

RADseq Works in Primates, Dammit.

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Abstract

... (This is the blurb from the email.) Our paper is an introduction to a 2nd generation sequencing technique for typing thousands of genome-wide markers from non-model organisms. Though it's been used in other taxa, this would be the first published application to primates. We demonstrate it with six Cercopithecoids and discuss its promise for doing mutli-locus population genetics in primates. ...

1 Introduction

- Next-gen sequencing revolution promises gains in primatology
- Still expensive
- Many genomes, but still tough doing genomics on non-model organisms
- What is RADseq?
- Previous RADseq studies
- Why would it be good for primates
- PRESENT STUDY
 - We did RADseq on 6 Cercopithecoids
 - Assessed how well it worked
 - Show it has promise for primates

2 Methods

Library Preparation and Sequencing Genomic DNA from 6 primates was digested with *PspXI* (New England Biolabs) and used to create a multiplexed RAD tag library. Our library preparation method followed that of Etter et al, 2011 with the following modifications: the P1 adapter top(?) oligonucleotide was modified to have an overhang corresponding to the cut site of *PspXI*, and a longer P2 adapter suitable for paired end sequencing was used (P2_top: 5'-SEQUENCEHERE-3'; P2_bottom: 5'-SEQUENCEHERE-3'). Individual-specific barcodes contained in the P1 adapter differed by at least three nucleotides. We chose *PspXI* based on the results of *in silico* digestion of the human, rhesus macaque, and baboon reference genomes using custom Perl scripts (refs). We sequenced the prepared library as one 150-cycle paired-end run of an Illumina MiSeq at the NYU Langone Medical Center's Genome Technology Center using a spike-in of 30% PhiX DNA to control for low diversity in the library at the barcode and restriction sites. Other individuals were sequenced alongside those of the present study. Sequences are available to download from the NCBI Short Read Archive (accession number SRXXXXXXX.X).

Sequence Analysis Sequence reads were demultiplexed, or separated by barcode, and reads without an expected barcode or an intact restriction enzyme cut site were excluded from the analysis. Reads were then aligned to the rhesus macaque reference genome (v.1.0, Mmul_051212/rheMac2, ref) using BWA with default parameters. Reads that were unmapped, unpaired, duplicates, or that had low mapping quality were removed after alignment using Picard (ref) and BamTools (ref).

After performing local realignment around indels with GATK (ref), SNPs and short indels were identified using SAMtools mpileup and BCFtools (ref). A minimum coverage of 3 reads and a maximum of 100 was required to call a SNP or an indel at a given location. Orthologous SNPs were tallied using VCFtools (ref).

To assess how many restriction sites were successfully sequenced and to analyze the degree of overlap between multiplexed individual's datasets, we first found all possible *PspXI* cut sites using the oligoMatch utility in the UCSC Genome Browser program (ref). This allowed us to calculate the coverage of these restriction site-associated regions using BEDtools' multiBamCov program (ref).

Analysis Pipeline - Inferring Phylogeny

- Using method like cichlid people?
- Using method like Rubin et al?
- Concatenated SNPs (not indels) into alignment.
- Dimensions = 6 x ???

3 Results

6.1 million sequencing reads with an intact barcode and restriction enzyme cut site could be assigned confidently to one of the six Old World monkeys in the present study. Roughly 2.0 million of those reads were successfully mapped to the rhesus macaque genome and passed all quality control filtration steps. Relative to the reference genome, our study identified 531,175 SNPs and 24,260 small indels among all samples. Information for each individual is summarized in Table 1.

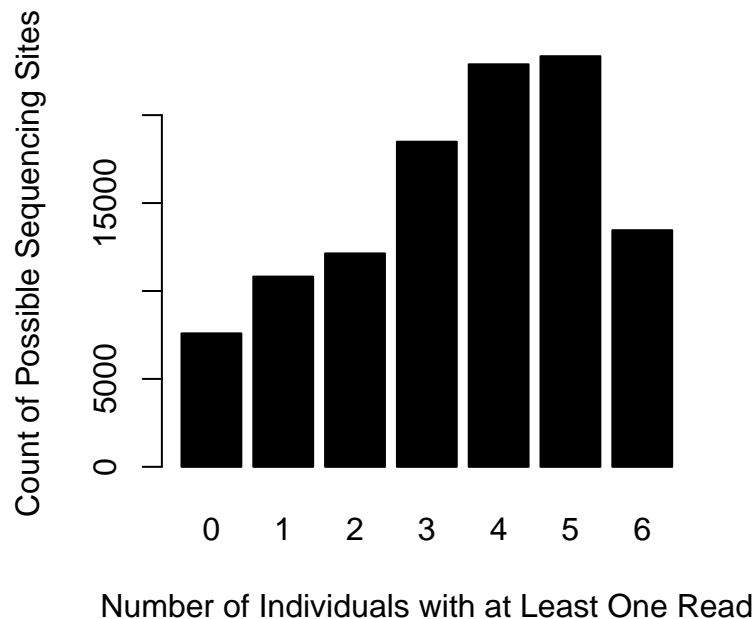
Table 1: Individual Sample Information

Taxon	Source	# Reads	# Filtered Reads	# Loci \geq 1 Read	# Loci \geq 3 Reads	# SNPs
<i>Semnopithecus entellus</i>	Unknown	1,367,050	306,248	47,516	24,052	363,501
<i>Allenopithecus nigroviridis</i>	Unknown	1,325,558	447,279	62,824	40,244	356,750
<i>Macaca mulatta</i>	Unknown	481,376	234,659	77,798	32,302	279,350
<i>Cercocebus torquatus atys</i>	Unknown	297,290	124,700	48,440	13,104	319,766
<i>Papio anubis?</i>	Unknown	743,556	313,676	74,582	39,062	316,118
<i>Theropithecus gelada</i>	Unknown	1,911,030	603,024	68,440	50,656	366,097

In the rhesus macaque genome, we found 54,364 possible cut sites for *PspXI* and 108,728 possible sequencing sites (two per cut site, one upstream

and one downstream). Of those, 101,138 sites (93.02%) were covered by at least one read in at least one individual, and for 13,456 sites (12.38%), all six individuals had at least one read (Figure 1). When we restrict the analysis to sites with at least three reads, 80,448 sites (73.99%) were covered in at least one individual and 1,354 sites (1.25%) had all six individuals present.

- Pairwise comparisons between individuals? Like X orthologous regions shared between Papio and Thero?



- SNP info from merged analysis
- SNP Venn diagram?
- Count orthologous SNPs shared between individuals. Pairwise?

4 Discussion

- RADseq is viable tool for researcher interested in primate phylogenetics, pop. gen.

- Cheap and easy to create libraries. Competent lab can do it in two days. (Ha!)
 - Multiplexing is great for pop gen studies.
 - Enzyme choice allows control over coverage, number of individuals, number of loci.
- Potential problems with RADseq method
 - Promise for primatology

5 Acknowledgements

- NYU Med School folk
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