Measuring Forest Biodiversity Status and Changes Globally

Samantha L. L. Hill¹,²*, Andy Arnell¹†, Calum Maney¹, Stuart H. M. Butchart³,⁴, Craig Hilton-Taylor⁵, Carolyn Ciciarelli⁶, Crystal Davis⁶, Eric Dinerstein⁷, Andy Purvis²,⁸ and Neil D. Burgess¹,⁴,⁹

¹ UN Environment World Conservation Monitoring Centre (UNEP-WCMC), Cambridge, United Kingdom, ² Department of Life Sciences, The Natural History Museum, London, United Kingdom, ³ BirdLife International, The David Attenborough Building, Cambridge, United Kingdom, ⁴ Department of Zoology, University of Cambridge, Cambridge, United Kingdom, ⁵ IUCN Global Species Programme, Cambridge, United Kingdom, ⁶ WRI, Washington, DC, United States, ⁷ RESOLVE, Washington, DC, United States, ⁸ Department of Life Sciences, Imperial College London, Ascot, United Kingdom, ⁹ CMEC, University of Copenhagen, Copenhagen, Denmark

The world’s forests are crucially important for both biodiversity conservation and climate mitigation. New forest status and forest change spatial layers using remotely sensed data have revolutionised forest monitoring globally, and provide fine-scale deforestation alerts that can be actioned in near-real time. However, existing products are restricted to representing tree cover and do not reflect the considerable spatial variation in the biological importance of forests. Here we link modelled biodiversity values to remotely sensed data on tree cover to develop global maps of forest biodiversity significance (based on the rarity-weighted richness of forest mammal, bird, amphibian and conifer species) and forest biodiversity intactness (based on the modelled relationship between anthropogenic pressures and community intactness). The strengths and weaknesses of these products for policy and local decision-making are reviewed and we map out future improvements and developments that are needed to enhance their usefulness.

Keywords: forest cover, remote sensing, biodiversity, Biodiversity Intactness Index (BII), IUCN Red List

INTRODUCTION

The world’s forested biomes are crucially important for terrestrial biodiversity but humanity’s growing demands for resources have led to the removal of natural forest for agriculture and the degradation of forest landscapes through hunting and timber removal, fragmentation, pollution, and other human impacts (Foley et al., 2005; Song et al., 2018). Such pressures are impacting forest biodiversity (Newbold et al., 2013, 2016; Phillips et al., 2017), as many sensitive species are reliant upon intact forest landscapes and “primary” forests (Gibson et al., 2011; Betts et al., 2017).

Remote sensing has been established for over 40 years as a vital tool for understanding how land cover and land use are changing over time. As satellite technology and analytical methodologies have improved, computing power has increased and data have become increasingly freely accessible, the available products have increased greatly in sophistication and spatial and temporal resolution. For forest biomes, these developments have resulted in the creation of the world’s first global 30-m resolution tree-cover status and change product (Hansen et al., 2013), which in turn facilitated the development of a suite of academic papers and freely available products within the “Global Forest Watch” partnership, covering forest status and trends (Hansen et al., 2013), forest carbon (Tyukavina et al., 2015), and forest tree height (Hansen et al., 2016).
However, measurements of tree presence and absence alone are a poor surrogate for biodiversity value and biodiversity loss (Tropek et al., 2014). Biodiversity, including measures such as species diversity, species endemism, or genetic diversity, is unequally distributed, with major biological differences among and within the forested ecoregions of the world (Olson et al., 2001; Dinerstein et al., 2017). This variation means that equally sized areas of tree cover, mapped through remote sensing, often differ dramatically in biological value (however this may be defined) among different continents, and at different locations, latitudes, and elevations. While these patterns have been explored for particular taxa by linking species distribution data to global tree cover loss data (e.g., Buchanan et al., 2011), until now there has been no global analysis of biological values of forests (covering a broad suite of taxa) linked to the newly available tree-cover data enabled by Hansen et al. (2013).

In this paper we present approaches that estimate how forest cover change affect two aspects of biodiversity value, through a combination of modelled biodiversity data that are spatially linked to remotely sensed data. The first approach uses data from the IUCN Red List (www.iucnredlist.org), a widely used dataset relating to species’ risk of extinction, including spatial distribution maps for each species. We use these maps to estimate and map biodiversity significance, based on rarity-weighted richness (through aggregate scores of range-size rarity), for all forest-dependent mammals, birds, amphibians and conifers across the forest regions of the world, highlighting locations that make a disproportionate total contribution to the global distributions of these species (Williams et al., 1996).

The second approach uses data from the PREDICTS database, a large taxonomically and geographically representative global database of the impact of anthropogenic pressures on local biodiversity (Hudson et al., 2017). This database is analysed statistically to estimate and map biodiversity intactness, following the framework outlined by Newbold et al. (2016) and Purvis et al. (2018), reflecting the proportion and abundance of a location’s original forest community that remain.

These two layers are both informative about how different facets of forest biodiversity are distributed; considering them together provides added information, such as highlighting areas that are potentially suitable for restoration or conservation.

**MATERIALS AND METHODS**

**Tree Cover Change**

Gridded tree cover, tree-cover loss and tree-cover gain data as described by Hansen et al. (2013) were accessed in December 2017 from the Google Earth Engine (Gorelick et al., 2017) data repository. However, tree cover for 2010 was accessed from the USGS Land Cover Institute (2017). Thresholds of minimum crown cover/closure (MCC), from here on referred to as tree cover, for delimiting forested land vary greatly within scientific literature (Lund, 2002, 2015; Magdon and Kleinn, 2013; Magdon et al., 2014). The US National Vegetation Classification System defines forests as areas with a 60% tree cover (Grossman et al., 1998), UNEP (2001) use a threshold of 40% tree cover to distinguish closed forests, Kohl and Päivinen (1996) use a threshold of 20% tree cover for distinguishing European forests and the Vegetation Resources Inventory (for Canadian forests) defines a treed area as having 10% tree cover (Sandvoss et al., 2005). The FAO uses a threshold of 10% MCC to determine whether an area has been deforested (FAO, 2000) in contrast to Hansen et al., 2010 who suggest that a value <25% MCC can be used for measuring global deforestation across all biomes, due to its ability to “identify tall woody vegetation unambiguously in multispectral imagery.” Tropical and more forested countries typically use higher tree cover in their national assessments relative to countries primarily outside of forest biomes, for example, Zimbabwe defines forest using a tree cover of 80% (Magugu and Chitiga, 2002) whereas Australia uses a tree cover of 20–50% (ABARES, 2018). The scale of the reference area is important, with forest area positively correlated to reference area size (Magdon and Kleinn, 2013). For this study we selected the more conservative value of 60% tree cover to indicate the presence of forest; however, for comparison, we also calculated biodiversity intactness and biodiversity significance maps using a forest definition of 25% tree cover and have included the results within the Supplementary Material (SM Figures 1–3).

Tree cover data may not distinguish between natural forests and forests that have been planted for human uses, yet the biodiversity value of such plantation forests is typically considerably lower (e.g., Gibson et al., 2011; Newbold et al., 2015; Phillips et al., 2017). If the rotation length of plantations is greater than the time span of the tree-cover data, and other criteria such as height and density of trees are met, then treating plantation forests as though they are natural would lead us to overestimate their biodiversity value. To distinguish between natural forests and plantation forests we used the Spatial database of Planted Trees (SDPT; Harris et al., 2019), a compilation of mapped and modelled plantation data from multiple countries, that focuses on including intensively managed plantations and excludes semi-natural forests with intensive natural regeneration. Plantations in China and Papua New Guinea in the SDPT could not be included due to data restrictions. Further details of spatial datasets used in this study can be found in SM Table 2.

**Biodiversity Significance**

Tree cover data for 2018 and annual tree-cover loss data between 2000 and 2018 were used in this analysis.

Using the IUCN Red List dataset (www.iucnredlist.org), we extracted spatial data on distributional boundaries and tabular data on habitat preferences and elevation limits for birds, mammals, and amphibians (provided by IUCN in October 2017) and conifers (in November 2017). Following Tracewski et al. (2016), we defined forest-dependent species as those birds coded by BirdLife International as having high or medium forest dependency (Buchanan et al., 2008; Bird et al., 2012), and those mammals and amphibians coded by Rondinini et al. (2011) and Ficetola et al. (2015), respectively, as having high forest dependency. Differences in forest-dependency selection criteria (i.e., high, medium etc.) between birds and other taxa reflect variation in how dependency is defined. We defined forest-dependent conifers as those coded for forest habitats only (ICUN,
This produced a list of forest-dependent mammals (n = 1,463), amphibians (n = 3,563), birds (n = 6,841), and conifers (n = 393), totalling 12,260 species for further analysis. These taxonomic classes are the only ones in which all or nearly all terrestrial species have been assessed for the IUCN Red List and for which spatial distribution maps are available (BirdLife International Handbook of the Birds of the World, 2016; IUCN, 2017).

### Biodiversity Significance of Remaining Forest in 2018

To calculate the significance of remaining forest habitat for each forest-dependent species, we produced an “Extent of Suitable Habitat” (ESH) map now known as Area of Habitat: (Brooks et al., 2019) through exclusion of areas within the species’ distribution with (1) <30% tree cover in 2000, or (2) any tree cover loss between 2000 and 2018, and/or (3) altitude outside the species’ elevational limits as defined by IUCN (2017). Spatially explicit elevation data was obtained from the GMTED2010 dataset (Danielson and Gesch, 2011). We also removed areas of plantations based on the SDPT dataset (Harris et al., 2019), except for ~20% of species that were listed with affiliations coded as “Suitable” for either Plantations or Subtropical/Tropical Heavily Degraded Former Forest habitats (IUCN, 2017). For such species, composed primarily of birds, including plantation areas in the ESH calculations represents their ability to utilise both habitats. Where these are present within the species’ range this will then lead to lower significance scores in natural forest areas, reflecting their lower dependence on natural forest habitat.

The range-size rarity for each species within a grid cell was calculated as the contribution of each ~30 m cell toward the global extent of suitable habitat for the species (i.e., the inverse of the ESH within each species’ distribution). For those species coded with having different seasonal distributions, we calculated range-size rarity scores for each of these distributions separately. Range-size rarity scores were summed across all species present within a grid cell to give an overall rarity-weighted richness score, or the “biodiversity significance” of the cell. Cells with high values for biodiversity significance typically contain more species for which the cell comprises a larger proportion of their global distribution. Loss of forest in such cells is therefore of disproportionate significance in terms of loss of biodiversity (at least for the taxonomic groups considered). We note that there are many alternative ways of estimating biodiversity importance, but we use the term “biodiversity significance” for this metric as “rarity-weighted richness,” “range rarity,” and related terms are not widely understood by non-specialists.

We converted species’ distribution polygons to raster format at ~1 km resolution. This resolution is more relevant to the accuracy of the species distribution data than the high resolution (~30 m) forest data. However, to calculate the area of forest habitat (ESH) per species, and for creating final outputs, we used the forest data at ~30 m resolution. Therefore, the coarser resolution of the underlying species data remains in the final outputs, i.e., biodiversity significance values for forest pixels do not vary within ~1 km cells. All analyses were completed in Google Earth Engine (Gorelick et al., 2017).

### Significance of Forest Loss 2000–2018

To calculate the significance of loss, we followed a similar approach, but instead calculated range-size rarity values based on ESH using forest cover from 2000. This shows how significant a pixel of forest was for a species in 2000. We then summed this value across all species per cell to calculate the biodiversity significance of forest cells lost during 2000–2018.

### Biodiversity Intactness

To model biodiversity intactness, we analysed the PREDICTS database which comprises well over 3 million rows of geographically and taxonomically representative data of land-use impacts to local terrestrial biodiversity derived from the primary literature and other databases (Hudson et al., 2017). As our intention was to explore the impacts of forest change, the database was first subset to sites within forested biomes (Olson et al., 2001) yielding a dataset from over 550 studies encompassing over 19,700 sites, 2.3 million observations and ~25,000 taxa. A generalised linear mixed-effects model framework was used to assess how community abundance was impacted by land use and human population density (extracted from HYDE 3.1: Klein Goldewijk et al., 2011) within forest biomes, following the methods outlined in Newbold et al. (2016). Briefly, the random-effects structure included a study-level random effect to account for the innate variability between samples collected using different methodology and focused upon varying taxa, and a biome-level random effect with human population density as a random slope to account for differences in the influence of human population density among biomes. A random slope of land use within study accounted for the study-level variation in the influence of land use on community abundance. The fixed-effects structure was selected using backwards stepwise selection using likelihood ratio tests to select the most appropriate model. The model of compositional similarity followed the methodology of De Palma et al. (2018) (see also Newbold et al., 2019, Nature Ecology and Evolution). In brief, a matrix of paired site-level comparisons was first prepared where all sites within a study were compared to all minimally-used primary vegetation sites within that same study. The asymmetric Jaccard Index was employed to calculate abundance-based compositional similarity for each paired comparison, and a mixed-effects model was fitted to predict the influence of land use, the environmental distance between sites and the geographic distance between sites, on logit-transformed compositional similarity. The community abundance and compositional similarity model coefficients were multiplied to produce the abundance-based Biodiversity Intactness Index (BII), our biodiversity intactness metric.

To estimate how the spatial patterns of forest change have affected biodiversity intactness, we used the layers of tree cover, loss, and gain during 2000–2014 (Hansen et al., 2013 updated) to produce a map of land-cover change. The downscaled land-use map produced by Hoskins et al. (2016) provides data on anthropogenic land uses (such as cropland, pasture and urban) at a spatial grain of ~1 km²; we used this map to infer the land use after deforestation. Each deforested pixel was allocated the proportions of the anthropogenic land-use categories within the corresponding grid cell of the downscaled land-use map.
We defined 12 Boolean conditions describing important boundaries within the input data layers (SM Table 1). Mutually exclusive expressions were then built to describe each of the land covers as a function of the Boolean variables (Equations 1–7). It was important to include a distinction between tropical and temperate areas due to the input data informing the model coefficients. The PREDICTS database, which populates and informs the models for BII, uses different classification schemas for tropical and temperate environments: specifically, tropical secondary forest characterised as "young" cannot be older than 10 years, whilst in temperate biomes "young" secondary forests can be up to 30 years old. This reflects the speed with which succession takes places in temperate vs. tropical areas, and especially how intactness can recover faster in the tropics. The expressions below reflect this by introducing variable I (SM Table 1), which differentiates tropical and temperate biomes.

\[ A_{\text{Primary, Mature Secondary}} = P' \cdot T' \cdot L' \cdot G' \cdot V' \] (1)

\[ A_{\text{Intermediate Secondary}} = P' \cdot (T' \cdot L' \cdot (G + V)) + ((T' + L) \cdot D' \cdot R' \cdot I) \] (2)

\[ A_{\text{Young Secondary}} = P' \cdot (T' + L) \cdot D' \cdot ((R - I) + I') \] (3)

\[ A_{\text{Cropland}} = P' \cdot (T' + L) \cdot D \cdot C \] (4)

\[ A_{\text{Pasture}} = P' \cdot (T' + L) \cdot D \cdot B \] (5)

\[ A_{\text{Urban}} = P' \cdot (T' + L) \cdot D \cdot U \] (6)

\[ A_{\text{Plantation}} = P \] (7)

A primary or mature secondary forest is defined as an area which is not a plantation, meets the forest cover in 2010 criteria, and which has not experienced forest loss (Equation 1). It must also not have recorded growth, and tree cover must have been stable (±20%) between 2000 and 2012 (Equation 1). An intermediate secondary forest is defined as an area which is not a plantation and which is either: an area which experienced loss or does not meet 2010 cover criteria, but which is not disturbed and is both tropical and over 10 years old; or which meets cover criteria and has not experienced loss but is either still growing or unstable in its cover between 2000 and 2012 (Equation 2). A young secondary forest is defined as an area which is not a plantation and has experienced loss or does not meet 2010 cover criteria, but which is not disturbed, and is either tropical and under 10 years old, or temperate (Equation 3). Cropland is defined as an area which is not a plantation and does not meet 2010 cover criteria or has experienced loss, which is also sufficiently disturbed, was not disturbed by natural fires, and is dominated by cropland (Equation 4). Pasture is defined as an area which is not a plantation and does not meet 2010 cover criteria or has experienced loss, which is also sufficiently disturbed, was not disturbed by natural fires, and is dominated by pasture (Equation 5). Urban land is defined as an area which is not a plantation and does not meet 2010 cover criteria or has experienced loss, which is also sufficiently disturbed, was not disturbed by natural fires, and is dominated by urban areas (Equation 6). Finally, plantations are defined as areas which are covered by the plantations layer, independent of all other variables (Equation 7). A description of the Boolean conditions used to define land use can be found in SM Table 1.

This resulted in a map within forested biomes with the following land use classes: Primary/mature secondary forest, Intermediate secondary forest, Young secondary forest, Cropland, Pasture, and Urban. The statistical models were crossed with global maps of biomes, land use, and human population density to make global spatial projections of both abundance and compositional similarity, which were then multiplied together to provide a map of modelled biodiversity intactness.

RESULTS

Forest Biodiversity Significance

Our forest biodiversity significance layer shows that the relative importance of forest locations—in terms of their contribution to the distributions of mammal, bird, amphibians and conifer species occurring in them—varies around the world (Figure 1). The areas with low values across most of the temperate region tend to support fewer species and these tend to have larger geographical distributions. While lowland tropical forests in the Amazon and Congo basins are species-rich, these species also tend to have large distributions, so the contribution of any individual location to the overall distributions of these species tends to be low. Conversely, montane forests in South America, Africa and SE Asia all contain many species with small geographical distributions, as do the lowland forests of insular SE Asia, coastal Brazil, Australia, Central American, and Caribbean islands. These regions all show high values for biodiversity significance on our map; as well as being species-rich these individual locations make a greater contribution to the overall distributions of the species occurring within them.

The forest biodiversity significance of tree-cover lost from 2000 to 2018 gives an indication of the impacts of removal of forested habitat (with the caveat that some forest loss may be from natural causes, such as hurricane damage). Our results (Figure 2) highlight those regions where tree cover loss (change from above 60% tree cover to 0% tree cover within a 1 km² pixel) has resulted in disproportionate loss of the distributions of the world's forest-dependent species (in the taxonomic groups considered): Madagascar, parts of eastern Brazil, central America, SE Asia, West Africa, Australia, and northern New Zealand. Intermediate levels of loss are seen across large regions of the forests of continental and insular SE Asia. Although areas of particularly significant forest biodiversity loss occurred in the tropics during this period (a reflection of the higher species richness as well as the density of endemic species within the tropics), it should be noted that the map also highlights the biodiversity significance of the substantial extent of deforestation that has occurred over the last 18 years within temperate areas. Scandinavia, Russia, Canada and the USA have all undergone considerable losses in forest cover, mainly due to large-scale logging and fires (Curtis et al., 2018). Although these areas support relatively fewer species and these species tend to have larger global distributions, the aggregate biodiversity impacts may be substantial. The layer does not distinguish between forest
Forest biodiversity significance

FIGURE 1 | Forest biodiversity significance in 2018, in terms of the contribution of each location to the distributions of forest mammal, bird, amphibian, and conifer species occurring in them. Grey shows low and dark purple shows high significance values. White shows areas not classified as forest (i.e., tree cover values were <60% in 2000, lost between 2000 and 2018, or mapped as plantations).

Forest biodiversity significance

FIGURE 2 | Forest biodiversity significance for areas of forest loss during 2000–2018, in terms of the contribution of each location to the distributions of forest mammal, bird, amphibian, and conifer species occurring in them. Values are for the year 2000 in areas where forest was subsequently lost. Grey shows low and purple shows high significance. White areas are not classified as forest in 2000 (i.e., tree cover was <25% in 2000, or area was mapped as plantation), or where forest remains in 2018 (i.e., no loss during 2000–2018).
loss that is likely to be permanent in the foreseeable future (e.g., conversion to agriculture) and forest loss that may only be temporary (e.g., resulting from fires or sustainable forestry practices in parts of Canada and Scandinavia). We did not consider forest gain in our assessment, but this is unlikely to overestimate biodiversity loss substantially during the period, as most forest-dependent species do not recolonise young regrowth.

**Forest Biodiversity Intactness**

The forest biodiversity intactness layer reveals the impact that forest change and human population density has had on species assemblages. Our models revealed that land use and human population density are significant predictors of community abundance ($\chi^2 = 38.04, df = 7, p < 0.001$ and $\chi^2 = 7.99, df = 1, p = 0.005$, respectively). Heavily utilised areas of the world are unsurprisingly less intact (Figure 3), for instance, much of Europe and the more densely populated areas of India, North America, Bangladesh, and China. In these areas, the impact of dense human populations together with the urban and agricultural land use required to support them has led to severe losses of biodiversity intactness. Madagascar, coastal Brazil, South Africa, southern Australia, and northern Africa, are also identified as areas with striking losses in biodiversity intactness. These regions have undergone intense removal of natural forest, but retain more of their native biodiversity due to lower levels of urbanisation.

**Comparison of Biodiversity Significance and Intactness**

Overlaying biodiversity intactness and significance provides insight into areas with high values for both, and areas that score highly for one but not the other. Regions with high values for both metrics include the Northern Andes and Central America, south-eastern Brazil, the western, and eastern parts of the Congo basin, southern Japan, the Himalayas, and various parts of Southeast Asia and New Guinea (Figures 4, 5). By contrast, Europe (Figure 5D) is dominated by large areas of biodiversity intactness in the north-east and areas of high biodiversity significance in the south, but lacks large areas where both are high.

**DISCUSSION**

We present two new biodiversity layers for the world's forests, derived from existing data but in novel ways that aim to add contextual meaning to forest data for use in conservation decision-making.

The layers describe two different dimensions of biodiversity and so are not expected to show the same geographic patterns. Biodiversity significance combines spatial variation in both species richness and levels of endemism, and hence shows the relative contribution of any location to the persistence of forest species. However, the degree to which the values in any particular location are driven by one of these characteristics or the other cannot be determined without further analysis. Furthermore, high endemism may result from either natural endemism, human-induced geographic restriction, or a combination. By contrast with biodiversity significance, biodiversity intactness is highest where ecological assemblages remain intact, irrespective of natural macroecological variations in species diversity or endemism; intermediate values can reflect a range of combinations of reduced overall abundance and reduced compositional similarity to an intact assemblage. In general, biodiversity significance is higher within the tropics (especially in topographically heterogeneous regions) and lower in northern, boreal regions, whereas biodiversity intactness is generally low across most of Europe, India and eastern Asia and high within wilderness areas in northern European and North American forests, as well as within tropical forest cores across Central America, South America, Africa, and Asia.

Safeguarding areas of high significance is important as their loss results in a disproportionate loss of species’ distributions, especially narrow-range endemics, elevating species’ risk of extinction. High intactness is important to safeguard in order to (a) maintain ecosystem functioning; (b) retain community resilience against pressures such as climate change; and, in the case of forest ecosystems, (c) help mitigate climate change through greenhouse gas regulation (Steffen et al., 2015). Biodiversity intactness is also relevant to efforts to define wilderness regions, intact forest landscapes, or areas that have been described as the “last of the wild” (Potapov et al., 2008; Watson et al., 2016, 2018).

At a more local scale, comparison of the layers may provide information relevant for conservation. For example, landscapes of high significance but low intactness may be appropriate targets for restoration efforts. Landscapes that contain both high intactness and high significance reveal locations with relatively high density of geographically restricted native species. Such areas may therefore be important to safeguard through broad-scale policy responses or site-scale conservation measures such as designation of protected areas. However, biodiversity has multiple dimensions and here we have chosen to focus on just two. When considering prioritisation of areas for conservation management other aspects of biodiversity may also be relevant such as phylogenetic diversity or the presence of charismatic species.

The biodiversity significance layer shows similar patterns to those revealed by the distribution of endemic bird areas and biodiversity hotspots that were identified in the 1990s (Myers, 1990; Mittermeier et al., 1998, 2004; Stattersfield et al., 1998). The advantage of our approach is that it is based on many more species than used in these earlier analyses, and the distribution of each species considered is spatially explicit, allowing much finer resolution maps. Furthermore, the analytical approach is repeatable and allows the layer to be updated as more species and taxonomic groups are added.

The biodiversity intactness layer broadly accords with Newbold et al.’s (2016) map, with areas of greatest loss in densely populated and heavily converted regions such as most of western Europe, northern China, and the southern coast of South Africa. However, our estimates for plantation-rich parts of Southeast Asia are notably lower than those of Newbold et al. (2016), which
were criticized by Martin et al. (2019) as being too high. Our forest biodiversity intactness map is also able to distinguish areas of recent (post-2005) forest change, such as lowland Mexico and regions in Southeast Asia.

The outputs we highlight here are relevant to international and national policy including the Convention on Biological Diversity (CBD) (Parties’ National Biodiversity Strategies and Action Plans, and National Reports), the Paris Agreement of
the UN Framework Convention on Climate Change, Bonn Challenge, and global environmental assessment processes such as the Intergovernmental Science-Policy Platform on Biodiversity and Ecosystem Services and the Global Biodiversity Outlook. The layers are potentially useful both for targeting policy responses and on-ground interventions, and for tracking progress towards goals and targets. For example, global, regional and national maps and indicators of the proportion of areas of high forest biodiversity significance or intactness lost over time are relevant to Aichi Target 5 in terms of measures of loss and degradation of habitats, Aichi Target 11 in terms of areas of biodiversity significance, and Aichi Target 12 in terms of preventing extinctions and declines of threatened species. Data on tree-cover loss linked to biodiversity can also be used for national REDD+ planning and monitoring other commitments to international, regional and nation agreements, policies, and laws. The layers are also relevant to the safeguard policies of investors, financial institutions, and companies.

The data that we have brought together here are the best available, but have a number of limitations. For instance, the Hansen et al. (2013) dataset does not allow for regional calibration, yet the height and density of natural tree cover will vary depending upon local variations in environmental conditions (Tropek et al., 2014). We were not able to account for the variation of natural tree cover on a local scale and, as no standard definition of the tree cover associated with natural forests exists at larger scales, our conservative literature-based choice of a 60% tree cover threshold is unlikely to optimally delimit natural forest across the entire area of our analysis. A comparison of the biodiversity significance and intactness maps derived using a 25% tree cover threshold to indicate forest presence (SM Figures 1–3) illustrates this issue. For instance, when considering the biodiversity significance layer, landscapes with high endemicity and a naturally low forest cover are highlighted in the layer at this lower threshold, including the Okavango Delta in Botswana, the South African coast, and the western coast of Madagascar (SM Figure 1). Likewise, the biodiversity intactness layer produced using the 25% threshold reveals intactness with northern boreal forests and dry forests in Zambia, which have a naturally sparse tree cover, that are not highlighted in the 60% threshold layer. However, the 25% threshold intactness layer does not show degraded areas within West Africa, including south-east Ghana, which would naturally be covered with dense tree cover (SM Figure 3). The Hansen et al. (2013) dataset does not provide gain data across all years of our analysis, which meant that it could not be used within the biodiversity significance layer. However, this is unlikely to bias our results substantially given that forests often take considerable time before becoming suitable for forest-specialist species (Newbold et al., 2014).

When producing the biodiversity significance layer, the forest species’ distribution maps were clipped by forest cover and suitable elevations to create maps of the Extent of Suitable Habitat (sensu Beresford et al., 2011), which are finer resolution representations of distribution. However, they do not map occupancy per se, and contain commission errors (e.g., owing
to extirpation caused by over-exploitation, invasive alien species
etc.), although this is not likely to bias the broad patterns across
>12,000 species that are shown in the resulting layer. The layer
does not take account of variation in abundance within species’
distributions, spatial data for which are not available for the
large set of species considered. The data used are not currently
taxonomically representative—for instance, no invertebrates are
included—and clades may have systematically different degrees
of spatial resolution in their distributions resulting in the species
with the most resolved maps obtaining higher values. The coding
of forest dependence in the IUCN Red List does not capture finer-
scale variation. In the coming years, the layer will be updated to
incorporate data on the distributions of all forest reptiles and
a number of forest plant groups (e.g., trees, gingers, rattans) and
invertebrate taxa (e.g., dragonfly, monarch and swallowtail
butterflies) as these are assessed and mapped for the IUCN
Red List.

Likewise, the biodiversity intactness layer has caveats (though
our analysis overcomes many of the issues raised by Martin
et al., 2019; see also Newbold et al., 2019). For instance, the
layer is based upon data extracted from the (Hansen et al., 2013
(updated) tree cover change dataset which only dates back to
2000. Therefore, we are not able to distinguish between forest
that had recovered by 2000 and pristine forest. The biodiversity
intactness layer reflects how species communities are impacted
by land use change and human population density. However,
we know that other anthropogenic pressures—such as climate
change, hunting and exploitation—are also important, but will
only be accommodated in our analysis to the extent that land use
or human population density serve as proxies. Although climate
change has a significant impact on biodiversity, it is not possible
to disaggregate the impacts of a changing climate from the
impacts of land use change and human populations over the short
time period on which our analysis focuses. Roads open forest
areas and affect biodiversity through harvesting (Sodhi et al.,
2004), the introduction of alien species (Hulme, 2009), alterations
in the microclimate and creating light gaps (Laurance et al., 2009)
but these subtle changes were not captured in our analysis.

We have used the plantation data for countries where such
data is available in the SDPT, but it is not possible to distinguish
all plantation forests from natural forests, notably for countries
not represented in the SDPT dataset. Furthermore, it should be
noted that China and Papua New Guinea are present in the
SDPT dataset but we were not able to obtain permission for
their data to be included in this analysis. This deficiency impacts
both approaches. In the forest biodiversity significance layer,
plantations may wrongly appear as highly significant if the forest
they replaced had high values (but not so otherwise). In the
intactness layer, plantations that contained mature trees in 2000
are indistinguishable from primary or mature secondary forests,
but those plantations composed of primarily non-native species
or high intensity, monoculture plantations will have markedly
lower intactness than indicated.

Technological revolutions over the last few years, including
in our ability to obtain and process satellite-derived data with
freely available supercomputer power, are opening up new areas
of opportunity for conservation science. We are moving closer
to near-real-time habitat and biodiversity-change products that
can ingest remotely sensed data and run algorithms to show both
areas of forest loss and the consequences for multiple facets of
biodiversity, within time periods that can lead to rapid responses
and interventions on the ground. Our work represents a further
contribution to this aim.

The layers described here have been integrated into the Global
Forest Watch platform (www.globalforestwatch.org), which aims
to provide the data necessary to document and conserve forests
worldwide. It provides information relevant to monitoring fires,
documenting illegal activities, screening estates for deforestation
and analysing trends in forest change.

Humanity has long relied upon forests, and the varied and
complex species assemblages they encompass and support, but in
recent times human impacts have become unsustainable, creating
areas depauperate in biodiversity. The layers presented here help
to evaluate and map how we have impacted forest biodiversity
and can inform what measures can be taken at a local scale to
conserve and restore forests.

DATA AVAILABILITY STATEMENT

The datasets analyzed for this study can be found in the
NHM data portal (data.nhm.ac.uk), and through request
at www.iucnredlist.org/resources/spatial-data-download,
and www.datazone.birdlife.org/species/requestdis, and the
resulting layers can be visualized on the Global Forest Watch
portal (data.globalforestwatch.org).

AUTHOR CONTRIBUTIONS

SH and AA led all analyses. CM carried out analyses. SH, AA,
CM, SB, CH-T, CC, CD, AP, and NB devised the methodology.
All co-authors assisted in the production of the manuscript.

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November 2019 | Volume 2 | Article 70
9
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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/ffgc.2019.00070/full#supplementary-material
Hill et al. | Forest Biodiversity Status and Changes


Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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