

## Mutation and polymorphism calling settings

- $\theta_1$  --consensus-frequency-cutoff  
(DEFAULT = consensus mode, 0.8; polymorphism mode, 0.0)
- $\theta_c$  --polymorphism-frequency-cutoff  
(DEFAULT = consensus mode, 0.2; polymorphism mode, 0.05)
- $S_1$  --consensus-score-cutoff  
(DEFAULT = 10)
- $S_\theta$  --polymorphism-score-cutoff  
(DEFAULT = consensus mode, 10; polymorphism mode, 2)
- $C_1$  --consensus-minimum-coverage-each-strand  
(DEFAULT = 0)
- $C_\theta$  --polymorphism-minimum-coverage-each-strand  
(DEFAULT = consensus mode, 0; polymorphism mode, 2)
- $\alpha_c$  --polymorphism-bias-cutoff  
(DEFAULT = OFF)

### Indel / homopolymer settings

- polymorphism-no-indels  
(DEFAULT = FALSE)
- polymorphism-reject-indel-homopolymer-length  
(DEFAULT = OFF; polymorphism mode, 3)
- polymorphism-reject-surrounding-homopolymer-length  
(DEFAULT = OFF; polymorphism mode, 2)

## Read Alignment (RA) evidence attributes

- $A_R$  Reference allele
- $A_1$  Major allele
- $A_\theta$  Minor allele
- $\theta$  Frequency of major allele
- Consensus score**
- $E_1$  E-value of posterior probability of consensus model ( $\theta_{\text{test}} = 100\%$ )
- Polymorphism score**
- $E_\theta$  E-value of LRT of polymorphic model with ( $\theta_{\text{test}} = \theta$ ) versus the consensus model
- $C_{1,\text{top}}/C_{1,\text{bot}}$  Reads with major allele on top/bottom strand
- $C_{\theta,\text{top}}/C_{\theta,\text{bot}}$  Reads with variant allele on top/bottom strand
- $P_s$  Read strand bias (Fisher's exact test)
- $P_q$  Base quality bias (Kolmogorov-Smirnov test)