Population structure and demography

Prepared by Claire Mérot & Anna Tigano Physalia Course

Why does population structure matter when studying adaptation?

Evolution (including adaptive evolution)

is the result of the interplay of

Selection

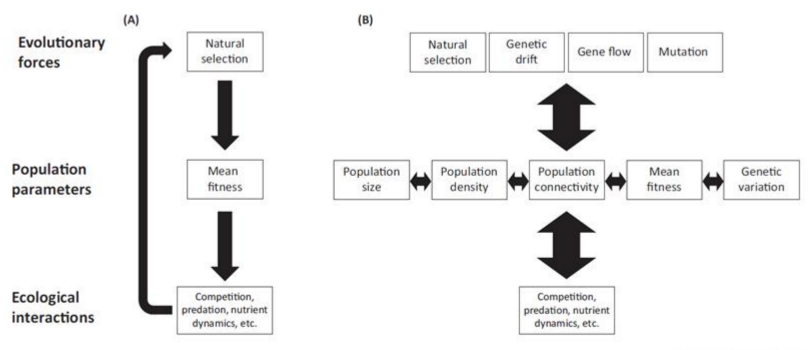
Drift

Mutation

Gene flow (migration + recombination)

Evolutionary, demographic and ecological processes are inseparable

Lowe, W. H., Kovach, R. P., & Allendorf, F. W. (2017). Population genetics and demography unite ecology and evolution. *Trends in Ecology & Evolution*, 32(2), 141-152.



Trends in Ecology & Evolution

Figure 1. Evolutionary and Ecological Processes Are Inseparable. Conceptual illustration of interconnections among evolutionary forces and ecological interactions (biotic and abiotic) through population-level demographic and genetic parameters. (A) represents those interconnections emphasized in current eco-evolutionary research. (B) represents a more comprehensive model of these interconnections, including the full suite of evolutionary forces and a range of population parameters that are themselves interdependent. We build our review around population demographic parameters (size, density, connectivity), but describe key interactions with genetic parameters (mean fitness, genetic variation). We define mean fitness according to population genetics theory as the sum of the fitnesses of genotypes in a population weighted by their proportions [88], thus representing the population-level effects of local adaptation.

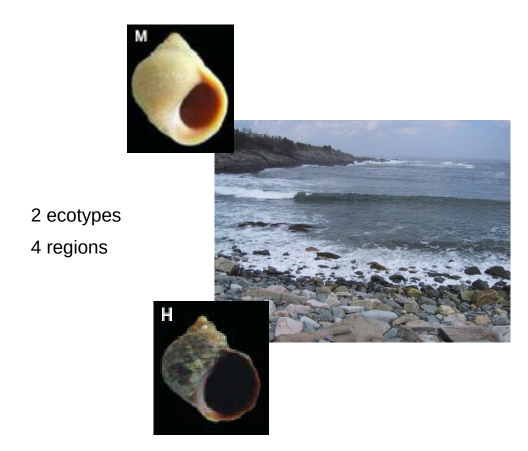
Complementary objectives:

	Study selection and adaptation	Demographic history and structure of populations
Actions	Focus on (putatively) adaptive loci	Focus on neutral loci
Use	. Study ecological/functional diversity	. Understand the past history of populations
	. Understand adaptative processes under divergent or balancing selection	. Describe population connectivity
	. Identify candidate genes	. Assess general genetic diversity

Different loci tells a different story...

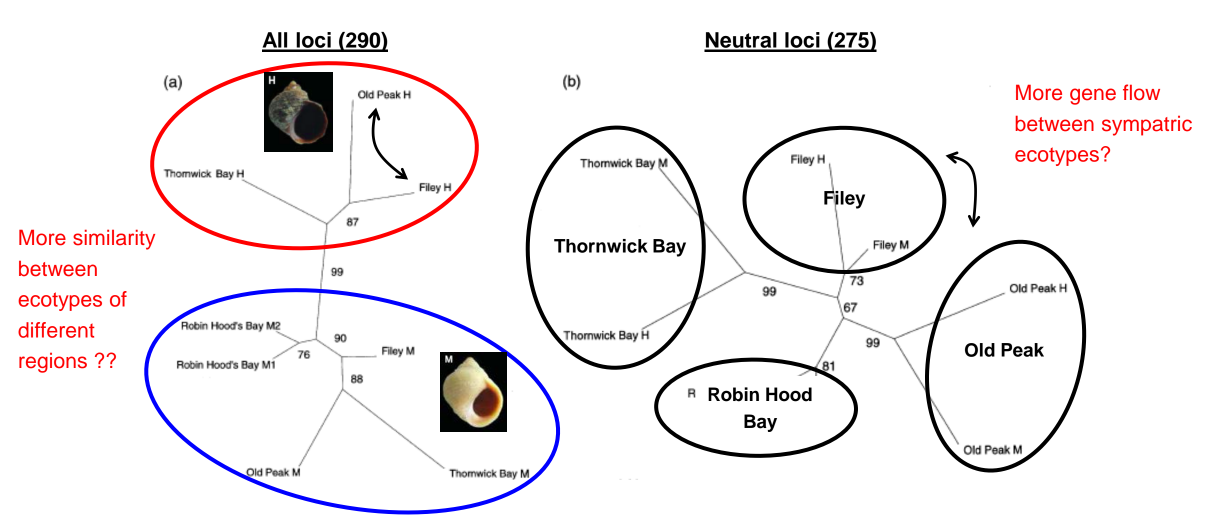
<u>AFLP</u>

. Parapatric ecotype *Littorina saxatilis*



Different loci tells a different story...

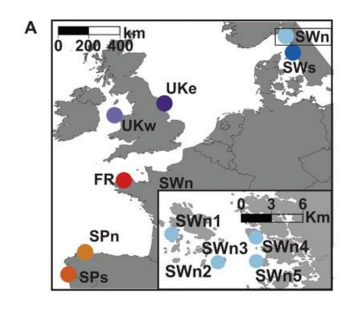
<u>AFLP</u>

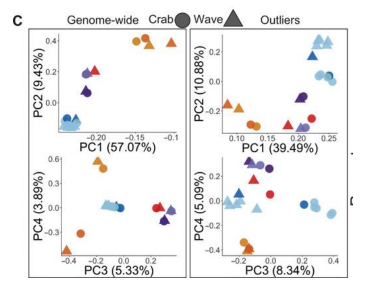


Wilding, C. S., Butlin, R. K., & Grahame, J. (2001). Differential gene exchange between parapatric morphs of Littorina saxatilis detected using AFLP markers. *Journal of Evolutionary Biology*, *14*(4), 611-619.

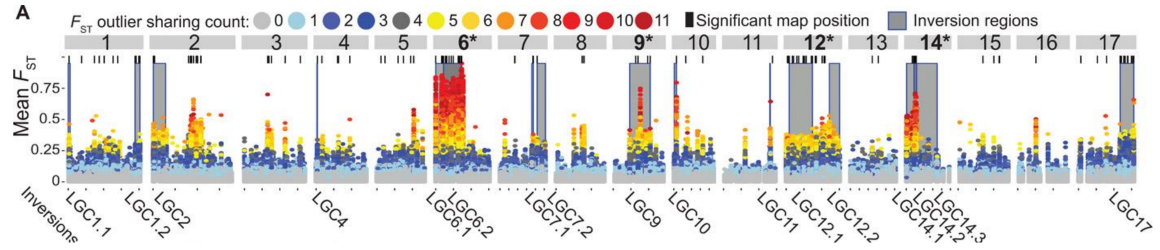
Different loci tells a different story...

Whole-genome sequencing (pool-seq)





Morales, H. E., Faria, R., Johannesson, K., Larsson, T., Panova, M., Westram, A. M., & Butlin, R. K. (2019). Genomic architecture of parallel ecological divergence: beyond a single environmental contrast. Science advances, 5(12), eaav9963.



Drift

= variation in allele frequency due to random processes

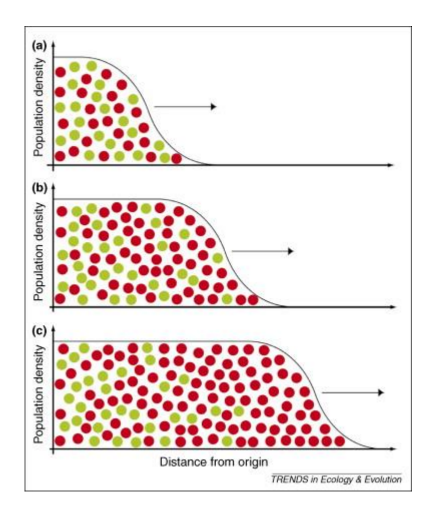
Drift is stronger in smaller populations and it can cause the loss or fixation of a variant due to random sampling of alleles.

Drift is the main driver of genetic population structure, and can generate a genetic footprint similar to that of selection.

Allele surfing

Populations on the leading edge of the expansion are small, and individuals from those populations contribute disproportionately to the propagating wave of expansion.

⇒Rapid drift of some alleles at the expanding edge and high differentiation in allele frequencies over the landscape for some loci, even in the absence of selection



Surfing during population expansions promotes genetic revolutions and structuration

Excoffier & Ray

TREE 2008 https://doi.org/10.1016/j.tree.2008.04.004

Spatial autocorrelation

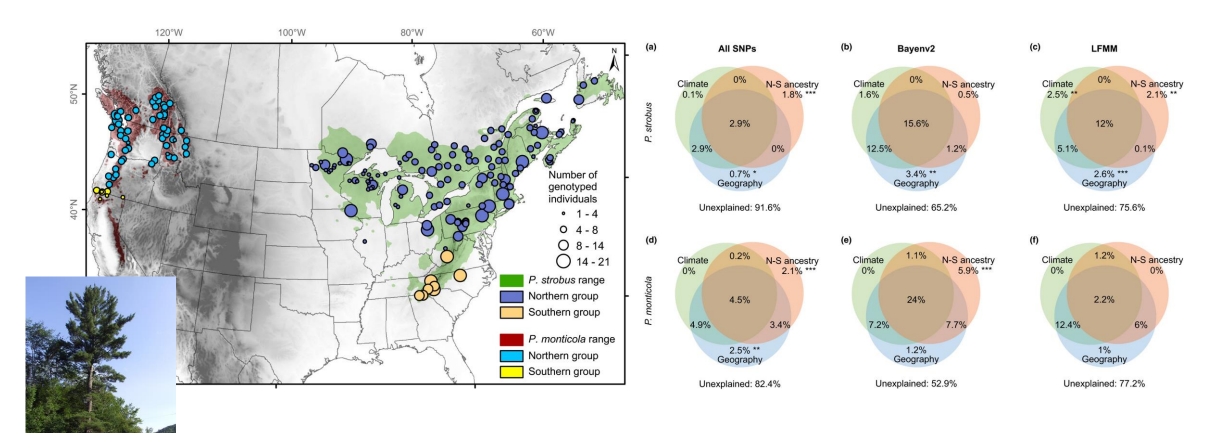
Correlation between environmental variation & geographic distances (e.g. climatic clines!)

⇒Which patterns results from the balance selection/migration? Which patterns result from the balance drift/migration?

- + Residuals of past range expansion out of glacial refugees...
- ⇒What is adaptation? What is the results of past history?

Nearby locations are not statistically independent, strong correlations between neutral alleles and environmental variables are more likely to occur by chance than expected with some null models

Isolation-by-distance or adaptation along a gradient ... or both?



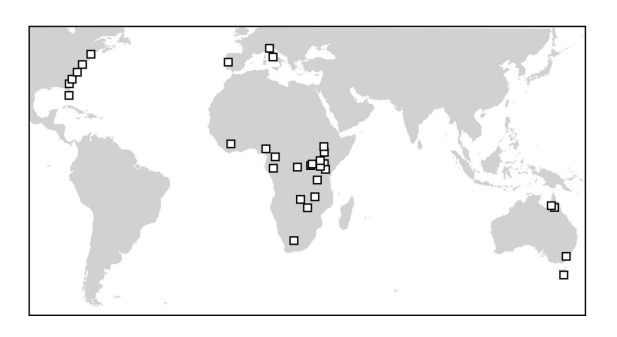
The challenge of separating signatures of local adaptation from those of isolation by distance and colonization history: The case of two white pines Nadeau et al, 2016 https://doi.org/10.1002/ece3.2550

Contact between different lineages / hybridization

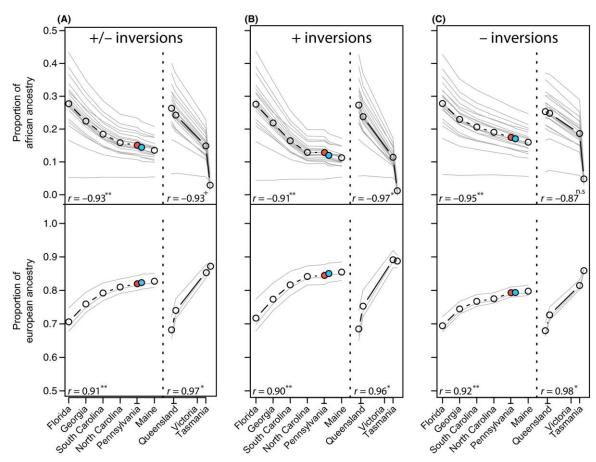
Signature of selection or of local adaptation are best detected in a context of (high) gene flow.

Any substructure (lineages, species, secondary contact, admixed populations) should be taken into account.

Clinal variation or secondary contact... Or both?



Bergland et al, 2015 MolEcol https://doi-org/10.1111/mec.13455



How to characterise population structure?

Unsupervised methods:

- PCA

Semi-supervised methods (K = number of expected clusters)

Bayesian clustering

Supervised methods (with location information for instance)

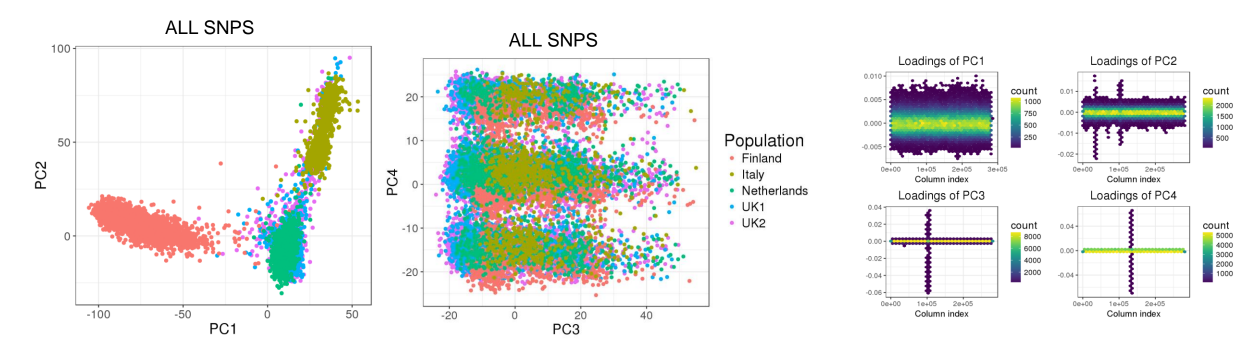
- DAPC
- Fst between pairs of populations

Principal Component Analysis (PCA)

- A common statistical tool that reduces matrix complexity by identifying the eigenvectors and ordering them
- The top PCs reflect axis of genetic variation along which individuals with same ancestry, or exchanging genetic material, are more similar to each other.
- Caution: can be strongly driven by few loci in linkage disequilibrium...
- For population structure purpose:
- -> compare PCA on all SNPs vs. PCA on LD-pruned SNPs
- -> look at loadings of the PCs: which fraction of the genome explains PC1? Explains PC2? Etc...
- There is lots of genetic variance, it can be relatively expected that even PC1 explains less than 1% of variance. (but it can also capture 20-50%... Depends on the dataset!

Principal Component Analysis (PCA)

Each individual is a point with coordinates along all PCs
Each genetic marker contribute to all PCs with a different strength (loadings)

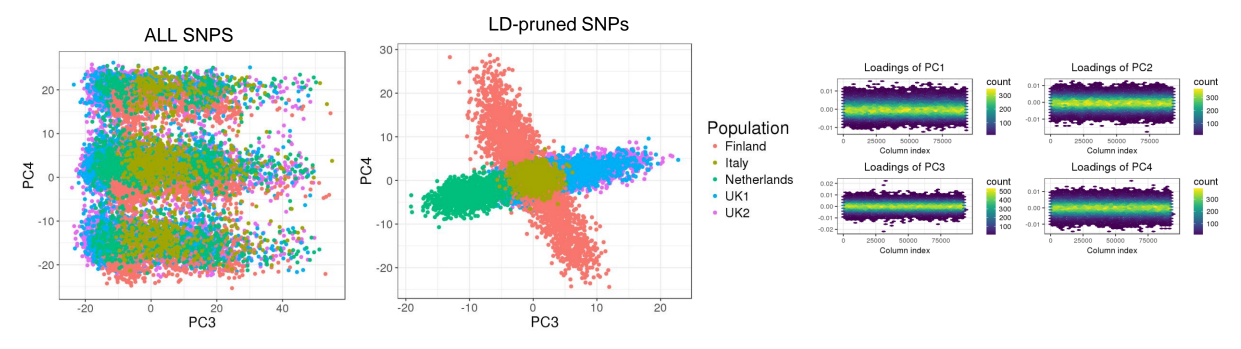


Packages *bigstatsr*, *bigsnpr* to remove short-range and long-range LD. Nice tutorial about PCA for pop genomics! Florian Privé https://privefl.github.io/bigsnpr/articles/how-to-PCA.html

Principal Component Analysis (PCA)

Each individual is a point with coordinates along all PCs

Each genetic marker contribute to all PCs with a different strength (loadings)



Packages *bigstatsr*, *bigsnpr* to remove short-range and long-range LD. Nice tutorial about PCA for pop genomics! Florian Privé https://privefl.github.io/bigsnpr/articles/how-to-PCA.html

Bayesian clustering (STRUCTURE, etc..)

- Aim to sort individuals into K clusters so as to minimize departures from Hardy-Weinberg equilibrium and linkage equilibrium
- Caution: can be strongly driven by few loci in linkage disequilibrium...
- For population structure purpose:
- -> compare results on all SNPs vs. results on LD-pruned SNPs
- -> explore many values of K report likelihood
- Admixture or FastSTRUCTURE replace STRUCTURE for genome-wide data
- Evaluate the fit of the model

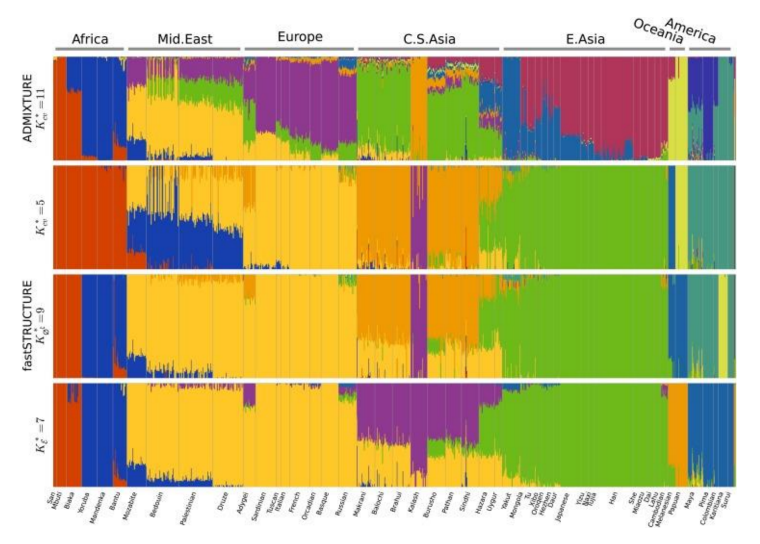
Evaluation of model fit of inferred admixture proportions

Genís Garcia-Erill Anders Albrechtsen

MER 2020

https://doi.org/10.1111/1755-0998.13171

Bayesian clustering (STRUCTURE, etc..)



Each individual is a thin vertical line that is partitioned into *K* colored segments according to its membership coefficients in *K* clusters.

fastSTRUCTURE: variational inference of population structure in large SNP data sets 2014 Genetics

Anil Raj¹, Matthew Stephens², Jonathan K Pritchard² 10.1534/genetics.114.164350

The advantage of unsupervised/semisupervised methods: => Other surprises!!

Species lineage substructure

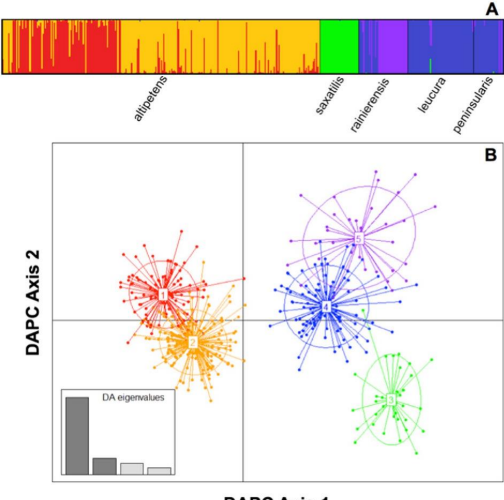
Hybridisation

Chromosomal rearrangements....

DAPC (discriminant PCA)

- A mix of a discriminant analysis and a PCA
- It will try very hard to find axis of variation that discriminate the groups given a priori
- WARNING: A dangerous analysis when we have much more markers (SNPs) than groups (populations)... Be well aware of not over-fitting and not-overinterpreting the output.

Miller, J.M., Cullingham, C.I. & Peery, R.M. The influence of a priori grouping on inference of genetic clusters: simulation study and literature review of the DAPC method. *Heredity* (2020). https://doi.org/10.1038/s41437-020-0348-2



DAPC Axis 1

Languin et al, 2018, Conservation genomics

Pairwise Fst

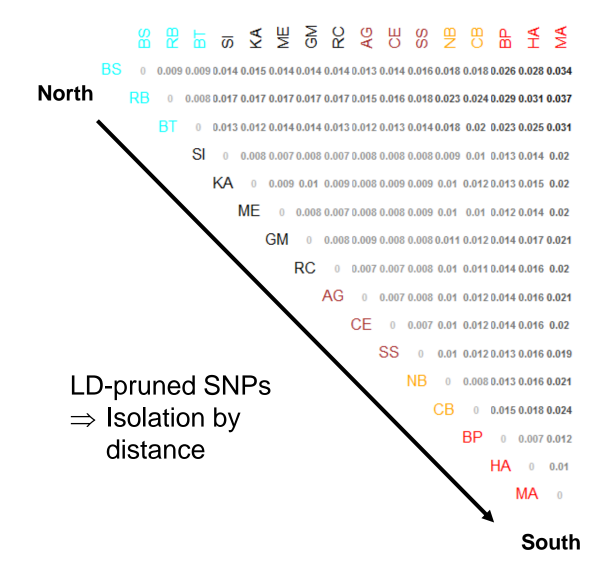
- Fst can only been computed between two groups.
- When sampling several populations, we will be interested in Fst between all pairs of individuals.
- A measure of genetic distance between all populations (does it correlate with ecological distances? Geographic distances? Etc?)
- Again, likely better on Ld-pruned SNPs to infer neutral structure...
- Absolute values are informative... (0,000x -> high gene flow, don't bother too much about looking for structure.
 0,01-0,1 -> consider carefully structure... Higher: do you really have one species?)

Pairwise Fst

ALL SNPs

M H M C S C A C M M A A W M A North 0.002 0.004 0.002 0.004 0.002 0.004 0.005 0.003 0.007 0.011 0.01 KΑ 0.009 0.02 .013 0.004 0.011 0.013 0.004 0.01 0.006 0.005 0.016 0.004 0.002 0.004 0.002 0.004 0.005 0.003 0.007 0.011 0.011 0.003 0.011 0.004 0.004 0.013 0.008 0.014 0.021 0.013 0.006 0.002 0.003 0.007 0.005 0.009 0.014 0.01 0.005 0.006 0.003 0.006 0.005 0.007 0.012 0.002 0.006 0.005 0.008 0.012 0.009 0.008 0.007 0.008 0.012 0.007 0 0.004 0.005 0.006 0.012 0.008 0.012 0.014 0 0.003 0.007 0.008 MA o South

LD-pruned SNPs



Case study

ORIGINAL ARTICLE

WILEY MOLECULAR ECOLOGY

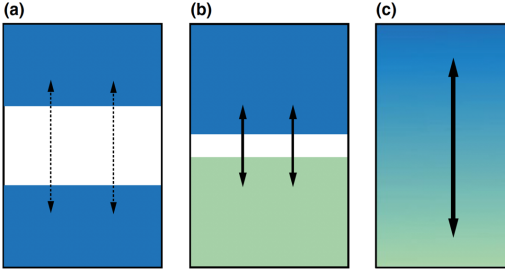
Genome-wide signals of drift and local adaptation during rapid lineage divergence in a songbird

```
Guillermo Friis<sup>1</sup> | Guillermo Fandos<sup>2</sup> | Amanda J. Zellmer<sup>3</sup> | John E. McCormack<sup>3,4</sup> | Brant C. Faircloth<sup>5</sup> | Borja Milá<sup>1</sup>
```

Set expectations

(a) Geographically isolated populations in similar habitats.

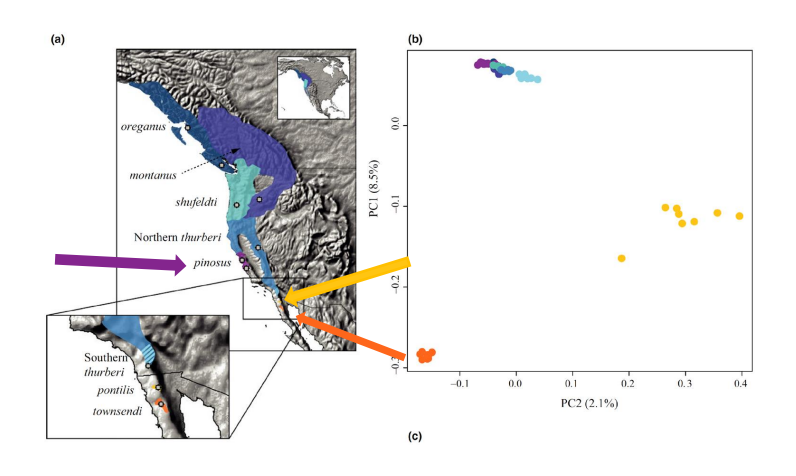
(b) Parapatric populations in ecologically divergent habitats.



(c) Population continuum across a selective gradient

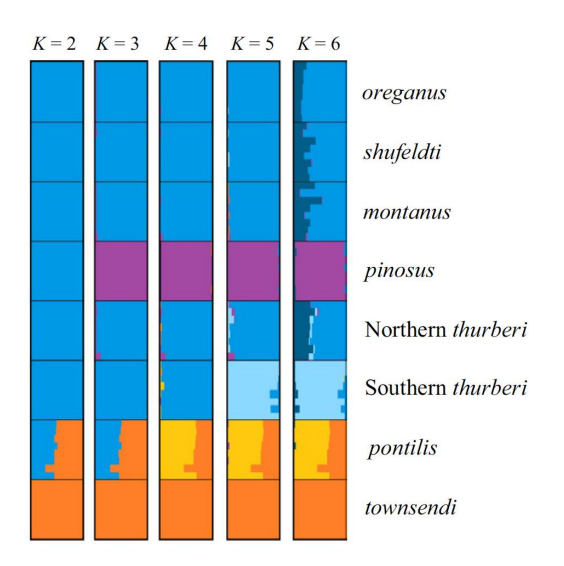
Habitat types: Gene flow: Neutral divergence: Adaptive divergence: ONE LOW HIGH LOW TWO MODERATE MODERATE HIGH GRADIENT HIGH LOW HIGH

PCA for data exploration



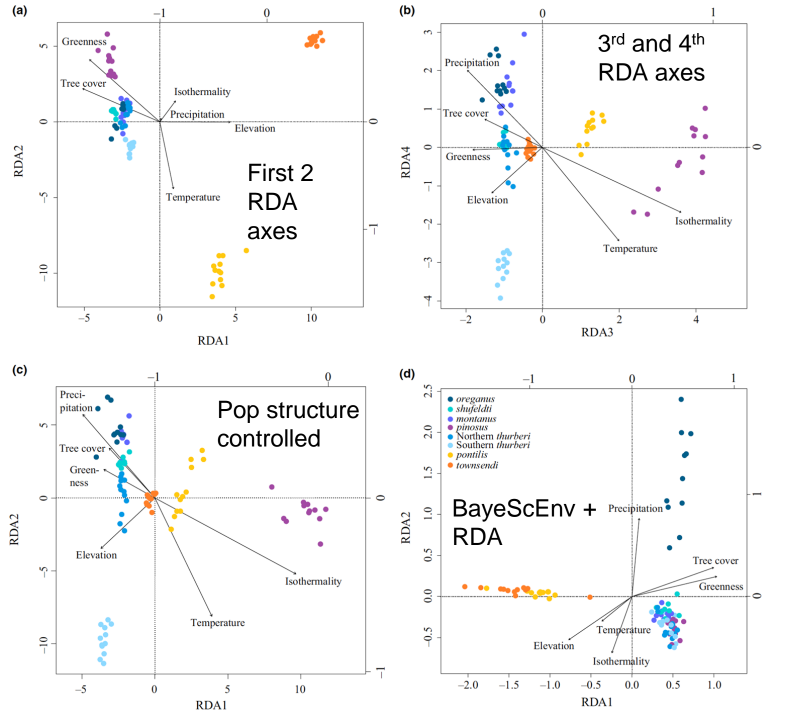
Prior-free data exploration

STRUCTURE for population structure



Always examine multiple K values as more than one K could be biologically informative

(STRUCTURE doesn't deal well with hierarchical population structure)



GEA

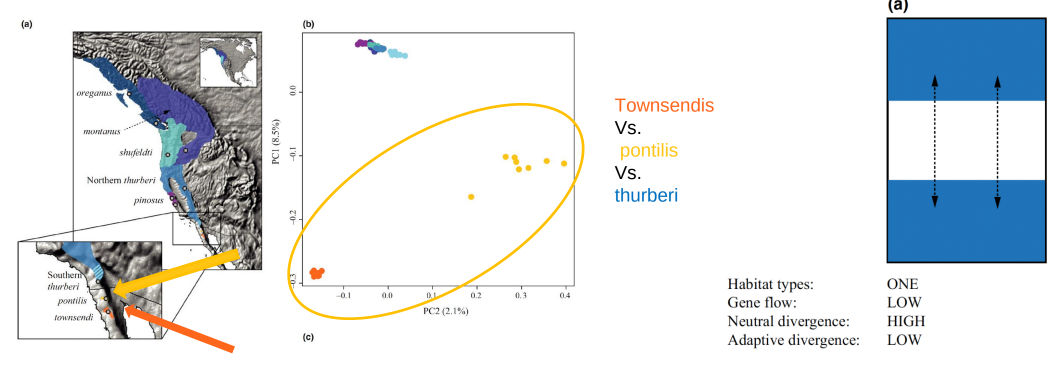
Genotype-environment associations with multiple approaches

Lecture tomorrow, we come back on that

Partition of genetic variation

- Environmental variables (controlling for population structure) 1.17%
- Environmental variable + pop structure 7.41%
- 92.59%?
 - Loci under balancing selection
 - Other selective pressures
 - Shared neutral variation due to relatedness and/or gene flow

Environment + geography + demography

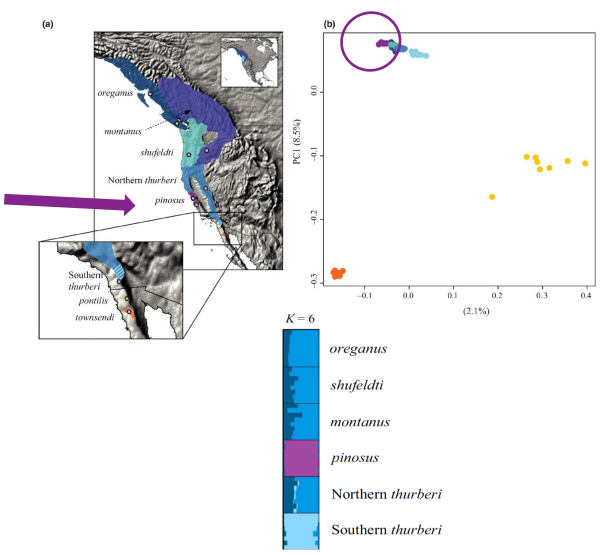


Strong population structure and weak GEA

= Isolation-by-resistance and drift

(small populations + desert between suitable habitats)

Environment + geography + demography



Weak pop structure and stronger differentiation on GEA

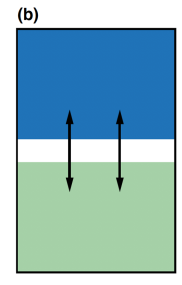
= Isolation-by-adaptation?

Or secondary contact zone?

Pinosus Vs. Thurberi

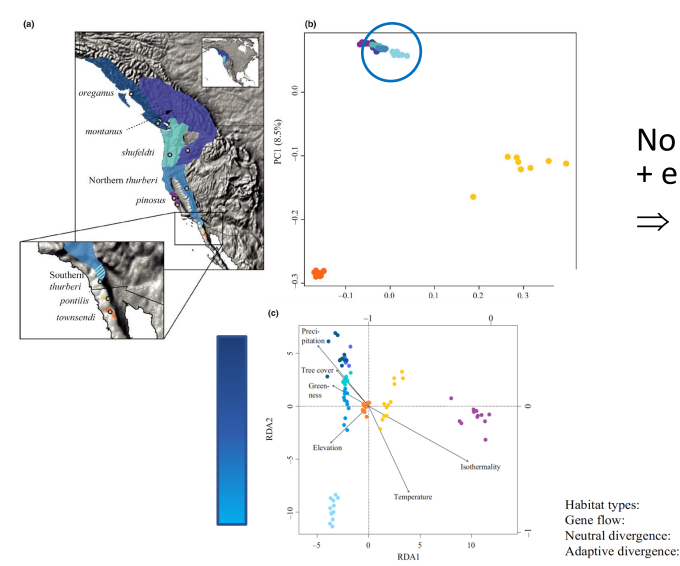
Barriers to gene flow due to niche differences?

Habitat types: Gene flow: Neutral divergence: Adaptive divergence:



TWO MODERATE MODERATE HIGH

Environment + geography + demography



No pop structure

- + environmental associations
- ⇒ Ongoing gene flow and local adaptation

GRADIENT
HIGH
LOW
te: HIGH

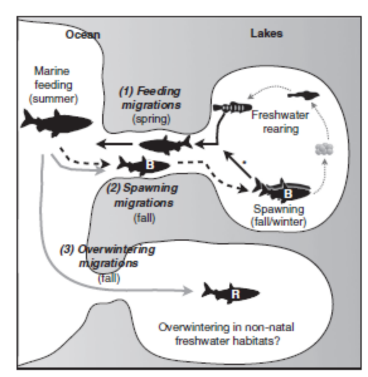
And help from experimental work /knowledge of natural history

Capture-Mark-Recapture

-> population size, dynamic and movement

Spatial ecology

-> tracking, etc..

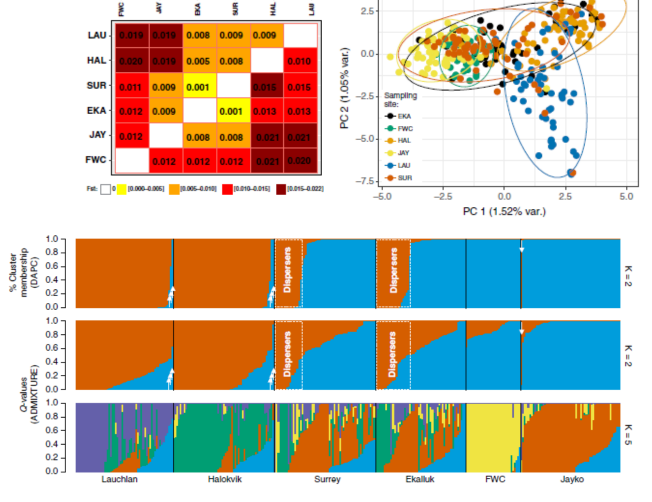


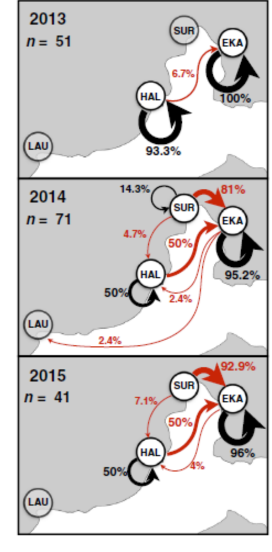


e.g: telemetry & genomics

Moore, J.-S., L.N. Harris, J. Le Luyer, B.J.G. Sutherland, Q. Rougemont, R.F. Tallman, A.T. Fisk & L. Bernatchez (2017) Genomics and telemetry suggest a role for migration harshness in determining overwintering habitat choice, but not gene flow, in anadromous Arctic Char. *Molecular Ecology*, 26(24): 6784-6800

And help from experimental work /knowledge of natural history





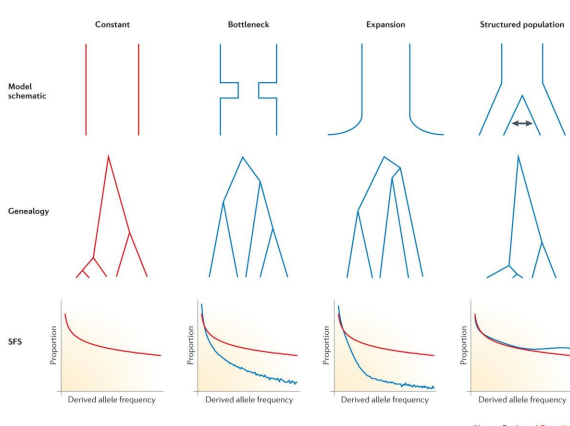
Genomics and telemetry suggest a role for migration harshness in determining overwintering habitat choice, but not gene flow, in anadromous Arctic Char

Moore, J.-S.,. *Molecular Ecology*, 26(24): 6784-6800

Beyond present structure... How to know population history and demography?

Models:

- to understand population history, bottleneck, gene flow...
- demography can set a null model against which one can look for the effect of selection



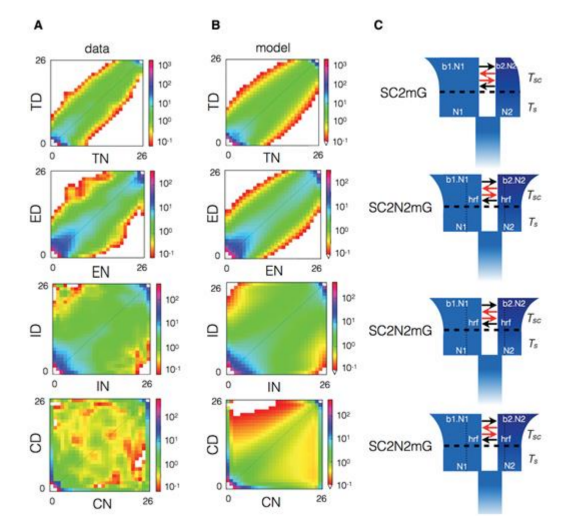
Nature Reviews | Genetics

Beyond present structure... How to know population history and demography?

Based on coalescence theory

Compare SFS (site frequency spectrum) between real data and modelled data under different scenario

Common tools: dadi, FastSimCoal, ABC...



Population structure and demography

A good overview

Schraiber, J., Akey, J. Methods and models for unravelling human evolutionary history. Nat Rev Genet 16, 727–740 (2015). https://doi.org/10.1038/nrg4005

