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## FIELD NOTE

**Biomass and floristics of a secondary forest in West Kalimantan, Indonesia****Mumammad Mansur<sup>1</sup> and Francis Q. Brearley<sup>2\*</sup>**<sup>1</sup> Research Center for Ecology and Ethnobiology, Indonesian National Research and Innovation Agency (BRIN), Cibinong Science Center, Jalan Raya Jakarta-Bogor Km 46, Cibinong 16911, Indonesia.<sup>2</sup> Department of Natural Sciences, Manchester Metropolitan University, Chester Street, Manchester, M1 5GD, United Kingdom.

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**ABSTRACT** To understand the biomass and floristics of secondary forests in Borneo better, we established a one-hectare plot on the lower slopes of Gunung Kelam in West Kalimantan, Indonesia. We recorded 683 stems ( $\geq 5$  cm dbh) representing 50 species, 44 genera and 27 families; the five species with the greatest Importance Value were *Artocarpus elasticus* (IV = 81.5), *Vitex pinnata* (40.2), *Cryptocarya ferrea* (14.2), *Polyscias elliptica* (12.7) and *Gordonia excelsa* (10.7); stem dbh distributions differed among species indicating that succession was still occurring. The stand basal area was 29.0 m<sup>2</sup>. We estimated biomass with eight different allometric equations. Four equations (Chave, Hashimoto, Kenzo and Manuri-DGH9) showed very close agreement at around 137 Mg ha<sup>-1</sup> suggesting they were all suitable for mid-aged secondary forest biomass estimation in this region. Despite tree diversity and biomass being lower than nearby primary forest, secondary forests will become increasingly prevalent in the future and this therefore necessitates their increased study and conservation.

**Key words:** biomass, diversity, Kalimantan, regeneration, secondary forest**INTRODUCTION**

The dipterocarp-dominated forests of Borneo are among the world's most species-rich (Kier et al., 2005; Ashton, 2014) and productive (Banin et al., 2014; Taylor et al., 2019). Studies in the forests of Kalimantan (Indonesian Borneo) have shown a rich tree flora (Suzuki et al., 1997; Kohyama et al., 2003; Brearley et al., 2004; Cannon & Leighton, 2004; Wilkie et al., 2004; Kartawinata et al., 2008; Sheil et al., 2010) yet these forests are under threat from multiple disturbances (Gaveau et al., 2018; Alamgir et al., 2019). An increasing proportion of forests are now secondary in nature (Chazdon, 2014) so understanding the diversity, structure and functioning of tropical secondary forests is becoming more important as secondary forests will play a greater role in biodiversity conservation and ecosystem service provision in the future (Chazdon et al., 2009). Furthermore, given their increasing prevalence and rapid growth rates, secondary forests have the potential to act as effective carbon sinks, but the magnitude and time frame of this is not well understood (Chazdon et al., 2016). A number of studies have examined biomass accumulation (Ewel et al., 1983; Chai, 1997; Hashimoto et al., 2000; Ohtsuka, 2001; Lawrence, 2005; Jepsen, 2006; Kenzo et al., 2010; Tanaka et al., 2021) and/or species diversity and composition (Ewel et al., 1983; Prajadinata, 1996; Lawrence et

al., 2005; Wasli et al., 2011; Labrière et al., 2015; Karyati et al., 2018; Ulfah & Sulistyawati, 2018; Tanaka et al., 2021) in secondary forests of Borneo but they have largely focused on younger secondary forests (<20 years) with only a few studies on older forests (Okimori & Matius, 2000; Brearley et al., 2004; Wasli et al., 2011).

When determining forest biomass, allometric equations are required as it is rarely possible, nor desirable, to directly measure and enumerate all the trees present through direct harvesting. Secondary forest trees have faster growth rates and thus lighter wood and a contrasting architecture to primary forest trees. Equations developed for primary forests may, therefore, not be appropriate for secondary forest biomass estimation, so comparing allometric equations derived from primary and secondary forests, as well as those based on multiple sites (pantropical equations) is important for accurate biomass estimation in these forests (van Breugel et al., 2011).

Gunung (= Mount) Kelam in West Kalimantan, Indonesian Borneo, is a spectacular granite dome, one of the largest monolithic mountains in the world, best known botanically for *Nepenthes clipeata* (Mansur et al., 2021) although we have recorded at least 300 additional species in a preliminary study (Mansur et al., in revision). Gunung Kelam is protected as a 'Taman Wisata Alam' (Nature Tourism Park) and has forests covering its lower slopes –

these forests were disturbed by fires brought about by El Niños in the 1980 s and 1990 s and are all now secondary. Our aims in this paper are to: i) describe the diversity and biomass of a plot within the secondary forest on the mountain, and ii) compare equations for estimating the secondary forest biomass.

## MATERIALS AND METHODS

In January 2018 we established a one-hectare (100 m × 100 m) sampling plot in the secondary forest at Gunung Kelam at 0°04'30.5" North and 111°37'59.2" East. The plot was based on a granite geology having steep topography and was laid out at *c.* 50 to 100 m altitude (Fig. 1). The plot was divided into 100 quadrats of 100 m<sup>2</sup> (10 m × 10 m) and within each quadrat all trees with dbh (diameter at breast height; 1.3 m) ≥ 5 cm were identified, their dbh was



Fig. 1. Internal view of secondary forest on the lower slopes of Gunung Kelam, Indonesian Borneo.

measured using a measuring tape, and tree heights were measured with a Senshin AT-15 measuring pole or estimated by comparing with the heights of selected trees that were used as a reference. Any trees with buttresses were measured 30 cm above the highest point; lianas were not measured. We collected herbarium specimens from each species within the plot and identified them at Herbarium Bogoriense (BO), Research Center for Ecology and Ethnobiology, National Research and Innovation Agency (BRIN), where they were deposited; only one single tree remained unidentified. We calculated species diversity metrics (Chao1, Fisher's  $\alpha$ , Shannon-Wiener) using the equations in Colwell (2009) and the importance value index (IVI), which is the sum of relative density, relative frequency and relative dominance, according to Mueller-Dombois & Ellenberg (1974). To estimate the above-ground biomass of the forest, we compared four equations derived from secondary forests in South-east Asia: Hashimoto et al. (2004), Kenzo et al. (2009), Karyati et al. (2019) (we used their equation derived from 20-year-old secondary forest) and Ketterings et al. (2001) as well as two derived from South-east Asian primary forests: Yamakura et al. (1986) and Manuri et al. (2016) (we used their DGH9 equation). In addition, we also used the pan-tropical allometric equations of Brown (1997) for moist forests and Chave et al. (2014). Each equation uses a combination, or a subset, of tree diameter, height and wood density to calculate the biomass of each tree (Table 1) which was then summed to calculate total tree biomass in the one-hectare plot; we used the Global Wood Density Database of Zanne et al. (2009) when wood density was required (72 % at species level, 22 % at genus level, 4 % at family level and 2 % as the plot mean). Equations are henceforward generally referred to by their first author name only.

Table 1. Eight equations used to estimate tropical tree above-ground biomass (AGB) from tree diameter at breast height (dbh), height (H) and wood density (WD) applied to trees in a secondary forest at Gunung Kelam, Indonesian Borneo.

Equation name	Equation	No. trees	Location	Reference
Brown-moist	$AGB = \exp(-2.134 + (2.53 \times \ln(\text{dbh})))$	170	Pantropical (5 study sites)	Brown (1997)
Chave	$AGB = 0.0673 \times (\text{WD} \times \text{dbh}^2 \times \text{H})^{0.976}$	4004	Pantropical (58 study sites)	Chave et al. (2014)
Hashimoto	$AGB = \exp(2.44 \times \ln(\text{dbh}) - 2.51)$	191	Secondary forest (Kalimantan)	Hashimoto et al. (2004)
Karyati-20	$AGB = \exp(2.3207 \times \ln(\text{dbh}) - 1.89)$	30	Secondary forest (Kalimantan)	Karyati et al. (2019)
Kenzo	$AGB = 0.0829 \times \text{dbh}^{2.43}$	136	Secondary forest (Sarawak)	Kenzo et al. (2009)
Ketterings	$AGB = 0.11 \times \text{WD} \times \text{dbh}^{2.62}$	29	Secondary forest (Sumatra)	Ketterings et al. (2001)
Manuri-DGH9	$AGB = 0.071 \times (\text{WD} \times \text{dbh}^2 \times \text{H})^{0.973}$	108	Primary forest (Kalimantan)	Manuri et al. (2016)
Yamakura	$AGB = (0.02903 \times (\text{dbh}^2 \times \text{H})^{0.9813}) + (0.1192 \times (0.02903 (\text{dbh}^2 \times \text{H})^{0.9813})^{1.059}) + (0.09146 \times ((0.02903 \times (\text{dbh}^2 \times \text{H})^{0.9813}) + (0.1192 \times (0.02903 (\text{dbh}^2 \times \text{H})^{0.9813})^{1.059})^{0.7266}))$	191	Primary forest (Kalimantan)	Yamakura et al. (1986)

## RESULTS

### Vegetation Composition

Within the one-hectare plot there were 683 stems of 50 species, 44 genera and 27 families (575 stems, 43 species, 37 genera and 27 families for stems  $\geq 10$  cm dbh). The five dominant tree species (dbh  $\geq 5$  cm) with the greatest Importance Values (IV) were *Artocarpus elasticus* (IV = 81.5), *Vitex pinnata* (IV = 40.2), *Cryptocarya ferrea* (IV = 14.2), *Polyscias elliptica* (IV = 12.7) and *Gordonia excelsa* (IV = 10.7) (Table 2). The abundances of all species found are listed in Table 2 along with their families and authorities. In terms of species richness, Fisher's  $\alpha$  was 12.4 and the Shannon-Wiener diversity index was 2.90. The Chao1 estimate of species richness was 55.1 indicating that we found around 90 % of the species predicted to be present. There was only one species of dipterocarp in the plot (*Anisoptera grossivenia*) although two other species were found outside the plot (*Shorea balangeran* and *S. parvifolia*; Mansur et al., in revision). Two species were endemic to Kalimantan (*Anisoptera grossivenia* and *Semecarpus glauca*) (Sidiyasa, 2015) and four were on the IUCN Red List: *Azelia rhomboidea* (VU), *Artocarpus anisophyllus* (VU), *Dimocarpus longan* subsp. *malesianus* (NT) and *Palaquium hexandrum* (NT).

### Vegetation Structure

The trees in the plot had a total basal area of 29.0 m<sup>2</sup> (28.5 m<sup>2</sup> for trees  $\geq 10$  cm dbh); there was a general decline in the number of stems with increasing size classes (Fig. 2) although trees in the smallest size class for a given species were often less numerous than the next larger class. Three species with the largest dbhs were *Gordonia excelsa* (82.9 cm), *Dacryodes rubiginosa* (69.7 cm) and *Palaquium hexandrum* (67.1 cm). Among the ten most abundant species, some tree species had a fairly even distribution among dbh size classes. Many species had few stems in the smallest size class indicative of poor regeneration: *Artocarpus elasticus* (Fig. 2a), *Vitex pinnata* (Fig. 2b), *Cryptocarya ferrea* (Fig. 2c), *Polyscias elliptica* (Fig. 2d) and *Cleistanthus oblongifolius* (Fig. 2g), whilst some other species were mostly of small dbh: *Antidesma tetrandum* (Fig. 2e) and *Leucosyke capitellata* (Fig. 2f), and others showed a fairly even distribution across size classes: *Neo-uvaria acuminatissima* (Fig. 2h) and *Azelia rhombodea* (Fig. 2i). Only 2 % of the trees were more than 20 m tall with the tallest tree (*Dacryodes rubiginosa*) reaching 26 m in height.

### Forest Biomass

The eight equations to estimate the forest biomass gave contrasting results (Fig. 3). The Brown-moist equation estimated the greatest biomass (228 Mg ha<sup>-1</sup>) and the Karyati-20 and Ketterings equations, both developed from trees in South-east Asian secondary forests, estimated forest biomass as 168 and 161 Mg ha<sup>-1</sup>, respectively, with the South-east Asian primary forest equation of Yamakura estimating 159 Mg ha<sup>-1</sup>. The other equations were either pan-tropical (Chave) or developed from other primary (Manuri-DGH9) or secondary forests in Borneo (Hashimoto, Kenzo) and all four gave remarkably similar estimates of between 135 and 141 Mg ha<sup>-1</sup> (Fig. 3).

## DISCUSSION

As anticipated, the diversity and biomass of trees in this area of secondary forest in West Kalimantan (Indonesian Borneo) was lower than other similar primary forests (Slik et al., 2010), in fact, both the biomass and number of species were about one-third of those found at Serimbu which is the nearest extensively studied primary forest site (Kohyama et al., 2003). The species composition of the Gunung Kelam secondary forest plot shares a number of commonalities in species composition with other secondary forests in Borneo and the broader region. It contained a similar set of key 'important' species to secondary forests of Sarawak where *Artocarpus elasticus* and *Vitex pubescens* were dominant species along with *Macaranga gigantea* (Nakagawa et al., 2013), and in East Kalimantan where a secondary forest plot was dominated by *Vitex pinnata* and *Geunsia furfuracea* (syn. *Callicarpa pentandra*) (Sukardjo, 1990). The low number of *Macaranga* trees (2 % of the stems), in contrast to other secondary forests in the region (Riswan & Abdulhadi, 1992; Prajadinata, 1996; Lawrence et al., 2005; Nakagawa et al., 2013; Tanaka et al., 2021), is possibly due to the low nutrient status of the granitic substrate (Chua et al., 2013) or the older age of the forest. The high diversity of edible fruit trees (e.g. *Artocarpus anisophyllus*, *A. elasticus*, *Durio zibenthinus*, *Dimocarpus longan*, *Nephelium lappaceum*) is unusual for secondary forests and this might be due to the closeness of a campground where human vectors may be responsible for discarding fruit tree seeds into the secondary forest. This could be leading to an inadvertent 'forest garden' being produced (Salafsky, 1994; Suzuki et al., 1997; Mansur, 2007) in the future. The lack of dipterocarps, that dominate primary forests of Borneo (Brearley et al., 2016), indicates

Table 1. Stem number, basal area and importance value index (IVI) of all tree species in a secondary forest plot (1-ha) at Gunung Kelam, Indonesian Borneo.

Latin name	Family	Local name	Number of stems	Basal area (m <sup>2</sup> ha <sup>-1</sup> )	IVI
<i>Azelia rhomboidea</i> (Blanco) S.Vidal	Leguminosae	Kayu gang	16	0.728	7.336
<i>Agrostistachys borneensis</i> Becc.	Euphorbiaceae	Lengkong	2	0.021	0.817
<i>Alstonia scholaris</i> (L.) R.Br.	Apocynaceae	Kayu pelai	10	0.717	5.970
<i>Anisoptera grossivenia</i> V.Sloot.	Dipterocarpaceae	Resak	3	0.197	1.571
<i>Antidesma tetrandrum</i> Blume	Phyllanthaceae	Remayan	32	0.298	10.45
<i>Archidendron fagifolium</i> (Miq.) I.C.Nielsen	Leguminosae	Jereng bukit	2	0.326	1.642
<i>Ardisia laevigata</i> Blume	Primulaceae	Sabang bubu	1	0.004	0.384
<i>Artocarpus anisophyllus</i> Miq.	Moraceae	Entawa	2	0.066	0.746
<i>Artocarpus dadah</i> Miq.	Moraceae	Dadak	7	0.592	4.422
<i>Artocarpus elasticus</i> Reinw. ex Blume	Moraceae	Tekalong	184	10.85	81.51
<i>Artocarpus gomezianus</i> Wall. ex Trécul	Moraceae	Cempedak	3	0.083	1.402
<i>Beilschmiedia madang</i> Blume	Lauraceae	Medang daun besar	7	0.096	2.484
<i>Chisocheton patens</i> Blume	Meliaceae	(Unknown)	1	0.012	0.412
<i>Cleistanthus oblongifolius</i> (Roxb.) Müll.Arg.	Phyllanthaceae	Papinang	23	0.904	10.32
<i>Clerodendrum villosum</i> Blume	Lamiaceae	Empait	1	0.003	0.383
<i>Cratoxylum cochinchinense</i> (Lour.) Blume	Hypericaceae	Mandieng	9	0.171	3.487
<i>Cryptocarya ferrea</i> Blume	Lauraceae	Medang	41	0.867	14.19
<i>Dacryodes rubiginosa</i> (A.W.Benn) H.J.Lam	Burseraceae	Kayu bunga	9	1.061	6.560
<i>Dillenia eximia</i> Miq.	Dilleniaceae	Simpur	6	0.035	2.128
<i>Dimocarpus longan</i> subsp. <i>malesianus</i> Leenh.	Sapindaceae	Lengkeng	2	0.149	1.257
<i>Diospyros borneensis</i> Hiern.	Ebenaceae	Kesepa/Merkuyung	2	0.101	1.093
<i>Durio zibethinus</i> L.	Malvaceae	Durian	10	0.214	4.459
<i>Endospermum diadenum</i> (Miq.) Airy Shaw	Moraceae	Rendong	5	0.271	2.795
<i>Falcataria moluccana</i> (Miq.) Barneby & J.W.Grimes	Leguminosae	Sepium	7	0.054	2.115
<i>Ficus obscura</i> Blume	Moraceae	Keruruh	7	0.079	2.426
<i>Ficus variegata</i> Blume	Moraceae	Kayu ara	15	0.610	7.688
<i>Garcinia rigida</i> Miq.	Clusiaceae	Kandis	4	0.228	2.276
<i>Gomphia serrata</i> (Gaertn.) Kanis	Ochnaceae	Bengkai	2	0.064	0.739
<i>Gordonia excelsa</i> (Blume) Blume	Theaceae	Kemidan	12	1.858	10.65
<i>Leucosyke capitellata</i> Wedd.	Urticaceae	Melangin	29	0.226	10.22
<i>Macaranga triloba</i> (Thunb.) Müll.Arg.	Euphorbiaceae	Purang	13	0.223	5.157
<i>Magnolia liliifera</i> (L.) Baill.	Magnoliaceae	Jambu batu	4	0.101	1.610
<i>Mallotus mollissimus</i> (Geiseler) Airy Shaw	Euphorbiaceae	Entali	1	0.003	0.382
<i>Mangifera laurina</i> Blume	Anacardiaceae	Mangga	1	0.005	0.389
<i>Nauclea subdita</i> (Korth.) Steud.	Rubiaceae	Ketap	1	0.049	0.541
<i>Neo-uvaria acuminatissima</i> (Miq.) Airy Shaw	Annonaceae	Kumpang	23	0.863	10.18
<i>Nephelium lappaceum</i> L.	Sapindaceae	Rambutan	2	0.010	0.779
<i>Palaquium hexandrum</i> (Griff.) Baill.	Sapotaceae	Tekam bukit	6	0.773	4.448
<i>Polyscias elliptica</i> (Blume) Lowry & G.M.Plunkett	Araliaceae	Pekulai	34	0.735	12.70
<i>Psydrax sumatranus</i> (Miq.) Mahyuni	Rubiaceae	Jelumpit	1	0.016	0.427
<i>Pyrenaria serrata</i> Blume	Theaceae	Mamere	12	0.398	4.710
<i>Santiria griffithii</i> Engl.	Burseraceae	(Unknown)	1	0.003	0.384
<i>Semecarpus glauca</i> Engl.	Anacardiaceae	Ketap hitam	2	0.030	1.074
<i>Sterculia cordata</i> Blume	Malvaceae	Gaus	5	0.240	2.690
<i>Syzygium nervosum</i> A.Cunn. ex DC	Myrtaceae	Bungkang	6	0.486	3.911
<i>Syzygium claviflorum</i> (Roxb.) Wall. ex A.M.Cowan & Cowan	Myrtaceae	Ubah	6	0.255	2.663
<i>Turpinia sphaerocarpa</i> Hassk	Staphyleaceae	Entili	6	0.644	4.453
<i>Urophyllum arboreum</i> (Reinw. ex Blume) Korth.	Rubiaceae	Tulang ular	2	0.007	0.769
<i>Vitex pinnata</i> L.	Lamiaceae	Leban	102	3.209	40.22
(Unidentified)	(Unidentified)	Maretebung	1	0.057	0.568

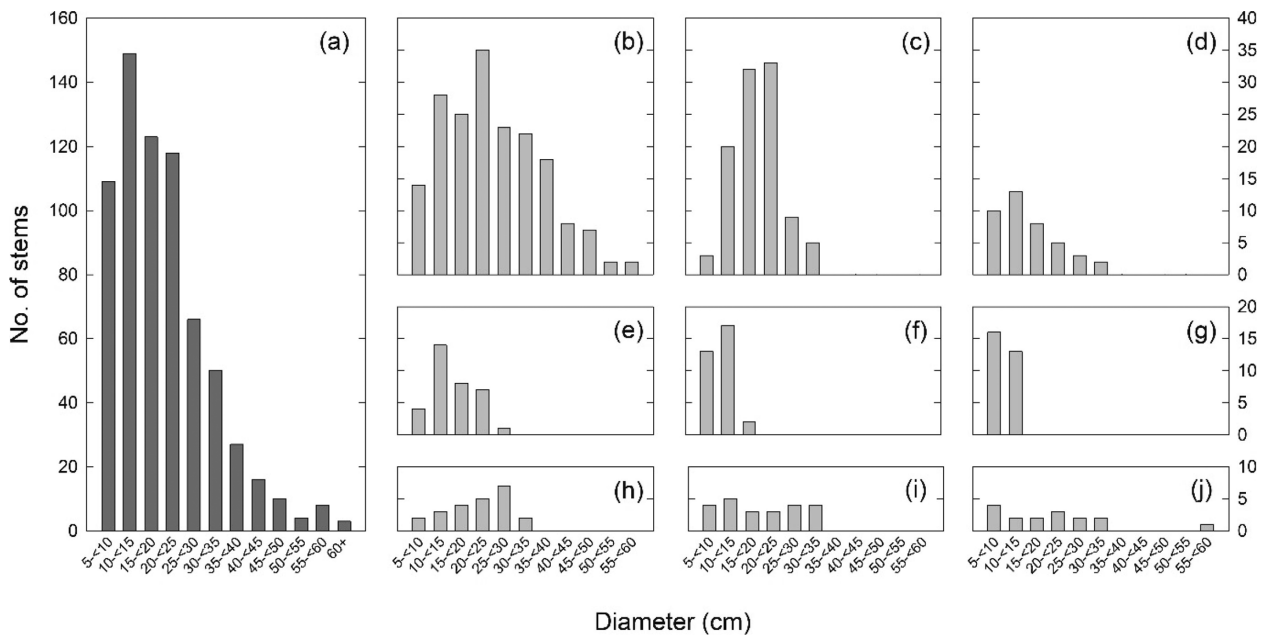


Fig. 2. Diameter at breast (dbh) distributions of (a) all the stems and (b–j) the nine most abundant species in a secondary forest at Gunung Kelam, Indonesian Borneo. (b) *Artocarpus elasticus*, (c) *Vitex pinnata*, (d) *Cryptocarya ferrea*, (e) *Polyscias elliptica*, (f) *Antidesma tetrandum*, (g) *Leucosyke capitellata*, (h) *Cleistanthus oblongifolius*, (i) *Neo-uvaria acuminatissima*, and (j) *Afzelia rhomboidea*. Note the different vertical axis scale between panel (a) showing all stems and panels (b) to (j) showing single species.

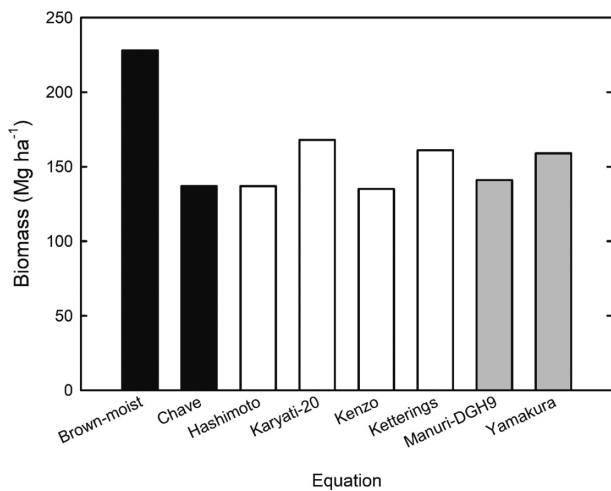


Fig. 3. Above-ground biomass estimation of a secondary forest at Gunung Kelam, Indonesian Borneo, using eight different allometric equations that were either pan-tropical (black bars), or derived from South-east Asian secondary (white bars) or primary (grey bars) forests.

that the plot is still at a moderately early stage of succession. Additionally, dispersal limitation through the matrix habitat that has a high proportion of oil palm plantations will be difficult for these heavy-seeded and wind-dispersed species. Based on comparisons with other forests in the region, we estimate that this forest is 35 to 40 years old and

suggest that it is possibly recovering from burning in the 1982/83 El Niño (Wooster et al., 2012) although we cannot be certain on this. Whilst parts of the forest on Gunung Kelam also burnt during the 1997/98 El Niño, the location of the plot did not (Kayoman, 2005). Although fires have been set by humans for many thousands of years in Borneo, they are becoming increasingly frequent, severe and problematic leading to loss of biodiversity of a range of organisms (e.g. van Nieuwstadt & Sheil, 2005). There were fewer stems in the 5–<10 cm dbh class than the next class above in size for a number of the abundant species. This suggests that current regeneration is poor, however, some of the species in the smaller size class are those that may take over as canopy dominants as succession proceeds, although others will remain as smaller understorey species. The current canopy is likely dominated by long-lived pioneers that are preventing establishment of shade-tolerant species. Additionally, the dense shrubby understorey may inhibit regeneration.

Above-ground biomass was lower than primary forests of Borneo at around only one-third of that of comparable forests (Slik et al., 2010). Poorter et al. (2016) found that mean biomass after 20 years of succession in South American forests was 122 Mg ha<sup>-1</sup> but there was an order of magnitude variation that was related, in part, to precipitation with wetter sites accumulating biomass more rapidly.

We used biomass equations specifically developed for secondary forests and it was interesting to note that two of the allometric equations that were derived from secondary forests in Borneo (Kenzo and Hashimoto) led to very similar biomass estimates but were also remarkably similar to the widely-used pan-tropical equation of Chave et al. (2014). While the Kenzo and Hashimoto equations are based on tree diameter alone, the Chave equation also includes tree height as well as wood density and therefore we would expect it to be better constrained with the inclusion of these parameters. The equation of Karyati-20, derived from a 20-year-old secondary forest in Kalimantan, was greater than the other secondary forest estimates, possibly because the mean wood density of trees ( $0.45 \text{ g cm}^{-2}$ ) included was greater than typical for secondary forests (e.g.  $0.35 \text{ g cm}^{-2}$  from Kenzo et al. (2009)). The equation of Ketterings produced greater biomass estimates than those noted already – although trees were harvested from a secondary forest, it appears to have been managed by local populations and the species composition was different. Among the equations including trees obtained from primary forest, those of Brown-moist and Yamakura provided the greatest biomass estimates, suggesting that location or forest-type specific equations would be more appropriate rather than those based on primary forest species with a large number of dipterocarps in the case of Yamakura. However, in contrast to that of Yamakura, the Manuri-DGH9 equation was very similar to that of Chave. Whilst derived from primary forest trees, it also included height and wood density indicating that some equations derived from primary forest trees could be appropriate for mid-aged secondary forests if well constrained. Further statistical comparison of these biomass estimates is not possible because we would need the raw data that was used to construct the allometric equations for error propagation and this data was not available in many cases.

In conclusion, we show here that this secondary forest plot at Gunung Kelam has a lower diversity and biomass than Bornean primary forests. In comparison with other secondary forests, successional dynamics may be influenced by the fragmented nature of the surrounding habitat and the infertile granitic substrate. The concurrence of four allometric equations for biomass estimation suggests their relevance for mid-aged secondary forests in this region.

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## CONFLICTS OF INTEREST

The authors declare that they have no competing interests.

## AVAILABILITY OF DATA

Data will be uploaded to ForestPlots.net

## CODE AVAILABILITY

Not applicable

## AUTHORS' CONTRIBUTIONS

MM and FQB have equal contributions to this work as the main contributors. MM designed the project and collected the data. MM and FQB performed the analyses and wrote the manuscript.

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