How cryptic is cryptic diversity? Machine learning approaches to classifying morphological variation in *Emys marmorata* (Testudinoidea, Emydidae).

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December 1, 2014

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Abstract

2

Introduction

MATERIALS AND METHODS

Specimens and sampling

- ⁶ We collected landmark-based morphometric data from SAMPLE adult *E. marmorata* museum specimens. These specimens include both newly sampled individuals and those sampled in
- previous studies of plastral shape variation (Angielczyk and Feldman 2013; Angielczyk et al. 2011; Angielczyk and Sheets 2007).
- Specimen classification was based on known specimen geographic information which was recorded from museum collection information. When precise latitude and longitude information
- was not available it was estimated from whatever locality information was present. Because the specimens used to define the subclades in Spinks and Shaffer (2005) and Spinks et al. (2010)
- were not available for study, all specimen classifications were based solely on this geographic

information and not from explicit assignment in previous studies. Instead, classification was based on matching museum locality data with the geographic boundaries of the molecularly-defined clades of Spinks and Shaffer (2005) and Spinks et al. (2010). Because the exact barriers between different biogeographic regions are unknown and unclear, two assignments for both the morphologically and molecularly based hypotheses were used. Each morphologically based hypothesis had three classes, while each molecular-based had four classes. In total, each specimen was given four different classifications.

Geometric morphometrics

22

Following previous work on plastral variation (Angielczyk and Feldman 2013; Angielczyk et al. 2011; Angielczyk and Sheets 2007), 19 landmarks were digitized using TpsDig 2.04 (Rohlf 2005). These landmarks were chosen to maximize the description of general plastral variation(Fig. 1). 17 of these landmarks are at the endpoints or intersection of the keratinous plastral scutes that cover the platron. 12 of these landmarks were chosen to be symmetrical across the axis of symmetry and, in order to prevent degrees of freedom and other concerns (Klingenberg et al. 2002), prior to analysis these landmarks were reflected across the axis of symmetry (i.e. midline) and the average position of each symmetrical pair was used. In cases where damage or incompleteness prevented symmetric landmarks from being determined, only the single member of the pair was used. Analysis was conducted on the resulting "half" plastra. Plastral landmark configurations were superimposed using generalized Procrustes analysis (Dryden and Mardia 1998) after which, the principal components (PC) of shape were calculated. This was done using the shapes package for R (Dryden 2013; R Core Team 2013).

Machine learning analyses

- Unsupervised learning.— In order to preserve the relationship between all landmark configurations in shape space, the dissimilarity between observations was measured using Kendall's
- Riemanninan shape distance or ρ (Dryden and Mardia 1998; Kendall 1984). This metric was chosen because shape space, or the set of all possible shape configurations following
- Procrustes superimposition, is a Riemannian manifold and thus non-Euclidean (Dryden and Mardia 1998). ρ varies between 0 and $\pi/2$ when there is no reflection invariance, which should
- 44 not be a concern in the case of the half plastral landmark configurations used in the study.
- The ρ dissimilarity matrix was divisively clustered using partitioning around mediods cluster-
- ing (PAM), a method similar to k-means clustering except that instead of minimizing the sum of squared Euclidean distances between observations and centroids, the sum of squared
- dissimilarities between observations and mediods is minimized (Kaufman and Rousseeuw 1990). Because the optimal number of clusters of shape configurations in the study was
- unknown, being possibly three, four, or some other value, clustering solutions were estimated with the number of clusters varied between one and NUMBER. Clustering solutions were

compared using the gap statistic, which is a measure of goodness of clustering (Tibshirani et al. 2001). The gap statistic is defined

$$Gap_n(k) = E_n^*[\log(W_k)] - \log(W_k)$$

where W_k is

$$W_k = \sum_{r=1}^{k} \frac{1}{2n_r} (\sum_{i,i' \in C_r} d_{ii'})$$

.

- $d_{ii'}$ is the dispersion of the clustering solution or the sum of the pairwise dissimilarities between observations in each cluster and their respective mediods (C) for all clusters r. This
- value is averaged and compared to the expected dispersion (E_n^*) of a sample n from a reference distribution. In this case, the reference distribution was estimated from NUMBER resamples.
- 60 This analysis was conducted using the cluster package for R (Maechler et al. 2013).
- Supervised learning.— Three different supervised learning, or classification, approaches were used: linear discriminate analysis, multinomial logistic regression, and random forests. Linear discriminate analysis, also known as canonical variate analysis, is commonly used in studies
- of geometric morphometric data (Mitteroecker and Bookstein 2011; Zelditch et al. 2004). The other two methods, however, are not. Each of these three methods has a different
- interpretation and reveal very different aspects of the data. In all cases, the optimal number of PCs used as predictors was chosen via max within-sample AUC value, explained below.
- Linear discriminate analysis (LDA) attempts to find a linear combination of predictors to best model two or more classes. LDA is very similar to PCA except that instead of finding
- the linear combination of features that maximize the amount of explained variance in the data, LDA maximizes the differences between classes. The results of this analysis produces a
- transformation matrix by which the original features can be transformed to reflect the best discrimination between the classes. In this analysis, LDA was applied on the eigenscores from
- a subset of the total number of PCs, ranging from two to NUMBER in increasing order of complexity. In total, this produced nine different LDA scaling matrices.
- Multinomial logistic regression is an extension of logistic regression, where instead of a binary response there are three or more response classes (Venables and Ripley 2002). Similar to
- the odds ratios calculated from the coefficients of a logistic regression, the relative risk of a classification can be determined from the coefficients of the model. Similar to LDA, the
- optimal number of PCs as predictors was determined by comparing within-sample AUC values across multiple models.
- Random forest models are an extension of classification and regression trees (CART) (Breiman 2001; Breiman et al. 1984). The goal of CARTs are to use a series of different features to

- estimate the class of an observation. In top-down induction of decision trees for each member of a given set of predictor variables, attribute value test are used to estimate the differences
- between classes. This process, called recursive partitioning, is then repeated on each subset.

 The recursion continues until the resulting observations all share the same class or no
- more meaningful partitions are possible. The resulting model is a tree structure by which observations are classified at each intersection via the estimated cutoff points from the
- ⁹⁰ attribute tests made during model fitting.

In a random forest model, many CARTs are built from a random subsample of both the features and the observations. This process is then repeated many times and the parameters of the final model was chosen as the mode of estimates from the distribution of CARTs

- 94 (Breiman 2001). In addition to classifying the observations, this procedure allows for the features to be ranked in order of importance. This is a generally important property that
- is useful for many other studies which want to describe and model the differences between classes and the relative importance of different predictors.
- In this analysis, random forest model parameters were estimated from 1000 subtrees. The best set of predictors necessary for each classification scheme was estimated using a recursive feature selection algorithm was used to choose the optimal number of PCs to include based on the AUC of the model. Following the backwards selection algorithm implemented in caret (Kuhn and Johnson 2013), the maximum number of features were included in the initial model, their importance ranked, and the AUC of the model calculated. The lowest ranked feature was then removed, and the AUC of the model recalculated. This was repeated until only one feature, remained. Because PCs were kept in order of importance and not in relation to the amount of variance each PC described, these means that the PCs are not included in assending eigenvalue.
- In classification studies, like this one, a common metric of performance is area under the receiver operating characteristic curve (AUC). AUC is an estimate of the relationship between the false positive and true positive rates, as opposed to just the true positive rate (accuracy). This relationship is especially useful in cases where misclassification needs to be minimized just as much as accurate classification, as in this study. AUC ranges between 0.5 and 1, with 0.5 indicating classification no better than random and 1 indicating perfect classification (Hastie et al. 2009). AUC was used for both LDA and random forest models while multinomail logistic model selection was done using the likelihood based AICc (Burnham and Anderson 2002).

The standard AUC calculation is defined for of binary classification, however in this application there are multiple categories. The alternative calculation used here follows an all-against-one strategy where the individual AUC values for each class versus all others are averaged to produce a multiclass AUC (Hand and Till 2001).

The ultimate measure of model fit is accurately predicting the values of unobserved samples (Hastie et al. 2009; Kuhn and Johnson 2013). Within-sample performance is inherently biased upwards, so model evaluation requires overcoming this bias. With very large sample sizes,

- as in this study, part of the sample can be used as the "training set" and the remainder acts as the "testing set." The former is used for fitting the model while the later is used for measuring model performance. This is called model generalization. In this analysis, 75% of samples were used as the training set while the remaining 25% were used as the testing set.
- In order to estimate confidence intervals on the out-of-sample AUC values, a nonparametric bootstrap was performed where the true and estimated classifications were resampled with replacement. This was done 1000 times.

It is common for some out of sample observations to be misclassified. This misclassification
may be due to the model not accurately representing shape variance, systematic differences
between the training and test sets, or systematic differences between the accurately and
inaccurately classified samples. Testing and training sets are determined completely at random
within each class and with respect to shape. Results were not effected by changes in testing
or training set assignment.

To determine if there were systematic differences between the correctly and incorrectly classified samples, the multivariate centroids of the correct and incorrect groups were compared to what would be expected by random chance. The group labels were permuted and the difference between the new centroids was calculated. This was repeated 1000 times. The number of permutations less than the empirical difference were counted and divided by 1000, giving a p-value. Significant results indicate that correctly and incorrectly classified specimens are systematically different. This was done only when there were 10 or of that

RESULTS

144

Unsupervised learning

Comparison of gap statistic values from PAM clustering show that the best number of clusters is 1. This means that there is no means of naturally partitioning plastron shape into distinct subgroups.

Supervised learning

- For multinomial logistic regression, the best number of predictors was always the first four PCs. This was not the case for either of the LDA or random forest approaches which varied from two to six different PCs. Note that the best random forest models were determined via recursive feature selection, so PCs were not included in order of percent variance explained.
- For both the LDA and multinomial logistic regression models, increasing model complexity increased cumulative percent variation necessary to best model the differing classification schemes FIGURE OF CUMMULATIVE. The high variance in optimal number of PCs used as predictors between schemes may represent either ease of fitting that particular model or that the data was too noising to accurately fit any model beyond the simplest.

As part of fitting a random forest model, a ranking of variable importance is also determined. Interestingly, the the order of variable importance is not the same as the order of principal components. This means that the principal components that best describe the differences between the various classes are not aligned with the principal components which describe the largest amount of variance. This result would be the case if variation between classes was extremely fine grained and not a part of the principal function or form of the plastron, which makes sense given that the plastron is involved in both protection and aquadynamics and not mate choice (Germano and Bury 2009; Holland 1992; Lubcke and Wilson 2007; Rivera 2008).

The observed AUC values for all of the optimal models are very low, values near 0.5 indicate that a model is no better than completely random assignment TABLE OR FIGURE. This means that in very few cases are any of the proposed classification schemes are generally poor descriptors of the observed variation. It appears that the data set is overwhelmed by noise, making any accurate classifications difficult at best. This observation is cemented with the generalizations of the models to the testing data set.

AUC values from model generalization, or estimating testing data set membership, does not indicate a clear "best" classification scheme (Fig. 2). While the scheme with two species has the greatest AUC point estimate for each modeling approach, this scheme is not significantly greater from any other except in some limited cases (e.g. LDA).

Mean AUC values for all model generalizations are, as expected, less than from the training set. This is expected because the low initial AUC values indicate little classification ability compared random, and the generalization to new data would only lead to loss in accuracy.

Differences in mean shape between correctly and incorrectly classified observations from test set were frequently statistically significant, though there are exceptions. The frequency of these results, however, is important because this means that the different models are poor predictors of class membership. This may be because differences in plastron shape do not align with the any of the hypothesized classification schemes.

DISCUSSION

Generalization and the unsupervised learning results are surprisingly congruous.

Plastron shape indicates no optimal classification scheme. A possible explanation for this that the genetic divergence associated with (sub)speciation is either not based on plastron morphology or local selective pressures due to hydrological regime overwhelming any possible morphological divergence.

BIBLIOGRAPHY

- Angielczyk, K. D. and C. R. Feldman. 2013. Are diminutive turtles miniaturized? The ontogeny of plastron shape in emydine turtles. Biological Journal of the Linnean Society 108:727–755.
- Angielczyk, K. D., C. R. Feldman, and G. R. Miller. 2011. Adaptive evolution of plastron shape in emydine turtles. Evolution 65:377–394.
- Angielczyk, K. D. and H. D. Sheets. 2007. Investigation of simulated tectonic deformation in fossils using geometric morphometrics. Paleobiology 33:125–148.
 - Breiman, L. 2001. Random Forests. Machine Learning 45:5–32.
- Breiman, L., J. Friedman, C. J. Stone, and R. A. Olshen. 1984. Classification and regression trees. Wadsworth International Group, Belmont.
- Burnham, K. P. and D. R. Anderson. 2002. Model selection and multi-model inference: a practical information-theoretic approach. 2nd ed. Springer, New York.
- Dryden, I. L. 2013. shapes: Statistical shape analysis. R package version 1.1-8.
 - Dryden, I. L. and K. Y. Mardia. 1998. Statistical shape analysis. Wiley, New York.
- Germano, D. J. and R. B. Bury. 2009. Variation in body size, growth, and population structure of Actinemys marmorata from lentic and lotic habitats in Southern Oregon. Journal of Herpetology 43:510–520.
- Hand, D. J. and R. J. Till. 2001. A Simple Generalisation of the Area Under the ROC Curve for Multiple Class Classification Problems. Machine Learning 45:171–186.
- Hastie, T., R. Tibshirani, and J. Friedman. 2009. The elements of statistical learning: data mining, inference, and prediction. 2nd ed. Springer, New York.
- Holland, D. C. 1992. Level and pattern in morphological variation: a phylogeographic study of the western pond turtle (Clemmys marmorata). Ph.D. thesis University of Southwestern Louisiana.
- Kaufman, L. and P. J. Rousseeuw. 1990. Finding groups in data: an introduction to cluster analysis. Wiley, New York.
- Kendall, D. G. 1984. Shape Manifolds, Procrustean Metrics, and Complex Projective Spaces. Bulletin of the London Mathematical Society 16:81–121.
- Klingenberg, C. P., M. Barluenga, and A. Meyer. 2002. Shape analysis of symetric structures: quantifying variation among individuals and asymmetry. Evolution 56:1909–1920.
- 222 Kuhn, M. and K. Johnson. 2013. Applied predictive modeling. Springer, New York, NY.

- Lubcke, G. M. and D. S. Wilson. 2007. Variation in shell morphology of the Western Pond
 Turtle (Actinemys marmorata Baird and Giarard) from three aquativ habitats in Northern
 California. Journal of Herpetology 41:107–114.
- Maechler, M., P. Rousseeuw, A. Struyf, M. Hubert, and K. Hornik. 2013. cluster: Cluster Analysis Basics and Extensions. R package version 1.14.4.
- ²²⁸ Mitteroecker, P. and F. Bookstein. 2011. Linear Discrimination, Ordination, and the Visualization of Selection Gradients in Modern Morphometrics. Evolutionary Biology 38:100–114.
- R Core Team. 2013. R: A Language and Environment for Statistical Computing. R Foundation for Statistical Computing Vienna, Austria.
- Rivera, G. 2008. Ecomorphological variation in shell shape of the freshwater turtle Pseudemys concinna inhabiting different aquatic flow regimes. Integrative and comparative biology 48:769–87.
 - Rohlf, F. J. 2005. TpsDig 2.04.
- Spinks, P. Q. and H. B. Shaffer. 2005. Range-wide molecular analysis of the western pond turtle (Emys marmorata): cryptic variation, isolation by distance, and their conservation implications. Molecular ecology 14:2047–64.
- Spinks, P. Q., R. C. Thomson, and H. B. Shaffer. 2010. Nuclear gene phylogeography reveals
 the historical legacy of an ancient inland sea on lineages of the western pond turtle, Emys
 marmorata in California. Molecular ecology 19:542–56.
- Tibshirani, R., G. Walther, and T. Hastie. 2001. Estimating the number of clusters in a data set via the gap statistic. Journal of the Royal Statistical Society: Series B (Statistical Methodology) 63:411–423.
- Venables, W. and B. D. Ripley. 2002. Modern applied statistics with S. 4th ed. Springer, New York.
- Zelditch, M. L., D. L. Swiderski, and H. D. Sheets. 2004. Geometric morphometrics for biologists: a primer. Elsevier Academic Press, Amsterdam.

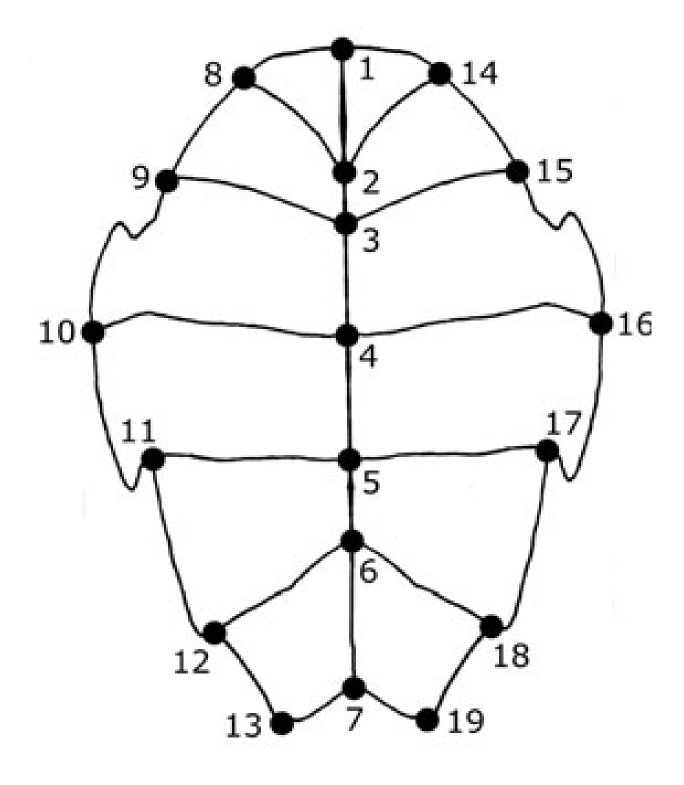


Figure 1: Depiction of general plastral shape of E. marmorata and position of the 19 landmark used in this study. Anterior is towards the top of the figure.

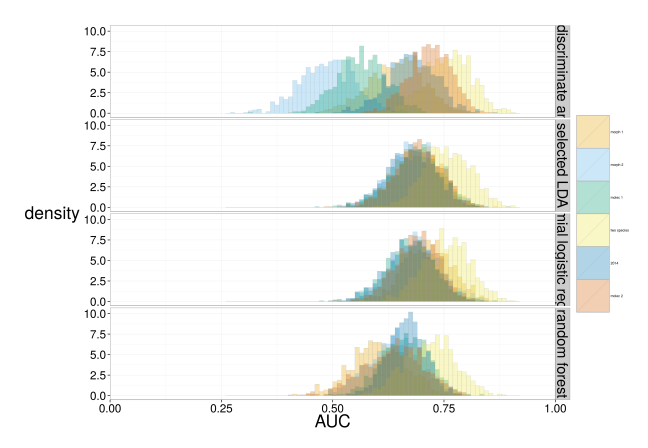


Figure 2: Bootstrap distributions for genearlized AUC values for each of the classification schemes. Each row corresponds to a different modeling approach: LDA, LDA using best variables from random forest, multinomial logistic regression, and random forest. Each distribution corresponds to 1000 bootstrap replicates.