

How much are wild vertebrate populations evolving right now?

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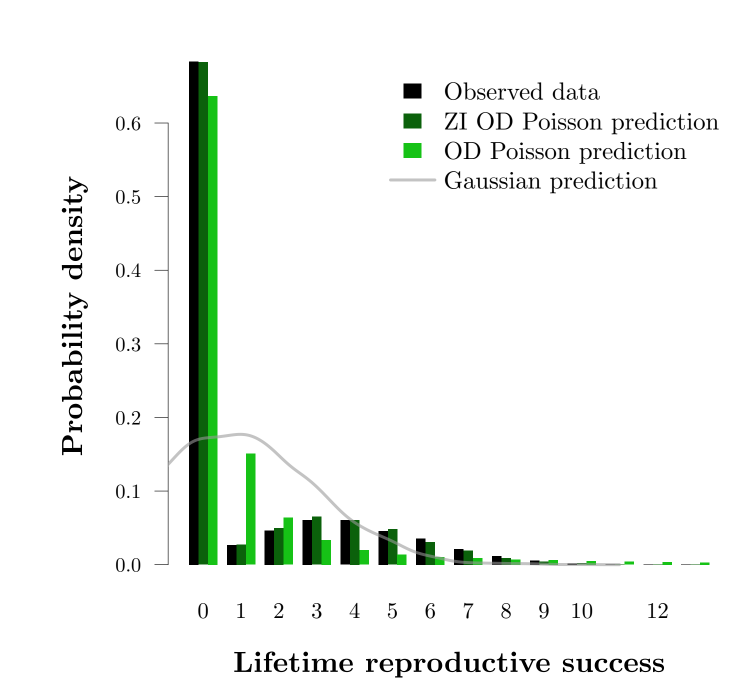


THE BIG PROBLEM: We do not know how much wild organisms are currently evolving!

Fisher's fundamental theorem of natural selection states that additive genetic variation in fitness measures evolution across all traits and all the genome. That is just what we need*! Yet, there are few estimates in free-ranging populations, and most may be unreliable. Indeed, it is difficult to measure fitness, difficult to estimate genetic variance, statistical models tend not to fit the data, and it is unclear how to interpret estimates from generalized linear models. We use a dozen long-term monitored populations with high quality fitness (measured as lifetime breeding success) and relatedness measurements to tackle these issues. How to make the most of these data? Are birds and mammals currently evolving?

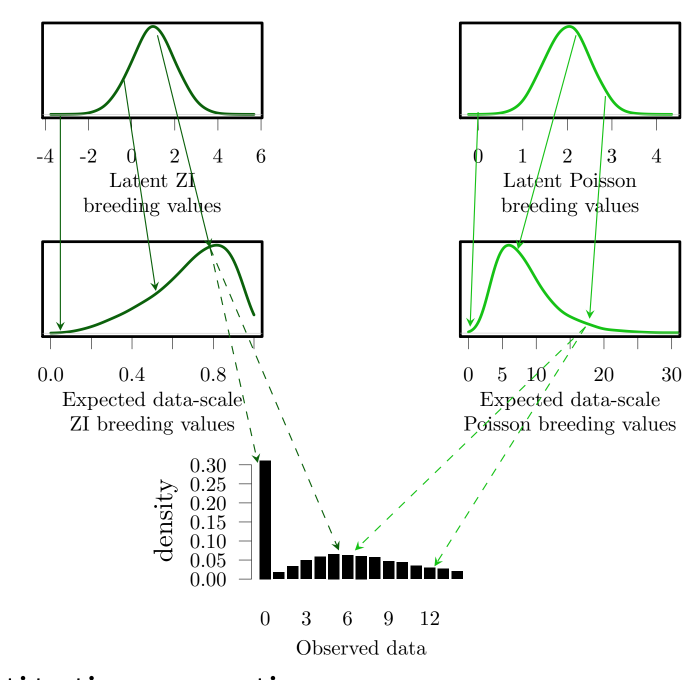
THEORY: How to estimate additive genetic variance in relative fitness ($V_A(\omega)$)?

Model distributions for fitness



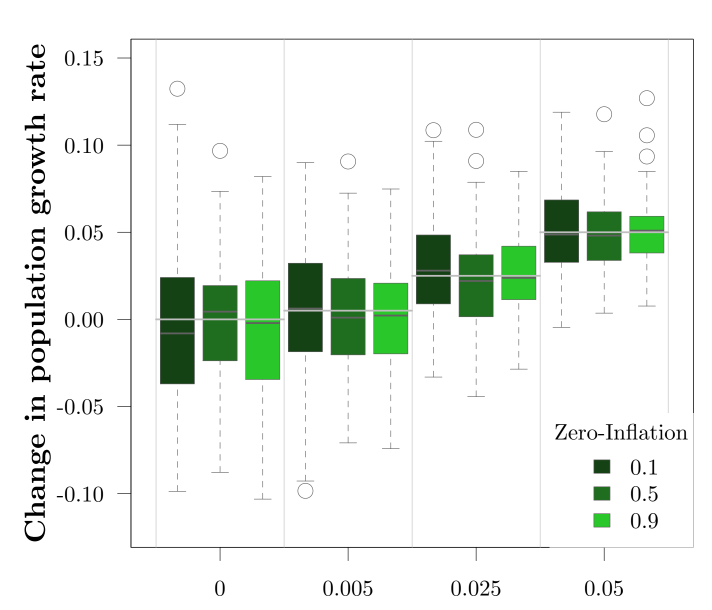
Zero-Inflated Over-Dispersed Poisson models tend to fit well lifetime fitness data. Quantitative genetic estimates may be more precise than with Gaussian or Poisson models.

ESTIMATE LATENT GENETIC VARIATION



Quantitative genetic ANIMAL MODELS use relatedness matrix to estimate G-matrix for the zero-inflation and the over-dispersed Poisson processes on two latent scales (log and logit).

Convert latent parameters to $V_{\mathcal{A}}(\omega)$



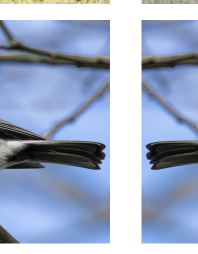
Simulated $V_A(\omega)$ after back-transformation

Monte-Carlo integration of latent breeding values to the data scale. Back-transformation works for individual-based simulations: $V_A(\omega)$ equals the increase in population growth rate.

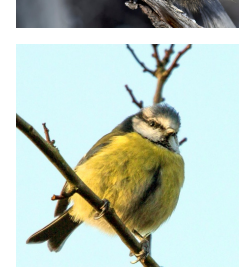
EMPERICAL RESULTS: Additive genetic variation in fitness detectable in most populations

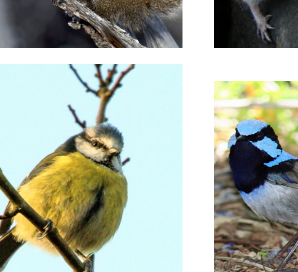




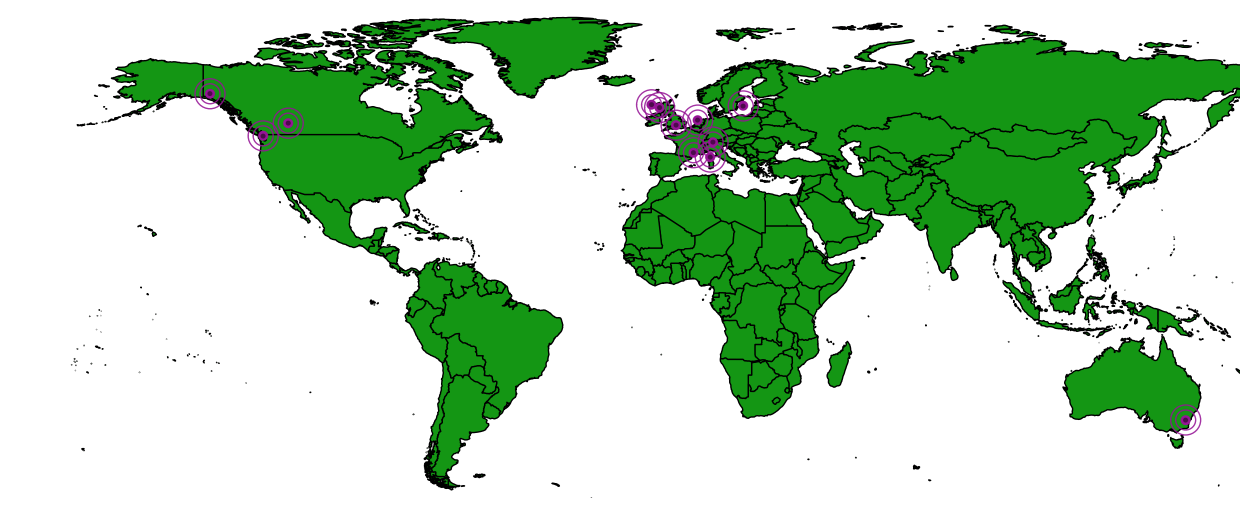




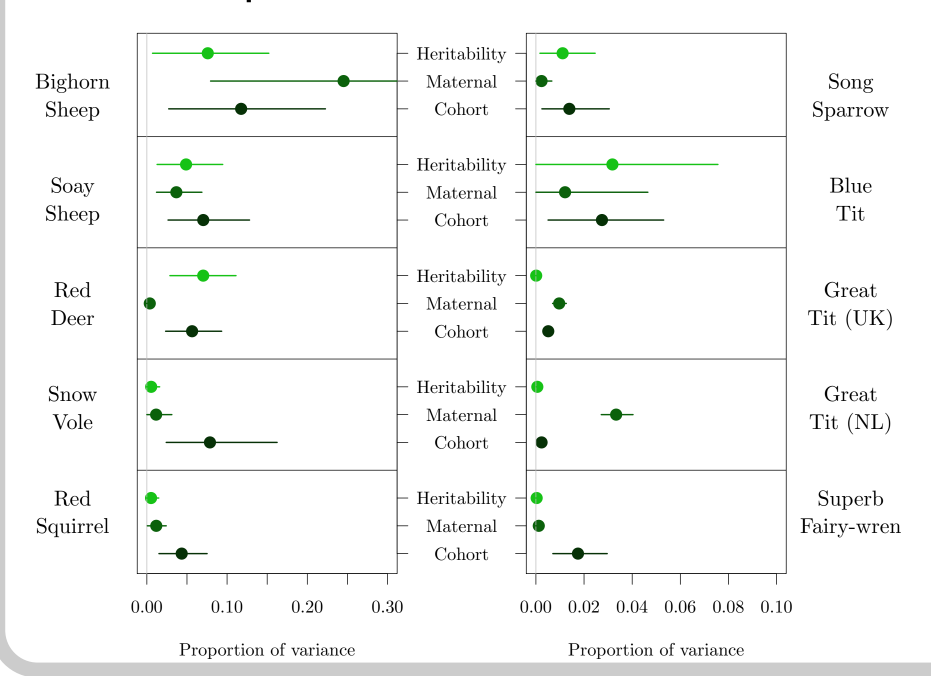


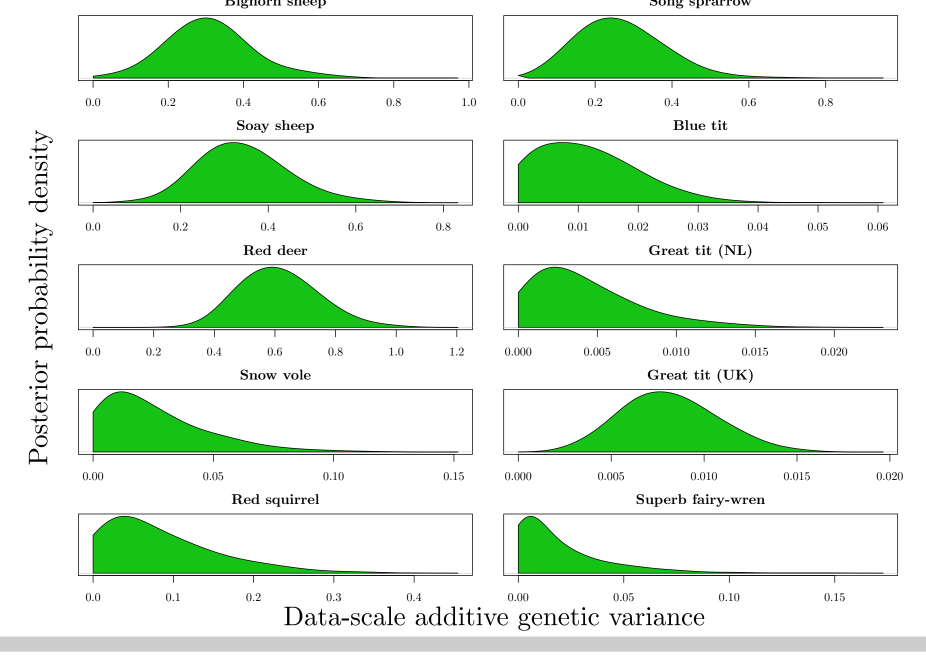






Preliminary models suggest small heritabilities of fitness on the data-scale (average 2.4%). Cohort effects, maternal effects and stochasticity dominate variation. However, credibility intervals of $V_A(\omega)$ exclude 0 in 5 out of 10 populations, and the posterior mode exceeds 0.1% in all of them, suggesting widespread on-going evolution.





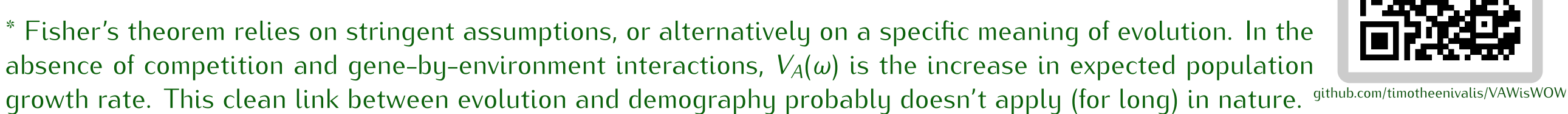
NEXT, WE WILL EXPLORE:

- Influence of priors on estimation
- Effect of missing data and immigration
- Sex-specific fitness. Does cross-sex genetic correlation constrain or enhance evolution?
- Relative importance of ZI, Poisson and their covariance
- Significance for population dynamics

Website

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R and LATEX code

timotheenivalis.github.io