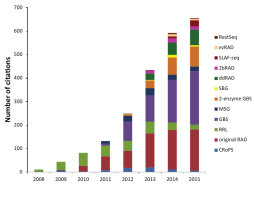
Intro

Alexander Jueterbock

2015-05-30

RAD-Seq - young and successful NGS methods

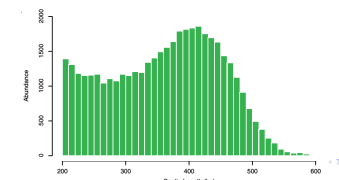


Purpose of RAD-seq

- Genome-reduction method to fragments adjacent to restriction enzyme recognition sites.
- Increases depth of coverage per locus compared to whole genome sequencing
- High-throughput genotyping of populations (multiplexing using barcoding) at relatively low cost.
- Makes genome-scale population genetic studies possible for non-model species lacking a reference genome.

Reductive *de novo* genome sequencing and SNP identification

- RAD-Seq of the sunflower genome (Illumina)
 - 44.7M reads (PE:40bpx80bp)
- De novo assembly of ca. 15.2 Mb in >42,000 contigs
- Identified >94,000 putative SNPs across six lines



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Genome-wide association study (GWAS)

■ No reference genome previously available

Intro

- identified >100,000 SNPs across 138 genotypes
- Related SNPs to 17 phenotypic traits in a field trial
- Increasing flexibility and speed of crop breeding



Figure: Miscanthus sinensis



References

Population genomics and parallel adaptive differentiation in threespine sticklebacks

■ Reference genome available

Intro

- >45,000 SNPs across 100 individuals ('genotyping by sequencing')
- Consistent signatures of selection between two oceanic and three freshwater populations
- Identified 31 candidate genes of evolutionary significance

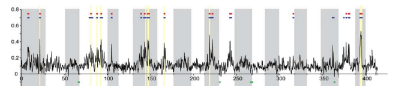


Figure: F_{ST} for SNPs in sliding windows across the genome between oceanic and freshwater populations

Original RAD-Seq protocol

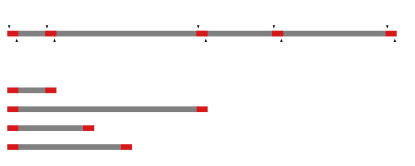
- Developed by (Baird et al., 2008; Miller et al., 2007).
- DNA fragments adjacent to restriction enzyme recognition sites



5' GAATTC 3' 3' CTTAAG 5'

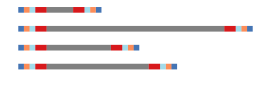
EcoRI recognition site

Step 1: cut DNA



 Note: Bias in GC content of restriction site samples the genome non-randomly

Step 2: ligate P1 adapter



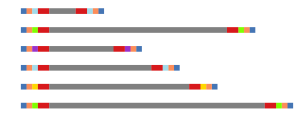
Amplification primer site

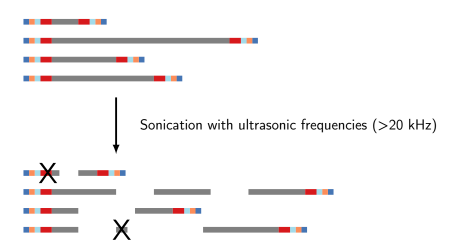
Sequencing primer site (Illumina-specific)

Barcode

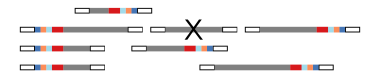
RAD-Seq methods

Barcoding allows to pool samples





Step 4: Ligation of P2 adapter with 'Y' structure



P2 adapter: AGATCG TCTAGCGTCCT

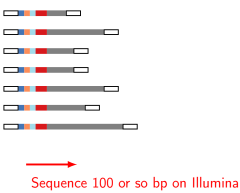
P2 primer: TCTAGCGTCCT

Intro

P2 primer binds only when P2 primer site was completed by amplification starting from the P1 adapter (removes Y-structure)

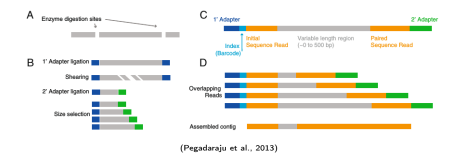
Intro

Step 5: Sequence amplified reads on Illumina

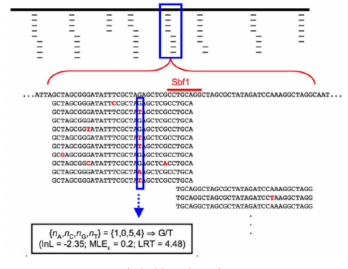


Random shearing of 3'ends helps to detect PCR duplicates

Paired-end sequencing of RAD-tags allows for *de novo* genome sequencing



Calling SNPs from RAD-tags

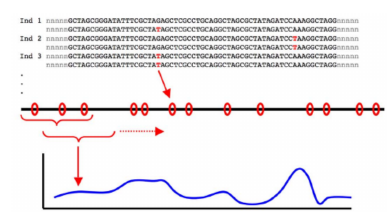


(Hohenlohe et al., 2010)

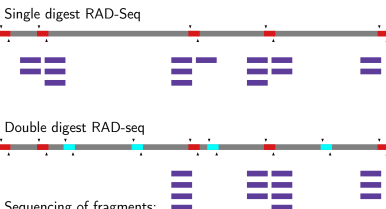


RAD-Seq methods

Summary statistics (e.g. population differentiation) along sliding windows



(Hohenlohe et al., 2010)



- Sequencing of fragments:
- within a specific size range - flanked by two different cutting sites
 - EcoRI recognition site
 - Sbfl recognition site



Pipelines

RAD-Seq methods

ddRAD compared to single-digest RAD sequencing

1 Rapid and 'cheap' protocol (8 hrs hands-on): Doesn't require difficult and high cost of shearing and enzymatic end-repair.

Pipelines

RAD-Seq methods

ddRAD compared to single-digest RAD sequencing

2 Lower number of loci but increased coverage and, thus, higher chance to target the same loci in different individuals.

Intro

RAD-Seq methods

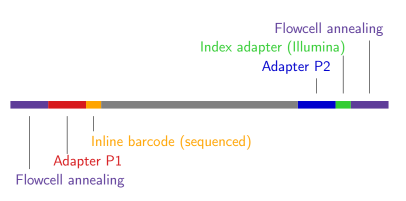
3 Coverage expected to be equal among individuals and highest for fragment lengths targeted by size selection.

RAD-Seq methods

ddRAD compared to single-digest RAD sequencing

Pipelines

4 Combinatorial indexing allows to multiplex more individuals (up to 12 barcodes were affordable for single-digest RAD-Seq).



 $48 \times 12 = 576$ (multiplexing level)

Intro

RAD-Sea methods

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added first, with ligation of adapters, allows to pool samples added second, with PCR primer, allows to combine multiple pools

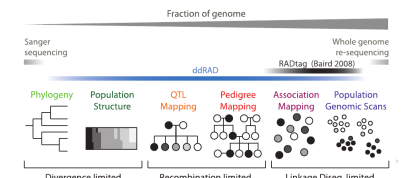
Pooling recommendations

- Critical: equimolar concentrations of individuals expected
- Recommended: >40 individuals/pool
 - Higher numbers
 - + decrease unequal representation of individuals in the pool
 - make it more more difficult to discriminate minor allele frequencies from sequencing errors

Great adjustability of the number of markers makes ddRAD suitable for a broader range of approaches than RAD-Seq

Number of markers adjusted by:

- Cutting frequency of restriction enzymes
- Size selection



How to predict the number of fragments

Based on our own study on Guppy

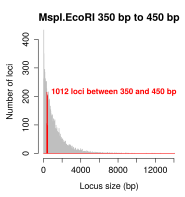
- Targeted coverage: 20x per individual
- Pooling: 60 individuals
- Sequencing output: 24M reads (12M fragments, minimum for Illumina v2 paired-end kits)
- Fragments per individual: 12M/60 = 200,000
- Target: 10,000 fragments (to reach a 20x coverage)

What combination of restriction enzymes to use to obtain the appropriate cutting frequency?

In silico genome digestion

Intro

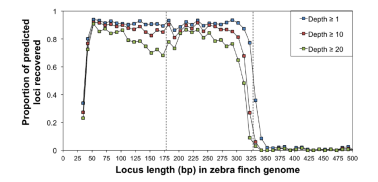
Simulate restriction enzyme digestion with the R package simRAD (Lepais and Weir, 2014)



Based on 10% of the entire genome size

Without reference genome: evaluate double-digest fragments on

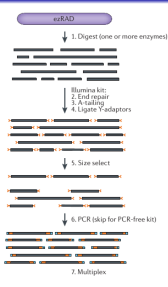
RAD-Seq methods



(DaCosta and Sorenson, 2014)

Targeted: 178-328bp, but short restriction fragments (38-178 bp) were carried through the agarose gel size selection step

ezRAD (Toonen2013)



ezRAD (Toonen2013)

Advantages

RAD-Seq methods

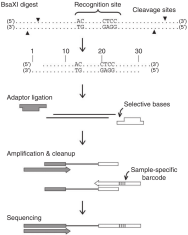
non-PCR kits can avoid PCR duplication and bypass any PCR bias.

Disadvantages

- All reads start with the same four bases (GATC).
 - Low diversity libraries can lead to poor read quality on Illumina sequencers. Use e.g. PhiX spiking or dark-cycling.

2bRAD (Wang2012)

- Type IIb restriction endonuclease to excise 36-bp fragments.
- Number of loci customized by base-selective adapters.



2bRAD (Wang2012)

Advantages

Intro

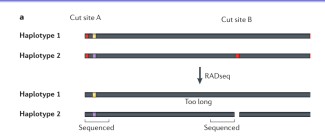
- Extremely simple and cost-effective: no purification or size selection.
- No biases due to fragment size selection.
- Sequencing either strand of the restriction fragments allows for the use of strand bias as a quality filtering criteria.

Disadvantages

- 36-bp tags could be too short to be non-ambiguously mapped in highly duplicated genomes.
- Likely not cross-mappable across large genetic distances.



Null alleles

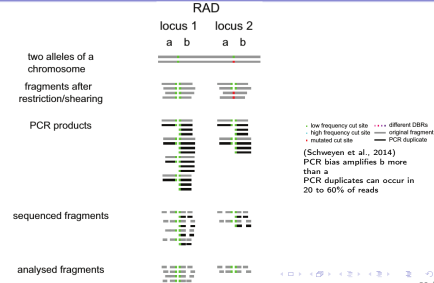


Null allelles due to allele dropbout (Andrews2016)

- Inflate homozygosity
- Underestimate genetic differentiation
- More frequent in longer cutting enzyme recognition sites
- More frequent in ddRAD than in original RADseq
- Identifiable by high variance in coverage among samples

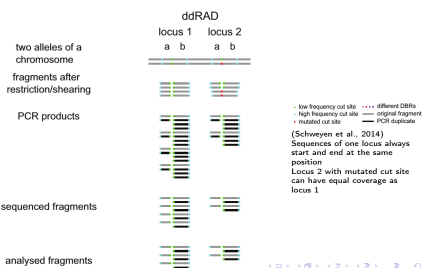
Detect PCR duplicates by length in paired-end RAD

sequencing



Intro

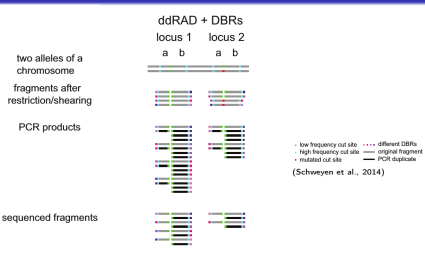
PCR duplicates in ddRAD and ezRAD - not detectable



References

analysed fragments

Degenerate base regions detect PCR duplicates in ddRAD ((Tin2014; Schweyen et al., 2014))

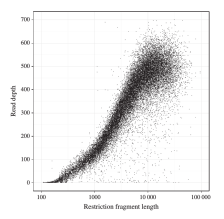


- Reduce occurrence by lowering PCR steps
- Avoid PCR duplicates in ezRAD with Illumina PCR-free kits

Shearing introduces bias in coverage

Intro

Bias in sequencing depth towards larger fragment sizes



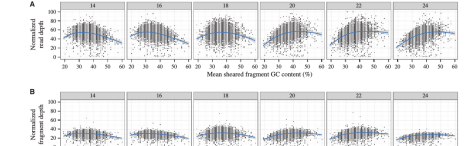
(Davey et al., 2013)

Reason: Fragments of <10 kb shear with lower efficiency



Amplification bias in favor of high GC content

Read depths are influenced by GC content and number of PCR cycles, with (A) or without PCR duplicates (B).



(Davey et al., 2013)

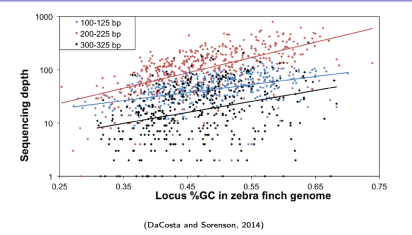
Mean sheared fragment GC content (%)

Modifications of PCR enrichment can help (see (Benjamini2012; Puritz et al.,



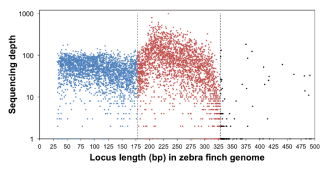


Sequencing depth bias in favor of loci with high GC content



 Combined with a GC-rich recognition sequence, this can result in an overrepresentation of GC-rich portions of the genome

Amplification and, thus, depth decreases with fragment length



(DaCosta and Sorenson, 2014)

- Affects ddRAD more than RAD-seq (each locus different fragment lengths) or 2bRAD (all loci same fragment length)
- Bias reduced by precise size selection (Pippin Prep instrument) (DaCosta and Sorenson, 2014).

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RAD-Sea methods

STACKS (Puritz et al., 2014b)

Stacks: Building and Genotyping Loci De Novo From Short-Read Sequences

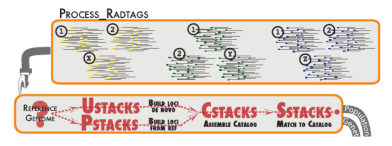
Julian M. Catchen,* Angel Amores, Paul Hohenlohe, William Cresko, and John H. Postlethwait, 1 *Center for Ecology and Evolutionary Biology and †Institute of Neuroscience, University of Oregon, Eugene, Oregon 97403

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RAD-Sea methods

STACKS - basic pipeline for RAD-Seq

STACKS - software pipleine to build loci from RADseg reads and use them for population genomics and phylogeographic analyses.



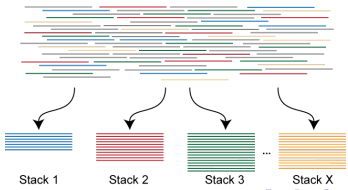
(Catchen et al., 2013)

STACKS - Ustacks de novo assembly step 1

- Only exact matches are assembled
- Secondary reads are set aside

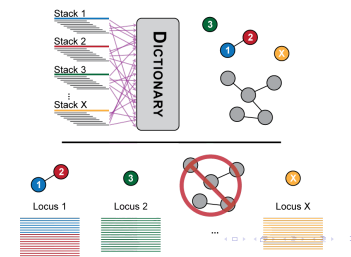
Intro

■ The minimum stack depth parameter controls the number of raw reads required to form an initial stack



STACKS - Ustacks de novo assembly step 2

- Stacks with few nucleotide differences are merged.
- Repetitive sequences with many alleles are excluded

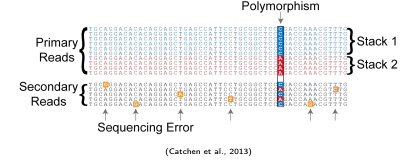


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STACKS - Ustacks de novo assembly step 3

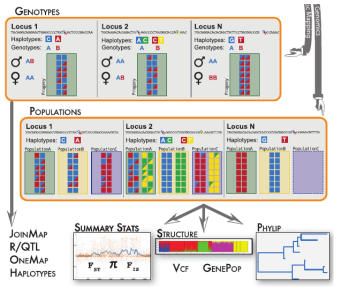
Intro

- Alignment of secondary reads (those not indcluded in stacks) against stacks.
- Alleles are discriminated from sequencing errors by their frequency.



Intro

STACKS - populations or genotypes pipeline



DDocent (Puritz et al., 2014b)

Peer.

Intro

dDocent: a RADseq, variant-calling pipeline designed for population genomics of non-model organisms

Jonathan B. Puritz, Christopher M. Hollenbeck and John R. Gold Marine Genomics Laboratory, Harte Research Institute, Texas A&M University-Corpus Christi, Corpus Christi, TX, USA

DDocent (Puritz et al., 2014b)

Uses stand-alone software packages to perform

quality trimming

Intro

- adapter removal
- de novo assembly of RAD loci
- read mapping
- SNP and InDel calling
- data filtering.

Identifies more SNPs at a higher coverage than STACKS, due to

- simulatneous use of forward and reverse reads during alignment to reference instead of clustering
- quality trimming instead of removing entire reads



MOLECULAR ECOLOGY RESOURCES

Intro

Molecular Ecology Resources (2015) 15, 1163-1171

doi: 10.1111/1755-0998.12378

AftrRAD: a pipeline for accurate and efficient *de novo* assembly of RADseq data

MICHAEL G. SOVIC.*+ ANTHONY C. FRIES* and H. LISLE GIBBS*+

*Department of Evolution, Ecology, and Organismal Biology, Aronoff Laboratory, The Ohio State University, 318 W. 12th Ave, Columbus, OH 43210, USA, †Ohio Biodiversity Conservation Partnership, Aronoff Laboratory, The Ohio State University, 318 W. 12th Ave, Columbus, OH 43210, USA

Bioinformatics Advance Access published March 20, 2014

BIOINFORMATICS ORIGINAL PAPER

2014, pages 1-6 doi:10.1093/bioinformatics/btu121

Phylogenetics

Intro

Advance Access publication March 5, 2014

PyRAD: assembly of de novo RADseq loci for phylogenetic analyses

Deren A. R. Faton 1,2

¹Committee on Evolutionary Biology, University of Chicago, 1025 E. 57th St. Chicago, IL 60637, USA and ²Botany Department, Field Museum of Natural History, 1400 S. Lake Shore Dr. Chicago, IL 60605, USA

Associate Editor: David Posada

RAD-Sea methods

Important considerations

Pipelines

- Degraded DNA interfers with cutted DNA in methods with enzyme-unspecific adaptors
- Higher amount of starting DNA can reduce number of PCR cycles and thus minimize PCR duplicates.
- RADseq libraries are low-diversity libraries as they all start with the same cutting site and can cause problems in cluster generation for Illumina sequencing.
 - Solution: Reduce cluster density and spike-in PhiX control

RAD-Sea methods

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RAD-Sea methods

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RAD-Sea methods

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