Energy In-Equivalence in Australian Marsupials: Evidence for Disruption of the Continent’s Mammal Assemblage, or Are Rules Meant to Be Broken?

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Abstract

The energy equivalence rule (EER) is a macroecological hypothesis that posits that total population energy use (PEU) should be independent of species body mass, because population densities and energy metabolisms scale with body mass in a directly inverse manner. However, evidence supporting the EER is equivocal, and the use of basal metabolic rate (BMR) in such studies has been questioned; ecologically-relevant indices like field metabolic rate (FMR) are probably more appropriate. In this regard, Australian marsupials present a novel test for the EER because, unlike eutherians, marsupial BMRs and FMRs scale differently with body mass. Based on either FMR or BMR, Australian marsupial PEU did not obey an EER, and scaled positively with body mass based on ordinary least squares (OLS) regressions. Importantly, the scaling of marsupial population density with body mass had a slope of −0.37, significantly shallower than the expected slope of −0.75, and not directly inverse of body-mass scaling exponents for BMR (0.72) or FMR (0.62). The findings suggest that the EER may not be a causal, universal rule, or that for reasons not yet clear, it is not operating for Australia’s unique native fauna.

Introduction

Identifying mechanistic associations between organism body size (mass), resource use and whole-ecosystem processes is central to predicting how different species and their ecosystems might respond to environmental challenges, or to other factors affecting body size and resource use (e.g. [1]). One theory that considers how organism size and resource needs affect abundance, and ultimately whole ecosystems processes, is the Energy Equivalence Rule (EER; [2,3,4]; also described as the Energetic Equivalence Rule). In short, the EER is a type of size-density relationship that states that total population energy-fluxes by different species should be equivalent, regardless of their respective body masses [2,3,5]. The EER has been used to explain a range of large-scale ecological phenomena, from community structuring to global biodiversity patterns (see [5,6,7,8]), but the validity of some of its underlying features have been questioned, and evidence supporting the idea as a general ‘rule’ is equivocal [2,3,5,6,9,10,11,12,13,14].

The EER was derived from the observation that individual energy requirements (or metabolic rate; kJ d⁻¹) apparently scale with animal body mass raised to a power close to 0.75 (i.e. mass⁰.⁷⁵), whereas the scaling of animal population densities (individuals km⁻²) scale with body mass raised to a power close to −0.75 (i.e. mass⁻⁰.⁷⁵; [15]), i.e. the direct inverse of metabolic rate [2,3]. Consequently, the EER states that whole-population energy fluxes (kJ per unit area) should be the same for differently-sized organisms, because total population energy use (PEU) equals energy turnover (or basal metabolic rate; BMR) multiplied by its density; i.e. [BMR•mass⁰.⁷⁵] • [Density•mass⁻⁰.⁷⁵] = PEU•mass⁰

There is empirical support for [2,3,4,12,19,20] and against [4,6,9,12,21] the EER. One concern is whether BMR is appropriate for deriving PEU, and that EERs should focus on ecologically-relevant indices like field metabolic rate (FMR; [11]); FMRs are typically 2-3 times BMR for mammals [15]. Axiomatically, using BMR may not be problematic provided it scales with the same slope as does FMR (e.g. mass⁰.⁷⁵), which is apparently the case for eutherian mammals [15]. However, comparable scaling of BMR and FMR is not apparent for all mammal groups [15], and one notable exception includes the marsupial fauna of continental Australia.

Australian marsupial BMRs scale with a body-mass exponent of 0.72 [15,22], but their FMRs scale with a body-mass exponent of 0.62 ( [15]; this study). Therefore, Australian marsupials present a novel group for testing the EER, partly because of their divergent body-mass scaling exponents for BMR and FMR, and also because they have largely evolved isolated from the eutherian ecological-analogues on other continents.
Materials and Methods

Density data were collected from published studies \([23,24]\) for \(n = 68\) species of Australian marsupial, spanning three orders of magnitude of body mass (Table S1) that encompassed the full spectrum of extant marsupial sizes. Data for marsupial BMRs were collated for \(n = 52\) species, and FMRs collated for \(n = 37\) species (Table S2 and Table S3). Specifically for FMR, we collated data for adult, non-reproductive (i.e. non-lactating/non-pregnant) animals covering three orders of magnitude (Table S3). When more than one measure of FMR was available for a species (e.g. seasonal studies) we used minimal values, usually representing dry season data (FMRs in other seasons or following rainfall are typically higher). By excluding data on juvenile (still growing) or lactating animals we present the most conservative dataset for marsupial FMRs to date, with the view to present the minimum free-living resource requirements of Australian marsupials generally.

We explored the scaling of marsupial population densities, BMRs, FMRs and PEUs against body mass using ordinary least squares (OLS) regressions on log10-transformed data (normality of respective residuals was tested using Shapiro-Wilk’s test). Log10-PEU was derived by multiplying raw BMR and FMR by density prior to log transformation. Not all species for which we had compiled density information (\(n = 68\)) were represented in the BMR (\(n = 52\)) or FMR (\(n = 37\)) datasets. Therefore, to test whether scaling patterns in the species subsets for BMR OR FMR were representative of slopes derived from each entire dataset. However, for the FMR dataset there were only two species for which estimates of density were not available, making formal density-scaling regressions for marsupials that included data on BMR or FMR were not significantly different from the entire dataset (DensityBMR interaction \(F_{1, 49} = 0.004, P = 0.95\); DensityFMR interaction \(F_{1, 49} = 0.001, P = 0.98\)).

Marsupial BMR (entire dataset) scaled with body mass with an exponent of \(-0.37\), and was significantly different from a slope of \(-0.75\) (Table 1; \(Z = 6.1, P < 0.0001\)). Importantly, the slopes for density-scaling regressions for marsupials that included data on BMR or FMR were not significantly different from the entire dataset (DensityBMR interaction \(F_{1, 49} = 2.1, P = 0.16\); Table 1). Marsupial FMR (entire dataset) scaled with body mass with an exponent of \(0.62\) (Table 1). Notably, the 95% confidence intervals for the exponents of either of the BMR or FMR did not include the reciprocal of the 95% confidence interval for the density scaling exponent; in other words, the scaling exponents of neither BMR nor FMR were directly inverse that for population density (Table 1). Consequently, marsupial population energy use based on measures of BMR (PEU\(_{\text{BMR}}\) scaled with a body mass exponent of \(0.21\) (\(P = 0.036\); Fig. 1). Similarly, marsupial population energy use based on measures of FMR (PEU\(_{\text{FMR}}\) scaled with a body mass exponent of \(0.20\) (\(P = 0.030\); Fig. 1).

Discussion

Australian marsupials do not follow an EER according to the OLS-regressions of PEU based on either BMR or FMR (Table 1). Most importantly, a key feature of the EER is that the body-mass scaling exponents for density and energy turnover (BMR or FMR) should be a direct inverse of one another, yet for both BMR and FMR the scaling exponents did not demonstrate a reciprocal overlap with the scaling exponent for population density (see also \([23]\)). We propose two alternative hypotheses to explain why Australian marsupials do not obey an EER. Firstly, and perhaps most parsimoniously, the EER may not be a general ecological ‘rule’, and as such it is not universal or predictive. Alternatively, the EER may indeed be causal, but for reasons that are not yet clear it is not operating at the continental scale for Australian marsupials. There is evidence to support both of these lines of argument.

That the EER formulated by Damuth \([2,3]\) is not in fact a ‘rule’ is somewhat supported by our data, and is comparable with other studies that refute EERs across a range of species, communities and trophic levels (e.g. \([6]\)). As such, global-scale EERs (sensu White et al. \([5]\)) may be emergent artefacts that are not be driven by bottom-up, local EERs (see also \([6,15]\)), but firm conclusions would require that all sympatric species be included in any analyses. Furthermore, it is worth noting that the composition of Australia’s extant marsupial fauna comprises of mainly small carnivores/omnivores, but medium-large herbivores (mainly the wombats and kangaroos). It is well-known that diet influences BMR and FMR (e.g. see \([15,18,25,26]\) and references therein). Nonetheless, local- and taxon-scale species-density patterns are rarely independent of body size \([5,6,14,27]\), but the mechanisms explaining such outcomes are unclear. Further, there is considerable heterogeneity in the scaling exponent of animal BMRs at least, with around 50% of orders displaying slopes for body-mass: BMR scaling that deviate from the expected (according to EER) slope 0.75 (\([8]\), see also \([28,29]\)). Therefore, it appears unrealistic to assume that global-EERs based on a single body-mass: density scaling exponent (e.g. 0.75) should apply locally. Instead, EERs should be tailored to the species-specific energy requirements and interactions for all species in a given area. Consequently, outright dismissal of an EER for Australian marsupials may be premature because we could not include information for all sympatric species, and particularly for the numerous introduced domestic and feral eutherians that have helped to transform Australia’s biomes since their introductions with Europeans some 200 years ago.

Dramatic changes to Australia’s landscapes since European arrival have precipitated major declines, even to extinction, of numerous small-medium sized marsupials, along with increases in the population sizes of the largest extant marsupials, particularly the grazing kangaroos \([30]\). Widespread land clearing and the establishment of permanent water sources for grazing ruminants, mainly sheep and cattle, have contributed to the declines of Australia’s small-medium sized marsupials, in addition to supporting the proliferation of some larger, grazing marsupials such as kangaroos \([30,31]\). Additionally, there has been widespread control and exclusion of Australia’s largest established predators, the mainland dingo/wild dog. This has apparently released some larger marsupial species from predation pressures (e.g. kangaroos), whilst concomitantly enabling introduced mesopredators (foxes and cats) to target small-medium marsupials, possibly driving down their population numbers or restricting them to sub-optimal refuges where their abundances are lower than might expected without such heavy predation pressures.
Moreover, the control of large predators and the spread of grasslands and artificial water sources (e.g. dams, tanks, bores) have supported extensive infiltrations of introduced herbivores like rabbits, goats and camel, as well as omnivores like pigs, which have further transformed Australia’s biomes and species compositions. Therefore, it is not wholly unexpected that the scaling of Australian marsupial density: body mass might differ from that of mammals generally, and particularly from that of eutherians on other continents [2,3,6]. Eutherians generally have higher energy requirements than marsupials [15,22], and the introduction of eutherians to Australia in high numbers as free-ranging domestic stock, and as extensive feral populations, could act to counter-balance, or even over-balance energy fluxes through Australian ecosystems. As such,

| Parameter | $a$ | $b$ | $R^2$ | $p$ |
|-----------|-----|-----|-------|-----|
| Density (n = 68) | $3.19 \pm 0.20$ | $-0.37 \pm 0.06$ | $0.394$ | $<0.0001$ |
| BMR (n = 52) | $0.169 \pm 0.051$ | $0.72 \pm 0.02$ | $0.967$ | $<0.0001$ |
| FMR (n = 37) | $0.877 \pm 0.057$ | $0.62 \pm 0.018$ | $0.972$ | $<0.0001$ |

Regressions were performed separately for all species for which density, BMR and FMR were available, in addition to those for which overlapping data were available (Note: overlap data for density, BMR and FMR are identified by respective subscripts, e.g. Density/BMR = density data for which BMR is also available; values in parentheses are 95% confidence limit ranges).

Table 1. Scaling (OLS regressions on log10-transformed data) of Australian marsupial population density (number of individuals km$^{-2}$), basal (BMR) and field (FMR) metabolic rate (kJ d$^{-1}$) with body mass (g).

doi:10.1371/journal.pone.0057449.t001

Figure 1. Scaling (OLS regressions on log10-transformed data) of Australian marsupial population energy use (PEU; kJ d$^{-1}$ km$^{-2}$) with body mass, based on (A) basal metabolic rate (BMR; kJ d$^{-1}$) and (B) field metabolic rate (FMR; kJ d$^{-1}$).

doi:10.1371/journal.pone.0057449.g001
Australia presents a unique opportunity to test EERs along gradients of disturbance at local, regional and continental scales. Unfortunately, there are presently too few data on the FMRs of Australian native and non-native eutherians to adequately compare the contributions of Australia’s marsupials with that of native and introduced eutherians to total-ecosystem energy fluxes.

Importantly, Australia’s extant marsupials are not representative of the continents’ pre-European assemblages, and numerous small-medium sized marsupials are now extinct [35,36]. Consequently, evaluating the role of phylogeny in the PEU patterns for Australian marsupials is complicated by the likely influence of phylogeny (and body mass) on the extinction-risk of Australia’s small-medium sized marsupials over the last 200 years, which may or may not be cross-correlated with their metabolic physiology (for further discussion on the risks of misinterpreting phylogenetic influences see [18,37]). For example, torpor and hibernation have apparently mitigated extinction risks for many small mammals, but larger mammals that maintain homeothermic body temperatures may suffer higher extinction rates, presumably because of their need to sustain high and relatively constant energy metabolism [38]; although to some degree large animals may ameliorate risks via migration to avoid climatic or other pressures. Phylogeny is therefore important, as has been demonstrated for mammalian BMRs across a range of taxa including marsupials [25,28], but other factors likely contribute to the patterns we observe [18,25,28]. Indeed, White et al. [29] has recommended that more parameter-rich models (that include phylogeny) are needed to fully appreciate the ecological patterns associated with mammalian BMR: body mass allometries, and we suggest that these ideas ought to extend to include FMR because that is the physiological level at which species operate ecologically [11]. Nonetheless, the specific hypothesis that we are testing here (that the EER holds true for extant Australian marsupials) does not directly concern phylogeny in that we are interested in present-day ecological patterns, rather than the evolution of energy metabolism or population density/energy use per se. Perhaps after sufficient data are collected for the energy metabolisms and population densities of Australia’s non-marsupial mammals, and indeed birds and reptiles, might we fully appreciate the extent to which phylogeny contributes to the macroecology of population-energy fluxes for Australia’s extant fauna and their ecosystems. Regardless, our continental-wide examination of marsupial density: body mass scaling reveals that Australia’s extant marsupials may be experiencing profound ecological imbalances, the consequences of which are probably still unfolding.

Acknowledgments

Sincere thanks to Professor Kris French, Dr Phil Byrne and Dr Terry O’Dwyer for thoughtful discussions and feedback on earlier drafts of this manuscript. Thanks also to Professors Chris Johnson and Don Bradshaw for providing raw data to assist with our collation of density and FMRs, along with Professor Mark Westoby and Dr Matt Symonds for discussion on regressions and phylogeny, and to Dr Matt Symonds and two anonymous reviewers for their comments and constructive criticism of earlier versions of this manuscript.

Supporting Information

Table S1 Australian marsupial population density (individual km$^{-2}$ and body mass (g)).

Table S2 Australian marsupial basal metabolic rate (BMR) and body mass (g).

Table S3 Australian marsupial field metabolic rate (FMR) and body mass (g).

Author Contributions

Conceived and designed the experiments: AM. Performed the experiments: AM CD. Analyzed the data: AM CD DM MC. Contributed reagents/materials/analysis tools: AM MC. Wrote the paper: AM MC.

References

1. Ozgul A, Tuljapurkar S, Benton TG, Pemberton JM, Clutton-Brock TH, et al. (2009) The dynamics of phenotypic change and the shrinking sheep of St. Kilda. Science 325: 464–467.
2. Damuth J (1981) Population density and body size in mammals. Nature 280.
3. Damuth J [1987] Interspecific allometry of population density in mammals and other animals: the independence of body mass and population energy-use. Biological Journal of the Linnean Society 31: 193–246.
4. Neve R, Read AF, Greenwood JJ, Harvey PH (1994) The relationship between abundance and body size in British birds. Nature 331: 312–313.
5. White EP, Ernst SKM, Kerrthof AJ, Enquist BJ (2007) Relationships between body size and abundance in ecology. Trends in Ecology & Evolution 22: 323–330.
6. Isaac JL, Storch D, Carbone C (2011) Taxonomic variation in size–density relationships challenges the notion of energy equivalence. Biological Letters 7: 615–618.
7. Carbone C, Rowcliffe JM, Cowlishaw G, Isaac Nick JB (2007) The scaling of abundance in consumers and their resources: Implications for the energy equivalence rule. The American Naturalist 170: 479–484.
8. Isaac NJB, Carbone C (2010) Why are metabolic scaling exponents so controversial? Quantifying variance and testing hypotheses. Ecology Letters 13: 728–735.
9. Marquet PA, Navarrete SA, Castilla JC (1995) Body size, population density, and the Energetic Equivalence Rule. Journal of Animal Ecology 64: 325–332.
10. Gaston KJ, Blackburn TM (2000) Patterns and Process in Macroecology: Blackwell Scientific, Oxford, UK.
11. Carbone C, Petorelli N (2009) Testing relationships between energy and vertebrate abundance. International Journal of Ecology 2009.
12. Ackerman JL, Bellwood DR, Brown JH (2004) The contribution of small individuals to density-body size relationships: examination of energetic equivalence in reef fishes. Oecologia 139: 568–571.
13. Ernst SKM, Enquist BJ, Brown JH, Charnov EL, Gillooly JF, et al. (2003) Thermodynamic and metabolic effects on the scaling of production and population energy use. Ecology Letters 6: 990–995.
14. Russo SE, Robinson SK, Terborgh J (2000) Size-abundance relationships in an American bird community: Implications for the Energetic Equivalence Rule. The American Naturalist 161: 267–283.
15. Capellini L, Venditti C, Burton RA (2010) Phylogeny and metabolic scaling in mammals. Ecology 91: 2783–2793.
16. White CR, Seymour RS (2003) Allometric scaling of mammalian metabolism. Journal of Experimental Biology 208: 1611–1619.
17. White CR, Seymour RS (2004) Does basal metabolic rate contain a useful signal? Mammalian BMR allometry and correlations with a selection of physiological, ecological, and life-history variables. Physiological and Biochemical Zoology 77: 929–941.
18. McNab BK (2008) An analysis of the factors that influence the level and scaling of mammalian BMR. Comparative Biochemistry and Physiology Part A: Molecular Integrative & Physiology 151: 5–28.
19. Brown JH (2005) Ecological food webs: High-quality data facilitate theoretical unification. Proceedings of the National Academy of Sciences 100: 1467–1468.
20. Allen AP (2002) Global biodiversity, biochemical kinetics, and the Energetic Equivalence Rule. Science 297: 1545–1548.
21. Blackburn TM, Gaston KJ (1997) A critical assessment of the form of the interspecific relationship between abundance and body size in animals. Journal of Animal Ecology 66: 233–249.
22. Withers PC, Cooper CE, Larcombe AN (2006) Environmental correlates of physiological variables in marsupials. Physiological and Biochemical Zoology 79: 437–453.
23. Johnson CN (1999) Relationships between body size and population density of animals: The problem of the scaling of study area in relation to body size. Oikos 85: 563–569.
24. Strahan R, van Dyck S, editors (2008) Mammals of Australia: CSIRO Publishing.
25. Lovegrove Barry G (2000) The zoogeography of mammalian basal metabolic rate. The American Naturalist 156: 201–219.
26. Anderson KJ, Jetz W (2005) The broad-scale ecology of energy expenditure of endotherms. Ecology Letters 8: 310–318.
27. Gotgreave P (1993) The relationship between body size and population abundance in animals. Trends in Ecology and Evolution 8: 244–248.
28. White CR (2011) Allometric estimation of metabolic rates in animals. Comparative Biochemistry and Physiology Part A: Molecular & Integrative Physiology 158: 346–357.
29. White CR, Frappell PB, Chown SL (2012) An information-theoretic approach to evaluating the size and temperature dependence of metabolic rate. Proceedings of the Royal Society B: Biological Sciences 279: 3616–3621.
30. Johnson CN (2006) Australia’s Mammal Extinctions: A 50,000-year History: Cambridge University Press, UK.
31. Dawson TJ (2012) Kangaroos. Collingwood: CSIRO Publishing.
32. Letnic M, Ritchie EG, Dickman CR (2012) Top predators as biodiversity regulators: the dingo Canis lupus dingo as a case study. Biological Reviews 87: 390–413.
33. Newsome AE, Catling PC, Cooke BD, Smyth R (2001) Two ecological universes separated by the Dingo Barrier Fence in semi-arid Australia: interactions between landscapes, herbivory and carnivory, with and without dingoes. The Rangeland Journal 23: 71–90.
34. Letnic M, Koch F, Gordon C, Crowther MS, Dickman CR (2009) Keystone effects of an alien top-predator stem extinctions of native mammals. Proceedings of the Royal Society B: Biological Sciences 276: 3249–3256.
35. Carrill M, Bromham L (2001) Body size and risks of extinction in Australian mammals. Conservation Biology 15: 1433–1440.
36. Johnson CN, Isaac JL (2009) Body mass and extinction risk in Australian marsupials: The ‘Critical Weight Range’ revisited. Austral Ecology 34: 35–40.
37. Westoby M, Leishman MR, Lord JM (1995) On misinterpreting the ‘phylogenetic correction’. Journal of Animal Ecology 83: 531–534.
38. Geiser F, Turbill C (2009) Hibernation and daily torpor minimize mammalian extinctions. Naturwissenschaften 96: 1235–1240.