Ecological Genetics – HS18

v0.3

Gleb Ebert

October 28, 2018

This document aims to summarize the lecture Ecological Genetics as it was taught in the autumn semester of 2018. Unfortunately I can't guarantee that it is complete and free of errors. You can contact me under glebert@student.ethz.ch if you have any suggestions for improvement. The newest version of this summary can always be found here: http://www.glebsite.ch

Contents

1	Introduction	2
2	Species	2
3	Molecular Markers	3
4	Sampling	4
5	Genomic Methods	5
6	Genetic variation	7

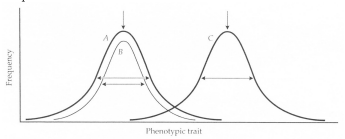
1 Introduction

"Nothing in biology makes sense except in the light of evolution"

— Theodosius Dobzhansky

In 1971 Ford wrote that ecological genetics deal with the adjustments and adaptations of wild populations to their environment. According to Conner and Hartl (2004) ecological genetics is the study of the process of phenotypic evolution occurring in present-day natural populations and is concerned with the genetics of ecologically important traits, that is, those traits related to fitness.

Phenotypic evolution is the change in the mean or variance of a trait across generations due to changes in allele frequencies.



Ecologically important traits are closely tied to fitness and are important in determining an organisms adaptation to its natural environment. Adaptation is a heritable phenotypic trait that has evolved in a population in response to a specific environmental factor and improves the survival or reproduction of its carriers. It can also be seen as a process whereby the members of a population become better suited to some feature of their environment through change in a characteristic that affects their survival or reproduction. Of the four key evolutionary processes, only natural selection consistently leads to adaptation (mutations, genetic drift and gene flow don't).

Uses of ecological genetics include

- agriculture (crop improvement)
- medicine (e.g. antibiotics)
- conservation measures (assisted migration)
- $\bullet\,$ geographical differences between populations
- changes in species composition
- habitat adaptation & speciation

Fields related to ecological genetics include

- population genetics
- ecology
- evolutionary biology
- phylogenetics
- quantitative genetics
- statistics
- molecular biology
- \bullet epigenetics
- genomics

2 Species

Species are the fundamental unit in ecology, evolution and conservation legislation. Depending on the research question, adequate species identification, assignment of samples to populations or discrimination of individuals may be of relevance.

2.1 Species concepts

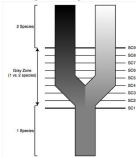
- Morphological / typological species concept: focus on similar morphology
- Biological species concept: group of potentially or actually interbreeding populations which are reproductively isolated from other such groups
- Phylogenetic species concept: focuses on monophyletic lineages

2.2 Operational Taxonomic Unit

An **OTU** is a group of organisms that is treated as a distinct evolutionary unit for the purposes of research underway. They are often applied when one or several species concepts fail. Once identified and research has been completed, OTUs should receive full taxonomic treatment and be given a scientific name if possible. OTUs are sometimes called **molecular operational taxonomic unit** (MOTU) when molecular methods are used.

2.3 Unified Species Concept

The only necessary property of species is that they form a separately evolving matepopulation (involves dynamics of gene flow and separation) lineage. The concept separates species conceptualization and separation. All criteria can be used for species delimination and any one of the properties is accepted as evidence for the existence of a species. More properties provide a higher degree of corroboration.



2.4 Identification of Species

Difficulties may include

- species-specific traits are not (always) visible
- differences are cryptic
- direct observation may be difficult and traces may be confused
- undescribed species may occur

Parataxonomy may be used when identification is difficult. It sorts the material to species on the basis of external morphology without considering taxonomy.

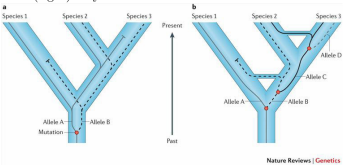
2.5 Species delimination

Species delimitation is the act of identifying species-level biological diversity (independent evolutionary lineages). Most methods fit models to collected data to make (often different) simplifying assumptions. Incongruence across methods may occur due to differences in the power to differentiate lineages or due to violations of one or several assumptions made by a given method. Fundamentally there are two approaches. Some models can assign samples to groups without being given information first (STRUCTURE, Structurama, Geneland). Others need the user to assign samples

to putative lineages (BPP, iBPP, spedeSTEM, DISSECT, tr2).

2.5.1 Problems of Species Trees

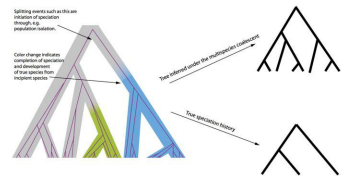
Problems like imcomplete lineage sorting (left) or gene flow (right) may occur.



2.5.2 Bayesian Species Identification under the Multispecies Coalescent

This method is currently the most used approach for species delimination. The **multispecies coalescent** describes the genealogical relationships between DNA samples from several species. Simplifying assumptions include:

- species phylogeny unknown
- complete isolation after divergence
- no recombination



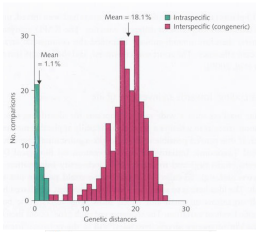
The above graph shows that the MSC approach identified populations as separate species from a simulated data set. MSC delimits structure, not species.

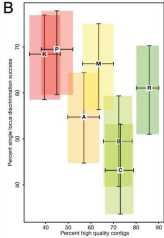
2.5.3 Recommendations for Species Delimination

- use at least 10 samples from all lineages
- simulate before analysis
- use several complementary methods
- combine genetic with nongenetic data
- define the used species concept
- be cautious

2.5.4 DNA Barcoding

DNA barcoding is a method for rapid identification of species through the analysis of short, standardized gene regions. These have to be universal for all animals or plants, have to be amenable to the production of bidirectional sequences with little ambiguity and allow for discrimination of most species. The barcoding gap defines the distance between species. Genetic distances are based on the extent of nucleotide sequence devergence. Some groups like orchirds lack the barcoding gap.





Examples of application are the identification of amphibian species diversity and abundance after epidemic diseases in Panama or restriction of shaving brushes to only use hair from the Hog badger instead of the Eurasian badger.

3 Molecular Markers

Molecular markers are polymorphic proteins or DNA sequences and reveal different alleles within individuals, populations or species. Ideally they can be used as **indicators** of genome-wide variation.

- Chromosome based markers
 - Numbers and staining patterns
- Enzyme based
 - Allozymes
- DNA based
 - Restriction fragment length polymorphisms (RFLPs)

• DNA & PCR based

- Random amplified polymorphic DNA (RAPD)
- Amplified fragment length polymorphism (AFLP)
- Microsatellites (SSRs)
- DNA sequencing and SNPs

3.1 Genome

The size of genomes can differ between species by large factors. Size and complexity are not coupled, as non-coding regions are the main reason for big genomes. These regions contain repeated sequences like tandem repeats or interspersed repeats (transposable elements). The number of chromosomes is limited, because at some point there would be problems with the spindle apparatus. Introns are spliced after transcription while intergenic regions aren't.

Mitochondria and chloroplasts have their own genome (mtDNA and cpDNA respectively). Depending on the species, different **organelle genomes** should be used for comparisons. There is much more organelle DNA in a cell than there is nuclear nDNA and it is easier to amplify because of its high conservation.

3.2 Widely used genetic markers

Marker	Advantages	Disadvantages
Allozymes	Cheap Universal protocols	Requirement for fresh or frozen material Potentially direct target of selection Limited number of available markers 'No longer used' (<1998)
Microsatellites	Informative (large number of alleles, high heterozygosity) Easy to isolate	High mutation rate Complex mutation behaviour Oifficult to automate Cross-study comparisons are difficult
AFLPs	Cheap Produces a large number of markers Easy to establish in the lab	Mainly dominant Oifficult to analyse Oifficult to automate Cross-study comparisons are difficult
DNA sequencing	Highest level of resolution possible Not biased Cross-study comparisons are easy Data repositories already exist (e.g. NCBI)	Sanger sequencing: significantly more expensive than the other techniques NGS: cost per base (bp) very low NGS: computational intense analyses
SNPs arrays	Low mutation rate High abundance Easy to type Cross-study comparisons are easy; data repositories already exist	Expensive to isolate Ascertainment bias Low information content of a single SNP

3.2.1 Microsatellites

- SSR (simple sequence repeat) and STR (short tandem repeat)
- highly polymorphic: mutation rates between 10^{-6} and 10^{-2} per locus per generation
- widely used to assess genetic variation in animals, plants and fungi as they are highly variable between individuals
- codominant
- mostly evolutionary neutral, as they are in intragenic regions
- PCR-based (primers, agarose-gel electrophoresis)
- capillary sequencers use fluorescence labelled primers
- $\bullet~$ 10-20 SSRs per study
- the mutation mechanism is called **slipped-strand mispairing**: polymerase slips of and when rejoining doesn't know which repeat it already copied; insertions and excisions happen, but the latter seem to be corrected in nature

3.2.2 Structural variation (SV)

- Microsatellite repeats
- 1bp indels
- More complex insertions and deletions
- Copy number variants (CNVs) (> 1kb)
- ...

3.2.3 Amplified fragment-length polymorphisms (AFLPs)

- \bullet 100-4'000 random markers
- dominant (homo- and heterozygotes indistinguishable)
- 1) DNA extraction
- 2) Digestion by restriction enzyme Msel and EcoRI
- 3) Ligation of the adaptors Msel and EcoRI
- 4) Selection amplification (+3/+3 bp)
- **5**) Sequencing
- 6) Binary data matrix (peak present/absent)

3.2.4 SNP microarrays

DNA is hybridised to predefined probes. Only common and known SNPs are called. This is the ascertainment bias.

4 Sampling

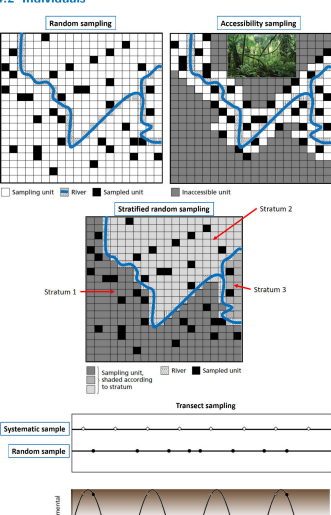
Sampling determines...

- The chances of answering your research questions
- The time you may have to spend to answer your questions
- Research costs
- The likelihood of obtaining research permits
- . . .

4.1 Populations

Population type	Definition
Genetic	All individuals which are connected by gene-flow
Ecological	Group of organisms occuring in a particular area at a particular time
Statistical	The universe of times that are under study

4.2 Individuals



Do you sample proportionally or equally across strata? Randomly or systematically? There is no single right solution. One should always consider the circumstances.

Roughly 20-30 individuals represent a population (> 80% of all alleles).

4.3 Important considerations

- Documentation: archived, reproducable, verifiable, new technologies
- Storage: adequate storage and transport, test in advance!
- Permits: sampling, handling animals, export and import

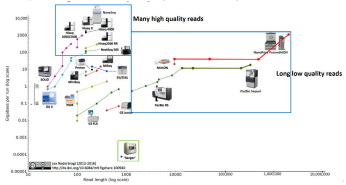
4.4 Nagoya protocol

The convention on biological diversity from 1993 had the objectives of conservation and sustainable use of biological diversity as well as the sharing to benefits arising from the utilisation of genetic resources. The Nagoya protocol from 2010 is a supplementary agreement that expands on the fair and equitable sharing of benefits arising out of the utilisation of genetic resources.

- Access obligations: Provide fair and non-arbitrary application procedures and issue permits when access is granted
- Benefit-sharing obligations: Share the value of genetic resources and traditional knowledge with developing countries
- Compliance obligations: Ensure that genetic resources and traditional knowledge have been accessed in accordance with prior informed consent

5 Genomic Methods

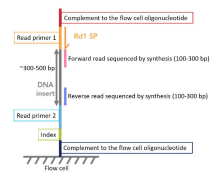
Sequencing methods evolved rapidly since the development of pyrosequencing in 1993. The cost per raw megabase is sinking rapidly as well. Data is being gathered faster and cheaper than it can be analysed nowadays.



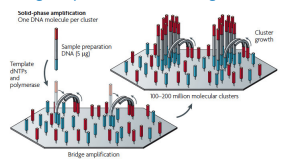
5.1 Illumina high-throughput sequencing

5.1.1 Sample / library preparation

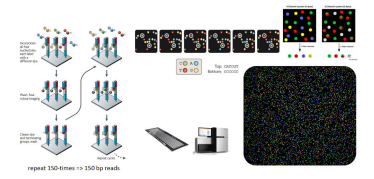
- 1) Fragmentation of genomic DNA
 - 1) Mechanical shearing: e.g. sonification
 - 2) Tagmentation: enzymes
- 2) Size selection: 300-500bp
- 3) Adapter ligation
 - 1) PCR amplification
 - 2) Individual barcoding



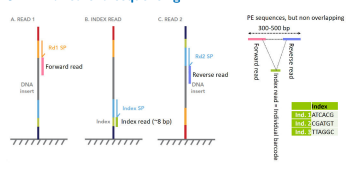
5.1.2 Bridge amplification and cluster generation



5.1.3 Sequencing by synthesis



5.1.4 Paired-end sequencing



5.2 3rd generation sequencing

3rd gen methods sequence single molecules and generates long reads of 10'000 - 2 million bp. They don't use PCR and thus avoid PCR artefacts. One player in the market is **Pacific Biosciences**. They use single molecule real-time analysis (SMART). Another one is **Oxford NanoPore**, whose **MinION** costs only around \$300 and generates even longer reads than PacBio is able to. Sequencing happens through voltage changes in the membrane when DNA passes the nano pore. NanoPore sequencing is useful for *de novo* genome assembly, CNV detection, real-time identification of samples and mRNA splicing variant identification.

5.3 Uses of NGS data

NGS inferences require fully annotated high quality reference genomes, draft reference genomes, reference transcriptomes or *de novo* assembled STACKS (RADseq). Methods used for high-throughput marker discovery include:

- Sequencing of individuals and populations
- Whole-genome re-sequencing
- Transcriptome sequencing (RNA-seq)
- Reduced representation sequencing (RADseq; no reference genome required)
- Target capture sequencing methods
 - Sequencing of ultra-conserved elements
 - Exome capture

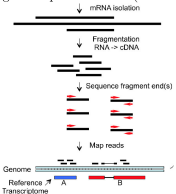
5.3.1 Whole genome re-sequencing

No bias from insufficient marker density or distribution occurs when re-sequencing whole genomes.

- Individual sequencing: 1 Flow cell NovaSeq S4 (~ 35'000CHF+ ~ 100CHF per individual library
- Sequencing pools of individuals (Pool-seq)
 - Cost effective: population libraries ($\sim 100 \text{CHF}$ per pool library)
 - Lower coverage per individual
 - Population allele frequencity estimates (no individuals genotypes)

5.3.2 Transcriptome sequencing (RNA-seq)

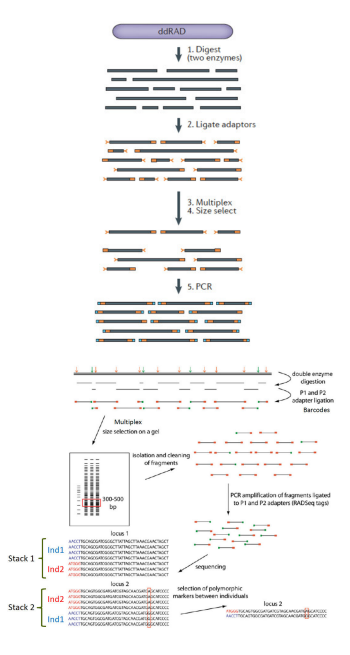
This method reduces the complexity of the genome through only sequencing the expressed exons (mRNA).



5.3.3 Reduced representation sequencing (RADseq)

One method of reduced representation sequencing is Restriction site-Associated DNA Sequencing (RADseq). Restriction enzymes are used to obtain DNA sequences adjacent to a large number of restriction cut sites. As these site are conserved and no reference genome is required, RADseq is used extensively for high-throughput SNP discovery and genotyping in ecological and evolutionary studies. The sequencing depth is considered to be high and the cost per sample lower. 1-200 loci per 1 Mb of genome with lengths of 100-150bp result in a $\frac{100 \log (*1000 \log)}{1'000'000} = 100 \times (*1000 \log)$ reduction of genome complexity.

ddRAD: double digest RADseq



5.3.4 Marker identification

If possible, reads are mapped to a genome (e.g. through Burrows-Wheeler Alignment). Other algorithms then do the **SNP calling**. They take into account the **coverage** (# of reads per base), base and mapping qualities as well as many other factors.

5.3.5 Gene expression differences

Differentially expressed genes (DEGs) can be inferred from RNAseq data. Expression is measured in FPKM, which stands for Fragments (reads) Per Kilobase per Million mapped fragments. It is then corrected for sequencing depth and gene / exon length.

5.4 The question of usefullness

The computational and storage needs when dealing with NGS data are enormous. Whether a few microsatellites are enough is a valid question. SSRs (see chapter 3.2.1) only reflect a limited portion of the genome and have very different mutation rates compared to SNPs and there's also a lot less of the former. Microsatellites in general suffer

from ascertainment bias. It is of statistical nature and is introduced during collection of the data, when only markers that were found to be polymorphic are used. When markers with little variance are ignored, genetic diversity is generally overestimated. Also, rare alleles are often missed which may lead to incorrect inferences of demographic parameters. Additional benefits of whole genome re-sequencing information include

- More than anonymous markers
- Candidate genes can be studied
- Signatures of adaptation / selection can be detected
- Genetic and genomic diversity (e.g. estimates through exome-wide diversity)
- Demographic history

5.5 NGS pro and contra

	Advantages	Disadvantages
Data quantity	huge amounts of data	huge amounts of data
Data quality	high quality	multiple sequencing required; storage costs
Costs	cost per bp relatively cheap	high costs for individual reads; expensive IT infrastructure
Data analyses	almost everything possible	lots of IT infrastructure

6 Genetic variation

The ultimate source of genetic variation are **mutations**. They occur at random positions in the genome but rates can vary across genomes. In sexual life cycles, existing genetic variation is being re-shuffled continuously through random gamete fertilization, random chromosome segregation and recombination.

6.1 Forms of genetic variation

6.1.1 Singe-nucleotide polymorphism

SNP refers to variation in a single nucleotide at a specific position in the genome. Adjacent nucleotides or indels (insertions and deletions) can have substantial effects on SNP mutation rates. Roughly 15% of all polymorphisms are small indels.

6.1.2 Structural variation

- Balanced nucleotide variation
 - Inversion
 - Intrachromosomal translocation
 - Interchromosomal translocation (up to whole arms of chromosomes)
- Unbalanced nucleotide variations
 - Duplication
 - Deletion

6.1.3 Sizes of variations

• SNP: 1bp

- Microsatellites and minisatellites: 14-200bp
- Indels: <1kb
- Copy number variations (CNVs): >1kb

6.1.4 Other forms of genetic variation

- Gene expression variation
- Methylation variation
- Post-transcriptional modification

6.2 Levels of genetic variation

The main axes of variation in diversity are among species and within genomes. The neutral theory of molecular evolutions postulates that the vast majority of evolutionary change at the molecular level is maintained by the interaction between mutation, which creates variation, and genetic drift, which eliminates it. It also predicts, that in a population of constant size, diversity should be proportional to N_e . The neutral theory is usefull as a null hypothesis for test whether natural selection is occuring.

6.2.1 Population size

In an idealized, panmictic population, also known as Wright-Fisher population, with an equal expected contribution of individuals to reproduction and equal survival, the strength of genetic drift is inversely proportional to the size of the population. Real populations depart from the concept. Therefore the following two concepts are used. The census population size N_c is the number of individuals in a population. It varies by several orders of magnitude across taxa. The effective population size N_e is the size of an idealized population that would show the same amount of genetic diversity as the population of interest. It varies over time, with long-term N_e explaining current levels of genetic diversity in populations but contemporary N_e explaining how strong drift currently is.

The observed differences in population size are expected to determine differences in genetic diversity across species. However, across-species variation in genetic diversity is much narrower than the variation in abundance. This conflict has been termed the **paradox of variation** by Lewontin and is also known as **Lewontin's paradox**. Possible solutions include

- Demographic fluctuations
- Natural selection and genetic hitchhiking
- Molecular constraints on heterozygosity (in yeast, recombination is impeded when heterozygosity is too high)
- Variation in mutation rate

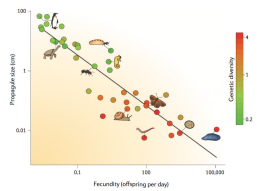


Figure 1 | Genetic diversity and the r/K gradient in animals. The average per-day fecundity is on the x axis and the average size of eggs or juveniles is on the y axis; each dot is for a family (one to four species each). The colour scale indicates the average nucleotide diversity at synonymous positions, expressed in per cent. The negative correlation reflects a trade-off between quantity and size of offspring, -strategists (bottom right; for example, blue mussels, heart urchins and lumbricid earthworms) are more polymorphic than K-strategists (top-left; for example, penguins, Galapagos tortoises and subterranean termites). Figure from REF. 37, Nature Publishing Group.

6.3 Determinants of genetic variation

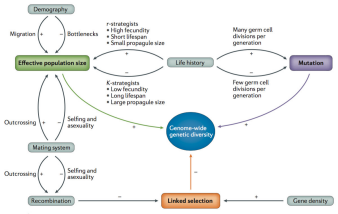


Figure 2 | **Overview of determinants of genetic diversity.** Effective population size, mutation rate and linked selection are the main factors affecting diversity. These factors are in turn governed by several other parameters. The direction of correlation is indicated by the + and - symbols. Selfing, self-fertilization.

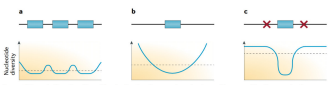
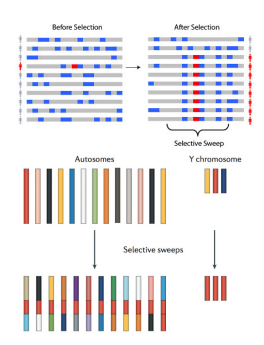


Figure 3 | Genetic diversity affected by the density of targets for selection and by recombination rate. A schematic illustration of the effects of linked selection on genetic (nucleotide) diversity around genes or other functional elements (boxes: upper panels). In the lower panels, solid lines indicate the local variation in diversity level and dashed lines indicate the average diversity in the whole region in question. In regions with a high density of targets of selection (part a), linked selection is pervasive and significantly reduces diversity compared with regions with a lower density of selection targets (part b). When the recombination rate is high (part c), the effect of linked selection becomes less prevalent, allowing maintenance of high diversity levels.

6.3.1 Selective sweeps

Selective sweeps happen, when a beneficial allele carries other neutral alleles close to it along through hitchhiking. It results in less diversity in the beneficial alleles vicinity.

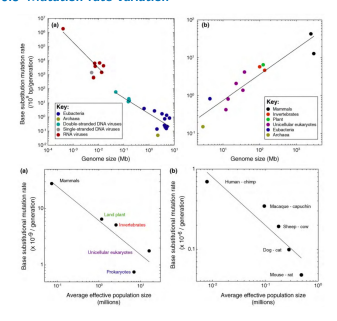


6.4 Loss of variation

Relative population size	Rate at which variation is lost each generation
Haploid	$1/N_e$
Diploid Tetraploid	$1/(2N_e)$
Tetraploid Plastid DNA	$\frac{1/(4N_e)}{1/N_{ef}^*}$
mtDNA	$1/N_{ef}^*$

*True for taxa in which plastid DNA (including cpDNA) and mtDNA are maternally inherited, since N_{ef} is the effective number of females in the population.

6.5 Mutation rate variation



Mutation accumulation experiments generate multiple lines of one acestor, that reproduce through selfing or inbreeding to accumulate mutations. They are useful to estimate mutation rates and mutation variation.

6.5.1 Direct sequencing of families

Analysis of Genetic Inheritance in a Family Quartet by Whole-Genome Sequencing (Roach et al, Science (2010), 328, 636-639.) We analyzed the whole-genome sequences of a family of four, consisting of two siblings and their parents. Family-based sequencing allowed us to delineate recombination sites precisely, identify 70% of the sequencing errors (resulting in > 99.999% accuracy), and identify very rare single-nucleotide polymorphisms. We also directly estimated a human intergeneration mutation rate of approximately 1.1×10^{-8} per position per haploid genome. Both offspring in this family have two recessive disorders: Miller syndrome, for which the gene was concurrently identified, and primary ciliary dyskinesia, for which causative genes have been previously identified. Family-based genome analysis enabled us to narrow the candidate genes for both of these Mendelian disorders to only four. Our results demonstrate the value of complete genome sequencing in families.