

What are SNPs?

- **S**ingle **N**ucleotide **P**olymorphisms
 - Differences between individuals of a species
 - EuPath: differences between strains / isolates (if clonal)
 - Some organisms are diploid so will also have allelic SNPs within strain
 - Genes that are different due to SNPs are alleles.
 - Our model does not include insertions/deletions (indels) currently

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tgondii_gt1_chr      ATTCGATGCGCAGAGGAGGAACTTACAGAGACGGAGCGGCACTGAAGCTTTTGCCAAAGAC
tgondii_veg_chr      ATTCGATGCGCAGAGGAGGAACTTACAGAGACGGAGCGGCACTGAAGCTTTTGCCAAAGAC
tgondii_me49_chr    ATTCGATGCGCAGAGGAGGAACTTACAGAGACGGAGCGGTACTGAAGCTTTTGCCAAAGAC 1129631
neospora_chr         ATTCGCTGCGCAGAAGAAGAGCTGCAAAGACGCAGCGGCACCGAGGCGTTCGCCAAAGAC
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tgondii_gt1_chr      TTACTTCTCCTCCTTGTCGGGGCTGAGGCCTCTTCCGCTGCGAAACAGGCTGGTAAGGCCG
tgondii_veg_chr      TTGCTTCTCCTCCTCGTCGGGGCTGAGGCCTCTTCCGCTGCGAAACAGGCTGGTAAGGCCG
tgondii_me49_chr    TTGCTTCTCCTCCTCGTCGGGGCTGAGGCCTCTTCCGCTGCGAAACAGGCTGGTAAGGCCG 1129571
neospora_chr         CTTCTCCTCCTCCTCGTCGGGGCAGACGCGTCGCCTGCTGCGAAACAGGCTGGTAAGCCA
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tgondii_gt1_chr      GCGGCGACGA---AGGGTGGCTCTGAA-----GAGC
tgondii_veg_chr      GCGGCGACGA---AGGGTGGCTCTGAA-----GAGC
tgondii_me49_chr    GCGGCGGGCGACGAAGGGTGGCTCTGAA-----GAGC 1129540
neospora_chr         CCCGCGGGCGGACGGACGTCGCGCGCACGCGAAGGCGAGAAAAAGGGGAAGCGTTTGAGC
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SNPs in EuPathDB are derived from two sources

- Chip based assays
 - Arrays are designed that allow identification of SNP alleles given a DNA sample. There are multiple different arrays in PlasmoDB. Also Barcode assays (24 SNPs)
 - Isolate DNA is then assayed on these arrays.
- Direct deep sequencing of DNA from isolates.
 - Reads are aligned to reference genome and SNPs called.
- What are Isolates – show on webpage

How are resequencing SNPs called in EuPathDB

- Reads (hopefully paired end) are retrieved from provider (best if from SRA)
- Reads are aligned to the reference using Bowtie2 (end-to-end)
- Reads realigned around indels using GATK
- SNPs, indels and consensus sequence generated using VarScan (min read depth 5, min read frequency 20%)
- SNPs based on this reference comparison are stored in the DB
- Every isolate alignment is checked for every SNP position and if sufficient evidence to make “like reference” call then this is indicated in the DB along with evidence for call.
- In the end, have allele(s) for every isolate at every position if the evidence from the alignment warrants it.
- <http://tinyurl.com/pebcdlz> (SNP record page - link to alignment)

Homozygous / Heterozygous SNPs

- Why do we care?
- Ploidy of organism is critical
 - Apicomplexans are haploid for majority of cycle
 - Tryps, Amoebas, etc are diploid (or worse)
- Why does this matter for SNP calling/queries?
 - Read frequency is the defining parameter
- What does a heterozygous SNP look like?
 - <http://tinyurl.com/o3tr9ly> – record page
 - <http://tinyurl.com/prffbf6>

What can we do with SNPs?

- SNPs are genetic markers
 - Distinguish specific strains / isolates.
 - Enable fine structure mapping of phenotypes in genetic crosses or association studies.
- Identify SNPs based on a variety of characteristics.
 - Within a group of isolates (includes allele frequency and confidence parameters)
 - Restrict to location (on chromosome or within genes)
 - Compare two groups of isolates to identify SNPs that distinguish the two groups.
- Identify Genes
 - Identify genes that appear to be under selection based on SNP characteristics.
 - Number of SNPs (coding, non-coding, synonymous etc)
 - Ratio of non-synonymous / synonymous indicates whether genes are under purifying or diversifying (balancing) selection.

Purifying vs. Diversifying selection

- Purifying selection (gene is evolutionarily constrained to maintain the primary amino acid sequence)
 - Genes that have a low Non-synonymous / Synonymous ratio
 - Tend to be genes critical for basic metabolic processes such as enzymes, cell cycle related etc.
 - Due to very high A/T bias in *falciparum*, the ratio of synonymous/non-synonymous can be skewed due to severe codon bias.
- Diversifying selection (it is evolutionarily advantageous to quickly change the amino acid sequence)
 - Genes that have a high Non-synonymous / Synonymous ratio.
 - Tend to be things like surface antigens that the organisms use to escape immune detection.
 - Use comparators high on the list (have more sequence coverage and thus more SNPs). *Reichenowi* frequently is not a good comparator because these genes are changing so rapidly that they may not be conserved well enough to call SNPs.