foieGras an R package for animal movement data: rapid quality control, behavioural estimation and simulation

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1 Abstract

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6 Keywords:

1 Introduction

The R package foieGras, pronounced "fwah grah," ...

19 **2 |** foieGras **overview**

The workflow for foieGras is deliberately simple, with much of the usual track data processing checks and formatting handled automatically. The main functions are listed in Table 1. When fitting a model, foieGras automatically detects the type of tracking data location quality classes designations that are typical of Argos data and that can be added to the data by the researcher for other types of track data. Based on the location quality classes and other, optional information on observation errors contained in the data, foieGras chooses an appropriate measurement error model for each observation. This capability allows for combinations of different tracking data types, e.g., Argos and GPS, in a single input data frame and to be fit in a single state-space model.

2.1 | Data preparation

Animal tracking data, consisting of a time-series of location coordinates, can be read into R as a data frame using standard functions such as read.csv. The canonical data format for Argos tracks consists of a data frame with 5 columns corresponding to the following named variables:
id (individual id), date (date and time), 1c (location class), 1on (longitude), 1at (latitude). Optionally, an additional 3 columns, smaj (semi-major axis), smin (semi-minor axis), eor (ellipse orientation), providing Argos error ellipse information may be included.

Other types of track data can be accommodated, for example, by including the 1c column where all 1c = "G" for GPS data. In this case, measurement error in the GPS locations is assumed to

Table 1: Main functions for the R package foieGras

Function	Description
fit_mpm	Fit a Move Persistence Model to location data
fit_ssm	Fit a State-Space Model to location data
fmap	Plot fitted/predicted locations on a map with or without a defined projection
grab	Extract fitted/predicted/observed locations from a foieGras model, with or without projection information
osar	Estimate One-Step-Ahead Residuals from a foieGras SSM
sim	Simulate individual animal tracks with Argos LS or KF errors
simfit	Simulate animal tracks from 'fG_ssm' fit objects
sim_filter	Filter tracks simulated with 'simfit' according to similarity criteria
plot.fG_ssm	Plot the fit of a foieGras SSM to data
plot.fG_osar	Plot One-Step-Ahead Residuals from a foieGras SSM
plot.fG_mpm	Plot move persistence estimates as 1-D or 2-D (along track) time-series
plot.fG_sim	Plot simulated animal tracks

have a standard deviation of 0.1 x Argos class 3 locations (approximately 30 m). Other types of track data can be considered in a similar manner (see the package vignette for further details).

9 2.2 | State-space model fitting - fit_ssm

State-space models are fit using fit_ssm. There are a large number of options that can be set in fit_ssm (see Suppl for details). We focus only the essential options here:

- data the input data structured as described in 2.1
- vmax a maximum threshold speed (ms⁻¹) to help identify potential outlier locations
- model the process model to be used

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• time.step the prediction time interval (h)

The function first invokes an automated data processing stage where the following occurs: 1)
data type (Argos Least-Squares, Argos Kalman Filter/Smoother, GPS, or General (e.g., processed light-level geolocations, acoustic telemetry, coded VHF telemetry) is determined; 2) datetimes are converted to POSIXt format, chronological order is ensured, and duplicate datetime records are removed; 3) observations occurring less than min.dt seconds after a prior observation are removed; 4) a speed filter [sda from the trip R package; Sumner et al. (2009)] is used to identify potential outlier locations; 5) locations are projected from spherical lon-lat coordinates to planar x,y coordinates in km.

The function then fits a state-space model to the processed data, where the process model (currently, either a continuous-time rw or a continuous-time crw) is specified by the user and the measurement model(s) are selected automatically (see I. D. Jonsen et al., 2020 for model details). The model is fit by numerical optimization of the likelihood using either the optim or nlminb R function. The R package TMB, Template Model Builder (Kristensen et al., 2016), is used to compute the gradient function in C++ via reverse-mode auto-differentiation and the Laplace Approximation is used to integrate out the latent states (random effects). Fits to a single versus multiple individuals are handled automatically, with sequential SSM fits occurring in the latter case. No hierarchical or

pooled estimation among individuals is currently available.

fit_ssm returns a foieGras fit object (a nested data frame with class fG_ssm). The outer data frame lists the individual id(s), basic convergence information and a list with class ssm. This list contains dense information on the model parameter and state estimates, predictions, processed data, optimizer results, and other diagnostic and contextual information. Users can extract a simple data frame of SSM fitted (location estimates corresponding to the, typically irregular, observation times) or predicted values (locations predicted at regular time.step intervals) using the grab function.

69 2.3 | Model checking and visualisation - osar, plot, fmap

Before using fitted or predicted locations, a model fit should be checked and visualised to confirm that the model adequately describes the data. In linear regression and a variety of analogous 71 methods, goodness-of-fit can be assessed by calculating standard residuals such as Pearson or 72 deviance residuals. There is no simple way to calculate residuals for latent variable models that have 73 non-finite state-spaces and that may be nonlinear, but they can be computed based on iterative 74 forecasts of the model (Thygesen et al., 2017). The osar function computes one-step-ahead 75 (prediction) residuals and uses the oneStepPredict function from the TMB R package to make this as efficient as possible. A set of residuals are calculated for the x and y values corresponding to 77 the fitted values from the SSM and returned as an fG_osar object. 78

A generic plot method provides an easy way to visualise the fG_osar residuals. Time-series plots of the prediction residuals can be used to detect temporal changes in goodness-of-fit. Quantile-quantile plots of residuals against standard normal quantiles can be used to detect departures from normality. Sample autocorrelation function plots of the residuals are useful for detecting autocorrelation not accounted for by the model. Assessing residual autocorrelation can be particularly important as Argos locations, for example, are themselves derived from a time-series model (Lopez et al., 2015) which can introduce additional autocorrelation in the location errors.

State-space model fits to data can also be visualised by using the generic plot function on an fG_ssm data frame. Options exist to plot fitted or predicted values along with observations as either paired, 1-D time-series or as 2-D tracks with confidence intervals or ellipses, respectively. These plots provide a more intuitive and rapid method for assessing SSM fits to data, however, they do not replace the residual diagnostics. Fitted fG_ssm data frames can be mapped using the fmap function for single or multiple individuals. Estimated tracks can be displayed with or without confidence ellipses, observations, and/or a projection and maps of single tracks can be coloured by date.

2.4 | Behavioural estimation - fit_mpm

The fit mpm function fits a simple move persistence model to estimate a continuous-valued, time-95 varying latent variable that indexes changes in movement behaviour (I. Jonsen et al., 2019). This variable measures the autocorrelation in speed and direction between consecutive pairs of movements such that high values correspond to fast, directed movements at one end of the continuum 98 and low values correspond to slow, tortuous movements at the other end. It's important to note 99 that this approach is unlike hidden Markov models (McClintock & Michelot, 2018; Michelot et al., 100 2016) and some state-space models (I. D. Jonsen, 2016) as there is no notion of discrete be-101 havioural states that animals periodically switch between. Nonetheless, move persistence can be used to identify objectively places where animals spend disproportionately more or less time, and with extensions be correlated with environment or other covariates (See Examples 3.x). 104

The move persistence model assumes that locations are absent of measurement error and can occur either irregularly or regularly in time. fit_mpm takes either a fG_ssm data frame as input or a

data frame with the follow variables: id, date, x, y, where x and y coordinates can be planar x, y 107 or spherical long, lat. This latter input format allows the model to be fit easily to GPS or other 108 tracking data with negligible measurement error. When the data contain multiple individuals, the 109 default model is fit jointly by assuming all individuals share the same move persistence variance 110 parameter. There is an option to fit the model separately to each individual. The time-series of 111 estimated move persistence with confidence intervals can be visualized by using the generic plot 112 function with the resulting fG mpm data frame. Visualization of move persistence along the 2-D 113 tracks can be plotted or mapped by using the plot or fmap functions, respectively, and supplying 114 both the fG_mpm and fG_ssm nested data frames. When using fit mpm on, for example, GPS 115 tracking data that do not require state-space filtering, the movement persistence estimates can be extracted from the fG mpm data frame using the grab function and subsequently merged with the 117 observed track data for visualization. 118

119 2.5 | Simulation - sim, simfit, sim filter

Track simulation can be a helpful, yet informal, way of evaluating the degree to which statistical 120 movement models capture essential features of animal movement data (Michelot et al., 2017). 121 Michelot et al. (2016) advocate comparison of simulated tracks from fitted hidden Markov models 122 to the observed tracks as a means of identifying potential weakness in the hidden Markov model 123 formulation. Here, we suggest that the rw and crw state-space models and the mpm model can be fit to track data simulated from different movement processes to evaluate robustness of location 125 and movement persistence estimates to model mis-specification. We illustrate this idea in section 126 3.x by drawing on flexibility in the sim function that allows a variety of movement processes to be 127 simulated. 128

Simulation is also used frequently to infer habitat availability, e.g., a null model of the distribution of foraging animals in the absence of external drivers, in habitat utilization studies (Hindell et
al., 2020; Raymond et al., 2015). The simfit function extracts movement parameters from a
fg_ssm fit object and simulates an arbitrary number of random tracks of the same duration from
these parameters. The argument cpf = TRUE ensures that the simulated tracks start and end
at approximately the same location, thereby simulating a central place forager. Something about
sim_filter here...

3 | Examples

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We illustrate the main capabilities of foieGras through a series of examples using real and simulated tracking data. These examples are purely for demonstration purposes and not intended as a comprehensive guide for conducting animal tracking data quality control or analysis with foieGras.

Complete code for reproducing the examples and in-depth understanding of foieGras functions are provided as supplements.

3.1 | Southern Elephant seal - SSM validation with prediction residuals

We use a subadult male southern elephant seal track included in foieGras (sese1), sourced 143 from from the Australian Integrated Marine Observing System (IMOS; data publicly available via 144 imos.aodn.org.au) deployments at Iles Kerquelen in collaboration with the French IPEV and SNO-MEMO programmes. We fit both the rw and crw models using fit_ssm with a speed filter threshold 146 (vmax) of 4 ms⁻¹ and a 12-h time step. We calculate prediction residuals using osar, and then use 147 the generic plot method for osar residuals to assess and compare the model fits (Fig. 1). The 148 plots of predicted states on top of the observations suggests both models yield similar fits (Fig. 149 1a,b), however, there are marked trends in the time-series of residuals for the rw model fit (Fig. 1c) 150 and the rw ACF's reveal consistent positive autocorrelation in the prediction residuals (Fig. 1e). The

corresponding crw prediction residuals show no apparent trends through time and have relatively little autocorrelation (Fig. 1d,f), implying that the crw provides a better fit to the data.

154 3.2 | Assessing SSM robustness with simulated data

Using the sim function, we simulate animal movement tracks with a variety of plausible movement patterns. We use these simulated data to examine the SSM's goodness-of-fit to tracks generated by processes that differ from the rw and crw process models within the SSM.

158 3.3 | Inferring movement persistence as an index of behaviour from Argos and GPS data

Drawing on an expanded version of the data used in 3.1, we quality control and infer movement 159 persistence along five southern elephant seal tracks using the fit ssm and fit mpm functions. 160 These data can be accessed in foieGras via the call: data(sese, package = 'foieGras'). To 161 illustrate how the method can accommodate other types of animal tracking data, we also infer movement persistence along five little penguin (Eudyptula minor) GPS tracks from Montague Is-163 land, NSW Australia, described in Phillips et al. (2021). The GPS tags were programmed such 164 that realized sampling rates were approximately 3 - 4 s, which results in extremely autocorrelated 165 location time-series that are not amenable for movement persistence estimation. To alleviate this, 166 we resampled the raw GPS data to approximately 5 min resolution using the resample track 167 function from the R package amt (Signer et al., 2019). We then compare move persistence estimates obtained by fitting directly to the resampled GPS tracks versus those obtained from tracks 169 that were interpolated from the raw GPS data to a fixed 5-min interval using the fit_ssm function. 170

71 3.4 | Simulating tracks from foieGras model fits



Figure 1: Selected diagnostic plots for assessing rw (a,c,e) and crw (b,d,f) state-space model fits to a southern elephant seal track. Top panels (a,b) are plots of predicted states (red; regular 12-h time intervals) and observations (blue) with pre-filtered observations (orange; ignored by the SSM), using the plot.fG_ssm function. Panels c,d are time-series plots of the prediction residuals for the x and y coordinates of each fitted state. Panels e,f are autocorrelation functions of the prediction residuals. All residual plots generated using the plot.fG_osar function.

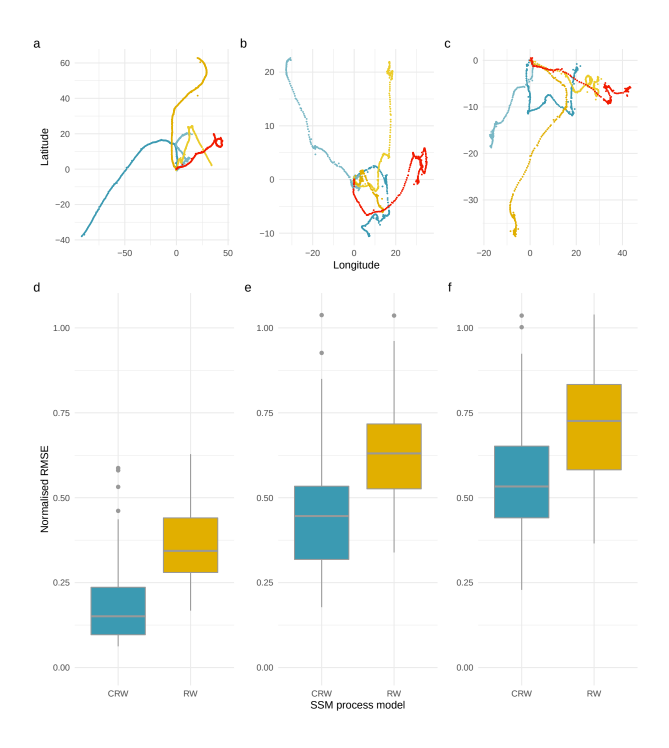
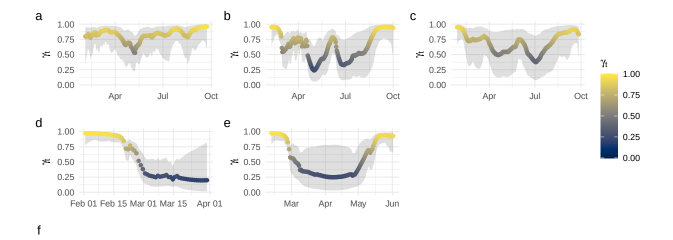


Figure 2: Five example tracks simulated from the correlated random walk process model (crw, a), a 2-state crw model (b), and the movement persistence model (mpm, c). Normalised Root Mean Squared Errors of state-space models fit with either the crw or random walk (rw) process model to 50 simulated crw tracks (d), 2-state crw tracks (e), and mpm tracks (f).



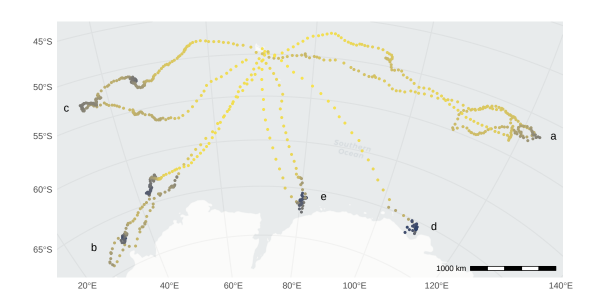


Figure 3: Inferred move persistence, γ_t , 1-D time-series for five southern elephant seals (a-e; grey envelopes are 95 % Cl's) and along their 2-D tracks (f; track labels, a-e, correspond to the 1-D time-series plots). Locations associated with low move persistence (blue) are indicative of slow, undirected movements, whereas high move persistence (yellow) is indicative of faster, directed movements. The lowest move persistence tends to occur at the distal end of foraging trips, furthest from the colony on lles Kerguelen, suggesting these bouts of low move persistence are associated with foraging activity.

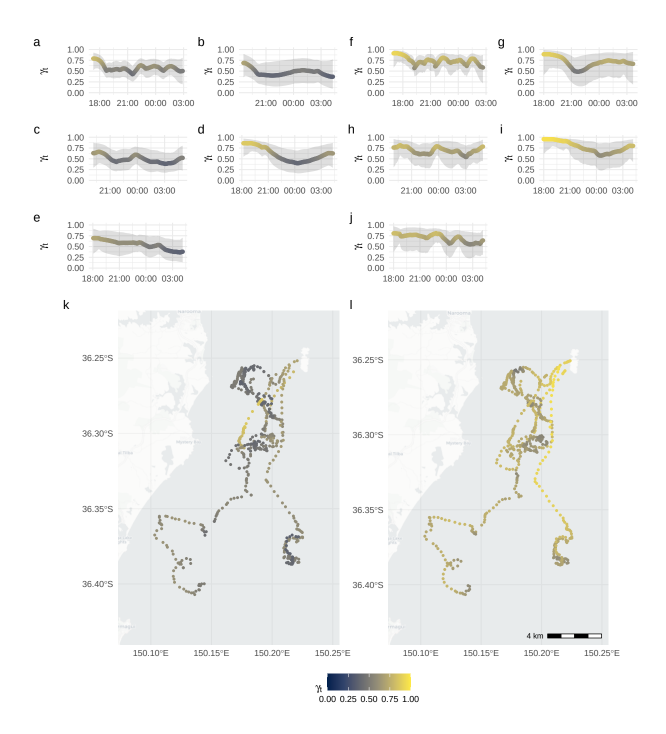


Figure 4: Inferred move persistence, γ_t , 1-D time-series (a-j; grey envelopes are 95 % Cl's) and along little penguin GPS tracks (k,l). Colour palette as in 3. Movement persistence was estimated directly from GPS data resampled to approximate 5-min intervals (a-e, k) and estimated from SSM-predicted locations with a regular 5-min interval (f-j, l). Overall movement persistence patterns are similar but note the consistently higher estimates obtained by fitting to the SSM-predicted locations.

172 4 | Discussion

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190 Author's Contributions

191 IDJ developed the R package; WJG contributed harp seal data and to the R package; LP, GC, and 192 RGH contributed little penguin data; IDJ and TAP developed the state-space models; IDJ wrote 193 an initial draft of the manuscript with a contribution from WJG; all authors edited the manuscript.

194 Data Accessibility

All code mentioned here is provided in the foieGras package for R available on CRAN at https: //CRAN.R-project.org/package=foieGras. The development version of the package is available on GitHub at https://github.com/ianjonsen/foieGras. Data used in the examples are available at...

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