Evolution of patterns on Conus shells

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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.

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. 1C, 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. 1C.

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

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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 $P = 0.0001$ for 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. aurissiacus, 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 ban- danus and Conus marmoratus 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.
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. furus*, *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. 6A and B show 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 networks) 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 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.
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-
ies. 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 to an absolute time scale. A more formal analysis would begin with the raw DNA sequence alignment and fossil calibration to an absolute time scale. A more formal analysis would begin with the raw DNA sequence alignment and fossil calibration to an absolute time scale. A more formal analysis would begin with the raw DNA sequence alignment and fossil calibration to an absolute time scale.

We found stronger 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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