Southern introgression increases adaptive immune gene variability in northern range margin populations of Fire-bellied toad

Binia De Cahsan, Katrin Kiemel, Michael Westbury, Maike Lauritsen, Marijke Autenrieth, Günter Gollmann, Silke Schweiger, Marika Stenberg, Per Nyström, Hauke Drews, and Ralph Tiedemann

1University of Copenhagen
2Affiliation not available
3University of Copenhagen Globe Institute
4Univ Potsdam

February 19, 2021

Abstract

Northern range margin populations of the European fire-bellied toad (Bombina bombina) have rapidly declined during recent decades. Extensive agricultural land use has fragmented the landscape, leading to habitat disruption and loss, as well as eutrophication of ponds. In Northern Germany (Schleswig-Holstein) and Southern Sweden, this decline resulted in decreased gene flow from surrounding populations, low genetic diversity, and a putative reduction in adaptive potential, leaving populations vulnerable to future environmental and climatic changes. Previous studies using mitochondrial control region and nuclear transcriptome-wide SNP data detected introgressive hybridization in multiple northern B. bombina populations after presumed illegal release of toads from Austria. Here, we determine the impact of this introgression by comparing the body conditions (as a proxy for fitness) of introgressed and non-introgressed populations, and the genetic consequences in two candidate genes for putative local adaptation (the MHC II gene as part of the adaptive immune system and the stress response gene HSP70 kDa). We detected regional differences in body condition. We observed significantly elevated levels of within individual MHC allele counts in introgressed Swedish populations, associated with a tendency towards higher body weight, relative to regional non-introgressed populations. These differences were not observed among introgressed and non-introgressed German populations. Genetic diversity in both MHC and HSP was generally lower in northern than southern populations. Our study sheds light on the potential benefits of translocations of more distantly related conspecifics as a means to increase adaptive genetic variability and fitness of struggling range margin populations without distortion of local adaptation.

Hosted file

MHCII alleles found in Austria and Germany

- HSP 70 kDa allele type 3
- HSP 70 kDa allele type 4
- HSP 70 kDa allele type 5
- HSP 70 kDa allele type 6
- HSP 70 kDa allele type 7
- HSP 70 kDa allele type 8
- HSP 70 kDa allele type 9

MHCII alleles found in Austria and Sweden

- HSP 70 kDa allele type 1
- HSP 70 kDa allele type 2

MHCII alleles found in Germany and Sweden

- HSP 70 kDa allele type 0

MHCII alleles found in all three countries

- HSP 70 kDa allele type 0