# RAD-seq in Roscoff

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# Mini-workshop about ddRAD

### Introduction about RAD-seq

- ► RAD? RAD-seq? ddRAD?
- Applications
- Workflow

### **Practicals**

- ▶ One complete project, from raw reads to final results
- Cherry-picking of some analysis steps
- Open questions

### **Objectives**

- Overview of RAD-seq
- Arouse curiosity
- ► Give useful pointers

## Disclaimer about the speaker!

- Not a population geneticist, not a bioinformatician
- Evolutionary biologist who dropped into a RAD-seq project when he was a small post-doc
- Some things said here are probably incorrect or plainly wrong!

## What are RAD markers?

### Miller et al. 2007

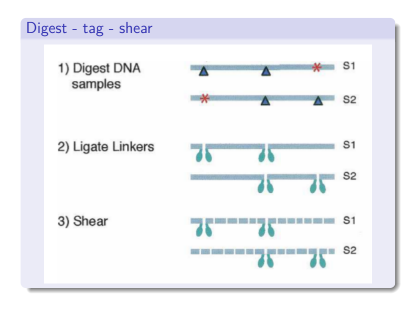
Rapid and cost-effective polymorphism identification and genotyping using restriction site associated DNA (RAD) markers

Michael R. Miller, <sup>1</sup> Joseph P. Dunham, <sup>2</sup> Angel Amores, <sup>3</sup> William A. Cresko, <sup>2</sup> and Eric A. Johnson<sup>1,4</sup>

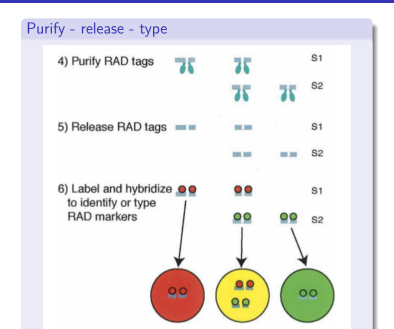
### Description of RAD markers

- Restriction site associated DNA fragments
- Used with micro-array systems
- ▶ Similar to RFLP or AFLP, but many more markers

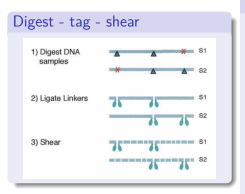
# RAD - Miller et al. 2007 (6 steps)

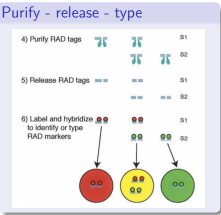


# RAD - Miller et al. 2007 (6 steps)



# RAD - Miller et al. 2007 (method summary)





### Demonstration

- Mapping breakpoint on a Drosophila chromosome
- ▶ Identification of the lateral plate locus in threespine stickleback

## RAD - Miller et al. 2007

## Advantage of the method

- ► Easy-to-produce genotyping resource for non-model species
- ► Moderate cost
- ► Genetic mapping possible (if markers location known)
- Bulk genotyping possible

### But note that...

- ▶ At this point the restriction site is the polymorphic marker
- ▶ One restriction enzyme only is used

## What is RAD-seq?

### Baird et al. 2008

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# Rapid SNP Discovery and Genetic Mapping Using Sequenced RAD Markers

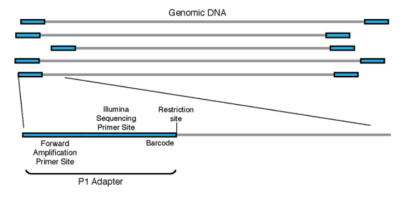
Nathan A. Baird<sup>1</sup>, Paul D. Etter<sup>1</sup>, Tressa S. Atwood<sup>2</sup>, Mark C. Currey<sup>3</sup>, Anthony L. Shiver<sup>1</sup>, Zachary A. Lewis<sup>1</sup>, Eric U. Selker<sup>1</sup>, William A. Cresko<sup>3</sup>, Eric A. Johnson<sup>1</sup>\*

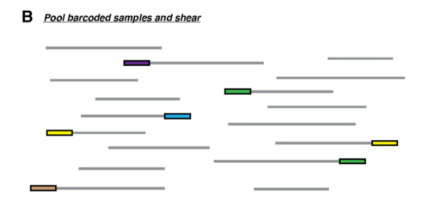
1 Institute of Molecular Biology, University of Oregon, Eugene, Oregon, United States of America, 2 Floragenex, Eugene, Oregon, United States of America, 3 The Center for Ecology and Evolutionary Biology, University of Oregon, Eugene, Oregon, United States of America

### RAD-seq

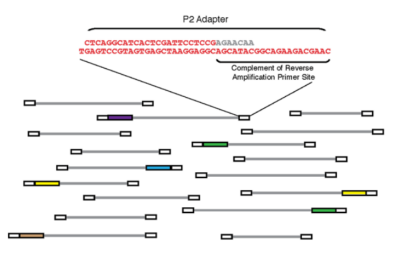
- ► RAD fragments with high-throughput sequencing (Illumina)
- ► SNP identified by sequence polymorphism and site disruption
- Can be used with or without reference genome

#### A Ligate P1 Adapter to digested genomic DNA

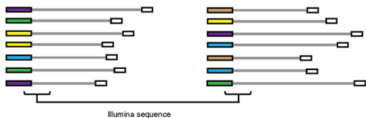




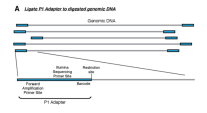
### C Ligate P2 Adapter to sheared fragments



D Selectively amplify RAD tags



llumina sequence read length

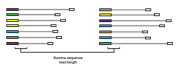






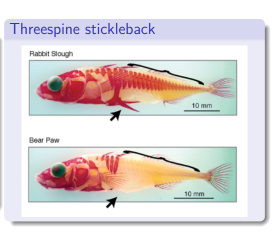






#### Demonstration

- Discover 13000 SNP in threespine stickleback and in Neurospora
- Barcoding system for multiplexing
- Marker density can be tuned by the choice of restriction enzyme



# Population genomics of parallel adaptation - Hohenlohe 2010

### A major paper

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PLOS GENETICS

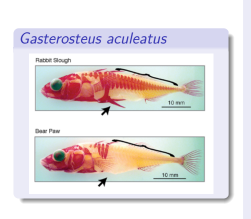
## Population Genomics of Parallel Adaptation in Threespine Stickleback using Sequenced RAD Tags

Paul A. Hohenlohe<sup>1</sup>\*, Susan Bassham<sup>1</sup>\*, Paul D. Etter<sup>2</sup>, Nicholas Stiffler<sup>3</sup>, Eric A. Johnson<sup>2</sup>, William A. Cresko<sup>1</sup>\*

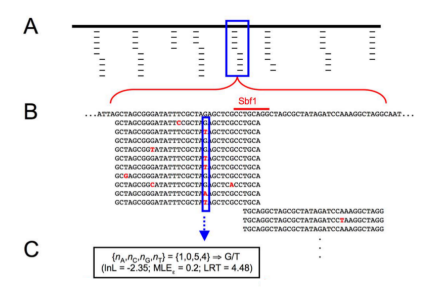
### Method

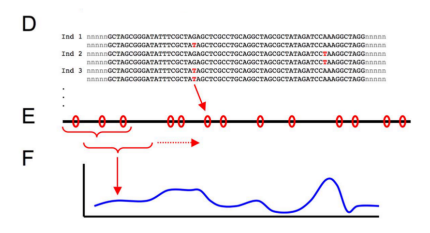
- ► Model: threespine stickleback
- Comparison of 3 freshwater and 2 marine populations
- 20 individuals per population, individual barcodes
- Single reads (not paired ends)

# Population genomics of parallel adaptation - Hohenlohe 2010

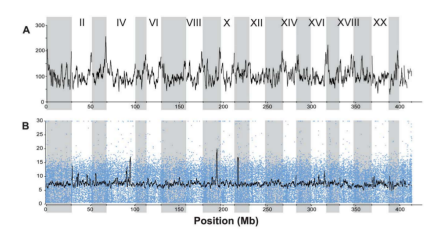




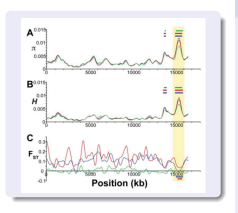




# Hohenlohe 2010 - Genome profiles



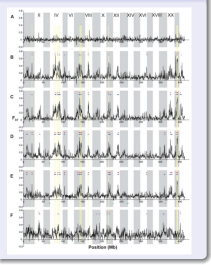
- ► A: number of RAD tags per 1Mb
- ▶ B: Coverage per RAD per individual in one run (16 individuals black line is average)



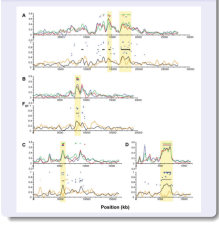
# Evidence for balancing selection

- A: Nucleotide diversity, B: heterozygosity across all five populations (blue), three FW (red) or two SW (green)
- C: Fst between FW and SW (blue), among FW (red) and among SW (green)
- Horizontal bars shows regions of significantly elevated or reduced values on the profile

# Genome-wide differentiation among populations



# Differentiation among SW and FW, zoom on LG



## Highlights

- ► RAD-seq on natural populations, 45000 SNPs in 100 individuals
- ► Barcoded samples
- ► Genome profiling, kernel smoothing and permutation testing

### But note that...

- ► Genome available
- ► Single reads

## What is paired-end RAD-seq?

#### Etter 2011

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# Local *De Novo* Assembly of RAD Paired-End Contigs Using Short Sequencing Reads

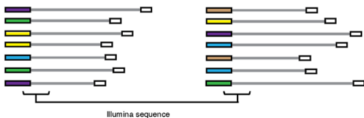
Paul D. Etter<sup>1</sup>, Jessica L. Preston<sup>1</sup>, Susan Bassham<sup>2</sup>, William A. Cresko<sup>2</sup>, Eric A. Johnson<sup>1</sup>\*

#### Method

- ▶ Paired-end sequencing of RAD fragments to build contigs on the randmoly sheared side
- ▶ Demonstration with threespine and *E. coli* sequencing
- Up to 5kb contigs with circularization step

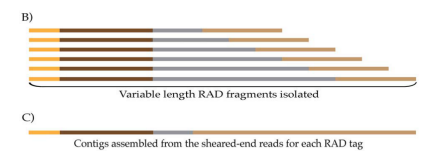
## Single-reads RAD-seq

D Selectively amplify RAD tags



llumina sequence read length

# Paired-ends RAD-seq



### **Notes**

- ► The stacked end is useful for high coverage work (SNP calling, allele frequency estimates)
- ► The echelon end is useful for contig building, but base coverage is lower

## What is double-digest RAD-seq?

### Peterson et al. 2012

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Double Digest RADseq: An Inexpensive Method for *De Novo* SNP Discovery and Genotyping in Model and Non-Model Species

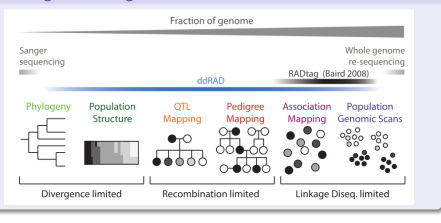
Brant K. Peterson\*, Jesse N. Weber, Emily H. Kay, Heidi S. Fisher, Hopi E. Hoekstra

### Method

- Two enzyme double digest followed by precise size selection
- Library contains only fragments close to target size
- Read counts across regions are expected to be correlated between individuals

## Peterson 2012

### Double digest RAD tag



## What is paired-end double RAD?

### Bruneaux et al. 2013

Molecular evolutionary and population genomic analysis of the nine-spined stickleback using a modified restriction-site-associated DNA tag approach

MATTHIEU BRUNEAUX,\*1 SUSAN E. JOHNSTON,\*1 GÁBOR HERCZEG,† JUHA MERILÄ,† CRAIG R. PRIMMER\* and ANTI VASEMÄGI\*‡

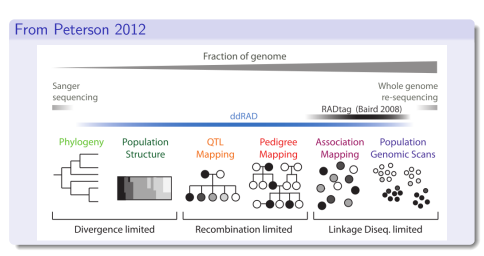
### Method

- Two enzyme double digestion
- Paired-end sequencing on after size-selection
- ► You will hear more about it soon (see practicals)

## Paired-end double RAD

Add a picture

## Uses of RAD tags



## Uses of RAD tags

## Population genomics

Hess 2013 - Pacific lamprey

Phylogeography

Emerson 2011

QTL mapping

Houston 2012

Phylogenies

Rubin 2012

## There are also some potential issues...

- PCR-duplicates
- individual vs pool genotyping for allele frequencies
- Comparison SNP vs microsat (deFaveri)

### Conclusion

#### In a nutshell

- ► RAD tags: versatile method of genome complexity reduction
- ▶ RAD-seq: large scale discovery of SNPs, affordable
- Useful for both model and non-model organisms
- Just a tool: the downstream analyses are still your expertise

## General workflow scheme

# Development of pipelines and tools

 ${\sf Rainbow,\ STACKS,\ GATK,\ dDocent}$ 

## Tools for NGS can be used for RAD

## Simple scripts can be used also

This is one thing I want to show during the practical Get a good grip and a good feeling/understanding about the data with simple, straightforward methods before choosing to apply more complex methods which rely on third-party scripts. It is important to understand what the third party scripts do!

# One complete project

## Tour of other tools and specific analyses

To illustrate some specific points (e.g. likelihood or bayesian based genotyping or allele frequency estimates or Fst calculations, . . . )