Evolution of patterns on Conus shells

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AUTHOR SUMMARY

Pigmentation patterns on gastropod (i.e., snail) shells vary widely among species, but the complexity of the patterns make it difficult to quantify these differences or understand their evolution. To answer these questions, we use a developmental model that reproduces the pigmentation patterns on 19 species of cone snails. Our model shows that evolutionary changes are generally slow, with a few episodes of rapid change, possibly indicating the action of natural selection. Our analysis allows the inference of ancestral shell patterns and represents an attempt to understand a complex developmental history by using phylogenetic methods on a developmental model.

Our neural-network model is based on a neurosecretory feedback loop; it has only a few parameters, and they correspond to physiologically measurable features of the gastropod nervous system. In our model, the pigmentation patterns are formed by secretory cells in the mantle (i.e., the part of the gastropod that creates and colors the shell). The modeled mantle output is controlled by a hypothesized network of excitatory and inhibitory neurons. The pigmentation pattern represents the history of the output of this network with "time" running along the axis of shell growth and "space" running

parallel to the aperture of the shell. We suppose that other neurons in the mantle are able to sense pigment that has been previously laid down. The output of these sensory neurons is transformed by the excitatory–inhibitory network that produces the next round of pigmentation. To determine the secretory output at a specific location for the current bout of pigmentation, the excitatory–inhibitory network performs a weighted average over the entire surface of the shell and then applies a saturating nonlinearity to this average. The weights of the summed activity are interpreted as excitatory for nearby spatial locations and recent bouts of activity or as inhibitory for spatially distant and earlier bouts of activity—a so-called center-surround dynamic. With these few parameters, our model can reproduce the diverse range of coloration seen on gastropod shells.

To understand the evolutionary history of the model parameters, we estimated parameter values for each of 19 species in the genus *Conus*, a group of marine snails whose shells display a wide range of pigmentation patterns. A well resolved phylogeny,



Fig. P1. Shells of living species are displayed at the tips of the tree. To the right of these are shells "grown" in the computer by using the neuralnetwork model. Based on a Brownian-motion model for the evolution of continuous traits, we estimated the model parameters for ancestors. Then, the neural network model was used to produce the ancestor shells by using the estimated parameters.

or map of evolutionary relationships, of these species had already been obtained from the analysis of DNA sequences of four genes from each species. By mapping our model parameter values for each species onto this phylogeny, we inferred the evolutionary trajectories of each parameter (Fig. P1). Specifically, we found that these parameters evolved roughly independently of one another in the ancestors of the extant species. Further, we show that the model parameters evolved slowly on most lineages, but with a few episodes of rapid change. These probably indicate the effect of natural selection on some parameters in some lineages. There is a strong "phylogenetic signal" for the parameters, which means that there is overall concordance between the model parameters and the phylogeny. This result contrasts with what we found for several observable features of the pigmentation patterns, including the presence of triangles or stripes, for which there is no phylogenetic signal. The overall fit of the model parameters to the phylogeny allowed us to infer the pigmentation patterns of ancestral cone snails, something that cannot be determined from the fossil record of most groups. Some of these inferred patterns lie outside the range of

phenotypic variation of the 19 species we analyzed, but are found in other existing gastropods.

Our analysis represents a unique attempt to understand the history of a complex developmental process by applying phylogenetic methods to an explicit developmental model.

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Data deposition: The computational parameters for the pattern formation model for each of the described species are available upon request.

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The pigmentation patterns of shells in the genus *Conus* can be generated by a neural-network model of the mantle. We fit model parameters to the shell pigmentation patterns of 19 living *Conus* species for which a well resolved phylogeny is available. We infer the evolutionary history of these parameters and use these results to infer the pigmentation patterns of ancestral species. The methods we use allow us to characterize the evolutionary history of a neural network, an organ that cannot be preserved in the fossil record. These results are also notable because the inferred patterns of ancestral species sometimes lie outside the range of patterns of their living descendants, and illustrate how development imposes constraints on the evolution of complex phenotypes.

pattern formation | developmental evolution | phylogenetics | ancestral inference

Pigmentation patterns on mollusk shells are typical complex phenotypes. They differ substantially among closely related species, but the complexity of the patterns makes it difficult to characterize their similarities and differences. Consequently, it has proven difficult to describe the evolution of pigmentation patterns or to draw inferences about how natural selection might affect them. In this report, we present an attempt to resolve this problem by combining phylogenetic methods with a realistic developmental model that can generate pigmentation patterns of shelled mollusks in the diverse cone snail genus *Conus*. The model is based on the interactions between pigment-secreting cells and a neuronal network whose parameters are measurable physiological quantities. The neural model used here is a generalization of models proposed earlier by Ermentrout et al. (1) and Boettiger et al. (2). Furthermore, the species have a well supported phylogeny that allows us to infer rates and patterns of parameter evolution.

We chose 19 species in the genus *Conus* for which Nam et al. have presented a resolved phylogeny (3). For each species, we found a model parameter set that matched the observed pigmentation pattern. Then we applied likelihood-based phylogenetic methods to measure phylogenetic signal in the model parameters, compare possible evolutionary models, estimate the model parameters of ancestral species, and then use these to infer the pigmentation patterns of ancestral species.

Neural Model

Fig. 1 shows a schematic of the mantle geometry and illustrates the basic principle of the neural model. The mathematical details are described in *SI Appendix*, Supplement A. The model is built on two general properties of neural networks: spatial lateral inhibition (also called center-surround), and "delayed temporal inhibition." The latter can be viewed as "lateral inhibition in time" (4–6), as illustrated in Fig. 1*C*, *Center*.

The neural field equations describe the local pattern of neuron spiking. Local activity of excitatory neurons induces the activity of inhibitory interneurons in the surrounding tissue. The net spatial activity has "Mexican hat" shape, as shown in Fig. 1C (5–7). As shell material and pigment are laid down in periodic bouts of secretion, the surface pigment pattern is a space–time record of the animal's secretory activity, in which distance from the shell aperture is proportional to the number of bouts of secretion. Excitation

of a cell during a bout inhibits its excitation for some future number of bouts, so that an active neuron will eventually be inhibited and remain inactive for a "refractory" period. Thus, "delayed inhibition" is equivalent to "half a Mexican hat backward in time." Finally, the secretory activity of pigment granule secretory cells depends sigmoidally on the difference between the activities of the excitatory and inhibitory cells, as shown in Fig. 1*C*. The logic of the model is that the sensory cells read the history of pigmentation and send this to the neural net that uses this history to "predict" the next increment of pigmentation and instruct the secretory cells to deposit accordingly. This feedback from output to input distinguishes the neural model from models whose future state depends only on their current state (e.g., diffusible morphogens and cellular automata).

The neural field model is characterized by 17 free parameters, each of which has a concrete physiological interpretation, as described in Fig. 2. In effect, there are four cell types: sensory cells, excitatory neurons, inhibitory neurons, and secretory cells; their effective connectivity relationship is shown in *SI Appendix*, Supplement A. The behavior of each cell type is given by its input/ output relationships, as shown in Fig. 2. Each excitatory and inhibitory neuron is described by a Gaussian spatial synaptic weight kernel described by two parameters (amplitude and width), and a temporal kernel described by four parameters. As several of the parameters appear in products with other parameters, we can normalize their magnitudes and thereby reduce them to three free parameters each describing the spatial and temporal ranges of excitation and inhibition. The precise parameter reduction procedure is described in *SI Appendix*, Supplement A.

Imbued with these properties, the neural network drives secretory cells to lay down both the shell material and pigment. Thus, the model can reproduce both the shell shape and the surface pattern for many shelled mollusk species, as described previously (2). The present model differs in several essential ways from that proposed previously (2); this is also discussed in *SI Appendix*, Supplement A.

The basic neural model consists of a simple feedback circuit that spatially and temporally filters previous activity and feeds the result through a nonlinear function to produce the next bout of pigment. One needs 17 parameters to specify the shape of the functions in Fig. 2. By varying the 17 parameters in the model, we were able to produce a wide variety of cone shell patterns. Some of these patterns are very sensitive to the initial conditions (i.e., "chaotic dynamics"), and thus small changes in the initial pattern

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Fig. 1. The neural-network model of the mantle. (*A*) Rough anatomy of a generic shelled mollusk. Note the "brain," where the neural patterns are processed consists of a ring of ganglia. (*B*) Cross-section of the mantle showing how the sensory cells "taste" the previously laid pigment patterns that are processed by the central ganglion and sent to the mantle network that controls the pigment-secreting cells. (*C*) Simple pattern on a *Conus* shell and how the model extrapolates the previous pattern to produce the current day's pigment secretion. The pigmentation pattern is read by the sensory cells in the mantle. This activity is then passed through the space-time filter of neural activation and inhibition. Here, time represents the pigmentation pattern that was laid down in previous bouts, whereas space is the dimension along the growing edge of the cell. The resulting filtered activity is passed through nonlinearities for excitation and inhibition, and this net activity drives the secretory cells that lay down the new pigmented shell material. The spatial filter, shown in top and perspective views, has the form of a Mexican hat, in which excitatory activity stimulates a surrounding inhibitory field. The temporal filter that implements delayed inhibition is half a Mexican hat. It generates a refractory period following a period of activity. The pigment secreting cells have a sigmoidal stimulus response curve. Feedback occurs as the current pigment deposition becomes part of the input to the sensory cells for the next secretion bout.

or small amounts of noise give rise to diversity among individuals while still maintaining the same qualitative pattern. Fig. 3A provides an example showing multiple instances of a simulation of *Conus crocatus* such that there are small differences in initial data or the addition of a small amount of noise. The overall look of the pattern is the same, but there are clear individual differences.

Somewhat surprisingly, the regions of parameter space that correspond to cone shell patterns are fairly restricted and almost always require that the effective spatial interaction be lateral inhibition. When we chose parameters outside this range, we produced shell patterns that do not correspond to any known species (Fig. 3*B*).

Although our basic model is capable of producing many of the observed patterns, there are some species (e.g., *Conus textile*) in which we had to assume that some of the parameters were modulated in space and "time" to specify prepatterns. The prepatterns generally are periodic or consist of a localized region where the parameter is greater or smaller than that of the surrounding region. Such prepatterns could be hard-wired into the network or could themselves be produced by another neural network in the central ganglia (further details are provided in *SI Appendix*, Supplement A).

Finally, we should point out some important differences between the morphogen models for shell patterns developed by Meinhardt and coworkers (8, 9) and the neural network model used here (1, 2). Structural studies provide strong evidence that shell patterns are a neurosecretory phenomenon rather than a diffusing morphogen phenomenon (2). However, from a theoretical viewpoint, morphogen models can be viewed as an approximation to the neural net model when the range of communication between neurons is short (9, 10). Therefore, in principle, morphogen models could have been used instead of the neural model (11). From a practical viewpoint, however, this would be considerably more difficult because a separate morphogen model is required for each shell pattern, whereas the neural model has a single set of parameters that are varied to match each pattern. Also, as the neural models are more general, they can generate a wider variety of patterns than can diffusible morphogen models. One other difference is fundamental. Morphogen models described by diffusion-reaction dynamics unfold with no "memory" of the system state other than the current state. The neural model, however, is a sensory feedback system in which the current secretion depends on sensing the history of the pattern before the current state.

Phylogenetic Analyses

Inferred Parameter Values for Each Species. We chose 19 species from the phylogeny published by Nam et al. (3) based on mitochondrial cytochrome C oxidase subunit I and rDNA sequences and on internal transcribed spacer 2 sequences from nuclear ribosomal DNA. There were sufficient data that the order of

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Fig. 2. Definition of cell specific model parameters. (A) Gaussian excitation and inhibition kernels whose difference creates the Mexican-hat spatial field. (*B*) Temporal filter implementing delayed inhibition. β_1 (β_2) is the strength of the temporal excitation (inhibition) and c_1 (c_2) is the decay in "time" of the excitation (inhibition), wherein time is measured discretely in secretory bouts, denoted by n ($0 < c_1 < c_2 < 1$, so that the inhibition decays more slowly in time; thus, the most recent activity is excitatory and more distant activity is inhibitory). (C) Sigmoid response function of the secretory cells; ν is the sharpness of the nonlinearity and θ is the midpoint (there is one nonlinearity for excitation and one for inhibition).

branching events in the phylogeny could be completely determined with a high degree of statistical confidence.

The neural network model was fit to each living species in the phylogenetic tree. Nine species can be reproduced using the basic model (i.e., a single neural network). Six species (*Conus tessulatus, Conus aurisiacus, Conus ammiralis, Conus orbignyi, Conus stercusmuscarum*, and *Conus laterculatus*) require a spatial prepattern (generated by a "hidden" network), and four species (*Conus dalli, C. textile, Conus aulicus,* and *Conus episcopatus*) require spatiotemporal prepatterns (generated by one or two hidden networks). In phylogenetic analyses of these shell parameters, we focus on the primary network, which can be compared across all species. The fitted parameters for each species are shown in *SI Appendix,* Supplement C. Images of real shells and their corresponding simulated ones are shown in Fig. 4.

Test for Phylogenetic Signal in Estimated Parameter Values. Phenotypic traits like body size and shape typically exhibit a substantial degree of "phylogenetic signal," meaning that they are inherited, and the phenotypes of closely related species are strongly correlated (12). One purpose of the present study is to determine whether parameters of the neural-network model exhibit a phylogenetic signal. They will if the construction of the model accurately approximates the real developmental process of shell patterning. Therefore, we tested for a phylogenetic signal when the model parameters are fitted to the observed pigmentation patterns. A basic test for phylogenetic signal in traits is to compare the observed data to a null model in which all phylogenetic signal are obliterated by randomly shuffling the species names or trait values at the tips of the phylogenic tree (13). To test for a phylogenetic signal in the neural network parameters, we constructed a neighbor-



Fig. 3. (A) Both noise and chaos generate within-species pattern diversity. *a*, Three real *C. crocatus* shells. *b*, Three shells generated with 1% noise only. *c*, Three shells generated with slightly different initial conditions, but no noise. *d*, Three shells with both 1% noise and slightly different initial conditions. (*B*) Two examples of "unknown" patterns having too-wide inhibition fields.



Fig. 4. Maximum-likelihood estimates of ancestral shell patterns. Shells of living species are displayed at the tips; to the right of these are shells "grown" in the computer by using the neural-network model and the fitted parameters. By using a Brownian motion model for the evolution of continuous traits, the maximum-likelihood value was estimated for each neural network parameter at each node. The neural network model was used to produce the shells using the estimated parameters for each node. Color is not part of the neural network model, so it was added independently to the models of living shells, and then mapped onto the phylogeny (using maximum likelihood) as a binary trait (black/white or brown/white). The text includes further details.

joining phylogeny of the 19 species based on the parameter values alone and compared it with the DNA phylogeny of Nam et al. (3). The parameter-based phylogeny was obtained as described in *SI Appendix*, Supplement B.

For each method of measuring distances between trees, we constructed a null distribution on tree-to-tree distances by taking the parameter-based tree and randomly reshuffling the species names. The distances between the randomized null-parameter tree and the DNA tree were then calculated. This procedure was repeated 10,000 times to produce the null distribution.

The trees are compared in Fig. 5. Despite several dissimilarities between the DNA- and parameter-based trees, the observed distance between the trees is much less than expected under the null hypothesis of only random similarity between the trees (*SI Appendix*, Fig. S7). The differences are statistically significant—P = 0.0146 for the topology-based distance measure and P = 0.0001 the branch-length-based distance measure—indicating that the observed distance was smaller than all the 10,000 null distances

generated. We conclude that there is a phylogenetic signal in the parameter values, despite the fact that they do not perfectly reflect the phylogenetic relationships of the group.

Similarity of DNA- and Parameter-Based Trees. Looking more closely at the parameter and DNA trees, we can see there is broad similarity but with notable exceptions. In both trees, there are two large clades, called arbitrarily clade 1 (*C. stercusmuscarum, C. aurisiacus, Conus pulicarius, Conus arenatus,* and *C. laterculatus*) and clade 2 (*C. gloriamaris, C. dalli, C. textile, Conus omaria, C. episcopatus,* and *C. aulicus*), that are nearly the same in both trees, although the detailed branching order differs slightly. In addition, *Conus bandanus* and *Conus marmoreus* are sister groups in both trees. There are some conspicuous differences, however. Most notably, *Conus furvus, C. tessulatus,* and *C. orbignyi* form a tight clade in the parameter tree yet are widely separated in the DNA tree. In fact, in the DNA tree, *C. orbignyi* is a well supported out-group to the other 18 species. *C. ammiralis* is part of clade 2 on the DNA tree



Fig. 5. Comparison of the DNA-based phylogeny of cone snails (*Left*, after Nam et al. (3), unrooted for display) and the parameter-based tree (*Right*, present study). Species labeled in blue exhibit major changes in topological position in the parameter-based tree. The observed tree-to-tree distances are significantly shorter than expected under a null hypothesis of random similarity (*SI Appendix*, Fig. S7).

but is quite separate on the parameter tree. *C. crocatus* is in clade 2 on the DNA tree and in clade 1 on the parameter tree (Fig. 5).

The overall similarity of the DNA-based and parameter-based trees is consistent with the hypothesis that the parameters of the developmental model evolved sufficiently slowly that sets of parameters in closely related species are similar. However, there are some exceptional lineages on which more rapid evolution of parameters seems to have occurred. The three species C. furvus, C. tessulatus, and C. orbignyi appear to have converged not only in pattern but in the developmental process that produces that pattern. C. crocatus appears to have shifted its pattern to become similar to species in clade 2, and both C. ammiralis and C. consors have undergone relatively rapid evolution that resulted in quite distinct patterns. The apparently higher rate of parameter evolution on these lineages is consistent with the action of natural selection either directly on pigmentation pattern or indirectly as a correlated response to selection on physiological processes that affect parameter values. In the absence of knowledge of the physiological basis of parameter values, we have no way to directly test for natural selection.

Parametric and nonparametric tests of the Brownian motion model. The estimation of parameter values for ancestral species in the phylogeny is most easily done if the Brownian motion model of continuous trait evolution can be used. Therefore, when we had established that detectable phylogenetic signal existed in the neural network parameters, we conducted a series of tests to assess the utility of Brownian motion versus other models for modeling the evolution of neural network parameters, as recommended by Blomberg et al. (14). We concluded that Brownian motion was an overall reasonable first approximation for the evolution of neural network parameters (*SI Appendix, Supplement B*).

Discrete Characters. Hidden Networks Treated as Discrete Characters.

We can treat the presence or absence of a hidden neural network as a binary discrete character. Then, the presence or absence of this character can be mapped onto the phylogeny by using parsimony and maximum-likelihood reconstruction for discrete characters. The two methods give identical results. The presence of hidden networks was restricted to small subclades of the full clade. The presence/absence of hidden networks (Fig. 6 A and Bshow the presence of a space-time-dependent hidden network and space-dependent hidden network, respectively) showed strong phylogenetic clustering. Relatively few transitions from simple models (i.e., no hidden network) to complex models (i.e., containing a hidden network) were needed for either character. For the space-time-dependent network, species in two small clades (*C. episcopatus/C. aulicus* and *C. textile/C. dalli*) are complex. For the space-dependent hidden network, a complex pattern is more dispersed in the phylogeny.

Discrete phenotypic characters. Other discrete characters were also mapped for comparison with the results for hidden networks. We mapped several discrete phenotypic characters on the phylogeny (*SI Appendix*, Supplement B). Cone shape is fairly scattered but shows some uniformity in small clades. Strikingly, prey preference shows extremely high conservation [as was clear in the discussion of Nam et al. (3)] compared with shell pattern characters. Each major clade is almost completely restricted to a certain prey, and the entire pattern is explained by the minimum possible number of transitions.

Fig. 6 shows the distributions of stripes and triangles in this group and the maximum-likelihood assignment of ancestral states. The presence and absence of stripes, in particular, is scattered throughout the phylogeny, indicating that they are evolutionarily labile, although triangle presence/absence shows some correlation with large clades. These observations are confirmed by standard parsimony statistics and their comparison with randomized-tip null models; presence/absence of stripes, despite these being visually striking patterns used in identification, appear to lack significant phylogenetic signal in that they do not show significantly more congruence with the phylogeny than is expected under the null model in which character states have been randomly shuffled among the phylogeny tips.

Inference of Ancestral Shell Patterns. We used a Brownian motion model to estimate parameter values in the species ancestral to the living species. We then ran the neural-network model with these estimated parameter values to predict the pigmentation patterns in the ancestral species. Those patterns are shown at the nodes in Fig. 4. Ancestral states for each parameter common to all species were estimated by using maximum-likelihood estimation on the tree inferred from DNA sequences, modeling the evolution of each parameter as an independent Brownian motion process (15, 16). Two other available methods—generalized least-squares and phylogenetically independent contrasts—gave similar estimates.

For the additional parameters used in the hidden networks, ancestral character estimation was performed as follows. Phylogenetically independent contrasts were applied to reconstruct the ancestral states of the hidden networks because it works from the tips downward, and so, unlike maximum likelihood, can be used when parameters for hidden networks are not available in the rest of the clade.



Fig. 6. Maximum-likelihood estimates of selected discrete characters. The relative simplicity of the inferred evolution of pattern complexity is in striking contrast to what can be inferred about the evolution of specific features of the patterns when they are described as discrete characters, as illustrated in A–D.

The ancestral shell patterns are shown in Fig. 4. Each estimate has an associated variance and confidence intervals. To test the robustness of the ancestral patterns to uncertainty in parameter estimates, we randomly generated sets of parameters from the distribution of each parameter and generated ancestral patterns from each set. We found that some ancestral patterns are quite robust to uncertainty in estimated parameters whereas others are not. Fig. 7 shows two examples of each kind. The ancestral patterns for nodes 25 and 29 are quite similar for different sets of estimated parameters, whereas those for nodes 27 and 31 differ greatly among sets of estimated parameters, although various detailed similarities can still be detected even among these shells because of the underlying similarity of neural network parameters.

Discussion and Conclusions

We have taken a step in applying modern phylogenetic methods to understanding the development of complex phenotypic characters. The pigmentation patterns of *Conus* shells can be generated by a neural-network model that has a sound anatomical and physiological basis. The model parameters fitted to observed patterns show a substantial phylogenetic signal, indicating that the processes governing evolutionary change in shell patterns are, to some extent, gradual across the phylogeny. Our analyses have allowed us to estimate the shell pigmentation patterns of ancestral species, identify lineages in which one or more parameters have evolved rapidly, and measure the degree to which different parameters correlate with the phylogeny.

Our results are summarized in Fig. 4. This figure shows that pigmentation patterns in living species are well approximated by the neural-network model presented in this study. It also shows the inferred ancestral shell patterns. Often, recent ancestors of sister species show recognizable similarity to the pigmentation patterns in living species (e.g., nodes 26-27 and 31-32). Nodes more remote from the present often show ancestors that are generally similar to the living species (nodes 21, 22, 24, and 33). However, some ancestors are strikingly different from any of the living species in the group we analyzed. Interestingly, such patterns can be found in other living species. For example, the strong striping perpendicular to the axis of coiling of the shell found in node 37 is quite similar to that of Conus hirasei, Conus papuensis, or Conus mucronatus (17). Striping parallel to the axis of coiling of the shell, observed in other estimated ancestors, can also be found in living species, for example in some specimens of Conus hyaena and Conus generalis (ref. 17, pp. 354 and 392).

A unique feature of our results is that the inferred pigmentation patterns of ancestors may be quite different from the patterns of their descendants. The patterns generated by the neural-network model are not necessarily smooth functions of the parameter val-

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Fig. 7. Possible realizations of ancestors. For each neural network parameter at each node, five values were drawn from their estimated distribution by using the predicted uncertainty around the maximum-likelihood estimate. Each set of parameter values for a node was then input into the neural-network model to produce a depiction of the possible ancestor. Some shell patterns appear to occupy larger regions of parameter space, and are thus more robust to perturbation. However, similarities can still be discerned even in shells that appear quite different at first glance.

ues. Instead, they can vary discontinuously when parameter values move into a different bifurcation region that produces qualitatively different patterns. The role of bifurcation boundaries in evolution was recognized in earlier studies of limb morphogenesis (18, 19). This feature of our results is quite different from what is usually found when inferring ancestral states of continuously variable characters. A well known limitation of methods for estimating ancestral states is that it is impossible for estimates to fall outside the range of the living species analyzed. This limitation does not apply to pigmentation patterns. Although the same averaging procedure is being used on each parameter of the neural-network model, it is possible, and even likely, that a set of estimated parameters will be in a region of parameter space not inhabited by any living species. In addition, the sensitivity of the neural network to perturbations means that small, gradual evolutionary shifts in one or a few parameters of the neural network can shift a shell from one pattern regime into an entirely dissimilar one.

We have necessarily made simplifying assumptions in our analysis to illustrate the overall logic of our method in a straightforward way. Although the DNA-based tree used in this study has strong statistical support, an important assumption is that the branch lengths inferred from the DNA sequence data are known without error, and that they have been accurately renormalized to an absolute time scale. A more formal analysis would begin with the raw DNA sequence alignment and fossil calibration points, and then integrate ancestral state estimates and parameters of evolutionary models, over the space of data-supported chronogram phylogenies (20).

A second assumption is that the set of parameter values for each species is unique and estimated without error. Given the number of parameters involved, a formal proof of uniqueness seems impossible; however, extensive experience with the numerical properties of the model suggests that each pattern is determined by a unique optimal (in the sense of a best fit to the observed pattern) set of parameters.

A third assumption is that the parameters evolved independently of one another on the phylogeny. That assumption is largely supported by our analysis of phylogenetically independent contrasts. Correlation in parameters could be accounted for by using a model of correlated Brownian motion on the phylogeny, but such a model was not needed for our analysis.

In estimating parameters of ancestral species and predicting their pigmentation patterns, we have not taken into account the range of parameters consistent with estimated values for living species. Parameter values estimated by using maximum likelihood and a Brownian motion model have associated confidence intervals that could make more than one qualitatively different pigmentation pattern for each ancestral species consistent with patterns in living species. Application of our method to a group of cone snails with a detailed fossil record-for example, those in southeastern North America (21)-might allow a more rigorous assessment of the accuracy of these techniques, and of what degree of uncertainty should be assigned to them. Usefully and remarkably, shell pigmentation patterns in fossil Conus can be visualized under UV light (21). Application of this technique to Conus fossils could provide a partial validation of our predicted ancestral patterns.

Our analysis is somewhat similar to that of Allen et al. (11), who examined spotted patterns in felids by using a morphogen-diffusion model of pattern formation. Allen et al. showed that there is little phylogenetic signal in the model parameters, indicating that spotting patterns in felids evolve convergently under ecological influences. One difference between their study (11) and the present one is that we found phylogenetic signal in most of the neural network parameters that produce shell pigmentation patterns. This allowed us to infer ancestral patterns and to identify lineages in which relatively rapid evolution of some parameters have taken place.

We found phylogenetic signal in the continuous parameters of the primary neural network and in the presence/absence of a hidden network, suggesting that the model reasonably approximates the developmental processes underlying pigmentation patterns in the *Conus* species we considered. In contrast, various features of the pigmentation patterns, such as the presence of stripes and dots, do not have significant phylogenetic signal (*SI Appendix*, Tables S2–S4). This is in agreement with the conclusion of Hendricks.*

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Supplemental Information for the evolution of patterns on *Conus* shells

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Supplement A: Neural Network Model for the Conus Shell Pigmentation Patterns

1 Mathematical Formulation

In this part, we derive a shell model from a general neural model. The shell model consists of three parts: the sensory cell model, the neural model and the secretory cell model. The sensory cell model and secretory cell model are derived from first-order dynamics, and neural model is derived from the Wilson-Cowan equation (1, 2). First, we derive a continuous model. Then we discretize it to get the discrete model that is used for simulations. Finally, we discuss its relations to diffusion-reaction models.

1.1 Deriving a Shell Model from a General Neural Model

In this section we derive the model equations starting from the Wilson-Cowan model for the firing rate pattern of a general excitatory/inhibitory network (1, 2).



Figure S1. Schematic representation of the coordinate systems used in the derivations.

To fix our notation, we denote by 0,1,2,3 the shell pattern, sensory, neural, and secretory cells, respectively, with coordinate systems shown in Figure S1. Assume the shell is a rectangle with coordinates (x_0 , y_0), where $0 \le x_0 \le L$ and $0 \le y_0 \le T_s$. $y_0 = 0$ is the growing edge. The mantle has its own coordinates (x_1 , y_1), where $0 \le x_1 \le L$ and $0 \le y_1 \le T_M$. We assume these coordinates do not change. The sensory cells are distributed on the mantle and 'taste' the pigments. The neural cells are aligned with the growing edge along the line $y_0 = 0$ or $y_1 = 0$.

1.1.1 The model for sensory cells

We construct models for the sensory cells, neurons and secretory cells separately (Figure S1). The sensory cells are distributed in the mantle with coordinates (x_1, y_1) . A sensory cell at (x_1, y_1) tastes the pigment on the shell at location (x_0, y_0)

In general, the activity of the sensory cells at (x_1, y_1) on the mantle satisfy the Wilson-Cowan equations:

Equation 1
$$\tau^{(1)} \frac{\partial u^{(1)}(x_1, y_1, t)}{\partial t} = -u^{(1)}(x_1, y_1, t) + S^{(1)}(\psi^{(1)}(x_1, y_1, t))$$

where $S^{(1)}(\psi^{(1)}(x_1,y_1,t))$ is the function computing the firing rate given the input:

Equation 2
$$\psi^{(1)}(x_1, y_1, t) = K^{(1)}(x_1, y_1) * *u^{(1)}(x_1, y_1, t) + M^{(1)}(x_1, y_1)$$

 $K^{(1)}(x_1, y_1)$ models the recurrent connections between the sensory cells. $M^{(1)}(x_1, y_1)$ is the input to the sensory cell located at (x_1, y_1) . On the neural time scale the pigment on the shell does not change, so $M^{(1)}(x_1, y_1)$ is independent of time, and is given by the double convolution:

Equation 3
$$M^{(1)}(x_1, y_1) = \int_0^L \int_0^{T_s} W^{(1)}(x_1 - x_1, y_1 - y_1) P(x_1, y_1) dx_1 dy_1$$

The recurrent connections make it difficult to compute the steady state of the sensory cells, so we assume that there are no recurrent connections between sensory cells. In the steady state, we can set $\partial u^{(1)}(x_1, y_1, t) / \partial t = 0$, so that

Equation 4
$$u^{(1)}(x_1, y_1) = S^{(1)}(M^{(1)}(x_1, y_1))$$

Next, we assume that the weight kernel $W^{(1)}(x - x_1, y - y_1)$ is a two-dimensional delta function, so that a sensory cell located at (x_1, y_1) tastes only the pigment, *P*, at location $(x_0, y_0) = (x_1, y_1)$. Then we have

Equation 5
$$u^{(1)}(x_1, y_1) = S^{(1)}(P(x_0, y_0))$$

This will become the sensory input to the neural net.

1.1.2 The model for the neural net

Starting again with the steady state Wilson-Cowan equations:

Equation 6	$u_e^{(2)}(x_2) = S_e^{(2)}(\psi_e^{(2)}(x_2))$
Equation 7	$\psi_{e}^{(2)}(x_{2}) = K_{ee}^{(2)}(x_{2}) * u_{e}^{(2)}(x_{2}) - K_{eh}^{(2)}(x_{2}) * u_{h}^{(2)}(x_{2}) + M_{e}^{(2)}(x_{2})$
Equation 8	$M_{e}^{(2)}(x_{2}) = \int_{0}^{L} \int_{0}^{T_{M}} W_{e}^{(2)}(x_{2} - x_{2}, y_{2} - y_{2}) P(x_{2}, y_{2}) dx_{2} dy_{2}$

For inhibitory cells, we have similar equations. $M_e^{(2)}(x_2)$ is the sensory input to the neuron located at x_2 .

Again, assume there are no recurrent connections. Then we have the steady state equation:

Equation 9
$$u_e^{(2)}(x_2) = S_e^{(2)}(M_e^{(2)}(x_2))$$

With a similar equation for the inhibitory neurons.

1.1.3 The model for the secretory cells

The secretory cells have first order temporal kinetics:

Equation 10
$$\tau^{(3)} \frac{\partial u^{(3)}(x_3,t)}{\partial t} = -u^{(3)}(x_3,t) + S^{(3)}(\psi^{(3)}(x_3,t))$$

Equation 11 $\psi^{(3)}(x_3,t) = K^{(3)}(x_3) * u^{(3)}(x_3,t) + M^{(3)}(x_3)$

Equation 12
$$M^{(3)}(x_3) = \int_0^L (W_e^{(3)}(x_3 - x_3)u_e^{(2)}(x_3) - W_h^{(3)}(x_3 - x_3)u_h^{(2)}(x_3))dx_3$$

The sensory inputs to the secretory cells are the weighted difference between excitatory neurons and inhibitory neurons. If there are no recurrent connections, then the steady state equation is:

Equation 13
$$u^{(3)}(x_3) = S^{(3)}(M^{(3)}(x_3))$$

If we assume that $W_e^{(3)}(x_3)$ and $W_h^{(3)}(x_3)$ are delta functions, then we obtain

Equation 14 $u^{(3)}(x_3) = S^{(3)}(u_e^{(2)}(x_3) - u_h^{(2)}(x_3))$

This is the pigment at the growing edge, i.e.

Equation 15 $P(x_0,0) = S^{(3)}(u_e^{(2)}(x_3) - u_h^{(2)}(x_3))$

1.1.4 The shell model

Combining the models for sensory, neural and secretory cells, we have the complete shell model:

Equation 16	Sensory cells:	$u^{(1)}(x_1, y_1) = S^{(1)}(P(x_1, y_1))$
Equation 17	Neural cells:	$u_{e,h}^{(2)}(x_2) = S_{e,h}^{(2)} \left(\int_0^L \int_0^{T_M} W_{e,h}^{(2)}(x_2 - x_2, -y_2) u^{(1)}(x_2, y_2) dx_2 dy_2 \right)$
Equation 18	Secretory cells:	$u^{(3)}(x_3) = S^{(3)}(u_e^{(2)}(x_3) - u_h^{(2)}(x_3))$
Equation 19	Pigment:	$P(x_0,0) = u^{(3)}(x_0,0)$

1.2 Discrete Shell Model

The *y*-axis is time in the past. Let Δ be the spatial thickness of a single bout of pigment. We also use Δ to discretize the sensory cells. Assume $T_M = Q\Delta$, which means the mantle can sense Qbouts of pigments into the past. Denote $P(x_0,t)$ as the pigment at position x_0 at bout time *t* (not in real time). $A^{(1)}(x_1,s\Delta,t)$ denotes the activity of sensory cell at position $(x_1,s\Delta)$ at bout time t. $A^{(2)}_{e,h}(x_2,t)$ denotes the activity of excitatory or inhibitory neurons at position x_2 at bout time t. $A^{(3)}(x_3,t)$ denotes the activity of secretory cell at position x_3 at bout time t. Then we have

Equation 20	$P(x_0, t) = A^{(3)}(x_0, t)$
Equation 21	$A^{(3)}(x_3,t) = S^{(3)}(A_e^{(2)}(x_3,t) - A_h^{(2)}(x_3,t))$
Equation 22	$A_{e,h}^{(2)}(x_2,t) = S_{e,h}^{(2)}(\int_0^L \int_1^Q W_{e,h}^{(2)}(x_2 - x_2, -s\Delta)A^{(1)}(x_2, s\Delta, t)dx_2^{'}ds)$
Equation 23	$A^{(1)}(x_1, s\Delta, t) = S^{(1)}(P(x_1, t-s))$

The discrete model is like:

Equation 24
$$P_{n+1}(x_0) = S^{(3)}(E_n(x_0) - H_n(x_0))$$

Equation 25
$$E_n(x_0) = S_e^{(2)}(\sum_{j=0}^{Q-1} \int_0^L W_e^{(2)}(x_0 - x_0^{'}, j) S^{(1)}(P_{n-j}(x_0^{'})) dx_0^{'})$$

Equation 26
$$H_n(x_0) = S_h^{(2)} (\sum_{j=0}^{Q-1} \int_0^L W_h^{(2)}(x_0 - x_0^{'}, j) S^{(1)}(P_{n-j}(x_0^{'})) dx_0^{'})$$

Figure S2 illustrates our discrete network structure. Our network is s simple feed forward network.



Figure S2. Illustration of network structure we use. Each neuron connects to all the sensory cells. Each secretory cell only connects to the excitatory neuron and inhibitory neuron at the same location.

1.3 Simulation Model

We assume the two dimensional space-time $W_{e,h}^{(2)}(x_2, j)$ filter is separable, i.e.

Equation 27
$$W_{e,h}^{(2)}(x_2, j) = w_{e,h}^{(2)}(x_2)v_{e,h}^{(2)}(j)$$

This widely adapted simplification is to accelerate the simulations.

In the simulation, we assume the spatial filter $w_{e,h}^{(2)}(x)$ is a Gaussian kernel

Equation 28
$$w_{e,h}^{(2)}(x) = \frac{\alpha_{e,h}^{(2)}}{2\pi\sigma_{e,h}^{(2)}} e^{-\frac{x^2}{2\sigma_{e,h}^{(2)}}}$$

The difference of Gaussian kernels can generate a 'Mexican Hat', which is necessary for pattern formation. In our model, both excitatory and inhibitory neurons have Gaussian kernels. The excitatory Gaussian kernel has a narrower variance than the inhibitory Gaussian kernel. Thus, the difference between them results in 'Mexican Hat'. The Mexican Hat kernel generates local activation and long-range inhibition that can generate periodic patterns. Similarly, the temporal

filter $v_{e,h}^{(2)}(j)$ is assumed to be a difference of exponential functions that generate a local activation and long-range inhibition in time:

Equation 29
Equation 29

$$v_{e}^{(2)}(j) = \beta_{e1}c_{e1}^{j} - \beta_{e2}c_{e2}^{j} \text{ for } j \ge 0, \ \beta_{e1} = \beta_{e2} + 1, \ c_{e1}^{j} < c_{e2}^{j}$$
Equation 30

$$v_{h}^{(2)}(j) = \beta_{h1}c_{h1}^{j} - \beta_{h2}c_{h2}^{j} \text{ for } j \ge 0, \ \beta_{h1} = \beta_{h2} + 1, \ c_{h1}^{j} < c_{h2}^{j}$$

This is equivalent to a refractory period that can generate temporal oscillations.

We assume Q = n, which means the mantle covers all of the previous pigment so that sensory cells can sense all the previous pigment. We define the temporal convolutions as follows:

Equation 31
$$R_{e,n}(x) = \sum_{j=0}^{n-1} (\beta_{e1} c_{e1}^j - \beta_{e2} c_{e2}^j) S^{(1)}(P_{n-j}(x))$$

Equation 32
$$R_{h,n}(x) = \sum_{j=0}^{n-1} (\beta_{h1} c_{h1}^j - \beta_{h2} c_{h2}^j) S^{(1)}(P_{n-j}(x))$$

Then we have

Equation 33
$$P_{n+1}(x) = S^{(3)}(S_e^{(2)}(w_e^{(2)}(x) * R_{e,n}(x)) - S_h^{(2)}(w_h^{(2)}(x) * R_{h,n}(x)))$$

In order to compute the temporal convolutions efficiently, we define notations:

Equation 34
$$R_{e1,n}(x) = \beta_{e1} \sum_{j=0}^{n-1} c_{e1}^{j} S^{(1)}(P_{n-j}(x))$$

Equation 35
$$R_{e2,n}(x) = \beta_{e2} \sum_{j=0}^{n-1} c_{e2}^{j} S^{(1)}(P_{n-j}(x))$$

Equation 36
$$R_{h1,n}(x) = \beta_{h1} \sum_{j=0}^{n-1} c_{h1}^{j} S^{(1)}(P_{n-j}(x))$$

Equation 37
$$R_{h2,n}(x) = \beta_{h2} \sum_{j=0}^{n-1} c_{h2}^{j} S^{(1)}(P_{n-j}(x))$$

Then we obtain the following recursive equations

Equation 38
Equation 38

$$R_{e,n+1}(x) = R_{e_{1,n+1}}(x) - R_{e_{2,n+1}}(x)$$

Equation 39
 $R_{h,n+1}(x) = R_{h_{1,n+1}}(x) - R_{h_{2,n+1}}(x)$

Equation 40	$R_{e_{1,n+1}}(x) = \beta_{e_{1}} S^{(1)}(P_{n+1}(x)) + c_{e_{1}} R_{e_{1,n}}(x)$
Equation 41	$R_{e2,n+1}(x) = \beta_{e2} S^{(1)}(P_{n+1}(x)) + c_{e2} R_{e2,n}(x)$
Equation 42	$R_{h1,n+1}(x) = \beta_{h1} S^{(1)}(P_{n+1}(x)) + c_{h1} R_{h1,n}(x)$
Equation 43	$R_{h2,n+1}(x) = \beta_{h2} S^{(1)}(P_{n+1}(x)) + c_{h2} R_{h2,n}(x)$

So, Equations 33, 38, 39, 40, 41, 42 and 43 are implemented in our Matlab code to generate the patterns.

1.4 Model Parameters

The mantle's length is *L*. When we fix the number of sensory cells in our discrete model, this length only influences the interval between cells, which does not affect the patterns. So, we let L = 1 in all our simulations.

There is a sigmoid function for the sensory cell, excitatory neuron, inhibitory neuron and secretory cell, respectively. So there are four sigmoid functions in our model. In this part, we assume all cells belonging to the same type have the same sigmoid function, i.e. all the sensory cells have the same sigmoid function, all the excitatory neurons have the same sigmoid function, etc. In later sections, we'll discuss the cases where the cells belonging to the same type have different sigmoid functions, which can generate complex patterns. The analytic form of sigmoid function is:

$$S(x) = \frac{\gamma}{1 + e^{-\nu(x-\theta)}}$$

Each sigmoid function has 3 parameters, i.e. θ , v, γ . θ is the middle point of the sigmoid function, v is proportional to the slope at the middle point, and γ is the magnitude of the sigmoid function. We set $\gamma = 1$ for all sigmoid functions in our simulations. So we have 8 free parameters for the 4 sigmoid functions.

The spatial kernels are Gaussians of the form

$$w_{e,h}^{(2)}(x) = \alpha_{e,h}^{(2)} e^{-\frac{x^2}{2\sigma_{e,h}^{(2)^2}}}$$

What's important for the pattern formation is the difference between the excitatory and inhibitory kernels, so we set the magnitude parameter $\alpha_h^{(2)} = 1$. This setting leaves us with 3 free parameters for the spatial kernels. Since there are 3 free parameters for each temporal kernel, we have 6 free parameters for temporal kernels. Thus our model is controlled by 17 free parameters, all of which have direct cellular interpretations. It turns out that the region in parameter space that generate realistic shell patterns is rather small. So the parameter search to match each shell pattern is not as difficult as the dimensionality of the parameter space might indicate.

1.5 Related Models

Cellular automata models were first used to reproduce shell patterns (3-5). Although they can generate some observed patterns, they cannot explain how these patterns arise in animal markings.

Meinhardt and his coworkers (6-10) used morphogen, or Diffusion-Reaction (DR) models to reproduce a wide variety of shell patterns. DR models are inspired by the chemical diffusion of morphogens, but there is no experimental evidence found for diffusing morphogens in pattern formation for shells. DR models can be viewed as an approximation of neural activity when only nearest neighbor neurons communicate (chapter 12.4 in (11)).

B. Ermentrout *et al* (12) and A. Boettiger *et al* (13) proposed neural models to reproduce shell patterns. Their models are somewhat different because they have different refractory terms. In (13), the refractory term is the temporal convolution of all previous pigment deposition, while refractory term in (12) is the temporal convolution of all previous pigments except the previous time period. Our model is inspired by these models, but is different in that we do not use an explicit refractory term. In the previous models, the pigment is the difference between secretory cells' activities and the refractory term. In our model, the pigment results from the net activity of the secretory cells. The previous models can only generate basic patterns, but the current model includes 'hidden' networks, and so can generate more complex shell patterns.

2 Pattern Generation

2.1 Basic Patterns

Mathematically, as stated in (13), a Turing bifurcation leads to spatial instability, which generates stripes perpendicular to the growing edge; a Hopf bifurcation generates temporal instability leading to oscillations, which generate parallel stripes. An infinite saddle-node bifurcation probably underlies the travelling waves, but we have not proven this.



Figure S3. Bifurcations. Black indicates pigment. (a) Turing stripes. (b) Hopf bifurcation with synchronizing phase. (c) Hopf bifurcation with spatially continuously varying phases (Time increases upward)

Figure S3 (a) illustrates the formation of Turing stripes. In the very beginning, all secretory cells have deposited a very small amount of pigment. Then due to the temporal inhibition, the cells go through an unpigmented period, except for two small groups of cells on the boundaries. The two groups have reached their steady states. The activities of the two groups have lateral inhibition on their lateral regions, so their neighboring regions have no pigments. After the unpigmented period ends, an array of cells have small pigments. Near the boundary of the array, the cells have local activation on both sides, but only have one side of long-range lateral inhibition, so stripes come into being near the boundaries. *Consors* has this kind of pattern.

Hopf bifurcations can generate two categories of patterns. First, if the cells at different locations have synchronizing phase, then we get parallel lines, as shown in Figure S3 (b); second, if the cells have spatially continuously varying phase, then we get oscillations, as shown in Figure S3 (c).

If Turing bifurcation and Hopf bifurcation happen together, then we get Turing-Hopf bifurcation. There are two kinds of Turing-Hopf bifurcations depending on what the Hopf bifurcation is. If the Hopf bifurcation generates parallel lines, then we get checkerboard like patterns. And the checkerboard pattern can be in phase or out phase. In-phase checkerboard means all the checkers have the same phase in time. Out-phase checkerboard means phases of spatially neighboring checkers have 180 degrees difference. *Furvus* has in-phase checkerboard pattern. The main pattern of *tessulatus* is out-phase checkerboard. If the Hopf bifurcation generates oscillations, then we get oscillating Turing-Hopf bifurcation, checkers of which have spatially varying phases. The main patterns of *orbignyi* and *stercusmuscarum* are oscillating Turing-Hopf bifurcation.

When the secretory cell's activity represses its future activity while exciting lateral cells, we get travelling waves. When two waves collide, they may reflect, singularly annihilate or mutually annihilate. When the secretory cell has slightly wider excitatory range, then waves with changing speed emerge. *Gloriamaris, omaria, textile, dalli, episcopatus* and *aulicus* have travelling waves.

Travelling waves can also generate triangles and dots. When the speed of waves is very big at some time, then there will be a sudden stop of pigment along an array of cells, which is the base of the triangle. Then waves starting from the boundaries of the array travel back to the unpigmented region, leaving a triangular region without pigment. *Ammiralis, marmoreus,* and *bandanus* have triangles. Two waves starting at the same point travel outward. At some time, they change to travel inward to the unpigmented region, leaving a spot region without pigment. *Pulicarius, crocatus, arenatus* and *aurisiacus* have spots.

For Turing stripes, wider excitatory neuron spatial kernels lead to wider pigmented stripes, and wider inhibitory neuron spatial kernels generate wider unpigmented stripes. For the Hopf bifurcation, a wider excitatory neuron spatial kernel leads to wider pigmented parallel lines or oscillations, and a wider inhibitory neuron spatial kernel generates wider unpigmented parallel lines or oscillations. For waves, narrower spatial kernels can generate more dense waves. For triangles, narrower spatial kernels can generate more and smaller triangles. For dots, narrower spatial kernels can generate more and smaller dots.

2.2 Patterns with Spatial Pre-pattern

Some shells have more than one basic pattern. For example, *ammiralis* has triangles and Turing stripes. This pattern can be generated using two independent networks, one secretes pigment over another. In the current model we need use only one network to generate this complex pattern. The *ammiralis* shell is shown in Figure S4 (a). We view the triangles as the main pattern, so we find parameters to generate them. The stripes imply some parameter is different in the stripe regions. The simplest way to do this is to spatially vary the sigmoid function of the secretory cells. That is, the sigmoid function is assigned a spatial pre-pattern, and this spatial pre-pattern can be generated by a hidden network that changes the middle points of the sigmoid function of the secretory cell has a spatial pre-pattern. The pre-pattern and generated shell are illustrated in Figure S4 (b). In the main pattern region, the parameters generate triangles. In the stripe region, the system reaches uniform steady states. Some of the spatial pre-patterns could be generated by a third hidden networ which generates Turing stripes. In our simulations, however, we simply set the parameter's spatial pre-pattern for convenience. *ammiralis, tessulatus, laterculatus, aurisiacus, stercusmuscarum,* and *orbignyi* are generated with spatial pre-patterns.



spatial pre-pattern of $\theta^{(3)}$



generated ammiralis (b)



2.3 Patterns with Spatio-temporal Pre-pattern

For some more complex shells, such as *textile* shown in Figure S5 (a), a spatial pre-pattern only is not sufficient. On the *textile* shell, there are travelling waves, Hopf oscillations and Turing strips. What's more interesting is that travelling waves appear occasionally in the stripe region. One may consider using three independent networks to generate the three patterns independently, and stack them to get the *textile* pattern. However, if we do it this way, it is impossible to get travelling waves in the stripe region. Therefore, we assume that the sigmoid functions of the secretory cells have spatio-temporal pre-pattern(s). For *textile*, we view the travelling waves as the main pattern, so that there are two spatio-temporal pre-patterns: Turing stripes and Hopf oscillations. Of course, one could view the Hopf oscillations as the main pattern, and the other two as pre-patterns. Any of these different assignment of the main pattern and pre-patterns can generate this shell.

Since we assume the parameter's spatio-temporal pre-pattern is controlled by hidden network(s), we need to discuss how this is generated by the hidden network. Assume there are N hidden networks. Each network has its own set of sensory cells, neurons and secretory cells. We cannot see activities of the hidden networks directly. But their activities are reflected by the pre-patterns on the shell. Besides the N hidden networks, there is one visible network whose activity is the pattern on the shell. Another assumption we use is that each network can only sense its own activity. This assumption is rational since there are different kinds of sensory cells on our tongue, and these cells can sense different stimulus, like spicy, sweet, etc. Based on these assumptions, we propose a simple but effective method to couple these networks.

Use $P_n(x,i)$ to denote the activity of the *i*th network's secretory cell located at *x* and during the *n*th bout time. Assume $P_n(x)$ is the pattern on the shell. We make the parameter $v_n^{(3)}(x)$, slops of the middle points of the secretory cells' sigmoid functions varied by the activities of the *N* hidden networks as follows:

$$V_n^{(3)}(x) = V^{(3)} - \sum_{i=1}^N f_i(P_n(x,i))$$

Where $v^{(3)}$ is the basic value for $v_n^{(3)}(x)$. With this basic value only, the visible network generates the main pattern on the shell. The threshold function f_i is

$$f_i(P_n(x,i)) = \begin{cases} a_i & \text{if } P_n(x,i) \ge thres_i \\ b_i & otherwise \end{cases}$$

This threshold function means that the *i*th hidden network has only two kinds of impact on the visible network.

Figure S5 (b) shows the pre-patterns of *textile*. This pre-pattern is generated by 2 hidden networks. One generates oscillations and the other generates Turing stripes. On the main pattern region, the effect of oscillations is not strong enough to change the travelling waves pattern. In the Turing region, the effect is strong enough to change the pattern to stripes. And because of the effect of the oscillations, there are also oscillations emerging in the stripe regions. Interestingly, travelling waves emerge occasionally in the stripe regions.



Figure S5. (a)An example showing the main pattern and pre-patterns of *Textile*. (b)Generated *textile* and the spatio-temporal pre-pattern of $v^{(3)}$.

3 Patterns Observed In Nature Correspond to A Small Region of Parameter Space

The 'Mexican hat', or 'center-surround' neural field is required for pattern formation. Thus the inhibition must be longer range than the activation, but with smaller amplitude than the excitation. If excitation is long range, but inhibition is short range, there will be no pattern. In addition, the strength of the excitation and inhibition must be roughly the same. Indeed, we find that real shells have excitatory and inhibitory spatial kernels of limited width, i.e. their range of excitation is fairly local, and the excitation and inhibition is roughly balanced.

Figure 3(Right) in the main text shows two examples of patterns generated by the neural model for which we have found no representative species. Interestingly, patterns such as these are generated when the neural net has highly 'unbalanced' excitation vs. inhibition. The shell patterns in Figure 3(Right) are unrealistic because the inhibition range is too large. We find that unrealistic patterns always have inhibition fields that are too long-range. The parameter region of realistic shells and the parameters of unknown A and B shown in Figure 3(Right) are illustrated in Table S1

Parameters	Ran	ige	Unknown A	Unknown B
$\boldsymbol{v}^{(1)}$	[6	15]	15	10
$oldsymbol{ heta}^{(1)}$	[0.25	0.42]	0.3	0.3
$V_e^{(2)}$	[5	17]	5	5
$ heta_e^{(2)}$	[0.01	0.3]	0.1	0.1
${oldsymbol{\mathcal{V}}}_h^{(2)}$	[5	20]	5	5
$oldsymbol{ heta}_h^{(2)}$	[0.06	0.35]	0.06	0.06
$v^{(3)}$	[5	40]	8	9
$ heta^{(3)}$	[0.03	0.37]	0.15	0.15
$\alpha_{_e}$	[1.3	7]	2.5	1.5
σ_{e}	[0.0044	0.06]	0.05	0.04
$\sigma_{_h}$	[0.007	0.18]	0.6	0.5
eta_{e1}	[1	5]	2.28	2.28
C _ e 1	[0	0.45]	0	0
C _{e2}	[0	0.56]	0.32	0.28
β_{h1}	[1	6.82]	1.78	1.08
C _{h1}	[0	0.15]	0	0
C _{h2}	[0	0.32]	0.32	0.25

Table S1 Parameters regions and the parameters for unknown pattern A and B shown in Figure 3(Right). The shaded row is the parameter that extends the inhibition to unrealistic size.

Supplement B: Statistical Methods and Phylogenetic Analyses

4 Software and phylogenetic data

The statistical and phylogenetic analysis was conducted using the R statistical language (14, 15) enhanced with the R phylogenetics packages APE (16), geiger (17), and phangorn (18) and custom scripts by N.J.M. (available upon request).

The phylogeny used for the analysis was that of Nam *et al.* (19). It is a well-supported phylogeny of the *Conus* species under study. The phylogeny, shown in Figure S6, is based on mitochondrial COI and rDNA sequences and on ITS2 sequences from nuclear ribosomal DNA. Nam *et al.* showed that ITS2 sequences resolved parts of the phylogeny that could not be resolved using only the mtDNA. The bootstrap values are sufficiently high that we will assume the phylogeny is correct.



Figure S6. (a) DNA-based tree used in this study, digitized from Nam et al. (2009). Taxa not used in this study have been excluded. (b) Ultrametric tree used in the study, calculated using NPRS. Absolute time information was not need for this study, so branch lengths are in units of relative time, with the root set to age 1.

The phylogeny was digitized to Newick format using GraphClick 3.0 (<u>http://www.arizona-software.ch/graphclick/</u>) and an in-house R script, TreeRogue 0.1 (available at: <u>https://stat.ethz.ch/pipermail/r-sig-phylo/2010-October/000816.html</u>). Correspondence of the topology and branch lengths of the digitized tree to the original was verified before use. Taxa

from Nam *et al.* that were not used in this analysis (*C. radiatus*, *C. parius*, *C. japideus*, *C. vimineus*, and outgroups) were dropped from the tree.

The time-calibrated ultrametric phylogeny was calculated using non-parametric rate scaling (NPRS) with the program r8s (20). Branch lengths are proportional to the total height of the tree; only branch lengths giving a relative measure of degree of shared ancestry were necessary for this study, rather than a tree calibrated to absolute time.

5 Test for phylogenetic signal in the parameter estimates

The estimation of neural network parameters is done manually through successive approximation. Before examining the matter, there was no guarantee *a priori* that there would one unique parameter solution to produce a specific shell pattern, or that the user will find, in a 19-dimensional parameter space, the single best match to the observed pattern in the living species. Additionally, it was conceivable that there might no phylogenetic signal in shell patterns of the living species either for biological reasons (rapid evolution to an equilibrium distribution of shell patterns) or technical ones (e.g., nonidentifiability of parameters of the neural network model).

To assess these assumptions, we tested whether or we could reject the null hypothesis of no phylogenetic signal in the 19 continuous parameters. A neighbor-joining phylogeny was constructed from the parameters, as follows. (1) each of the neural network parameters was normalized to a 0-1 scale; (2) the normalized parameters were used to calculate the pairwise Euclidean distance between each pair of species; (3) a phylogeny was inferred from the resulting distance matrix via neighboring-joining (16). The parameter-based tree was then compared to the DNA-based tree using tree-to-tree distance metrics. Two metrics were used. A measure of topological distance (considering just tree topology, and ignoring branch length) was provided by Robinson-Foulds topological distance, also known as symmetric difference (dist.topo, PH85 option in APE). This measures the number of partitions found in each tree which are not found in the other (21-23). A measure of distance that takes branch lengths into account is provided by Robinson-Foulds branch-length difference, which is the sum of changes in branch length that would have to be made to made two trees identical (24); dist.topo in APE, BHV01 option).





The null distribution of the of DNA-tree-to-parameter-tree distances was constructed by randomly shuffling the tip labels on the parameter tree and measuring the distance between it and the DNA-based tree. This was repeated 10,000 times. For each distance metric, the one-tailed p-value was obtained by comparing the rank of the observed distance between the parameter tree and the DNA tree to the 10,000 ranked distances from the null distribution. For branch-length difference, it is possible that the non-ultrametric nature of the DNA tree and the parameter tree, and the different total lengths of the trees, could bias the results, so the test (including generation of the null trees) was repeated on ultrametricized versions of the DNA tree calculated with

nonparametric rate smoothing (NPRS) in r8s (20) and the parameter tree (which was midpointrooted, and then rescaled with NPRS), with no substantial differences in results.

The observed tree-to-tree distances, and the null distributions on tree-to-tree distances, are shown in Figure S7 for the case where both compared trees were ultrametric. For both distance metrics, the observed distance between the DNA tree and the parameter-based tree was significantly closer than expected under the null of random similarity between the trees.

6 Phylogenetic signal in discrete characters

In order to compare our results to those that might be obtained using more traditional cladistic methods where the shell patterns are described with discrete character states, several prominent discrete shell-pattern characters were scored in the traditional cladistic manner based on photographs of the species. The characters were scored as follows: stripes: 0=absent, 1=weak, 2=strong; triangles: 0=present, 1=absent; dots: 0=absent, 1=present; color: 0=black and white only, 1=brown/orange and white. One shell shape character was also scored: conical shape: 0=strongly rounded, 1=weakly rounded, 2=no rounding (triangular cone). Additionally, prey preference was scored according to Nam et al.'s descriptions (0=piscivore, 1=verminivore, 2=molluscivore, ?=unknown). These characters were mapped onto the phylogeny using parsimony and maximum likelihood, which gave similar results. The congruence of these characters with the phylogeny was measured with consistency index (CI), retention index (RI), and rescaled consistency index (RCI), and the significance of these results was assessed by comparison to null distributions of these statistics generated by 1000 reshufflings of the tip data.

Phylogenetic signal in discrete characters. Parsimony-based summary statistics for the 8 discrete characters are shown in Table S2-S5.

character	Min. # of steps	Max # of steps	Obs. # of steps	CI	RI	RCI
space-time hidden network	1	4	2	0.50	0.67	0.33
spatial hidden network	1	6	4	0.25	0.40	0.10
stripes	2	12	9	0.22	0.30	0.07
triangles	1	6	4	0.25	0.40	0.10
dots	1	8	6	0.17	0.29	0.05
color	1	7	4	0.25	0.50	0.13
conical	2	10	5	0.40	0.63	0.25
food	2	8	2	1.00	1.00	1.00
total for all	11	61	36	0.31	0.50	0.15

Table S2 Parsimony summary statistics (CI, RI, RCI) for discrete characters along with the inputs to these statistics. Values closer to 1 indicate more congruence between the character and the phylogeny. However, these must be compared to null distributions to determine if the observed values of the statistics are higher than would be expected under the null hypothesis of no phylogenetic signal (see Table S3 and Table S4).

character	Min. # of steps	Max # of steps	Obs. # of steps	CI	RI	RCI
space-time hidden network	1	4	3.74	0.27	0.09	0.03
spatial hidden network	1	6	5.27	0.20	0.15	0.04
stripes	2	12	9.38	0.22	0.26	0.06
triangles	1	6	5.38	0.19	0.12	0.03
dots	1	8	6.48	0.16	0.22	0.04
color	1	7	5.95	0.17	0.17	0.03
conical	2	10	8.50	0.24	0.19	0.05
food	2	8	7.29	0.28	0.12	0.05
total for all	11	61	51.98	0.21	0.18	0.04

Table S3 Means of summary statistics calculated on 1000 draws from the null hypothesis where character states have been randomly shuffled on the tips (no phylogenetic signal).

character	Obs. # of steps	CI	RI	RCI
space-time hidden network	0.030	0.000	0.000	0.000
spatial hidden network	0.139	0.030	0.030	0.030
stripes	0.555	0.168	0.168	0.168
triangles	0.139	0.010	0.010	0.010
dots	0.485	0.129	0.129	0.129
color	0.050	0.010	0.010	0.010
conical	0.010	0.000	0.000	0.000
food	0.010	0.000	0.000	0.000
total for all	0.010	0.000	0.000	0.000

Table S4 Non-parametric p-values (one-tailed) of summary statistics, based on the rank of the observed statistic amongst the ranks of the 1000 null draws of the statistic summarized in Table S3. The null hypothesis of only random congruence with the phylogeny cannot be rejected at the p=0.05 level for the stripes and dots characters. No Bonferroni correction for multiple tests has been applied; if it is, then s3n_spd, triangles, and color also fail to reject the null at the p<0.05 significance level.

Two discrete characters, stripes and dots, fail to reject the null hypothesis of only random congruence with the phylogeny, with one-tailed non-parametric p-values > 0.05 (Table S4). For each statistic, a value of 1 indicates perfect congruence with the phylogeny, 0 means no congruence. The statistics for prey preference are all 1, indicating perfect congruence with phylogeny (Table S4). And the discrete characters, all taken together, also exhibit significant congruence (Table S4). However, the variability in stripes and dots indicates substantial homoplasy in these characters, despite their obviousness to human observers. This is likely an indication that multiple convergent pathways able to produce these patterns.

7 Test of character independence

Before estimation of ancestral states was attempted, the parameter estimates for living species were examined for correlation structure, as the simplest methods of ancestral character estimation assume that each character is independent. Using standard correlation analysis (Pearson's product-moment correlation, p-values produced with t-test), only 10 of 171 parameter pairs exhibited statistically significant correlation (p < 0.05 after Bonferroni correction for multiple tests).

Standard correlation analysis is dubious when the data may have correlation due to shared phylogenetic structure, so the correlation analysis on the 19 species was repeated on the 18 available phylogenetically-independent contrasts (PICs) (25). PICs were calculated using an ultrametricized phylogeny derived from the Nam *et al.* molecular tree by nonparametric rate smoothing (NPRS) (20). Using PICs, only 7 of 171 parameter pairs were statistically significantly correlated. Of these, two pairs were the β parameters of the temporal kernels of the excitatory and inhibitory neurons (β_{e1} and β_{e2} formed one pair, and β_{i1} and β_{i2} the other), which were always perfectly correlated in the estimates made on living species (the first parameter of the pair is always 1 unit higher than the second). The others were correlations of about 0.75-0.9 between parameters of the response functions of the excitatory and inhibitory neurons and the parameters of the temporal kernels for the excitatory and inhibitory neurons. A more elaborate analysis might take these correlations into account, or even take into account weak correlation structure that exists despite statistical insignificance, but for the present purpose of assessing the feasibility of integrating the developmental model with phylogenetic methods, statistical simplicity was preferred, and was judged to be a reasonable approximation given the overall weak correlation structure.

8 Model Selection

Tests for trait correlations using Phylogenetic Independent Constrasts (PIC), and the estimation of ancestral states, are most easily performed if the trait data can be modeled as evolving under a Brownian motion process. Under Brownian motion, traits wander without limit such that the expected variance (σ^2) between lineages increases as a linear function of phylogenetic distance.

This may be a valid approximation within a relatively closely-related clade where the traits have not yet run up against intrinsic limits (26-28). The Brownian motion model was tested against a variety of other models of continuous trait evolution using parametric methods (likelihood ratio test (LRT) and Akaike Information Criterion (AIC), (29) as well as nonparametric methods; the latter are expected to be more robust in the situation where the low number of taxa (here, 19) mean that the asymptotic assumptions of maximum likelihood inference may not be met in full. (30)

Seven models for the evolution of continuous traits were compared. The likelihood of the trait data under of each of the models was calculated using the R package geiger. (17) The Brownian motion model has two parameters (global mean and rate of variance increase); all of the others, except white noise, add one additional parameter so as to model stabilizing selection (Ornstein-Uhlenbeck, OU), lack of phylogenetic signal on internal branches (lambda), speciational or punctuated evolution (kappa), increases or declines in rate across the tree (delta), or an early burst of evolutionary change which then declines. White noise assumes no phylogenetic signal and simply models the data as a normal distribution with a mean and variance. (17, 27, 29, 30) The log-likelihoods for the neural network parameter data under each model are given in Table S5.

Model	s1n	s1t	s2en	s2et	s2in	s2it	s3n	s3t	sf2ea	sf2ed	sf2id	tf2eb1	tf2ec1	tf2eb2	tf2ec2	tf2ib1	tf2ic1	tf2ib2	tf2ic2	kz	kr	turn	Ν	T
Brownian	-46.9	28.6	-50.3	24.6	-54.7	11.5	-68.1	14.1	-30.1	55.5	29.1	-23.8	13.4	-23.8	10.1	-35.2	30.4	-35.2	17.5	61.0	40.8	-44.7	-100.4	-136.1
0-U	-46.9	28.6	-50.3	24.8	-54.3	13.3	-68.1	14.3	-30.1	56.8	31.4	-23.8	13.4	-23.8	10.1	-31.4	30.4	-31.4	17.9	61.5	43.5	-42.2	-100.4	-134.7
lambda	-46.9	28.6	-50.3	24.6	-54.7	12.2	-68.1	14.1	-30.1	56.7	30.0	-23.8	13.4	-23.8	10.1	-31.5	30.4	-31.5	17.5	61.0	43.1	-42.3	-100.4	-134.7
kappa	-46.9	28.6	-49.1	24.6	-54.7	11.5	-68.1	14.1	-29.7	55.5	29.5	-23.8	13.4	-23.8	10.1	-35.1	30.4	-35.1	17.5	61.0	41.4	-44.5	-100.4	-136.1
delta	-46.8	28.7	-50.1	25.0	-54.1	12.9	-91.2	14.4	-29.9	56.7	30.7	-23.8	13.7	-23.8	10.2	-33.2	30.5	-33.2	18.1	61.7	42.5	-43.1	-1444.7	-51452.1
early burt	-46.9	28.6	-50.3	24.6	-54.7	11.5	-68.1	14.1	-30.1	55.5	29.1	-23.5	13.4	-23.5	10.1	-35.2	30.5	-35.2	17.5	61.0	40.8	-44.7	-99.9	-136.1
white noise	-48.8	27.0	-54.4	23.5	-57.0	12.2	-71.9	13.9	-33.8	56.7	30.0	-27.8	9.4	-27.8	7.9	-31.5	28.1	-31.5	17.2	58.8	43.1	-42.3	-105.9	-134.7

Table S5 Log-likelihoods of the observed shell pattern parameters as explained under different models. Parameter abbreviations on the x-axis correspond to those in Table S9, with underscores removed and with the last character being the Latin equivalent of the relevant Greek symbol.

Likelihood-Ratio Tests (LRT). All of the models except for white noise contain Brownian motion as a special case and thus can be compared to Brownian motion via a LRT, which is chi-squared distributed with 1 degree of freedom. (31) The LRT fails to reject the Brownian motion model at the 0.05 significance level for 16/19 neural network parameters. When Brownian motion loses, it loses to the O-U model, which suggests that stabilizing selection is removing phylogenetic signal (Table S6).

Model	s1n	s1t	s2en	s2et	s2in	s2it	s3n	s3t	sf2ea	sf2ed	sf2id	tf2eb1	tf2ec1	tf2eb2	tf2ec2	tf2ib1	tf2ic1	tf2ib2	tf2ic2	kz	kr	turn	Ν	T
0-U	1.000	0.900	0.741	0.515	0.335	0.055	1.000	0.556	0.864	0.107	0.031	1.000	0.771	1.000	0.938	0.006	1.000	0.006	0.331	0.303	0.018	0.027	0.999	0.090
lambda	1.000	1.000	1.000	1.000	1.000	0.232	1.000	1.000	1.000	0.134	0.177	1.000	1.000	1.000	1.000	0.006	1.000	0.006	1.000	1.000	0.029	0.028	1.000	0.090
kappa	1.000	1.000	0.118	1.000	1.000	0.837	1.000	1.000	0.397	1.000	0.360	1.000	1.000	1.000	1.000	0.747	1.000	0.747	1.000	1.000	0.243	0.513	1.000	1.000
delta	0.670	0.684	0.475	0.357	0.246	0.089	1.000	0.485	0.611	0.129	0.071	0.815	0.470	0.815	0.629	0.045	0.913	0.045	0.252	0.215	0.065	0.076	1.000	1.000
early burst	0.884	1.000	1.000	1.000	1.000	1.000	0.787	1.000	1.000	1.000	1.000	0.395	1.000	0.395	1.000	1.000	0.878	1.000	1.000	1.000	1.000	1.000	0.324	1.000

Table S6. P-values of likelihood-ratio tests (chi-squared, 1 d.f.) for each model compared to the Brownian motion model. Tests that are significant at the 0.05 level are bolded. No correction for multiple testing was made here; a Bonferroni correction for 95 tests makes all results non-significant at the 0.05 level. Parameter abbreviations on the x-axis correspond to those in Table S9, with underscores removed and with the last character being the Latin equivalent of the relevant Greek symbol.



Figure S8. Akaike Information Criterion weights of 7 models for the evolution of neural network parameters. The models compared are: Brownian motion (BM, brown); the Ornstein–Uhlenbeck (OU, red) stabilizing selection model; the lambda model (orange) which rescales internal branch lengths by a linear fraction; the kappa model (green) which rescales each branch length by a power equal to the kappa parameter, and which becomes a speciational model as kappa approaches 0; delta (yellow) which focuses change towards the base or tips; early burst (EB, cyan) which has an initial high rate of change that then declines; and white noise (white), where observations are produced by a normal distribution with no tree structure, which represents the situation of no phylogenetic signal. Brownian motion (BM) has the highest AIC weight for 68% of the parameters (13/19). White noise (no phylogenetic signal) is superior for 16% (3/24) of the parameters. Parameter abbreviations on the x–axis correspond to those in Table S9, with underscores removed and with the last character being the Latin equivalent of the relevant Greek symbol.

Akaike Information Criterion. All of the models can be compared to each other at once using AIC weights (29), shown in Figure S8. Here, Brownian motion has the heaviest weight for 13/19 parameters. Typically this is a plurality rather than a majority weight, which is not surprising considering that the weight is being apportioned among seven models, the limited number of taxa limits the power to distinguish models, and the fact that the additional parameter of the non-Brownian models often converges upon parameter value that produces the Brownian model.

Non-parametric tests. Blomberg *et al.*'s (32) K is a measure of phylogenetic signal. K is the ratio of the observed MSE₀/MSE and expected MSE₀/MSE, where MSE₀ is the mean squared error between the phylogenetically correct mean and the tip data, and MSE is the mean squared error derived from the variance-covariance matrix calculated from the phylogenetic tree. The observed MSE₀/MSE is calculated from the data, and the expected MSE₀/MSE is calculated from 1000 nulls generated by reshuffling the tip data. K=1 indicates that a trait is evolving via Brownian motion and that trait data has a variance-covariance matrix that mirrors the phylogenetic structure. K<1 indicates less phylogenetic signal than expected under Brownian motion; K > 1 indicates phylogenetic overdispersion in the trait data (28).

Parameter	Blomberg's	mean null K	2.5% null	97.5% null	p-value	signifi-
s1n	1 20	0.50	0.27	1.00	0.99	sia hiah
s1t	1.10	0.48	0.25	0.87	0.99	sia hiah
s2en	1.18	0.47	0.28	0.76	1.00	sia hiah
s2et	1.17	0.48	0.25	0.91	0.98	sia hiah
s2in	0.65	0.48	0.24	0.87	0.89	e. <u>9</u> <u>9</u>
s2it	0.77	0.52	0.21	1.48	0.92	
s3n	0.89	0.48	0.26	0.85	0.99	sig high
s3t	0.79	0.48	0.26	0.91	0.95	5- 5
sf2ea	0.99	0.48	0.24	0.86	0.97	
sf2ed	0.45	0.58	0.18	3.14	0.55	
sf2id	0.55	0.49	0.23	1.06	0.75	
tf2eb1	1.01	0.48	0.25	0.85	0.95	
tf2ec1	0.83	0.48	0.24	0.93	0.98	sig_high
tf2eb2	0.86	0.48	0.25	0.88	0.94	
tf2ec2	0.94	0.48	0.25	0.89	0.96	
tf2ib1	0.49	0.55	0.18	2.91	0.64	
tf2ic1	0.84	0.48	0.27	0.86	0.99	sig_high
tf2ib2	0.55	0.60	0.18	3.30	0.76	
tf2ic2	0.56	0.48	0.25	0.84	0.76	
kz	1.11	0.50	0.26	1.05	0.95	
kr	0.46	0.48	0.24	0.85	0.55	
turn	0.52	0.48	0.26	0.79	0.67	
N	1.12	0.52	0.24	1.32	0.95	
т	0.61	0.62	0.18	3.68	0.85	

Table S7. Observed values of Blomberg's K for each neural network parameter, compared to a distribution generated under a null hypothesis of no phylogenetic signal (tip data randomly reshuffled 1000 times). The proportion of the null distribution beneath each observed K is reported; all proportions are above 0.5; 7 are significant at the 0.025 level. Parameter abbreviations as in previous tables.

The K statistics calculated for observed trait data were compared to two null hypotheses, one generating K statistics under a null hypothesis of no phylogenetic signal (tip data reshuffled 1000

times), and a second generating K statistics by simulating data under the Brownian motion fit to each trait. The proportion of the null distribution less than the observed K was calculated for each paramter. Under the no-signal null hypothesis, all observed K statistics are in the top 50% of the null distribution, indicating that the observed K is always higher than the mean of the null distribution. 13/19 observed Ks are in the top 10% of their nulls, 11/19 in the top 5%, and 7/19 in the top 2.5%. This indicates that most traits exhibit more phylogenetic signal than expected by chance (Table S7).

Comparison of the observed K statistic to K statistics generated by simulating under the best-fit Brownian model for each trait allow the detection of traits that show significantly more or less phylogenetic signal than expected under Brownian motion. Results are presented in . Only one parameter has a significantly lower K (and thus, less phylogenetic signal) than is expected under this null distribution (SF_2_e, σ_e , p=0.019), although the 95% confidence interval is quite broad and two other parameters would have significantly low Ks in a one-tailed test (Table S8).

Para- meter	Obser- ved K	sim. K mean	2.5% percen- tile	97.5% percen- tile	prop. below obs. K	signif- icance
s1n	1.20	0.999	0.4897	2.2348	0.774	
s1t	1.10	0.9993	0.4788	2.27	0.705	
s2en	1.18	0.9986	0.4751	2.2459	0.747	
s2et	1.17	0.9984	0.4667	2.169	0.726	
s2in	0.65	1.0036	0.4747	2.1932	0.179	
s2it	0.77	1.0084	0.4917	2.3036	0.353	
s3n	0.89	1.0053	0.4731	2.3132	0.517	
s3t	0.79	1.0327	0.4737	2.3383	0.383	
sf2ea	0.99	0.9955	0.4838	2.139	0.616	
sf2ed	0.45	1.0036	0.4724	2.2135	0.019	sig. low
sf2id	0.55	1.0061	0.4955	2.1998	0.075	
tf2eb1	1.01	0.9969	0.471	2.2203	0.63	
tf2ec1	0.83	1.0121	0.4958	2.3892	0.44	
tf2eb2	0.86	1.01	0.4709	2.2598	0.47	
tf2ec2	0.94	1.002	0.4689	2.179	0.567	
tf2ib1	0.49	0.9876	0.4621	2.1637	0.041	
tf2ic1	0.84	0.9892	0.4621	2.1357	0.472	
tf2ib2	0.55	1.0217	0.4701	2.2572	0.072	
tf2ic2	0.56	0.9881	0.4803	2.2479	0.09	

Table S8. Observed values of Blomberg's K for each neural network parameter, compared to a distribution generated by simulating each trait under a pure Brownian motion model, at the rate estimated by ML for each parameter. The proportion of the null distribution beneath the observed K statistic is reported; Brownian motion can be rejected in only 1 case at p<0.025 level (two-tailed non-parametric test). Parameter abbreviations as in previous tables.

As one additional check that the Brownian motion assumption was reasonable to use for PIC and ancestral state reconstruction, a variety of diagnostic plots were generated as suggested by (30) and D. Ackerly (personal communication). These confirmed that in general the data were roughly normally distributed, and that the PICs did not show significant correlations with node depth or node value.

Brownian motion models are often run on ln-transformed data, e.g. if the trait distribution is highly skewed or varies over several orders of magnitude (as can happen with e.g. body size or genome size), or if there is concern that having a lower trait boundary of 0 could violate the assumptions of Brownian motion. Therefore, all of the above tests, and subsequent ancestral state reconstruction, were also replicated on a ln-transformed version of the neural net parameters. (For a few parameters, this required setting 0 values to 10% of the minimum nonzero value for that parameter, before ln-transformation.) However, this did not produce substantial change in results or interpretation and the results of ancestral state estimation were highly correlated, as might be expected given that the raw parameter data is roughly normally distributed and typically not close to 0.



9 Ancestral State Reconstruction



Ancestral states were estimated for each continuous parameter using maximum likelihood estimation under the Brownian motion model, as implemented in the ace function of the R package APE (16). After ancestral parameter values were estimated, shell patterns were generated from them using the same model as used for the living species and plotted on the

phylogeny. Discrete characters were also estimated using ML; the color estimation was incorporated into the illustrations of the ancestors.

Results

ML estimation of continuous parameters. Examples of maximum likelihood estimations of ancestral parameter values are shown in Figure S9 and Figure S10.



Figure S10. Maximum likelihood ancestral parameter estimates for neural net parameter S_2_e, θ (excitatory neuron response function, θ parameter; abbreviated s2et).

Inspection of the estimated history of these parameters and the others indicates that many clades do show similarities in parameter values with nearby species. However, as expected, estimates for each parameter tend towards the overall (phylogenetically corrected) average as estimates are made for ancestors further and further back in time, and uncertainty increases.

ML estimation of discrete characters.

In addition to the character mapping described in the main text, other discrete characters were also mapped for comparison (Figure S11). Cone shape is fairly scattered but shows some uniformity in small clades. Strikingly, prey preference shows extremely high conservation (as was clear in the discussion of Nam *et al.*) compared to shell pattern characters. Each major clade is almost completely restricted to a certain prey, and the entire pattern is explained by the minimum possible number of transitions.

Figure S11 also shows the distributions of dots and color (black versus orange/brown) in this group and the maximum likelihood assignment of ancestral states. The presence and absence of dots and cone shape are scattered throughout the phylogeny, indicating that they are evolutionarily labile, although orange/brown color shows some correlation with large clades.



Figure S11. ML estimation of history of discrete characters. Character coding: dots 0/1 present/absent; color: 0=black and white, 1=orange/brown and white; conical: 0=rounded, 1=slightly rounded, 2=conical; food: 0=piscivorous, 1=vermivorous, 2=molluscivorous. See text for details.

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Supplement C: Fitted Parameters

In this Supplement, we analyze how to generate the shells one by one. And list their parameters. In order to do that, we first discuss the notation for the parameters. Take the parameters of *Furvus* as an example. Table S1 shows the vector notation for the parameters used in MatlabTM. In the following, we first give the parameters for shells with basic patterns, then shells with spatial pre-pattern, and finally shells with spatio-temporal pre-patterns. We use hidden networks to generate spatio-temporal pre-patterns, so we also list the parameters of the hidden network(s).

S_1_e=[1 8 0.42]	$[\gamma^{(1)}, v^{(1)}, \theta^{(1)}]$	Sensory cell's sigmoid function
$S_2_e = [1 \ 6.7 \ 0.033]$	$[\gamma_{e}^{(2)}, v_{e}^{(2)}, heta_{e}^{(2)}]$	Excitatory neuron's sigmoid function
S_2_i=[1 8.3 0.074]	$[\gamma_h^{(2)}, \ m{v}_h^{(2)}, \ m{ heta}_h^{(2)}]$	Inhibitory neuron's sigmoid function
S_3_e=[1 31 0.37]	$[\gamma^{(3)}, v^{(3)}, \theta^{(3)}]$	Secretory cell's sigmoid function
SF_2_e=[5 0.006]	$[\alpha_e, \sigma_e]$	Excitatory neuron's spatial kernel
SF_2_i=[1 0.018]	$[\alpha_h, \sigma_h]$	Inhibitory neuron's spatial kernel
$TF_2_e=[5 0.02 4 0.01]$	$[\beta_{e_1}, c_{e_1}, \overline{\beta}_{e_2}, c_{e_2}]$	Excitatory neuron's temporal kernel
TF_2_i=[1.2 0.15 0.2 0.14]	$[\beta_{h1}, c_{h1}, \beta_{h2}, c_{h2}]$	Inhibitory neuron's temporal kernel

Table S1 The notations of the parameters. We always set $\gamma^{(1)} = \gamma_e^{(2)} = \gamma_h^{(2)} = \gamma_h^{(3)} = \alpha_h = 1$, and $\beta_{e1} = \beta_{e2} + 1$, $\beta_{h1} = \beta_{h2} + 1$ in our simulations. So we have 17 free parameters.

1.1 Shells with Basic Patterns

Furvus has in-phase checkerboard pattern.

S_1_e S_2_e S_2_i S_3_e SF_2_e SF_2_i TF_2_e TF_2_e TF_2_i		[1 [1 [1 [5 [1 [5 [1.20	8. 6.7 8.3 31 0.006]; 0.018]; 0.02 0.15	0.42]; 0.033]; 0.074]; 0.37]; 4 0.20	0.01]; 0.14];
Consors 1	nas	Turing strips.			
S_1_e S_2_e S_2_i S_3_e SF_2_e SF_2_i TF_2_e TF_2_i		[1 [1 [1 [1.5 [1 [1 [1]	10 5 10 6 0.06]; 0.13]; 0 0	0.35]; 0.2]; 0.15]; 0.1]; 0	0]; 0];

Marmoreus has triangles.

S_1_e	= [1	15	0.25];
S_2_e	= [1	10	0.17];
S_2_i	= [1	20	0.35];
S_3_e	= [1	40	0.1];

$SF_2_e = [6]$	0.01];		
SF_2_i = [1	0.18];		
TF_2_e = [3.79	0.45	2.79	0.56];
$TF^{2}i = [2.47]$	0.03	1.47	0.25];

Bandanus has more and smaller triangles than *marmoreus* does. So *bandanus* has narrower spatial kernels.

S_1_e	=	[1	15	0.25];	
S_2_e	=	[1	10	0.17];	
S_2_i	=	[1	20	0.35];	
S_3_e	=	[1	40	0.1];	
SF_2_e	=	[6	0.007];		
SF_2_i	=	[1	0.12];		
TF_2_e	=	[3.79	0.45	2.79	0.56];
TF_2_i	=	[2.47	0.03	1.47	0.25];

Omaria has travelling waves.

S_1_e	=	[1	15	0.28];	
S_2_e	=	[1	14	0.17];	
S_2_i	=	[1	12.7	0.1];	
S_3_e	=	[1	25	0.1];	
SF_2_e	=	[5	0.0065]	;	
SF_2_i	=	[1	0.007];		
TF_2_e	=	[3.665	0.21	2.665	0.33];
TF_2_i	=	[1.2	0.085	0.2	0.18];

Gloriamaris has more dense travelling waves, so its spatial kernel is narrower than that of omaria.

S 1 A	= [1	15	0 281.	
b_1_c	[-	15	0.20],	
S_2_e	= [1	14	0.17];	
S_2_i	= [1	12.7	0.1];	
S_3_e	= [1	19.5	0.1];	
SF_2_e	= [4.5	0.005];		
SF_2_i	= [1	0.007];		
TF_2_e	= [3.665	0.21	2.665	0.33];
TF_2_i	= [1.2	0.085	0.2	0.18];
Pulicarius	s has dots.			
S 1 e	= [1	15	0.31:	

<u> </u>		ι-	10	0.01/	
S_2_e	=	[1	5	0.1];	
S_2_i	=	[1	5	0.06];	
S_3_e	=	[1	8	0.15];	
SF_2_e	=	[3.5	0.02];		
SF_2_i	=	[1	0.055];		
TF_2_e	=	[2.28	0	1.28	0.32];
TF_2_i	=	[1.78	0	0.78	0.32];

Arenatus has smaller and more dots than *pulicarius* does. So *arenatus* has narrower spatial kernels.

S_1_e	=	[1	15	0.3];	
S_2_e	=	[1	5	0.1];	
S_2_i	=	[1	5	0.06];	
S_3_e	=	[1	8	0.15];	
SF_2_e	=	[3.5	0.0044];	;	
SF_2_i	=	[1	0.012];		
TF_2_e	=	[2.28	0	1.28	0.32];
TF_2_i	=	[1.78	0	0.78	0.32];

Crocatus has fewer dots than pulicarius does. So crocatus has wider spatial kernels.

S_1_e	=	[1	15	0.3];	
S_2_e	=	[1	5	0.1];	
S_2_i	=	[1	5	0.06];	
S_3_e	=	[1	7.9	0.14];	
SF_2_e	=	[3.5	0.02];		
SF_2_i	=	[1	0.1];		
TF_2_e	=	[2.28	0	1.28	0.32];
TF_2_i	=	[1.78	0	0.78	0.32];

1.2 Shells with Spatial Pre-pattern

The parameter $\theta^{(3)}$, the middle point of the secretory cells' sigmoid functions, shown in this section has a spatial pre-pattern. We show its basic value as did in previous section. Moreover, we show the spatial pre-pattern of $\theta^{(3)}$.

Tessulatus's main pattern is out-phase checkerboard. But at some stripe regions, the checker has different color. At these strip regions, parameter $\theta^{(3)}$ is different.

S_1_e S_2_e S_2_i S_3_e SF_2_e SF_2_i TF_2_e TF_2_i Pre-patter	= = = = = =	$\begin{bmatrix} 1 \\ [1] \\ [1] \\ [5] \\ [1] \\ [5] \\ [1.20] \\ f \theta^{(3)} \end{bmatrix}$	8 6.7 8.3 30 0.015]; 0.029]; 0.02 0.15	0.42]; 0.033]; 0.074]; 0.4]; 4 0.20	0.01]; 0.14];		
00 0.31 0.31 0.32 0.32 0.33 0.35 0.35 0.35					÷0	- 	250

Aurisiacus's main pattern is dots. But there are also stripes. At these strip regions, parameter $\theta^{(3)}$ is different.

S_1_e S_2_e S_2_i S_3_e SF_2_e SF_2_i TF_2_e TF_2_i Pre-patter	= = = = = =	$\begin{bmatrix} 1 \\ 1 \\ 1 \\ 3.1 \\ 2.28 \\ 1.78 \\ f \theta^{(3)} \end{bmatrix}$	15 5 8 0.015]; 0.041]; 0	0.3]; 0.1]; 0.06]; 0.15]; 1.28 0.78	0.3]; 0.29];	
0.10 0.16 0.45 0.44 0.44 0.44 0.44 0.43 0.43 0.43 0.43		- VV	 		· ••	

Ammiralis's main pattern is triangles. And there are stripes. At these strip regions, parameter $\theta^{(3)}$ is different.

S_1_e	=	[1	15	0.19];	
S_2_e	=	[1	17	0.3];	
S_2_i	=	[1	3	0.54];	
S_3_e	=	[1	20	0.37];	
SF_2_e	=	[5	0.008];		
SF_2_i	=	[1	0.041];		
TF_2_e	=	[3.96	0	2.96	0.17];
TF_2_i	=	[6.82	0	5.82	0.04];

Pre-pattern of $\theta^{(3)}$:

0.4		· ·			
0.315 -					-
	1	1 (-
0.25 -					-
0.2 -	1				-
		160	190	200	250

Orbignyi's main pattern is oscillating Turing-Hopf bifurcation. But at some stripe regions, the checkers have different color. At these strip regions, parameter $\theta^{(3)}$ is different.

S_1_e	= [1	8	0.42];
S_2_e	= [1	6.7	0.033];
S_2_i	= [1	8.3	0.074];

S_3_e = [1	30	0.4];	
$SF_2_e = [5]$	0.008];	;	
SF_2_i = [1	0.015];	;	
$TF_2_e = [5]$	0.02	4	0.01];
$TF_2_i = [1.20]$	0.15	0.20	0.14];

Pre-pattern of $\theta^{(3)}$:



Laterculatus's main pattern is oscillations. But at some stripe regions, the oscillations disappear. At these strip regions, parameter $\theta^{(3)}$ is different. We may also view this pattern as oscillating Turing-Hopf bifurcation, but the parameter region is relatively small, because the Turing bifurcation in this case is not regular.

S_1_e	=	[1	6	0.3];	
S_2_e	=	[1	5	0.01];	
S_2_i	=	[1	5	0.06];	
S_3_e	=	[1	5	0.1];	
SF_2_e	=	[2.5	0.009];		
SF_2_i	=	[1	0.03];		
TF_2_e	=	[3.28	0	2.28	0.3]
TF_2_i	=	[2.58	0	1.58	0.4]

Pre-pattern of $\theta^{(3)}$:



;;

Stercusmuscarum's main pattern is oscillating Turing-Hopf bifurcation. At some stripe regions, Turing bifurcation disappears, so only oscillations remain.

S_1_e	=	[1	15	0.3];	
S_2_e	=	[1	5	0.1];	
S_2_i	=	[1	5	0.06];	
S_3_e	=	[1	22	0.03];	
SF_2_e	=	[1.3	0.0067]	;	
SF_2_i	=	[1	0.009];		
TF_2_e	=	[2.28	0	1.28	0.3];
TF_2_i	=	[1.78	0	0.78	0.29];

Pre-pattern of $\theta^{(3)}$:

0.00					1
0.020					
0.020	1				
5.024 -		1	- L		
0.02 -					-
	-			300	

1.3 Shells with Spatio-temporal Pre-pattern

The parameter $v^{(3)}$, the slope of the middle point of the secretory cells' sigmoid functions, shown in this section has a spatio-temporal prepattern. We show its basic value as did in previous section. The pre-pattern of $v^{(3)}$ is generated by hidden network(s). So we also show the parameters of the hidden network(s), and the threshold functions used to couple visible networks and hidden networks.

Episcopatus's main pattern is travelling waves. And there are some patches.

Visible network (generate travelling waves and patches):

S_1_e	= [[1	15	0.28];
S_2_e	= [[1	14	0.17];
S_2_i	= [[1	12.7	0.1];
S_3_e	= [[1	19.5	0.1];
SF_2_e	= [[4	0.0065];	
SF_2_i	= [[1	0.007];	

TF_	_2_	_e	=	[3.665	0.21	2.665	0.33];
TF	2	i	=	[1.2	0.085	0.2	0.18];

Hidden network (generate patches):

S_1_e	=	[1	15	0.3];	
S_2_e	=	[1	5	0.1];	
S_2_i	=	[1	5	0.06];	
S_3_e	=	[1	8	0.15];	
SF_2_e	=	[2.5	0.01];		
SF_2_i	=	[1	0.3];		
TF 2 e	=	[2.28	0	1.28	0.33];
TF_2_i	=	[1.78	0	0.78	0.37];

Threshold function:

*thres*₁ = 0.2, $a_1 = 0$, $b_1 = 5.7$

Aulicus's main pattern is travelling waves.

Visible network (generate travelling waves and patches):

S_1_e	=	[1	15	0.28];	
S_2_e	=	[1	14	0.17];	
S_2_i	=	[1	12.7	0.1];	
S_3_e	=	[1	19.5	0.1];	
SF_2_e	=	[4	0.006];		
SF_2_i	=	[1	0.007];		
TF_2_e	=	[3.665	0.21	2.665	0.33];
TF_2_i	=	[1.2	0.085	0.2	0.18];

Hidden network (generate patches):

S_1_e	=	[1	15	0.3];	
S_2_e	=	[1	5	0.1];	
S_2_i	=	[1	5	0.06];	
S_3_e	=	[1	8	0.15];	
SF_2_e	=	[2.5	0.01];		
SF_2_i	=	[1	0.3];		
TF_2_e	=	[2.28	0	1.28	0.33];
TF_2_i	=	[1.78	0	0.78	0.33];

Threshold function:

*thres*₁ = 0.2, $a_1 = 0$, $b_1 = 7$

Dalli's main pattern is travelling waves. There are two spatio-temporal pre-patterns, i.e. oscillations and Turing stripes. They are generated by two independent hidden networks.

Visible network (generate travelling waves, Turing stripes and oscillations):

S_1_e	=	[1	15	0.28];	
S_2_e	=	[1	14	0.17];	
S_2_i	=	[1	12.7	0.1];	
S_3_e	=	[1	19.5	0.1];	
SF_2_e	=	[7	0.006];		
SF_2_i	=	[1	0.007];		
TF_2_e	=	[3.665	0.21	2.665	0.33];
TF_2_i	=	[1.2	0.085	0.2	0.18];

Hidden network1 (generate oscillations):

S_1_e	=	[1	6	0.3];	
S_2_e	=	[1	5	0.01];	
S_2_i	=	[1	4	0.06];	
S_3_e	=	[1	3.5	0.1];	
SF_2_e	=	[2.5	0.005];		
SF_2_i	=	[1	0.2];		
TF_2_e	=	[3.28	0	2.28	0.3];
TF_2_i	=	[2.58	0	1.58	0.4];

Hidden network2 (generate Turing stripes):

S_1_e =	[1	10	0.4];	
S_2_e =	[1	5	0.2];	
S_2_i =	[1	10	0.15];	
S_3_e =	[1	6	0.1];	
$SF_2_e =$	[1.5	0.06];		
SF_2_i =	[1	0.241];		
$TF_2_e =$	[1	0	0	0];
TF_2_i =	[1	0	0	0];

Threshold functions:

*thres*₁ = 0.3, $a_1 = 0$, $b_1 = 5.5$

*thres*₂ = 0.3, $a_2 = 0$, $b_2 = 7$

Textile is similar to *dalli*.

Visible network (generate travelling waves, Turing stripes and oscillations):

S_1_e	=	[1	15	0.28];	
S_2_e	=	[1	14	0.17];	
S_2_i	=	[1	12.7	0.1];	
S_3_e	=	[1	19.5	0.1];	
SF 2 e	=	[4.5	0.006];		
SF ² i	=	[1	0.007];		
TF 2 e	=	[3.665	0.21	2.665	0.331;
TF_2_i	=	[1.2	0.085	0.2	0.18];
Hidden ne	etw	ork1 (generat	e oscillation	ns):	
S_1_e	=	[1	6	0.3];	
S_2_e	=	[1	5	0.01];	
S_2_i	=	[1	4	0.06];	
S_3_e	=	[1	3.5	0.1];	
SF_2_e	=	[2.5	0.005];		
SF_2_i	=	[1	0.2];		
TF_2_e	=	[3.28	0	2.28	0.3];
TF_2_i	=	[2.58	0	1.58	0.4];
Hidden ne	etw	ork2 (generat	e Turing str	ipes):	
S_1_e	=	[1	10	0.4];	
S_2_e	=	[1	5	0.2];	
S_2_i	=	[1	10	0.15];	
S_3_e	=	[1	6	0.1];	
SF_2_e	=	[1.5	0.06];		
SF_2_i	=	[1	0.241];		
TF_2_e	=	[1	0	0	0];
TF_2_i	=	[1	0	0	0];
Threshold	l fu	nctions:			

*thres*₁ = 0.3, $a_1 = 0, b_1 = 5.5$

 $thres_2 = 0.2, a_2 = 0, b_2 = 4.5$

1.4 Inferred Ancestral Shells' Parameters

Number 20:				
Number 20.				
S_1_e =	[1	10.799	0.35];	
S_2_e =	[1	5.936	0.085];	
S_2_i =	[1	-7.855	0.107];	
s_3_e =	[1	20.447	0.248];	
$SF_2_e =$	[4.244	0.012];		
SF_2_i =	[1	0.037];		
TF_2_e =	[3.836	0.065	2.836	0.17];
TF_2_i =	[1.621	0.081	0.621	0.195];
Number 21:				
S_1_e =	[1	12.017	0.319];	

S_2_e = S_2_i = S_3_e = SF_2_e = SF_2_i = TF_2_e = TF_2_i =	[1 [1 [3.915 [1 [3.33 [1.804	9.377 5.901 20.314 0.014]; 0.047]; 0.084 0.051	0.108]; 0.121]; 0.181]; 2.33 0.804	0.24]; 0.219];
Number 22: $S_1_e =$ $S_2_e =$ $S_2_i =$ $S_3_e =$ $SF_2_e =$ $SF_2_i =$ $TF_2_e =$ $TF_2_i =$ $TF_2_i =$	[1 [1 [1 [4.304 [1 [3.583 [1.821	12.236 10.675 8.032 22.89 0.013]; 0.048]; 0.113 0.062	0.318]; 0.115]; 0.137]; 0.193]; 2.583 0.821	0.247]; 0.206];
Number 23: $S_1 = =$ $S_2 = =$ $S_2 = =$ $S_3 = =$ $SF_2 = =$ $SF_2 = =$ $TF_2 = =$ $TF_2 = =$ $TF_2 = =$	[1 [1 [1 [4.551 [1 [3.643 [1.9	12.856 10.488 9.367 24.307 0.011]; 0.051]; 0.147 0.063	0.307]; 0.129]; 0.157]; 0.182]; 2.643 0.9	0.277]; 0.202];
Number 24: $S_1_e =$ $S_2_e =$ $S_2_i =$ $S_3_e =$ $SF_2_e =$ $SF_2_i =$ $TF_2_e =$ $TF_2_i =$ $TF_2_i =$	[1 [1 [1 [4.644 [1 [3.702 [1.909	13.354 12.035 7.041 22.283 0.009]; 0.037]; 0.141 0.069	0.3]; 0.142]; 0.151]; 0.18]; 2.702 0.909	0.269]; 0.187];
Number 25: S_1_e = S_2_e = S_2_i = S_3_e = SF_2_e = SF_2_i = TF_2_e = TF_2_i =	[1 [1 [1 [4.848 [1 [3.717 [2.193	14.026 12.541 2.056 20.813 0.008]; 0.028]; 0.148 0.067	0.283]; 0.167]; 0.175]; 0.175]; 2.717 1.193	0.277]; 0.17];
Number 26: $S_1_e =$ $S_2_e =$ $S_2_i =$ $SF_2_e =$ $SF_2_e =$ $SF_2_i =$ $TF_2_e =$ $TF_2_i =$ $TF_2_i =$	[1 [1 [1 [5.102 [1 [3.741 [2.651]	14.425 13.657 4.523 20.985 0.008]; 0.025]; 0.141 0.061	0.268]; 0.188]; 0.212]; 0.186]; 2.741 1.651	0.274]; 0.153];
Number 27: $S_1_e =$ $S_2_e =$ $S_2_i =$ $S_3_e =$ $SF_2_e =$ $SF_2_i =$ $TF_2_e =$ $TF_2_i =$ $TF_2_i =$	[1 [1 [5.322 [1 [3.715 [2.158]	14.62 13.074 9.359 19.664 0.007]; 0.019]; 0.164 0.069	0.272]; 0.182]; 0.174]; 0.157]; 2.715 1.158	0.293]; 0.162];

Number 27 is assumed to have two hidden networks as its decedents do. We use phylogenetically independent contrasts method to infer the parameters of the two hidden networks.

Hidden network1:

$S_1_e = S_2_e = S_2_i = S_3_e = SF_2_e = SF_2_i = SF_2_i = SF_2_i = TF_2_i = TF_2_i = TF_2_i = Hidden networks and the set of the $	[1 [1 [1 [2.5 [1 [3.28 [2.58 work2:	6 5 4 3.5 0.005]; 0.2]; 0	0.3]; 0.01]; 0.06]; 0.1]; 2.28 1.58	0.3]; 0.4];
$S_1_e = S_2_e = S_2_i = S_3_e = SF_2_e = SF_2_i = SF_2_i = TF_2_i = TF_2_$	[1 [1 [1 [1.5 [1 [1 [1]	10 5 10 6 0.06]; 0.241]; 0 0	0.4]; 0.2]; 0.15]; 0.1]; 0	0]; 0];

Threshold functions:

thres₁ = 0.3,
$$a_1 = 0$$
, $b_1 = 5.5$
thres₂ = 0.25, $a_2 = 0$, $b_2 = 5.75$

Number 28:			
S_1_e = [1	13.185	5 0.31];	
$S_2 = [1]$	11.499	0.132];	
$S_2_i = [1]$	12.518	0 . 125];	
$S_3 = [1]$	21.853	3 0 . 183];	
$\overline{SF}_2 = [4]$.546 0.009	;	
$SF_2_i = [1]$	0.034	;	
$TF_2 = [3]$.738 0.13	2.738	0.255];
$TF_2_i = [1]$.675 0.077	0.675	0.19];
Number 29:	14 01/		
$S_1_e = [1]$	14.012	0.299];	
$S_2_e = [1]$	11.978	3 0.141];	
$S_2_i = [1]$	14.718	3 0.108];	
$S_3_e = [1]$	18.892	2 0.15];	
$SF_2_e = [4]$.359 0.01];		
$SF_2_i = [1]$	0.033	;	
$TF_2_e = [3]$.532 0.14	2.532	0.288];
$TF_2_i = [1]$.531 0.07	0.531	0.203];
Number 30:			
S 1 ρ = [1	14 08	0 2981.	
$5_1 = [1]$	11 781	0.141.	
$5_2_c = [1]$	15 005	0 1061.	
$S_2 = [1]$	18 560	0 1481	
$SE^{2} = [1]$	352 0 011	, 0.140],	
$SF_2 i = [1]$	0 035	,	
$TF_2 = [3]$.493 0.138	2.493	0.291.
$TT_2_c = [3]$	528 0.068	0 528	0 2061.
11_2_1 [1	.520 0.000	0.520	0.200],
Number 31:			
S_1_e = [1	14.417	0.291];	
S_2_e = [1	11.844	0.153];	
$S_2_i = [1]$	10.032	2 0.105];	
S_3_e = [1	18.977	0.13];	
$SF_2_e = [4]$.212 0.008	;	
$SF_2_i = [1]$	0.022	;	

TF_	_2_	_e	=	[3.587	0.169	2.587	0.305];
TF	2	i	=	[1.395	0.076	0.395	0.194];

Number 31 is assumed to have one hidden network.

Hidden network (generate patches):

S_1_e	=	[1	15	0.3];	
S_2_e	=	[1	5	0.1];	
S_2_i	=	[1	5	0.06];	
S_3_e	=	[1	8	0.15];	
SF_2_e	=	[2.5	0.01];		
SF_2_i	=	[1	0.3];		
TF_2_e	=	[2.28	0	1.28	0.33];
TF_2_i	=	[1.78	0	0.78	0.35];

Threshold function:

*thres*₁ =
$$0.2, a_1 = 0, b_1 = 6.35$$

Number 32: $S_1 = =$ $S_2 = =$	[1 [1 [1 [5.88 [1 [3.778 [2.423	14.823 9.807 19.971 38.701 0.009]; 0.142]; 0.425 0.033	0.255]; 0.167]; 0.334]; 0.107]; 2.778 1.423	0.537]; 0.246];
Number 33: $S_1_e =$ $S_2_e =$ $S_2_i =$ $S_3_e =$ $SF_2_e =$ $SF_2_i =$ $TF_2_e =$ $TF_2_i =$ $TF_2_i =$	[1 [1 [1 [3.519 [1 [3.034 [1.813	11.98 8.451 5.623 18.181 0.016]; 0.047]; 0.06 0.036	0.316]; 0.105]; 0.108]; 0.161]; 2.034 0.813	0.243]; 0.235];
Number 34: $S_1 = =$ $S_2 = =$ $S_2 = =$ $S_3 = =$ $SF_2 = =$ $SF_2 = =$ $TF_2 = =$ $TF_2 = =$ $TF_2 = =$	[1 [1 [1 [3.401 [1 [2.96 [1.895	11.711 8.779 7.91 16.032 0.014]; 0.043]; 0.045 0.027	0.312]; 0.093]; 0.096]; 0.153]; 1.96 0.895	0.26]; 0.265];
Number 35: $S_1_e =$ $S_2_e =$ $S_2_i =$ $S_3_e =$ $SF_2_e =$ $SF_2_i =$ $TF_2_e =$ $TF_2_i =$ $TF_2_i =$	[1 [1 [1 [3.481 [1 [2.413 [1.803	14.359 7.307 5.719 9.967 0.013]; 0.035]; 0.009 0.005	0.302]; 0.099]; 0.067]; 0.151]; 1.413 0.803	0.308]; 0.309];
Number 36: $S_1_e = S_2_e = S_2_i = S_3_e = SF_2_e = SF_2_i = SF_2_i = TF_2_e = TF_2_e = TF_2_i = TF$	[1 [1 [1 [2.572 [1 [2.287 [1.611	12.609 4.275 1.819 12.908 0.024]; 0.028]; 0.024 0.015	0.318]; 0.125]; 0.101]; 0.121]; 1.287 0.611	0.207]; 0.2];
Number 37: s_1_e =	[1	13.691	0.31];	

S_2_e =	[1	3.745	0.114];	
S_2_i =	[1	8.045	0.082];	
S_3_e =	[1	13.409	0.107];	
SF_2_e =	[2.404	0.018];		
SF_2_i =	[1	0.043];		
TF_2_e =	[2.284	0.013	1.284	0.249];
TF_2_i =	[1.688	0.008	0.688	0.241];