



**The covariance between metabolic rate and behaviour
varies across behaviours and thermal types: meta-analytic
insights**

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3 **1 The covariance between metabolic rate and behaviour varies across behaviours and thermal types:**
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5 **2 meta-analytic insights**
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51 19 **Authorship statement:** KJM, NJD and SN conceived of the study. KJM performed the literature review
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53 20 and extracted data estimates. SN assisted with data extractions and performed the meta-analysis. KJM
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55 21 and SN co-wrote the paper. All co-authors contributed to revisions.
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For Review Only

Abstract

Energy metabolism has received much attention as a potential driver of repeatable among-individual differences in behaviour (animal personality). Several factors have been hypothesized to mediate this relationship. We performed a meta-analysis of >70 studies comprised of >8000 individuals reporting relationships between measures of maintenance metabolic rates (i.e., basal metabolic rate, resting metabolic rate, and standard metabolic rate) and behaviour. We evaluated support for three hypothesized mediators: 1) type of behaviour, 2) opportunities for energy re-allocation, and 3) magnitude of energetic constraints. Relationships between measures of maintenance metabolic rate (MR) and behaviour are predicted to be strongest for behaviours with strong consequences for energy turnover (acquisition or expenditure). Consistent with this, we found that behaviours with known consequences for energy gain (e.g., foraging, dominance, boldness) or expenditure (e.g., maximum sprint speed, sustained running speed, maximum distance travelled, etc.) had strong positive correlations with MR, while behaviours with putatively weak and/or inconsistent associations with net energy gain or loss (e.g., exploration, activity, sociability) were not correlated with MR. Greater opportunities for energy reallocation are predicted to weaken relationships between MR and behaviour by creating alternative pathways to balance energy budgets. We tested this by contrasting relationships between MR and behaviour in ectotherms versus endotherms, as thermoregulation in endotherms creates additional opportunities for energy reallocation compared with ectotherms. As predicted, the relationship between behaviour and metabolic rate (MR) was stronger in ectotherms compared with endotherms. However, statistical analyses of heterogeneity among effect sizes from different species did not support energy re-allocation as the main driver of these differences. Finally, we tested whether conditions where animals face greater constraints in meeting their energy budgets (e.g., field versus lab, breeding versus non-breeding) increased the strength of relationship between MR and behaviour. We found that the relationship between MR and behaviour was unaffected by either of these modifiers. This meta-analysis

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3 49 provides two key insights. First, we observed positive relationships of similar magnitude between MR
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5 50 and behaviours that bring in net energy, and behaviours that cost net energy. This result is only
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7 51 consistent with a performance energy management model. Given that the studies included in our meta-
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9 52 analysis represent a wide range of taxa, this suggests that the performance model may be the most
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11 53 common model in general. Second, we found that behaviours with putatively weak or inconsistent
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13 54 consequences for net energy gain or expenditure (exploration, activity, sociability) show no relationship
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15 55 with MR. This provides the first systematic demonstration of the central importance of the ecological
16
17 56 function of traits in mediating relationships between MR and behaviour.
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22 57 **Keywords:** among-individual differences, animal personality, basal metabolic rate, resting metabolic
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24 58 rate, routine metabolic rate, standard metabolic rate, energetic constraints, energy management
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83 I. Introduction

84 The last decade has seen a surge of interest in understanding the causes and consequences of among-
85 individual differences in behaviour (i.e. animal personality). The majority of adaptive explanations for
86 animal personality are based on state-dependence (Dingemanse & Wolf, 2010; Sih *et al.*, 2015; Wolf &
87 Weissing, 2010), where state is broadly defined as any factor that influences the payoffs of a given
88 behavioural action (Houston & McNamara, 1999). State variables related to energy metabolism in
89 particular have been the subject of much attention in this burgeoning area of research (e.g., Biro *et al.*,
90 2010; Careau & Garland, 2012; Careau *et al.*, 2008; Houston, 2010; Mathot & Dall, 2013; Mathot &
91 Dingemanse, 2015; Wolf & McNamara, 2012), and the number of empirical studies investigating links
92 between metabolism and behaviour has grown dramatically in the last several years (reviewed in Biro &
93 Stamps, 2010; Careau & Garland, 2012; Mathot & Dingemanse, 2015; Niemelä & Dingemanse, 2018;
94 Royauté *et al.*, 2018).

95 Several papers have called attention to the fact that relationships between measures of maintenance
96 MR (i.e., basal metabolic rate, resting metabolic rate, and standard metabolic rate) and behaviour are
97 likely mediated by interacting factors (Biro & Stamps, 2010; Careau & Garland, 2012; Careau *et al.*, 2008;
98 Killen *et al.*, 2013; Mathot & Dingemanse, 2015; Speakman *et al.*, 2004). For example, the energy
99 management model of an organism (i.e., how maintenance MR covaries with total energy expenditure)
100 (Biro & Stamps, 2010; Careau & Garland, 2012; Careau *et al.*, 2008; Mathot & Dingemanse, 2015;
101 Speakman *et al.*, 2004), the type of behaviour (Biro & Stamps, 2010; Mathot & Dingemanse, 2015), the
102 extent to which there are alternative allocation opportunities (Careau & Garland, 2012; Mathot &
103 Dingemanse, 2015), and environmental stressors (Biro & Stamps, 2010; Killen *et al.*, 2013) are all
104 expected to shape the relationships between MR and behaviour.

105 Here, we performed a meta-analysis of existing studies to assess the overall support for a relationship
106 between MR and behaviour. In addition, we evaluated the importance of several of the factors that have

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3 107 been proposed to mediate these relationships: 1) the type of behaviour, 2) the opportunities for energy
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5 108 re-allocation, and 3) the degree of energetic stress. Below, we detail the rationale and predictions for
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7 109 each of these proposed mediators.
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11 **(1) Does covariation between MR and behaviour depend on the type of behaviour?**
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14 111 Assuming that variation in MR reflects variation in energy requirements, then to balance their energy
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16 112 budgets, variation in MR can be logically expected to be associated with variation in either behaviours
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18 113 that bring in net energy (e.g. foraging, food defense, foraging boldness), or cost net energy (e.g.
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20 114 movement, mate defense, etc.) (Biro & Stamps, 2010; Careau & Garland, 2012; Careau *et al.*, 2008;
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22 115 Mathot & Dingemanse, 2015; Speakman *et al.*, 2004). However, variation in MR inferred from measures
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24 116 of oxygen consumption may not reflect true differences in energetic requirements if organisms differ in
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26 117 their mitochondrial efficiency (i.e., the amount of adenosine triphosphate, ATP, generated per molecule
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28 118 of oxygen consumed) (Salin *et al.*, 2015). Although there is evidence for among-individual differences in
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30 119 mitochondrial efficiency (reviewed in Salin *et al.*, 2015), we do not address this here given that paucity of
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32 120 studies simultaneously recording among-individual differences in MR, mitochondrial efficiency, and
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34 121 behaviour.
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39 122 Assuming variation in oxygen consumption does reflect variation in energy requirements, the direction
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41 123 of the relationship between MR and behaviours that cost net energy depends on the energy
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43 124 management model of the organism. Three energy management models have been described which
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45 125 reflect three distinct relationships between MR and daily energy expenditure (DEE). Under the allocation
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47 126 model (also called the compensation model), DEE does not vary as a function of MR (Careau & Garland,
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49 127 2012; Careau *et al.*, 2008; Mathot & Dingemanse, 2015). Thus, logically, variation in MR is not predicted
50
51 128 to be associated with variation in behaviours that bring in net energy (e.g., foraging) (Mathot &
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53 129 Dingemanse, 2015). However, because organisms work with a fixed energy budget, high MR means a
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3 130 relatively smaller fraction of energy available for energetically costly behaviours (e.g., sustained sprint
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5 131 speed) (Careau & Garland, 2012; Careau *et al.*, 2008; Mathot & Dingemanse, 2015). Under the
6
7 132 independent model (also referred to as compensation model), the energy devoted to activity is
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9 133 independent of MR, i.e., the amount of energy expended on activities above basic maintenance does not
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11 134 vary with MR (Careau & Garland, 2012; Mathot & Dingemanse, 2015). However, this still has the
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13 135 consequence that total energy required, DEE, does increase with increasing MR, and therefore, a positive
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15 136 relationship is predicted between MR and behaviours such as foraging (Mathot & Dingemanse, 2015).
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17 137 Finally, under the performance model (also referred to as increased intake), the capacity of an organism
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19 138 to bring in energy is positively correlated with MR (Careau & Garland, 2012; Careau *et al.*, 2008; Mathot
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21 139 & Dingemanse, 2015). Thus, organisms adopting a performance model would be predicted to have
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23 140 higher expressions of both behaviours that bring in net energy, and behaviours that cost net energy.
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28 141 Few studies directly assess the relationship between MR and DEE (but see Portugal *et al.*, 2016), thus the
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30 142 energy management model is typically an untested assumption (Mathot & Dingemanse, 2015). Assuming
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32 143 that existing studies comprise a mix of study species that adopting each of the three potential energy
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34 144 management models, we would predict that there should generally be stronger support for a positive
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36 145 relationship between behaviours that are associated with increased intake rate or increased access to
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38 146 resources (e.g. foraging, foraging boldness, foraging dominance) compared with behaviours that cost net
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40 147 energy (e.g. courtship/mating, stress responses and performance measures such as maximum sprint
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42 148 speed). This is because two of the three energy management models predict a positive relationship
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44 149 between MR and resource acquisition (independent and performance models) while the third predicts
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46 150 no relationship (allocation model). In contrast, only one of the three energy management models
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48 151 predicts a positive relationship between MR and behaviours that consume net energy (performance
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50 152 model), the others predict either no relationship (independent model), or a negative relationship
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52 153 (allocation model).
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3 154 Notably, many studies investigating relationships between behaviour and metabolic rate quantify
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5 155 behaviours for which the functional significance in terms of net energy gain/loss is unclear. For example,
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7 156 exploration behaviour in novel environmental conditions and general activity, are two commonly
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9
10 157 measured behaviours in studies of repeatable among-individual variation. Activity measures the speed
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12 158 with which an organism moves through a familiar environment, and exploration is typically defined as
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14 159 the speed with which an individual moves through a novel environment (Réale *et al.*, 2007). Although the
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16 160 expression of these behaviours undoubtedly involve energetic costs, they are often also assumed to
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18 161 determine the rate with which organisms encounter resources. Thus, depending on the intensity of
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20 162 activity and exploration and the extent to which they determine encounter rates with food, a high
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22 163 expression of activity or exploration may be associated with large net energy costs, large net energy
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24 164 gains, or anything in between (Biro & Stamps, 2010; Careau & Garland, 2012; Carter *et al.*, 2013). Thus,
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26 165 relationships between activity or exploration and MR will likely differ for different species, or under
27
28 166 different ecological contexts, therefore, overall relationships between MR and exploration or activity are
29
30 167 predicted to be weak or non-existent (Careau & Garland, 2012; Carter *et al.*, 2013; Mathot &
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32 168 Dingemanse, 2015). Similarly, it is unclear how sociability (the propensity to affiliate with congeners)
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34 169 should influence net energy gain or loss. On the one hand, being more strongly affiliated with congeners
35
36 170 may decrease expected intake through resource competition. On the other hand, it may allow for
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38 171 increased feeding rates if sociable animals can reduce their relative investment in vigilance due to
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40 172 dilution or group vigilance effects. Thus, we do not predict a consistent net effect of sociability on energy
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42 173 gain or energy loss, and therefore, predict either no or weak relationships between sociability and MR.
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49 174 ***(2) Do opportunities for alternative energy allocation decisions weaken relationships between MR and***
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51 175 ***behaviour?***
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54 176 Assuming that the energy management model of a study system is known, we can predict how metabolic
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56 177 rate should covary with behaviours that facilitate net energy gain versus net energy loss *in general*.
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3 178 However, animals are able to allocate the energy available for such behaviours to a variety of different
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5 179 behaviours (e.g. foraging, foraging boldness, food defense). The ability to allocate among multiple
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7 180 behaviours means that even if metabolic rate is predicted to covary positively with behaviours that bring
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9 181 in net energy, it need not covary positively with *all* behaviours that bring in net energy (Mathot &
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11 182 Dingemanse, 2015). Opportunities to reallocate energy across different behaviours that have the same
12
13 183 net consequence for energy balance means that associations between MR and specific behaviours may
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15 184 not match the relationship between MR and behaviours in general. Consider the following, very
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17 185 simplified, example. In a system with an independent energy management model, higher MR is
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19 186 predicted to be associated with greater expression of behaviours that bring in net energy (see Figure 1,
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21 187 Mathot & Dingemanse, 2015). Imagine that an organism has the ability to allocate between just two
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23 188 behaviours that bring in net energy; foraging boldness (green) or resource defense (blue). Note that
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25 189 resource defense is only predicted to occur when organism experience net benefits from defense (i.e.,
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27 190 the resource is economically defensible), thus, despite costs associated with territory defense, the
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29 191 behaviour would still be associated with net energetic gains (Davies *et al.*, 2012). In this simplified
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31 192 scenario, alternative allocation decisions between these two behaviours may create scenarios where
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33 193 both behaviours show the predicted positive correlation with MR (Figure 1a), or only one behaviour
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35 194 shows the predicted relationship (i.e., resource defense, Figures 1b-c). Note that it is even possible for
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37 195 specific behaviours (i.e., foraging boldness) to show relationships with MR that are opposite to the
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39 196 predicted relationship for a given energy management model (in this case, the independent model), so
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41 197 long as other behaviours (i.e., resource defense) are sufficiently upregulated to offset this (Figure 1c).
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49 198 We test the idea that a greater number of alternative options for energy allocation will weaken the
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51 199 strength of relationships between MR and behaviour by contrasting endotherms with ectotherms.
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53 200 Thermoregulation introduces additional allocation opportunities for endotherms, for example, because
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55 201 heat produced through activity can be used to offset thermoregulation costs (Careau & Garland, 2012;
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202 Careau *et al.*, 2014; Humphries & Careau, 2011). Thus, we predict that the relationships between MR and
203 behaviour will be weaker in endotherms compared with ectotherms.

204 **(3) Does energetic stress promote stronger relationships between metabolic rate and behaviour?**

205 Environmental stressors can alter the relationship between MR and behaviour, though the direction of
206 the effect may differ for different types of stressors (Killen *et al.*, 2013). However, the rationale for
207 relationships between MR and behaviour is based in part on the assumption that balancing one's energy
208 budget is challenging. Environmental stressors that create greater challenges for organisms in terms of
209 balancing their energy budgets (e.g., reduced food availability, breeding), are expected to strengthen the
210 relationships between MR and behaviour (Mathot & Dingemanse, 2015; Ricklefs & Wikelski, 2002). We
211 test this prediction in two ways: 1) by testing for an effect of reproductive status (reproductive versus
212 non-reproductive), and 2) by testing for an effect of captivity (free-living versus wild-caught captive
213 animal versus lab-reared captive animal). We assume that animals face greater challenges in balancing
214 their energy budgets during reproduction due to increased energetic costs (e.g., production of offspring,
215 parental care). For animals that engage in parental care, reproduction may also decrease time available
216 for self-feeding, and thereby further challenge animals to balance their energy budgets. Consequently,
217 we predicted that the strength of the relationship between MR and behaviour would be greater in
218 reproductive versus non-reproductive animals. We also assume that free-living animals face greater
219 challenges in balancing their energy budgets compared with captive animals, because captive animals
220 are typically provided with predictable, *ad libitum*, access to food. Therefore, we predicted that
221 relationships between MR and behaviour would be stronger in free-living animals compared with lab-
222 reared animals. We also distinguished wild-caught animals that were tested in captivity, as they may be
223 expected to be intermediate to free-living and lab-reared animals depending on the time-scale across
224 which animals adjust to *ad-libitum* food conditions in captivity.

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56 226 **II. Methods**7
8 227 **(1) Data collection and inclusion/exclusion criteria**
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11 228 We followed the steps outlined in the Preferred Reporting Items for Systematic Reviews and Meta-
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13 229 Analyses (PRISMA) protocol (Moher *et al.*, 2009) for our meta-analysis as closely as possible, as
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15 230 recommended by Nakagawa and Poulin (2012). We performed a literature search using two online
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17 231 databases; *Scopus* (subject area: Life Sciences) and Web of Science. We included all available years up to
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19 232 August 2016 (when the search was last updated). We had both behavioural and metabolism search
20
21 233 terms. The behavioural search terms used were: behavio*, “coping strategy”, “coping style”, personality,
22
23 234 and temperament. The metabolism search terms were: “basal metabolic rate”, BMR, “resting metabolic
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25 235 rate”, RMR, “standard metabolic rate”, SMR and “routine metabolic rate”. We searched for articles
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27 236 including these terms in the “Topic” field. Articles had to include at least one of the behavioural search
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29 237 terms and one of the metabolism search terms. Our meta-analysis was focused on non-human animals
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31 238 thus, to eliminate human studies, we additionally excluded articles that had any of the following terms in
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33 239 the Topic field: child*, infant, baby, patient, women, men, student, person, elderly, boy or girl.
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39 240 We performed the initial literature search on 30 June 2014, and later updated the search on 24 August
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41 241 2016. The second search was restricted to articles published after 2013 to reduce the number of
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43 242 references duplicated from the initial search. These literature searches produced a total of 2614 unique
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45 243 references. These references were screened by reading the titles and abstract to assess their relevance
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47 244 to the meta-analyses. We selected studies according the following five criteria:
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51 245 (1) The study had to include one of the following measures of metabolic rate: resting metabolic rate,
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53 246 standard metabolic rate, basal metabolic rate, routine metabolic rate. We excluded studies that only
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55 247 reported metabolic rates in active animals (except in fish, because water must flow over the gills for fish
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3 248 to respire), such as summit metabolic rate, daily energy expenditure, sustained metabolic rate, etc. This
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5 249 was to reduce the extent to which the measured MR was confounded with behaviour.
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8 250 (2) The behaviour had to be measured outside of the MR measurements. For example, activity during
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10 251 MR was not considered a valid behaviour. Again, this was to avoid scenarios where the MR measure and
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12 252 behaviour were confounded.
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16 253 (3) The study had to present a correlation between the appropriate measure of MR and a behaviour, or
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18 254 present statistics that allowed for the correlation to be estimated indirectly. Thus, studies which only
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20 255 presented categorical data on MR or behaviour were not included with the exception of studies
21
22 256 investigating links between dominance and MR. This is because dominance is not an individual trait, but
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24 257 a relative trait, thus paired comparisons of MR rate in relation to dominance status (categorical:
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26 258 dominant or subordinate) were also included.
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30 259 (4) The correlation provided had to be either a raw (un-partitioned) phenotypic correlation or an among-
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32 260 individual correlation. We did not consider within-individual correlations (not the targeted hierarchical
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34 261 level) or genetic correlations. Although genetic correlations represent the relevant hierarchical level (i.e.,
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36 262 among-individual), there were insufficient studies providing such estimates (N = 2 studies, Careau *et al.*,
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38 263 2011; Mathot *et al.*, 2013) to allow us to compare the effects sizes of genetic correlations with either
39
40 264 among-individual or un-partitioned phenotypic correlations. One of these studies was nonetheless
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42 265 included in the meta-analysis as it additionally provided estimates of raw un-partitioned phenotypic
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44 266 correlations (Careau *et al.*, 2011). One study (Gifford *et al.*, 2014) presented both among-individual and
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46 267 un-partitioned phenotypic correlations for two behavioural traits (foraging and exploration). Both types
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48 268 of correlations were included in the meta-analysis and coded appropriately. The resultant non-
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50 269 independence was accounted for by additionally assigning a group ID to estimates derived from the
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52 270 same sample of individuals.
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3 271 (5) The study had to provide sufficient information to allow for sample size, estimated effect size via
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5 272 descriptive or inferential statistics, and uncertainty to be extracted. In some cases, the study did not
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7 273 directly report the relevant information, but the information could be extracted from data provided in
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9 274 the supplementary material, from the published data set, or from data presented in figures.
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13 275 These selection criteria resulted in a list of 71 papers and 5 published datasets that were appropriate for
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15 276 our meta-analysis (ESM Table S1). Studies that did not fulfill our selection criteria are listed in the
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17 277 electronic supplementary material (ESM Table S2) along with the reason for their exclusion. When the
18
19 278 relevant data was presented in figures, we extracted the data using WebPlotDigitizer 3.8
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21
22 279 (<http://arohatgi.info/WebPlotDigitizer/>). The 76 sources (71 articles and 5 published data sets) for
23
24 280 relevant effect size estimates produced a total of 163 estimates from 48 species (Table 1). The full
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26 281 PRISMA flow chart is provided in Figure 2.
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28
29 282 Our meta-analysis was intended to focus on among-individual correlations between MR and behaviour.
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31 283 We nonetheless included studies that reported both among-individual correlations and raw (i.e., un-
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33 284 partitioned phenotypic) correlations. Raw correlations represent a mix of within- and among-individual
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35 285 correlations and may not accurately reflect the correlation structure at the level of interest (in our case,
36
37 286 among-individual) when within- and among-individual correlations are qualitatively different (e.g.,
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39 287 positive versus negative) (Dingemanse *et al.*, 2012). We tested whether estimates derived from among-
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41 288 individual (N = 41 estimates) versus phenotypic correlations (N = 122 estimates) differed and found no
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43 289 support for a difference. Importantly, the estimated difference was close to zero (the contrast between
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45 290 the two: $\beta_{[\text{difference in } Z_r]} = 0.003$, 95% confidence interval, CI = [-0.165, 0.170]; see the section '*Meta-*
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47 291 *analysis and meta-regression analysis*' below for the details of the statistical model). This suggests that in
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49
50 292 our data set, within- and among-individual correlations are quantitatively and qualitatively similar,
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52 293 otherwise, estimated phenotypic correlations would not be identical to estimated among-individual
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54 294 correlations.
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3 295 A previous meta-analysis found that within-individual correlations between state and behaviour were
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5 296 significantly weaker than among-individual correlations (Niemelä & Dingemanse, 2018). However, when
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7 297 considering only correlations between MR and behaviour, the contrast between among- and within-
8
9 298 individual correlations were not significantly different (P. Niemelä, personal comment). Further, based
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11 299 on the strong support for lack of difference between phenotypic and among-individual correlations in
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13 300 the present dataset, we did not treat among-individual correlations and un-partitioned phenotypic
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15 301 correlations separately in any subsequent analyses.
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19 302 **(2) Data coding and calculation of effect sizes**

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23 303 We collected and transformed relevant study results into the standardised effect statistic, Fisher's z-
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25 304 transformation statistic (Z_r); Z_r (a transformation of correlation) was chosen because we were interested
26
27 305 in the relationship between two variables, metabolism and behaviour (c.f. Nakagawa *et al.*, 2017).
28
29 306 Behaviours were grouped into nine broad categories and were always coded such that higher values
30
31 307 represented higher expression of behaviours associated with greater energy expenditure and/or
32
33 308 acquisition (see ESM Table S3 for behaviours and definitions). Thus, effect size estimates obtained for
34
35 309 relationships between MR and behaviours such as latency scores (e.g., latency to resume feeding) were
36
37 310 multiplied by -1 so that positive estimates indicated that higher MR was associated with higher
38
39 311 expression of behaviours that bring in net energy (resuming feeding). For behaviours that we presumed
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41 312 to have weak or inconsistent relationships with energy turnover (activity, exploration, sociability), data
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43 313 were coded such that higher values indicated higher expression of those traits.
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48 314 Following recommendations from Noble *et al.* (2017) we preferentially collected descriptive statistics
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50 315 (e.g. correlations, group means, standard deviations and standard errors) over inferential statistics (t , F ,
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52 316 χ^2 , U and p). Both descriptive and inferential statistics were transformed using formulas found in the
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54 317 following references (Krishnamoorthy, 2006; Lipsey & Wilson, 2001; Nakagawa & Cuthill, 2007) and using
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3 318 an effect size determination program (Wilson, 2001). Note that we only retained effect sizes from
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5 319 inferential statistics when directions could be determined (e.g., F , χ^2 , and p do not have directional
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7 320 information), and directions of effect sizes were adjusted in a way that all positive effect sizes meant
8
9 321 strong correlations between metabolism and behaviour that either bring in or consume net energy (see
10
11 322 above). For the calculation of Zr 's sampling variance, we use the number of independent subjects (i.e.
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13 323 $1/(i - 3)$; i = the number of subjects). For phenotypic correlations, this provides a more conservative
14
15 324 estimate of sampling variance compared with using the total number of observations. We also collected
16
17 325 more information for each data point for use as moderators to explain potential heterogeneity and bias
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19 326 in the data (e.g., species information, ectothermic vs. endothermic, types of behaviour, publication year).
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21 327 The details of these potential moderators are summarized in the published dataset
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23 328 (<https://osf.io/gmvab/>).

29 329 **(3) Meta-analysis and meta-regression analysis**

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32 330 We employed phylogenetic multilevel meta-analysis (PMMA) to control the effect of phylogeny and
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34 331 other types of non-independence (Hadfield & Nakagawa, 2010; Nakagawa & Santos, 2012). We used the
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36 332 function, *rma.mv* from the R package, *metafor* (Viechtbauer, 2010). A PMMA requires a phylogenetic
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38 333 tree so we constructed a tree using the R package *rotl* (Michonneau *et al.*, 2016) which uses the Open
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40 334 Tree of Life data (Hinchliff *et al.*, 2015). The resulting phylogenetic tree did not have tree branch length
41
42 335 estimates. Therefore, we used the function *compute.brLen*, in the R package *ape* (Paradis *et al.*, 2004) to
43
44 336 estimate branch lengths. With this function, we used the method of Grafen (1989), setting the power to
45
46 337 1, to create an ultrametric tree closely imitating the Brownian motion model of evolution. We used this
47
48 338 ultrametric tree for PMMA by converting it to a correlation matrix (using the *vcv* function from *ape*).
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53 339 We first constructed a meta-analytic model (only with the intercept) with four random factors: 1) species
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55 340 identity (a non-phylogenetic component of species), 2) phylogeny (a phylogenetic component of

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3 341 species), 3) group (i.e., unique set of individuals because the same individuals could be used to estimate
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5 342 multiple relationships between MR and behaviour) and 4) observation (effect size) level random effect
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7 343 (equivalent to the residual term in a normal linear model). The species term captures the similarities of
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9 344 effect sizes within the same species, while the phylogenetic term represents the similarity due to
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11 345 common ancestors (Hadfield & Nakagawa, 2010). For a meta-analysis, the quantification of variance not
12
13 346 due to sampling errors, known as heterogeneity or I^2 (Higgins & Thompson, 2002). We calculated the
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15 347 multilevel-model version of heterogeneity, which quantify I^2 for each random effect as well as the total
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17 348 heterogeneity, following Nakagawa and Santos (2012).

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21
22 349 We then created a set of meta-regression models, which address our three main questions (see
23
24 350 Introduction). The first model added nine behavioural types (activity, boldness, courtship, dominance,
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26 351 exploration, foraging, performance, sociability, and stress response) as a categorical moderator to the
27
28 352 meta-analytic model above (see ESM Table S3 for definitions). The second model had the thermal types
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30 353 (endotherms and ectotherms) as a moderator along with an extra random factor, behavioural types
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32 354 (because the first model showed that this categorical variable was important; we also note that the same
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34 355 random-factor structure was used for the model comparing among-individual and phenotypic
35
36 356 correlations, above). We also created meta-analytic models for endotherms and ectotherms separately
37
38 357 to quantify heterogeneity for both thermal types (total I^2 and I^2 for the four different random terms).
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40 358 Finally, we tested the effects of the breeding statuses (breeding versus non-breeding) and the testing
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42 359 conditions (free-living, wild-caught and tested in captivity, captive reared and tested) by constructing
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44 360 two separate models with the same random factors as the second model. For meta-regression models,
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46 361 we calculated R^2 as the marginal R^2 in mixed models described in Nakagawa and Schielzeth (2013); in
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48 362 meta-analysis, R^2 consisted of variance accounted for after taking away sampling error. Complete details
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51 363 of meta-analytic and meta-regression models are available through the Open Science Framework
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3 364 repository (R-markdown file along with our data set: DOI 10.17605/OSF.IO/GMVAB; web address:
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5 365 <https://osf.io/gmvab/>).

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8 366 **(4) Publication bias analysis and sensitivity analysis**

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11 367 In the past, meta-analyses often identified temporal trends of declining effect size over time, known as
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13 368 the time-lag effect (Jennions & Møller, 2002; Trikalinos & Ioannidis, 2006). We tested this as a part of
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15 369 our publication bias analysis, by including the publication year as a moderator; we created a uni-
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17 370 moderator model with only the publication year as a moderator (along with the five random factors as
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19 371 described above) and a full model with publication year, thermal type, breeding status, and place of
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21 372 origin/testing conditions fitted as fixed effects. The analysis of the time-lag effect showed an important
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23 373 effect of the publication year (see below). Thus, we decided to add a set of sensitivity analyses where we
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25 374 repeated the analysis using meta-analyses and meta-regression with the data since 2007, because this
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27 375 represented the year with a marked increase in the number of studies per year that met the criteria for
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29 376 inclusion in our meta-analysis and because the time trend was not significant after 2007 (see Results for
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31 377 further details).

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33
34 378 Publication bias analysis often includes Egger regression tests along with funnel plots (Egger *et al.*, 1997)
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36 379 and trim-and-fill tests (Duval & Tweedie, 2000a; Duval & Tweedie, 2000b). However, multilevel (non-
37
38 380 independent) data are not amenable to these methods in their original forms. We applied these two
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40 381 methods to the meta-analytic residuals, which consist of the effect-size level effects (equivalent to
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42 382 normal residuals) and sampling errors (*sensu* Nakagawa & Santos, 2012). The meta-analytic residuals
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44 383 were taken from the full model above as this model should have accounted for most heterogeneity in
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46 384 our data set.
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385 III. Results

386 Overall, a phylogenetic multilevel meta-analytic model revealed that MR and behaviour were
387 significantly and moderately correlated ($\beta_0 = 0.261$, 95% confidence interval, CI = [0.053, 0.469]; Table 1
388 and Figure 3A). However, the observed total heterogeneity was high ($I^2_{[total]} = 93.00\%$), implying that this
389 correlation was contingent upon moderators; this set the stage for our meta-regression models (see
390 Table 2). Notably, approximately 21% of variation in the data were explained by phylogeny (Table 2). The
391 first meta-regression model showed that the type of behaviour accounted around 20% of the variation
392 ($R^2 = 19.06\%$), with boldness, dominance, foraging and stress showing moderate to large, significant
393 correlations with metabolic rate, MR (Figure 3A), while activity, exploration and sociability showed little
394 or zero correlation with MR. Though not significant, courtship and performance showed moderate
395 correlations with MR (complete contrasts among the behaviour types are in the ESM, Figure S1).

396 The second meta-regression on the thermal types showed that the mean effect for ectotherms was
397 significant and moderate ($\beta_{[ectotherm]} = 0.301$, 95% CI = [0.102, 0.500]), and that for endotherms the effect
398 was non-significant and weak ($\beta_{[endotherm]} = 0.151$, 95% CI = [-0.076, 0.378]; Figure 4; also see Figure S2),
399 although the contrast between the two was not significant ($\beta_{[contrast]} = -0.150$, 95% CI = [-0.355, 0.056],
400 $Q_{M[df=68]} = 2.031$, $p = 0.15$; $R^2 = 3.88\%$; Figure 4). The separate meta-analytic models mirrored the results
401 from the meta-regression, apart from the mean effect for endotherms being significant ($\beta_{[ectotherm]} =$
402 0.297 , 95% IC = [0.0445, 0.548]; Figure 3B; $\beta_{[endotherm]} = 0.101$, 95% CI = [0.033, 0.200]; Table 1, Figure 3C).

403 In the corresponding meta-regression models incorporating the different behaviour types, we found
404 similar patterns observed in the meta-regression model with all data, but the patterns were, in general,
405 stronger for ectotherms and weaker for endotherms than for all species together (compare panels A, B
406 and C in Figure 3). The magnitude of heterogeneity was high for both meta-analytic models ($I^2_{[total]} =$
407 94.16% for ectotherms and $I^2_{[total]} = 81.22\%$ for endotherms). Notably, much of heterogeneity in the
408 ectotherm data came from the phylogenetic and non-phylogenetic effects of species ($I^2_{[phylogeny]} = 18.20\%$

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3 409 and $I^2_{[\text{species}]} = 44.52\%$, respectively) whereas there was virtually zero variation resulting from phylogeny
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5 410 or species in the endotherm data (Table 2). In our final meta-regression models, in contrast to our
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7 411 predictions, we did not find significant effects of the breeding status ($Q_{M[df=2]} = 1.047$, $p = 0.592$; $R^2 =$
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9 412 0.70%) or testing conditions (i.e., free-living, wild-caught and tested in captivity, captive reared and
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11 413 tested) ($Q_{M[df=2]} = 0.592$, $p = 0.744$, $R^2 = 0.45\%$; Figure 4; Figure S2).

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15 414 We found a time-lag effect in our data set: there was a significant trend of decreasing effect size with
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17 415 publication year ($\beta_{[\text{year}]} = -0.0132$, 95% IC = $[-0.0250, -0.0015]$; $Q_{M[df=1]} = 4.8644$, $p = 0.027$; $R^2 = 4.62\%$;
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19 416 Figure 5). As a sensitivity analysis, we repeated the main analyses (the analyses shown in Figure 3 and
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21 417 Table 2) using only data points published from 2007 onwards, because 2007 was the year in which the
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23 418 number of studies on this topic showed a sharp increase and because the time trend was not significant
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25 419 after 2007 ($\beta_{[\text{year}]} = 0.0123$, 95% IC = $[-0.0185, 0.0431]$; $Q_{M[df=1]} = 0.610$, $p = 0.434$; see Figure 5 & Figure
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27 420 S3). The results of these analyses were quantitatively very similar to the original analyses (see ESM
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29 421 Figure S4, Table S4 & Table S5). An Egger's regression test on the meta-analytic residuals from our full
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31 422 model suggested no evidence for funnel asymmetry ($\underline{t}_{[df=161]} = 0.225$, $p = 0.822$). However, a visual
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33 423 inspection of the funnel plot showed some hint of asymmetry (Figure 6) and this was corroborated by
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35 424 the results from a trim-and-fill test. The trim-and-fill test indicated that seven data points were
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37 425 potentially missing ($p = 0.0039$ for the hypothesis for no missing data points). The meta-analytic mean
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39 426 incorporating these seven filled points was -0.022 (95% CI = $[-0.066, 0.022]$; Figure 6B). This result means
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41 427 we could have overestimate the meta-analytic mean in the original model by 0.022 , but this amount is
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43 428 negligible (see Figure 3 and Table 2). Taken together, we conclude that our results are robust against
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45 429 publication bias.
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5 431 **IV. Discussion**
6 432 We used meta-analyses to test for general relationships between MR and behaviour, as well as the
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8 433 importance of several putative modifiers of the relationship. We demonstrate the importance of two
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10 434 moderators of the magnitude of the relationship between MR and behaviour; the type of behaviour, and
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12 435 the thermal type of the organism (endotherm versus ectotherm). However, we did not find any support
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14 436 for the notion that conditions that impose greater energetic stress on organisms (breeding versus non-
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16 437 breeding or captive vs. free-living), strengthen the relationship between MR and behaviour.
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18 438 Unexpectedly, despite identifying several important moderators for the relationship between MR and
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20 439 behaviour, we found support for an overall relationship between MR and behaviour even when not
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22 440 accounting for any moderators. We discuss the implications of these findings for understanding the
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24 441 functional significance MR and its relationship with behaviour.
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30 442 ***(1) Covariation between MR and behaviour depends on the type of behaviour***

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32 443 Several recent conceptual papers have highlighted the fact that both the magnitude and direction of the
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34 444 relationship between MR and behaviour are likely to be influenced by several moderators (Careau &
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36 445 Garland, 2012; Killen *et al.*, 2013; Mathot & Dingemanse, 2015). For example, the strength of the
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38 446 relationship should differ for different types of behaviour. In particular, behaviours associated with net
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40 447 energy expenditure or net energy gain are expected to have the strongest functional linkage with MR
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42 448 (Biro & Stamps, 2010; Careau & Garland, 2012; Mathot & Dingemanse, 2015). Our results support this
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44 449 prediction. Behaviours that cost net energy or brought in net energy were positively correlated with MR,
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46 450 consistent with a performance energy management model.
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51 451 The strongest relationship between MR and behaviour was found for stress response (response to
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53 452 human handling), a putatively energetically costly trait (Koolhaas *et al.*, 2011; Manzur *et al.*, 2014).
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55 453 Higher metabolic rates were associated with quicker escape latencies from observers, greater struggle
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3 454 rates and higher breath rates. There was also a moderately strong, albeit non-significant, correlation
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5 455 between 'performance' traits (e.g., maximum sprint speed, maximum sustained activity, etc.), which are
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7 456 also energetically demanding behaviours. Although courtship behaviour is often assumed to be
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9 457 energetically costly, we found a weak and non-significant relationship between MR and
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11 458 courtship/mating behaviours. However, our analysis of courtship behaviour was based on only three
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13 459 studies (Figure 3); two studies of calling rates, and one study of copulation duration. This is not a
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15 460 comprehensive sample of courtship behaviours. Given that the relationship between courtship
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17 461 behaviours and net energy expenditure may vary for different types of courtship behaviours and as a
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19 462 function of both the intensity and duration of the behaviour (Clark, 2012), the number of existing studies
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21 463 ($N = 3$), is insufficient to draw strong conclusions at this time.
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26 464 We also found moderate to strong and significant correlations between MR and each of the traits
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28 465 assumed to be associated with net energy gain (dominance, boldness, foraging). Animals with higher MR
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30 466 were more dominant (i.e., had priority access to food), bolder (i.e., resumed feeding more quickly after a
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32 467 disturbance), and foraged more intensively.
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36 468 In contrast, there was little or no support for significantly positive correlations between MR and
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38 469 behaviours with unknown or putatively weak and/or inconsistent relationships with net energy gain or
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40 470 net energy expenditure. As predicted, there was no support for a relationship between sociability and
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42 471 MR. Although the sociability estimate was derived from only three studies, the point estimate lies at
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44 472 zero, as predicted. There was also no support for a relationship between MR and either activity or
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46 473 exploration. Activity and exploration are two commonly measured behaviours in animal personality
47
48 474 research; both provide some measure of the movement behaviour of animals, and so may be assumed
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50 475 to involve some net energetic costs. However, these movements can range from very low (e.g., periodic
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52 476 slow walking) to very high gross energetic costs (e.g., continuous running, flying). Further, these
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54 477 behaviours are assumed to increase encounter rates with resources. Thus, the net consequences of
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3 478 these behaviours could range from high net energetic costs to high net gains. Thus, the lack of
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5 479 correlation between MR and exploration/activity is not surprising (Mathot & Dingemanse, 2015), and
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7 480 further substantiates recent criticism of the use of standardized behavioural assays in animal personality
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9 481 research without careful consideration of the functional significance of the traits being studied (Carter *et*
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11 482 *al.*, 2013). We are not suggesting that exploration and activity are not associated with net energetic costs
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14 483 or gains, but rather that the relationship likely differs in different organisms or under different conditions
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16 484 (Carter *et al.*, 2013; Mathot *et al.*, 2012). The consequences of greater activity or exploration on net
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18 485 energy expenditure or gain should not remain untested assumptions in studies aimed at understanding
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20 486 functional linkages between metabolism and behaviour, but should be evaluated directly if we are to
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22 487 better understand why they are (or are not) associated with MR.
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26 488 **(2) Relationships between MR and behaviour differ across thermal types**

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28 489 We also tested the idea that greater opportunities for energy (re-)allocation (Humphries & Careau, 2011)
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30 490 could obscure relationships between MR and behaviour (Careau & Garland, 2012; Mathot &
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32 491 Dingemanse, 2015) by contrasting ectotherms with endotherms. Because endotherms can offset
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34 492 thermoregulation costs by substituting heat produced through activity (Careau & Garland, 2012;
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36 493 Humphries & Careau, 2011), we predicted that endotherms would show weaker relationships between
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38 494 MR and behaviour compared with ectotherms. Although our meta-analysis confirmed this prediction, we
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40 495 suggest that thermal substitution alone cannot account for this observed differences. Within
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42 496 endotherms, opportunities for heat substitution vary both taxonomically and allometrically (Humphries
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44 497 & Careau, 2011). Thus, if heat substitution was a primary reason for a weaker relationship between MR
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46 498 and behaviours in endotherms, we would expect to observe strong phylogenetic- or species-related
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48 499 heterogeneity. However, this was not the case. In fact, the proportion of heterogeneity associated with
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50 500 phylogeny and species in endotherms was close to zero (Table 2). Thus, although the strength of the
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3 501 relationships between MR and behaviour appear to differ for endotherms and ectotherms, the
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5 502 mechanism underlying this variation is unclear.
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9 503 **(3) No evidence that greater energetic stress strengthens relationships between MR and behaviour**

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11 504 The rationale for predicting relationships between MR and behaviour hinges on the assumption that
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13 505 animals face constraints in balancing their energy budgets. We tested whether conditions that create
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15 506 greater energetic stress for organisms (reduced access to food and reproduction) strengthen the
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17 507 relationship between MR and behaviour but found no support for this. There was no support for
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19 508 differences in mean effect sizes for breeding versus non-breeding animals, nor for contrasts between
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21 509 studies in captive versus free-living organisms (which we assume covaries with access to resource, as
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23 510 captive studies typically provide ad libitum access to food) (Figure 4, Figure S2). The lack of an effect of
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25 511 breeding status may reflect that breeding does not actually impose greater challenges on organisms in
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27 512 terms of balancing their energy budgets. This could occur if animals adjust their metabolic profiles during
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29 513 breeding, for example, suppressing resting MR to offset energetic costs associated with breeding
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31 514 behaviour (Welcker *et al.*, 2015), if increased energetic costs of breeding are easily offset by increased
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33 515 food availability, or if breeders represent non-random sample of the populations (i.e., animals that are
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35 516 the least energetically stressed).
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41 517 We also found little effect of captivity on the strength of the relationship between MR and behaviour,
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43 518 despite the fact that food availability and predictability are almost always higher under captive
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45 519 compared with free-living conditions. Animals with increased access to food for prolonged periods may
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47 520 exhibit metabolic adjustments (e.g., increased metabolic rate under increased food availability, Mueller
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49 521 & Diamond, 2001), such that energetic constraints are similar in the field versus the lab. However, there
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51 522 were also no differences between wild-caught animals tested in captivity shortly after capture and either
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53 523 free-living or lab-reared animals. Thus, although some studies have found that increased energetic
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55 524 constraints strengthen the relationship between MR and behaviour (e.g., Killen *et al.*, 2011), our analyses
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3 525 suggest that this may not generally true. Although unexpected, this result is also promising in that it
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5 526 implies that the results from studies investigating links between MR and behaviour in the lab can be
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7 527 generalizable to free-living organisms.
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10 528 **(4) Energy management model inferences**

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13 529 The energy management model of the organism is believed to be critical in shaping both the magnitude
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15 530 and direction of relationships between MR and behaviour (Careau & Garland, 2012; Mathot &
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17 531 Dingemanse, 2015). Predicted correlations between MR and specific behaviours are contingent on the
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19 532 energy management strategy, with positive, negative and zero correlations all possible depending on the
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21 533 type of behaviour being considered, and the energy management strategy. Alternative energy
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23 534 management models do not always make exclusive predictions (e.g., both the independent model and
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25 535 the performance model predict positive correlations between behaviours that bring in net energy and
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27 536 MR). Further, where relationships are predicted between MR and a class of behaviours such as
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29 537 behaviours that bring in net energy, the relationship need not exist for every type of behaviour that
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31 538 brings in net energy, but may be present for only a subset of them. Thus, in an earlier opinion (Mathot &
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33 539 Dingemanse, 2015), we pointed out that a single estimate of the relationship between MR and a single
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35 540 behaviour provides weak inference. In this meta-analytical review, by combining estimates from multiple
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37 541 studies and explicitly taking into account the support for relationships between MR and different types
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39 542 of behaviours, we can establish the overall support for the alternative energy management strategies
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41 543 across all published studies.
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47 544 We found support for an overall positive relationship between MR and behaviour. This is consistent with
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49 545 a previous meta-analysis focused exclusively on among-individual correlations between state (including
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51 546 MR measures) and behaviour (Niemelä & Dingemanse, 2018). More specifically, we found the strongest
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53 547 support for positive relationship between MR and behaviours that bring in net energy or cost net energy,
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55 548 with no support for relationships between MR and behaviours with putatively weak or inconsistent
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3 549 consequences for energy gain/expenditure. Further, effect sizes were comparable for behaviours that
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5 550 bring in net energy and behaviours that cost net energy. The performance model is the only one to
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7 551 predict similar effects for both types of behaviour, suggesting that our sample of studies is comprised
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9 552 primarily of organisms with a performance energy management model. As this meta-analysis includes a
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11 553 taxonomically diverse set of organism, this suggests that the performance model may be most common.
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14 554 In a recent paper, Portugal *et al.* (2016) compiled estimates of MR and daily energy expenditure in 7
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16 555 birds and 4 mammals to evaluate support for alternative energy management models. Although they
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18 556 observed heterogeneity in estimates across species, a meta-analysis of the reported slopes reveals that
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20 557 the best supported model is in fact the performance model (see ESM Text S1), as the overall slope of the
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22 558 relationship between MR and DEE is > 1 ($\beta = 1.11$, 95% CI = 1.00, 1.22).

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26 559 Finally, we assessed evidence for publication biases in the studies compiled for this meta-analysis using
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28 560 several tests (e.g., Egger's regression and trim-and-fill tests). Overall, the evidence for publication bias
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30 561 distorting our main results was weak (see Figure 6). However, our time-lag analysis revealed two notable
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32 562 patterns (Figure 5). First, the number of studies published per year increased dramatically from 2007
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34 563 onwards, and second, there was a significant trend towards decrease effect sizes estimates over time.
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36 564 We propose that both of these patterns may be explained by the rapid growth of the field of animal
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38 565 personality research in the last two decades. A major aim of the field of animal personality research is to
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40 566 understand the factors that promote consistent among-individual differences in behaviour, with several
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42 567 influential papers proposing links between among-individual differences in MR and behaviour around
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44 568 this time (Biro & Stamps, 2008; Biro & Stamps, 2010; Careau *et al.*, 2008). The second consequence of
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46 569 the boom of animal personality studies is that there was a shift in the types of behaviours studied. In
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48 570 particular, there was a rapid increase in the number of studies using standardized assays of exploration
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50 571 and activity behaviour, two of the behaviours revealed by our current analyses to exhibit no relationship
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52 572 with MR.
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56 574 **V. Conclusions**
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8 575 (1) The likelihood of any overarching relationship between MR and behaviour has been questioned given
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10 576 that such relationships are likely to be shaped by a variety of factors (Careau & Garland, 2012; Killen *et*
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12 577 *al.*, 2013; Mathot & Dingemanse, 2015). Our current analysis identified two important moderators; the
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14 578 thermal type of the organism, and the type of behaviour.

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18 579 (2) Relationships between MR and behaviour were in the same direction for endotherms and ectotherms
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20 580 (positive relationship between MR and behaviours that either bring in or cost net energy), but the
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22 581 strength of the relationship was weaker in endotherms compared with ectotherms.

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26 582 (3) We also found that the types of behaviours that showed the strongest (positive) associations with MR
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28 583 were behaviours with clear consequences for net energy expenditure (e.g., maximum sprint speed,
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30 584 sustained running speed, maximum distance travelled, etc.) or gain (e.g., foraging, dominance, boldness).

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33 585 (4) In contrast, common behavioural assays used in animal personality research (exploration and activity)
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35 586 which have unknown or putatively weak consequences for net energy cost or gain showed no
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37 587 relationship with MR.
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41 588 (5) Taken together, the results of our meta-analysis highlight the importance of studying behaviours that
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43 589 are functionally relevant in the context of the research question.
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48 591 **VI. Acknowledgements**

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52
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734 **VIII. Supporting information**

735 Additional supporting information may be found in the online version of this article.

Table 1: Summaries and results from phylogenetic multilevel meta-analyses of the overall data set, and the data set separated for ectotherms and endotherms. Note that the number of individuals ($N_{\text{[individuals]}}$) represents the sum of individuals making up each effect size estimate, not the total number of unique individuals (which is over 8000 for the whole data set). This non-independence is accounted for in our analyses (see the text). LCI and UCI denotes the lower and upper confidence limits, respectively, for 95% confidence intervals.

Data	$N_{\text{[individuals]}}$	$N_{\text{[effect sizes]}}$	$N_{\text{[groups]}}$	$N_{\text{[species]}}$	Estimate	SE	z value	p value	LCI	UCI
Overall	11849	163	87	48	0.261	0.106	2.459	0.014	0.053	0.469
Ectotherm	7539	69	47	28	0.297	0.128	2.314	0.021	0.045	0.548
Endotherm	4310	94	40	20	0.101	0.050	2.026	0.043	0.003	0.199

Table 2: Total heterogeneities (I^2 , which can vary between 0 and 1) and heterogeneities at each hierarchical level (fitted as random effects; see the text) for meta-analytic models with phylogeny for the overall dataset, and separated for ectotherms and endotherms.

Data	$I^2_{[\text{species}]}$	$I^2_{[\text{phylogeny}]}$	$I^2_{[\text{group}]}$	$I^2_{[\text{effect size}]}$	$I^2_{[\text{total}]}$
Overall	0.138	0.209	0.068	0.517	0.930
Ectotherm	0.445	0.182	<0.001	0.311	0.942
Endotherm	<0.001	<0.001	0.186	0.626	0.812

For Review Only

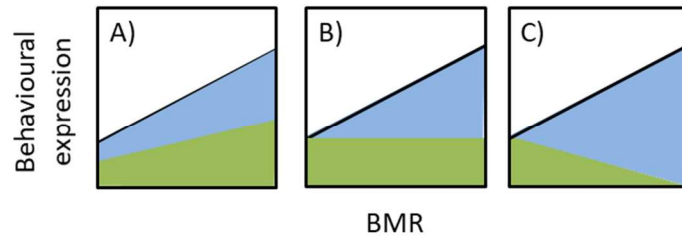
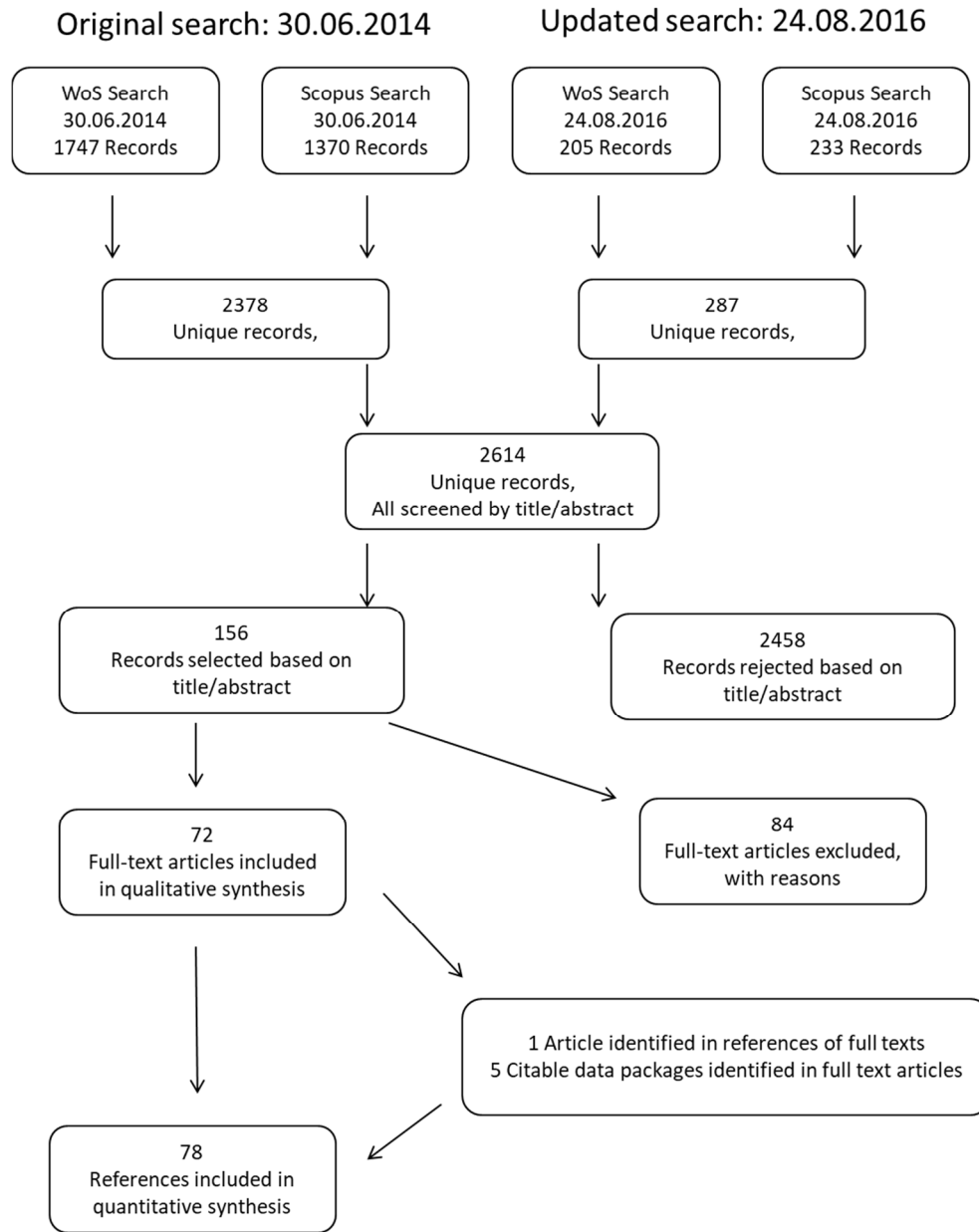


Figure 1: Illustration of how opportunities to allocate among alternative options can create scenarios where the relationship between specific behaviours and MR do not match the more general predictions for the energy management model. Consider an example where an organism can express two behaviours that both have net positive effects on energy intake: foraging boldness (green) or resource defense (blue). The total expression of each behaviour is indicated by their area under the black line. In a) expression of both behaviours increases with increasing MR (i.e., the area occupied by both green and blue increases from left to right across the x-axis), matching the more general prediction. However, in b) only one behaviour shows the predicted relationship with MR (blue, but not green), and in c) one behaviour shows the predicted relationship (blue) while the other behaviour shows the opposite relationship to the one predicted (green).



44 **Figure 2:** PRISMA flow chart indicating articles identified during different phases of the systematic
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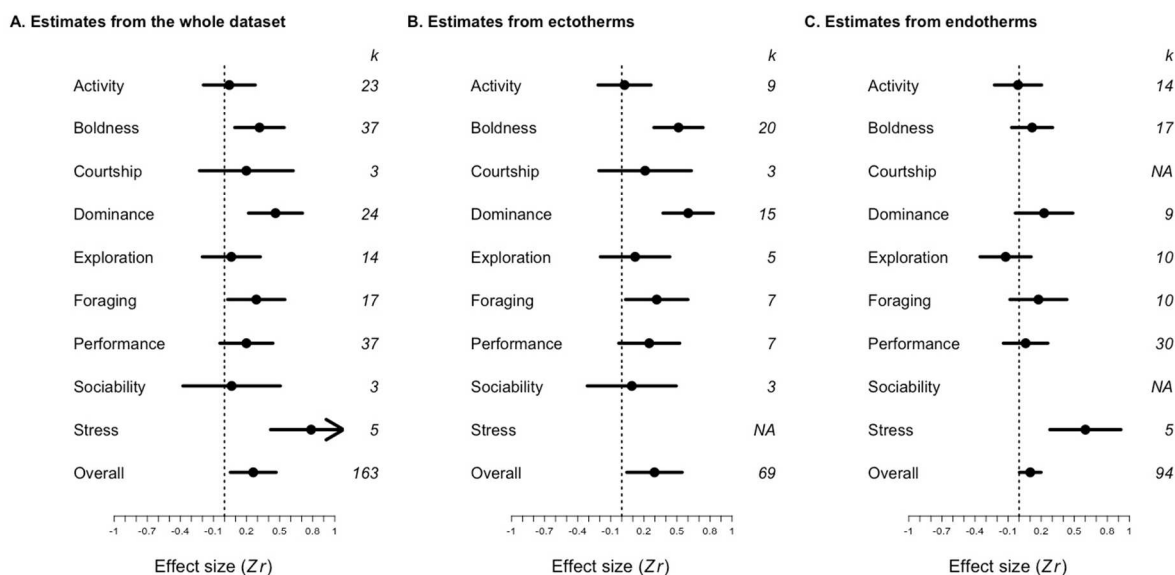


Figure 3: Effect sizes for the relationship between metabolic rate and behaviour separated for different types of behaviours, including an overall estimate (bottom). Panel (A) combines the full data set, panel (B) presents estimates from ectotherms, and panel (C) presents estimates from endotherms. Circles denote point estimates, whiskers denote 95% confidence intervals (the arrow tip indicates the end point is beyond the scale), and k is the number of effect sizes (NA = not available; see Table 1). Note that estimates derived from identical data (e.g., stress behaviours in panel A and panel C) may not be identical because model structures differed for each analysis.

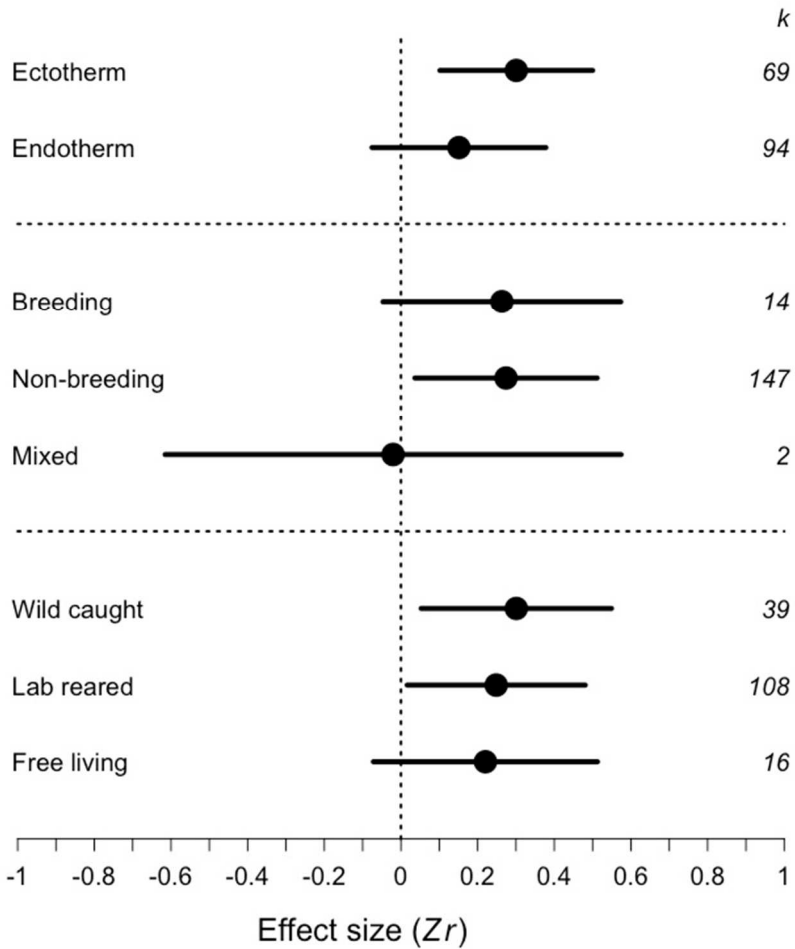


Figure 4: Effect sizes for the relationship between metabolic rate and behaviour for three potential moderators: thermal type of the organism (ectotherm or endotherm), breeding context (breeding, non-breeding, or mixed/unspecified), and testing condition (wild-caught animals tested in the lab, lab-reared animals tested in the lab, free-living animals tested under natural conditions). Circles denote point estimates, whiskers denote 95% confidence intervals and k is the number of effect sizes.

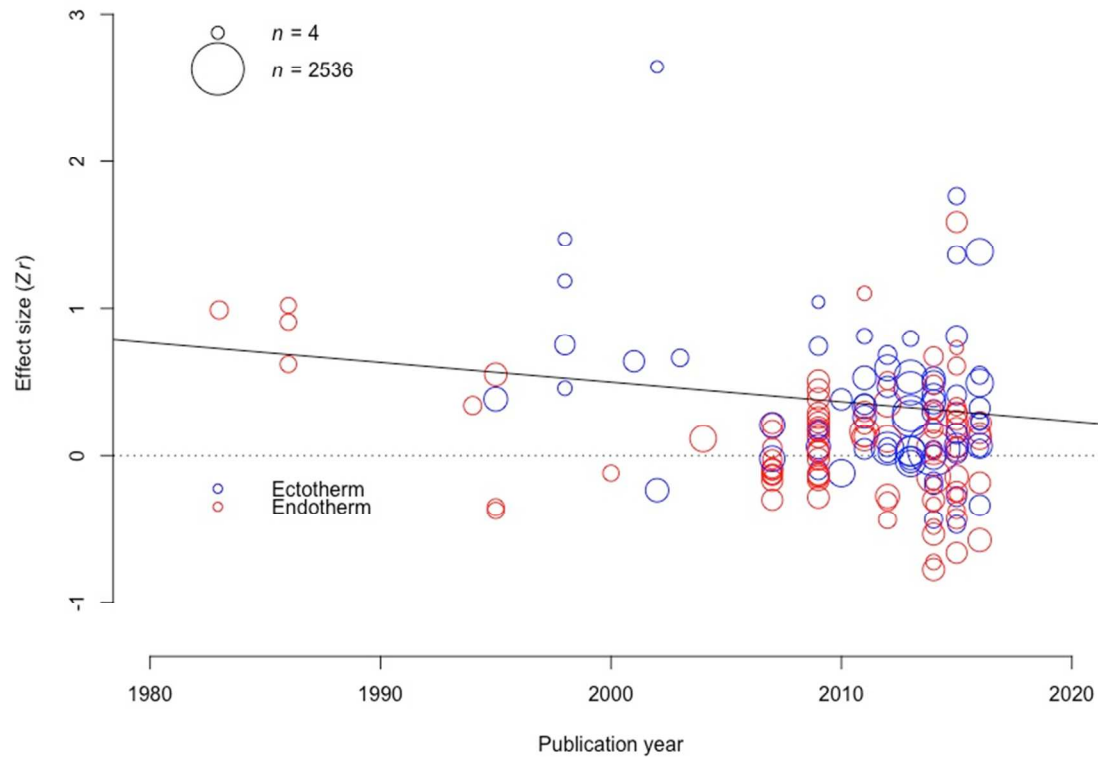


Figure 5: A bubble plot showing a time-lag effect characterised by effect sizes as a function of publication year. After the first study (for an endotherm) in the early 80's, the number of studies remains sparse until 2007, when there is a notable increase in the number of studies for both ectotherms and endotherms (see the text). The size of the circle represents the sample size (on a linear scale), with the largest circle representing the sample size of 2536 animals and the smallest circle representing a sample size of 4 animals.

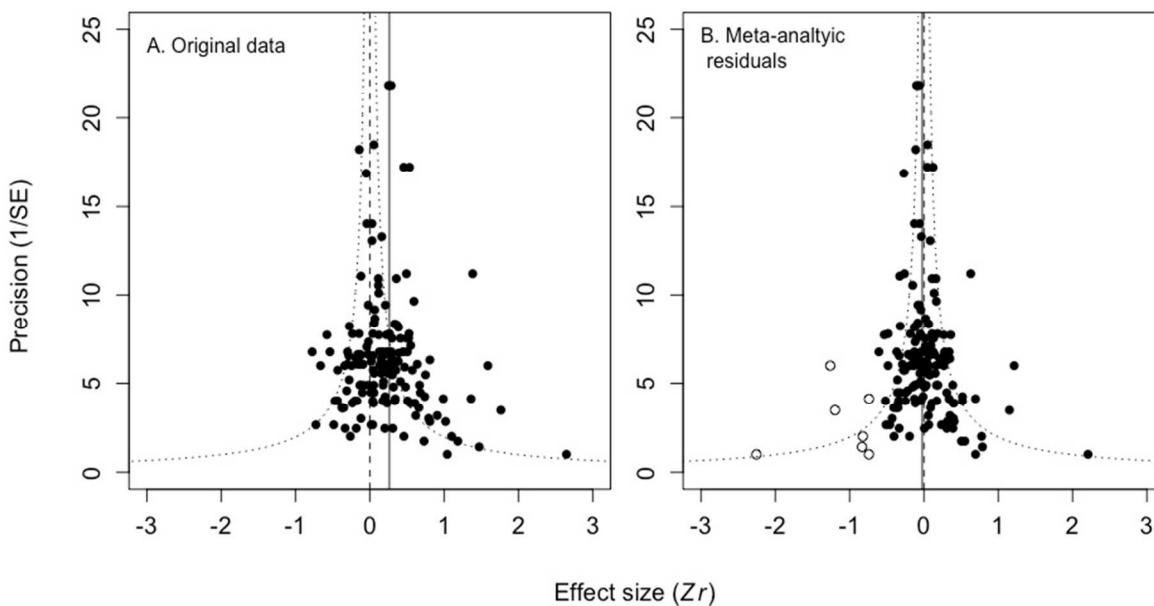


Figure 6: Funnel plots showing effect size and its precision which is the inverse of the square-root of the sampling variance (or standard error, which is the standard deviation of the estimate/effect size). Panel (A) shows the original (raw) data and the meta-analytic mean (the solid vertical line), but note that the original data, which has a non-independent data structure is not appropriate for assessment of funnel asymmetry. Panel (B) shows the meta-analytic residuals (see the text) and data points added by a trim-and-fill test (empty dots) along with the meta-analytic mean (note that the meta-analytic residuals have a mean of zero).