12/15/2019 GEDmatch

GEDmatch® Autosomal One-to-one Comparison - V1.0

Software Version Nov 12 2019 14:54:30

Here is a link to a useful YouTube video on using the One to One DNA comparison tool.

			tions:

Base Pairs with Full Match	
Base Pairs with Half Match	
Match with Phased data	
Base Pairs with No Match	

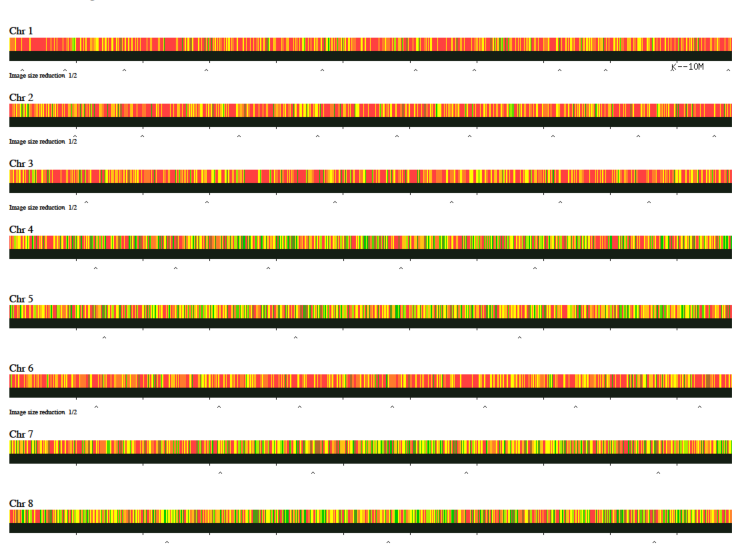
Validity of segments:

Significant	
Moderate	
Low	
Insignificant	
Large gap between adjacent SNPs	
No Match	
Centromere	

Comparing Kir (*target.stripy.genome.2) [23andMe] and (*22stripy.genome.missing) [23andMe]

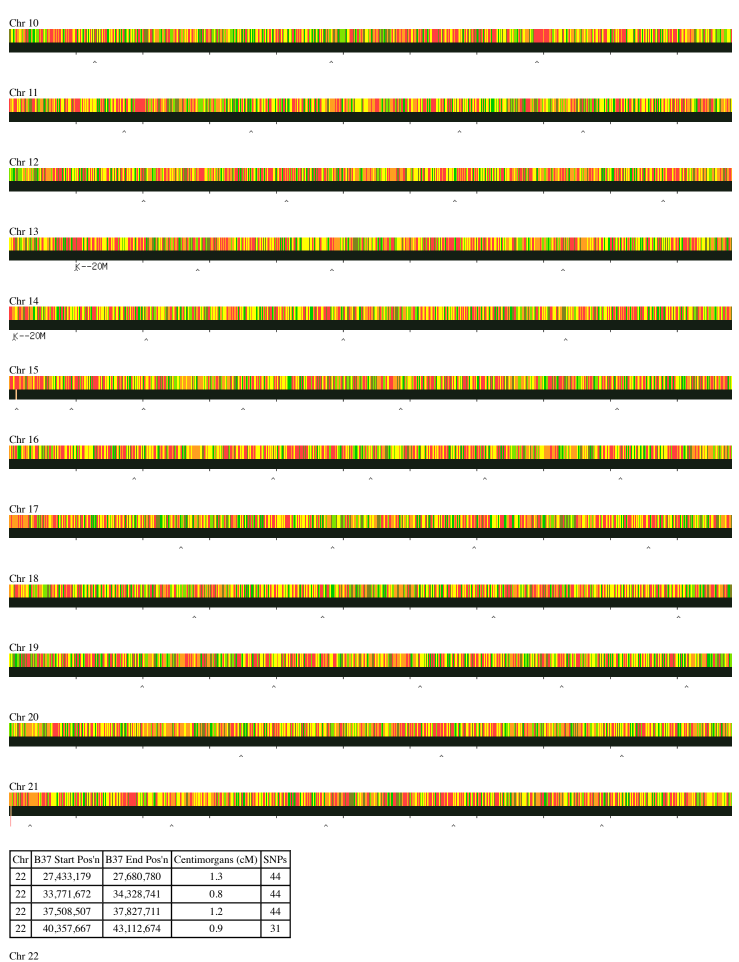
Minimum segment threshold size to be included in total = 25 SNPs Minimum segment cM to be included in total = 0.7 cM

Mismatch-bunching Limit = 25 SNPs



Chr 9

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Largest segment = 1.3 cM

Total Half-Match segments (HIR) = 4.1 cM (0.115 Pct)

4 shared segments found for this comparison.

514759 SNPs used for this comparison.

18.410 Pct SNPs are full identical

Comparison took 0.343 seconds. CPU time used: 0.156 cpu seconds.

Ver: Nov 12 2019 14:54:30