Comparative studies need to rely both on sound natural history data and on excellent statistical analysis

Dieter Lukas and Tim Clutton-Brock

Department of Zoology, University of Cambridge, Downing Street, Cambridge CB2 3EJ, UK

DL, 0000-0002-7141-3545; TCB, 0000-0001-8110-8969

We certainly agree with Schradin [1] that comparative studies need to use the best information available—as well as clear definitions, up-to-date statistical techniques and common sense. We have divided our response into two sections: the first dealing with Schradin’s critique of our 2017 paper and the classification of the social organization of shrews; the second dealing with three general issues that his commentary raises.

1. The social organization of shrews

The paper that Schradin is commenting on [2] asks whether cooperative breeders (which we define as species where non-breeding helpers assist breeders to raise their young) in mammals are more commonly found in arid environments than monogamous species (defined as species where breeding pairs of males and females remain together for more than one season) from which cooperative breeders appear to be derived [3]. It shows that, as in birds, cooperative breeders tend to live in relatively arid habitats compared to monogamous species. As we describe below, these conclusions are unaffected by differences between our classification of the social organization of shrews and the classification suggested by Schradin.

Schradin’s first criticism is that the categorizations of shrews that we have used in this analysis were based on one paper on a single species. This is incorrect, for our categorization of shrews was based on more than 70 separate sources (see electronic supplementary material, appendix S1). When we published our dataset [4], we were required to reduce the associated reference list by the editors of the journal and, to comply, we removed around 75% of our 2000+ sources, often listing a single reference per species or group of species, and frequently using generic reviews for particular groups where these were available: for
example, for many rodent species, we simply listed Wolff and Sherman’s *Rodent Societies* [5], and for many carnivores, we listed the review by Dalenrum [6]. To warn readers that our dataset was not derived from these generic references alone, we inserted a comment in the electronic supplementary material of the paper that ‘although the supplement provides a single reference for each species for further information, the classification of most species was based on information from several sources’ [4]. The traditional view of social organization in shrews and the consensus of most overviews of the social organization of shrews is that breeding females are intolerant of each other and that multiple breeding females do not form coherent social groups (see [7–11]). Although the generic reference that we cited in connection with our classifications of shrews [12] contained a statement that shrews are solitary in its Introduction, it was not a general review and we agree with Schradin that it was not a suitable source to quote in this connection and that it would have been more appropriate to have listed one or more sources that provided extensive reviews or primary information, like many of those listed in electronic supplementary material, appendix S1.

Schradin’s second criticism of our 2017 paper concerns differences in the classification of eight shrew species between our 2017 paper and his 2015 paper with Valomy et al. [13]. Three species that Valomy et al. classify as group living in their table (*Cryptotis parva*, *Sorex cinereus* and *Sorex ornatus*), we classify as solitary breeders (see table 1). These differences are related to contrasts in definitions. We classify species as social breeders only if multiple breeding females aggregate in coherent groups during the breeding season and classify them as solitary breeders if breeding females are intolerant of each other and do not aggregate with each other in the breeding season (the rationale for this is described in box 1). The papers that Valomy et al. refer to in their 2015 paper provide no evidence that female shrews are social during the breeding season and so do not allow us to classify them as social breeders. Valomy et al. [13] do not provide definitions of their categories but evidently used a less restrictive definition of group living. *Cryptotis parva* are reported to aggregate during the winter for thermo-regulatory reasons, but it is not clear whether females aggregate during the breeding season [22,23]: reports of sociality in *S. cinereus* describe parties of males competing to gain access to single females during the mating season but provide no evidence that multiple breeding females aggregate in coherent groups or share a common range [15]; and, while aggregations of *S. ornatus* (*simouosus*) occur outside the breeding season, they appear to be loose and unstable and there is no indication that breeding females form coherent groups during the breeding season [14,16] (table 1).

A further four species that we have classified as solitary breeders are classified as pair-living by Valomy et al. Contrasts in definitions are involved here, too. We classify species as socially monogamous (or pair-living) if there is evidence that they form stable mixed sex pairs which persist across breeding attempts or breeding seasons (the rationale for this is described in box 1 and is related to the need to distinguish between pair-living species and the large number of mammals where individual males guard single females for short periods during the breeding season before moving on to search for other partners) and regard socially monogamous species as a subset of species where breeding females do not aggregate with each other. Although the four species that Valomy et al. list as pair-living (*Crocidura leucodon*, *Crocidura russula*, *Sorex coronatus* and *Suncus varilla*) are commonly seen in mixed sex pairs during the breeding season, none of the studies they cite tracked the movements of substantive samples of identifiable individuals over time and it is not clear whether the same individuals associate with each other either throughout particular breeding attempts or across breeding attempts in the same season—and some observations suggest that they break up at the end of each breeding season. Moreover, all four species are short-lived and few pairs, if any, can persist across years. In this respect, shrews differ from some murid rodents, including species of *Microtus* and *Peromyscus*, where there is firm evidence that individuals form lifelong pairs which persist across breeding attempts and seasons (when new individuals start to reproduce) if both partners survive (e.g. [24–26]). As there is no firm evidence that individuals form pairs that persist across breeding attempts or seasons or that breeding females aggregate during the breeding season in these four shrew species, we classified these species as solitary breeders, too (table 1). A final difference is that we classified one species, *Suncus etruscus*, as monogamous because it was among the species listed in an early review by Kleiman & Malcolm [27] as being monogamous. However, as Schradin points out, this information is not sufficient to classify its social system and we now regard this as an error, as there is no evidence that pairs remain together across breeding attempts or seasons and we have found new reports that females are solitary and aggressive towards males outside the breeding season [28] and shall change its categorization in any future analysis. To check whether contrasts in the classifications of shrews between Valomy et al. and ourselves affected the conclusions of our analysis, we have re-run the comparisons included in our 2017 paper using Valomy et al.’s classifications and found that they do not (see electronic supplementary material, appendix S3).
Table 1. Differences in the classification of shrews.

| species            | Do breeding females aggregate in groups? | Are aggregations during the breeding season stable and cohesive? | Are individuals seen in mixed sex pairs during the breeding season? | Do males remain with the same female throughout the breeding season? | Do pairs remain stable across breeding seasons? | classification Lukas & Clutton-Brock | classification Valomy et al. |
|--------------------|------------------------------------------|-----------------------------------------------------------------|------------------------------------------------------------------|---------------------------------------------------------------------|------------------------------------------------|---------------------------------|---------------------------------|
| Cryptotis parva    | no: aggregations dissolve and females build territories [14] | no                                                              | no                                                               | no                                                                   | no                                              | solitary                        | group                           |
| Sorex cinereus     | no: aggregations appear to be of males chasing females [15] | no                                                              | no                                                               | no                                                                   | no                                              | solitary                        | group                           |
| Sorex ornatus      | yes                                       | no: frequent changes [16], and females intolerant [14]           | no                                                               | no                                                                   | no                                              | solitary                        | group                           |
| Crocidura leucodon | no                                        | no                                                              | yes                                                              | potentially                                                          | no: in autumn, shift to a gregarious way of life [17]          | solitary                        | pair                            |
| Crocidura russula  | no                                        | no                                                              | uncertain: pairs occur, but high frequency of polygyny [18]      | uncertain                                                            | no: during winter, shift to mixed sex aggregations [19]         | solitary                        | pair                            |
| Sorex coronatus    | no                                        | no                                                              | yes                                                              | no: male home-ranges change during breeding season [20]            | no: in autumn, males separate from females [20]                 | solitary                        | pair                            |
| Suncus varilla     | no                                        | no                                                              | yes                                                              | possibly, but there is only a single breeding attempt per year [21] | uncertain: partners appear to separate after breeding season [21] | solitary                        | pair                            |
Box 1. Definitions of categories of social organization used in our 2013 and 2017 papers.

The definition of the categories that we used to classify social organization in shrews and other mammals are described in the main text of our 2013 paper, with more detail in the associated electronic supplementary material [4]. We first considered the behaviour of breeding females, classifying species as social breeders if, during the course of the breeding season, several females that breed regularly share a common range, tolerate each other and form coherent groups—and as solitary breeders if they do not do so. This definition is intended to distinguish between species where females are solitary, accompany dependent young or aggregate outside the breeding season or only at foraging grounds and the relatively small number of mammals where social groups include multiple breeding females which share all areas of their range.

We classify species as socially monogamous if breeding females do not aggregate or share a common range and there is evidence that individual males and individual females form breeding pairs that persist for more than a single breeding season (in species where individuals reach maturity quickly, multiple breeding seasons might occur within a single year, though in most species pairs persist for more than 1 year). Here, our definition is intended to distinguish between the relatively large number of polygynous mammals where individual males guard single receptive females for minutes, hours or days before moving on to search for other partners and species where the sexes form bonded breeding pairs that persist for more than a single breeding season. Most monogamous mammals are iteroparous and we have not yet found firm evidence of any mammal which forms mixed sex pairs that share a common range and last throughout a breeding attempt but are not maintained across breeding attempts within years or across seasons (as is the case in many birds). However, two of the shrews listed in Table 1 (Cro. leucodon and Su. varilla) may possibly do so and firm evidence of this would cause us to consider modifying our definition to include them and any other species where mixed sex pairs persist across individual breeding attempts but not across seasons.

Finally, we classify species as cooperative breeders where breeding adults are assisted in rearing young by non-breeders of either or both sexes. This is intended to distinguish between species where several breeding females share care of their offspring and are likely to gain direct fitness benefits from doing so or where non-breeders contribute to activities likely to generate mutualistic benefits (such as nest maintenance and predator defence) from species where non-breeding subordinates engage in activities that are unlikely to increase their own direct fitness, like feeding or carrying young born to other group members.

In retrospect, it would have been useful to have included a more detailed description and discussion of our classification of these eight species in our 2017 paper and to have explained the reasons why we did not follow Valomy et al.’s classifications.

Schradin’s final criticism is that our 2017 paper draws on our 2013 dataset where we list all shrew species as solitary breeders. This is in line with most reviews of the social organization of shrews, which describe them as solitary breeders (see above); with the absence of evidence that breeding females aggregate in any of the 68 species for which we were able to find published information; and with the absence of group breeding (on our definition) in any other small, terrestrial insectivores or carnivores in our dataset. The principal comparisons in our 2017 paper that Schradin is commenting on did not include species whose social organization we inferred, as we compared the habitats of socially monogamous species and cooperative breeders (where all categorizations were based on published information). However, the paper contains a subsidiary comparison between the habitats occupied by cooperative breeders and those occupied by all other mammals, including 51 shrews for which we could find published information and 87 whose social organization we inferred to be similar (see electronic supplementary material, appendix S2). Here, we included data for the second group of species because the consensus view is that all shrews are solitary breeders and their inclusion helped to compensate for the under-representation of shrews as a result of the relative scarcity of studies. We agree with Schradin that there is always a danger that inclusion of species whose social organization is inferred may introduce error—though their exclusion has disadvantages, too (see below). A sensible course in situations of this kind would seem to be to repeat analyses with and without inferred data and we have consequently checked whether the inclusion of the shrews listed in our dataset for which no published reports were available could have affected the results of our comparisons. A re-run of our analyses shows that our results are unchanged if they are excluded (see electronic supplementary material, appendix S3).
also re-ran the comparisons using only Valomy et al.’s categorizations and found that, here too, our results are unaffected by these differences (see electronic supplementary material, appendix S3). As a case could be made that all shrew species should be omitted from comparative studies of the distribution of contrasting forms of social organization until their social behaviour has been studied in more detail, we also investigated whether the removal of all shrews from our comparisons affected their outcome. Here, too, we found that the outcome of the analysis was unchanged (see electronic supplementary material, appendix S3). We also checked the results of previous analyses that have included these categorizations and found that they, too, are unaffected by whether or not these species are included.

2. Broader issues

Schradin’s commentary on our paper raises three general methodological issues regarding comparisons of major taxonomic groups based on categorizations of large samples of species. First, it emphasizes the importance of clear definitions and the need to be aware that contrasting definitions are often likely to cause the same species to be classified in different ways (table 1). Where the aim is to explore the distribution of traits across major taxonomic groups, the same definitions have to be used to classify very different species. This may require categories to be more specific and more restrictive in order to be applicable to diverse species [29,30], so definitions may diverge from those that might be used to compare more closely related species. Categories and definitions also need to be adjusted to the taxa involved and to the questions that are being asked—so they often vary between analyses of different taxa. For example, in our 2017 paper, we were principally interested in comparing the distribution of cooperative breeders with that of monogamous species where pairs persist for more than one season because phylogenetic reconstructions suggest that cooperative breeding has usually evolved in ancestors that formed long-lasting breeding pairs [3]. These issues are relevant to the use of published datasets by others: few datasets are constructed for general use and their purpose is usually to allow particular analyses to be checked and investigated. As a result, those that use published datasets for other analyses or add them to comparative databases they have compiled need to pay close attention to the definitions of categories that are being used in order to avoid combining information based on contrasting definitions. It is consequently important that comparative studies should provide clear and detailed descriptions of the definitions of their categories or variables.

A second point concerns the common problem that the distribution of information across species is often strongly affected by biases in the species that have been studied. Among mammals, the proportion of species that have been studied in any detail is much higher for diurnal, terrestrial and social species than it is for nocturnal, arboreal or solitary ones: for example, we have been able to locate published information on social behaviour for 62% of all diurnal mammals but only for 36% of all nocturnal species, including nocturnal shrews (see [31]). Data availability is also frequently geographically biased, with tropical and Southern Hemisphere species being less well represented than palaearctic ones [32,33]. As a result, comparisons of the incidence of female sociality across major taxonomic groups may need to compensate for biases in the distribution of species that have been studied. Where species from under-represented taxa, like shrews, show a high degree of conformity in their behaviour, a common approach has been to use the available information on the distribution of traits across taxa, combined with the number of species per taxon, to infer the traits of related species for which no direct data are available. In some cases, inference of this kind is based simply on the assumption that related species are likely to show the same traits, sometimes potentially supported by phylogenetic reconstructions—for example, a number of comparative studies of social behaviour (including two of the three studies that Schradin cites in his Commentary) have been based on datasets that assume that congeneric species share the same traits (e.g. [34–44]). In others, the categorization of species for which no direct data are available is inferred using ecological or life-history information in addition to phylogenetic proximity (for discussion, see [45]). At the end of his Commentary, Schradin argues against the use of inference and suggests that many comparisons may need to wait until enough detailed, high-quality datasets are available to allow quantitative comparisons to be made that do not rely on inference. We have much sympathy with his argument and, like him, believe that only long-term, individual-based studies provide reliable insights into the processes responsible for the diversity of social behaviour in mammals [46,47]. However, the proportion of species for which detailed, individual-based data are available from wild populations is likely to remain low for a long time—and we do not believe that it is sensible to delay asking questions about the distribution of major traits across different animal groups. As a result, we believe that, in some comparisons, it will be necessary to infer the traits of some species in order to reduce
Data availability. Data are available in electronic supplementary material, appendix S2 as part of the electronic supplementary material (a table listing all shrew species with observed or inferred social system included in our 2017 analyses and additional species with observed social system included in our 2013 paper).

Authors’ contributions. D.L. collected the data and carried out the statistical analyses; both authors wrote the manuscript and gave final approval for publication.

Competing interests. We declare we have no competing interests.

Funding. During the writing of this manuscript, D.L. was employed by the Max Planck Society and T.C.-B. was employed by the University of Cambridge.

References

1. Schradin C. 2017 Comparative studies need to rely both on sound natural history data and on excellent statistical analyses. R. Soc. open sci. 4, 170346. (doi:10.1098/rsos.170346)
2. Lukas D, Clutton-Brock T. 2017 Climate and the distribution of cooperative breeding in mammals. R. Soc. open sci. 4, 160897. (doi:10.1098/rsos.160897)
3. Lukas D, Clutton-Brock T 2012 Cooperative breeding and monogamy in mammalian societies. Proc. R. Soc. B 279, 2531–2536. (doi:10.1098/rspb.2011.2468)
4. Lukas D, Clutton-Brock T. 2017 The evolution of social monogamy in mammals. Science 341, 526–530. (doi:10.1126/science.1238677)
5. Wolff JD, Sherman PW. 2007 Rodent societies: an ecological and evolutionary perspective. Chicago, IL: University of Chicago Press.
6. Daleur F. 2007 Phylogenetic reconstruction of carnivore social organizations. J. Zool. 273, 90–97. (doi:10.1111/j.1469-7998.2007.00303.x)
7. Churchill S. 1990 The natural history of shrews. London, UK: Christopher Helm/A & C Black.
8. Kirkland GL, Merritt JF, Rose RK (eds). 1994 Advances in the biology of shrews. Pittsburgh, PA: Carnegie Museum of Natural History.
9. Habel W. 1996 Fostering in European common shrews Sorex araneus (Soricidae, Insectivora). Acta Theriol. 41, 433–438. (doi:10.4098/ATarch.96-43)
10. Gliwicz J, Taylor JR. 2002 Comparing life histories of shrews and rodents. Acta Theriol. 47, 185–208. (doi:10.1007/BF0292487)
11. Rychlik L, Zwołek R. 2005 Behavioural mechanisms of conflict avoidance among shrews. Acta Theriol. 50, 289–308. (doi:10.1007/BF0292627)
12. Muraru D, Benedek AM. 2005 New reports on the presence of Sorex alpinus Schinz, 1867 (Insectivora: Soricidae) in the southern Carpathians (Romania). Tranz. du Mus. Natl. Hist. Nat. ‘Grigore Antipa’ 48, 395–405.
13. Valomy M, Hayes LD, Schradin C. 2015 Social organization in Eulipotyphla: evidence for a social shrew. Biol. Lett. 11, 20150825. (doi:10.1098/rsbl.2015.0825)
14. Rychlik L. 1996 Evolution of social systems in shrews. In Evolution of shreds (eds JW Wojcik, M Wielanska), pp. 347–406. Białowieża, Poland: Mammal Research Institute, Polish Academy of Sciences.
15. Maier TJ, Doyle KL. 2006 Aggregations of masked shrews (Sorex cinereus): density-related mating behavior? Agregatien de muscaragine'es masques: (Sorex cinereus): accouplement relatif à la densité de la population? Mammalia 70, 86–89. (doi:10.1515/mamm.70.1-2.86)
16. Hays WS, Lidicker WJ. 2000 Winter aggregations, Dehnel effect, and habitat relations in the Eurasian shrew Sorex minutus minutus. Acta Theriol. 45, 433–442. (doi:10.4098/AT.arch.00-44)
17. Simeonovska-Nikolova DM. 2004 Seasonal changes in social behaviour and spatial structure of Crocidura leucodon in north-western Bulgaria. Acta Theriol. 49, 167–179. (doi:10.1007/BF03192627)
18. Boutelleur C, Perrin N. 2000 Individual reproductive success and effective population size in the greater white-toothed shrew. Crocidura rufa. Proc. R. Soc. B 267, 701–705. (doi:10.1098/rspb.2000.1059)
19. Cantoni D, Vogel P. 1989 Social organization and mating system of free-ranging, greater white-toothed shrews, Crocidura rufa. Anim. Behav. 38, 205–214. (doi:10.1006/sobu.1997.0083)
20. Cantoni D. 1993 Social and spatial organization of free-ranging shrews, Sorex coronatus and Neomys fodiens (Insectivora, Mammalia). Annu. Behav. 45, 975–995. (doi:10.1006/beha.1993.1116)
21. Lynch CD. 1991 Population dynamics in the Lesser dwarf shrew, Suncus vallilo (Mammalia: Soricidae): results. Nauvörung van de Nasionale Museum: Researches of the National Museum 7, 467–470. (doi:0526/JAK000679208_893)
22. McCallor WH. 1959 An unusually large nest of Crotoporus parvus. J. Mammal. 40, 245. (doi:10.2307/1376442)
23. Choate JR, Jones JK, Jones C. 1994 Handbook of mammals of the south-central states. London, UK: Louisiana State University Press.
24. Jannett F. 1992 Nesting patterns of adult voles, Microtus montanus, in field populations. J. Mammal. 63, 495–498. (doi:10.2307/1380450)
25. Fitzgerald RW, Madison DM. 1983 Social organization of a free-ranging population of pine voles, Microtus pinetorum. Behav. Ecol. Sociobiol. 13, 183–187. (doi:10.1007/BF00299291)
26. Ruble DD. 1992 Lifetime reproductive success and its correlates in the monogamous rodent, Peromyscus californicus. J. Anim. Ecol. 61, 457–468. (doi:10.2307/5316)
27. Kleiman DG, Malcolm JR. 1981 The evolution of male parental investment in mammals. In Parental care in mammals (eds DJ Gubernick, PH Kloper), pp. 347–387. New York, NY: Springer.
28. Kingdon J, Happold OCO, Butynski TM, Hoffmann M, Happold M, Kalmy A. 2013 Mammals of Africa. London, UK: Bloomsbury.
29. Feckleton RP. 2009 The seven deadly sins of comparative analysis. J. Ecol. Biol. 22, 1367–1375. (doi:10.1111/j.1462-9070.2009.01075.x)
30. Barresi C, Sandel AA, Koenig A, Fernandez-Duque E, Kamilar JM, Amoros CR, Gordon AD. 2016 Transparency, usability, and reproducibility: guiding principles for improving comparative databases using primate as examples. Evol. Anthropol. 25, 232–238. (doi:10.1002/evan.21002)
31. Wilman H, Beltmaker J, Simpson J, de la Rosa C, Rivadeneira MM, Jetz W. 2014 EltonTraits 1.0: species-level foraging attributes of the world’s birds and mammals. Ecology 95, 2017. (doi:10.1890/13-1917.1)
32. Cockburn A. 2003 Cooperative breeding in oscine passerines: does sociality inhibit speciation? Proc. R.
33. Amano T, González-Varo JP, Sutherland WJ. 2016 Languages are still a major barrier to global science. *PLoS Biol.* 14, e1002093. (doi:10.1371/journal.pbio.2002093)

34. Di Fiore A, Rendall D. 1994 Evolution of social organization: a reappraisal for primates by using phylogenetic methods. *Proc. Natl Acad. Sci. USA* 91, 9941–9945. (doi:10.1073/pnas.91.21.9941)

35. Dunn PO, Whittingham LA, Pitcher TE. 2001 Mating systems, sperm competition, and the evolution of sexual dimorphism in birds. *Evolution* 55, 161–175. (doi:10.1111/j.0014-3820.2001.tb00781.x)

36. Reynolds JD, Goodwin NB, Freckleton RP. 2002 Evolutionary transitions in parental care and live bearing in vertebrates. *Phil. Trans. R. Soc. Lond. B* 357, 269–281. (doi:10.1098/rstb.2001.0930)

37. Cockburn A. 2006 Prevalence of different modes of parental care in birds. *Proc. R. Soc. B* 273, 1373–1383. (doi:10.1098/rspb.2005.3458)

38. Shultz S, Dunbar RIM. 2006 Both social and ecological factors predict ungulate brain size. *Proc. R. Soc. B* 273, 207–215. (doi:10.1098/rspb.2005.3283)

39. Wenseleers T, Ratnieks FL. 2006 Comparative analysis of worker reproduction and policing in eusocial Hymenoptera supports relatedness theory. *Am. Nat.* 168, E163–E179. (doi:10.1086/508619)

40. Blumstein DT, Möller AP. 2008 Is socality associated with high longevity in North American birds? *Biol. Lett.* 4, 146–148. (doi:10.1098/rsbl.2007.0606)

41. Jetz W, Rubenstein DR. 2011 Environmental uncertainty and the global biogeography of cooperative breeding in birds. *Curr. Biol.* 21, 72–78. (doi:10.1016/j.cub.2010.11.075)

42. Shultz S, Opie C, Atkinson QD. 2011 Stepwise evolution of stable socality in primates. *Nature* 479, 219. (doi:10.1038/nature06061)

43. Gonzalez JCT, Sheldon BC, Tobias JA. 2013 Environmental stability and the evolution of cooperative breeding in hirbills. *Proc. R. Soc. B* 280, 20131297. (doi:10.1098/rspb.2013.1297)

44. Liker A, Freckleton RP, Székely T. 2013 The evolution of sex roles in birds is related to adult sex ratio. *Nat. Commun.* 4, 1587. (doi:10.1038/ncomms2660)

45. Nakagawa S, Freckleton RP. 2011 Model averaging, missing data and multiple imputation: acase study for behavioural ecology. *Behav. Ecol. Sociobiol.* 65, 103–116. (doi:10.1007/s00265-010-1044-7)

46. Clutton-Brock T, Sheldon BC. 2010 Individuals and populations: the role of long-term, individual-based studies of animals in ecology and evolutionary biology. *Trends Ecol. Evol.* 25, 562–573. (doi:10.1016/j.tree.2010.08.002)

47. Lukas D, Clutton-Brock T. 2014 Costs of mating competition limit male lifetime breeding success in polygynous mammals. *Proc. R. Soc. B* 281, 20140418. (doi:10.1098/rspb.2014.0418)

48. Jones KE et al. 2009 PanTHERIA: a species-level database of life history, ecology, and geography of extant and recently extinct mammals. *Ecology* 90, 2648. (doi:10.1890/08-1494.1)

49. Kissling WD, Dalby L, Fløggaard C, Lenoir J, Sandel B, Sandom C, Svenning JC. 2014 Establishing macroecological trait datasets: digitalization, extrapolation, and validation of diet preferences in terrestrial mammals worldwide. *Ecol. Evol.* 4, 2913–2930. (doi:10.1002/ece3.1936)