Inferring the selective pressures acting on insertions and deletions in the great tit genome

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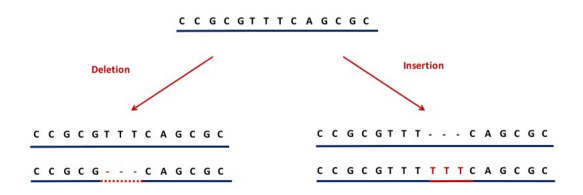
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Insertions and deletions

▶ short INDELs: sections of DNA < 50bp that are deleted or inserted in a genome



INDELs often overlooked

- Disproportionately occur in repetitive sequence
- Hard to align
- ▶ Often occur in hotspots
- ▶ 1/8 as frequent as SNPs in humans

(Earl et al., 2014; Montgomery et al., 2013)

The importance of INDELs in genome evolution

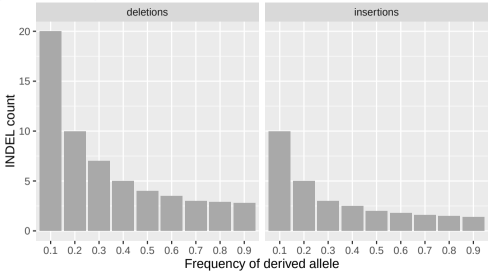
- ► Contribute more to sequence divergence, in terms of the number of base differences, than SNPs
- ► Influence genome size:
 - ▶ low deletion rate → large genomes?
 - ▶ high deletion rate → compact genomes?

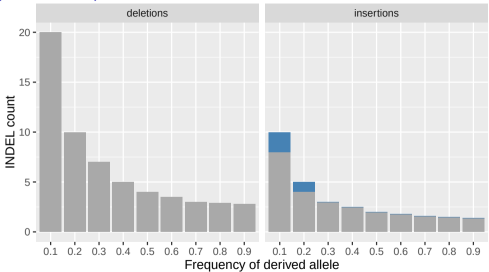
(Britten, 2002; Nam and Ellegren, 2012; Ometto et al., 2005; Sun et al., 2012)

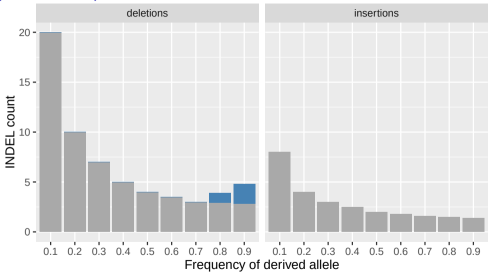
INDEL selection

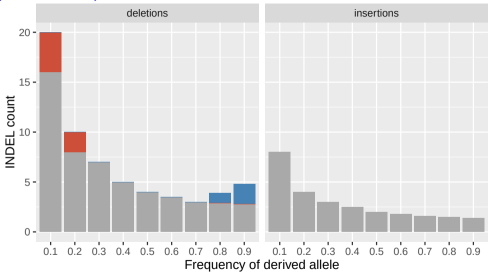
- Deletions
 - lower mean allele frequency
 - more deleterious
 - two breakpoints
- ▶ Insertions may be favoured:
 - elevated fixation probability
 - biased gene conversion
 - minimum intron size
- Polarisation error

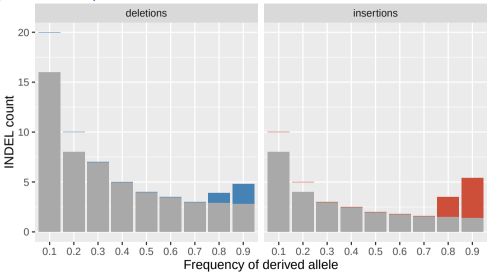
(Leushkin and Bazykin, 2013; Ometto et al., 2005; Petrov, 2002; Sjödin et al., 2010)

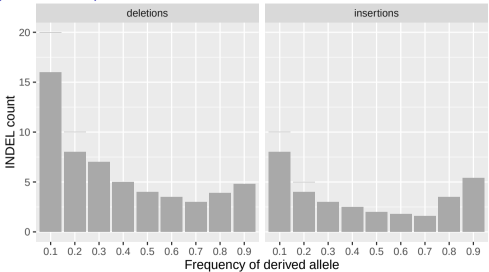












Aims

Overcome confounding affect of polarisation error

Quantify how natural selection shapes INDEL diversity in the great tit (Parus major)

- 1. within coding regions
- 2. in non-coding regions

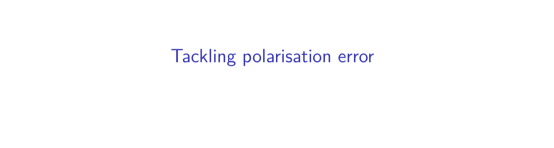


Advantages of an avian system

- Conserved karyotype and synteny good for alignments
- Genomes consist of few large macrochromosomes and many small microchromosomes
- ▶ Results in a highly dynamic recombination landscape power to associations with recombination



(van Oers et al., 2014; Stapley et al., 2008)

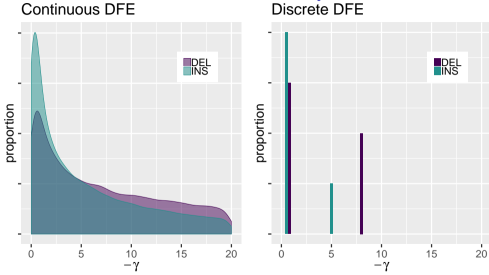


A novel maximum likelihood approach

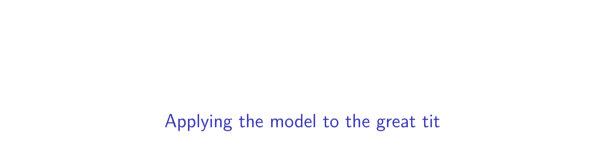
- 'anavar'
- takes the unfolded site frequency spectrum
- estimates for both insertions and deletions:
 - mutation rate $(\theta = 4N_e\mu)$
 - the distribution of fitness effects (DFE)
 - polarisation error (ϵ)
- ► Controls for demography using neutral sites (Eyre-Walker et al., 2006)
- ► Applicable to both INDELs and SNPs or a combination

(Barton and Zeng, MBE, 2018)

The model can describe the DFE in two ways



(Barton and Zeng, MBE, 2018)

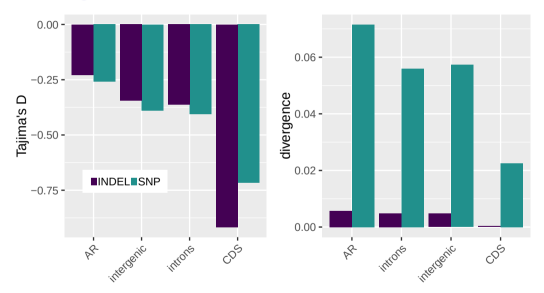


Sample and pipeline

- ▶ 10 european great tit males (Corcoran et al., 2017)
- ► high coverage (44x)
- ▶ variant calling with GATK
- multispecies alignment between zebra finch, flycatcher and great tit
- parsimony based polarisation

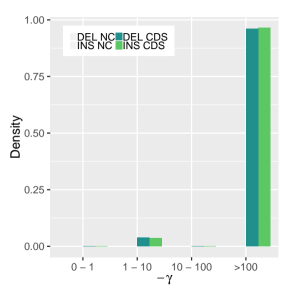


Summarising the data set

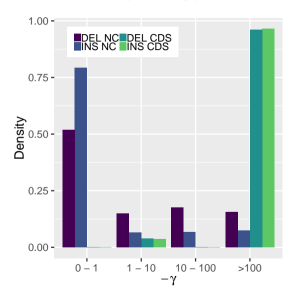




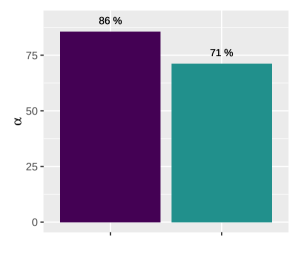
Polymorphic INDELs predominantly strongly deleterious

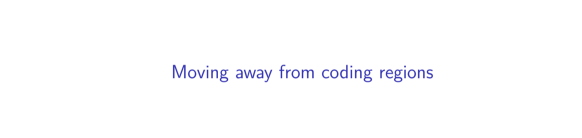


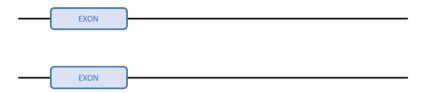
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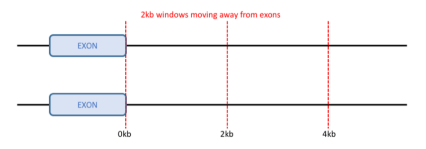


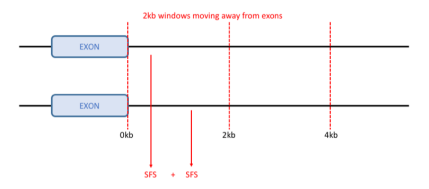
Coding INDEL fixations largely beneficial

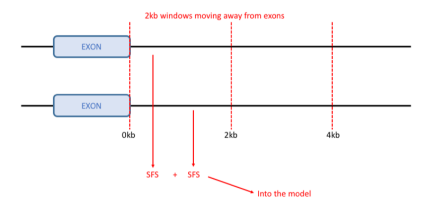




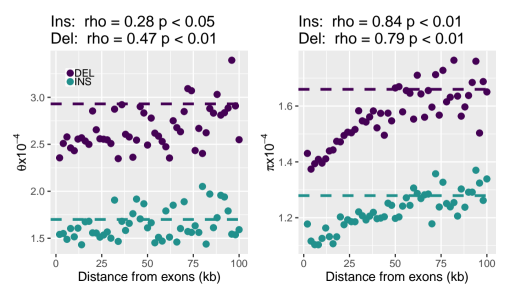


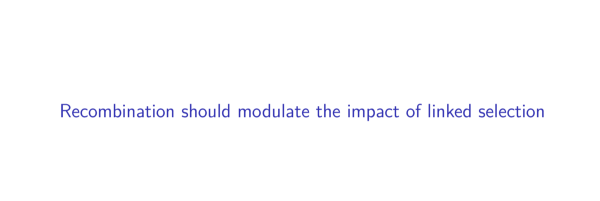




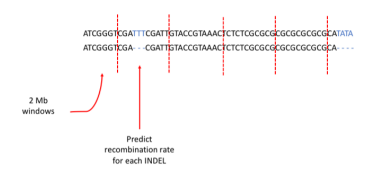


Diversity increases with distance

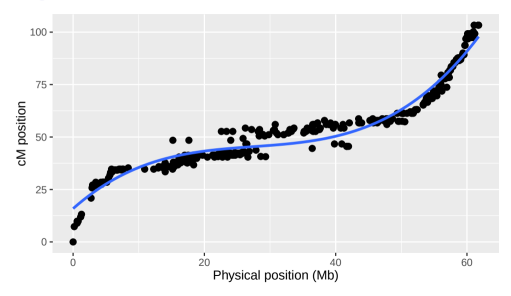


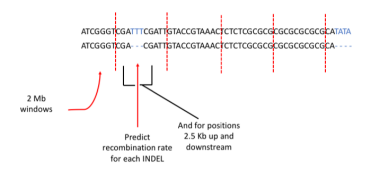


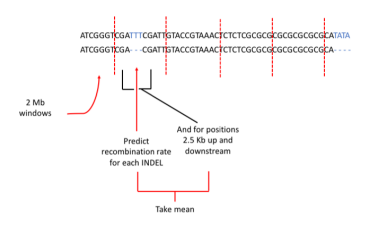


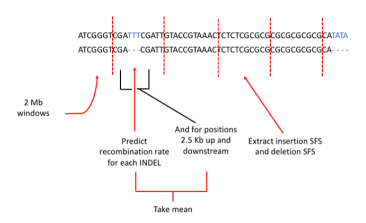


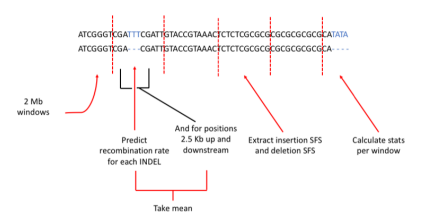
Getting the data - recombination rate



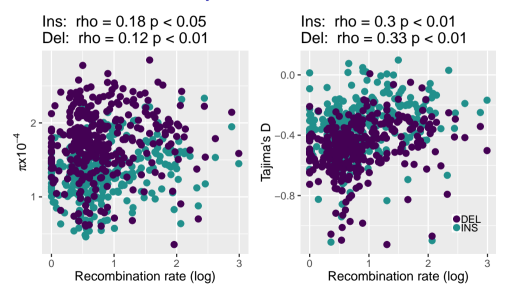








Association between diversity and recombination





Conclusion

- ▶ INDELs in genes mostly extremely deleterious 96%
- ► Remainder are weakly deleterious
- ightharpoonup lpha estimate at 71% and 86% for insertions and deletions
- Regions adjacent to exons, and areas of low recombination, have reduced INDEL diversity - linked selection
- ► Extends over relatively large distance up to ~50kb

Next steps

- ► Interesting to investigate if reduced diversity is due to positive selection or purifying selection

 ► Lock at whather efficient of selection on INDEL a is higher in regions with higher N
- ightharpoonup Look at whether efficacy of selection on INDELs is higher in regions with higher $N_{
 m e}$

Plug for the model

- ▶ User friendly computer package anavar http://zeng-lab.group.shef.ac.uk
- ▶ Methods are applicable to both INDELs and SNPs or a combination
- Code for integration with python https://henryjuho.github.io/anavar_utils/

