An ecophysiologically informed model of seed dispersal by orangutans: linking animal movement with gut passage across time and space

Esther Tarszisz1,2,*, Sean Tomlinson3,4, Mark E. Harrison2,5, Helen C. Morrogh-Bernard2,6 and Adam J. Munn1,7

1School of Biological Sciences, University of Wollongong, Wollongong, NSW 2522, Australia
2Borneo Nature Foundation, Jl. Bukit Raya 82, Palangka Raya 73112, Central Kalimantan, Indonesia
3School of Molecular & Life Sciences, Curtin University of Technology, Kent Street Bentley, WA 6102, Australia
4School of Geography, Geology and the Environment, University of Leicester, University Road, Leicester LE1 7RH, UK
5Centre for Ecology & Conservation, College of Life and Environmental Sciences, University of Exeter, Cornwall Campus, Penryn, Cornwall TR10 9EZ, UK
6School of Biological, Earth and Environmental Sciences, The University of New South Wales, NSW 2052, Australia
7*Corresponding author: School of Biological Sciences, University of Wollongong, Wollongong, NSW 2522, Australia. Tel: +61411079741. Email: ebt707@uow.edu.au; etarszisz@hotmail.com

Fauna-mediated ecosystem service provision (e.g. seed dispersal) can be difficult to quantify and predict because it is underpinned by the shifting niches of multiple interacting organisms. Such interactions are especially complex in tropical ecosystems, including endangered peat forests of Central Borneo, a biodiversity hot spot and home to the critically endangered orangutan (Pongo pygmaeus wurmbii). We combined studies of the digestive physiology of captive orangutans in Australia with detailed field studies of wild orangutans in the Natural Laboratory of Peat-Swamp Forest of Sabangau, Central Kalimantan, Indonesia. By measuring the gut transit time (TT) of indigestible seed mimics (beads) in captivity and applying this as a temporal constraint to movement data of wild orangutans, we developed a mechanistic, time-explicit spatial model to project the seed dispersal patterns by these large-bodied, arboreal frugivores. We followed seven orangutans and established home range kernels using Time Local Convex Hull (T-LoCoH) modelling. This allowed us to model individual orangutan movements and to adjust these models according to gut transit times to estimate seed dispersal kernels. Female movements were conservative (core ranges of 55 and 52 ha in the wet and dry seasons, respectively) and revisitation rates to the same location of \( n = 4 \) in each 24-h block. Male movements were more unpredictable, yielding fragmented core ranges and revisitation rates to the same location of only 1.2 times each 24 h; males also demonstrated large disjunctions where they moved rapidly over long distances and were frequently lost from view. Seed dispersal kernels were nested predictably within the core ranges of females, but not males. We used the T-LoCoH approach to analyse movement ecology, which offered a powerful tool to predict the primary deposition of seeds by orangutans, thereby providing a reliable method for making \textit{a priori} predictions of seed dispersal dynamics by other frugivores in novel ecosystems.

Key words: ecological service provision, endozoochory, home range estimates, kernel modelling, orangutan, plant–animal interactions, T-LoCoH

Editor: Steven Cooke

Received 29 November 2017; Revised 14 February 2018; Editorial Decision 16 February 2018; accepted 23 February 2018

Cite as: Tarszisz E, Tomlinson S, Harrison ME, Morrogh-Bernard HC, Munn AJ (2018) An ecophysiologically informed model of seed dispersal by orangutans: linking animal movement with gut passage across time and space. Conserv Physiol 6(1): coy013; doi:10.1093/conphys/coy013.
Introduction

The ecosystem services provided by animal-plant interactions are complicated and are governed by numerous co-evolved ecological processes (Tylianakis et al., 2010). Consequently, many of these services and their persistence may be sensitive to disruption (McCauley et al., 2012). In many ways, these associations are contingent upon overlaps in the biotic elements of each species’ realized niche (Soberón and Nakamura, 2009), and small changes in the niche and/or behaviour of either the plant or the animal participant can have substantial influences on the other. Consequently, fauna-mediated ecosystem service provision can be highly context-specific, particularly in especially biodiverse systems where there are numerous biotic and abiotic interactions. One such region includes the tropical peat forests of southeastern Asia, which are notably biodiverse and also represent major carbon sinks (Posa et al., 2011). These peat swamp forests support some of the last remaining populations of one of the world’s largest arboreal frugivores, the orangutan. Recent research has strongly highlighted the role of orangutans as seed-dispersing agents in these peat swamps, and they may be especially important for the dispersal of large-seeded tree species, typical of other tropical forests (Peres and van Roosmalen, 2002; Wotton and Kelly, 2011; Vidal et al., 2013; Donoso et al., 2017; Tarszisz et al., 2017). Consequently, with ongoing pressures of logging and fragmentation, details concerning orangutans and their seed dispersal capacity are much needed.

Animal-mediated seed dispersal, or zoochory, is a crucial component of plant population dynamics, influencing plants and their communities through both short- and long-distance dispersal (Howe and Miriti, 2000; Nathan and Muller-Landau, 2000; Wang and Smith, 2002; Russo et al., 2006; Cousens et al., 2010; McConkey et al., 2012). Zoochory is an important limiting factor for animal-dispersed seeds in several respects. Zoochory can determine the seed deposition location where plants have a potential to establish (Schupp et al., 2010) and may remove the seeds from competition with the parent plant (Howe and Miriti, 2000; Levin et al., 2003; Muller-Landau, 2007; Nathan et al., 2008b; Ruxton and Schaefer, 2012; Schupp et al., 2010), protect seeds from pathogens and predators (Levin et al., 2003; Nathan et al., 2008b; Schupp et al., 2010; Ruxton and Schaefer, 2012) and has the potential to deposit the seeds in beneficial microsites (Nathan and Muller-Landau, 2000; Schupp et al., 2010; Ruxton and Schaefer, 2012). Additionally, in the case of endozoochory, faecal deposition potentially provides a fertilizer (Traveset and Verdu, 2002; Robertson et al., 2006; Traveset et al., 2007; Fuzessy et al., 2016).

The influence of zoochory, and disruptions to this, has recently been powerful inferred on the basis of population genetic structures of plants, even though their dependence upon zoochory is, in some cases, poorly substantiated (Nathan and Muller-Landau, 2000; Wang and Smith, 2002; He et al., 2009; Krauss et al., 2009; Hamrick and Trapnell, 2011; Pascov et al., 2015), but see Fuzessy et al. (2017). There have been advances with respect to connecting seed dispersal studies to movement in recent years, for example with the use of GPS trackers on the animal vectors (Kays et al., 2011; Lenz et al., 2011; Abedi-Lartey et al., 2016; Stevenson et al., 2014), or detailed on-the-ground study of animal movements (Culot et al., 2010; Albert et al., 2013), which this study aims to build on, especially with regard to possible discrepancies between the animal movement and their defecation patterns.

Broadly, movement ecology aims to understand the underlying processes and systems that govern the animal movements in their natural habitats, along with predicting the ecological consequences of those movements (Nathan et al., 2008a; Cagnacci et al., 2010; Hebblewhite and Haydon, 2010; Kie et al., 2010; Morales et al., 2010). A key part of such predictions concerns the spatio-temporal pattern over which animals move, broadly considered as an animal’s home range (HR): the area traversed by an animal in its normal activities of feeding, mating and caring for young, i.e. the entirety of its lifetime (Burt, 1943; Anderson, 1982; Quin et al., 1992). Where we could not follow orangutans for their entire lives, we revert to the concept of a utilization area (Bradshaw and Bradshaw, 2002; Bradshaw et al., 2007) or an occurrence distribution (Fleming et al., 2014), which represents the projection of movement patterns of an animal from observations made for less than an individual’s complete adult life, but which still encompasses enough time to be ecologically relevant. In our interpretation, the seasonal comparisons of male and female orangutans meet this definition best, but for reasons of generality, we still refer to this as HR.

Traditionally, animal HRs have been estimated using location point data to construct minimum convex polygons (MCP) or kernel density estimates (KDE) to define core or larger space-use areas (Laver and Kelly, 2008). However, recent work indicates that the typical MCP methods may overestimate animal space-use and may not adequately reflect patterns of space-use, particularly in complex environments (Getz and Salz, 2008; Munn et al., 2013). Consequently, predictions of seed dispersal based on traditional metrics of HR may be inaccurate and may not adequately reflect the action of seed-dispersing agents within their habitat.

More recently, two methods have been proposed that more realistically model animal space-use using point data, particularly for complex environments where movements may be governed by physical features, such as that of the largely arboreal orangutan in peat swamps, that of local convex hulls [LoCoH (Getz and Wilmers, 2004; Getz et al., 2007; Lyons et al., 2013)] and the outlier-restricted edge polygons [OREP (Kenward et al., 2008; Munn et al., 2013)] methods. As these approaches are essentially the same, we shall use the term LoCoH hereafter. The sophistication of the LoCoH spatial statistics provide insight not only for
where animals go but also how they use space (resources) within their range (Börger et al., 2008; Morales et al., 2010; Jachowski and Singh, 2015), as metrics of time-use such as revisitation and duration of stay are able to be established (Lyons et al., 2013). Consequently, the LoCoH method offers notable advantages for describing the seed dispersal potential of zoochorus agents, which is intricately bound with their movement ecology (Nathan and Muller-Landau, 2000).

In addition to more accurately describing an animal’s movement ecology, the LoCoH methods used herein offer advantages for exploring aspects of animal-mediated seed dispersal, specifically concerning the timing of seed ingestion and elimination in faeces from frugivores, which is an inherently time-based metric. In this regard, the physiology of the dispersal agent can have important consequences for the dependent animal–plant interactions, such as seed dispersal or pollination (Abrol, 2005; McCallum et al., 2013; Seltzer et al., 2013; Tomlinson et al., 2014). The capacity to disperse seeds by endozoochory represents an interaction between the animal, its movement patterns and seed movement from ingestion to elimination, i.e. defecation (Cousens et al., 2010; Fuzessy et al., 2017). Despite a reasonable body of research in other locales, such as the Neotropics (Fuzessy et al., 2017), and past research of orangutans in dipterocarp forests (Galdikas, 1982), there is a paucity of information about present-day orangutan populations (Corlett, 2017) and orangutans in peat swamp forest in general [with the exception of a pilot germination study at the site (Nielsen et al., 2011)]. Our aim is to develop a technique to model home ranges of orangutans that was flexible enough to also estimate their provision of seed dispersal. Understanding and prediction of seed dispersal patterns necessitate an ecophysiological informed spatial model and our hypothesis was that the increased timescale over which movements were modelled would result in larger seed shadows than predicted by home ranges, but where these shadows would specifically fall was unpredictable. Our ultimate aim in presenting this model is to be able to use it to predict some of the potential alteration of floral diversity in tropical peat swamp, with loss/change of its largest seed dispersal vector, the orangutan. Furthermore, this model could be used to make predictions about the potential impact of orangutans in logged/degraded areas (Morrogh-Bernard et al., 2014; Corlett, 2017).

Materials and methods

Captive gut-retention studies

Study animals

In total, six orangutans informed these studies: two adult hybrid Sumatran–Bornean orangutans at Taronga Zoo (AEC #4a/11/11), one male (27 years old, 113.5 kg) and one female (29 years old, 66 kg), along with three adult females with infants of varying ages (Female 1: 22 years old, 50.4 kg; Female 2: 24 years old, 40.95 kg; Female 3: 44 years old 42.5 kg) and one adult flanged male (27 years old, 119.6 kg) at Perth Zoo (PZ; AR&E ZA/4991-4 #59404). All animals were fed their regular diet and maintained in their regular enclosures, which consisted of three concrete pens and two separate outdoor areas. Additional banana was added to the regular diet of all the orangutans to hide the seed mimics used (below); additionally, Perth Zoo animals were provided diet cordial (an intermittent dietary ‘treat’). See Supplementary Material for further husbandry details of captive orangutans.

Passage times of seed mimics

On Day 1 of each feed trial, the orangutans were each fed different-coloured spheroid, non-toxic polyethylene seed mimics of 2, 4 and 6 mm diameter, with average ± SD masses (n = 15 beads per size range) of 22.5 ± 4.6, 28.5 ± 1.5 and 103 ± 2.4 mg (OHAUS Adventurer Analytical, AX423). These seed sizes were chosen as they represent the size range of seeds found intact in faeces from 13 of the wild orangutans followed at the field site during comprehensive studies of the fruits eaten, gut passage of seeds and germination success of gut-passed seeds from orangutans at the field site (Nielsen et al., 2011; Tarszisz et al., 2017). Of note, attempts to disguise larger seed mimics of 8 mm and 10 mm in soft food were unsuccessful. The number of seed mimics ingested by the captive orangutans was comparable with that number of similarly sized seeds found eliminated in the faeces of wild orangutans at the study site (Tarszisz et al., 2017 and see also Nielsen et al., 2011).

Throughout the entire experiment, the orangutans were observed during daylight hours between 0530 and 1730 h. Following ingestion of seed mimics, faeces were collected regularly over 10 days. The enclosure design did not allow for camera placement to observe animals overnight; however, faeces could be distinguished by the presence of different-coloured seed mimics during unobserved times. Faecal elimination in the orangutan is noted as occurring mostly in the morning, with reduced production by afternoon and none overnight (Caton et al., 1999). Preliminary observation of faecal production in wild orangutans agrees with this (Tarszisz unpublished data), and the defecations were observed by the primary investigator (E Tarszisz). When they occurred, however, night samples were considered to have occurred at the midpoint of the sampling interval. Coprophagy was not observed. Faeces were frozen immediately after collection before later thawing to extract eliminated beads.

Faeces were washed through mesh sieves of decreasing diameter (down to 1mm) until all faeces had been examined and all seed mimics collected. As an indicator of the potential seed passage time applied to our mechanistic seed dispersal model, we used the transit time of seed mimics as they first appeared in faeces (i.e. time in hours from ingestion to first appearance in faeces).
 Movements of free-ranging orangutans

The field program was conducted within the Natural Laboratory of Peat-Swamp Forest (NLPSF), a 500 km² area contiguous within the wider 9 200 km² of peat swamp forest in the Sabangau ecosystem, Central Kalimantan, Indonesia (Page et al., 1999; Morrogh-Bernard et al., 2003). This area is managed as part of the multidisciplinary research partnership of the Borneo Nature Foundation (BNF) and their Indonesian counterparts, the Centre for the International Cooperation in Sustainable Management of Tropical Peatlands (UPT LLG CIMTROP) at the University of Palangka Raya. The climate is tropical, with high annual rainfall, separated into distinct wet and dry seasons that last from October to May and June to September, respectively. Daily weather observations at our study site recorded precipitation of 67.31 mm per month on average between October 2012 and May 2013, and 53.13 mm per month on average in June 2013 to September 2013. Unlike the region’s lowland dipterocarp forests, peat swamp forests such as Sabangau are non-masting and thus produce fruit relatively consistently throughout the year (Cannon et al., 2007a, 2007b). The relative homogeneity of the TPSF environment (Singleton and van Schaik, 2001; Singleton et al., 2009), as well as limited secondary seed dispersers (such as rodents) and seed predators (rodents and invertebrates) (D’Arcy and Graham, 2008), makes this an ideal location for modelling overall seed dispersal in TPSF by orangutans because there are likely to be very few effects on dispersal of large seeds other than those related to orangutan movement ecology. Previously home range estimates for orangutans at this site were >560 ha for adult (flanged) male and 250–300 ha for adult females using minimum convex polygons (Morrogh-Bernard et al., 2003, Utami Atmoko et al., 2009) and 1900 ha for males using KDE with the least square cross-validation method (Buckley, 2014). However, these modelling approaches do not allow for deeper interrogation of time-space usage or seed dispersal capacity, and so provide limited capacity for in-depth interpretation of orangutan movements and spatial ecology, let alone of seed dispersal. LoCoH methodology

LoCoH uses a non-parametric approach to HR estimation and thereby circumvents assumptions about the distribution form of the point data that is inherent to parametric kernel methods, thereby reducing HR overestimates (Getz and Wilmers, 2004; Getz et al., 2007; Getz and Saltz, 2008; Munn et al., 2013). This enables that LoCoHs produce a set of non-parametric kernels constructed by aggregating local minimum convex polygons and computing a density estimate distribution for all locations based on nearest neighbour linkages (Getz and Wilmers, 2004; Getz et al., 2007; Getz and Saltz, 2008; Lyons et al., 2013, 2015), the union of which estimates HR (Getz and Wilmers, 2004; Getz et al., 2007). Data handling

Focal orangutan follows were conducted following standardized data collection protocols (Martin and Bateson, 1986; Morrogh-Bernard et al., 2002). Activity data (including that of feeding duration and food species) and location data were recorded at 5-min intervals, whereas diet data were collected continuously (Morrogh-Bernard, 2009; Harrison et al., 2010). To ensure that full daily travel patterns were accurately represented, only full-day (nest-to-nest) data were used in this study. Seven individuals, three adult flanged males and four adult females (with juveniles) were followed throughout the study period from October 2012 to December 2013. All point locations were standardized from longitude and latitude into UTM zone 49M coordinates using Earth Point (Clark, 2016) and were transformed into Coordinated Universal Time (UTC) prior to analysis.

Importantly, we have incorporated time into the LoCoH model, described as the T-LoCoH method and incorporates timestamps of each point in both nearest neighbour selection and in the sorting of hulls (Lyons et al., 2013). Of three possible methods, we pursued the a-LoCoH method, which reduces the number of nearest neighbours in areas with thin, scattered points, to better homogenize potential sampling bias. The ‘a’ method adds cumulative distance from the parent point up to an ‘a’ value and determines nearest neighbours whose aggregate distance is ≤ a (Lyons et al., 2013, 2015) and can be superior to other T-LoCoH methods for reducing the minimum spurious hole covering (Getz et al., 2007; Lyons et al., 2013).

Since time is a critical factor contributing to space usage in T-LoCoH, the first step is to determine an appropriate value by which to scale the maximum theoretical velocity, denoted by Lyons et al. (2013) as the dimensionless factor s. To construct the home range kernels of the orangutans, we selected 24-h intervals because orangutans are largely diurnally active, sleeping from dusk to dawn (Mitra Setia et al., 2009). When modelling seed dispersal hulls, s was chosen based on the transit time for seed mimics determined in captive orangutans. As there was no significant difference in gut passage times between seed mimic sizes of 2, 4 and 6 mm (see below), an average passage time of 76 h was applied as the intervisit gap (IVG). To project the largest possible seed dispersal kernels, we also used a maximum passage time of 133h as the IVG. Kernel model refinement

In applying the a-LoCoH approach, the most appropriate value of a was established by examining the differing density of isopleths, overlaid on GIS data to reduce both type I (including areas that are not part of the home range) and type II (overlooking areas that are part of the home range) errors. We checked the validity of the initial value of a by visually assessing whether the ‘a’th isopleth encompassed...
95% of the data, which is often used as definition of the home range (Laver and Kelly, 2008; Lyons et al., 2013). We used the relationships between isopleth area and edge:area ratios and \( a \) to determine the least erroneous values for each individual’s movement patterns following the guidelines suggested by Lyons et al. (2013). As each animal had a different \( V_{\text{max}} \) which was the maximum observed velocity between any two consecutive points, and different movement patterns, the \( a \) value differed between each animal.

**Temporal effects**

We computed revisitation rate (the number of visits to the same GPS location) and duration of use by first specifying an IVG of 24 h. This means that observations were only recognized by the T-LoCoH model as separate visits if at least 24 h had elapsed between them. Secondly, an IVG of 76 h, the average time for a transit of an undigested seed and an IVG of 133 h, the longest transit time for a seed, were specified, creating metrics for revisitation and duration of use over these larger time scales. In effect, for each individual, we modelled two ‘animals’ separately: the orangutan that moved in ‘real time’ and the average seed in their gut passage, which was approximately three times ‘slower’ on average, and over five times ‘slower’ at its slowest. Seed dispersal was therefore explored by interrogating the differences in revisitation, duration of stay and space-use between these first and second ‘animals’.

Spatially explicit projections (Fig. 1) were generated by exporting the probability kernels as shape files and displaying them using the GIS package qGIS v2.4.0-Chugiai. The home range estimates resulting from the T-LoCoH approach were compared against MCP estimates computed using the ‘convex hulls’ command in qGIS that is consistent with previous studies of orangutan home ranges at NLPSF (Morrogh-Bernard et al., 2003; Utami Atmoko et al., 2009; Buckley, 2014).

**Statistical analyses**

In order to test the capacity of the kernel models to predict defecation, known locations of defecation were recorded in the field and not used to train the model. These locations were intersected with the kernel models of defecation/seed dispersal in qGIS. The expected proportions of defecation points falling into each kernel were tested against the observed proportion falling into each kernel using Pearson’s chi-squared test for all animals, and also for males and females only.

We explored the effect of sex and season on orangutan movements and seed dispersal capability by constructing generalized linear models (GLMs) of several modelled elements of orangutan movement, including step length, 80% kernel area, residency (revisitation rate) and duration of stay. Step length refers to the use of the Pythagorean Theorem to calculate the Euclidean step length distances between subsequent GPS fixes.

The core range was initially defined by examining the distribution of hulls in time-use space, choosing a value of \( a \) that filled core areas and minimized spurious crossovers (Lyons et al. 2013) and is defined here as the 20% likelihood kernel (i.e. 80% kernel area). This describes locations that are the most heavily used, which encompass a small proportion of known locations.

Tests were constructed using a fully factorial design of sex and season. All analyses were conducted using R v3.2.2 [R Core (R Core Team, 2015)] in the R studio shell v0.99.48 (R Studio Team, 2015), and all data are reported as means ± 1SEM unless stated otherwise.

**Results**

**Gut passage times**

From our captive feeding trials, the TTs were of 70.6 ± 7.1, 72.5 ± 6.8 and 86.2 ± 16.6 h for the 2, 4, and 6 mm seed mimics, respectively. The maximum TTs were 159.3 ± 14.2, 118 ± 18.4, and 118.0 ± 19.6 h for the 2, 4, and 6 mm seed mimics, respectively. (see Supplementary Material for further details). There were no significant differences in the TTs between any seed mimic (Supplementary Material, \( F = 0.36, \) d.f. = 2; \( P = 0.54 \)), and the common TT\(_{\text{max}}\) averaged 133 h, while the average TT for all seed mimics was 76 h.

**Orangutan movement ecology**

Kernel models showed that, with the exception of two related females, the focal orangutans were semi-solitary, with very little overlap between models of the same sex. Males tended to have much more disjunct movement patterns than females and also tended to overlap several females’ home ranges within their own. Total female orangutan home ranges overlapped by 21.9 ± 11.2 ha, heavily influenced by the large degree of overlap between two related females (approximately 65 ha). Total male orangutan home ranges overlapped by 3.4 ± 1.9 ha. The total home ranges of our focal males overlay the home ranges of all our focal females, averaging 216.9 ± 9.2 ha overlap.

Home ranges were characterized by significantly higher revisitation rates for females (4.01 ± 0.02 visits per day for females compared to 1.24 ± 0.01 visits per day for males, see Table 1) in the core range. Females furthermore had long loops of short duration and low revisitation around the edges of their home ranges. This pattern appeared potentially true for the males, but the data were not extensive enough to state this definitively.

The core range was initially defined by examining the distribution of hulls in time-use space, defined here as the 20% likelihood kernel, describing locations that are the most heavily used. The hullsets that resulted for male orangutans were classified into different movement patterns following the guidelines suggested by Lyons et al. (2013). As each animal had a different \( V_{\text{max}} \) which was the maximum observed velocity between any two consecutive points, and different movement patterns, the \( a \) value differed between each animal.
ha, with an average step length of 8.89 ± 0.11 m, a revisitation rate of 3.43 ± 0.02 visits each day and average visit duration of 41.00 ± 20.18 min. The T-LoCoH home range estimated for females in the dry and wet seasons were 55.31 ± 6.97 ha and 52.38 ± 8.35 ha, respectively. The minimum convex polygons for females in the dry and wet seasons were 149.00 ha and 160.84 ha, respectively.

There were differences in all of the movement parameters of the orangutans between the sexes, while season had no influence. Sex had a significant effect on all indices including the step length ($F_{1,10} = 13.0; P = 0.0047$), revisitation rate ($F_{1,10} = 70.9; P = 7.51 \times 10^{-6}$) and duration of visit ($F_{1,10} = 22.0; P = 8.50 \times 10^{-4}$). For the 80% kernel area, the influence of sex was significant ($F_{1,5} = 16.78; P = 0.009$), although season did not significantly influence the home range area of females ($F_{2,5} = 0.70; P = 0.540$). Revisitation rates were higher and intervisit duration shorter for females than for males in both seasons (Table 1).

Figure 1: Likelihood distribution kernels and revisitation points (dots), as determined by T-LoCoH analysis period = 24 h for females (a–d) and for males (e–g).
### Table 1: Effects of season and sex on the measures of orangutan movement at NLPSF extracted from T-LoCoH kernel modelling.

|                   | Step length (m) | Revisitation rate | Duration of visit |
|-------------------|-----------------|-------------------|-------------------|
|                   | Mean (S.E.)     | Mean (S.E.)       | Mean (S.E.)       |
|                   | $F_{1,10}$      | $P$               | $F_{1,10}$        | $P$               | $F_{1,10}$ | $P$               |
| IVG = 24 h        |                 |                   |                   |
| Season            |                 |                   |                   |
| Dry               | 8.61 (0.15)     | 4.23              | 0.0667            | 3.37 (0.02)       | 0.001     | 0.974             | 43.06 (0.30)     | 0.344 | 0.571          |
| Wet               | 9.12 (0.16)     |                   |                   |
| Sex               |                 |                   |                   |
| M                 | 7.28 (0.26)     | 13.1              | $0.00474$         | 1.24 (0.01)       | 70.9      | $7.51 \times 10^{-6}$ | 61.69 (0.40)     | 22.0 | $8.51 \times 10^{-4}$ |
| F                 | 9.32 (0.12)     |                   |                   |
| Season x sex      |                 |                   |                   |
| DM                | 6.34 (0.39)     | 0.001             | 1.18 (0.01)       | 1.27 (0.01)       | 0.0167    | 0.900             | 65.57 (0.68)     | 0.323 | 0.582          |
| WM                | 7.90 (0.35)     |                   |                   |
| DF                | 8.57 (0.17)     |                   |                   |
| WF                | 9.98 (0.18)     |                   |                   |
| IVG = 76 h        |                 |                   |                   |
| Season            |                 |                   |                   |
| Dry               | –               |                   |                   |
| Wet               | –               |                   |                   |
| Sex               |                 |                   |                   |
| M                 | –               |                   |                   |
| F                 | –               |                   |                   |
| Season x sex      |                 |                   |                   |
| DM                | –               |                   |                   |
| WM                | –               |                   |                   |
| DF                | –               |                   |                   |
| WF                | –               |                   |                   |

Revisitation rate here is the number of visits to the same location per 24 h and the duration of visit gives the average number of minutes spent at each location. Note: Step lengths were not directly calculable for seed dispersal estimates at IVG = 76h. Bold $P$-values are statistically significant at $P < 0.01$.

### Seed dispersal projections

The average 76-hr 80% seed shadow estimated for a female orangutan at NLPSF by T-LoCoH was 52.4 ± 6.44 ha (Fig. 2), with an average revisitation rate of 2.17 ± 0.244 visits every 76 h. The T-LoCoH seed shadow estimated for females in the dry and wet seasons was 57.3 ± 10.46 ha and 47.5 ± 8.24 ha, respectively. The average 133-hr 80% likelihood seed shadow estimated for a female orangutan at NLPSF by T-LoCoH was 94.2 ± 7.49 ha. Due to their disjunct movement patterns, the seed shadows projected for males were much less certain and could not be projected for all individuals beyond 76 h, nor in all seasons. The average 76-hr 80% seed shadow estimated for a male orangutan at NLPSF by T-LoCoH was 17.3 ± 3.93 ha (Fig. 2).

When the seed dispersal kernels projected at the average 76-hr gut passage time were analysed, sex alone influenced the area of the seed shadows ($F_{1,8} = 11.4; P = 0.0097$), revisitation rate ($F_{1,8} = 11.2; P = 0.0102$) and duration of stay ($F_{1,8} = 11.2; P = 0.010$). Season became marginally significant for duration of stay only ($F_{1,8} = 4.56; P = 0.065$). Seeds were likely to be dispersed over a larger area by females, but revisitation was higher, and duration of stay shorter; this was also the case across both seasons (Table 2a,b). Essentially, dispersal by males resulted in smaller, more discrete dispersal ‘islands’ than dispersal by females because their movement patterns were projected over longer, narrower corridors (Fig. 2).

### Model validations

There was no significant difference between the proportion of defecation events observed in each seed dispersal kernel and the likelihood of seed dispersal predicted by the model projections (Pearson’s $r^2 = 0.893; P = 0.151$). The mean average percent error (MA%E) of model predictions was 3.86 ± 0.97 %, ranging from 1.05% to 7.89%. The model fit was stronger for females only ($r^2 = 0.229; P = 0.999$), but marginally less so for males only, although they were not statistically significant ($r^2 = 8.28; P = 0.141$).

### Discussion

To the best of our knowledge, this is the first study to incorporate a time constraint in the construction of kernels, thereby presenting the first ecophysiological informed kernel...
models to predict the spatial consequences of animal–plant interactions via seed dispersal. Importantly, the diet of the wild orangutans observed, from this and earlier studies (Morrogh-Bernard et al., 2009), was comparable with that of the captive animals used to measure the seed mimic gut passage times, in that fruits and some vegetation made up the bulk of the orangutan diet. As such, we are confident that the seed mimic TTs that we measured from captive animals presents realistic temporal constraints defining the likely seed dispersal patterns by the wild animals, at least within the peat swamp habitat reported here. Overall, only sex influenced the measures of orangutan movement, and males tended to move further and more erratically than females. The data that we collected for males, however, were less consistent than for females because we were less able to repeatedly find and follow the males, reducing our confidence in the analysis of their data. Females tended to spend all their time foraging, likely moving mainly in search of fluctuating

Figure 2: Seed shadow kernels projected by T-LoCoH at analysis period = 76 h estimated as the average gut passage time of seeds by orangutans. Generally, most likely seed shadow is more localized than home range for females (a–d), but becomes much less predictable for males (e–g).
Table 2: Effects of sex-season concatenate on revisitation rate for WF = female wet season, DF = female dry season, WM = male wet season, DM = male dry season.

|       | Group 1 | Group 2 | Difference | Adjusted P value |
|-------|---------|---------|------------|-----------------|
| A) IVG = 24h |         |         |            |                 |
| WF—DF | 2.05 ± 0.02 | 3.05 ± 0.02 | 0.993      | <0.01           |
| DM—DF | 1.18 ± 0.01 | 1.18 ± 0.01 | 0.027      | <0.01           |
| WM—DF | 1.27 ± 0.01 | 1.27 ± 0.01 | 0.993      | <0.01           |
| DM—WF | 1.18 ± 0.01 | 1.18 ± 0.01 | 0.027      | <0.01           |
| WM—WF | 1.27 ± 0.01 | 1.27 ± 0.01 | 0.993      | <0.01           |
| WM—DM | 1.27 ± 0.01 | 1.18 ± 0.01 | 0.996      | 0.344           |
| B) IVG = 76h |         |         |            |                 |
| WF—DF | 3.87 ± 0.02 | 3.42 ± 0.02 | 0.444      | <0.01           |
| DM—DF | 1.16 ± 0.01 | 1.23 ± 0.01 | 0.075      | <0.01           |
| WM—DF | 1.23 ± 0.01 | 1.23 ± 0.01 | 0.075      | <0.01           |
| DM—WF | 1.16 ± 0.01 | 1.23 ± 0.01 | 0.075      | <0.01           |
| WM—WF | 1.23 ± 0.01 | 1.23 ± 0.01 | 0.075      | <0.01           |
| WM—DM | 1.23 ± 0.01 | 1.16 ± 0.01 | 0.075      | 0.522           |

Data are presented as mean ± S.E.M. Adjusted P values represent the probability of differences offset against the effects of multiple comparisons, representing the smallest ‘family’ error rate at which the null is rejected. Bold P-values are statistically significant at P < 0.01.

food resources typical of peat swamp forest (Cannon et al., 2007a, 2007b), while male movements may be more motivated by the search for mating opportunities. Kernel models constrained on the basis of gut transit times resulted in a longer time interval, implying a ‘slower’ rate of movement for seeds than for their orangutan dispersal vectors. To the best of our knowledge, the only other study to investigate feed passage rates in orangutans was by Caton et al. (1999). However, Caton et al.’s (1999) study examined the gut passage of small particle (size) and fluid markers; thus, we felt that our study had merit as we are primarily interested in the passage of larger indigestible seed markers in order to extrapolate reliable information on seed passage relevant for broader scale seed dispersal studies. Of note, the seed mimic elimination patterns were more staggered and less smooth than the typical elimination pattern of finer particles and, combined with the much smaller number of seeds typically ingested, the standard measures of particle MRT may not adequately describe seed passage patterns.

Orangutan movement, sexes and seasons

The MCP estimates of home range that we generated for females in each season (150 ha in the dry and 160 ha in the wet) are consistent with previous reports at this study site (Morrogh-Bernard, 2009; Singleton et al., 2009). MCP estimates for males were even larger due to their greater and more erratic movement patterns, consistent with reports by Buckley (2014), who followed orangutans in the same location from 2010 to 2012. Although our MCP home range estimates more closely approximated previous findings, our T-LoCoH estimates are approximately 36% of our MCP projections for females across both seasons. Of note, the kernel areas we have described (Fig. 1) gave an integrated time–space view of orangutan home range use for females, as opposed to previous kernel areas based on space alone (Morrogh-Bernard, 2009; Singleton et al., 2009). As a result, these more precise estimates produced home ranges that were, on average, 10% of the previously published estimates at NLPSF (Morrogh-Bernard, 2009; Singleton et al., 2009; Buckley, 2014). Large discrepancies between LoCoH methods and more traditional methods (MCP, KDE and alpha-hull) have been reported in other studies (Getz and Wilmers, 2004; Getz et al., 2007; Munn et al., 2013). LoCoH approaches tend to produce smaller, more refined estimates than MCP or KDE with fewer type I and II errors (Getz and Wilmers, 2004; Getz et al., 2007; Munn et al., 2013), and our data further confirmed that traditional home range methods such as MCP can substantially overestimate home range and space-use. The incorporation of time aims to take the concept of home range from a static spatial construct, such as the MCP where all known locations are considered equally, towards a more realistic evaluation of space-use, weighting areas where greater time is spent with greater importance.

The use of T-LoCoH generated several informative parameters that described the movement ecology of orangutans at NLPSF: kernel area, revisitation rate, step length and duration of stay. Revisitation rates and duration of stay can
illustrate the importance of different locations between sexes. Our integration of time has shown significant interactions between both how and where space is used between sexes (Table 1). The movement parameters generated by T-LoCoH (step length, revisit rate and duration of stay) for orangutans at the NLPSF were all influenced by sex and all suggested that males ranged over greater areas than females, but were resident for less time, and visited each location less often than females, similar to previous studies (Utami Atmoko et al., 2009). Females had more predictable movement patterns within a more structured core area, and from this, we infer that the females were most likely moving order to meet their ecological energetic requirements.

We also did not detect any seasonal patterns in the movements of male orangutans, and their more unstable core ranges suggested that they had more fluid home ranges that did not fluctuate in accordance with patterns of fruiting at NLPSF. Rather than moving principally or only to forage, males were potentially moving in relation to another powerful imperative—that of mating and/or avoiding (or aggressing) other conspecifics, as Utami Atmoko et al. (2009) have suggested previously.

By modelling the movements of orangutans using T-LoCoH, and specifically incorporating different time and space-use metrics to estimate behaviour patterns, we have both refined the projected home ranges and uncovered possible differences in the motivations of habitat use between males and females. These models are replicable for other individuals and can be readily remodelled as additional data are gathered at the study site (of BNF/CIMPTROP) in ongoing orangutan monitoring projects. Furthermore, due to the malleability of this model, we have been able to extend this to the prediction of downstream ecological patterns resulting from orangutan movement in the form of their likely seed dispersal activity.

**Implications for predicting seed dispersal**

When temporally constrained on the basis of known gut transit of seeds by orangutans, the dispersal kernels created were similar to the 24-h movement kernels of the orangutans themselves, but the ‘seed kernels’ at 76 h are more likely to ‘travel’ through a circuit of the home range and return (or rather, be deposited) in the core utilization area. Essentially, projected defecation points were more closely distributed in space, clustering more closely within the core home range of the focal orangutan, particularly for females, whereas the more predictable movement patterns. Primary endozoochorous seed dispersal can be effectively predicted on the basis of where an animal, in this case an orangutan, will defecate (Wang and Smith, 2002; Cousens et al., 2010). Our model predictions of defecation patterns were well supported by the χ²-test of actual defecation data, with only a small (<10%) error, suggesting that physiologically informed T-LoCoH models should provide accurate estimates of primary seed dispersal.

The movement of seeds can powerfully contribute to tree species’ colonization, succession and post-disturbance recovery, and consequently therefore ecological restoration and management (Wang and Smith, 2002; Bascompte and Jordano, 2007; Schupp et al., 2010; Ruxton and Schaefer, 2012; Côrtes and Uriarte, 2013). Seed dispersal also represents half of the gene flow pattern of plant populations [the other half being pollination (Abrol, 2005; Krauss et al., 2009; Menz et al., 2011; McCallum et al., 2013)], and so is a powerful contributor to population genetic structure. As a critical element of ecological and evolutionary processes, the mechanistic estimation of passive seed dispersal has made considerable strides (Wright et al., 2008, Nathan et al., 2011, 2002). The modelling of plant–animal interactions in a mechanistic manner has, however, remained somewhat elusive, with most zoochory studies applicable only to the time and place of their model training (Cousens et al., 2010; Schupp et al., 2010; Côrtes and Uriarte, 2013). This is largely due to the plethora of stochastic influences on zoochory, such as sex, season, reproductive patterns and ecological energetics (Nathan et al., 2008a), all of which make prediction of animal movements difficult, even in a hypothetically stable ecological system (Guisan and Zimmermann, 2000; Guisan and Thuiller, 2005). In novel ecological ‘hyperspace’ represented by areas of changing land-use and/or climate, the changing patterns of ecological cascades that influence spatial population structure are rendered unpredictable (Dornam et al., 2012; Mesgaran et al., 2014). Thus, while we have developed a unique set of mechanistically informed models of likely seed dispersal patterns for the NLPSF, extrapolating from these into different tropical peat forests or into other orangutan habitats, such as dipterocarp forest, may require further model training. Nonetheless, this study firmly demonstrates how movement and gut transit times of female orangutans influence seed deposition shadows. Furthermore, it suggests that seed dispersal by female orangutans is linked to their foraging activity and that their movement and seed dispersal patterns will change in relation to food availability. This has potentially serious implications for forest structure and genetic isolation if the habitat is disturbed or population levels decrease, particularly for the large-seeded tree species they were found to have endozoochorously transported (Tarszisz et al., 2017).

**Limitations of this study**

Time and logistical constraints made continuous monitoring of the same animals difficult, perturbing the internal consistency of our data. In particular, there is a paucity of data on adult males, compared with adult females, due to their increased space-use requirements (Utami Atmoko et al., 2009, Buckley, 2014), their fast movement on the ground, causing increased ‘loss’ of males during follows compared to females, and their more labile home ranges, based on competition with both flanged and unflanged males. While these home range models are partially indicative of male orangutan movements, they do not give as complete or refined a picture as emerges for the females. It is entirely possible that
with more data for males, we may have found some stability and connectivity of male home ranges. Although Buckley (2014) addressed some of these issues, the consistency of our results with previous research suggests that this problem is a general constraint on the orangutan movement ecology literature (Utami Atmoko et al., 2009). Affixing remote sensing (GPS tags) could have facilitated data collection without any risk of the presence of human observers disturbing the orangutans and altering their movements. Remote sensing would also guarantee consistent survey effort, regardless of the constraints of manpower and inclement conditions (Kie et al., 2010; Tomkiewicz et al., 2010; Lyons et al., 2013; Munn et al., 2013). However, the application of different technology must be considered in the light of other data that would be lost in remote sensing, such as defecation locations and feeding observations, in addition to ethical and logistical issues.

The timescale we followed orangutans was only a relatively short period, when compared with their life history. Longer observations could yield more accurate ranging and space-use data and should produce more accurate models. Furthermore, continual incorporation of faeces location should yield more accurate data for seed dispersal. This would provide the opportunity to build on the data we have collected here.

Future directions

A T-LoCoH approach appears to provide a method to accurately predict (estimate) orangutan movement within TPSF, and we suggest that it is likely that seed dispersal cascades will be similar in other TPSF landscapes, both within and outside of Sabangau. TPSF is an important orangutan habitat that is considerably less studied than the region’s dipterocarp forests, although this has begun to be redressed in recent years (Rieley et al., 1997; Page et al., 1999, 2011; Jauhiainen et al., 2005; Harrison et al., 2010; Hooijer et al., 2010; Morrogh-Bernard et al., 2011, 2014; Beaudrot et al., 2013). Ecological processes may differ considerably between TPSF and non-peat tropical forests (Cannon et al., 2007a; Harrison, 2013), and the ultimate goal of a modelling approach should be the generation of models that can produce context-specific projections that capture these differences.

Our models of orangutan movement and seed dispersal provide projective capacity for novel locations or ecosystems by being data-referential. While our model was not completely static, allowing for extrapolation to other TPSF areas, the next step towards a fully predictive model would be using such models to identify training areas, overlaying T-LoCoH models with mechanistic niche envelope estimates (Austin, 2007; Kearney and Porter, 2009; Kearney et al., 2010, 2012; Mesgaran et al., 2014), making it possible to project orangutan movements and seed dispersal without a priori expectations in novel habitats.

A major criticism of modelling focused research programs is that the model represents a set of evidence-based hypotheses that are rarely tested (Tomlinson et al. 2014). Our internal statistical tests notwithstanding, it should be noted that we have not provided any empirical tests of our model hypotheses here. The modelling of seed dispersal, whilst being a process that contributes to the population structures of the plants dispersed (McConkey, 2000; Wang and Smith, 2002; Jordano et al., 2007; Couzens et al., 2010; Cortes and Uriarte, 2013), and the community that results (Howe and Miriti, 2000; Wang and Smith, 2002; Bascompte and Jordano, 2007; McConkey et al., 2012), is also a model prediction of plant maternal gene flow (Wang and Smith, 2002; Jordano et al., 2007; Hamrick and Trapnell, 2011). This implies that measurements of plant maternal gene flow could be used to test these models. These could be carried out using parentage assignment of seeds collected from orangutan defecation within the bounds of the models constructed herein, using an array of emerging next-generation sequencing technologies (Pritchard et al., 2000; Chen et al., 2007; Poland et al., 2012; Grabowski et al., 2014).

Conclusions

- Our data provide a mechanistic link between animal movements and the provision of endozoochory. The approach offers a powerful tool to reliably begin predicting the primary deposition of seeds by a large-bodied species such as the orangutan in contiguous TPSF.
- We developed a method with the ability to model and predict seed movements with changing orangutan populations by modelling the ecological cascade of endozoochory mechanistically. This is applicable to the continued study of orangutans at this study site.
- Changes to orangutan population structure and number, particularly female populations, has a potential flow-on effect to floristic composition heterogeneity in TPSF. Furthermore, changes to vegetation structure and productivity may initiate a feedback loop on seed dispersal, since female movement patterns and seed dispersal shadows appear to be dependent upon foraging patterns.
- This is the first objective tool of its kind in orangutan ecological research in TPSF and the first application of T-LoCoH to ecological service provision anywhere. We believe that this process is useful for establishing a training region for mechanistic models to make a priori projections of seed dispersal dynamics in novel ecosystems.

Supplementary material

Supplementary material is available at Conservation Physiology online.

Author contribution statement

E.T., A.J.M., M.E.H. and H.M.B. conceived the ideas and designed methodology. E.T. collected the data, using the
study site and orangutan follow protocols set up by H.M.B. and managed by both H.M.B. and M.E.H. E.T., A.M. and S.T. analysed the data and led the writing of the manuscript. All authors contributed critically to the drafts and gave final approval for publication.

Acknowledgements

This work was conducted in Australia with animal ethics approval for Taronga Zoo (AEC #4a/11/11) and Perth Zoo (Ar&EZA/4991-4 #59404) and in Indonesia with approval from the Indonesian Ministry of Research and Technology (RISTEK) SIP#: 349/SIP/FRP/SM/IX/2012-13. The zoo component of this study was made possible with support from many members of both Taronga and Perth Zoo. At Taronga Zoo, we would like to thank Larry Voglenest, Rebecca Spindler, Paul Thompson; Louise Grossman; Michael Shiels and Katie Hooker in particular, as well as all the other wonderful and helpful Taronga Zoo primate keepers and Jo Wiszniewski in the research office. At Perth Zoo, we would like to thank Katja Geschke, Holly Thompson, Kylie Bullo and all the other fantastic Perth Zoo primate keepers, and Caroline Lawrence in the research office. We would like to thank all our sponsors in the field in Indonesia, particularly Dr Suwido Limin, Director of UPT LLG CIMTROP (the Centre for the International Cooperation in Sustainable Management of Tropical Peatlands) at the University of Palangka Raya, who passed away in June 2016, Napa J. Awat, former Rektor of Tropical Peatlands) at the University of Palangka Raya, for supporting us for the International Cooperation in Sustainable Management of Tropical Peatlands) at the University of Palangka Raya, for supporting us for the last 15 years and supporting research in the Laboratory for the Study of Peat-Swamp Forest (LAHG), and the Indonesian Institute of Sciences, LIPI, and RISTEK. We thank our financial supporters: The US Fish and Wildlife Service Great Ape Conservation Fund, Borneo Nature Foundation, Primate Conservation Inc., the L.S.B. Leakey Foundation. Special thanks to all our field assistants who helped collecting data, Susan Cheyne contributed supporting data on seed morphology. Lastly, our sincere thanks to an anonymous reviewer and to Dr Mariëlle van Toor for their insightful and helpful comments, it is much appreciated.

Funding

E. Tarszisz was supported by an Australian Postgraduate Award (APA) from the Australian Research Council 2010-2015 as well as a DIISRTE-Funded Short-term Mobility Scholarship in 2012. S. Tomlinson was initially financially supported in this work by an Australian Government Department of Human Services Newstart unemployment allowance and latterly by the Australian Research Council grant IC150100041.

References

Abidi-Lartey M, Dechmann DK, Wikelski M, Scharf AK, Fahr J (2016) Long-distance seed dispersal by straw-coloured fruit bats varies by season and landscape. Global Ecol Conserv 7: 12–24.

Albert A, Hambuckers A, Culver L, Savini T, Huynen M-C (2013) Frugivory and Seed Dispersal by Northern Pigtailed Macaques (Macaca leonina), in Thailand. Int J Primatol 34: 170–193.

Anderson DJ (1982) The home range: a new nonparametric estimation technique. Ecology 63: 103–112.

Austin M (2007) Species distribution models and ecological theory: a critical assessment and some possible new approaches. Ecol Modell 200: 1–19.

Bascompte J, Jordano P (2007) Plant-animal mutualistic networks: the architecture of biodiversity. Ann Rev Ecol Evol Syst 38: 567–593.

Beaudrot L, Struiebig MJ, Meijaard E, Van Balen S, Husson SJ, Young CF, Marshall AJ (2013) Interspecific interactions between primates, birds, bats, and squirrels may affect community composition on Borneo. Am J Primatol 75: 170–185.

Bradshaw SD, Bradshaw FJ (2002) Short-term movements and habitat use of the marsupial honey possum (Tarsipes rostratus). J Zool 258: 343–348.

Bradshaw SD, Phillips RD, Tomlinson S, Holley RJ, Jennings S, Bradshaw FJ (2007) Ecology of the honey possum, Tarsipes rostratus, in Scott National Park, Western Australia. Austral Mammal 29: 25–38.

Buckley BJ (2014) Ranging behaviour of wild male orang-utans in an unfragmented Bornean habitat: Implications for mating-system mechanics, Cambridge, UK PhD thesis University of Cambridge

Burt WH (1943) Territoriality and home range concepts as applied to mammals. J Mammal 24: 346–352.

Börger L, Dalziel BD, Fryxell JM (2008) Are there general mechanisms of animal home range behaviour? A review and prospects for future research. Ecol Lett 11: 637.

Cagnacci F, Boitani L, Powell RA, Boyce MS (2010) Animal ecology meets GPS-based radiotelemetry: a perfect storm of opportunities and challenges. Philos Trans R Soc B 365: 2157–2162.

Cannon CH, Curran LM, Marshall AJ, Leighton M (2007a) Beyond mast-fruited events: community asynchrony and individual dormancy dominate woody plant reproductive behavior across seven Bornean forest types. Curr Sci 93: 1558–1566.

Cannon CH, Curran LM, Marshall AJ, Leighton M (2007b) Long-term reproductive behaviour of woody plants across seven Bornean forest types in the Gunung Palung National Park (Indonesia): supranual synchrony, temporal productivity and fruiting diversity. Ecol Lett 10: 956–969.

Caton J, Hume I, Hill D, Harper P (1999) Digesta retention in the gastro-intestinal tract of the orang utan (Pongo pygmaeus), Primates 40: 551–558.

Chen C, Durand E, Forbes F, François O (2007) Bayesian clustering algorithms ascertaining spatial population structure: a new computer program and a comparison study. Mol Ecol Notes 7: 747–756.

Clark B (2016) Earth Point: tools for Google Earth http://www.earthpoint.us
Corlett RT (2017) Frugivory and seed dispersal by vertebrates in tropical and subtropical Asia: an update. Global Ecol Conserv 11: 1–22.

Cousens RD, Hill J, French K, Bishop ID (2010) Towards better prediction of seed dispersal by animals. Funct Ecol 24: 1163–1170.

Culot L, Muñoz Lazo FJJ, Huynen MC, Poncin P, Heymann EW (2010) Seasonal variation in seed dispersal by tamarins alters seed rain in a secondary rain forest. Int J Primatol 31: 553–569.

Côrtes MC, Uriarte M (2013) Integrating frugivory and animal movement: a review of the evidence and implications for scaling seed dispersal. Biol Rev 88: 255–272.

Donoso I, Schleuning M, García D, Fründ J (2017) Defaunation effects on plant recruitment depend on size matching and size trade-offs in seed-dispersal networks. Proc R Soc B 284: 20162664. The Royal Society.

Dormann CF, Schymanski SJ, Cabral J, Chuine I, Graham C, Hartig F, Kearney M, Morin X, Römermann C, Schröder B (2012) Correlation and process in species distribution models: bridging a dichotomy. J Biogeogr 39: 2119–2131.

D’Arcy LJ, Graham LLB (2008) The potential effects of naturally low rates of secondary seed dispersal, coupled with a reduction in densities of primary seed dispersers on forest tree species diversity in regenerating peat swamp forest. In Wösten JHM, Rieley JO, Page SE, eds. Restoration of Tropical Peatlands. Alterra—Wageningen University and Research Centre and the EU INCO—RESTORPEAT, Wageningen, The Netherlands, pp 45–55.

Fleming CH, Calabrese JM, Mueller T, Olson KA, Leimgruber P, Fagan WF (2014) Non-Markovian maximum likelihood estimation of auto-correlated movement processes. Methods Ecol Evol 5: 462–472.

Fuzessy LF, Cornelissen TG, Janson C, Silveira FAO (2016) How do primates affect seed germination? A meta-analysis of gut passage effects on neotropical plants. Oikos 125: 1069–1080.

Fuzessy LF, Janson CH, Silveira FAO (2017) How far do Neotropical primates disperse seeds? Am J Primatol. DOI:10.1002/ajp.22659.

Galdikas BMF (1982) Orangutans as seed dispersers at Tanjung Puting, Central Kalimantan: implications for conservation. In de Boer LEM, ed, The Orang Utan: Its biology and conservation, Vol. 2. Dr W Junk Publishers, The Hague, Netherlands, p 285–298.

Getz WM, Fortmann-Roe S, Cross PC, Lyons AJ, Ryan SJ, Wilmers CC (2007) LoCoH: nonparametric kernel methods for constructing home ranges and utilization distributions. PLoS One 2: e207.

Getz WM, Saltz D (2008) A framework for generating and analyzing movement paths on ecological landscapes. Proc Natl Acad Sci U S A 105: 19066–19071.

Getz WM, Wilmers CC (2004) A local nearest-neighbor convex-hull construction of home ranges and utilization distributions. Ecography 27: 489–505.

Grabowski PP, Morris GP, Casler MD, Borevitz JO (2014) Population genomic variation reveals roles of history, adaptation and ploidy in switchgrass. Mol Ecol 23: 4059–4073.

Guisan A, Thuiller W (2005) Predicting species distribution: offering more than simple habitat models. Ecol Lett 8: 993–1009.

Guisan A, Zimmerman NE (2000) Predictive habitat distribution models in ecology. Ecol Model 135: 147–186.

Hamrick JL, Trapnell DW (2011) Using population genetic analyses to understand seed dispersal patterns. Acta Oecol 37: 641–649.

Harrison ME (2013) Using conceptual models to understand ecosystem function and impacts of human activities in tropical peat-swamp forests. Wetlands 33: 257–267.

Harrison ME, Morrogh-Bernard HC, Chivers DJ (2010) Orangutan energetics and the influence of fruit availability in the nonmasting peat-swamp forest of Sabangau, Indonesian Borneo. Int J Primatol 31: 585–607.

He T, Lamont BB, Krauss SL, Enright NJ, Miller BP, Gove AD (2009) Ants cannot account for interpopulation dispersal of the arillate pea Daviesia triflora. New Phytol 181: 725–733.

Hebblewhite M, Haydon DT (2010) Distinguishing technology from biology: a critical review of the use of GPS telemetry data in ecology. Philos Trans R Soc London B Biol Sci 365: 2303–2312.

Hooijer A, Page S, Canadell JG, Silvius M, Kwadijk J, Wösten H, Jauhiainen J (2010) Current and future CO2 emissions from drained peatlands in Southeast Asia. Ecosystems 12: 888–905.

Howe HF, Miriti MN (2000) No question: seed dispersal matters. Trends Ecol Evol 15: 434–436.

Jachowski DS, Singh NJ (2015) Toward a mechanistic understanding of animal migration: incorporating physiological measurements in the study of animal movement. Conserv Physiol 3: cov035.

Jauhiainen J, Takahashi H, Heikkinen JEP, Martikainen PJ, Vasander H (2005) Carbon fluxes from a tropical peat swamp forest floor. Global Change Biol 11: 1788–1797.

Jordano P, Garcia C, Godoy JA, Garcia-Castano JL (2007) Differential contribution of frugivores to complex seed dispersal patterns. Proc Natl Acad Sci U S A 104: 3278–3282.

Kays R, Jansen PA, Knecht EMH, Vohwinkel R, Wikelski M (2011) The effect of feeding time on dispersal of Viola seeds by toucans determined from GPS tracking and accelerometers. Acta Oecol Int J Ecol 37: 625–631.

Kearnery MR, Matzelle A, Helmuth B (2012) Biomechanics meets the ecological niche: the importance of temporal data resolution. J Exp Biol 215: 922–933.

Kearnery M, Porter W (2009) Mechanistic niche modelling: combining physiological and spatial data to predict species ranges. Ecol Lett 12: 334–350.

Kearnery MR, Wintle BA, Porter WP (2010) Correlative and mechanistic models of species distribution provide congruent forecasts under climate change: congruence of correlative and mechanistic distribution models. Conserv Lett 3: 203–213.

Kenward RE, Walls SS, South AB, Casey N (2008) Ranges8: For the analysis of tracking and location data. Anatrack Ltd, Wareham, UK.
Kie JG, Matthiopoulos J, Fieberg J, Powell RA, Cagnacci F, Mitchell MS, Gaillard J-M, Moorcroft PR (2010) The home-range concept: are traditional estimators still relevant with modern telemetry technology? Philos Trans R Soc London B Biol Sci 365: 2221–2231.

Krauss SL, He T, Barrett LG, Lamont BB, Enright NJ, Miller BP, Hanley ME (2009) Contrasting impacts of pollen and seed dispersal on spatial genetic structure in the bird-pollinated Banksia hookeriana. Heredity 102: 274–285.

Laver PN, Kelly MJ (2008) A critical review of home range studies. J Wildlife Manag 72: 290–298.

Lenz J, Fiedler W, Caprano T, Friedrichs W, Gaese BH, Wikelski M, Boehning-Gaese K (2011) Seed-dispersal distributions by trumpeter hornbills in fragmented landscapes. Proc R Soc -Biul Sci 278: 2257–2264.

Levin SA, Muller-Landau HC, Nathan R, Chave J (2003) The ecology and evolution of seed dispersal: a theoretical perspective. Annual Rev Ecol Evolution System 34: 575–604.

Lyons A, Getz WG, R DCT (2015). T-LoCoH: Time Local Convex Hull homerange and time use analysis Vol. 1.37.00, R Core Team

Lyons AJ, Turner WC, Getz WM (2013) Home range plus: a space-time characterization of movement over real landscapes. Movement Ecol 1: 1–14.

Martin P, Bateson P (1986) Measuring behaviour: an introductory guide. Cambridge University Press, New York, USA.

McCallum KP, McDougall FO, Seymour RS (2013) A review of the energetics of pollination biology. J Comp Physiol B 183: 867–876.

McCaughey DJ, DeSalles PA, Young HS, Dunbar RB, Dirzo R, Mills MM, Micheli F (2012) From wing to wing: the persistence of long ecological interaction chains in less-disturbed ecosystems. Sci Rep 2: 409.

McConkey KR (2000) Primary seed shadow generated by gibbons in the rain forests of Barito Ulu, central Borneo. Am J Primatol 52: 13–29.

McConkey KR, Prasad S, Corlett RT, Campos-Arceiz A, Brodie JF, Rogers H, Santamaria L (2012) Seed dispersal in changing landscapes. Biol Conserv 146: 1–13.

Menz MHM, Phillips RD, Winfree R, Kremen C, Aizen MA, Johnson SD, Dixon KW (2011) Reconnecting plants and pollinators: challenges in the restoration of pollination mutualisms. Trends Plant Sci 16: 4–12.

Mesgaran MB, Cousens RD, Webber BL (2014) Here be dragons: a tool for quantifying novelty due to covariate range and correlation change when projecting species distribution models. Divers Distributions 20: 1147–1159.

Mitra Setia T, Delgado RA, Utami Atmoko SS, van Schaik CP (2009) Social organization and male-female relationships. In Wich SA, Utami Atmoko SS, Mitra Setia T, vSC P, eds. Orangutans: Geographic Variation in Behavioral Ecology and Conservation. Oxford University Press, Oxford, UK, pp 245–253.

Morales JM, Moorcroft PR, Matthiopoulos J, Frair JL, Kie JG, Powell RA, Merrill EH, Haydon DT (2010) Building the bridge between animal movement and population dynamics. Philos Trans Royal Soc London B Biol Sci 365: 2289–2301.

Morrogh-Bernard HC (2009) Orang-utan behavioural ecology in the Sabangau peat-swamp forest, Borneo, Unpublished D. Phil, University of Cambridge

Morrogh-Bernard HC, Husson SJ, Haranto FA, Chivers DJ (2014) Fine-scale habitat use by orang-utans in a disturbed peat swamp forest, Central Kalimantan, and implications for conservation management. Folia Primatol 85: 135–153.

Morrogh-Bernard HC, Husson SJ, Knott CD, Wich SA, van Schaik CP, van Noordwijk MA, Lackman-Ancrenaz I, Marshall AJ, Kanamori T, Kuze N, et al (2009) Orangutan activity budget and diet. In Wich SA, Utami Atmoko SS, Mitra Setia T, van Schaik CP, eds. Orangutans: Geographic Variation in Behavioral Ecology and Conservation. Oxford University Press, Oxford, UK, pp 119–134.

Morrogh-Bernard H, Husson S, McLardy CS (2002). Orang-utan data collection standardisation, LSJ Leakey Foundation, Orangutan culture workshop, February, San Anselmo, California, USA

Morrogh-Bernard HC, Husson SJ, Page SE, Rieley JO (2003) Population status of the Bornean orang-utan (Pongo pygmaeus) in the Sebangau peat swamp forest, Central Kalimantan, Indonesia. Biol Conserv 110: 141–152.

Morrogh-Bernard HC, Morf NV, Chivers DJ, Krützen M (2011) Dispersal patterns of orang-utans (Pongo spp.) in Bornean peat-swamp forest. Int J Primatol 32: 362–376.

Muller-Landau HC (2007) Predicting the long-term effects of hunting on plant species composition and diversity in tropical forests. Biotropica 39: 372–384.

Munn AJ, Dawson TJ, McLeod SR, Dennis T, Maloney SK (2013) Energy, water and space use by free-living red kangaroos Macropus rufus and domestic sheep Ovis aries in an Australian rangeland. J Comp Phys 183: 843–858.

Nathan R, Getz WM, Revilla E, Holyoak M, Kadmon R, Saltz D, Smouse PE (2008a) A movement ecology paradigm for unifying organismal movement research. Proc Natl Acad Sci U S A 105: 19052–19059.

Nathan R, Katul GG, Bohrer G, Kuparinne A, Soons MB, Thompson SE, Trakhtenbrot A, Horn HS (2011) Mechanistic models of seed dispersal by wind. Theoretical Ecol 4: 113–132.

Nathan R, Katul GG, Horn HS, Thomas SM, Oren R, Avisser R, Pacala SW, Levin SA (2002) Mechanisms of long-distance dispersal of seeds by wind. Nature 418: 409–413.

Nathan R, Muller-Landau HC (2000) Spatial patterns of seed dispersal, their determinants and consequences for recruitment. Trends Ecol Evol 15: 278–285.

Nathan R, Schurr FM, Spiegel O, Steinitz O, Trakhtenbrot A, Tsar A (2008b) Mechanisms of long-distance seed dispersal. Trends Ecol Evol 23: 638–647.
Nielsen NH, Jacobsen MW, Graham L, Morrogh-Bernard HC, D’Arcy LJ, Harrison ME (2011) Successful germination of seeds following passage through orang-utan guts. *J Trop Ecol* 27: 433–435.

Page SE, Rieley JO, Banks CJ (2011) Global and regional importance of the tropical peatland carbon pool. *Global Change Biol* 17: 798–818.

Page SE, Rieley JO, Shotyk W, Weiss D (1999) Interdependence of peat and vegetation in a tropical peat swamp forest. *Philos Trans R Soc London B Biol Sci* 354: 1885–1897.

Pascoal CM, Nevill PG, Elliott CP, Major JD, Anthony JM, Krauss SL (2015) The critical role of ants in the extensive dispersal of Acacia seeds revealed by genetic parentage assignment. *Oecologia* 179: 1123–1134.

Peres CA, van Roosmalen M (2002) Primate frugivory in two species-rich neotropical forests: implications for the demography of large-seeded plants in overhunted areas. In Levey DJ, Silva WR, Galetti M, eds. *Seed Dispersal and Frugivory: Ecology, Evolution, and Conservation*. CAB International, New York, pp 407–422.

Poland JA, Brown PJ, Sorrells ME, Jannink JL (2012) Development of high-density genetic maps for barley and wheat using a novel two-enzyme genotyping-by-sequencing approach. *PLoS One* 7:e32253.

Posa MR, Wijedasa LS, Corlett RT (2011) Biodiversity and conservation of tropical peat swamp forests. *BioScience* 61: 49–57.

Pritchard JK, Stephens M, Donnelly P (2000) Inference of population structure using multilocus genotype data. *Genetics* 155: 945–959.

Quin DG, Smith AP, Green SW, Hines HB (1992) Estimating the home ranges of sugar gliders (*Petaurus breviceps*) (Marsupialia: Petauridae), from grid-trapping and radiotelemetry. *Wildlife Res* 19: 471–487.

R Core Team (2015) *R*: a language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria. Ed 2015.

R Studio Team (2015) *RStudio: integrated Development for R*. RStudio, Inc, Boston, MA, USA.

Rieley JO, Page SE, Limin SH, Winarti S (1997) The peatland resources of Indonesia and the Kalimantan peat swamp forest research project. In Rieley JO, Page SE, eds. *Biodiversity and Sustainability of Tropical Peatlands*. Samara Publishing, Cardigan, UK, pp 37–44.

Robertson AW, Trass A, Ladley JJ, Kelly D (2006) Assessing the benefits of frugivory for seed germination: the importance of the deinhibiting effect. *Funct Ecol* 20: 58–66.

Russo SE, Portnoy S, Augspurger CK (2006) Incorporating animal behavior into seed dispersal models: implications for seed shadows. *Ecology* 87: 3160–3174.

Ruxton GD, Schaefer HM (2012) The conservation physiology of seed dispersal. *Philos Trans R Soc B* 367: 1708–1718.

Schupp EW, Jordano P, Gómez JM (2010) Seed dispersal effectiveness revisited: a conceptual review. *New Phytol* 188: 333–353.

Seltzer CE, Ndangalasi HJ, Cordeiro NJ (2013) Seed dispersal in the dark: shading light on the role of fruit bats in Africa. *Biotropica* 45: 450–456.

Singleton I, Knott CD, Morrogh-Bernard HC, Wich S, van Schaik CP (2009) Ranging behavior of orangutan females and social organisation. In Wich SA, Utami Atomoko SS, Mitra Setia T, van Schaik CP, eds. *Orangutans: Geographic Variation in Behavioral Ecology and Conservation*. Oxford University Press, UK, pp 205–213.

Singleton I, van Schaik CP (2001) Orangutan home range size and its determinants in a Sumatran swamp forest. *Int J Primatol* 22: 877–911.

Soberón J, Nakamura M (2009) Niches and distributional areas: concepts, methods, and assumptions. *Proc Natl Acad Sci* 106: 19644–19650.

Stevenson PR, Link A, Onshuus A, Quiroz AJ, Velasco M (2014) Estimation of seed shadows generated by Andean woolly monkeys (*Lagothrix lagothricha luguens*). I. *Int J Primatol* 35: 1021–1036.

Tarszisz E, Tomlinson S, Harrison ME, Morrogh-Bernard HC, Munn AJ (2017) Gardeners of the forest: effects of seed handling and ingestion by orangutans on germination success of peat forest plants. * Biol J Linnean Soc* 123: 125–134.

Tomkiewicz SM, Fuller MR, Kie JB, Gates K (2010) Global positioning system and associated technologies in animal behaviour and ecological research. *Philos Trans R Soc B* 365: 2163–2176.

Tomlinson S, Arnall S, Munn AJ, Bradshaw SD, Maloney SK, Dixon KW, Didham RK (2014) Applications and implications of ecological energetics. *Trends Ecol Evol* 29: 280–290.

Traveset A, Robertson AW, Rodríguez-Perez J (2007) A review on the role of endozoochory in seed germination. In Dennis AJ, Green RJ, Schupp EW, eds. *Seed dispersal: Theory and its Application in a Changing World*. CAB International, Wallingford, Oxon, UK, pp 78–103.

Traveset A, Verdú M (2002) A meta-analysis of the effect of gut treatment on seed germination. In Levey DJ, Galetti M, eds. *Seed Dispersal and Frugivory: Ecology, Evolution and Conservation*. CAB International, Wallingford, UK, pp 339–350.

Tylianakis JM, Laliberté É, Nielsen A, Bascompte J (2010) Conservation of species interaction networks. * Biol Conserv* 143: 2270–2279.

Utami Atomoko SS, Singleton I, van Noordwijk MA, van Schaik CP, Mitra Setia T (2009) Male-male relationships in orangutans. In Wich SA, Utami Atomoko SS, Mitra Setia T, van Schaik CP, eds. *Orangutans: Geographic Variation in Behavioral Ecology and Conservation*. Oxford University Press, UK, pp 225–233.

Vidal MM, Pires MM, Guimarães PR Jr (2013) Large vertebrates as the missing components of seed-dispersal networks. * Biol Conserv* 163: 42–48.

Wang BC, Smith TB (2002) Closing the seed dispersal loop. *Trends Ecol Evol* 17: 379–386.

Wotton DM, Kelly D (2011) Frugivore loss limits recruitment of large-seeded trees. *Proc. R Soc B Biol Sci* 278: 3345–3354.

Wright SJ, Trakhtenbrot A, Bohrer G, Detto M, Katul GG, Horvitz N, Muller-Landau HC, Jones FA, Nathan R (2008) Understanding strategies for seed dispersal by wind under contrasting atmospheric conditions. *Proc Natl Acad Sci USA* 105: 19084–19089.