Reconstructing CNV genotypes using segregation analysis: combining pedigree information with CNV assay

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Abstract

Background: Repeated blocks of genome sequence have been shown to be associated with genetic diversity and disease risk in humans, and with phenotypic diversity in model organisms and domestic animals. Reliable tests are desirable to determine whether individuals are carriers of copy number variants associated with disease risk in humans and livestock, or associated with economically important traits in livestock. In some cases, copy number variants affect the phenotype through a dosage effect but in other cases, allele combinations have non-additive effects. In the latter cases, it has been difficult to develop tests because assays typically return an estimate of the sum of the copy number counts on the maternally and paternally inherited chromosome segments, and this sum does not uniquely determine the allele configuration. In this study, we show that there is an old solution to this new problem: segregation analysis, which has been used for many years to infer alleles in pedigreed populations.

Methods: Segregation analysis was used to estimate copy number alleles from assay data on simulated half-sib sheep populations. Copy number variation at the Agouti locus, known to be responsible for the recessive self-colour black phenotype, was used as a model for the simulation and an appropriate penetrance function was derived. The precision with which carriers and non-carriers of the undesirable single copy allele could be identified, was used to evaluate the method for various family sizes, assay strategies and assay accuracies.

Results: Using relationship data and segregation analysis, the probabilities of carrying the copy number alleles responsible for black or white fleece were estimated with much greater precision than by analyzing assay results for animals individually. The proportion of lambs correctly identified as non-carriers of the undesirable allele increased from 7% when the lambs were analysed alone to 80% when the lambs were analysed in half-sib families.

Conclusions: When a quantitative assay is used to estimate copy number alleles, segregation analysis of related individuals can greatly improve the precision of the estimates. Existing software for segregation analysis would require little if any change to accommodate the penetrance function for copy number assay data.

Background

With the increasing resolution at which genomes can be examined has come the recognition that variation in genome structure is common and affects more nucleotides per genome than the sequence variation found in single nucleotide polymorphisms (SNP) [1-3]. Copy number variation (CNV) in DNA, defined as insertions, deletions and duplications larger than 1 kb, is an important component of this structural variation. Recent publications document the contribution of CNV to genetic diversity in humans [2,4-6] and human disease [7-9]. CNV has been shown to contribute to phenotype in model organisms [9-11] and to important production and disease traits in domesticated livestock species [12-15]. Current technologies to assay a CNV genotype (genome copy number) and its corresponding alleles have limitations [8,16]. Distinguishing among genomes that have multiple DNA copies (> 4-5 copies) is imprecise while SNP that might ‘tag’ copy number alleles through linkage disequilibrium are usually only found for relatively simple diallelic CNV [2,3,6,17,18]. Typically, assays attempt to quantify the total number of
copies in diploid DNA and cannot discriminate between, for example, an individual homozygous for a two-copy allele and a heterozygous individual carrying one- and three-copy alleles. In many cases, this is not an important limitation as copy number alleles have additive dosage effects, but in other cases it is important, for example when only one copy number allele is associated with disease. To resolve individual alleles, data on related individuals can be analyzed concurrently. Pedigree information applies Mendelian constraints to the allowable sets of copy number alleles in related individuals. These constraints have been exploited using Bayesian graphical models [19] to infer copy number alleles, and hidden Markov model based methods [20,21] to find de novo CNV and infer copy number alleles.

The Mendelian constraints underlying analyses such as those noted above have been well studied in the area of segregation analysis. Originally the term “segregation analysis” referred to the determination of the mode of inheritance of a phenotype but in recent decades it has come to include the inference of genotypic probabilities in pedigreed populations. The peeling algorithm of Elston and Stewart [22] is applicable in both cases. In small pedigrees, and in those without inbreeding loops, this algorithm produces unbiased estimates of allelic probabilities. A number of approaches have been described that are computationally feasible for larger pedigrees with inbreeding loops, including iterative peeling [23,24], cutting loops [25] and Markov Chain Monte Carlo (MCMC) methods [26,27]. MCMC methods remain an active area of research (e.g. [28]).

The data required for single locus segregation analysis are four-fold: 1) a pedigree relating individuals to each other; 2) phenotypes on some individuals in the population; 3) a penetrance function expressing the probabilities of phenotype based on genotype; and 4) estimates of the frequencies of the genotype alleles in the population. Probabilities of mutation can also be incorporated. Assays that return estimates of the number of copies in diploid DNA can be thought of as a phenotypic measurement, and if assumptions are made about the distribution of the variation around the expected value of the assay then it is relatively straightforward to derive the appropriate penetrance function. Software to perform computations for segregation analysis is available and may include options to estimate penetrance functions as well as to estimate allelic probabilities (e.g. [29]).

In this paper, we demonstrate that segregation analysis methods are well suited to the inference of copy number alleles, and that if knowledge on the actual allele configurations is sufficiently important then the utility of a quantitative assay can be greatly improved through the incorporation of relationship data. On small datasets, our methods can be implemented in readily available software such as Mendel [29], accounting for the uncertainty in the individual assay results through the penetrance function. With minor modifications, existing software for large datasets could also be used. While we restrict our analyses to half-sib families, our approach is general and could be applied to any data set with both pedigree and quantitative copy number assay components.

**Methods**

**Assay parameters and allele frequency estimation**

The characteristics of the assay are reported in Norris and Whan [13]. In an asymmetric competitive PCR assay, DNA amplified from the junctions between copies is compared to the DNA amplified from the junctions and from the 5' breakpoint region. For diploid DNA with a copy number count $n$, the expected value of the assay is the ratio $n - 2n$, as there are $n - 2$ junctions and two copies of the 5’ breakpoint. As diploid DNA has at least two copies (at least one on each chromosome) this ratio takes values from the series $0/2, 1/3, 2/4, 3/5 ...$ which asymptotically approaches unity. It is important to note that since the assay is quantitative, variation occurs around the expected ratios. The magnitude of the variation can differ among laboratories, and even among batches of samples in one laboratory, so we have treated it as unknown and conducted analyses for a range of values. As a lower limit, we chose a CV of 3% (which equates to a standard deviation of 0.01 for the class with an expected ratio of 1/3). This figure is a little smaller than that derived empirically from ranking 111 assay samples and estimating the mean and standard deviation.
deviation for the 35 samples that were extremely unlikely to be from a class other than that with an expected ratio of 1/3, the most easily distinguished class. The CV of 3% was used as the lower limit since it is achievable with real data, but we also conducted analyses assuming CV values of 6% and 9%. The expected ratios and standard deviations considered are summarized in Table 1.

From the expected ratios and standard deviations, and assuming a distribution for the variation around the expected value, the probability of each copy number count can be estimated for each observed assay value. We assumed a normal distribution and estimated copy number count probabilities for a population of 87 phenotypically white Merino ewes coming from commercial sale yards. These are summarized in Table 2A. To estimate the allele frequencies that produced this distribution of copy number counts we minimized the $X^2$ statistic obtained by comparing the vector of copy number count frequencies with the frequencies expected given a vector of frequencies for alleles segregating in Hardy-Weinberg equilibrium. Black Merino sheep in wool flocks are commonly culled soon after birth and thus are never presented in a commercial sale yard, so to avoid ascertainment bias we excluded the $A^{c1}/A^{c1}$ class when calculating the $X^2$ statistic. Table 2B contains the estimated allele frequencies.

### Penetrance function

For black sheep, the genotype is known to be $A^{c1}/A^{c1}$ with certainty and conversely, for sheep with genotype $A^{c2}/A^{c2}$ the phenotype is known to be black wool with certainty. The assay is not relevant for these animals. For white sheep, the penetrance function relates to the assay rather than to the phenotype. That is, the penetrance function is the probability of returning a particular assay value given the genotype. Again under the assumption of normality, this is proportional to the height of the normal distribution relevant to the underlying genotype. Table 3 contains an example of the mapping of an assay ratio of 0.675 onto a grid representing the penetrance function for an individual, using standard deviations for a CV of 3%.

### Simulation

As the stud Merino sheep industry is likely to use a test for the genotype at the Agouti locus, small, highly selected and relatively closed flocks were simulated. In each replicate, 10 studs were simulated, each mating five rams to 200 ewes each year. Selection was on a trait uncorrelated to Agouti, except for the last generation when no homozygous $A^{c1}/A^{c1}$ animal was selected. In the absence of evidence to the contrary, we assumed that the genotype at the Agouti locus had no effect on fitness other than through artificial selection. The simulation was run for 20 years. In the founder populations, allele frequencies at the Agouti locus were as in Table 2B. There was limited exchange of genetics between

### Table 1 Expected ratios and standard deviations of the assay

| Copies | Expected Ratio | Standard Deviation of Assay |
|--------|----------------|-----------------------------|
|        |                | CV = 3% | CV = 6% | CV = 9% |
| 2      | 0.000          | NA      | NA      | NA      |
| 3      | 0.333          | 0.010   | 0.020   | 0.030   |
| 4      | 0.500          | 0.015   | 0.030   | 0.045   |
| 5      | 0.600          | 0.018   | 0.036   | 0.054   |
| 6      | 0.667          | 0.020   | 0.040   | 0.060   |
| 7      | 0.714          | 0.021   | 0.043   | 0.064   |
| 8      | 0.750          | 0.023   | 0.045   | 0.068   |
| 9      | 0.778          | 0.023   | 0.047   | 0.070   |
| 10     | 0.800          | 0.024   | 0.048   | 0.072   |
| 11     | 0.818          | 0.025   | 0.049   | 0.074   |
| 12     | 0.833          | 0.025   | 0.050   | 0.075   |

The assay is not required for sheep with two copies as they are phenotypically black and easily identified.

### Table 2 Estimated copy number frequencies (A) and allele frequencies (B) derived from assay results

| Copies | Frequency | Allele | Frequency |
|--------|-----------|--------|-----------|
| 2      | 0.000     | $A^{c1}$ | 0.100     |
| 3      | 0.029     | $A^{c2}$ | 0.430     |
| 4      | 0.344     | $A^{c3}$ | 0.410     |
| 5      | 0.023     | $A^{c4}$ | 0.000     |
| 6      | 0.004     | $A^{c5}$ | 0.020     |
| 7      | 0.002     | $A^{c6}$ | 0.000     |

Data are from 87 white Merino ewes, and a CV of 3.0% for the assay scores was assumed.

### Table 3 Copy number probabilities and penetrance values for an assay value of 0.675

| Copies | Probability |
|--------|-------------|
| 3      | 0.000       |
| 4      | 0.000       |
| 5      | 0.000       |
| 6      | 0.838       |
| 7      | 0.159       |
| 8      | 0.003       |
| 9      | 0.000       |

Penetrance Values

| Copy number allele from dam | $A^{c1}$ | $A^{c2}$ | $A^{c3}$ | $A^{c4}$ | $A^{c5}$ | $A^{c6}$ |
|-----------------------------|----------|----------|----------|----------|----------|----------|
| $A^{c1}$ from sire          | 0.000    | 0.000    | 0.000    | 0.000    | 0.838    | 0.159    |
| $A^{c2}$ from sire          | 0.000    | 0.000    | 0.000    | 0.000    | 0.159    | 0.000    |
| $A^{c3}$ from sire          | 0.000    | 0.000    | 0.838    | 0.159    | 0.030    | 0.000    |
| $A^{c4}$ from sire          | 0.000    | 0.838    | 0.159    | 0.000    | 0.000    | 0.000    |
| $A^{c5}$ from sire          | 0.838    | 0.159    | 0.000    | 0.000    | 0.000    | 0.000    |
| $A^{c6}$ from sire          | 0.159    | 0.003    | 0.000    | 0.000    | 0.000    | 0.000    |

The CV of the assay was assumed to be 3%; the penetrance values are proportional to the probability of an assay value of 0.675 given the genotype.
studs, with one outside ram and four home-bred rams used by each stud each year. The intention behind this relatively complicated structure was to simulate populations in which cohorts of ewes were related, with variations in the frequencies of the various Agouti alleles between studs. Parents and progeny of the last generation were assumed to be available for assay. Using the known simulated copy number count, an assay value was simulated using the means and standard deviations described above. The whole simulation was repeated 20 times, producing 1,000 half-sib families (20 replicates × 10 studs × 5 rams) for analysis. The distributions of allele frequencies in the final generations of the simulated populations are displayed in Table 4.

Relationships between assayed individuals
In our simulation study, we performed the assay on half-sib groups of animals and their parents, ignoring relationships between parents. This is equivalent to “cutting” inbreeding loops in a relatively naive but systematic way, rather than the more sophisticated approaches such as in [25]. We chose this simple approach for a number of reasons, but primarily because we performed many analyses on replicated datasets and therefore needed a fast execution time. However, we believe that the design is justified. In commercial and stud sheep flocks, large half-sib families are usual and although deeper pedigree data is usually available (especially on the male side), it may not be without errors. In half-sib families, most of the power in the method comes from the confidence with which the sire’s genotype can be estimated, which then adds confidence to estimates for progeny.

Software and analysis
Genotype probabilities were estimated using a restricted implementation of the Elston-Stewart algorithm [22], restricted in that it operated only on half-sib families. This restriction allowed a very fast execution time enabling the investigation of a wide range of scenarios. The software was validated by comparing the estimated probabilities for test half-sib families to probabilities estimated using the software package Mendel [29]. The allele frequencies assumed for the parents are those shown in Table 2B, that is, in the analysis we used the frequencies that were used in the simulation. Each dataset was evaluated six times, with one of the three formulas for the standard deviation of the assay (CV = 3%, 6%, 9%) used to simulate the assay values, and either the correct CV or one half of the correct CV used for the analysis. From the final generation of the simulated populations, families of 1, 2, 4, 8, 16 or 32 half-sib progeny were chosen. Assay results were made available for either progeny only, progeny and sire, or progeny, sire and dams. Genotype probabilities were estimated and compared to the simulated copy number alleles.

Results
Table 5 contains the results for a lamb analyzed without pedigree data. This is not quite the same as relying on assay results alone to estimate the genotype, as population allele frequencies are still used in the analysis. Without these there is no power at all to declare an animal non-carrier. In all cases, when a lamb was declared to be a carrier with greater than 95% certainty it had the genotype Ac1/Ac2 and in all cases when a lamb had the genotype Ac1/Ag it was declared to be a carrier with over 99.9% certainty.

The effect of jointly analyzing half-sib families of varying size is displayed in Table 6. The analyses correspond to a situation where the sire and his half-sib progeny are assayed, and the coefficient of variation (CV) of the assay output is 3%. The most important application of a test in an industrial genetic improvement program is the identification of sheep that are non-carriers. We found that with a family size of one, only 46% of non-carriers were identified as non-carriers with a 99% probability. However, this value is almost twice that achieved without genotyping the sire, as in Table 5. Increasing the family size improved the power, and with family sizes of 16, 75% of non-carriers can be declared clear with a 99% probability. Increasing the family size beyond 16 has only a small effect on power, unless the goal is to achieve 99.9% probabilities of being clear.

False positives (declaring non-carriers to be carriers) are absent and false negatives (declaring carriers to be non-carriers) are less frequent as family size increases. This is an important result since in industrial

| Table 4 Frequencies of alleles in the final generation of the simulated populations |
|----------------------------------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|
| Frequency Allele | 0.0:0.1 | 0.1:0.2 | 0.2:0.3 | 0.3:0.4 | 0.4:0.5 | 0.5:0.6 | 0.6:0.7 | 0.7:0.8 | 0.8:0.9 |
| Ac1                  | 0.79     | 0.05    | 0.08    | 0.08    | 0.01    | 0.00    | 0.00    | 0.00    | 0.00    |
| Ac2                  | 0.07     | 0.14    | 0.14    | 0.11    | 0.18    | 0.16    | 0.07    | 0.08    | 0.04    |
| Ac3                  | 0.04     | 0.13    | 0.14    | 0.10    | 0.19    | 0.16    | 0.07    | 0.11    | 0.06    |
| Ac4                  | 0.89     | 0.04    | 0.03    | 0.03    | 0.00    | 0.00    | 0.00    | 0.00    | 0.00    |
| Ac5                  | 1.00     | 0.00    | 0.00    | 0.00    | 0.00    | 0.00    | 0.00    | 0.00    | 0.00    |
| Ac6                  | 0.96     | 0.01    | 0.02    | 0.01    | 0.00    | 0.00    | 0.00    | 0.00    | 0.00    |
applications, this second form of error has a much greater adverse impact. The fact that we do not find proportions of type II errors in accordance with appropriate p-value threshold (for example, 5% of type II errors in the case of a P > 0.95 threshold) is not unexpected, just as we do not expect to identify 95% of carriers with a 95% probability.

Table 7 displays results obtained when exploring the effect of assaying progeny only, or progeny and the sire, or progeny, sire and dam. A family size of 16 half-sibs and a CV of 3% for the assay were used. Even if the sire is not assayed, in 62% of replicates a carrier sire is identified with greater than 99% probability. This improves when the sire is assayed, and if the dams are also assayed almost all the carrier sires are detected. The results are more or less symmetrical, in that the probabilities of declaring a non-carrier sire to be a non-carrier are similar to the probabilities of declaring a carrier sire to be a carrier. For the progeny, the power to detect carriers and non-carriers is lower than for sires and even lower for dams, and if the dam is not assayed there is no power at all to declare her a non-carrier.

In Table 8, the effect of the precision of the assay is examined, for family sizes of 16 half-sibs assayed along with their sire. Considering first the situation where the CV used in the analysis is the same as the CV used in the simulation, and again focusing on non-carriers, increasing the CV of the assay decreases the power to declare non-carriers to be clear. This is particularly true if the goal is to achieve a less than 0.1% probability of being a carrier. When an underestimate of the CV is used in the analyses it is not unexpected that the power to declare non-carriers to be clear is improved. This is because reducing the CV reduces the proportion of individuals with ambiguous results, and as most individuals are non-carriers, in most cases overconfidence in the precision of the assay does not result in an error. However, for individuals that are carriers, underestimating the CV results in an increased probability of being declared clear by error.

Discussion
At the Agouti locus in sheep, colour phenotype is affected by a recessive, single copy allele, which cannot always be uniquely identified using the available assay. Furthermore, the assay has almost no power to determine that an animal is free of the undesirable single copy allele. Exploiting the family structures common in sheep flocks, across the scenarios examined, the joint analysis of half-sib families resulted in a modest increase in the power to declare individuals as carriers of the undesirable Ac1 allele. More importantly in a selective breeding environment, including family data resulted in a large increase in the power to declare individuals as non-carriers of the Ac1 allele.

The improvement is dramatic: from 7% of non-carrier lambs being identified as clear at the 99.9% level when a
lamb is assayed and analyzed alone (last row, Table 5), to as many as 80% of non-carrier lambs being declared clear at the same threshold when the sire, dams and lambs for a half-sib family of 16 are assayed and analyzed together (4th last row, Table 7). This is achieved at a cost of assaying 33 related individuals, and looks a better strategy than assaying 32 half-sib lambs and their sire but not their dams (last row, Table 6), especially as ewes are generally used over a number of years and would not need to be re-assayed.

Provided that the precision of the assay is not overestimated (i.e. the CV underestimated) in formulating the penetrance function, the joint analysis of half-sib families does not increase the proportion of false negatives. On the contrary, it reduces the proportion, from 7% at a 5% threshold for lambs analyzed alone, to as few as 1% if 16 half-sibs are analyzed together. As noted earlier, the assignment of a non-carrier status to carriers is most undesirable in industrial applications of the assay. A ram, sold as a non-carrier, that subsequently produces progeny exhibiting the self colour black condition, can adversely affect the reputation of the stud selling the ram, and the reputation of the assay. Furthermore, if the ram is used only in flocks that are clear of the $A^{c2}$ allele, then the undesirable allele can become established in the ewe population. Thus it might take

| Table 7 Frequencies of estimated probabilities of being a carrier |
|----------------------|------------------|------------------|------------------|------------------|------------------|------------------|------------------|------------------|
| Status | Pedigree | Assay | P < 0.001 | P < 0.01 | P < 0.05 | P > 0.95 | P > 0.99 | P > 0.999 |
| carrier sire | p | 0.00 | 0.01 | 0.02 | 0.71 | 0.62 | 0.58 |
| carrier sire | ps | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 |
| carrier sire psd | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 |
| carrier progeny | p | 0.00 | 0.01 | 0.03 | 0.56 | 0.52 | 0.50 |
| carrier progeny | ps | 0.00 | 0.00 | 0.01 | 0.66 | 0.62 | 0.57 |
| carrier progeny psd | 0.00 | 0.00 | 0.02 | 0.73 | 0.71 | 0.68 |
| carrier dam | p | 0.00 | 0.00 | 0.00 | 0.15 | 0.09 | 0.06 |
| carrier dam | ps | 0.00 | 0.00 | 0.00 | 0.26 | 0.22 | 0.15 |
| carrier dam | psd | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 |
| non-carrier sire | p | 0.37 | 0.67 | 0.84 | 0.00 | 0.00 | 0.00 |
| non-carrier sire | ps | 0.84 | 0.93 | 0.96 | 0.00 | 0.00 | 0.00 |
| non-carrier sire psd | 0.98 | 0.99 | 0.99 | 0.00 | 0.00 | 0.00 |
| non-carrier progeny | p | 0.40 | 0.61 | 0.77 | 0.00 | 0.00 | 0.00 |
| non-carrier progeny | ps | 0.63 | 0.75 | 0.81 | 0.00 | 0.00 | 0.00 |
| non-carrier progeny psd | 0.80 | 0.83 | 0.90 | 0.00 | 0.00 | 0.00 |
| non-carrier dam | p | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 |
| non-carrier dam | ps | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 |
| non-carrier dam | psd | 0.58 | 0.69 | 0.85 | 0.00 | 0.00 | 0.00 |

Given actual status (carrier or non-carrier), position in pedigree (sire, dam or progeny), and assay strategy (p = progeny only, ps = progeny and sires, psd = progeny, sires and dams); the assay CV simulated and CV used in the analysis were both 3% and 16 half-sibs in each family were assayed.

| Table 8 Frequencies of estimated probabilities of being a carrier |
|----------------------|------------------|------------------|------------------|------------------|------------------|------------------|------------------|
| Status | CVSim | CVan | P < 0.001 | P < 0.01 | P < 0.05 | P > 0.95 | P > 0.99 | P > 0.999 |
| carrier | 0.030 | 0.015 | 0.00 | 0.01 | 0.02 | 0.68 | 0.65 | 0.61 |
| carrier | 0.030 | 0.030 | 0.00 | 0.00 | 0.01 | 0.66 | 0.62 | 0.57 |
| carrier | 0.060 | 0.030 | 0.01 | 0.02 | 0.04 | 0.61 | 0.58 | 0.55 |
| carrier | 0.060 | 0.060 | 0.00 | 0.01 | 0.03 | 0.55 | 0.51 | 0.48 |
| carrier | 0.090 | 0.045 | 0.01 | 0.02 | 0.06 | 0.56 | 0.53 | 0.51 |
| carrier | 0.090 | 0.090 | 0.00 | 0.01 | 0.04 | 0.46 | 0.38 | 0.25 |
| non-carrier | 0.030 | 0.015 | 0.68 | 0.76 | 0.80 | 0.00 | 0.00 | 0.00 |
| non-carrier | 0.030 | 0.030 | 0.63 | 0.75 | 0.81 | 0.00 | 0.00 | 0.00 |
| non-carrier | 0.060 | 0.030 | 0.48 | 0.64 | 0.74 | 0.00 | 0.00 | 0.00 |
| non-carrier | 0.060 | 0.060 | 0.33 | 0.59 | 0.76 | 0.00 | 0.00 | 0.00 |
| non-carrier | 0.090 | 0.045 | 0.32 | 0.52 | 0.69 | 0.01 | 0.00 | 0.00 |
| non-carrier | 0.090 | 0.090 | 0.14 | 0.44 | 0.69 | 0.00 | 0.00 | 0.00 |

For progeny, given actual status (carrier or non-carrier), CV used in the simulation (CVsim) and CV used in the analysis (CVan); the sire and 16 progeny in each family were assayed.
several generations before this is detected when another carrier ram is used.

In this paper, we have focused on an example from livestock, the self colour black condition in sheep which means we have assumed large half-sib families and restricted ourselves to analyzing these in isolation. These analyses are very quick and the analysis of high-throughput quantitative CNV assay data for half-sib families would be feasible using this method. However, a half-sib family structure is not a requirement for a segregation analysis, and even for family sizes of one or two half-sibs with the sire being also assayed, power was increased to a higher value than when analyzing an individual alone. Readily available segregation analysis software makes optimal use of all pedigree links in estimating allele probabilities, and is suitable for copy number allele calling in human pedigrees and other pedigrees with smaller family sizes.

In our simulation study, we assumed that parameters relating to penetrance function were known. Specifically, we assumed allele frequencies and parameters for a normal distribution for the error associated with the quantitative trait. These were estimated from a sample of unrelated sheep, but could be estimated from the population of interest. This aspect of segregation analysis was not applied in this study, but software such as Mendel [29] can be used for this purpose. In our study, we also investigated an already known copy number variant, identified from an experimental population designed to uncover the cause of the recessive black condition in Merino sheep. In most cases an experimental population will not be available. Using segregation analysis to search for de novo CNV affecting quantitative traits would be similar to using hidden Markov model based approaches [20,21], but from a different statistical perspective. To apply segregation analysis software for general pedigrees to high-density genomic data would likely be computationally prohibitive but software such as that used here for half-sib families might be feasible.

Conclusions
The precision of copy number allele estimates from quantitative assay data in pedigree populations is greatly increased if the pedigree information is used in the estimation, and segregation analysis methods based on the peeling algorithm are well suited to this application. In the case of the Agouti locus and the recessive self-colour black condition, where the purpose of the test is to identify animals to use as parents, the proportion of lambs correctly identified (at the 99.9% level) as non-carriers increased from 7% when the lambs were analysed alone to 80% when the lambs were analysed in families. Any segregation analysis software can be used provided that the appropriate penetrance function is specified.

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Authors’ contributions
JMH wrote the software for simulating and analysing the datasets, carried out the analyses and drafted the manuscript. VAW conducted the molecular genetic studies and contributed to the manuscript. BJN contributed to the design of the study and to the manuscript. All authors have read and approved the final manuscript.

Competing interests
The authors declare that they have no competing interests.

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