HOW Q-MATCH COMPARES WITH 2021 AND 2013 GEDMATCH ANALYTICS IN THE CONTEXT OF 3CM MATCHES

In our 8 by 8 study, we noted that between 2021 and 2023, the default GEDmatch analytics for autosomal matching at the 3cM threshold changed significantly. GEDmatch then suggested to us that its Q-match technique should be used for small matches. Here we report our experience of using Q-match for small matches.

Q-match has a parameter P (for 'Precision') which can be adjusted downward to get more matches, albeit of lower quality, than the default setting P = 7 provides. Accordingly, for purposes of comparison, we used the same core 6 relatives and randoms we identified in our 8 by 8 study, which is where the lower half of the following table came from. In its totality the table illustrates what we found thus allowing a comparison between Q-match and its precursors.

	Qmatch P=7		Qmatch P=3		Comparison	
	matches #	multiples #	matches #	multiples #	matches	multiples
6 core relatives 6 core randoms	115 120	12 10	405 337	83 67	up 3.5 times up 2.8 times	up 6.9 times up 6.7 times
	2021 analytics		2023 analytics			
6 core relatives 6 core randoms	103 118	17 13	171 150	34 18	up 66% up 27%	up 2 times up 1.4 times

The first pair of columns shows 2021 analytics beneath P = 7 results. We can see that while Q-match selects matches better in that there are more matches but fewer multiples – most of the multiples are still inconclusive with just 2 multiples - the Cr2 and Cr8 indicator segments - being the difference between the family and the random sets. The inconclusive multiples are mainly weak and strong triples for both relatives and randoms. However, this result still clashes with John Griffiths' intuition in that the family set continues to yield fewer matches than the random set.

Selecting lower Precision rectifies this, for in the second pair of columns, which show 2023 analytics beneath P = 3 results, we see that Q-match produces many more matches with the family set ahead in both matches and multiples. But here, the large number of multiples presents a challenge in sorting out the useful from the inconclusive. In fact, at P = 3 we produce so many matches that the number of multiples is constrained by their coverage of a large fraction of the genome – the third pair of columns shows this slowdown in rate of increase of frequency.

Our method for doing this sorting is to look for rare coincident segment boundaries (RCSBs) and abutting segment boundaries (ASBs). During the present study we considered afresh the frequency at which RCSBs and ASBs occur at random.

These events occur when a terminal SNP of one match coincides with a terminal SNP of another match. Lefthand RSBCs, righthand RSBCs, and ASB's are generated at the same rate.

On average there are 50,000 SNPs per chromosome. If there are N matches on a chromosome, there are N*(N-1)/2 pairs of matches. Thus, there are N*(N-1)/100,000 possible instances of coincident segment boundary SNPs on a chromosome. Using the observed average number of matches on each chromosome, we find the estimated frequency of random occurrence of boundary SNPs to be:

Group	Matches on a chromosome	Frequency on a chromosome	Frequency on the genome
6 core randoms	15.3	0.0022	0.049
6 core relatives	18.4	0.0032	0.070
12 relatives	89.2	0.0786	1.730

In this study we found 1 RSBC and 2 ASBs in the random set – probably reflecting close family relationships previously unnoticed. We found zero in the 6-relative set, but 22 lefthand RSBCs, 19 righthand RSBCs, and 17 ASBs in the 12 relative set. While these 3 latter incidences seem only about 10 times expectation, it should be remembered that the 3-person version of RSBC is quite common (but worthless genealogically), making the useful 4-person version much less likely than the 1.73 figure indicates.