Rapid species recognition favors greater avian-perceived plumage dichromatism in true thrushes (genus: *Turdus*)

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# Abstract

## Keywords

*dichromatism*, *plumage*, *species recognition*

# Background

# Methods

## *Plumage sexual dichromatism*

A total of N=77 *Turdus* thrush species (approximately ~89% of all known true thrush species) were sampled for plumage spectral reflectance using prepared bird skin specimens at the American Museum of Natural History in New York City and the Field Museum in Chicago. Reflectance measurements spanning 300-700nm were taken in triplicate from the belly, breast, throat, crown and mantle plumage patches [[1](#ref-andersson2006)] of each individual. N=3 male and N=3 female individuals were measured for most species (exceptions: *T. lawrencii*, N=2 males and N=2 females; *T. swalesi*, N=1 male and N=1 female). Reflectance spectra were measured using a 400 μm fiber optic reflection probe fitted with a rubber stopper to maintain a consistent measuring distance of 3 mm and area of 2 mm2 at a 90° angle to the surface of the feather patch. Measurements were taken using a JAZ spectrometer with a pulsed-xenon light source (Ocean Optics, Dunedin, USA) and we used a diffuse 99% reflectance white standard (Spectralon WS-1-SL, Labsphere, North Sutton NH, USA).

We applieda receptor-noise limited visual model [[2](#ref-vorobyev1998)] of the European Blackbird (*T. merula*) visual system [[3](#ref-hart2000)] in the *pavo* [[4](#ref-maia2019)]⁠ package in R v4.0.0 [[5](#ref-rcoreteam2020)]⁠ to calculate avian-perceived chromatic and achromatic visual contrast (in units of “Just-Noticeable Differences”,or JNDs) of male vs. female plumage patches for all sampled *Turdus* species. Chromatic and achromatic JNDs were calculated for male-female pairs within each species (i.e., N=9 JND values calculated per patch for each species where N=3 males and N=3 females sampled), and then JND values were averaged for each species’ respective plumage patches. Under ideal laboratory conditions, 1 JND is generally considered to be the discriminable threshold past which an observer is predicted to be able to perceive the two colors as different. However, natural light environments vary both spatially and temporally [[6](#ref-endler1993)]⁠, bringing into question the accuracy of a 1 JND threshold for generalizing visual contrast under natural conditions. Therefore, we calculated the total number of sexually-dichromatic plumage patches per species (out of N=5 measured patches) as the number of plumage patches with average JND values > 1, 2, or 3 to account for uncertainty in visual discrimination thresholds due to variation in psychophysical and ambient lighting conditions affecting the strength of between-sex plumage visual contrast [[7](#ref-kemp2015)]⁠.

## *Life History Data*

### *Breeding Timing Model*

We collected data on migration behavior and breeding season length from *Thrushes* [[8](#ref-clement2000)] and the *Handbook of the Birds of the World* [[9](#ref-delhoyo2017)]⁠. We assigned three different kinds of migratory behavior: 1) *full migration* when a species description clearly stated that a species “migrates”, 2) *partial migration* when a species was described to have “altitudinal migration”, “latitudinal migration” or “movement during non-breeding season”, or 3) *sedentary* when when a species was described as “resident” or “sedentary”. Breeding season length was defined as the number of months the species breeds each year.

### *Breeding Sympatry Model*

Species’ breeding ranges were acquired from *BirdLife International* [[10](#X6c896e2b80dd0fca0e8ee32fce3f4251147131f)]⁠. We calculated congener breeding range overlaps (as percentages) using the *letsR* package in R [[11](#ref-vilela2015)]⁠. We then calculated the number of sympatric species as the number of congeners with breeding ranges that overlap >30% with the focal species’ breeding range [[12](#ref-cooney2017)].

### *Breeding Spacing Model*

Species’ breeding range sizes (in km2) were acquired using the *BirdLife International* breeding range maps. Species’ island vs. mainland residence was also determined using breeding ranges from *BirdLife International*. Mainland residence was assigned if the species had a breeding range on any continent and Japan. Island residence was assigned to species having a breeding range limited to a non-continental landmass entirely surrounded by an oceanic body of water.

## *Statistical Modeling*

We used phylogenetically-corrected Bayesian multilevel logistic regression models using the *brms* v2.13.0 package [[13](#ref-burkner2017)] in R v4.0.0 [[5](#ref-rcoreteam2020)]⁠ where responses, the number of sexually-dichromatic patches >1, 2, and 3 chromatic and achromatic JNDs, were modeled as binomial trials (N=5 plumage patch “trials”) to test for associations with breeding timing, breeding sympatry and breeding spacing. For all phylogenetically-corrected models, we used the *Turdus* phylogeny from Nylander et al. (2008) [[14](#ref-nylander2008)]to create a covariance matrix of species’ phylogenetic relationships. All models used a dataset of N=67 out of the *Turdus* species for which all the types of data (see above) were available.

Our *breeding timing* models included the following predictors: z-scores of breeding season length (mean centered and divided by one standard deviation), migratory behavior (full migration as the reference category versus partial migration or sedentary), and their interaction. *Breeding sympatry* models included the number of sympatric species with greater than 30% breeding range overlap as the only predictor of the number of sexually-dichromatic plumage patches. *Breeding spacing* models included transformed breeding range size (km2) and breeding landmass (mainland as the reference category versus island). We also ran null models (intercept only) for all responses. All models’ intercepts and response standard deviations were assigned a weak prior (Student T: df = 3, location = 0, scale = 10), and predictor coefficients were assigned flat priors. We ran each model for 6,000 iterations across 6 chains and assessed Markov Chain Monte Carlo (MCMC) convergence using the Gelman-Rubin diagnostic (Rhat) [[15](#ref-gelman2013)]. We then performed k-fold cross-validation [[16](#ref-vehtari2017)] to refit each model *K*=16 times. For each k-fold, the training dataset included a randomly selected set of or N≈63 species, and the testing dataset included or N≈4 species not included in the training dataset. Finally, we compared differences between the models’ expected log pointwise predictive densities (ELPD) to assess which model(s) best predicted the number of sexually-dichromatic plumage patches [[16](#ref-vehtari2017)]⁠.

# Results

We obtained N ≥ 4000 effective samples for each model parameter and all models’ Markov Chains (MCMC) successfully converged (Rhat = 1 for all models’ parameters) (Supplementary Figure). All *breeding sympatry*, *breeding timing*, and *breeding spacing* models performed similarly well and substantially better than *intercept only* models in predicting the number of sexually dimorphic plumage patches with achromatic JND values > 1, 2, or 3 (Table 1; all models predicting achromatic plumage patches had ELPD values within 4, following the convention of Burnham and Anderson (2002)[[17](#ref-burnham2002)]). Among models predicting the number of sexually-dichromatic plumage patches with chromatic JND values >1, 2, or 3, all models performed much better than *intercept only* models, and *breeding sympatry* models had the best predictive performance (Table 1; *breeding sympatry* models all have ELPD =0, only the *breeding spacing* models predicting dichromatic plumage patches with had similar predictive performance).

# Discussion

# Conclusions

# Acknowledgements

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