

1 **Modelling activity patterns of wild animals - an application of the Multiple Discrete-Continuous Extreme**
2 **Value (MDCEV) model**

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5 **Abstract**

6 Advanced econometric models used in the field of transport or marketing are becoming increasingly
7 sophisticated and able to capture complex decision making and outcomes. In this paper, we apply state-
8 of-the-art discrete-continuous choice models to the field of Ecology, in particular to model activity
9 engagement of the population of Swedish Brown bears. Using data from GPS collars that track wild animals
10 over time, we estimate a Multiple Discrete-Continuous Extreme Value (MDCEV) model to understand
11 activity engagement and duration as a function of both bear characteristics and other external factors. We
12 show that the methodology is not only suitable to address this aim, but also allows us to produce insights
13 into the connection between the animal's age and gender and activity engagement as well as the links
14 with climate variables (temperature and precipitation) and human activity (hunting).
15

16 **1. Introduction**

17 The understanding of the patterns of movement of living organisms is a prominent area of study in animal
18 biology and ecology. Indeed, the presence and distribution of different animals in space and time are a
19 product of the underlying process of animal movement (Turchin 1998; Nathan et al. 2008; Kays et al. 2015).
20 Identifying behavioural states along an animal's movement path is straightforward when visual
21 observation is possible (Bates and Byrne 2009; Hayward et al. 2009). Direct observation is an effective
22 method for investigating animal behaviour and the least prone to errors when it comes to identifying the
23 specific behavioural state (Loettker et al. 2009; Shamoun-Baranes et al. 2012). However, in most cases it
24 is challenging if not impossible to constantly observe animals as they go through their lives in their natural
25 habitat. Wild animals are often elusive and can reside in remote areas with challenging terrain. Many
26 species minimize exposure to perceived threats, which often include human encroachments. Due to the
27 difficulties in locating and observing animals, early studies have been marred by small sample sizes, often
28 resulting in insufficient data for statistical inference (Caro 2007). In addition, investigating wild animal
29 behaviour via direct observation may pose threats to researcher safety.
30

31 The development and diffusion of animal telemetry devices has revolutionized the ability to study animal
32 movements and behaviour in the wild (Cagnacci et al. 2010; Kays et al. 2015), helping to overcome many
33 of the practical, logistical, and financial challenges of direct field observation. Recent advances in GPS
34 radiocollar technologies allow tracking of animals for long sampling periods, providing large data sets of
35 locations at flexible time intervals (Cagnacci et al. 2010). Telemetry data have created possibilities that
36 allow to address some of the most fundamental ecological hypotheses about space use, movement,
37 resource selection and behaviour. The GPS locations can be used to investigate, among others, habitat
38 selection (Signer et al. 2019), spatiotemporal movements (Nathan et al. 2008) and habitat influences on
39 animal movement (Patterson et al. 2008). One line of research specifically relevant to the present work is
40 the use of GPS radio-collar data to infer and analyse animal behavioural states. A common assumption in
41 the ecology literature is that individual animals have a small set of movement strategies (Nathan et al.
42 2008), and the time allocation to different behaviours (or "activity budgets") depends on environmental,
43 individual animals' characteristics and external factors (Hooten et al. 2017). There are many approaches
44 to infer animals' behaviour from radio-collar data. Some studies have decomposed an individual animal's
45 movement trajectory into a broad set of movement bouts based on rates of movement (Johnson et al.
46 2002). Others have inferred behavioural states based on the time required for an animal to first move out
47 of a circle centred on a location along their path (Frair et al. 2005) or total time spent in the vicinity of a
48 location (Barraquand and Benhamou 2008). For example, in carnivore studies in which authors attempt to
49 derive behaviour from GPS relocation data, the main goal has often been to identify GPS location clusters
50 indicative of predation or bedding events (Merrill et al. 2010; Ordiz et al. 2011; Rauset et al. 2012). Clusters
51 form when an animal spends a certain amount of time within a site of a given radius, where time and

52 radius are specified by the researcher and should be tailored to the behaviour of the study species and
53 field conditions.

54 Movement trajectory and cluster analysis are common techniques for identifying behavioural states, such
55 as resting (Ordiz et al. 2011), predation (Rauset et al. 2012) or foraging (Bastille-Rousseau et al. 2011;
56 Hertel et al. 2016a; Hertel et al. 2016b). Frequently used methodologies to study patterns of animal
57 movement and behaviour are autocorrelation analysis (Boyce et al. 2010) or generalized additive models
58 for either net squared displacement or step length (Ciuti et al. 2012). The detailed overview of these
59 techniques can be found in (Gurarie et al. 2016; Hooten et al. 2017). Simple random-utility based models
60 (such as logit models) have also been used to study the behavioural states (e.g. Ordiz et al. 2011).

61 While the above-mentioned studies provide interesting insights into animal behavioural states, they are
62 generally narrow in scope because they focus on the analysis of a single type of behaviour (also referred
63 to as “activity” from here on) in isolation. This can limit the insights produced due to the lack of the overall
64 picture: for example, an animal could move in a given direction so that they can conduct another activity,
65 or because they have been conducting another activity/experienced certain circumstance.

66 The present work represents an advancement in this field of research not only by analysing engagement
67 in multiple activities within a given time frame, but by jointly modelling activity choice and its duration.
68 We refer to such choice processes as discrete-continuous. Many studies involving humans have recognised
69 the importance of accommodating for the joint nature of multiple discrete-continuous decisions in
70 econometric models (e.g. Bhat et al. 2005, Calastri et al. 2022, Van Nostrand et al. 2013). In this work we
71 aim to test whether models accounting for the discrete-continuous nature of activity choice can be used
72 to infer more detailed insights about animals’ behaviour by acknowledging the connection between the
73 choice of activity and its duration. For example, an important topic in brown bear research, highly relevant
74 from the management and conservation perspective, is the impact of hunting on bears behaviour. A few
75 studies have demonstrated that apex predators may perceive and respond to human-caused risk like prey
76 responds to a natural predator (Ordiz et al. 2011). However, these studies explore the impact of hunting
77 on a single behavioural state, for example Hertel et al. (2016a) analysed the impact of hunting on foraging,
78 Ordiz et al. (2011) analysed the impact of hunting on the choice of places for bedding, whereas in this work
79 we are able to document the impact of hunting on engagement in multiple activities and their durations.
80 Our approach allows for better understanding of the full cost in terms of energy expenditure and intake
81 for bears resulting from hunting. These topics lay in the heart of budget activity (Christiansen et al. 2013)
82 and foraging ecology (Pyke, 2019).

83
84 Multiple discrete-continuous models have not been previously applied in Ecology, and this paper aims to
85 present a proof-of-concept of their potential usefulness in this discipline.

86 In particular, by combining movement trajectory and cluster analysis we identify distinct behavioural
87 states of brown bears (*Ursus arctos*) (i.e., rest, forage, move). We then apply the state-of-the-art Multiple
88 Discrete-Continuous Extreme Value (MDCEV) model to jointly represent the engagement in and duration
89 of activities. The long-term individual-animal based dataset used in this study allows us to better
90 understand the effect of individual bear characteristics, environmental variables and external factors on
91 bear behaviour over time.

92 The remainder of this paper is organised as follows. The next section presents the data, before we discuss
93 the model structure. This is followed by the results of the empirical analysis, and finally, the study
94 conclusions.

95 96 **2. Methods**

2.1. Species description, study area and data collection

The GPS telemetry data from brown bears are collected as part of the Scandinavian Brown Bear Research Project (www.bearproject.info), an individual-bear level, long-term population monitoring program in south-central Sweden. The bear population in the study area has been continuously monitored since 1985 (Swenson et al. 1994). As part of this project, bears are captured by darting (Dan-Inject, Børkop, Denmark) with an immobilizing drug from a helicopter soon after hibernation and den emergence in late April. Captured bears are equipped with a GPS collar (GPS Plus; Vectronic Aerospace, Germany) prior to re-release. For more information on capture and handling procedures, see Arnemo et al. (2011).

The study area is situated in Dalarna and Gävleborg counties in south-central Sweden. The terrain is hilly, with elevations between 250 and 650m above sea level and mostly covered by intensively managed coniferous forests mainly composed of Scots pine (*Pinus sylvestris*), Norway spruce (*Picea abies*), and birch (*Betula* spp.). The human settlement in the area consist of 24 scattered small villages and a few seasonally used independent houses. Human population density is low and ranges from 4.1 to 7.1 habitants per km² (Ordiz et al. 2012), but there is an extensive network of forestry roads (Frank et al. 2015). Bear density is about 30 bears per 1,000 km² (Bellemain et al. 2005), and bears are hunted in the study area (Frank et al. 2017). The annual brown bear hunting season in Sweden starts on 21 August and lasts for approximately 2 months or until predefined quotas are filled. Hunting is allowed from 1h after sunrise until 2h before sunset (Bischof et al. 2008, Bischof et al. 2018).

In this study, we have focussed on GPS relocation data from adult (≥ 4 years) bears (Zedrosser et al. 2006, Zedrosser et al. 2009) collected on an hourly basis between 2008 and 2015. We removed GPS locations with a dilution of precision $>10^1$ from the data (D'eon and Delparte 2005). We focussed our analysis on the *berry season* (15th July – 15th September) (Ordiz et al. 2011, Hertel et al. 2018), i.e., the time period when bears build adipose tissue reserves in preparation for hibernation (Manchi and Swenson 2005). During this time period, the bears in our study area forage almost exclusively on berries, mainly *Vaccinium* spp. (Stenset et al. 2016). As the main purpose of this paper is to showcase MDCEV modelling framework in ecology, to reduce ecological complexity, we focused on solitary adult males (n=24) and adult females (n=40). Low satellite coverage may lead to failed GPS fixes (Moe et al. 2007). In only 30.9% of days for which fixes were recorded, all 24 fixes were available. In order to preserve data quality, we resorted to retaining observations (at the day level) for which at least 22 hourly fixes had been recorded (50.8% of all days). While data with up to two discontinuous missing fixes were retained, if the two missing fixes were contiguous, the whole day of data was discarded. In the case of such discontinuous missing fixes, the position for the missing hour was approximated as the midpoint between the two nearest available relocations.

2.2. Behavioural classifications

Bear behaviour was classified into three activities based on GPS data: *foraging*, *resting*, and *moving*. The classification of the behavioural states was based on the existing literature for brown bears in the study area (i.e. Moe et al. 2007, Ordiz et al. 2011, Hertel et al. 2016a, Hertel et al. 2016b). A bear was defined as exhibiting resting behaviour when a minimum of three consecutive GPS locations within a radius of 30m were recorded based on 30-minute GPS-relocation data (Ordiz et al. 2011). Given the coarser temporal

¹ Dilution of precision (DOP) is a term used in satellite navigation and geomatics engineering to specify the error propagation as a mathematical effect of navigation satellite geometry on positional measurement precision. Observations with DOP larger than 10 are considered as having too large error to be used for analysis (D'eon and Delparte 2005).

140 resolution in our data (60 min time intervals), we defined resting behaviour (*Rest*) as an activity in which
 141 an animal stayed at least 1hour within a radius of 30m. Berry foraging by bears is characterized by slow
 142 and meandering movements (Stelmock and Dean 1988). Hertel et al. (2016a, 2016b) defined berry
 143 foraging in our study population as continuous movements in which a bear covered a distance of 25–300
 144 m over at least three consecutive 30 min intervals. Field validation by Hertel et al. (2016a) confirmed that
 145 bears were foraging on berries at 80% of the locations classified as forage based on GPS-relocation data.
 146 For the purposes of this study, we slightly modified the criterion used by Hertel et al. (2018) and defined
 147 feeding behaviour (*Feed*) as relocations within a distance of 30–300 m in 60 min (2 consecutive GPS fixes).
 148 Any behaviour with movements longer than 300 m in 60min was classified as *Move*. This resulted in a very
 149 wide range of travelled distances in this category (i.e., mean=911 m, std dev=592, max distance=8,504m).
 150 Given the temporal resolution of our data, very heterogenous behaviour is likely within a 60 min interval,
 151 including the whole spectrum from mostly feeding to pure travel. We tested two approaches to deal with
 152 the *Move* category; first, we classified all observations with travelled distances longer than 300m/h as
 153 *Move*; second, we further divided *Move* into two subcategories, *Short move* – including all relocations
 154 ranging between 300m/h and 600m/h, and *Move* comprising all relocations for which the covered distance
 155 was larger than 600m. The second approach resulted in significantly better model performance and was
 156 used in the final model specification.

157
 158 Table 1 presents a summary of the sample characteristics based on the defined criteria. As can be seen,
 159 all four types of activities are conducted on the vast majority of all days, with *Rest* being conducted every
 160 day.

161
 162 Table 1. Summary statistics of GPS relocation data of radio-collared brown bears during the berry season
 163 (15th July – 15th September) in south-central Sweden, 2008-2016.

	Daily activity duration (hours)			
	Mean	Std. dev.	Min.	Max.
Rest	8.72	2.57	2	21
Feed	6.00	2.81	0	18
Short move	5.47	3.02	0	18
Move	3.79	2.06	0	11
	Bear characteristics			
	Mean	Std. dev.	Min.	Max.
Sex	40 – females 24 – males			
Age (years, average in sample)	9.36	4.32	5	22
Number of observations (days) (% of the sample)				
July (15 th – 30)	1,321 (28%)			
August (1 st – 31)	2,399 (51%)			
Sept (1 st – 15 th)	967 (21%)			
Days with Rest > 0h	4,687 (100%)			
Days with Feed > 0h	4,647 (99.1%)			
Days with Short move > 0h	4,508 (96.2%)			
Days with Move > 0	4,505 (96.1%)			
	Mean	Std. dev.	Min.	Max.
Number of obs (days) per bear	73.25	60.36	1	266
Total number of observations	4,687			

164

165 3. Modelling framework

166 3.1. Overview

167 The family of Multiple Discrete-Continuous Extreme Value (MDCEV) models initially developed by Bhat
168 (2005) and subsequently extended in different directions (Bhat, 2008; Castro et al., 2012; Pinjari and Bhat,
169 2010, Mondal and Bhat, 2021, Palma and Hess, 2022), represents the current state of the art in modelling
170 multiple discrete-continuous choices. Human travel behaviour has been the main field of application of
171 this modelling framework, for example in the study of the choice of vehicle type and mileage (Bhat and
172 Sen, 2006), and to type and duration of activities (Bhat, 2005; Calastri et al., 2022). Application beyond
173 transport choices include those analysing residential energy consumption (Iraganaboina, & Eluru, 2021),
174 social interactions (Calastri et al., 2017) and consumer purchase behaviour (Lu et al., 2017). To the best of
175 our knowledge, MDCEV has never been applied to the study of animal behaviour.

176
177 The model is derived coherently with the random utility maximisation theory in economics, but
178 relaxes the mutual exclusivity assumption inherent in traditional discrete choice models. This means that
179 subjects are not constrained to selecting one option (in our case one activity) but allocate their available
180 resources to a combination of these. While the model has been developed in the generic context of
181 consumption choices, we describe it in terms of time allocation decisions, in line with the application
182 presented in this paper.

183 The MDCEV model is based on a direct utility function $U(x)$ that individuals (in this case bears)
184 maximise by choosing to allocate a vector x of non-negative time intervals to each of the K possible
185 activities, $x = (x_1, \dots, x_K)$. The choice of total time allocation is subject to a time “budget” constraint $\sum x = E$,
186 where E is the total time available.

187 The vector x generally includes a so-called “outside activity” to represent allocation to an activity
188 that is always undertaken by all the individuals (bears) in the sample, in our case *Rest*. A decision needs to
189 be made on the unit of measurement. In our case we work with data at the day level, implying a time
190 budget of 24 hours per observation. The time budget takes the following form:

$$\sum_{k=1}^K x_k = E, x_k > 0, x_k \geq 0 \forall k (k = 2, \dots, K) \quad (1)$$

191 where in our model activity 1 is *Rest*, i.e. the *outside activity*, and $K=4$ (*Rest*, *Feed*, *Move* and *Short move*)
192 and $E=24$ hours.

193 194 3.2. Econometrics

195 The utility formulation, introduced by Bhat (2008) is given by:

$$196 U(x) = \frac{1}{\alpha_1} \psi_1 x_1^{\alpha_1} + \sum_{k=2}^K \frac{\gamma_k}{\alpha_k} \psi_k \left(\left(\frac{x_k}{\gamma_k} + 1 \right)^{\alpha_k} - 1 \right), \quad (2)$$

197 such that $U(x)$ is quasi-concave, increasing and continuously differentiable with respect to x and ψ . ψ_k is
198 the baseline utility accrued from activity k . It is a function of observed characteristics of the individual
199 (bear) and of activity k , z_k , which also includes a constant δ_k representing the generic preference for activity
200 k , so that $\psi_k = \psi_k(z_k) * e^{\delta_k}$. The parameters γ_k and α_k relate to activity k . The γ_k parameters are
201 translation parameters that allow the model to accommodate corner solutions, i.e. observations for which
202 no time is allocated to a given activity k . They also affect satiation, as a higher γ_k implies that more time
203 invested in the corresponding activity (x_k) is needed to reach saturation. The α_k parameter is solely
204 associated with the satiation effect, i.e., decreasing marginal utilities.

205 Empirical identification requires some constraints for normalisation, and in our work, we make
206 use of the implementation of the MDCEV model in which we have product-specific γ parameters, i.e. we
207 estimate γ_k parameters for $k = 2, 3, 4$, along with a generic satiation parameter α . As in most of the work in

208 the literature, all the model specifications that we estimated displayed an extremely small value of α for
 209 which we could not reject the null hypothesis that it was equal to zero, where, with $\alpha \rightarrow 0$, the utility form
 210 collapses to a log utility formulation (cf. Bhat, 2008) with:

$$212 \quad U(x) = \psi_1 \ln(x_1) + \sum_{k=2}^K \gamma_k \psi_k \left(\frac{x_k}{\gamma_k} + 1 \right) \quad (3)$$

213 This formulation implies that direct utility increases with additional units of consumption in a logarithmic
 214 fashion, i.e. with diminishing returns. The only parameters relating to satiation that we estimate are the
 215 γ_k terms, which can be interpreted in terms of how long the activities of *Feed*, *Move* and *Short move* are
 216 performed for.

217 The probability that an individual (bear) chooses a specific vector of time allocations $x_1^*, x_2^*, \dots, x_M^*, 0, \dots, 0$,
 218 where M of the K activities are performed in a given day, is given by:

$$219 \quad P(x_1^*, x_2^*, \dots, x_M^*, 0, \dots, 0) = \frac{1}{p_1} \frac{1}{\sigma^{M-1}} \left(\prod_{m=1}^M f_m \right) \left(\sum_{m=1}^M \frac{p_m}{f_m} \right) \left(\frac{\prod_{m=1}^M e^{v_i/\sigma}}{\left(\sum_{k=1}^K e^{v_k/\sigma} \right)^M} \right) (M-1)!, \quad (4)$$

220 where σ is an estimated scale parameter and where $f_m = \left(\frac{1-\alpha}{x_m^* + \gamma_m} \right)$.

221

222 **3.3. Specification for our study**

223

224 3.3.1 The discrete choice

225 As mentioned above, the baseline utility ψ_k broadly captures the “discrete choice”, i.e., the likelihood of
 226 performing an activity. In the present study, this is composed of a constant δ_k and additive shifts for each
 227 of the covariates we consider. Differently from discrete choice models, the δ constants in the MDCEV
 228 model are influenced by both the discrete and the continuous parts of the model, and this explains the
 229 fact that the values for the constants for the non-base activities can be positive even though the base
 230 alternative (*Rest*) is conducted more often.

231

232 3.3.2 The continuous choice

233 Analogously, in the case of the satiation component, we estimate one γ_k for each activity but allow for
 234 heterogeneity as a function of covariates, i.e., measuring the impact of the covariates on the continuous
 235 choice. Differently from the case of the baseline utility, the parameterisation of γ_k was operationalised in
 236 a multiplicative fashion. For example, the overall satiation from activity k was expressed as:

$$237 \quad \gamma_k = \gamma_{k,base} * \prod_{i=1}^I f_k(z_i) \quad (5)$$

238 where $\gamma_{k,base}$ is a constant for activity k (reported as “Core parameters” in Table 3), z_i is one of a set of I
 239 covariates. For categorical variables, such as sex, we set a base category, and estimate a multiplier for
 240 others, meaning that for a covariate with L levels, we use $f_k(z_i) = \sum_{l=1}^L \kappa_{ikl} \cdot (z_i == l)$, where $(z_i == l)$
 241 is equal to 1 if z_i takes the l^{th} level, and zero otherwise, and where we fix $\kappa_{ikl} = 1$ for one category. For

242 continuous variables, we use $f(z_i) = \left(\frac{z_i}{\bar{z}_i} \right)^{\lambda_{ik}}$, where the estimate of λ_{ik} captures the non-linearity.

243 We retained those effects in the model where the κ_{ikl} multipliers are significantly different from 1, or
 244 where the elasticity parameters λ_{ik} were different from 0, implying that $f(z_i)$ is different from 1.

245

246 3.3.3 Explanatory variables

247 In our MDCEV application, we have tested the effect of a range of variables that have been demonstrated
 248 by a large amount of ecological literature to be related to brown bears’ behaviour. These mainly relate to
 249 environmental/climatic factors, characteristics of the animal and potential human impacts. Here and in
 250 the Results section, we only describe the variables for which statistically significant effects on activity

251 choice and duration were found. These include the bears age, sex, and two climatic variables. The full list
 252 of the variables used in the final model specification is reported in Table 2.

253
 254

Table 2 – Explanatory variables used in the model

Variable	Type of variable	Description
Sex	Categorical	1= Solitary adult female 0= Solitary adult male
Age	Categorical	1= 4-8 years old 2= 9-15 years old 3= Older than 15
Average daily temperature	Continuous	Range: 4.63-22.1 °C Mean: 13.12 °C
Average daily precipitation	Continuous	Range: 0-33.47 mm Mean: 2.94 mm
Daily duration of daylight*	Continuous	Range: 12.89-18.36 h Mean: 15.73 h
Daily duration of twilight*	Continuous	Range: 3.24-7.23 h Mean: 4.86 h
Daily duration of night*	Continuous	Range: 0-7.86 h Mean: 3.40 h
Hunting season	Categorical	1= hunting season 0= not hunting season

255 * Calculated for the central location of our study area (Tackåsen, Sweden: 61.5N, 15.05E)

256
 257 The climatic variables related to temperature and precipitation were obtained from the Swedish
 258 Meteorological and Hydrological Institute (SMHI). In particular, station-specific time series within the study
 259 area were converted using the R package “mba” (Finley & Banerjee, 2014) to interpolated raster series
 260 with a 5-day temporal resolution and a 5 km spatial resolution. To account for variation across the study
 261 area, the raster values of these climate variables associated with the area inhabited by each bear were
 262 averaged using a circular home range with a sex-specific average home range diameter (Bischof et al.,
 263 2018). In our models, we tested minimum, maximum and average temperature. Since the focus of our
 264 study is to understand how animals allocate time to different activities across 24 hours, the mean
 265 temperature seemed to be the most appropriate measure. The R (R Core Team, 2020) library “mapproj”
 266 (Bivand and Lewin-Koh, 2020) was used to determine the length of day, night, and twilight for every day
 267 during the berry season. We consider as “Daylight” the period between sunrise and sunset, “twilight” as
 268 the time period between sunset and nautical dusk (i.e., when the sun moves to 12 degrees below the
 269 horizon in the evening) and from nautical dawn (when the sun moves to 12 degrees below the horizon) to
 270 sunrise. The remaining time (i.e., between nautical dusk and dawn) is defined as “night”. Due to high
 271 correlation between *Daylight* and *Night duration* only the former variable has been retained in the model.
 272 This variable is also a good proxy for the time elapsed so far in the berry season, as the *Daylight duration*
 273 changes linearly in the study period. The dummy variable taking value 1 for the time between 21 August –
 274 15 September is used to test whether the hunting season affects the bears’ behaviour. The MDCEV model
 275 was estimated using the “apollo” package (Hess & Palma, 2019) in R (R Core Team, 2020).

276
 277 **4. Model results**

278

279 The final model results are presented in Table 3. We look first at the baseline parameters that relate
 280 primarily to the discrete choice, before looking at the satiation parameters that relate more to the
 281 continuous choice. The baseline parameters have limited interpretation (Bhat, 2018).
 282

283 Table 3 – Model results

	Coefficient	Estimate	Rob. t-ratio (0)	Rob. t-ratio(1)
Baseline parameters	α	0 (fixed)	NA	NA
	γ_{feed}	4.099	15.174	11.472
	γ_{move}	7.062	12.609	10.824
	$\gamma_{short\ move}$	4.107	19.403	14.679
	δ_{feed}	0.668	2.289	-1.135
	δ_{move}	-2.568	-9.067	-12.598
	$\delta_{short\ move}$	-2.175	-17.939	-26.186
Shifts in the δ parameters	Age 16+ <i>feed</i>	0.081	2.729	-30.781
	Age 16+ <i>move</i>	-0.361	-9.928	-37.394
	Female <i>short move</i>	0.263	7.419	-20.713
	Daylight duration <i>feed</i>	-1.294	-7.649	-13.556
	Daylight duration <i>move</i>	0.705	4.195	-1.75
	Daylight duration <i>short move</i>	0.342	4.572	-8.814
	Hunting period <i>feed</i>	-0.142	-4.788	-38.401
	Hunting period <i>move</i>	0.047	1.051	-21.048
	Temperature <i>feed</i>	0.008	1.832	-208.204
	Temperature <i>move</i>	-0.008	-3.783	-429.586
	Precipitation <i>move</i>	-0.052	-3.856	-77.558
Multipliers of the γ parameters (κ_{ik})	Female <i>feed (base=male)</i>	1.256	16.827	3.431
	Female <i>short move (base=male)</i>	0.908	17.357	-1.738
	Hunting period <i>move (base=outside hunting period)</i>	0.943	10.18	-0.608
Exponential term in γ multipliers (λ_{ik})	Daylight duration <i>feed</i>	1.804	5.071	2.261
	Daylight duration <i>move</i>	-0.695	-1.415	-3.451
	Precipitation <i>move</i>	0.013	2.414	-182.821
	Temperature <i>feed</i>	-0.045	-0.375	-8.624
Scale	Scale parameter (σ)	0.272	64.741	-173.423

284 *Model performance:* Final LL: -32116.1; AIC: 64282.21; BIC: 64443.52

285
 286 **4.1. Shifts in the δ parameters**

287 We found find that within a 24-hour time span, older bears (aged 16 years or over) are more likely to *Feed*
 288 and less likely to *Move* compared to younger ones (cf. Table 3). The coefficient “Female short move” points
 289 to the fact that females are more likely to move than solitary males, suggesting that they are more likely
 290 to do a combination of activities in the given time frame. When the day lasts longer (i.e., at the beginning
 291 of the berry season), the baseline utility of moving and short moving increases, while that of foraging
 292 decreases.

293 The start of the hunting season has a significant effect on bear behaviour, i.e., the bears are more likely to
 294 move and less likely to feed (cf. Table 3). Interestingly, since in our application all activities are modelled
 295 jointly, we see that decreased probability of feeding is accompanied by increased probability of *moving*,
 296 with *short move* being unaffected.

297 We tested for the effect of daily temperature in different forms (i.e., min, max, range, average), and daily
 298 average temperature had the highest explanatory power and was retained in the final model specification.
 299 A higher likelihood of *Feed* and a lower likelihood of *Move* are associated with increases in temperature
 300 (Table 3).

301 The shift of the δ parameters as a consequence of precipitation (in mm) shows that an increase in this
 302 variable is linked with a lower likelihood to *Move*.

303

304 *4.2. Multipliers of the γ parameters*

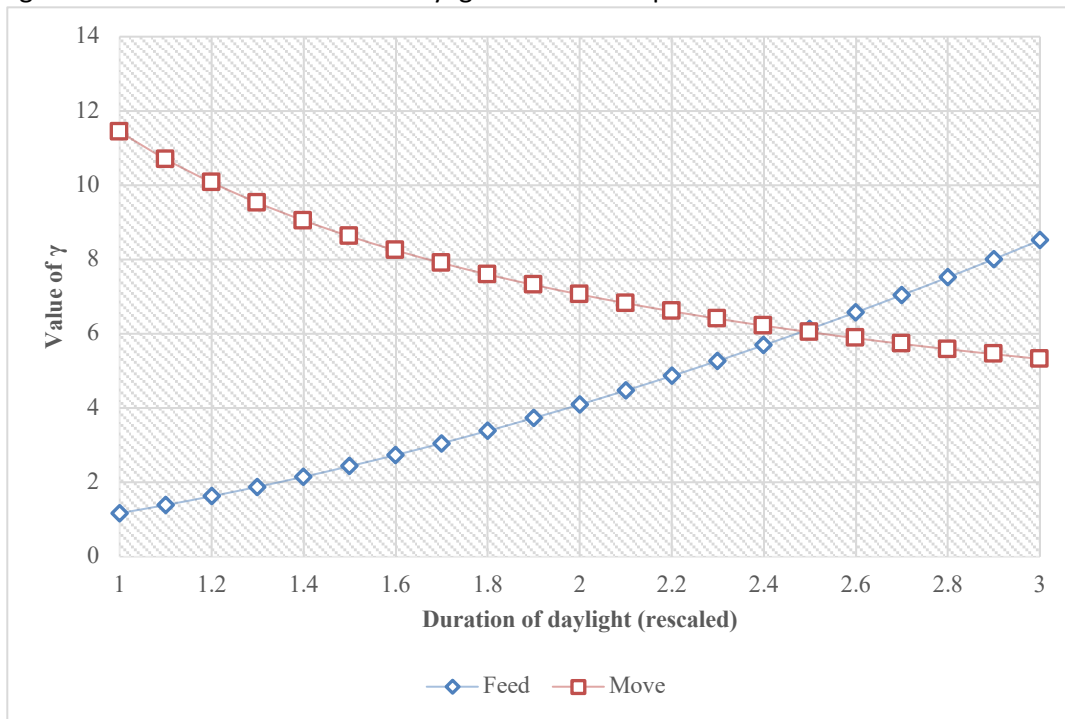
305

306 The results related to the parameterisation of the γ parameters are shown in the bottom part of Table 3.
 307 Due to the fact that they enter the satiation equation multiplicatively, these parameters have a significant
 308 impact if they are significantly different from 1. Female bears are found to spend longer time in *Feed* and
 309 *Short move* as opposed to males. We also observe that during the hunting period, bears spend more time
 310 moving as opposed to outside of the hunting season. This indicates that not only the probability of *Move*
 311 increases due to hunting but also time spent in this activity increases.

312 The last set of parameters in Table 3 measure the sensitivity of the overall satiation to changes in
 313 continuous variables. Figure 1 shows the impact of the duration of daylight on the satiation from *Feed* and
 314 *Move*, given the estimated values of λ_{ik} .

315

316 Figure 1: Effect of the duration of daylight on satiation parameters



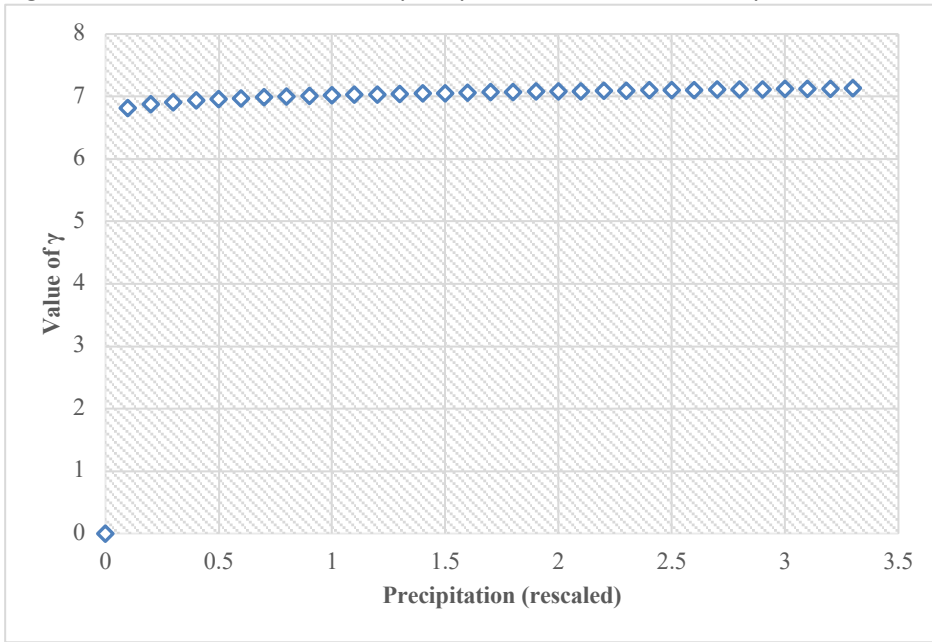
317

318 A positive value of this exponential term (cf. "Daylight duration *feed*") implies that as the duration of
 319 daylight increases, the activity (*Feed*) will be performed for longer. The opposite is true for *Move*. This
 320 implies that at the beginning of the berry season, if bears engage in *feeding*, they will do so for longer than

321 at the end of the berry season. The opposite is true for *Move*. This is likely related to the fact that the end
322 of the berry season corresponds with the hunting season, when bears feed less and move more.

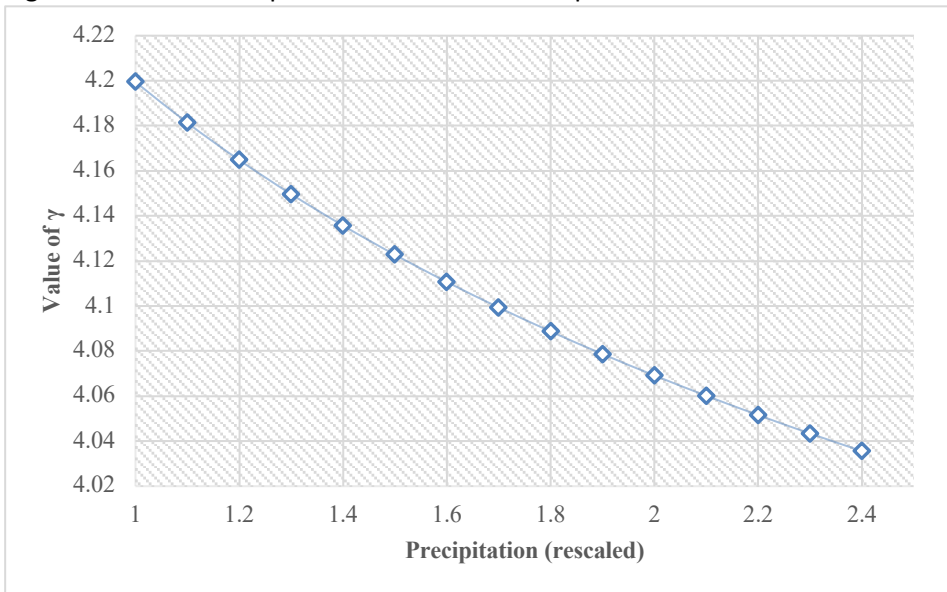
323 The magnitude of the changes in satiation is determined by the value of the estimated parameters and
324 the baseline value ($\gamma_k base$). Precipitation positively affects the value of the satiation parameter for *Move*,
325 although its small value results in a slowly increasing trend (cf. Figure 2). Higher temperatures imply a
326 smaller amount of time spent foraging (cf. Figure 3).
327

Figure 2. Effect of the amount of precipitation on the satiation parameter for *move*



328
329
330

Figure 3. Effect of temperature on the satiation parameter for *Feed*



331
332
333

334 4.3 Model validation

335 In order to obtain a measure of how well our proposed model performs at capturing the trends in the data,
336 we have carried out a prediction of the shares of days on which a given activity is carried out, as well as of
337 the activity duration and compared it with the data. This is what is sometimes referred to as “base”
338 prediction, i.e. applying the model without introducing any changes to the data. As shown in Table 4, the
339 differences between the discrete and continuous choice statistics in the data and in the prediction are
340 limited, and the RMSE is low in both cases, indicating that the model adequately replicates the behaviour
341 in the data.
342

Discrete choice (share of days with non-zero time in activity)			Continuous choice (hrs spent)	
Activity	Share in data	Predicted share	Average duration in data	Average predicted duration
Rest	1.00	1.00	8.73	8.56
Feed	0.99	0.99	6.00	5.95
Short Move	0.96	0.97	3.80	3.92
Move	0.96	0.95	5.47	5.58
RMSE= 0.12			RMSE=0.01	

343

344 5. Discussion

345 In this paper, we present a proof-of-concept for the application of advanced econometrics models to
346 understand the behaviour of wild animals. Addressing the limitations of previous studies, we produced a
347 model of brown bear behaviour where we model engagement in multiple activities, considering not only
348 which activities are conducted but also the amount of time invested in each. We showed how both the
349 discrete and continuous outcomes are related to the characteristics of the animals themselves as well as
350 to environmental variables and external factors (i.e. hunting). While the MDCEV modelling framework
351 adopted was not previously used to improve the understanding of animal behaviour, we demonstrated
352 how this tool can be suitable for our investigation by obtaining results in line with existing work as well as
353 providing new insights made possible by the specific model structure.
354

355 To start with, our classifications of the different activities inferred via the GPS relocation data allowed us
356 to unveil behavioural patterns correlated with bear characteristics. We showed that female bears are more
357 likely to engage in the *Short Move* activity and do so for longer compared to males. This is an example of
358 a finding allowed by the MDCEV model, i.e., capturing discrete and continuous behaviour contextually. As
359 explained in the Methods section, this activity is most likely to be made up of a mixture of different
360 activities, implying that female bears have a more varied pattern of behaviour within each observed time
361 slot. We have also shown that older bears are more likely to engage in foraging rather than travelling long
362 distances.
363

364 Capturing both the discrete and the continuous outcomes allowed us to gain a more complete picture of
365 behaviour, as we understand which characteristics are associated with activity duration instead of only
366 providing insights on which activities are performed. A key result of our model is that during the hunting
367 season, bears are less likely to feed and more likely to move – and do so for longer periods of time. This
368 finding is in line with existing ecological literature which demonstrated that apex predators may perceive

369 and respond to hunting risk similar to that of prey responding to a natural predator (Ordiz et al. 2011;
370 Brook et al. 2012) , forcing them to increase vigilance at the expense of foraging.

371
372 Our results conform well with Hertel et al. (2016b), who found that bears effectively responded to
373 variation in risk during the day by decreasing their foraging activity in the morning hours of the hunting
374 season and not changing their behaviour in the afternoon foraging bout, indicating that bears did not
375 allocate antipredator behaviour to times of comparably lower risk. On the other hand, contrary to the
376 existing literature, which has focused on modelling one activity (Hertel et al. 2016a, Hertel et al. 2016b,
377 McLellan and McLellan 2015, Steyaert et al. 2016), the MDCEV framework allows us to directly identify the
378 trade-offs in bears behaviour. Bears reduce risk from hunting by increasing probability of Move and its
379 duration, this happens at the cost of the forage activity. This shows that hunting affects the energy budget
380 of bears in a more complex way than identified in the literature (Sahlen et al. 2015, Steyaert et al. 2016,
381 Hertel et al. 2016b), that is, not just by lowering energy intake by decreasing foraging activity, but also by
382 increasing energy expenditure as they are more likely to Move and do it for a longer period. This shows
383 that hunting season is highly costly for bears in terms of energy balance, as they are less likely to feed and
384 more likely to move at this crucial time for their energy intake, i.e., before hibernation. The additive effect
385 of reduced forage intake and higher engagement in movement is likely to result in poorer body condition
386 upon den entry and may ultimately reduce winter hibernation survival and reproductive success. This
387 result potentially enables managers and conservationists to better understand the ecology of this species
388 as well as how people affect its behaviours and what are the consequences on energy expenditure and
389 survival.

390 The sign and magnitude of the λ_{ik} parameters combined with the shift in the δ parameters (i.e. *Daylight*
391 *duration Feed*) showed that as *Daylight duration* decreases, bears are more likely to *Feed* but will carry out
392 this activity for a shorter period of time. Exactly the opposite pattern is true for *Move*, that is as *Daylight*
393 *duration* decreases bears are less likely to *Move* but will carry out this activity for a longer period of time
394 if it is conducted. The opposing trends displayed as time advances in the berry season indicates that bears
395 constantly change (adapt) their foraging strategy during the studied period. A possible explanation of the
396 observed phenomena is that at the beginning of the berry season, food is scarcer (or of lower quality) and
397 once bears find a feeding place, they will carry out the *Feed* activity for longer, as there are no good
398 alternatives available; whereas later in the berry season, when more berry species are available and food
399 is more abundant, bears will travel between high quality spots. This interpretation is in line with what has
400 been reported by Hertel et al. (2016b), who showed that in the berry season, bears are selective and
401 navigate in the forest landscapes by using areas of higher than average berry abundance.

402
403 Overall, this study is innovative in two ways: it demonstrated the application of a state-of-the art
404 discrete-continuous model to a new field of research, and it shows not only that results which are intuitive
405 and in line with the literature can be obtained, but also that new insights can be added due to the more
406 comprehensive approach looking at different activities and at both choice of activities and activity
407 duration. In particular, this approach allowed us to distinguish cases of zero time in an activity (i.e. a corner
408 solution) from small non-zero times, and that is allows for a study of satiation, i.e. non-linear gains in utility
409 from additional consumption. In terms of specific conclusions for our case study, the proposed approach
410 allowed us to observe that the disturbance due to hunting affects energy expenditure as it impacts on
411 multiple activities, namely feeding and moving. It also allowed us to understand key differences across the
412 animals. For example, we found that female bears not only engage in different types of activities (e.g.
413 Short Move) but also do so for different durations with respect to their male counterpart.

414
415 Like any study, we acknowledge that our work has a number of limitations. The classification of
416 the moving activity in two separate activities is subjective and while it helps with the interpretation of

417 results, it might not be ecologically accurate, and could impact our findings. Especially in the case of *Feed*
418 and *Move*, our activity classification is rather coarse with respect to studies using human data collected
419 with smartphone or GPS trackers that can virtually capture any movement. This is a result of working at
420 the level of one-hour data. Finer temporal resolution (i.e., GPS fixes every 30min or 15min) would allow
421 for more reliable classification and understanding of behaviours, as it is likely that bears undertake a
422 mixture of activities during 1h intervals. At the same time, this would be very battery-intensive and would
423 require capturing and re-releasing bears more often to change the batteries, which is expensive and may
424 not be possible due to animal welfare concerns. Additional improvement of the method could be achieved
425 by linking the GPS data with other high-resolution data such as, 3D accelerometers, heart-rate sensors.
426 Moreover, as this study aims to be a proof-of-concept, we only used a sub-set of the available data. Using
427 a larger and more comprehensive dataset could unveil further patterns of behaviour, for example the ones
428 of younger animals and females with dependent offspring. We leave these developments to future work,
429 with the main aim of the present paper being to present the method and its potential.

430

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437

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