14]. According to our species distribution models, the average deforestation loss for the 17 tree species reached 71 km²/year (± 43.9 SD). Our results support previous direct and indirect measurements that identified deforestation as the greatest
threat to dry forest ecosystems in Ecuador [6,17]. Conversely, our area loss calculations resulting from climate change projections reached 21 km²/year (± 26.4 SD) for the same tree species without accounting for potential distribution gain (i.e., potential new favorable areas). For instance, the difference in area-changes attributed to deforestation and climate remained statistically significant when we limited the comparison to potential area loss (excluding area gain) from the future climate scenario. Area gain can be disregarded owing to low regeneration processes and conflicting land uses that might strongly inhibit natural succession.

In summary, in terms of relative area change, deforestation and climate change represented average annual area reductions of 1.4% and 0.6%, respectively. Thus, in terms of spatial coverage and effect severity, deforestation may pose a higher threat to species distributions than climate change. However, the spatial disaggregation of both processes indicates that climate change may affect a greater number of forest areas, which are not subject to conversion.

Moreover, our species overlap results in the future climate scenario (Figure D in S1 Appendix) indicate reductions in high overlap (13–17 species) areas and expansions in low and medium overlap areas (1–6 and 7–12 species), in particular, at elevations below 1000 m in southwestern Ecuador. The apparent progression into the lowlands can be explained by the future increase in the precipitation predictor variables (e.g., bio16 and bio17), considering that other climate models also predict a precipitation increment in this area [20,22]. Finally, the magnitude and direction of the core distributional shifts differ by threat and species; this fact supports the notion of climate change pushing towards novel species compositions [72,73]. One explanation for the observed divergent directions of deforestation and climate change may be that each threat has different underlying mechanisms; deforestation pressure is determined by land use, access roads, and other density-dependent human factors, while climate change pressure is estimated solely by changes in environmental variables and responses will depend on species-specific evolutionary histories and physiological requirements. Furthermore, northward or southward species migrations might be the only possible routes for species in southwestern Ecuador given the altitude in these paths gradually increases, while two major geographic barriers, the Andes and the Pacific Ocean, lie to its East and West, respectively.

Furthermore, as our projections predict range expansions for some species and range contractions for others (in addition to the range shifts induced by deforestation and climate), tree species distributions are potentially susceptible to a myriad of unpredictable community-level effects. Thus, as an alternative to evaluating the effects of species re-assemble on the dry forests, we propose a functional traits approach that emphasizes at the level of response traits and effect traits. As derived by Suding et al. [29], when the geographical displacement of each response group contains species from each effect group, ecosystem function resilience is expected. Conversely, the occurrence of new species assemblages under future climate scenarios may have consequences for the provision of supporting services, as well as regulating services. For instance, Sakschewski et al. [74] provided theoretical evidence that plant trait diversity may enable large-scale ecosystems to adjust to new climate conditions through competition. Moreover, in the particular case of novel combination of species, Lugo [75] asserted that tropical novel forests might behave similarly to native forests, specifically in terms of soil protection, nutrient cycles, wildlife support, carbon storage, and watershed function maintenance. Thus, we believe that changes in species composition in the dry deciduous forests of Ecuador represent challenges for forest management and requires exploration of new strategies to maintain the long-term provision of ecosystem services.

Measures to avoid deforestation should be promoted in a deforestation hotspot in the southernmost coastal region [76] and below 1000 m of elevation according to our study. Concurrently, conservation should be encouraged in the southwestern border with Peru [77], where our results predict the highest concentration of species of this forest type. Further
research must incorporate deforestation data and modeling scenarios for northern Peru where the underlying causes for deforestation are similar [18,78], and social processes such as demographics, economics, and policy play an important role [76]. Therefore, strategies to avoid deforestation and promote conservation include expanding protected areas, biological corridors, species-specific timber use regulations and flagship species valorizations (e.g., *Cavanillesia platanifolia*). However, regardless of the conservation strategy, successfully achieving these goals will require both private landowner and local community participation [79].

Less than 10% of the original extent of dry deciduous forests remains in the neotropics [3] and only 2.3% is under conservation in Ecuador [6]. Therefore, it is likely that the Aichi target Nr. 11, which aims to protect 17% of terrestrial land by 2020 [80], may not be achieved for this biome in Ecuador. The results of this study can be used as an additional resource for decision-making regarding the improvement or expansion of existing protected areas and biological corridors. For instance, there is almost no mid to high altitude (100–1000 m) land under conservation in southwestern Ecuador with the exception of scarce private reserves such as La Ceiba and Laipuna. Our study highlights conservation area weaknesses in southern Ecuador, in agreement with Cuesta et al. [77]

If the protection of individual species is prioritized, attention should be given to *Albizia multiflora*, *Ceiba trichistandra*, and *Cochlospermum vitifolium* because they are heavily threatened by both deforestation and climate change. Moreover, because these species can be used for timber and in agroforestry systems [43], we recommend a special focus on sustainable management practices and gene pool conservation. With regard to species heavily threatened by deforestation and with high and valuable usage as timber and wood products (i.e., *Cordia macrantha*, *Guazuma ulmifolia*, *Handroanthus chrysanthus* and *Prosopis juliflora*), we recommend species-specific measures for their conservation, such as potential genecological zone identification, potential subpopulation evaluation, and gene pool variability tests. In combination with enrichment planting efforts, additional measures for sustainable management of these species include the establishment of seed orchards and in vitro propagation.

Conclusion

Using presence-only modeling and native forest masks from the Ecuadorian Ministry of Environment, we obtained approximations of characteristic tree species distributions in the dry deciduous forest of southwestern Ecuador, which are threatened by deforestation and climate change. Our estimates indicate that deforestation affects more spatial range than climate change, even under an extreme climate change scenario. Despite this result, climate change may cause additional stress at the species and community levels. Special attention to *Albizia multiflora*, *Ceiba trichistandra*, and *Cochlospermum vitifolium* populations may be required because our results reveal that these species are vulnerable to both deforestation and climate change. The diverging displacement shifts of a large number of species may indicate the commencement of plant community disaggregation and a transition toward novel ecosystems. However, further research is required to discern the effects on the synecology, resilience, and ecosystem services of dry forest ecosystems.

Our results indicate that the effects of climate change result bigger in terms of distributional shifts but that deforestation affects more surface area. Therefore, as climate change adaptation and deforestation reduction measures in Ecuador likely will not match spatially, deforestation reduction should be prioritized over climate change adaptation. Similar cases of habitat loss due to these two threats may be occurring in other ecosystems in the Tropics, where annual deforestation rates are considerably far more severe [81] and novel species assemblages (i.e., redistribution) are not sufficiently considered in policies and international agreements [82].
Supporting information
S1 Dataset. Occurrence records.
(XLS)
S1 Appendix. Additional information.
(DOCX)

Acknowledgments
We thank Z. Aguirre at the Universidad Nacional de Loja for complementing data on species occurrences and the Ecuadorian Ministry of Environment for providing the research permit 017-2013-DPL-MA. We also acknowledge the cooperation of Nature and Culture International.

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