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#### RH: ESTIMATING MACROEVOLUTIONARY LANDSCAPES

# A GENERAL MODEL FOR ESTIMATING MACROEVOLUTIONARY LANDSCAPES

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Abstract.

The evolution of quantitative characters over long timescales is often studied using 20 stochastic diffusion models. The current toolbox available to students of macroevolution is 21 however limited to two main models: Brownian motion and the Ornstein-Uhlenbeck process, 22 plus some of their extensions. Here we present a very general model for inferring the dynamics 23 of quantitative characters evolving under both random diffusion and deterministic forces of 24 any possible shape and strength, which can accommodate interesting evolutionary scenarios 25 like directional trends, disruptive selection, or macroevolutionary landscapes with multiple peaks. This model, which we call FPK, is based on a general partial differential equation 27 widely used in statistical mechanics: the Fokker-Planck equation. We first explain how this model can be used to describe macroevolutionary landscapes over which quantitative traits evolve and, more importantly, we detail how it can be fitted to empirical data. Using simulations, we show that the model has good behavior both in terms of discrimination from alternative models and in terms of parameter inference. We provide R code to fit the model to empirical data using either maximum-likelihood or Bayesian estimation, and illustrate 33 the use of this code with two empirical examples of body mass evolution in mammals. FPK 34 should greatly expand the set macroevolutionary scenarios that can be studied since it opens 35 the way to estimating macroevolutionary landscapes of any conceivable shape. (Keywords: adaptation, bounds, macroevolution, maximum-likelihood estimation, MCMC 37

methods, selection, phylogenetic comparative data)

Understanding the evolution of phenotypes over geological timescales is one of the fundamental goals of macroevolution (Simpson (1944)). Phenotypic evolution is typically inferred either from time series of measurements obtained in the fossil record (Hunt (2007)) or from the distribution of phenotypic characters at the tips of a phylogenetic tree (O'Meara (2012)). In both cases, one then fits stochastic models for the evolution of traits on single lineages or on phylogenies, all of which treat trait evolution as a diffusion process that may or may not be influenced by deterministic forces (O'Meara (2012)).

Many approaches attempt to bridge the gap between microevolutionary process and macroevolutionary pattern by interpreting model parameters in the context of the dynamics of evolution on adaptive landscapes (Wright (1932); Simpson (1944); Arnold et al. (2001); Arnold (2014)). Recent years have revitalized this connection with the development of nu-51 merous methodological tools specifically aimed at inferring 'macroevolutionary landscapes' 52 (Hansen et al. (2008); Ingram and Mahler (2013); Eastman et al. (2013); Uyeda and Har-53 mon (2014)). Such landscapes almost certainly do not reflect static landscapes upon which 54 populations evolve over long time scales; instead, these landscapes are most productively 55 described as representing the movements of adaptive peaks over million-year time scales (Hansen (1997); Uyeda et al. (2011); Uyeda and Harmon (2014)). In particular, a peak 57 on such a landscape might not be a phenotypic optimum in any particular generation of 58 evolution of a lineage, but instead might represent a long-term average peak location on a 59 dynamic landscape. Throughout this paper, we will refer to these as 'macroevolutionary landscapes', which summarize patterns of trait evolution averaged over many generations.

Comparative methods to infer macroevolutionary landscapes are all based on the
Ornstein-Ulhenbeck (OU) process (Hansen (1997)), which was itself strongly inspired by
the original concept of adaptive landscape in population genetics (Lande (1976)). Under

the OU process, a continuous trait evolves under both random diffusion (i.e. Brownian motion, Edwards and Cavalli-Sforza (1964)) and a force that brings back the trait close to an optimal value. Following models from quantitative genetics (e.g. Lande (1976)), these two 67 components of the macroevolutionary OU process are sometimes interpreted as genetic drift 68 and stabilizing selection. However, such an interpretation is almost always overly simplistic. 69 First, many other processes can generate evolution following an OU model. For example, the shape of the peak and, in turn, the dynamics of selection and drift within populations may 71 be less important for long-term patterns than the movement of the peak itself. Under such a scenario, both the diffusion and deterministic components of OU reflect peak movement, and both are strongly dependent on the dynamics of both selection and drift. Second, the actual parameters of OU models are almost always incompatible with Lande's model of evolution on a static adaptive landscape (Estes and Arnold (2007); Uyeda and Harmon (2014)). However, even if we do not interpret diffusion as drift and determinism as selection, it is still useful to divide macroevolutionary dynamics into these two components. Any factor that leads to trait change that is random in direction from one generation to the next (e.g. drift, randomly varying selection, plasticity due to random environmental noise) will affect the diffusion component, and any factor that leads to predictable change towards some particular value 81 (e.g. selection, predictable patterns of peak movement over time, developmental constraints 82 towards certain values) will be seen in the deterministic component. Various extensions of 83 the OU model have been proposed in recent years, including different optima in different clades, either determined a priori (Hansen (1997); Butler and King (2004)) or not (Ingram and Mahler (2013); Uyeda and Harmon (2014)), varying rates of diffusion and intensities of attraction towards optima in different clades (Beaulieu et al. (2012)), evolution of the optimum itself (Hansen et al. (2008)), or the possibility to study multivariate evolution (Bartoszek et al. (2012)). The complexity of all of these extensions, however, leads to difficulties in model identifiability and parameter estimation (e.g. Khabbazian et al. (2016)).

While these models cover a wide range of possible scenarios, they are restricted to two main kinds of macroevolutionary landscapes: (i) macroevolutionary landscapes with a single peak continuously moving in time (Hansen et al. (2008)) and (ii) macroevolutionary landscapes 93 with one or several peaks, eventually of varying heights (varying attraction strengths) and 94 widths (varying ratios of diffusion rate to attraction strength). Importantly, in the second 95 case the different peaks are experienced at different epochs or by different lineages, so that no single lineage evolves in an macroevolutionary landscape with multiple peaks (see below). In 97 addition, another type of macroevolutionary landscape might be described by the bounded Brownian motion model (BBM, Boucher and Démery (2016)). BBM was developed as a model of neutral evolution between bounds, but it could also describe macroevolutionary 100 landscapes in which one part of phenotypic space (i.e., traits values between the bounds) 101 has high but constant fitness while other phenotypes have null fitness, a scenario related to 102 holey adaptive landscapes (Gavrilets (1997)). 103

As can be seen from this short overview of existing methods, the types of macroevo-104 lutionary landscapes that can be estimated from comparative data are still rather limited. 105 For example, disruptive selection is central to the theories of ecological speciation (Doe-106 beli (1996); Nosil (2012)) and adaptive radiation (Schluter (2000)), in which diverging lin-107 eages adapt to different ecological niches. These theories are being increasingly appreci-108 ated in the macroevolution community and ecological speciation and/or adaptive radiation 109 are frequently invoked as explanations for the diversity of extant clades (Soulebeau et al. 110 (2015)). However, despite recent theoretical advances in modeling interspecific interactions 111 over macroevolutionary timescales (e.g. Nuismer and Harmon (2015); Drury et al. (2016)), 112 we still lack proper tools to infer disruptive selection on phylogenies since macroevolutionary landscapes that contain multiple local optima cannot be inferred from phylogenetic 114 comparative data. The current alternative is to model phenotypic evolution in multimodal 115 macroevolutionary landscapes using OU models with multiple optima (Butler and King 116

(2004); Uyeda and Harmon (2014)). In this framework, cases in which transitions between peaks are frequent can be interpreted as evidence of a changing of the adaptive environment, 118 rather than the existence of multiple, simultaneously existing peaks among which lineages 119 alternate. Such a model in which several adaptive peaks are simultaneously present in the 120 adaptive landscape experienced by all species in the clade would be a step towards a more explicit model for disruptive selection. 122

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In this paper we introduce a general model for the evolution of continuous characters 123 on phylogenies that can accommodate macroevolutionary landscapes of any shape and thus 124 attempts to provide solutions to the limitations mentioned above. In this model the con-125 tinuous trait of interest evolves under random diffusion but is also subject to deterministic 126 change following a macroevolutionary landscape of any possible shape and strength. We 127 present algorithms for both maximum-likelihood and Bayesian estimation of model param-128 eters, i.e. the value of the trait at the root of the tree, the diffusion rate and the shape 129 of the macroevolutionary landscape. Using simulations, we show that this model is easily 130 distinguishable from other models of trait evolution like BM, OU, and BBM. Parameter 131 estimation is also generally reliable, and in particular the shape of the macroevolutionary 132 landscape can be accurately estimated as long as it has been fully explored by the clade 133 evolving on it. We also show how alternative hypotheses can be statistically tested in empir-134 ical datasets. Our approach opens new avenues for macroevolutionary research: it renders 135 possible the detection of evolutionary trends from neontological data only, but also inference 136 of disruptive selection or of even more complex scenarios in which the macroevolutionary 137 landscape contains multiple peaks. 138

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#### General presentation of the Fokker-Planck-Kolmogorov model

We introduce a general model for the evolution of continuous traits on phylogenies, 141 in which a trait x undergoes a random walk (i.e., BM) that is biased by a deterministic 142 force that can be of any shape and strength. The force biasing trait evolution derives from 143 a potential V(x): differences in the values of the potential generate a force that attracts the trait towards regions of trait space in which the potential is the lowest and at each point x, the BM process is biased by a force proportional to -V'(x). This process can be modeled 146 using the Fokker-Planck equation, a partial differential equation widely used in statistical 147 mechanics to describe the time evolution of the probability density of an observable under the 148 influence of both random and deterministic forces (Risken (1984)). In population genetics, 149 the Fokker-Planck equation has been used to model changes in allele frequencies and is 150 better known as the Kolmogorov forward equation (Wright (1945)). This is why we label 151 the present model FPK. Under FPK, the probability density  $p(x, x_0, t)$  of the position of a 152 trait x with initial value  $x_0$  after time t has elapsed follows: 153

$$\frac{\partial p}{\partial t}(x, x_0, t) = \frac{\sigma^2}{2} \frac{\partial}{\partial x} \left[ \frac{\partial p}{\partial x}(x, x_0, t) + p(x, x_0, t) \frac{\partial V}{\partial x}(x) \right]. \tag{1}$$

In this equation, the evolution of the probability density of the trait (left hand side) is determined by both random diffusion (*i.e.*, BM; first term on the right hand side) and a deterministic force set by the derivative of the potential (second term on the right hand side). The factor  $\sigma^2/2$  on the front of the second term in the right hand side is chosen so that the stationary distribution for the probability density is

$$p^*(x) = \lim_{t \to \infty} p(x, x_0, t) = \mathcal{N} \exp(-V(x)), \tag{2}$$

where  $\mathcal{N}$  is a normalization factor. Finally, the initial position of the trait,  $x_0$ , gives the initial condition for this partial differential equation.

In summary, the potential V(x) determines the force  $-\sigma^2 V'(x)/2$  that is exerted on 161 the trait over the interval, and the process has a stationary distribution, which is propor-162 tional to  $\exp(-V(x))$  but does not depend on  $\sigma^2$ . The force represents the deterministic 163 component of trait evolution, since it pulls traits towards specific values and the stationary 164 distribution can be interpreted as a macroevolutionary landscape because trait values are 165 attracted towards regions of trait space with the lowest potential, which themselves corre-166 sponds to peaks of  $\exp(-V(x))$ . Fig. 1 shows how the potential, the deterministic force, and 167 the macroevolutionary landscape are related. In the remainder of this article, we will use 168 the term macroevolutionary landscape to refer to  $\mathcal{N}\exp(-V(x))$  and will avoid mentioning 169 the potential (V(x)) as much as possible. It is important to note here that the evolutionary 170 rate,  $\sigma^2$ , is not a measure of the strength of the random component of the process. Indeed, 171  $\sigma^2$  determines both the intensity of the random diffusion component (first term in the right 172 side of Eq. 1) and of the deterministic force exerted on the trait (second term in the right 173 side of Eq. 1). The relative strengths of the random and deterministic components of the 174 process are better captured by the variations in the stationary distribution of the process, 175  $p_{\text{max}}^*/p_{\text{min}}^* = \exp(V_{\text{max}} - V_{\text{min}})$ : a ratio close to one means that diffusion dominates, while 176 deterministic forces are important if this ratio is large. 177

We can also define a characteristic time  $T_c$  representing the time it takes for the trait to explore the macroevolutionary landscape. Online Appendix I shows how this characteristic time can be calculated. Over short time periods, *i.e.* for  $t \ll T_c$ , the trait has no time to fully explore the macroevolutionary landscape. Over long time periods, *i.e.* for  $t \gg T_c$ , the trait explores the whole macroevolutionary landscape; the probability density thus reaches its stationary distribution (Eq. 2). An important feature of the characteristic time is that very low values of the macroevolutionary landscape can considerably slow down the exploration.

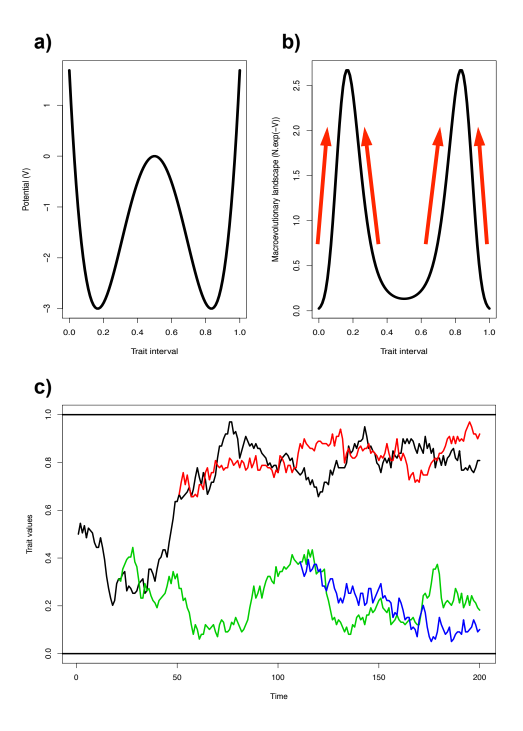


Figure 1: Behaviour of the FPK model. a) Example of a potential with  $V(x) = 3x^4 - 6x^2$ . b) The stationary distribution of the process,  $\mathcal{N} \exp(-V(x))$ , can be interpreted as a macroevolutionary landscape. Wells of the potential become peaks in the macroevolutionary landscape. Differences in the potential generate a force -V'(x), which attracts trait values towards the two peaks of the macroevolutionary landscape, as indicated by the arrows. c) One simulation of the evolution of a trait in a clade of four species in this macroevolutionary landscape: the x-axis shows time and the y-axis the trait value of each species, in different colors. The ancestral trait value lies between the two peaks, and species' traits get attracted towards one of the peaks, which have equal heights. The bounds of the trait interval, represented by thick horizontal lines, are not reached during the process and thus do not influence the evolution of the trait in this case.

For instance, if two maxima are separated by a local minimum with potential difference  $\Delta V$ , the characteristic time follows the Arrhenius law for large  $\Delta V$ :  $T_c \sim \exp(\Delta V)$  (Gardiner (1985)). If there is a single peak in the macroevolutionary landscape, like in the OU process, the characteristic time is proportional to the phylogenetic half-life of the process (Online Appendix I).

### Calculation of the likelihood

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The likelihood of the FPK model given a phylogenetic tree and observed values of the trait at the tips of the tree is obtained by multiplying the probability densities along each branch of the tree and integrating over all possible trait values at the internal nodes:

$$\mathcal{L} = \int \left( \prod_{i \in I \cup T} p(x_i, x_{parent(i)}, t_i - t_{parent(i)}) \right) \prod_{i \in I} dx_i, \tag{3}$$

where I is the set of internal nodes (excluding the root), T is the set of tips,  $x_i$  is the value of the trait at the node i,  $t_i$  is the time at node i and parent(i) is the parent of the node i. Computing the likelihood numerically thus requires integrating over all possible values of the trait at interior nodes of the tree, which makes it computationally challenging. Since the distribution of the trait at the tips of the tree is often not multivariate normal, fast methods like Generalized Least-Squares (Grafen (1989)), phylogenetic independent contrasts (Felsenstein (1985); Freckleton (2012)), or the 3-point algorithm (Ho and Ané (2014b)) cannot be used either.

To compute the likelihood of FPK we instead discretize the trait interval by considering only a set of n points equally spaced between two extreme values,  $B_{\min}$  and  $B_{\max}$ , a procedure already used for the BBM model (Boucher and Démery (2016)). In the following of this article, we call this regular set of points the *grid*. Online Appendix II shows how the continuous evolution equation for the probability density (1) can then be cast in a matrix form and that these discretized equations converge to the continuous one (1) as we increase
the number of points used to discretize the trait interval  $(i.e., n \to \infty)$ .

The use of this discretization procedure imposes that bounds on the trait interval exist: these two bounds are denoted  $B_{\min}$  and  $B_{\max}$ . As done for the BBM model, we make the hypothesis that these two bounds are reflective and calculate the probability density of a trait evolving under FPK using the method of images (Jackson (1998)), i.e. by cutting and reflecting the probability density of the unbounded model an infinite number of times at each one of the two bounds (Boucher and Démery (2016)). However, these bounds need not be reached by the trait and need not even influence the evolutionary process. We thus distinguish two different cases: 

- 1. In situations where the stationary distribution of the FPK model converges to zero when trait values tend to  $+\infty$  or  $-\infty$ , bounds located far apart from the observed trait interval will most likely never be reached: there is a strong force opposing trait evolution towards low-lying regions of the macroevolutionary landscape. Such scenarios can be seen on the top row of Fig. 2. In this case the discretization procedure only introduces a very slight approximation to the likelihood function, and the FPK model is unbounded in practice.
- 2. While we need to introduce bounds for technical reasons, we can also actually make use of them. The same model can thus be used to model situations in which reflective bounds on each side of the trait interval are actually reached during trait evolution. Such scenarios can be seen on the bottom row of Fig. 2. For clarity's sake, in these situations, we will call the model BBMV, for bounded BM with an evolutionary potential. BBMV is actually the most general model, since FPK is a special case of it with bounds set to  $-\infty$  and  $+\infty$ .

The discretization procedure that we use enables calculating the transition matrix

between different points on the grid. Once this matrix is obtained, we calculate the likelihood of the model as is done for the evolution of discrete characters (*i.e.*, the Mk model, Lewis (2001)). We use the pruning algorithm (Felsenstein (1973)) to calculate the likelihood of the model, starting from the probability density of the trait at the tips, and propagating it down to the root of the tree. Finally, we treat the value of the trait at the root of the phylogenetic tree,  $x_0$ , as a parameter of the FPK model. This makes comparison with other models of evolution possible, since most implementations include the root value as a parameter.

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The precision of the discretization procedure (i.e., the number of points used to discretize the trait interval) naturally influences the precision of the numerical calculation of the likelihood (Boucher and Démery (2016)) and the accuracy in the estimation of the shape of the macroevolutionary landscape, but on the other hand calculation grows quadratically with the number of points (the transition matrix M has  $n^2$  terms). In the rest of this article all simulations have been run with n = 50 points on the trait grid. This value was used because of the large number of simulations we ran, but we generally recommend people working on a single empirical case to increase this number if they have the computational facilities to do so.

# Shape of the macroevolutionary landscape

The model we have presented above can accommodate any shape of the macroevolu-249 tionary landscape. However, in order to infer macroevolutionary landscapes from empirical 250 data we need to specify a parametric shape for it and optimize its parameters. One pos-251 sibility to do so would have been to use step functions with different values at each point 252 in the trait grid, but this would have lead to a very large number of parameters to esti-253 mate. Using combinations of sine functions of various periods and amplitudes would also 254 have been possible, but their periodicity renders optimization difficult. Instead we chose a 255 polynomial function with only three terms, such that  $V(x) = ax^4 + bx^2 + cx$ . We discard 256

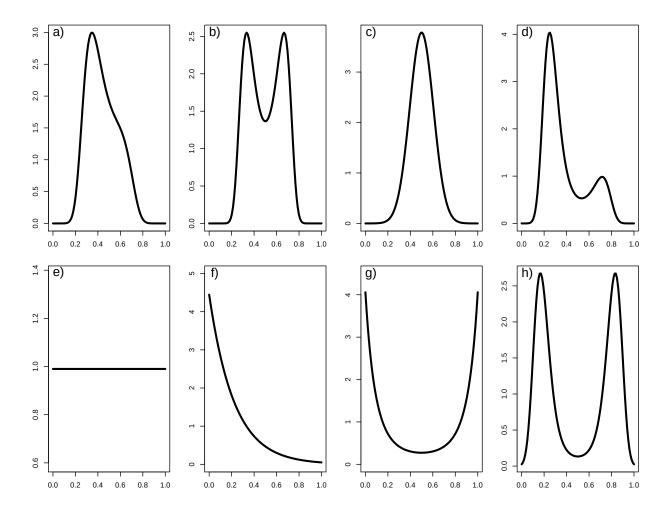


Figure 2: Contrasted shapes of macroevolutionary landscapes that can be obtained using a parametric form of the potential with only three parameters:  $V(x) = ax^4 + bx^2 + cx$ . The x-axis show trait values, always bounded by 0 and 1, and the y-axis the associated density of the stationary distribution of the FPK process. The top row shows situations in which the bounds on the trait interval do not influence the process since the density of the stationary distribution is null at and close to them, corresponding to the pure FPK model. The bottom row shows situations in which the density of the stationary distribution is strictly positive at the bounds and in which bounds can thus be reached, corresponding to what we label the BBMV model. These height scenarios are the ones used in our simulations, with corresponding letter codes.

the term  $x^3$  because any function of the form  $f(x) = ax^4 + bx^3 + cx^2 + dx$  can be written as  $f(x) = a(x - x_0)^4 + b'(x - x_0)^2 + c'(x - x_0) + d'$  with  $x_0 = -b/(4a)$ . This means that 258 adding a term proportional to  $x^3$  amounts to a translation of the potential. No constant term 259 needs to be added to this polynomial function either, since it is the derivative of V(x) that 260 controls the dynamics of the model (see above). This shape of the potential can approximate 261 a variety of scenarios, including flat landscapes (i.e., BBM, V(x) = 0), linear trends (e.g., 262 V(x)=x), domed (e.g.,  $V(x)=-x^2$ ) or U-shaped (e.g.,  $V(x)=x^2$ ) macroevolutionary 263 landscapes, but also macroevolutionary landscapes with two central peaks of equal (e.g., 264  $V(x) = x^4 - x^2$ ) or different heights (e.g.,  $V(x) = x^4 - x^2 + x$ ). Fig. 2 shows a variety of 265 shapes of the macroevolutionary landscape that can be obtained with this parametric func-266 tion, either under the pure FPK (no bounds in practice) or the BBMV model (bounds are 267 actually reachable). Finally, note that both BM and the OU model are special cases of the 268 FPK model: BM corresponds to V(x)=0 and OU to  $V(x)=(\alpha/\sigma^2)x^2-(2\alpha\theta/\sigma^2)x$ , where 269  $\alpha$  and  $\theta$  are the attraction strength and optimum of the OU model, respectively (Hansen 270 (1997)). 271

# Maximum-likelihood inference of model parameters

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The FPK model has five parameters: the value of the trait at the root of the tree  $x_0$ , 273 the evolutionary rate  $\sigma^2$ , and the three coefficients determining the shape of the macroevo-274 lutionary landscape. We have implemented maximum-likelihood estimation of the FPK 275 model in the R statistical environment (R Core Team (2016)). All functions needed to 276 fit the model to empirical data are freely available from the following Github repository: 277 https://github.com/fcboucher/BBMV, and only depend on functions from the ape 278 package (Paradis et al. (2004)). We have verified that the likelihoods obtained from our 279 code are compatible with likelihoods for other models of trait evolution implemented in the 280 fitContinuous function of package geiger (Pennell et al. (2014)). This makes comparison 281

between FPK and other evolutionary models like BM or OU possible, using the Akaike information critetion (AIC) for example. Maximum-likelihood inference of model parameters 283 is conducted using the optim function in R with the Nelder-Mead optimization routine (i.e., 284 the simplex method, although our code also allows for optimization using the BFGS method 285 with box constraints), which was found to perform better than other optimization routines 286 following preliminary tests. For better numerical precision, we do not directly optimize the 287 evolutionary rate,  $\sigma^2$ , but rather  $\log(\sigma^2/2)$ . In cases where obvious bounds on the trait values 288 exist, for example if the trait under study is a proportion or a probability, the user can spec-289 ify the values of the bounds on the trait interval and fit the BBMV model. In cases where no 290 actual bounds are suspected to exist and the user wants to fit the pure FPK model, we place 291 artificial bounds far away from the observed trait interval (i.e.,  $B_{min} = x_{min} - (x_{max} - x_{min})/2$ 292 and  $B_{max} = x_{max} + (x_{max} - x_{min})/2$ . 293

In order to test complex macroevolutionary landscapes against simpler ones, we have 294 also written R functions for fitting the FPK and BBMV models with simpler macroevolution-295 ary landscapes, i.e.  $V(x) = bx^2 + cx$ , V(x) = cx, and V(x) = 0. Alternatives shapes of the 296 macroevolutionary landscape can then be statistically compared based on their likelihoods, 297 using likelihood ratio tests or any information criterion. Note that the pure FPK model does 298 not make sense in cases where exp(-V(x)) does not converge to 0 when x tends to  $+\infty$  or 299  $-\infty$ : this is the case for V(x) = cx and V(x) = 0, as well as for both  $V(x) = ax^4 + bx^2 + cx$ 300 and  $V(x) = bx^2 + cx$  when the dominant polynomial coefficient is negative. This does not 301 mean that evolutionary trends cannot be estimated from comparative data when the trait 302 interval is not bounded: a macroevolutionary landscape which quickly raises from low to 303 high probabilities, then decreases slowly until a given trait value, and finally quickly drops to low probabilities again would fit a scenario with a trend towards small trait values with 305 soft bounds on trait values: such a scenario is shown on Fig. 2a. 306

Finally, for all versions of the macroevolutionary landscape, confidence intervals con-

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taining the 95% highest probability density around parameter estimates while fixing other parameters to their maximum likelihood estimate can be calculated. This is technically done by removing the lowest 2.5% density regions on each side of the MLE for  $\sigma^2$ , the parameters describing the shape of the macroevolutionary landscape, and the root value when its MLE does not lie in one of the bounds of the trait interval. If the MLE of the root value lies in one of the bounds of the trait interval, then the lowest 5% density region on the other side is removed. These confidence intervals can be returned along with likelihood profile plots around parameter estimates. 

## MCMC algorithm

In addition to maximum-likelihood optimization, we present a MCMC algorithm to estimate parameters of the FPK model, which is also written in R. Since the aim of this MCMC algorithm will often be to get an idea of the distribution of parameter estimates, we have focused on the full model with three polynomial terms. However, nested models with simpler macroevolutionary landscapes can also be fit by setting the probability of update for unnecessary parameters to zero. Numerical calculation of the likelihood of FPK in our MCMC implementation is done as for the maximum-likelihood case and we use the Metropolis-Hastings algorithm to create a Markov chain of parameter estimates.

Parameters of the FPK model have different natures: the three coefficients determining the shape of the macroevolutionary landscape (a, b, and c) as well as the diffusion coefficient  $\log(\sigma^2/2)$  are continuous variables, while the root value of the trait,  $x_0$ , is only allowed to vary on a regular grid of points (see above). These different parameters thus have different kinds of prior and proposal functions.

We have implemented two prior distributions for continuously varying parameters: either normal or uniform ones. However, one should keep in mind that very large values of a and b in particular (i.e., the coefficients of the  $x^4$  and  $x^2$  terms) can lead to extremely

steep macroevolutionary landscapes, which will be unrealistic in most cases. For these two parameters at least, a normal prior centered on zero thus seems to be the most sensible choice. For  $x_0$  we have only implemented a discrete uniform prior on all points of the trait grid.

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As for proposal functions, both normal deviates and sliding windows are available for continuously distributed parameters (but other proposals could easily be implemented by modifying our R code). Note that since we actually update  $\log(\sigma^2/2)$ , this corresponds to a multiplier proposal for  $\sigma^2$ . Only a discrete sliding windows is possible for  $x_0$  and a move on the trait grid is forced to occur each time this parameter is updated. The sensitivity of these proposal functions can be set by the user, but we recommend that the discrete sliding window for  $x_0$  only allows for moves of one step on the grid. Parameters of the model are updated independently, but in order to speed up convergence of the Markov chain to its stationary distribution the relative frequencies of update of the different parameters can be modified.

In practice, we have observed two contrasting behaviors for convergence: (i) when 347  $T_c$  is larger than the depth of the phylogenetic tree,  $\sigma^2$  will converge rapidly in the MCMC 348 chain while parameters setting the shape of the macroevolutionary landscape will not; (ii) 349 on the opposite, when  $T_c$  is smaller than the depth of the phylogenetic tree, parameters 350 setting the shape of the macroevolutionary landscape will converge rapidly while  $\sigma^2$  will 351 be slow to converge. These two behaviors simply reflect the fact that when  $T_c$  is large 352 relative to the total time span of trait evolution, the distribution of traits at the tips of the 353 phylogeny will not have converged to the stationary distribution of the FPK model: the 354 macroevolutionary landscape is poorly explored and tip values retain very little information regarding its shape. On the contrary, when  $T_c$  is small, the macroevolutionary landscape 356 will have been thoroughly explored by the clade, but it is difficult to determine the value 357 of the evolutionary rate with precision (see below). Manipulating the relative frequencies 358

at which these different parameters are updated can have dramatic effects on the speed of 359 convergence of the MCMC chain. A good way to set this tuning parameter would be either 360 to first run a maximum-likelihood estimation of the model or to do an initial quick MCMC 361 run (for example with a very low number of points to discretize the trait interval) in order 362 to get an idea of the values of  $\sigma^2$  and V. From our experience, convergence of our MCMC 363 algorithm takes a long time, even on rather small datasets (c. 200,000 to 1 million steps). 364 We recommend running at least two independent chains in order to make sure that they 365 have converged to the same stationary distribution. 366

In order to assess the performance of FPK in terms of parameter inference and model discrimination, we have conducted a large number of simulations. Our focus was on the like-lihood of FPK, thus we restricted our simulations to the maximum-likelihood optimization procedure since it is much faster than MCMC estimation. We ran simulations under height contrasted scenarios, four of them corresponding to the pure FPK model and four to the BBMV model. The macroevolutionary landscapes corresponding to these height scenarios are pictured on Fig. 2.

The four scenarios of the FPK model that we simulated were the following: (a) a 375 directional trend limited to a portion of trait space  $(V(x) = 5x^4 - x^2 + x)$ , (b) a macroevo-376 lutionary landscape with two peaks of equal height  $(V(x) = 10x^4 - 5x^2)$ , (c) a macroevo-377 lutionary landscape with two peaks of different heights  $(V(x) = 5x^4 - 5x^2 + x)$ , and (d) 378 a single peak (i.e., an OU model,  $V(x) = 5x^2$ ). The four scenarios simulated under the 379 BBMV model, i.e. in which trait evolution was actually bounded, cover a broad range of 380 interesting cases: (e) a flat macroevolutionary landscape (BBM, V(x) = 0), (f) a directional 381 trend (V(x) = 1.5x), (g) disruptive selection (i.e., a U-shaped macroevolutionary landscape 382 with extreme trait values being favored,  $V(x) = -1.2x^2$ , and (h) two peaks of the same 383 height  $(V(x) = 3x^4 - 6x^2)$ . 384

For each one of these height scenarios, we fit four different versions of FPK: the full model  $(V(x) = ax^4 + bx^2 + cx)$ , a model with only quadratic and linear terms  $(V(x) = bx^2 + cx)$ , a model with only a linear term (V(x) = cx), and a model with a flat macroevolutionary landscape (i.e., BBM, V(x) = 0). In simulations of the FPK model (scenarios a-d), bounds were placed far apart from the observed trait interval (see above) for inference, while in simulations of the BBMV model (scenarios e-h) the true bounds used in simulations were specified. In addition, we also fit BM and an OU model with a single optimum to each

simulated dataset using the fitContinuous function in the geiger package (Pennell et al. (2014)).

Phylogenetic trees were simulated under a pure birth model, with unit birth rate. 394 Trees were grown until the desired number of tips plus one was obtained and one of the two 395 sister tips originating from the last speciation event was trimmed. Trees were then rescaled to 396 a total depth of 100 arbitrary time units in order to enable comparison between simulations. 397 All simulations were done with  $B_{\min} = 0$ ,  $B_{\max} = 1$ , and  $x_0 = 0.5$ . However, for each of 398 the four scenarios described above we used two different values of  $\sigma^2$  which were calculated 399 so that: (i)  $T_c = 5$  (i.e., 1/20 of tree depth), which should ensure that the distribution of 400 the trait at the tips of the tree has converged to the stationary distribution of the model, 401 and (ii)  $T_c = 2,000$  (i.e., 20 times tree depth), in which stationarity should not have been 402 reached at the end of the simulation. In addition, we also explored the effect of tree size 403 on parameter estimation and model discrimination using trees of 50, 100, and 200 tips. For 404 each combination of the shape of the potential, the value of  $\sigma^2$ , and tree size, we conducted 405 20 different simulations (960 simulations in total). 406

#### Model discrimination

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We first focused on whether FPK and BBMV can be distinguished from other classic 408 models of evolution using relative Akaike weights (Burnham and Anderson (2002)). Our 409 simulations showed that when stationarity was reached  $(T_c = 5)$ , all four scenarios of the 410 FPK model that we simulated could easily be discriminated from BM, which always received 411 less than 0.001% Akaike weight (Fig. 3). Discrimination from OU was also easily achieved, 412 this model always receiving less than 13% Akaike weight, except in the case where it was 413 the model which was actually simulated (scenario d, Fig. 3). Discrimination was even better 414 under the four scenarios of the BBMV model that we simulated, with both BM and OU 415 always receiving less than 0.01% Akaike weight (Fig. 3). In simulations where stationarity 416

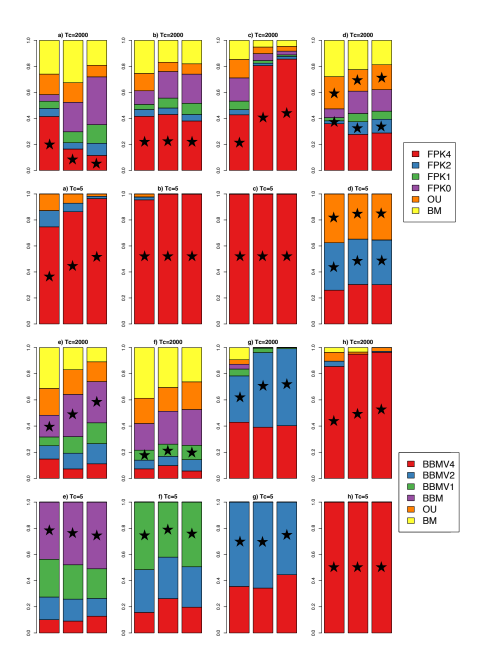


Figure 3: Model discrimination in simulations. Each panel shows AIC weights of each of the six models fitted to the simulated data: 4 versions of the FPK or BBMV models, plus BM and OU, averaged over 20 simulations. In each case, black stars indicate the model that was used for simulations and blocks of three columns show results for trees of 50, 100 and 200 tips, from left to right. The top height panels show simulations of the pure FPK model, the height bottom ones simulations of the BBMV model. Numbers after FPK or BBMV in the legends indicate the degree of the leading polynomial term that was used for fitting: for example, FPK4 stands for the FPK model with  $V(x) = ax^4 + bx^2 + cx$ , and BBMV1 for the BBMV model with V(x) = cx.

was not reached ( $T_c = 2,000$ ), only scenarios c, g, and h) could still be discriminated from BM and OU (Fig. 3). All other five scenarios indeed gave relatively high Akaike weights to either BM or OU (Fig. 3). The number of tips in the tree did influence discriminatory power between FPK and BM or OU positively, but its effect was moderate (Fig. 3).

We then looked at whether FPK or BBMV models with different shapes of the 421 macroevolutionary landscape can be statistically distinguished. Discrimination between al-422 ternative versions of the model with different shapes of the macroevolutionary landscape 423 was also generally satisfactory. In cases where stationarity was reached  $(T_c = 5)$ , the version 424 of the model that was used to simulate the data was always the one to receive the high-425 est AIC weight (Fig. 3). Increasing tree size generally lead to an increase in the Akaike 426 weight of the generating model, the effect being substantial this time (Fig. 3). Scenar-427 ios with macroevolutionary landscapes containing two peaks (i.e., scenarios b, c, and h), 428 which can only be accommodated by the most complex form of the macroevolutionary land-429 scape  $(V(x) = ax^4 + bx^2 + cx)$  always led to more than 95% Akaike weight to this model 430 (Fig. 3). Importantly, for all four scenarios simulated under the FPK model, models with a 431 flat (V(x) = 0) and a linear potential (V(x) = cx) always received less than 2.3e - 8 Akaike 432 weight. We have seen above that this two shapes of the potential do not make sense in 433 the case of the FPK model (i.e., when there are no bounds in practice): these simulations 434 confirm that both of these models are strongly rejected statistically in these situations. In 435 contrast, when stationarity was not reached  $(T_c = 2,000)$  different shapes of the macroevolu-436 tionary landscape were difficult to discriminate and simpler forms were often preferred over 437 more complex ones, especially so in trees with few tips. This is normal since in these cases, traits only had time to explore a small fraction of the macroevolutionary landscape. The only notable exception to this general observation was for scenario h (a BBMV model with 440 two peaks), in which the full BBMV model always received over 85% Akaike weight, even 441 when  $T_c = 2,000$  and with trees of 50 tips only (Table 1). 442

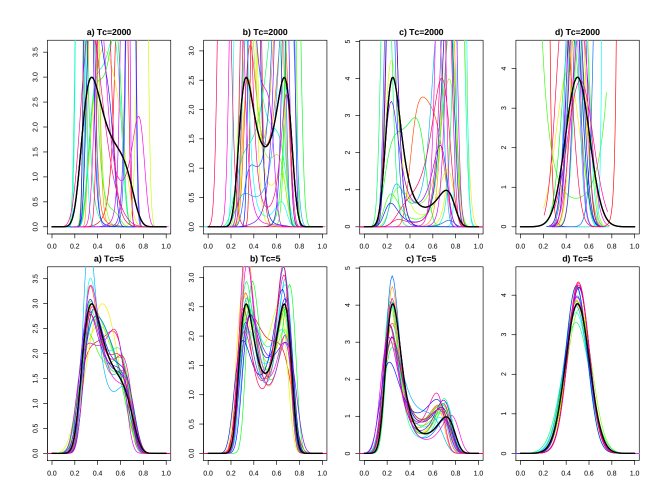


Figure 4: Estimation of the macroevolutionary landscape in different versions of the FPK model. Thin lines in each plot show the macroevolutionary landscapes estimated in 20 simulations, each one in a different color, while the simulated macroevolutionary landscape is shown by the thick black line. Only results for trees with 100 tips are shown. Top row: simulations with  $T_c = 2,000$ , in which stationarity was not reached. Bottom row: simulations with  $T_c = 5$ , in which stationarity was reached. From left to right, columns show simulation for scenarios a to d.

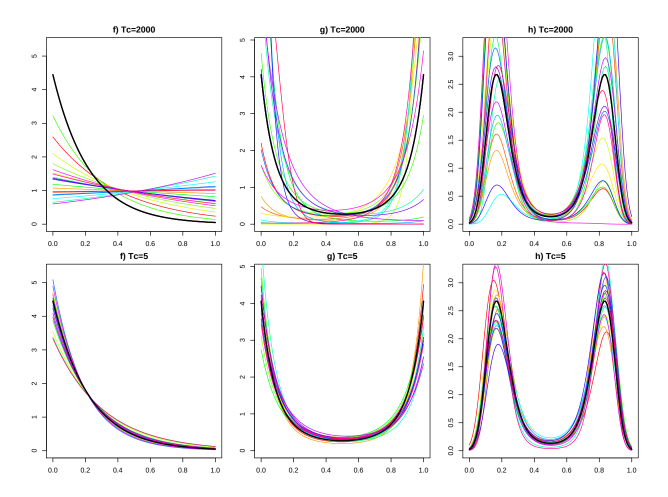


Figure 5: Estimation of the macroevolutionary landscape in different versions of the BBMV model. Thin lines in each plot show the macroevolutionary landscapes estimated in 20 simulations, each one in a different color, while the simulated macroevolutionary landscape is shown by the thick black line. Only results for trees with 100 tips are shown. Top row: simulations with  $T_c = 2,000$ , in which stationarity was not reached. Bottom row: simulations with  $T_c = 5$ , in which stationarity was reached. From left to right, columns show simulation for scenarios f to g. Results for simulations of the flat landscape (BBM, scenario e) are not shown since the macroevolutionary landscape is fixed in this case.

Accuracy of FPK models in parameter estimation was assessed by comparing the 444 maximum-likelihood estimates of parameters with values used in simulations. We first com-445 pared the precision in the estimation of the macroevolutionary landscape as a whole, and 446 not in the estimation of a, b, and c separately. This is because these three coefficients can 447 sometimes be redundant and lead to very similar shapes of the macroevolutionary landscape: 448 for example, a (the coefficient of the  $x^4$  term) and b (the coefficient of the  $x^2$  term) are highly 449 correlated. Simulations showed that macroevolutionary landscapes are generally accurately 450 estimated in cases where stationarity has been reached since the actual shape that was sim-451 ulated is most often recovered (Fig. 4 & 5). Estimation of the macroevolutionary landscape 452 was even better in simulations of the BBMV model (Fig. 5) compared to simulations the 453 pure FPK model (Fig. 4), probably because in the former case the actual bounds used in 454 simulations were specified when inferring parameters. In all height scenarios, accuracy in 455 the estimation of the macroevolutionary landscape increased with the number of tips in the phylogeny (Online Appendix III). Accuracy was much worse in simulations that had not 457 reached stationarity (Fig. 4 & 5). 458

As for the other two parameters of the FPK model, accuracy in the estimation of  $x_0$  was much less satisfactory and usually had a huge variance, especially so when  $T_c$  was small (Online Appendix Figures 1 & 2). The estimation of  $\sigma^2$  was very accurate for large values of  $T_c$  but had much larger variance when  $T_c$  was small (Online Appendix Figures 3 & 4). No bias in the estimation of  $\sigma^2$  was apparent for the FPK model, but it seemed that the estimation of  $\sigma^2$  was slightly biased towards larger values in the four scenarios of the BBMV model that we simulated and for  $T_c = 5$ .

Empirical example: body size evolution in North-American watersnakes (tribe

Thamnophiini)

We demonstrate the utility of FPK using an example of body size evolution in snakes.

We decided to study North-American watersnakes (Colubridea, subfamily Natricinae, tribe

Thamnophiini) because the distribution of their body length shows a slight bimodality (Burbrink and Myers (2014)). A time-calibrated phylogeny as well as measurements of total
length (hereafter, TL) for 45 species included in this group were obtained from Burbrink

and Myers (2014), and TL was log<sub>10</sub>-transformed prior to analysis.

We first fit three alternative models for the evolution of TL along the watersnake phylogeny using maximum-likelihood: BM, an OU model with a single optimum, and the FPK model (with  $V(x) = ax^4 + bx^2 + cx$ ). In addition, we used our MCMC algorithm to obtain posterior estimates of the shape of the macroevolutionary landscape in this clade (detailed methods can be found in Online Appendix IV). Convergence of MCMC chains was assessed both visually by looking at the trace plots of the parameters, likelihood, prior, and posterior, and by measuring the effective sample sizes of these different quantities using the R package coda (Plummer et al. (2006)).

Among the three models compared using maximum-likelihood, the FPK model had by 482 far the lowest AIC, followed by the OU model ( $\triangle$ AIC=13.2), and finally BM ( $\triangle$ AIC=15.1). 483 The macroevolutionary landscape estimated by the FPK model contained two distinct peaks, 484 the peak corresponding to longer TLs being the highest (Fig. 6). Confidence intervals on 485 the maximum-likelihood estimates of model parameters confirmed that the coefficient for 486 the  $x^4$  term of the potential, a, was positive (ML estimate: 9.2, 95% CI: [5.8;14.3]), while 487 the coefficient for the  $x^2$  term, b, was negative (ML estimate: -3.4, 95% CI: [-4.9;-1.0]), 488 which is typical of macroevolutionary landscapes with two peaks. The linear coefficient 489 of the potential, c, was not significantly different from 0 (ML estimate: -0.34, 95% CI: [-1.9;1.8]). There was large uncertainty as to the value of TL for the ancestor of watersnakes, 491 the confidence interval spanning almost the whole distribution of TL in extent species (ML 492 estimate: 72.9cm, 95% CI: [39.9;118.1]). We estimated a characteristic time of 22.4 Myrs 493

for the FPK process, which is slightly higher than the crown age of Thamnophiini (16.6 Myrs, Burbrink and Myers (2014)). Results obtained using maximum-likelihood estimation were supported by the two MCMC chains that we ran: the posterior distribution of the macroevolutionary landscape also had two peaks of unequal heights, and the mode of this distribution closely matched the macroevolutionary landscape estimated using maximum-likelihood (Fig. 6).

These results suggest that TL might have evolved toward two different optima in
North-American watersnakes, the first one roughly corresponding to 50cm and the second,
higher optimum, to 130cm (Fig. 6). In their original publication Burbrink and Myers (2014)
had proposed that the skewness in the distribution of TL in watersnakes could be due to
higher diversification rates for longer species. Comparing both explanations would prove
especially interesting, but would require extending the FPK model to account for diversification rates depending on the value of the evolutionary potential.

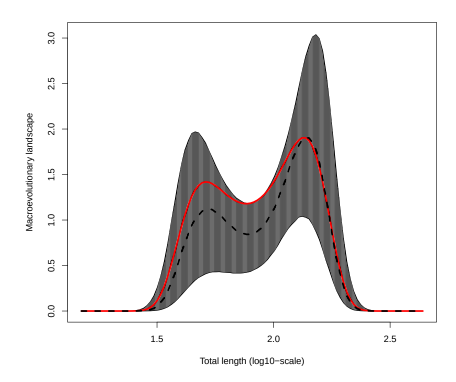


Figure 6: Posterior distribution of the macroevolutionary landscape estimated for body length evolution in watersnakes (tribe Thamnophiini). This posterior distribution was obtained by concatenating the two MCMC chains after the first 10% of samples were discarded as burnin (800,000 MCMC steps in total). The figure shows the value of the macroevolutionary landscape (N.exp(-V(x))) on the y-axis as a function of  $log_{10}(total\ length)$  measured in centimeters. The dashed black line shows the median value of the macroevolutionary landscape over the posterior, while the grey area ranges from the 25% to the 75% quantiles. The solid red line shows the maximum-likelihood estimate of the macroevolutionary landscape.

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In this article we have presented equations for a very general model of evolution for continuous traits, as well as its implementation. This opens new possibilities for estimating macroevolutionary landscapes from phylogenetic comparative data. Below we discuss the strengths and weaknesses of this model. We note that Blomberg (2016) has recently introduced a family of essentially similar models for continuous trait evolution, but no framework exists yet to infer parameters of these models from phylogenetic comparative data.

New avenues for studying phenotypic evolution from comparative data

The flexibility of the new model that we propose is its greatest strength. FPK and its bounded version BBMV can indeed be used to infer macroevolutionary landscapes of any conceivable shape, and thus recover processes such as evolution between bounds, evolutionary trends, disruptive selection, or diversifying selection leading to macroevolutionary landscapes with several peaks. These last three scenarios lie at the core of modern (macro) evolutionary theory, but could not yet be inferred from phylogenetic comparative data (O'Meara (2012)). FPK is a model that generally retains very little phylogenetic signal: over all simulations that we ran in which stationarity had been reached, the median value of the  $\lambda$  index of phylogenetic signal (Pagel (1997)) was 1.6e-109 (see Online Appendix V). This absence of phylogenetic signal stems from the strong deterministic component of the FPK model in all scenarios that we simulated, a result already known for the OU model (Münkemüller et al. (2015)), which is a special case of FPK. However, some scenarios in which the deterministic component of FPK is small or even absent can yield high level of phylogenetic signal: this is the case of the BM model, also a special case of FPK. As a result, most of the information needed to infer the shape of the macroevolutionary landscape ultimately comes from the distribution of the trait for extant species. Indeed, the trait distribution is expected to converge to the stationary distribution of FPK, set aside the fact that recently diverged species will
have similar trait values. For example, evolutionary trends can be recovered in the absence
of fossil data from a highly skewed trait distribution for contemporaneous species. In the
same vein, the simultaneous presence of two peaks in the macroevolutionary landscape can
be inferred from a bimodal trait distribution.

The characteristic time of the FPK process,  $T_c$ , gives the typical time needed to 536 reach stationarity and should always be compared to the total depth of the phylogenetic 537 tree for the clade under study,  $T_{tot}$ . Unsurprisingly, model performance will increase with 538  $T_{tot}/T_c$  and in the extreme case where  $T_{tot} \ll T_c$ , traits will not have explored much of 539 the macroevolutionary landscape.  $T_c$  bears much similarity with the phylogenetic half-life of the OU process, which describes the time necessary for the trait value to move halfway from its initial position to the optimum. Indeed, the characteristic time of an OU process is directly proportional to its phylogenetic half-life (Online Appendix I). In agreement with our findings, previous studies have shown that accuracy in parameter estimation of the OU 544 model increases with decreasing phylogenetic half-lives (Uyeda and Harmon (2014); Ho and 545 Ané (2014a)). 546

Using simulations, we have demonstrated that FPK can be distinguished from other 547 classic models of trait evolution, and also that distinct shapes of the macroevolutionary 548 landscape can be distinguished from each other based on their likelihoods. 549 that alternative versions of the FPK model can be used for testing evolutionary hypotheses 550 about trait evolution. Our simulations also show that the danger of over-fitting is quite 551 low with FPK since simpler models will often be preferred when stationarity has not been 552 reached. Our focus on AIC to discriminate between alternative models was motivated by the fact that it is the most commonly used in the macroevolutionary community. However, 554 AIC might be prone to overfitting in parameter rich macroevolutionary models and other 555 measures that penalize more for extra parameters, like the Bayesian Information Criterion 556

or its modified version (Zhang and Siegmund (2007)), might be preferable (Ho and Ané (2014a)). Another solution to diagnose overfitting would be to use parametric bootstrapping techniques (Boettiger et al. (2012)), which is readily implementable for FPK since we provide an R function to simulate the model.

Our results also show that estimation accuracy under FPK drastically differs between 561 parameters (Online Appendix III). Estimation is generally accurate when estimating the 562 shape of the macroevolutionary landscape and increases with  $T_{tot}/T_c$ . Estimation accuracy 563 also increases with tree size: our results suggest that trees with 50 tips lead to reasonable 564 estimation of the general shape of the macroevolutionary landscape (Online Appendix III), 565 while trees with 100 tips should most often be large enough to obtain very reliable estimates 566 (Fig. 4 & 5). Importantly the three coefficients that determine the shape of the macroevolu-567 tionary landscape should not be analyzed separately since numerous different combinations 568 of a, b, and c can give similar shapes. Rather, we recommend to interpret the general shape 569 of the macroevolutionary landscape by focusing on a few important features: (i) whether the 570 macroevolutionary landscape is flat or not, (ii) whether it features a single trend towards 571 one of the bounds of the trait interval, (iii) whether it contains one or several peaks, and if 572 relevant (iv) where are these peaks located in the trait interval. In contrast, the evolutionary 573 rate  $\sigma^2$  has high estimation variance, and especially so when  $T_{tot}/T_c$  is large. This same ef-574 fect had already been observed for BBM (Boucher and Démery (2016)) and probably reflects 575 the fact that when the macroevolutionary landscape has been fully explored by the clade 576 its shape and extent are easily estimated but the speed at which the landscape is travelled 577 is not. Given this limitation we don't see much value in interpreting the estimate of  $\sigma^2$ 578 when fitting FPK to an empirical dataset. Rather, the characteristic time of the process,  $T_c$ , should be the quantity that is interpreted in comparison with tree depth. Finally, we 580 found that the estimation of the trait value at the root of the tree,  $x_0$ , is poor as soon as 581 the macroevolutionary landscape has been moderately explored, a result that generalizes the 582

one already obtained for BBM (Boucher and Démery (2016)). This stems from the fact that FPK is a model that retains low phylogenetic signal since it includes strong deterministic forces and wipes out any hope of confidently inferring ancestral trait values in empirical datasets.

These differences in the quality of the estimation of the shape of the macroevolutionary landscape vs.  $\sigma^2$  generalize results that have been obtained for the OU model. Indeed, in this model  $\sigma^2$  and especially  $\alpha$ , the attraction strength, are difficult to estimate (Butler and King (2004); Ho and Ané (2014a)) while it seems that the stationary variance of the OU process,  $\sigma^2/2\alpha$ , and the trait optimum,  $\mu$ , generally have higher estimation accuracy (Münkemüller et al. (2015); Ho and Ané (2014a)). In light of our results, this is normal since  $\mu$  and  $\sigma^2/2\alpha$  respectively determine the mean and variance of the stationary distribution of the OU process, what we have called the macroevolutionary landscape in this article.

# Interpretation of macroevolutionary landscapes inferred from FPK

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We have introduced FPK as a method for the inference of macroevolutionary land-596 scapes from phylogenetic comparative data. The adaptive landscape is a fruitful metaphor to understand phenotypic evolution on a variety of evolutionary scales (Wright (1932); Simp-598 son (1944); Arnold et al. (2001)) but care must be taken when interpreting inferences made 599 from phylogenetic comparative data in the light of adaptive landscape theory, which was 600 mainly developed for population genetics (Uyeda and Harmon (2014)). Some of the models 601 of trait evolution on phylogenies include deterministic forces that influence trait evolution in 602 an attempt to mimic selection: the OU model includes a term that was designed to resemble 603 stabilizing selection towards a given trait value (Hansen (1997)) and FPK can imitate a 604 large variety of selection shapes. However, all macroevolutionary models for trait evolution, 605 including OU and FPK, are phenomenological by nature since they rely on probabilistic dif-606 fusion equations and since they model evolution over long time-scales (typically thousands 607

to million years) that are not amenable to observation. In other words, these models recover patterns from the data, which researchers have to interpret in terms of evolutionary processes. One example of such a confusion between long-term patterns and short-term processes has been highlighted when interpreting the good fit of an OU model to empirical datasets: sev-eral studies have indeed shown that neutral evolution between bounds produces patterns closely resembling the ones obtained under an OU process (Revell et al. (2008); Boucher et al. (2014)). Development of the BBM model has now rendered possible to distinguish be-tween these two scenarios (Boucher and Démery (2016)), but many such cases in which two different microevolutionary processes produce the same macroevolutionary pattern remain. 

Macroevolutionary landscapes estimated using FPK should thus be interpreted with extreme caution: they reflect the general shape of deterministic forces that have been acting on the evolution of a continuous trait in a clade, but are agnostic regarding the nature of these deterministic forces, *i.e.*, they are not actual measurements of the relation between individuals' traits and fitness. The most obvious limitation of FPK is that it makes the hypothesis that the macroevolutionary landscape is constant through time. This is likely to be wrong in a majority of cases since environmental change or interactions with other species will lead to changes in the intensity and shape of the selection gradient acting on one trait in a given clade (Simpson (1944); Hansen (2012)). Macroevolutionary landscapes inferred using FPK will thus necessarily reflect some kind of average macroevolutionary landscape experienced by the clade throughout its evolutionary history.

Notwithstanding these limitations, which are inherent to all macroevolutionary models, FPK offers the opportunity to estimate a variety of macroevolutionary landscapes from phylogenetic comparative data. One kind of possible landscapes deserves particular mention here: macroevolutionary landscapes in which multiple peaks exist. This scenario would be especially interesting to compare to a situation in which these multiple peaks are available for different lineages, which is what is implemented in OU models with multiple optima. In this

latter class of models, each lineage is indeed subject to attraction towards a single peak at a time, but lineages may shift between different peaks. These shifts are either determined a 635 priori (Butler and King (2004); Beaulieu et al. (2012)) or inferred directly from phylogenetic 636 patterns of trait evolution (Ingram and Mahler (2013); Khabbazian et al. (2016); Uyeda and 637 Harmon (2014)). In FPK with multiple peaks on the contrary, each lineage is always in-638 fluenced by the different peaks in its macroevolutionary landscape, and transitions between 630 peaks might be frequent if the traits of most species in the clade are located in a valley of 640 the macroevolutionary landscape. More theoretically, these two alternatives would repre-641 sent two different scenarios: OU models with several optima might be better at describing 642 situations in which a lineage shifts to another adaptive zone (Simpson (1944)), while FPK 643 with multiple peaks might represent more genuine diversifying selection towards alternative phenotypic optima. Whether these alternatives can be discriminated using empirical data 645 would be of uttermost interest but remains a totally open question.

Even though we have emphasized that it is difficult to connect microevolutionary 647 processes to macroevolutionary patterns, there is one promising way in which FPK could 648 be used to do so. Indeed, the Bayesian implementation of the model enables the use of 649 informative priors based on quantitative genetic parameters. Uyeda and Harmon (2014) 650 have demonstrated how this could be done for the OU model: using the quantitative genetic 651 model of Lande (1976), they showed how measurements of heritability, phenotypic variance, 652 and effective population size can inform priors on the parameters of the OU model. By 653 connecting the parameters in Eq. 1 to quantitative genetic models, the same procedure 654 could be carried out for FPK. 655

Finally, FPK need not be restricted to infer macroevolutionary landscapes. This model indeed has its roots in spatial diffusion theory and as such could be used in phylogeographic studies to model the dispersal of a set of individuals or populations for which the phylogeny is known (e.g. Grollemund et al. (2015)). This field of research has indeed

seen huge methodological advances in recent years (Lemey et al. (2010); Bloomquist et al. (2012)). In this context, FPK could be used to infer preferred directions of dispersal (*i.e.*, directional trends) or even to infer particular regions that act as geographic attractors for the taxon under study (*i.e.*, one or several peaks), while explicitly taking into account hard boundaries on the distribution of organisms (e.g. oceans for terrestrial organisms). Hard bounds on the distribution of organisms (e.g., coastlines for terrestrial organisms) could even be taken into account explicitly using the BBMV model.

#### Limitations of the FPK model

Our implementation of FPK does not come without limitations. The main technical limitation is that our implementation of the model is restricted to single traits. We are fully aware that extending it to multivariate datasets would be very convenient, since multiple traits are expected to often evolve in a correlated fashion (Arnold (1992)). However, this is for the moment hampered by computational time. Indeed, the most time-consuming part in the calculation of the likelihood is to invert the instantaneous transition matrix, M. If we were to study two traits simultaneously, we would need to discretize the plane that they define into a regular grid of points, and computing time would not be multiplied by two but rather raised to the power two. The only possible solution that we can envision would be to use algorithmic tricks that avoid inverting the entire transition matrix, but rather a matrix describing transitions between a given point on the grid and its immediate neighbors, as recently proposed for inference of ancestral areas (Landis et al. (2013)). This would require much development and is out of the scope of this article.

The fact that our implementation of FPK can only accommodate a maximum of two peaks in the macroevolutionary landscape might also seem frustrating for some users. Extending the model so that three peaks or more can be simultaneously present is rather straightforward: the most obvious solution would be to use polynomial functions with more

terms for V(x). However, this would increase computational time, and more importantly would probably yield likelihood functions that are extremely difficult to optimize. This is why we have not implemented it yet. We however note that in our code to infer the FPK model, we have left open the possibility to specify a given shape for V(x): users can thus experiment with more complex macroevolutionary landscapes if they feel this is relevant to their specific study system.

As already discussed above, the fact that the macroevolutionary landscape is constant across time and across different clades in the phylogeny is another limitation. Further developments of the FPK model could aim at extending it to cases in which the macroevolutionary landscape differs among clades or among specified time periods across the whole phylogeny.

The last limitation of FPK perhaps lies in its very formulation. FPK is indeed based on a constant-rate diffusion model and as such cannot model accelerating or decelerating trait evolution (Harmon et al. (2010)) or sudden jumps in the value of the trait, as would be expected under quantum evolution (Simpson (1944); Kirkpatrick (1982)) or punctuated equilibrium (Gould and Eldredge (1977)).

701 Conclusion

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Our development and implementation of FPK greatly expands the set of models 702 available for studying the evolution of continuous characters on phylogenies and enables 703 the estimation of macroevolutionary landscapes of various shapes. We have shown that the 704 model generally achieves good performance both in terms of parameter estimation and in 705 terms of discrimination from alternative macroevolutionary models. R code for fitting FPK 706 (and its special case BBMV) to empirical data is freely available from https://github. 707 com/fcboucher/BBMV, and this repository also contains a detailed tutorial to the different 708 functions for simulating and inferring the model. 709

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Online appendices can be found in the Dryad repository associated with this submission.

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