

1 Genotype Likelihoods from Reads

$$\begin{aligned}
L(g = 1) &= p(d | g = 1) \\
&= p(d, g = ra | g = 1) + p(d, g = rb | g = 1) + p(d, g = rc | g = 1) \\
&= \sum_a p(d | g = ra) \times p(g = ra | g = 1)
\end{aligned}$$

Here we do not favour any heterozygous genotype, and all have likelihood 1/3. This may be changed to reflect empirical or dbSNP data

$$\begin{aligned}
L(g = 1) &= 1/3 \sum_a p(d | g = ra) \\
&= 1/3 \sum_a p(d | g = ra, \text{ado}) \times p(\text{ado}) + p(d | g = ra, \text{no ado})(1 - p(\text{ado})) \\
p(d | g = ra, \text{ado}) &= p(d | g = ra, \text{drop r}) * p(\text{drop r}) + p(d | g = ra, \text{drop a}) * p(\text{drop a})
\end{aligned}$$

Here we assume either allele is equally likely to be dropped in an ado event and $p(\text{drop r}) = p(\text{drop a}) = 0.5$. This is unlikely to change.

$$\begin{aligned}
p(d | g = ra, \text{ado, drop a}) &= \prod_i p(d_i | g = rr) \\
p(d | g = ra, \text{ado, drop r}) &= \prod_i p(d_i | g = aa)
\end{aligned}$$