

# MoveFormer: a Transformer-based model for step-selection animal movement modelling

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### Abstract

The movement of animals is a central component of their behavioural strategies. Statistical tools for movement data analysis, however, have long been limited, and in particular, unable to account for past movement information except in a very simplified way. In this work, we propose MoveFormer, a new step-based model of movement capable of learning directly from full animal trajectories. While inspired by the classical step-selection framework and previous work on the quantification of uncertainty in movement predictions, MoveFormer also builds upon recent developments in deep learning, such as the Transformer architecture, allowing it to incorporate long temporal contexts. The model predicts an animal’s next movement step given its past movement history, including not only purely positional and temporal information, but also any available environmental covariates such as land cover or temperature. We apply our model to a diverse dataset made up of over 1550 trajectories from over 100 studies, and show how it can be used to gain insights about the importance of the provided context features, including the extent of past movement history. Our software, along with the trained model weights, is released as open source.

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## 62    1 Introduction

63    The movement of animals is a central component of their behavioural strategies  
64    to best exploit the landscape they live in, to find a mate or to avoid predators,  
65    for instance. The role that these movements have beyond the individual,  
66    for instance in shaping animals' ecosystem impacts, is clear. Accordingly, and  
67    thanks to the technological developments that are allowing to collect more de-  
68    tailed movement data on more individuals and species each day, the study of  
69    animal movement has become an important goal of ecology [1].

70    For a long time, however, statistical tools to analyze movement data were  
71    lacking or limited. Over time, though, purely pattern-based descriptions (e.g.  
72    home-range analyses) have been complemented by regression models allowing  
73    to infer the effects of spatio-temporal features on movement. *Step-selection*  
74    *function* (SSF) models, which compare actual movement steps with realistic  
75    candidate ones, are ~~one of such models and have de facto become the established~~  
76    ~~approach to analyse animal trajectories [2–4]. They are now routinely used to~~  
77    ~~infer and quantify the effect of environmental variables such as, for instance,~~  
78    ~~such as~~ land cover or temperature, on animal trajectories [2–4].

79    However, an animal's movement is likely to be driven not only by spatio-  
80    temporal environmental features, but also by some internal knowledge and rules  
81    that are unobservable directly. The importance of memory and of an animal's  
82    familiarity with places is increasingly recognized [5–7], and familiarity is usu-  
83    ally incorporated into SSF models using a *previously visited* yes/no variable,  
84    or a *time-spent* variable, often calculated over an arbitrary time window [8, 9].  
85    Memory of places and their characteristics can also lead to routine movement  
86    behaviours. Traplining, in which an individual travels to the same places in the  
87    same order, is rare, but it is clear from visual inspection of animal trajectories  
88    that many animals display some form of routine movement behaviours. But  
89    for traplining, which has received a lot of interest, the study of routine move-  
90    ment behaviour has remained extremely limited [10]. Riotte-Lambert et al. [11]  
91    showed how conditional entropy, calculated using the information on visits to  
92    patches, could be used as a metric of routine in movement. That metric has  
93    not been used much since then, possibly because the need to determine sites  
94    may render its application difficult on data collected in nature, where patches  
95    can be difficult to determine, be diffuse, or not exist at all. Further work is  
96    needed to describe and explain routine movements, which result from the in-  
97    teraction between memorized knowledge, movement rules and environmental  
98    context. Additionally, we are not aware of any work that has focused on how  
99    to incorporate complex information about past movement and environmental  
100   context into predictive models of animal movement, although it should, by def-  
101  inition, improve predictions. The question: 'To what extent past movements  
102   inform where an animal is likely to go next?' remains open.

103   The classic implementation of the SSF framework appears unsuitable to  
104   address this difficult question. We therefore developed a new type of model that  
105   we named MoveFormer. We conserved the conceptual attractiveness of SSF, but  
106   built on the most recent developments in deep learning to embed the information  
107   about current and past animal location, movement and environmental context.

108   Our contribution is threefold. First, we propose a model that learns to best  
109   predict the next step of a movement trajectory based on a given context length,  
110   i.e. a given time-window of information about the past. Second, the proposed

111 approach is flexible enough to allow each step in the context to be defined not  
112 only by the locations of the start and end points, but also by any kind of fea-  
113 tures that could be relevant, in particular environmental variables. Third, we  
114 show how the model can be used to gain insights about the importance of the  
115 provided context, both in terms of the extent of the past that it is useful to  
116 know, and in terms of what kinds of information are most ecologically relevant  
117 to predict an animal’s movement. We demonstrate this by comparing the pre-  
118 dictions, via information-theoretic metrics and prediction accuracy, for different  
119 context lengths or with randomized features. Model training and analyses are  
120 conducted on a dataset made up of over 1550 trajectories from over 100 studies,  
121 encompassing various species within mammals, birds and reptiles.

122 ~~The We release the~~ MoveFormer source code,<sup>1</sup> including code for data ~~pre-processing~~  
123 ~~pre-processing~~ and evaluation, as well as complete hyperparameter settings ~~,is~~  
124 ~~available online.~~<sup>2</sup> We also release and the weights of the trained models.<sup>2</sup> See  
125 ~~Section 7 for more information.~~

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<sup>1</sup><https://github.com/cifikao/moveformer>  
<sup>2</sup>

# <sup>126</sup> Materials and methods

## <sup>127</sup> 2 Data

<sup>128</sup> In this section, we describe our sources of data, specifically: movement data (tra-  
<sup>129</sup> jectories consisting of latitudes, longitudes and timestamps), geospatial variables  
<sup>130</sup> (associated with locations), and taxonomic classification information (associated  
<sup>131</sup> with each animal).

### <sup>132</sup> 2.1 GPS location data

<sup>133</sup> Our main source of location data is Movebank<sup>2</sup> [12], an online repository for an-  
<sup>134</sup> imal movement data. The location data in Movebank is presented as latitude/-  
<sup>135</sup> longitude pairs along with UTC timestamps and is grouped into trajectories  
<sup>136</sup> (*deployments*) and associated with (occasionally missing) metadata such as a  
<sup>137</sup> taxon name, sex, and date of birth. We used the Movebank API to retrieve data  
<sup>138</sup> from GPS sensors for all 269 studies that were available<sup>3</sup> for download under a  
<sup>139</sup> Creative Commons<sup>4</sup> license (CC0, CC BY and CC BY-NC), obtaining 13 577  
<sup>140</sup> trajectories comprising a total of 197 million observations (location events). We  
<sup>141</sup> subsampled the trajectories (splitting them into segments when necessary) so  
<sup>142</sup> that observations occur at midnight and at noon (according to local mean time)  
<sup>143</sup> with a tolerance of  $\pm 3$  h and so that the time difference between consecutive  
<sup>144</sup> observations is 9 to 15 h. We discarded trajectory segments shorter than 120  
<sup>145</sup> observations, leaving us with 1440 trajectories from 98 different studies [13–  
<sup>146</sup> 165]. See Table 4 in [the appendix](#) [Supplementary information](#) for the full list of  
<sup>147</sup> studies and their licenses.

<sup>148</sup> We added [unpublished](#) data from 4 more studies, collected by one of us (S.C.-  
<sup>149</sup> J). These are GPS data from plains zebras [\[166\]](#) and African elephants [\[167\]](#),  
<sup>150</sup> collected in Hwange National Park ([Zimbabwe](#))[in Zimbabwe](#), and GPS data from  
<sup>151</sup> plains zebras and blue wildebeest, collected in Hluhluwe-iMfolozi Park ([South](#)  
<sup>152</sup> [Africa](#))[in South Africa and yet unpublished](#). After subsampling and filtering  
<sup>153</sup> as in the case of the Movebank data, we obtained 73 trajectories.

<sup>154</sup> [See Section 7 for information about data availability.](#)

<sup>155</sup> The final dataset contains about 1 million observations from 1506 individu-  
<sup>156</sup> als, grouped into 1513 trajectories with a median length of 408 [observations](#)  
<sup>157</sup> ([corresponding to around 204 days](#)). We performed a train/validation/test  
<sup>158</sup> split, making sure that 1) the validation and test sections contain only frequent  
<sup>159</sup> species([with at least 10 members in the full dataset](#)), and 2) each individual  
<sup>160</sup> appears in exactly one split.<sup>5</sup> Table 1 details the amounts of data by section  
<sup>161</sup> and by taxonomic classification and Fig. 1 shows the geographical distribution.

<sup>162</sup> During training and evaluation, we additionally split each trajectory into  
<sup>163</sup> segments of length  $N_{\max} = 500$  and subsequently consider each of these segments

<sup>2</sup>[www.movebank.org](http://www.movebank.org)

<sup>3</sup>as of [15-15](#) February 2022

<sup>4</sup><https://creativecommons.org/>

<sup>5</sup>[Specifically, after filtering out infrequent species \(with less than 10 members in the full dataset\), we randomly assign 4% of the remaining individuals to the validation set, another 5% to the test set, and the rest to the training set.](#)

section	#spec	#ind	#traj	#obs	
train	61	1383	2786	915618	
val	17	53	133	50984	
test	21	70	133	40334	
class	order	#spec	#ind	#traj	#obs
<i>Aves</i>		39	895	1915	640420
Accipitriformes		10	138	315	65189
Anseriformes		10	169	210	69386
Bucerotiformes		1	5	6	4448
Cathartiformes		1	12	34	8965
Charadriiformes		10	310	585	235722
Ciconiiformes		2	249	751	253541
Gruiformes		1	1	1	189
Passeriformes		1	3	3	474
Pelecaniformes		1	6	6	1764
Struthioniformes		1	1	1	221
Suliformes		1	1	3	521
<i>Mammalia</i>		15	439	845	290789
Artiodactyla		7	329	660	218698
Carnivora		6	41	51	12404
Perissodactyla		1	26	34	25582
Proboscidea		1	43	100	34105
<i>Reptilia</i>		6	58	116	40353
Testudines		6	58	116	40353

Table 1: Number of species (#spec), individuals (#ind), trajectories (#traj), and observations (#obs) in the dataset. The upper part of the table displays the counts for each section of the sections of the dataset (training, validation, and a breakdown by taxon). The lower part details the counts for each taxonomic class and order (aggregated over all 3 sections).

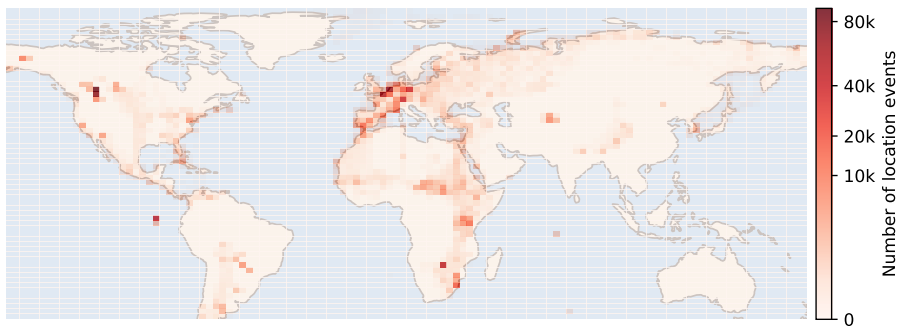


Figure 1: The geographical distribution of the observations in the dataset.

164 as a separate trajectory.<sup>6</sup>

## 165 2.2 Taxon vectors

166 Each trajectory in our data is associated with a taxon name (most commonly  
167 the animal's species). To obtain a dense vector representation of the taxon,  
168 we look up its Wikipedia article and retrieve the associated 100-dimensional  
169 embedding vector from Wikipedia2Vec [168].

170 A property of Wikipedia2Vec is embeddings, derived from the text of Wikipedia  
171 articles as well as the link structure of Wikipedia, have the property that em-  
172 beddings of semantically similar related entities are placed close together in the  
173 embedding space. To illustrate that this extends, to some degree, to similarity  
174 between relationships between biological species, we display in Fig. 2 the PCA  
175 (principal component analysis) projections of species embeddings, labeled by  
176 higher taxonomic ranks. We also measured the cosine similarity between all  
177 pairs of embeddings and found it to be correlated with the number of common  
178 ancestors of the two species in the taxonomic hierarchy (Spearman  $\rho = 0.68$ ).

179 Overall, the Wikipedia2Vec embeddings appear to meaningfully encode a  
180 species' position in the phylogeny. Hence, we speculate (though we do not test  
181 this in the present work) that their inclusion in the model should help this model  
182 to generalize to species that are not present in the training data, at least as long  
183 as they are sufficiently similar to those that are.

## 184 2.3 Geospatial variables

185 The proposed model is powerful enough to account not only for each trajectory  
186 intrinsic dynamics but also for any *third-party* additional information that may  
187 be available as covariates. In order to illustrate this, we augment each trajectory  
188 data point with exogenous information. For each location, we retrieve the fol-  
189 lowing geospatial variables, which could be ecologically relevant, from publicly  
190 available raster data:

- 191 • 2009 Human Footprint, 2018 Release [169, 170] (resolution:  $\sim 1 \text{ km}$ ); we  
192 normalize the values between 0 and 1 and sample them with bilinear  
193 interpolation-use linear interpolation when retrieving the values by location;
- 194 • 19 bioclimatic variables from WorldClim 2.1 [171] (resolution:  $\sim 1 \text{ km}$ ),  
195 listed in Table 6 in Supplementary information; we standardize the values  
196 (zero mean, unit variance) and use nearest neighbor interpolation;
- 197 • land cover classification (23 classes) from Copernicus Global Land Service,  
198 version 3.0.1, epoch 2015 [172] (resolution:  $\sim 100 \text{ m}$ ); we use a one-hot  
199 encoding and nearest neighbor interpolation.

6 A Transformer typically supports sequences up to a maximum length, determined by the lengths of the sequences encountered during training. Training on full-length sequences is usually not feasible due to its computational cost, but also because the training dataset is likely to only contain a single sequence of exactly the full length, preventing successful generalization up to that length. Setting an upper length limit close to the median trajectory length is a convenient way to avoid these issues.

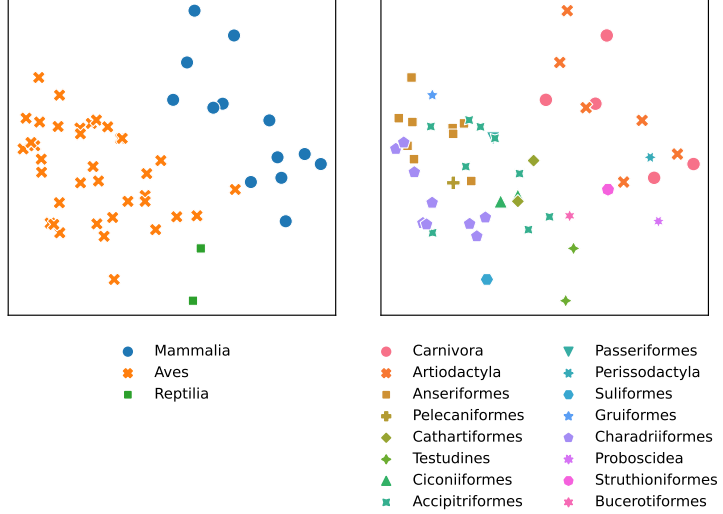


Figure 2: A PCA projection of Wikipedia2Vec embeddings of species, labeled by class (left) and order (right). The  $x$  and  $y$  axis correspond to the first two principal components, respectively. Note that since Wikipedia2Vec is a distributed representation, its dimensions are not easily interpretable and have no meaningful units.

### 200 3 Model

201 Formally, we consider the dataset as composed of *trajectories*, where a trajectory<sup>7</sup>  $\xi_{1\dots N}$  of length  $N$  consists of locations  $x_{1\dots N}$ , corresponding to timestamps  
202  $t_{1\dots N}$ , and any associated variables  $z_{1\dots N}$ , i.e.  $\xi_n = (x_n, z_n, t_n)$  as described  
203 above. Our main goal is to estimate a model for the next-step prediction task,  
204 i.e. for any given  $n \in \{1, \dots, N\}$ , predict the next location  $x_{n+1}$  from the tra-  
205 jectory prefix  $\xi_{1\dots n}$  and the next timestamp  $t_{n+1}$ .

206 As a fundamental use case, we are interested in analyzing the effect of avail-  
207 able past context on the prediction of  $x_{n+1}$ . Specifically, for a varying *context*  
208 *length*  $c \in \{1, \dots, c_{\max}\}$  (where  $c_{\max}$  is an arbitrary constant), we wish to study  
209 the behavior of the prediction of  $x_{n+1}$  given  $\xi_{n-c+1\dots n}$  and  $t_{n+1}$ . Hence, we are  
210 in fact interested in a model accepting as input any *trajectory segment* of length  
211 at most  $c_{\max}$ , and predicting the next location.

212 We adopt a step-selection function modelling approach [2, 4], based on se-  
213 lecting the end-point location of a step from a set of candidates. Specifically, for  
214 a position  $n+1$  within a trajectory, given an associated timestamp  $t_{n+1}$ , a set of  
215 candidate locations  $x_{n+1}^{(1\dots K)}$  and associated variables  $z_{n+1}^{(1\dots K)}$ , we are interested  
216 in estimating a probability distribution over the candidates:

$$P(y_{n+1} = i \mid \xi_{1\dots n}, t_{n+1}, x_{n+1}^{(1\dots K)}, z_{n+1}^{(1\dots K)}), \quad (1)$$

218 where  $i \in \{1, \dots, K\}$ .

---

<sup>7</sup>We use  $\xi_{1\dots N}$  as shorthand notation for the sequence  $\xi_1, \xi_2, \dots, \xi_N$ . Note that  $N$  may be different for each trajectory in the dataset.

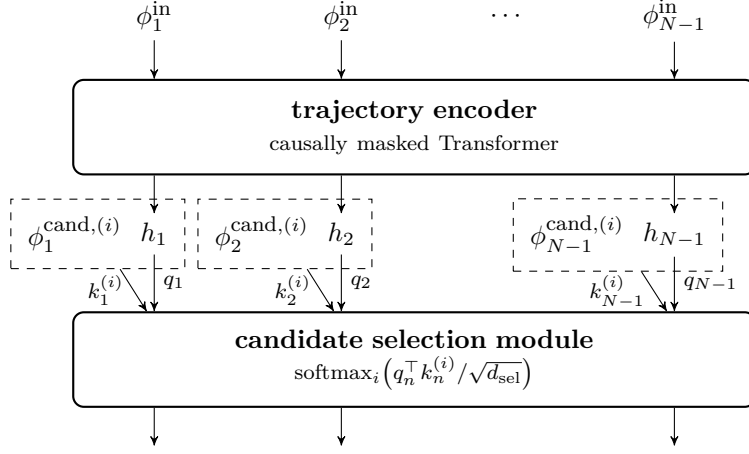


Figure 3: The high-level architecture of MoveFormer. The input to the trajectory encoder is a sequence of embedding vectors  $\phi_1^{\text{in}}, \phi_2^{\text{in}}, \dots, \phi_{N-1}^{\text{in}}$ , each corresponding to a different data point (location-timestamp pair) in the trajectory. The encoder outputs a sequence of vectors  $h_1, \dots, h_{N-1}$ ; the causal masking in the encoder causes each  $h_n$  to encode only the inputs up to position  $n$ , i.e.  $\phi_1^{\text{in}}, \dots, \phi_n^{\text{in}}$ . This representation is then fed to the candidate selection module, which uses it as *queries* in an attention mechanism that assigns probabilities to different candidate locations. Both the input embeddings  $\phi_1^{\text{in}}, \dots, \phi_{N-1}^{\text{in}}$  and the candidate embeddings  $\phi_n^{\text{cand},(i)}$  are computed through embedding layers which are not displayed here but described in Section 3.1.1.

219 We propose to model this distribution using a deep neural network, consisting  
 220 of a *Transformer* [173] *encoder* and a *candidate selection module*, as depicted  
 221 in Fig. 3. The role of the Transformer is to encode the trajectory up until posi-  
 222 tion  $n$ , i.e.  $\xi_1, \dots, \xi_n$  along with the timestamp for the next observation,  $t_{n+1}$ . The  
 223 candidate selection module then encodes each candidate  $x_{n+1}^{(i)}$  and employs an  
 224 attention mechanism to compute a probability distribution over the candidates.  
 225 The model is described in detail in Section 3.1, followed by our choice of input  
 226 representation in Section 3.2.

227 In order to train and evaluate this model, we also need a way to generate  
 228 suitable candidate locations  $x_{n+1}^{(i)}$ . We use a simple but general method em-  
 229 ploying quantile-based modelling of turning angles and movement distances, as  
 230 detailed in Section 3.3.

### 231 3.1 Step-selection model

#### 232 3.1.1 Input embeddings

233 We build two sets of embeddings  $\phi_n^{\text{in}}, \phi_n^{\text{cand},(i)} \in \mathbb{R}^{d_{\text{emb}}}$ ,  $n \in \{1, \dots, N-1\}$ ,  
 234  $i \in \{1, \dots, K\}$  such that:

- 235 •  $\phi_n^{\text{in}}$ , input for the trajectory encoder, depends on  $x_{n-1}, x_n, t_n, t_{n+1}, z_n$ ;
- 236 •  $\phi_n^{\text{cand},(i)}$ , input for the candidate selection module, depends on  $x_n, x_{n+1}^{(i)}, z_{n+1}^{(i)}$ .

237 The inputs are represented as collections of carefully engineered continuous and  
 238 discrete features that we will describe later (see Section 3.2). Missing (NaN)  
 239 values are replaced with a special embedding vector learned as an additional  
 240 parameter. In each case, we project each feature vector to a common embedding  
 241 space  $\mathbb{R}^{d_{\text{emb}}}$ , then linearly combine them (with different learnable coefficients in  
 242 each of the two cases).

243 More precisely, for  $\phi^{\text{in}}$ :

$$\phi_n^{\text{in}} = \sum_{j=1}^F w^{\text{in},(j)} \left( W^{(j)} f_n^{(j)} + b^{(j)} \right), \quad (2)$$

244 where  $f_n^{(j)}$  is the  $j$ -th out of all  $F$  feature vectors at step  $n$ , and the learnable  
 245 parameters are coefficients  $w^{\text{in},(j)} \in \mathbb{R}$  (we set  $w^{\text{in},(j)} = 0$  for features we do  
 246 not wish to consider), biases  $b^{(j)} \in \mathbb{R}^{d_{\text{emb}}}$  and weight matrices  $W^{(j)} \in \mathbb{R}^{d_{\text{emb}} \times d_j}$ .  
 247 The formula for  $\phi_n^{\text{cand},(i)}$  is analogous. As can be seen from Eq. (2), the chosen  
 248 method for constructing input embeddings allows features to have different di-  
 249 mensions and automatically projects them to the desired embedding dimension  
 250 (via  $W^{(j)}$  and  $b^{(j)}$ ) before applying scaling through  $w^{\text{in},(j)}$ .

### 251 3.1.2 Trajectory encoder

252 The trajectory encoder is a Transformer encoder with causally masked attention.  
 253 It receives the embedding sequence  $\phi_{1\dots N-1}^{\text{in}}$  and outputs a sequence of vectors  
 254  $h_{1\dots N-1}$  where  $h_n$  is a representation of  $\xi_{1\dots n}$ . The encoder does not use any  
 255 positional encoding in the conventional sense (encoding the indices  $1, \dots, N-1$ ,  
 256 as is commonly done in Transformers), but position information is conveyed by  
 257 the feature representations of the timestamps  $t_{1\dots N-1}$ .

### 258 3.1.3 Candidate selection

259 The candidate selection module is used to select the next location out of a list of  
 260 candidates. We build upon the common approach that models the probability of  
 261 an individual being present at a given candidate location via conditional logistic  
 262 regression [3]; expressed in our notation:

$$\frac{\exp(\beta^\top \phi_n^{\text{cand},(i)})}{\sum_{i'=1}^K \exp(\beta^\top \phi_n^{\text{cand},(i')} )}, \quad (3)$$

263 where  $\beta$  is a parameter vector.

264 In this work, in order to incorporate the context representation computed  
 265 by the trajectory encoder, we replace the global parameter vector  $\beta$  with a  
 266 context-dependent *query vector*  $q_n \in \mathbb{R}^{d_{\text{sel}}}$ , which is a linear projection of the  
 267 trajectory encoder output  $h_n$ . We also do not use the raw candidate features  
 268  $\phi_n^{\text{cand},(i)}$  but replace them with a *key vector*  $k_n^{(i)} \in \mathbb{R}^{d_{\text{sel}}}$ , which is computed  
 269 by concatenating the feature vector with the corresponding encoder output  $h_n$   
 270 and passing the result through a *candidate encoder* (a fully-connected network):  
 271  $k_n^{(i)} = E_{\text{cand}}([\phi_n^{\text{cand},(i)}, h_n])$ . Thus, we arrive at a *dot-product attention mecha-*  
 272 *nism*; scaling the dot products by  $1/\sqrt{d_{\text{sel}}}$  as in Transformer attention [173], we  
 273 have:

$$P(y_{n+1} = i \mid \xi_{1\dots n}, t_{n+1}, x_{n+1}^{(1\dots K)}, z_{n+1}^{(1\dots K)}) = \frac{\exp(q_n^\top k_n^{(i)}/\sqrt{d_{\text{sel}}})}{\sum_{i'=1}^K \exp(q_n^\top k_n^{(i')}/\sqrt{d_{\text{sel}}})}. \quad (4)$$

During training, the first candidate location  $x_{n+1}^{(1)}$  is taken as the true next location  $x_{n+1}$ ; the rest of the candidates are randomly sampled around the current location  $x_n$  (we detail this process below). This allows us to define a cross entropy loss, which we minimize through stochastic gradient descent using the Adam optimizer:

$$\mathcal{L} = -\frac{1}{N-1} \sum_{n=1}^{N-1} \log_K P(y_{n+1} = 1 \mid \xi_{1\dots n}, t_{n+1}, x_{n+1}^{(1\dots K)}, z_{n+1}^{(1\dots K)}) \quad (5)$$

### 3.1.4 Variable receptive field training

As mentioned above, we aim to evaluate our model on arbitrary trajectory segments up to some maximum length  $c_{\max}$  (this procedure is detailed below in Section 4.1). As can be seen from Eq. (5), our model is effectively being simultaneously trained on all prefixes of the trajectory  $\xi_{1\dots N}$ . Hence, the model is able to accept segments of variable length as desired, but being only trained on trajectory prefixes may bias it, leading to incorrect predictions on segments that are not prefixes. To alleviate this, we propose a training scheme that intervenes on the attention weights to randomly vary the past context available for each prediction.

In each training batch, we sample a random integer  $B$  uniformly from  $\{1, \dots, N_{\max}\}$  and apply a block-diagonal attention mask to the attention matrix (on top of the causal mask) with blocks of size  $B$  (with the last block truncated if  $B \nmid N$ ). As a result, the ranges of positions  $\{1, \dots, B\}$ ,  $\{B+1, \dots, 2B\}$ , etc. are prevented from attending to each other, and the corresponding segments are therefore effectively considered as separate trajectories.

## 3.2 Data representation

Let us now describe the feature mappings used for location and time, as well as associated features.

### 3.2.1 Location

In the raw data, each location  $x_n$  is represented as a GPS coordinate pair (latitude, longitude). We represent it as a geodetic normal vector (*n-vector*)  $\nu(x_n) \in \mathbb{R}^3$ .

Additionally, we encode the position relative to the previous location  $x_{n-1}$  as a *movement vector*  $\mu(x_{n-1}, x_n) \in \mathbb{R}^2$ , obtained by computing the bearing and distance from  $x_{n-1}$  to  $x_n$  and converting them to cartesian coordinates. We apply scaling to make the overall root-mean-square (RMS) of the norms of movement vectors computed on the training dataset equal to 1.

Analogously, we encode each candidate location  $x_{n+1}^{(i)}$  as an n-vector  $\nu(x_{n+1}^{(i)})$  and as a movement vector  $\mu(x_n, x_{n+1}^{(i)})$ .

309    **3.2.2 Time**

310    We encode a timestamp  $t_n$  as:

- 311    • a 10-dimensional vector of sines and cosines with a period of 1 second, 1  
312    minute, 1 hour, 1 day, and 1 tropical (solar) year, respectively, such that  
313    their phase synchronizes on January 1st, 2000, at 00:00:00 UTC;  
314    •  $\sin(\text{LMT}/24 \cdot 2\pi)$  and  $\cos(\text{LMT}/24 \cdot 2\pi)$ , where LMT is the local mean  
315    time (i.e. UTC adjusted by longitude) in (fractional) hours;  
316    • 3 integer values (one-hot-encoded) representing the calendar month (0–  
317    11), the day of the month (0–30), and the day of the week (0–6) in UTC.

318    We also encode the time difference w.r.t. the next timestamp  $t_{n+1}$  as a 12-  
319    dimensional vector of sines and cosines with the same periods as above, plus a  
320    period of 25 years.

321    While this multi-scale encoding may not be necessary in our case (where the  
322    time differences are between 9 and 15 h), we propose it as a generic representa-  
323    tion suitable for any time scale from seconds to years (and hence for virtually  
324    all existing animal movement data).

325    **3.2.3 Associated variables**

326    For each input and candidate location, we retrieve and pre-process geospatial  
327    variables as described in Section 2.3. We also include the taxon vectors (as de-  
328    scribed in Section 2.2) as an additional encoder feature vector for every element  
329    of the input sequence.

330    **3.3 Candidate sampling**

331    We sample each candidate location  $x_{n+1}^{(i)}$  as follows:

- 332    • we estimate the current bearing  $\beta$  of the animal from the positions  $x_n$  and  
333     $x_{n-1}$ ;  
334    • we independently sample a *turning angle*  $\theta \sim \hat{P}(\theta)$  and a *log-distance*  
335     $\log d \sim \hat{P}(\log d)$ ;  $\beta' \leftarrow \beta + \theta$ ;  
336    • we compute  $x_{n+1}^{(i)}$  by moving  $x_n$  according to  $\beta'$  and  $d$ .

337     $\hat{P}(\theta)$  and  $\hat{P}(\log d)$  are estimated on the training set as follows:

- 338    • We collect all turning angles from the training set and compute the quan-  
339    tiles (estimated using linear interpolation) at 101 equally spaced points  
340     $0 = q_0, q_1, \dots, q_{100} = 1$ . We use them to construct the quantile function  
341    of  $\hat{P}(\theta)$  as a piecewise linear function with knots at  $q_0, q_1, \dots, q_{100}$ .  
342    • We collect the natural logarithms of all non-zero distances between consec-  
343    utive points in the dataset; we construct the quantile function of  $\hat{P}(\log d)$   
344    analogously.

345 We sample from each distribution by drawing a sample from  $\mathcal{U}[0, 1]$  and  
346 passing it through the estimated quantile function; this is sometimes called the  
347 *increasing rearrangement* [174].

348 In our experiments, we condition the distributions on the taxon, i.e. we  
349 estimate a separate pair of distributions on the section of the training dataset  
350 corresponding to each taxon.

### 351 3.4 Implementation details and hyperparameters

352 Our implementation of MoveFormer, available as open source software,<sup>8</sup> is writ-  
353 ten in Python using the PyTorch framework<sup>8</sup> and the `x-transformers`<sup>9</sup> pack-  
354 age. The code for efficient geospatial variable loading relies on the `rasterio`<sup>10</sup>  
355 library and is released as a separate package, `gps2var`.<sup>11</sup> See Section 7 for  
356 information about code availability.

357 The trajectory encoder is a 6-layer Transformer with 8 attention heads per  
358 layer and a feature dimension of 128. The candidate encoder is a fully-connected  
359 neural network with one hidden layer of size 256 and a GELU activation [175].  
360 The candidate selection module has  $d_{\text{sel}} = 128$ . The total number of param-  
361 eters of the model is around 2.6 million – several orders of magnitude smaller  
362 than current state-of-the-art Transformer language models, for instance, but  
363 appropriate for the limited-size dataset that we are working with.

364 The Adam optimizer uses a learning rate of  $5 \times 10^{-5}$  with linear warm-up  
365 and exponential decay. We train for 180 epochs with a batch size of 24, taking  
366 7.5 h on a Tesla V100 GPU (note that GPU utilization was only about 20 % and  
367 the performance bottleneck appeared to be the geospatial variable loading). We  
368 validate on the validation set twice per epoch and use the checkpoint with the  
369 lowest validation loss.

370 The complete hyperparameter settings are included with the source code.

## 371 4 Analysis methods

### 372 4.1 Context length analysis

373 Riotte-Lambert et al. [11] propose to use *conditional entropy* as a measure of  
374 uncertainty in predicting the next location given the  $c$  previous locations. Specif-  
375 ically, given a distribution  $P$  over sequences of locations, conditional entropy of  
376 order  $c$  can be written as

$$377 H_c = -\mathbb{E}_{P(s_1, \dots, s_c)}[\log P(s_{c+1} | s_1, \dots, s_c)], \quad (6)$$

377 where  $P(s_1, \dots, s_c)$  is understood as the probability of  $c$  consecutive locations in  
378 a sequence being equal to  $s_1, \dots, s_c$ , and  $P(s_{c+1} | s_1, \dots, s_c)$  as the conditional  
379 probability of  $s_{c+1}$  immediately following the sequence  $s_1, \dots, s_c$ . Considering  
380 this uncertainty measure as a function of the context length  $c$ , it may be used  
381 to study routine movement behavior.

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382 8

383 8<sup>8</sup><https://pytorch.org/>

384 9<sup>9</sup><https://github.com/lucidrains/x-transformers>

385 10<sup>10</sup><https://github.com/rasterio/rasterio>

386 11

382 Riotte-Lambert et al. [11] work with a finite set of discrete locations, allowing  
383 them to evaluate the expression (6) empirically on a given trajectory. However,  
384 the probability estimates quickly become unreliable with increasing  $c$  due to  
385 data sparsity. Moreover, the method is inapplicable when locations are unique,  
386 as in our case.

387 We propose an alternative way, which is to approximate  $\log P$  using a suitable  
388 machine learning model (e.g. our proposed step selection model), so that  
389 Eq. (6) becomes *cross entropy* computed on trajectory *segments* of appropriate  
390 length. In our case:<sup>11</sup>

$$\begin{aligned} H_c &\approx -\frac{1}{N-1} \sum_{n=1}^{N-1} \log_K P(y_{n+1} = 1 \mid \xi_{n-c+1 \dots n}, t_{n+1}, x_{n+1}^{(1 \dots K)}, z_{n+1}^{(1 \dots K)}) \\ &= -\frac{1}{N-1} \sum_{n=1}^{N-1} \log_K P(y_{n+1} = 1 \mid \psi_{n,c}), \end{aligned} \quad (7)$$

391 where we collapse all the conditioning variables to  $\psi_{n,c}$  for brevity. For more fine-  
392 grained analysis, we may be interested not in the sequence-level cross entropy,  
393 but rather in the “pointwise” values, i.e.  $-\log_K P(y_{n+1} = 1 \mid \psi_{n,c})$ .

394 More generally, we may alternatively choose to examine any metric that can  
395 be computed from the probabilities. We adopt the *relative entropy* (also known  
396 as the *Kullback-Leibler divergence*) of the prediction with the maximum context  
397 length  $c_{\max}$  with respect to the one at context length  $c$  (as proposed by Cifka  
398 and Liutkus [176] in the context of causal language models for text):

$$D_{\text{KL}}[P(y_{n+1} \mid \psi_{n,c_{\max}}) \parallel P(y_{n+1} \mid \psi_{n,c})] = \sum_{i=1}^K P(y_{n+1} = i \mid \psi_{n,c_{\max}}) \log \frac{P(y_{n+1} = i \mid \psi_{n,c_{\max}})}{P(y_{n+1} = i \mid \psi_{n,c})}. \quad (8)$$

399 Note that this metric does not depend on the ground truth location, but measures  
400 the amount of information gained by considering the maximal context  
401 instead of the limited one.

#### 402 4.1.1 Relevant context length

403 We may expect that there would be a critical context length  $C$  after which  
404 the above metrics stop improving, as further extending the context does not  
405 result in significant information gain. Similarly to Riotte-Lambert et al. [11],  
406 we define the *relevant context length*  $C_m$  – for a given metric  $m$  – as the smallest  
407 context length for which the metric reaches its optimum, with a 5% tolerance  
408 for robustness to noise:

$$C_m = \min \left\{ c: \frac{m(c) - \min_{c'} m(c')}{\max_{c'} m(c') - \min_{c'} m(c')} \leq 0.05 \right\}. \quad (9)$$

#### 409 4.1.2 Efficient evaluation

410 We now discuss how to efficiently compute the probabilities needed to calculate  
411 the above metrics, following the procedure proposed for causal language models

---

<sup>11</sup>We use the number of candidates  $K$  as the base of the logarithm for consistency with Eq. (5) and noting that this amounts to a multiplicative constant ( $1/\log K$ ).

<sup>412</sup> by Cífká and Liutkus [176]. We may collect all the probabilities in a tensor  
<sup>413</sup>  $\mathbf{P} \in \mathbb{R}^{N \times c_{\max} \times K}$  such that

$$\mathbf{P}_{n,c,i} = P(y_{n+1} = i \mid \psi_{n,c}). \quad (10)$$

<sup>414</sup> Observe that by running the model on a segment of the trajectory corresponding  
<sup>415</sup> to indices  $n, \dots, n + c_{\max} - 1$  for a given  $n$ , we obtain all the values  $\mathbf{P}_{n+c-1,c,*}$  for  
<sup>416</sup>  $c \in \{1, \dots, c_{\max}\}$ . We may also notice that  $\mathbf{P}_{n,n,*} = \mathbf{P}_{n,n+1,*} = \dots = \mathbf{P}_{n,c_{\max},*}$   
<sup>417</sup> for any  $n < c_{\max}$ . Hence, we can efficiently fill in the tensor  $\mathbf{P}$  using  $N$  runs of  
<sup>418</sup> the model on segments of length at most  $c_{\max}$ .

## <sup>419</sup> 4.2 Candidate feature importance

<sup>420</sup> While the parameters of step-selection models fitted by conditional logistic re-  
<sup>421</sup> gressions or point-process models are directly interpretable [4], deep learning  
<sup>422</sup> models are known as “black boxes” that require special techniques to be inter-  
<sup>423</sup> preted post-hoc. A simple but popular technique [177, 178] is based on testing  
<sup>424</sup> the model on a dataset with the values of a given feature randomly permuted.  
<sup>425</sup> While aware of the caveats related to using this technique with correlated fea-  
<sup>426</sup> tures [179], we employ it here to demonstrate the possibility of interpretation,  
<sup>427</sup> and leave more advanced techniques for future work.

<sup>428</sup> Specifically, we study how individual *candidate features* (components of  
<sup>429</sup>  $\phi_n^{\text{cand},(i)}$ ) influence the selection of candidates. We pick a feature (or a group  
<sup>430</sup> of features), and for every observation in the dataset, we randomly shuffle the  
<sup>431</sup> feature’s values among the  $K$  candidates (in contrast to Fisher et al. [178], who  
<sup>432</sup> shuffle values across the entire dataset). The aim is to make the feature com-  
<sup>433</sup> pletely uninformative while maintaining its values plausible in the given context.  
<sup>434</sup> We evaluate the model on both the permuted and the original dataset, and use  
<sup>435</sup> the difference in performance as a measure of the importance of the selected  
<sup>436</sup> feature.

# <sup>437</sup> Results and discussion

## <sup>438</sup> 5 Results

### <sup>439</sup> 5.1 Validation

<sup>440</sup> We evaluate the proposed model (here dubbed VARCTX) against variants to  
<sup>441</sup> serve as baselines:

- <sup>442</sup> • FULLCTX is a variant without the variable receptive field training (see  
<sup>443</sup> Section 3.1.4);
- <sup>444</sup> • NOATT is a model where all the attention layers are removed from the  
<sup>445</sup> Transformer encoder, so that information is not allowed to flow between  
<sup>446</sup> different positions in the sequence;
- <sup>447</sup> • NOENC is a model where the Tranformer encoder is removed, i.e. we have  
<sup>448</sup>  $h_n = \phi_n^{\text{in}}$ .

<sup>449</sup> Note that the last two variants have a receptive field of 1 (i.e. only the features  
<sup>450</sup> at position  $n$  are available for predicting the location at  $n + 1$ ). To simulate  
<sup>451</sup> this for VARCTX and FULLCTX in a comparable way, we test these in a regime  
<sup>452</sup> (denoted by +DIAG) where the attention matrices are restricted to an identity  
<sup>453</sup> matrix, i.e. each position can only attend to itself.

<sup>454</sup> After running each of the above models on the test set, we compute the  
<sup>455</sup> following metrics:

- <sup>456</sup> • xent@16: cross entropy (Eq. (5)) computed with 16 candidates;
- <sup>457</sup> • xent@100: cross entropy computed with 100 candidates;
- <sup>458</sup> • acc 1/16: accuracy (i.e. how often the top scoring candidate is the ground  
<sup>459</sup> truth) with 16 candidates,
- <sup>460</sup> • acc 10/100: top-10 accuracy (i.e. how often the ground truth is among the  
<sup>461</sup> 10 top scoring candidates) with 100 candidates.

<sup>462</sup> The results, averaged over all trajectories, are presented in Table 2. We note  
<sup>463</sup> that the results are very consistent across all metrics, and we found all pairs of  
<sup>464</sup> metrics to be strongly correlated (Pearson  $\rho > 0.87$ , computed over all models  
<sup>465</sup> and trajectories).

<sup>466</sup> Both FULLCTX and VARCTX outperform the rest of the models, which  
<sup>467</sup> have a receptive field length of 1. This is evidence that providing past move-  
<sup>468</sup> ment as context is beneficial. Interestingly, VARCTX yields better results than  
<sup>469</sup> FULLCTX, possibly because the variable receptive field training scheme effec-  
<sup>470</sup> tively makes the training data more diverse, alleviating overfitting.

<sup>471</sup> We can also observe that the results of VARCTX+DIAG are closest to those  
<sup>472</sup> of the models trained with minimum context (NOATT, NOENC). This sug-  
<sup>473</sup> gests that the performance of VARCTX is not strongly degraded by limiting its  
<sup>474</sup> receptive field at test time (unlike that of FULLCTX), validating our variable  
<sup>475</sup> receptive field training approach.

<sup>476</sup> Finally, we noticed large performance differences between species. For the  
<sup>477</sup> VARCTX model, we calculated the average cross entropy for each taxonomic

model	xent@16 ↓	xent@100 ↓	acc 1/16 ↑	acc 10/100 ↑
FULLCTX	0.869	0.909	0.198	0.293
FULLCTX+DIAG	0.990	0.998	0.102	0.157
VARCTX	<b>0.847</b>	<b>0.894</b>	<b>0.221</b>	<b>0.323</b>
VARCTX+DIAG	0.932	0.954	0.136	0.204
NOATT	0.919	0.945	0.157	0.231
NOENC	0.928	0.950	0.148	0.217

Table 2: Results for different variants of the model. FULLCTX: trained on full trajectories (max. length 500); VARCTX: trained with variable receptive field; NOATT: no attention layers; NOENC: no encoder; DIAG: attention restricted to diagonal matrix during inference. Xent: cross entropy (lower is better), acc: accuracy (higher is better).

class	order	xent@16	#train
Aves	Accipitriformes	0.815	58 994
	Anseriformes	0.827	64 008
	Cathartiformes	1.057	8653
	Charadriiformes	0.815	205 602
	Ciconiiformes	0.697	237 304
Mammalia	Artiodactyla	0.928	201 464
	Carnivora	0.986	12 282
	Proboscidea	0.980	24 870
Reptilia	Testudines	0.998	34 577

Table 3: VARCTX validation cross entropies by taxonomic order, along with numbers of observations in the training data.

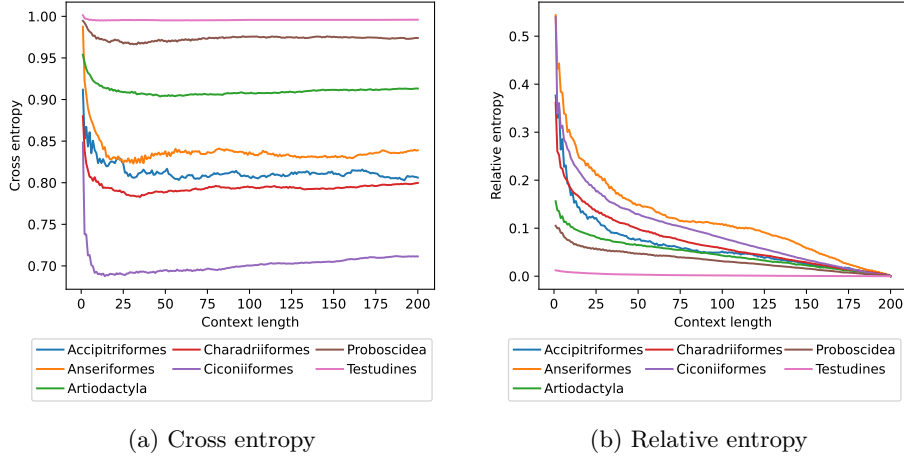
478 order (see Table 3) and found that it tends to be lower (i.e. better) for orders  
 479 with a higher number of observations in the training set (Pearson  $\rho = -0.71$ ).

## 480 5.2 Context length analysis

481 In this section, we demonstrate the ability to use the VARCTX to study the  
 482 dependence of the predictions on the length  $c$  of the available past context, as  
 483 described in Section 4.1. We set  $c_{\max} = 200$  and  $K = 16$ .

484 First, we display in Fig. 5 the average cross entropy and relative entropy as  
 485 a function of context length and by taxonomic order, and in Fig. 5 examples  
 486 for concrete observations, with the relevant context length  $C$  highlighted. We  
 487 observe that the best predictions tend to be achieved around context lengths  
 488 10–50, which corresponds to 5–25 days.

489 Apart from the clear inter-species differences in cross entropy already noted  
 490 in the previous section (Table 3), we also observe some differences in *relative*  
 491 entropy, though less marked. For example, while *Ciconiiformes*' movements are  
 492 substantially easier (in terms of cross entropy) for our model to predict than  
 493 those of *Anseriformes*, both have a similar relative entropy profile, indicating  
 494 that the amount of information contributed by each time scale is similar for both  
 495 taxa. On the other hand, note that the flat relative entropy profile of *Testudines*



(a) Cross entropy

(b) Relative entropy

Figure 4: Metric value averages by context length and taxonomic order, computed on the test set (only positions  $n > c_{\max} = 200$ ).

simply reflects a failure of our model to accurately predict their movements at any time scale – as evidenced by the cross entropy values being close to 1 –, which is possibly due to an insufficient amount of reptile training data.

Fig. 6 shows the distribution of the relevant context length  $C$  for each taxon in the test set. There are apparent differences between taxa, but we also note the large variability *within* each taxon that could be of interest in itself.

### 5.3 Candidate feature importance

We present in Fig. 7 the results of the feature importance experiment. Vector features (location) are treated as groups; bioclimatic variables are tested both individually and as a group.

The most important features found by this method are movement vector and land cover, followed by human footprint. The bioclimatic variables appear to have relatively low impact, with the most important ones being BIO2 (mean diurnal range), BIO14 and BIO17 (both related to precipitation). Interestingly, global location (represented as n-vectors) seems to be the least important feature, possibly because it is difficult to exploit for candidate selection compared to the relative location information provided by the movement vectors.

Note that only *candidate* features  $\phi_n^{\text{cand},(i)}$  are tested here, and the results do not say anything about the *input* (past observation) features  $\phi_{1\dots n}^{\text{in}}$ . For example, global location, which we found to be unimportant as a candidate feature, may well turn out to be an important past context feature.

## 6 Discussion

In this work, we propose a new model to learn from animal trajectories. Inspired by the classical step-selection framework [2] and previous work on the quantification of uncertainty in movement predictions [11], we designed MoveFormer, a step-based model of movement that builds upon recent developments in deep

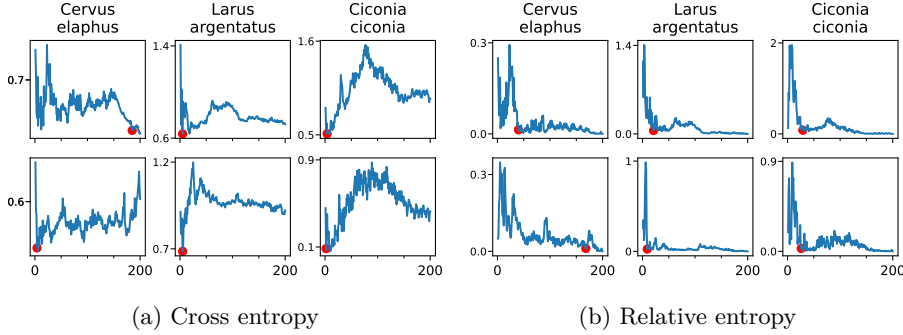


Figure 5: Examples of metric values (pointwise, i.e. for a single observation within a given trajectory) plotted as a function of context length. Top and bottom correspond to different (random) positions within the same trajectory. The red dot marks the relevant context length (where the metric reaches 5 % of its min-max range).

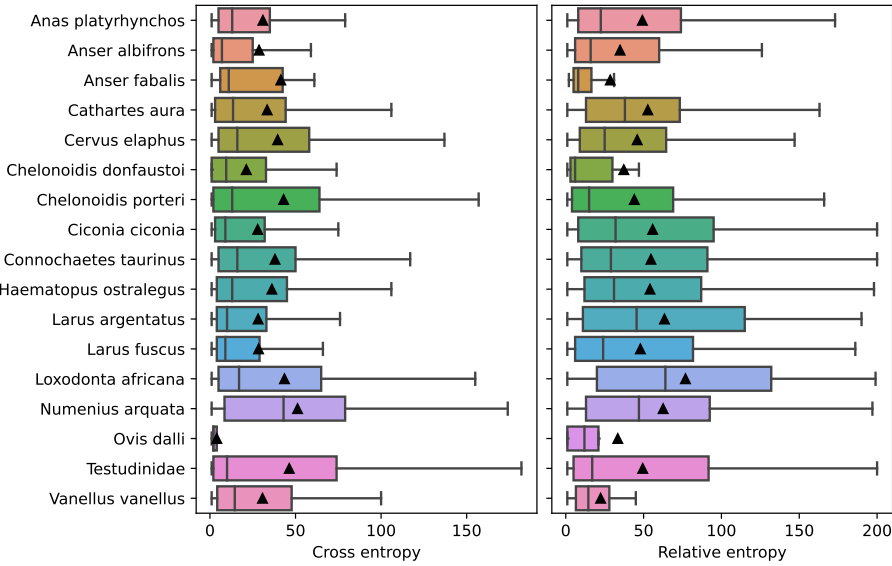


Figure 6: Relevant context length  $C$  by taxon, computed using cross entropy and relative entropy (pointwise values, as shown in Fig. 5), respectively. The black triangles indicate means.

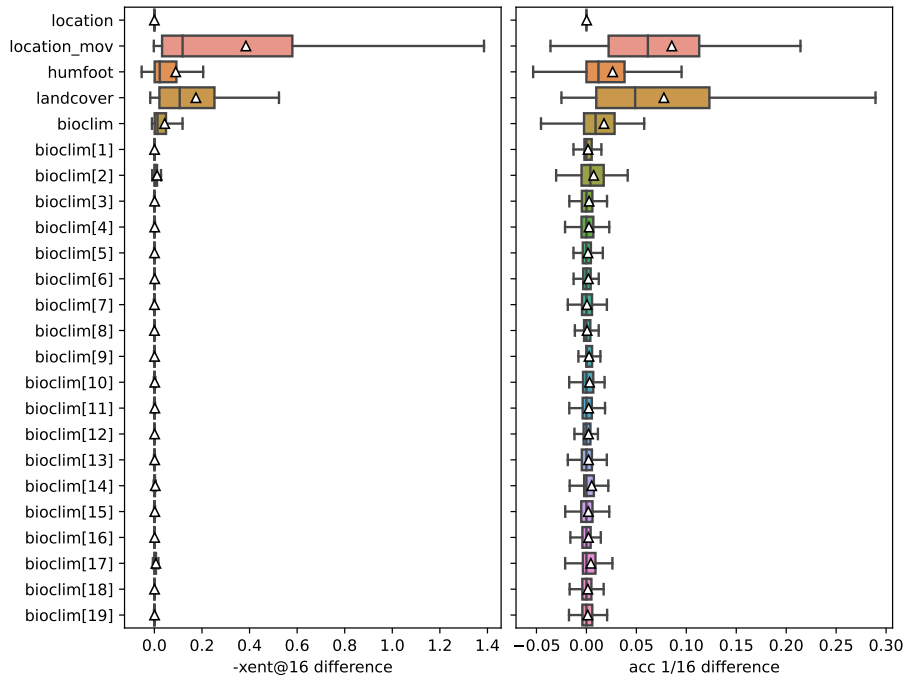


Figure 7: Candidate feature importances computed as differences between performance (measured using negative cross entropy and accuracy with 16 candidates) with and without permuted feature values. The plot shows the distribution over test trajectories. The features are, from top to bottom: location n-vector, movement vector, human footprint, land cover, and finally the 19 bioclimatic variables (BIO1 to BIO19; [see Table 6 in Supplementary information](#)), first as a group and then each individually.

learning, such as the Transformer architecture. This allowed us to meet our initial goal to endow the model with a unique ability to learn how past movements influence current and future ones. Although this is an important question in movement ecology, it has remained poorly addressed so far because classical step-selection functions or other movements models are unable to account for past information except in a very simplified way (e.g. by including a feature indicating whether or not the animal has previously visited a given site).

An important contribution of this work is also to generalize the suggestion of Riotte-Lambert et al. [11] to use conditional entropy calculated over visits to discrete sites as a way to measure movement uncertainty. Although attractive, the difficulty of discretizing trajectories to meaningful ‘sites’ has slowed down the application of this idea. Here, we extend it to locations acquired in continuous space and propose cross-entropy and relative entropy, estimated through the movement model, as a more general approach. This allows to estimate the *relevant context length* (‘relevant order of dependency’ in Riotte-Lambert et al. [11]), i.e. the amount of the past that significantly improves the predictions about further movements. We did so in this study, and to the best of our knowledge, our study therefore provides the first estimation of how much of the past one needs to know to improve predictions of animal movements.

Our results suggest that for most datasets, predictions are improved when integrating the information from about a few days to two or three weeks before the movement to be predicted. Why this is the case, and why these results are broadly consistent between species, with possibly significant within-species variability, remains to be investigated further as it was beyond the goal of this methodological work. We note that, possibly, these results are affected by our choice to alternate sampling at midnight and at noon and to limit the length of trajectories to 500 locations, restricting the receptive field of our model to about 250 days. This may have weakened or excluded the influence of migration, which commonly leads to seasonal back-and-forth movement patterns~~and that, when accounted for, could help improve predictions about future movements.~~. Therefore, in future work, it would be interesting to study how the temporal scale and resolution of trajectories affects the results. It is possible that using data with a higher temporal resolution and/or longer temporal context would reveal a second peak in context length, indicative of nested scale-patterns in the trajectories.

The effect of *spatial* scale and resolution of the geospatial variables may also be worth investigating. Instead of only considering the features at a given location, future work should explore including information from a local neighborhood. Apart from improving predictions in general, this additional context information may enable dealing with lower-accuracy (e.g. Argos) tracking data, where the variables associated with exact location may become too noisy.

One obvious limitation of our approach is the data requirement. As with all deep learning approaches, learning is limited by the data available in the training set, and enough data should also be available for validation and test sets. The whole dataset we gathered here, despite being rather large ( $> 1500$  trajectories) compared to movement datasets currently analyzed in ecology, is likely close to the minimal size required to obtain a robust model and avoid severe overfitting issues. Currently, there are probably very few, if any, single-species datasets large enough to fit this model. For this reason, we aggregated data from

572 numerous species; as a benefit, this allowed us to demonstrate that comparative  
573 analyses could be conducted with the model, for instance by comparing the  
574 distribution of relevant context lengths between species or higher-order taxa.

575 An important characteristic of the proposed approach is that the model not  
576 only accounts for past movements to predict new ones, but can also account for  
577 environmental predictors. First, this is crucial for realistic predictions, as the  
578 step-selection literature has well demonstrated that step selection by animals is  
579 critically linked to habitats to be traversed or reached. Second, this allows to  
580 evaluate the relative importance of predictors in improving predictions. Interest-  
581 ingly, we found that purely relative positional information (movement vec-  
582 tors) could be more important than environmental variables for future location  
583 prediction. We tentatively suggest that this result might be linked to the fact  
584 that most animals favor familiar places and by doing so restrict themselves to  
585 well-established home-ranges [180]. We however also found, without surprise,  
586 that among the environmental variables tested, land cover and human footprint  
587 significantly affected animal movements [181].

588 To summarize, in the present work, we provide a new, state-of-the art model  
589 to analyze and predict animal movement data. The novelty of the model lies in  
590 the fact that it leverages the power of deep learning approaches and can account  
591 for past movements in the predictions. However, we emphasize, and have shown  
592 above, that the model is not only a tool for prediction, but can also be used to  
593 test hypotheses about the intrinsic and extrinsic drivers of animal movements.

## 594 **7 AcknowledgmentsData, script and code availability**

595 Our code used for data retrieval, data pre-processing, model training, and  
596 analyses is available online as open source software<sup>12</sup> [182]; the code for efficient  
597 geospatial variable loading is released as a separate open source package, `gps2var`<sup>13</sup>  
598 [183]. We also release the weights of the trained models [184].

599 The data used in this work was compiled from 102 tracking studies, out of  
600 which:

- 601 • 98 are publicly available and were retrieved on 15 February 2022 via the  
602 Movebank API;
- 603 • 2 more are additionally made publicly available in Movebank [166, 167]  
604 at the time of writing;
- 605 • the remaining 2 (plains zebras and blue wildebeest in Hluhluwe-iMfolozi  
606 Park) are under restricted use imposed by the institution managing the  
607 study area.

608 For a list of the Movebank studies, see Table 4.

## 609 **8 Funding and acknowledgments**

610 This work was supported by the LabEx NUMEV (ANR-10-LABX-0020) and  
611 the REPOS project, both funded by the I-Site MUSE (ANR-16-IDEX-0006).

<sup>12</sup><https://github.com/cifikao/moveformer>

<sup>13</sup><https://github.com/cifikao/gps2var>

612 Computations were performed using HPC/AI resources from GENCI-IDRIS  
613 (Grant AD011012019R1).

614 We would like to thank all authors who made their data available through  
615 Movebank under Creative Commons licenses(see Table 4 in the appendix for  
616 the list of datasets used in this work).

## 617 9 Conflict of interest disclosure

618 The authors declare they have no conflict of interest relating to the content of  
619 this article.

620 **A Appendix**

621 **Supplementary information**

Table 4: The list of all Movebank datasets used in this work.

ID	Name and references	License
446579	MPIAB Lake Constance Mallards GPS [13]	CC BY
481458	Vultures Acopian Center USA GPS [14–17]	CC BY
1764627	Kruger African Buffalo, GPS tracking, South Africa [18–21]	CC0
2928116	Galapagos Tortoise Movement Ecology Programme [22–28]	CC BY
2988333	Navigation experiments in lesser black-backed gulls (data from Wikelski et al. 2015) [29, 30]	CC0
6770990	MPIAB PNIC hurricane frigate tracking [127]	CC BY
7002955	HUJ MPIAB White Stork GSM E-Obs [128]	CC BY
7431347	MPIAB Argos white stork tracking (1991–2018) [31]	CC BY
8019591	Dunn Ranch Bison Tracking Project [129]	CC BY
8849813	LifeTrack - Great Egrets [130]	CC0
8863543	HUJ MPIAB White Stork E-Obs [131]	CC BY
9493881	LifeTrack White Stork Uzbekistan [132]	CC BY
9651291	Egyptian vultures in the Middle East and East Africa [32–35]	CC BY
10157679	LifeTrack White Stork Tunisia [133]	CC BY
10204361	Pandion haliaetus Osprey - SouthEast Michigan [134]	CC0
10236270	LifeTrack White Stork Armenia [36–38]	CC BY
10449318	LifeTrack White Stork Loburg [135]	CC BY
10449535	LifeTrack White Stork Greece Evros Delta [36–38]	CC BY
10449698	HUJ MPIAB White Stork GSM 2013 [136]	CC BY
10596067	LifeTrack White Stork Moscow [36–38]	CC BY
10763606	LifeTrack White Stork Poland [37]	CC BY
14671003	Hooded Vulture Africa [137]	CC BY
16880941	Turkey vultures in North and South America (data from Dodge et al. 2014) [17, 39]	CC0
19411459	Movement ecology of the jaguar in the largest floodplain of the world, the Brazilian Pantanal [40–45]	CC0
20202974	e-Obs GPRS Himalayan Griffon - Bhutan-MPIAB [46–49]	CC BY
21231406	LifeTrack White Stork SW Germany [37, 38, 50, 51]	CC BY
24442409	LifeTrack White Stork Bavaria [38, 50, 52]	CC BY
69724677	FTZ Geese Wadden Sea [138]	CC0
74496970	MPIAB white stork lifetime tracking data (2013–2014) [36, 37, 53]	CC BY
92261778	LifeTrack Whooper Swan Latvia [139]	CC BY
133992043	Migration timing in white-fronted geese (data from Kölzsch et al. 2016) [54, 55]	CC BY
173641633	LifeTrack White Stork Vorarlberg [50, 56]	CC BY
178979729	Latham Alberta Wolves [57, 58]	CC BY-NC
178994931	Peters Hebblewhite Alberta-BC Moose [59]	CC BY
182746263	High-altitude flights of Himalayan vultures (data from Sherub et al. 2016) [47, 49]	CC0
190490326	Movement strategies of Galapagos tortoises (data from Bastille-Rousseau et al. 2016) [24, 26–28]	CC BY-NC
208413731	White-bearded wildebeest in Kenya [60]	CC BY
209824313	Hebblewhite Alberta-BC Wolves [61, 62]	CC BY
212096177	LifeTrack White Stork Oberschwaben [38, 50, 63]	CC BY

622 Continued on next page

Table 4: The list of all Movebank datasets used in this work. (Continued)

ID	Name and references	License
217784323	Vultures Acopian Center USA 2003-2016 [16, 17, 64]	CC BY
236953686	LifeTrack Ducks Lake Constance [140]	CC0
329155299	Canada geese ( <i>Branta canadensis</i> ) [141]	CC0
384868221	White-tailed Eagle Poland. [142]	CC BY-NC
475878514	Coyote Valley Bobcat Habitat Connectivity Study [143]	CC BY-NC
501787846	Aromas Hills Bobcat Habitat Connectivity Study [144]	CC BY-NC
505156776	Graugans Zugverhalten Neusiedler See [145]	CC BY-NC
560041066	Eastern flyway spring migration of adult white storks (data from Rotics et al. 2018) [65, 66]	CC BY
604806671	MH_WATERLAND - Western marsh harriers ( <i>Circus aeruginosus</i> , <i>Accipitridae</i> ) breeding near the Belgium-Netherlands border [67]	CC0
657674643	North Sea population tracks of greater white-fronted geese 2014-2017 (data from Kölzsch et al. 2019) [68, 69]	CC BY
657965212	Pannonic population tracks of greater white-fronted geese 2013-2017 (data from Kölzsch et al. 2019) [69, 70]	CC BY
672882373	<i>Milvus_milvus_atlantismarcuard</i> [146]	CC BY-NC
673728219	NPS Dall Sheep in Yukon-Charley Rivers National Preserve [147]	CC BY
736029750	ThermochronTracking Elephants Kruger 2007 [71, 72]	CC BY-NC
892924356	<i>Milvus migrans</i> [148]	CC0
897981076	Ya Ha Tinda elk project, Banff National Park, 2001-2020 (females) [61, 62, 73-78]	CC0
918219824	ECOPATH, Brown skua, Boulinier et al., Amsterdam Island [149]	CC BY-NC
922263102	H_GRONINGEN - Western marsh harriers ( <i>Circus aeruginosus</i> , <i>Accipitridae</i> ) breeding in Groningen (the Netherlands) [79]	CC0
933711994	Elk in southwestern Alberta [80-91]	CC BY
938783961	MH_ANTWERPEN - Western marsh harriers ( <i>Circus aeruginosus</i> , <i>Accipitridae</i> ) breeding near Antwerp (Belgium) [92]	CC0
985143423	LBBG_ZEEBRUGGE - Lesser black-backed gulls ( <i>Larus fuscus</i> , <i>Laridae</i> ) breeding at the southern North Sea coast (Belgium and the Netherlands) [93]	CC0
986040562	HG_OOSTENDE - Herring gulls ( <i>Larus argentatus</i> , <i>Laridae</i> ) breeding at the southern North Sea coast (Belgium) [94]	CC0
1030734949	Biotelemetry of Bewick's swans [95, 96]	CC0
1049685237	Greater white-fronted goose family migration flight [97, 98]	CC0
1071134107	Herring Gulls ( <i>Larus Argentatus</i> ); Ronconi; Brier Island, Canada [99, 100]	CC0
1077731101	Eurasian Curlews [ID_PROG 1083] [150]	CC BY-NC
1080341217	Herring Gulls ( <i>Larus Argentatus</i> ); Clark; Massachussets, United States [100, 101]	CC0
1080341737	Herring Gulls ( <i>Larus Argentatus</i> ); Ronconi; Sable Island, Canada [100, 102]	CC0
1087068449	Von der Decken's hornbill (Jetz Kenya) [103, 104]	CC0
1088836380	Carnivore movements near Black Rock Forest New York [151]	CC BY
1091848505	gullSpecies_USGS_ASC_argosGPS [105]	CC0
1092737859	GPS calibration data (global) [152]	CC BY
1099562810	O_WESTERSCHELDE - Eurasian oystercatchers ( <i>Haematopus ostralegus</i> , <i>Haematopodidae</i> ) breeding in East Flanders (Belgium) [106]	CC0
1123149708	Ivory gull N Greenland 2018/19 [153]	CC BY-NC
1208105916	Caspian Gulls - Poland [154]	CC0

Continued on next page

Table 4: The list of all Movebank datasets used in this work. (Continued)

ID	Name and references	License
1229945587	Common Crane 2020 (Lithuanian University of Educational Studies; LEU) [155]	CC0
1241071371	Arctic fox Bylot - GPS tracking [156]	CC0
1259686571	LBBG_JUVENILE - Juvenile lesser black-backed gulls ( <i>Larus fuscus</i> , Laridae) hatched in Zeebrugge (Belgium) [107]	CC0
1260886163	Cheetah Pilanesberg National Park, South Africa, 2014-2015 [108, 109]	CC0
1266784970	<i>Corvus corone</i> [ID_PROG 883] [157]	CC BY-NC
1278021460	BOP_RODENT - Rodent specialized birds of prey ( <i>Circus, Asio, Buteo</i> ) in Flanders (Belgium) [110]	CC0
1285079529	Monitoring of <i>Capra ibex</i> (Bovidae) populations in the western alps (project ALCOTRA LEMED-IBEX) [158]	CC BY
1393954358	<i>Cathartes aura</i> MPIAB Cuba [159]	CC BY-NC
1395952585	FTZ: Migrating curlews (data from Schwemmer et al. 2021) [111, 112]	CC0
1410035327	HUJ MPIAB White Stork E-Obs (subset for Carlson et al. 2021) [113, 114]	CC0
1415844328	Moult migration of taiga bean geese to Novaya Zemlya [115, 116]	CC BY
1448377103	Wood stork ( <i>Mycteria americana</i> ) Southeastern US 2004-2019 [117, 118]	CC0
1448409403	Lapwing NFW <i>Vanellus Vanellus</i> [160]	CC BY-NC
1498452485	Variability of White Stork flight patterns prior to earthquakes [161]	CC0
1562253659	LifeTrack White Stork Sarralbe [ID_PROG 1093] [162]	CC0
1605798640	O_BALGZAND - Eurasian oystercatchers ( <i>Haematopus ostralegus</i> , Haematopodidae) wintering on Balgzand (the Netherlands) [119]	CC0
1605799506	O_SCHIERMONNIKOOG - Eurasian oystercatchers ( <i>Haematopus ostralegus</i> , Haematopodidae) breeding on Schiermonnikoog (the Netherlands) [120]	CC0
1605802367	O_VLIELAND - Eurasian oystercatchers ( <i>Haematopus ostralegus</i> , Haematopodidae) breeding and wintering on Vlieland (the Netherlands) [121]	CC0
1605803389	O_AMELAND - Eurasian oystercatchers ( <i>Haematopus ostralegus</i> , Haematopodidae) breeding on Ameland (the Netherlands) [122]	CC0
1606812667	Hawksbill/green turtles Chagos Archipelago Western Indian Ocean [123, 124]	CC0
1671751878	Tchad Redneck Ostrich [163]	CC BY-NC
1841261165	Eurasian wigeon ( <i>Mareca penelope</i> ) Netherlands Lithuania 2018-2019 [125, 126]	CC BY
1907973121	Lowland tapirs, <i>Tapirus terrestris</i> , in Southern Brazil [164]	CC BY-NC
1907974323	Vega gull ( <i>Larus vegae</i> ) - GPS - Russia South Korea Japan [165]	CC BY-NC
295134472	Plains zebra Chamaillé-Jammes Hwange NP [166]	CC BY-NC
307786785	African elephant (Migration) Chamaillé-Jammes Hwange NP [167]	CC BY-NC

Table 5: Number of observations of each taxon in each section of the dataset.

class	order	taxon	#obs		
			train	val	test
Aves	Accipitriformes	Cathartes aura	37609	2186	1703
		Circus aeruginosus	5537	—	—
		Circus cyaneus	540	—	—
		Circus pygargus	1088	—	—
		Gyps himalayensis	4047	736	312
		Haliaeetus albicilla	409	—	—
		Milvus milvus	323	—	—
		Necrosyrtes monachus	4337	951	126
		Neophron percnopterus	2138	—	—
		Pandion haliaetus	2966	181	—
	Anseriformes	Anas penelope	8297	—	—
		Anas platyrhynchos	22207	1970	1484
		Anser albifrons	18871	1122	558
		Anser anser	3549	—	—
		Anser fabalis	6258	—	244
Bucerotiformes	Charadriiformes	Anseriformes	1123	—	—
		Branta bernicla	1566	—	—
		Branta leucopsis	628	—	—
		Cygnus columbianus	602	—	—
		Cygnus cygnus	907	—	—
		Tockus deckeni	4448	—	—
		Coragyps atratus	8653	—	312
		Haematopus ostralegus	44582	5439	3960
		Larus	196	—	—
		Larus argentatus	39655	8237	909
		Larus cachinnans	1021	—	—
		Larus fuscus	91380	8202	1534
		Larus glaucescens	293	—	—
		Larus smithsonianus	900	—	—
Mammalia	Artiodactyla	Larus vegae	10018	—	—
		Numenius arquata	14896	—	1588
		Vanellus vanellus	2661	—	251
		Ciconiiformes	230170	5835	10402
		Ciconia ciconia	7134	—	—
		Gruiformes	Grus grus	189	—
		Passeriformes	Corvus corone	474	—
		Pelecaniformes	Ardea alba	1764	—
		Struthioniformes	Struthio camelus	221	—
		Suliformes	Fregata magnificens	521	—
		Alces alces	3061	—	—
		Bison bison	130	—	—
		Cervus elaphus	158108	6648	3849
		Connochaetes taurinus	30036	755	4065
Reptilia	Carnivora	Ovis dalli	9075	934	983
		Sus scrofa	556	—	—
		Syncerus caffer	498	—	—
		Acinonyx jubatus	239	—	—
		Canis lupus	4811	—	122
		Lynx	426	—	—
		Lynx rufus	4353	—	—
		Panthera onca	1873	—	—
		Vulpes lagopus	580	—	—
		Perissodactyla	Equus quagga	24873	709
		Proboscidea	Loxodonta africana	24870	5696
		Testudines	Chelonoidis	546	—
		Chelonoidis donfaustoi	9386	594	251
		Chelonoidis hoodensis	2251	—	—
		Chelonoidis porteri	12560	789	2283
		Eretmochelys imbricata	1286	—	—
		Testudinidae	8548	—	1859

Table 6: WorldClim bioclimatic variables as listed at <https://www.worldclim.org/data/bioclim.html>.

<u>Variable</u>	<u>Description</u>
BIO1	Annual Mean Temperature
BIO2	Mean Diurnal Range (mean of monthly (max temp – min temp))
BIO3	Isothermality ( $BIO2/BIO7 \times 100$ )
BIO4	Temperature Seasonality (standard deviation $\times 100$ )
BIO5	Max Temperature of Warmest Month
BIO6	Min Temperature of Coldest Month
BIO7	Temperature Annual Range ( $BIO5 - BIO6$ )
BIO8	Mean Temperature of Wettest Quarter
BIO9	Mean Temperature of Driest Quarter
BIO10	Mean Temperature of Warmest Quarter
BIO11	Mean Temperature of Coldest Quarter
BIO12	Annual Precipitation
BIO13	Precipitation of Wettest Month
BIO14	Precipitation of Driest Month
BIO15	Precipitation Seasonality (Coefficient of Variation)
BIO16	Precipitation of Wettest Quarter
BIO17	Precipitation of Driest Quarter
BIO18	Precipitation of Warmest Quarter
BIO19	Precipitation of Coldest Quarter

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