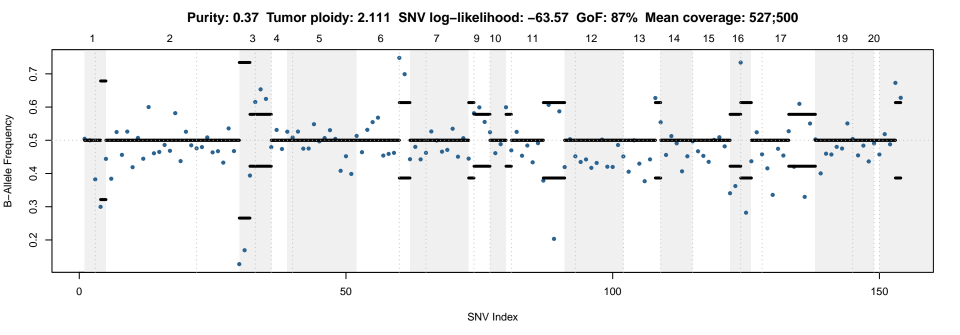
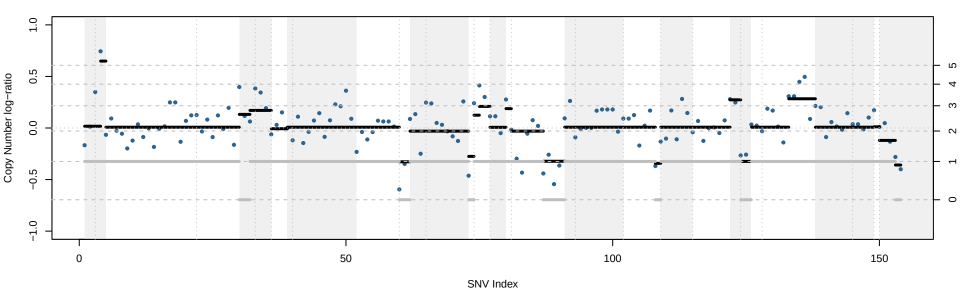
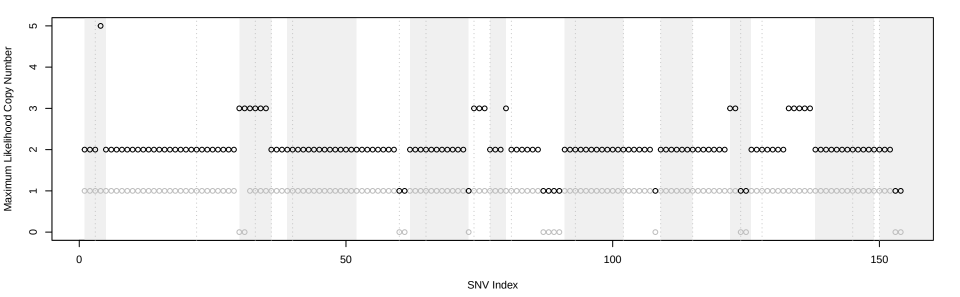
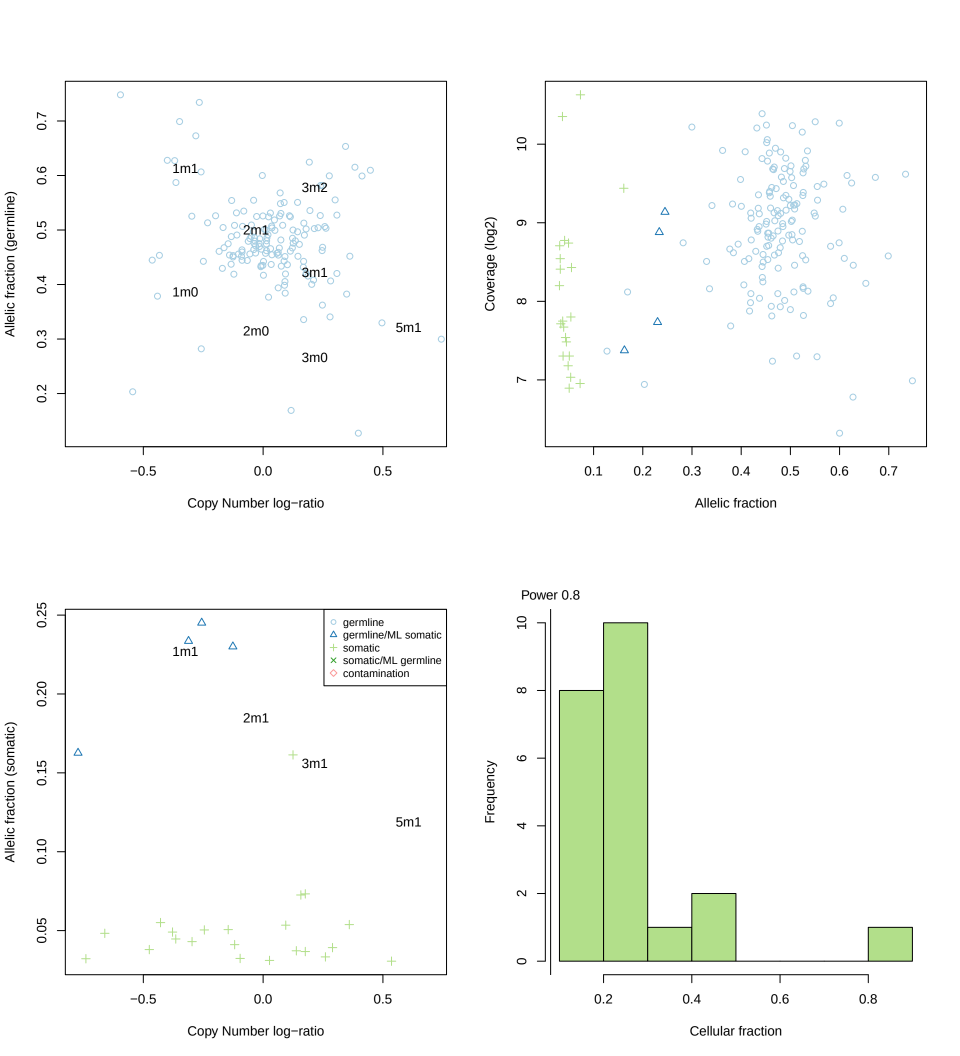
Purity: 0.37 Tumor ploidy: 2.111 3 0 5 0.3 Fraction Genome 0.2 0.1 0.0 -0.5 0.0 0.5 1.0 log2 ratio



SCNA-fit log-likelihood: -4569.95







Purity: 0.46 Tumor ploidy: 4.094 2 3 5 6

0.0

log2 ratio

0.5

1.0

0.3

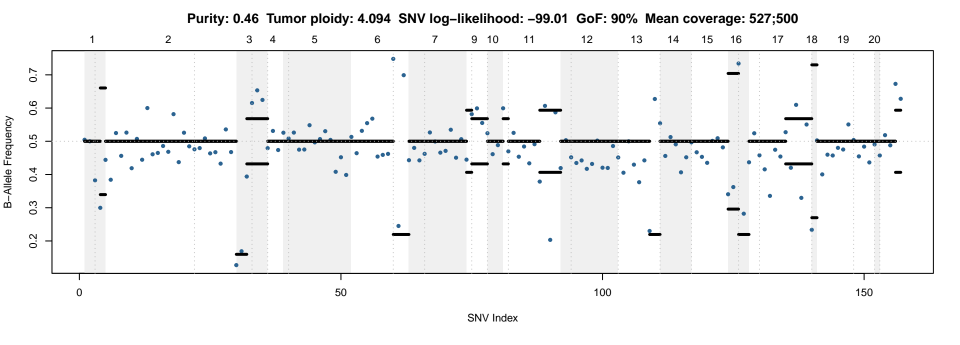
0.2

0.1

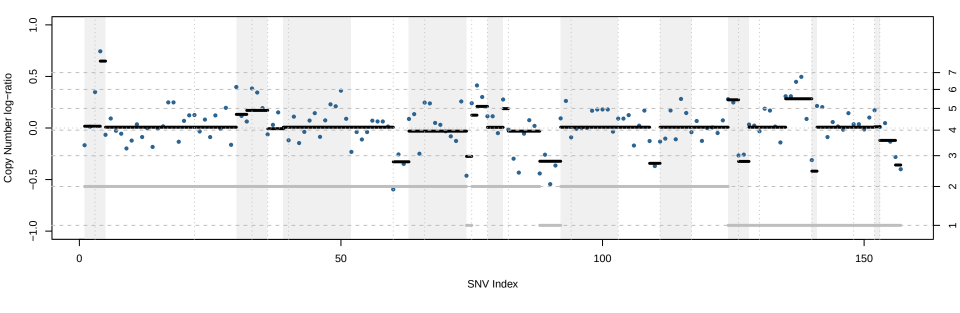
0.0

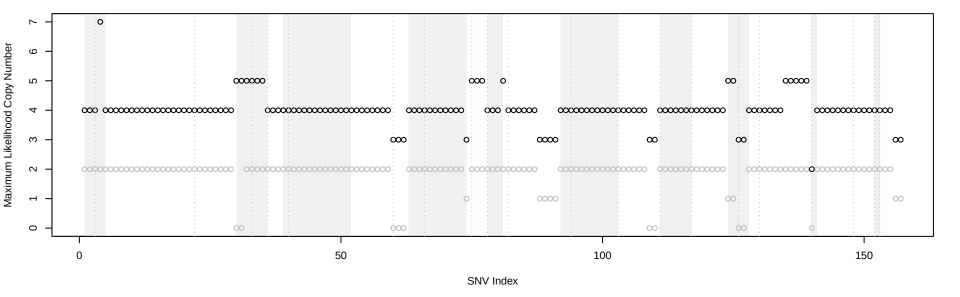
-0.5

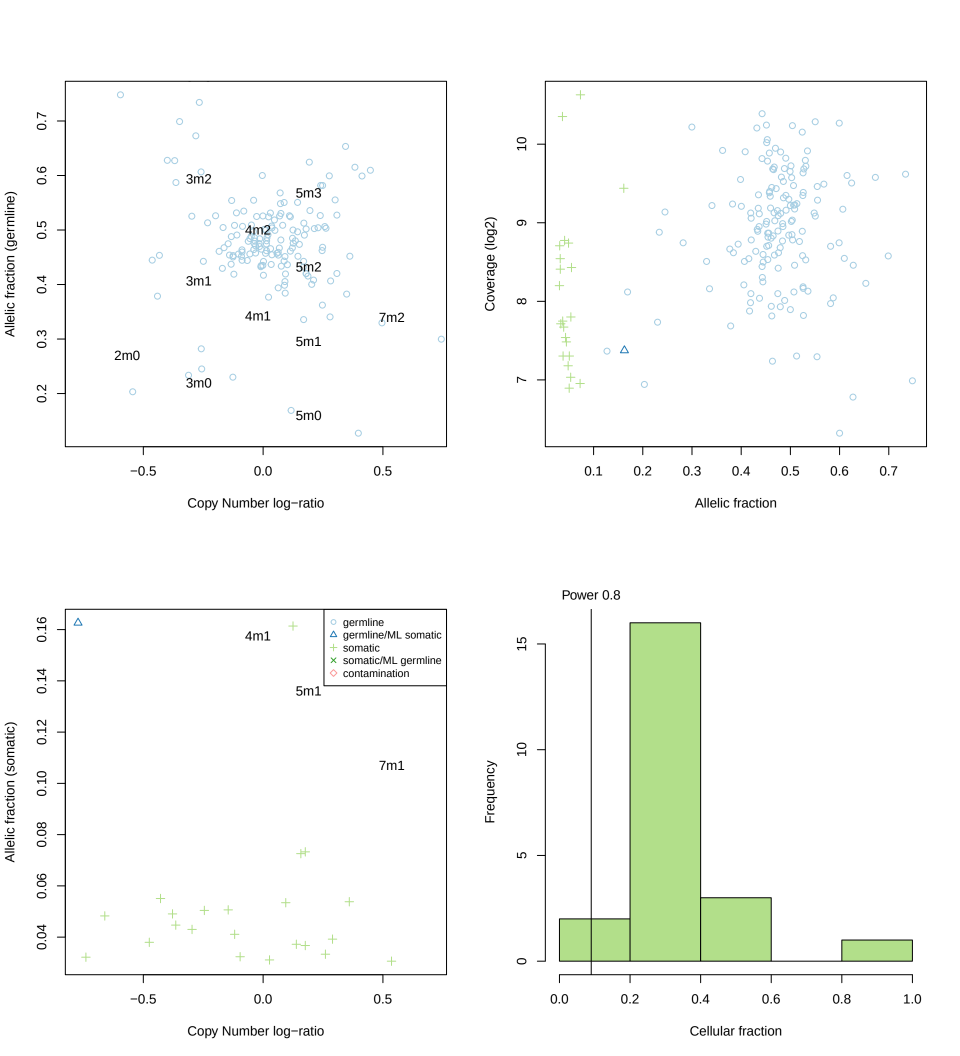
Fraction Genome



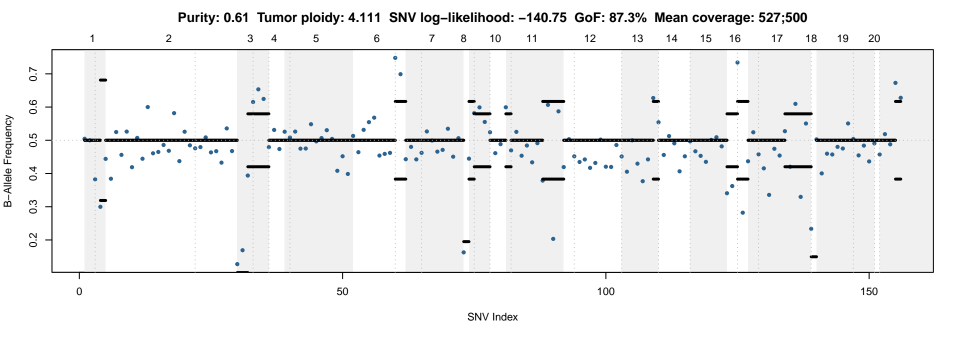
SCNA-fit log-likelihood: -4593.01



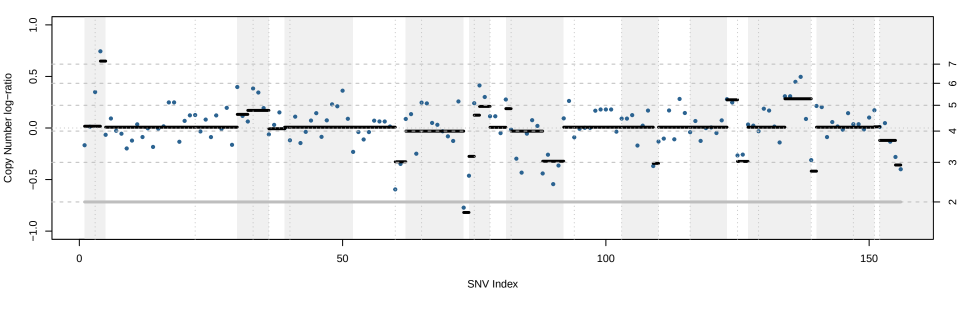


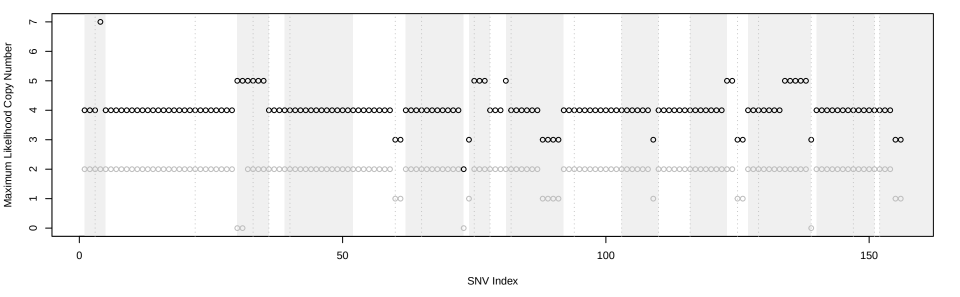


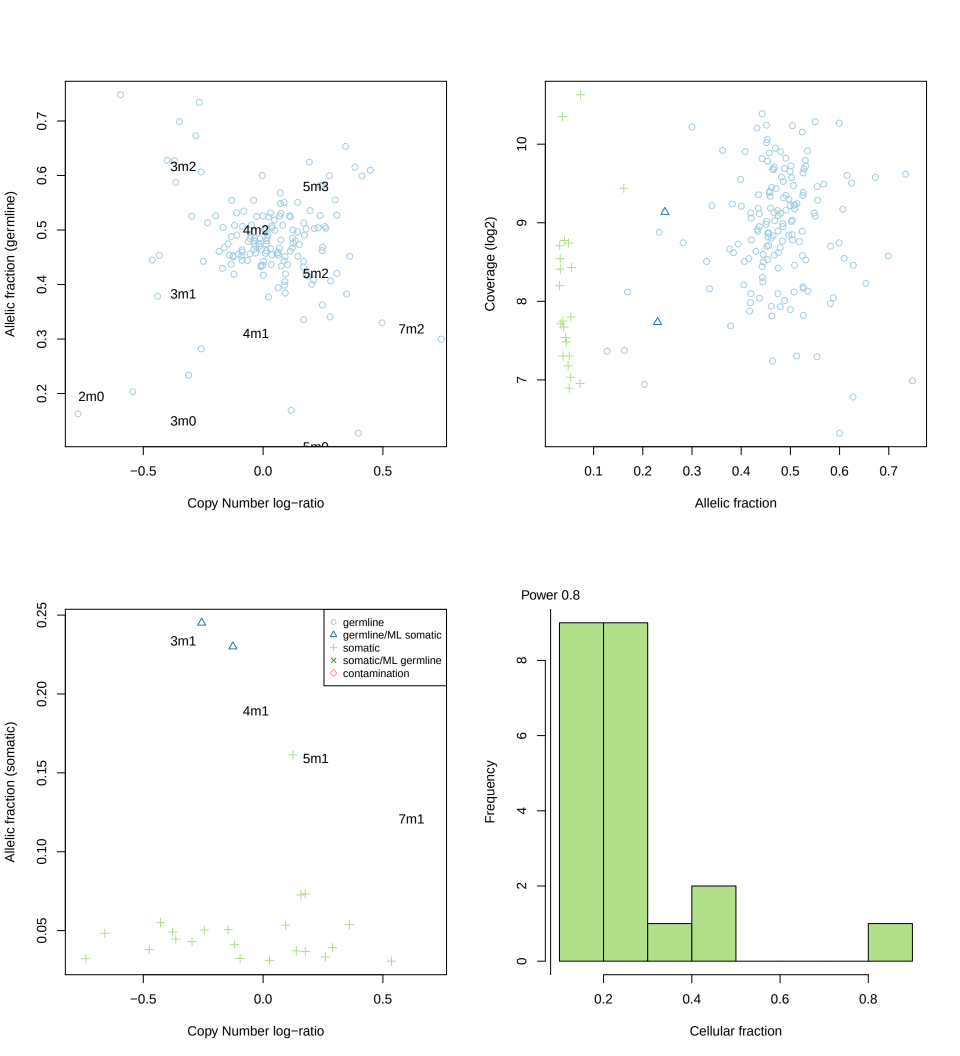
Purity: 0.61 Tumor ploidy: 4.111 6 2 3 5 0.3 Fraction Genome 0.2 0.1 0.0 -0.5 0.0 0.5 1.0 log2 ratio



SCNA-fit log-likelihood: -4571.35

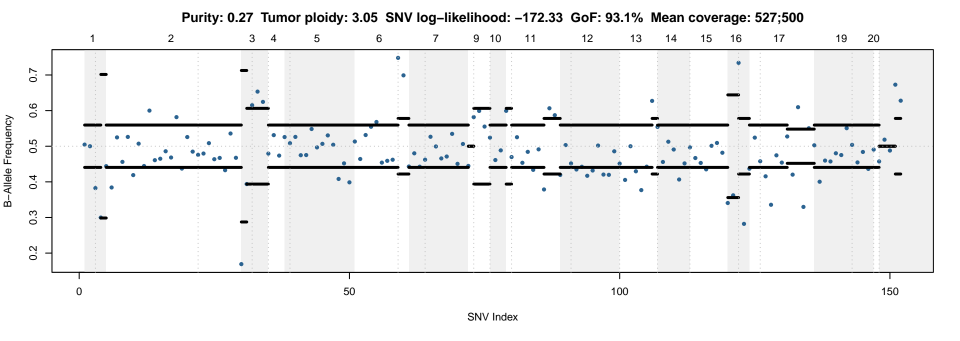




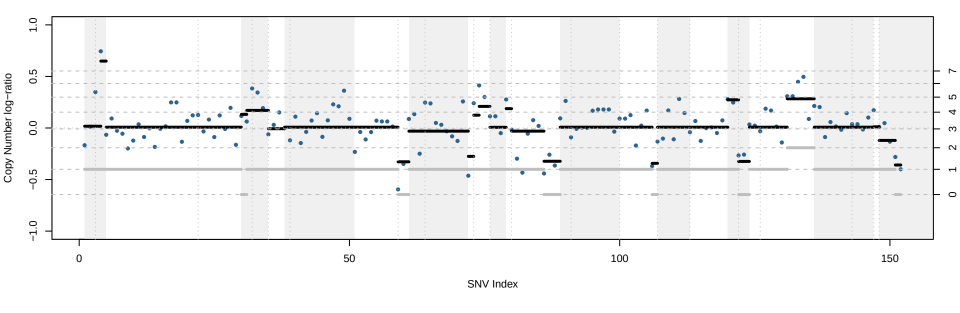


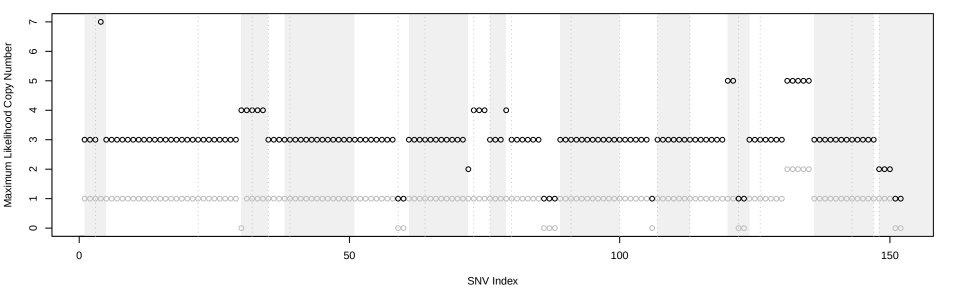
Purity: 0.27 Tumor ploidy: 3.05 0 2 6 3 5 0.3 Fraction Genome 0.2 0.1 0.0 -0.5 0.0 0.5 1.0

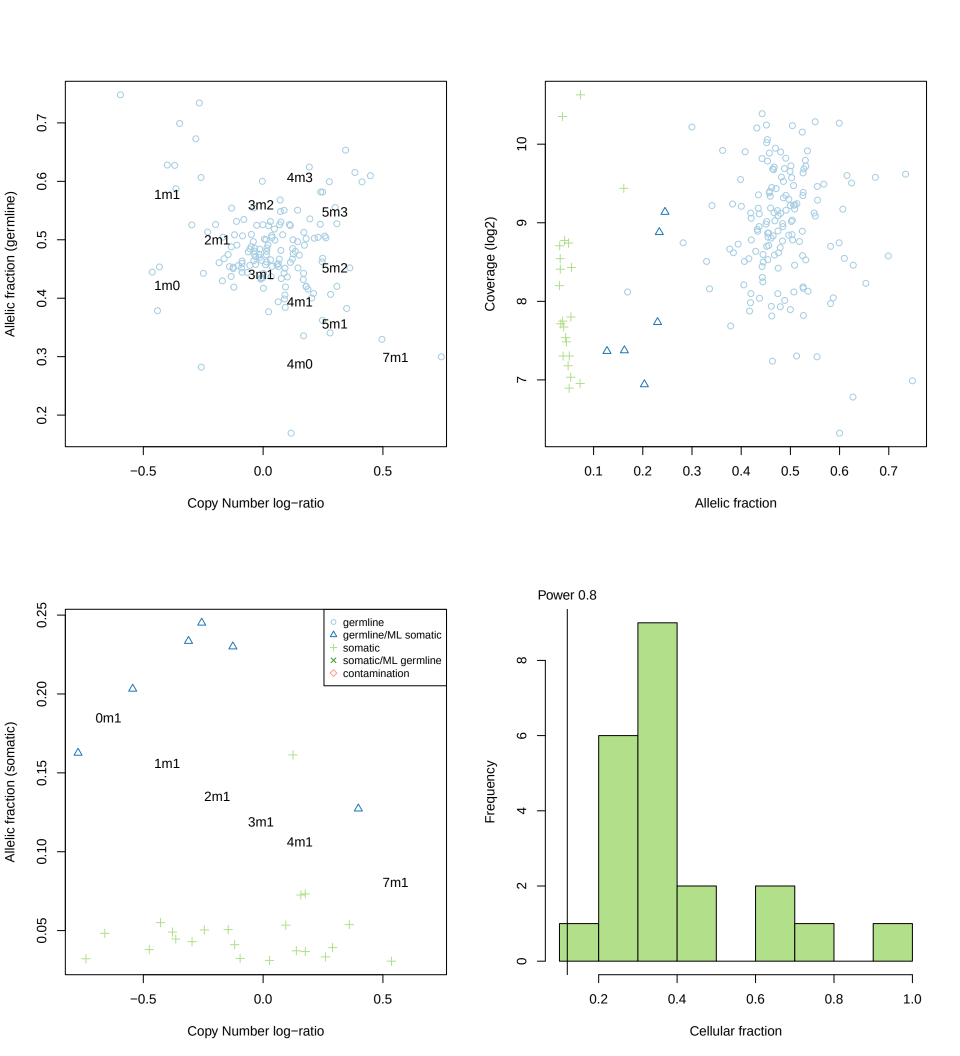
log2 ratio



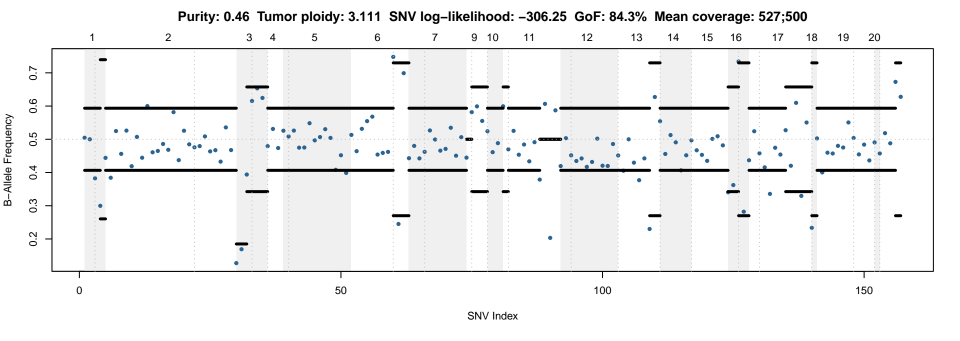
SCNA-fit log-likelihood: -4541.33



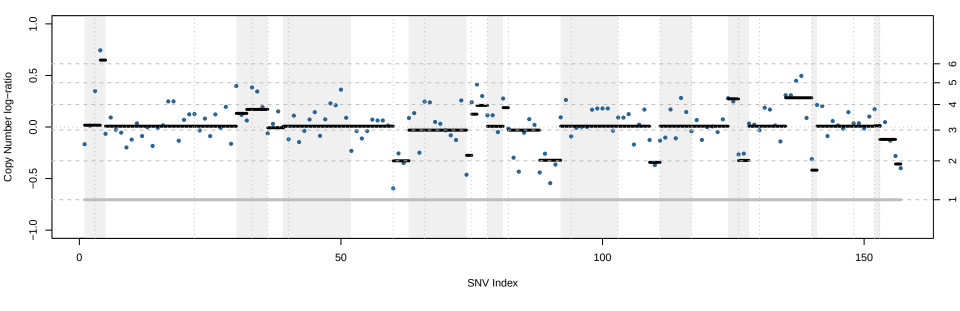


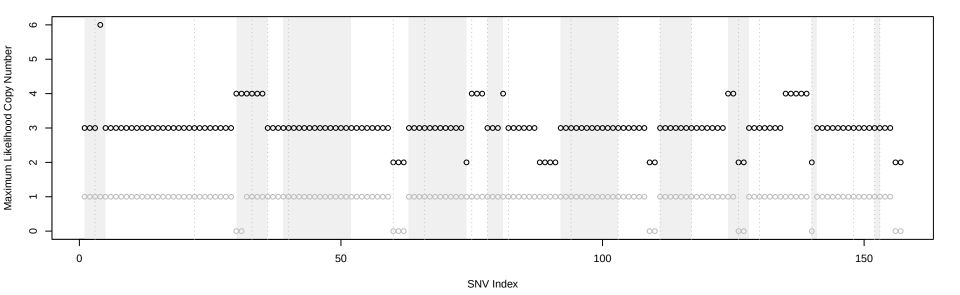


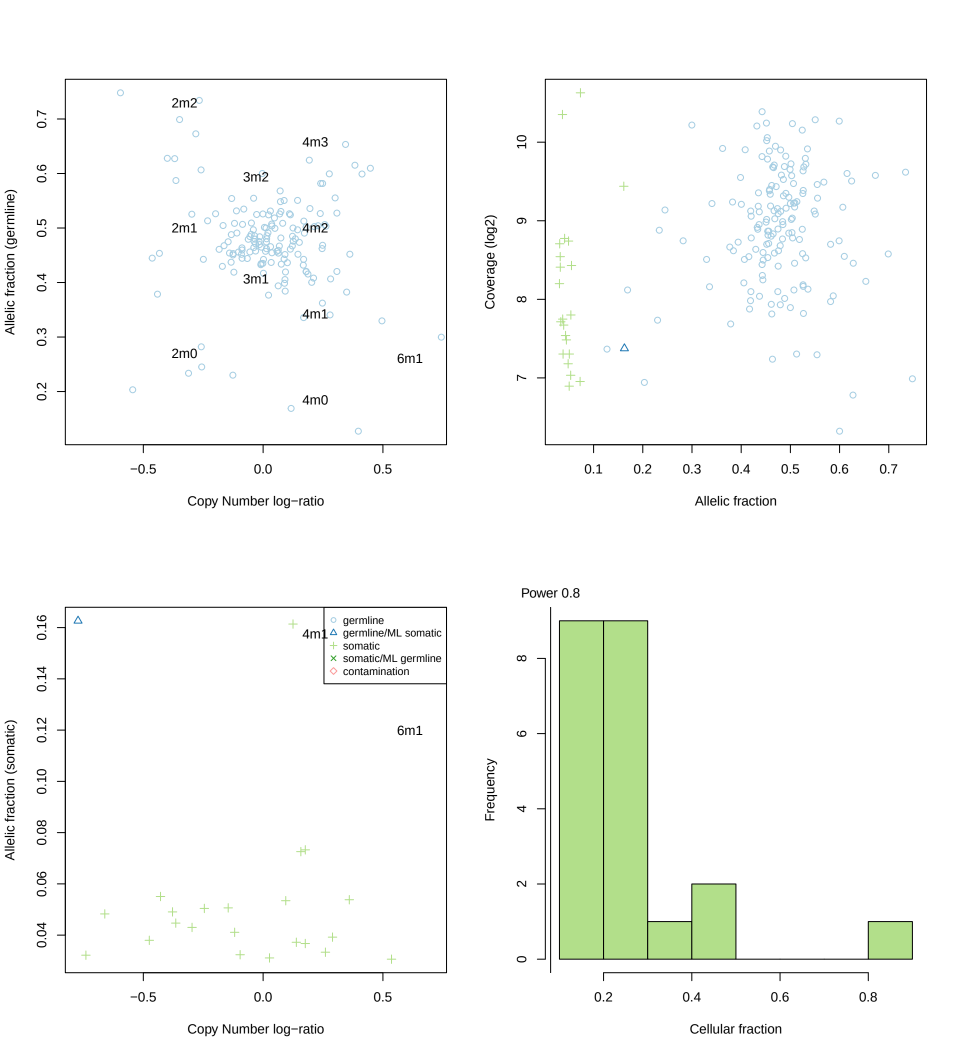
Purity: 0.46 Tumor ploidy: 3.111 5 2 6 0.3 Fraction Genome 0.2 0.1 0.0 -0.5 0.0 0.5 1.0 log2 ratio



SCNA-fit log-likelihood: -4570.25

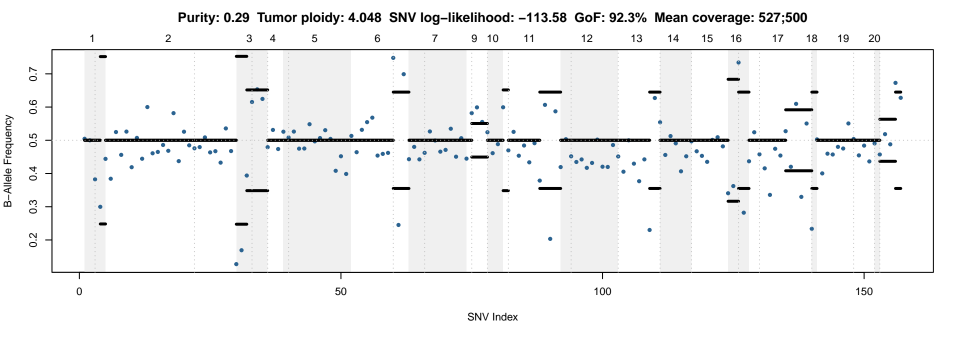




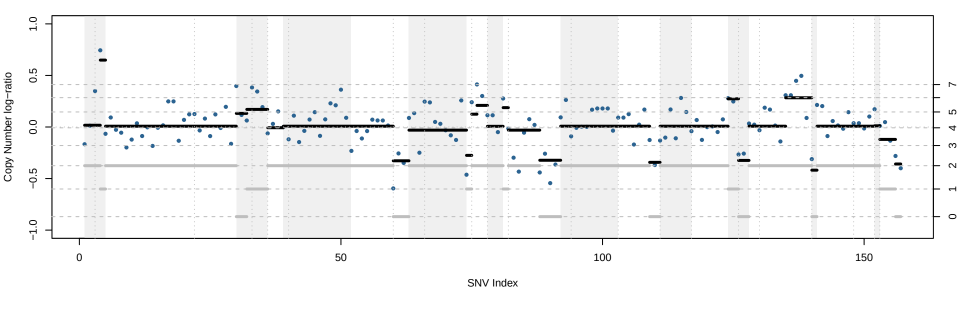


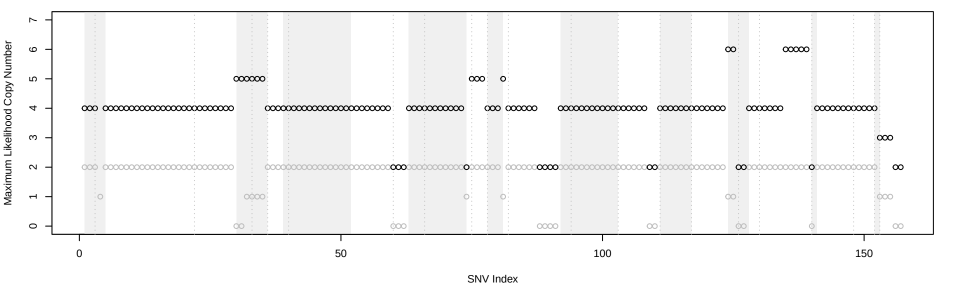
Purity: 0.29 Tumor ploidy: 4.048 2 0 3 5 6 0.4 0.3 Fraction Genome 0.2 0.1 0.0 -0.5 0.0 0.5 1.0

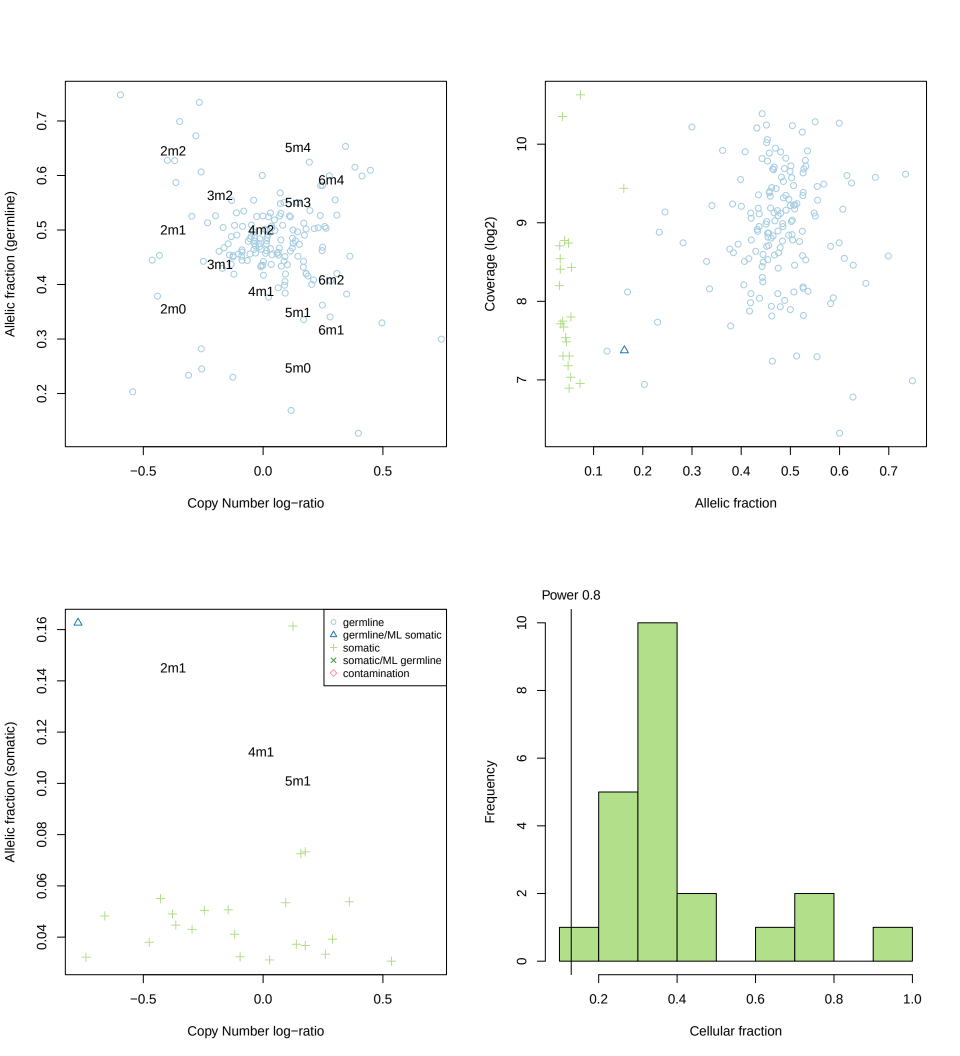
log2 ratio



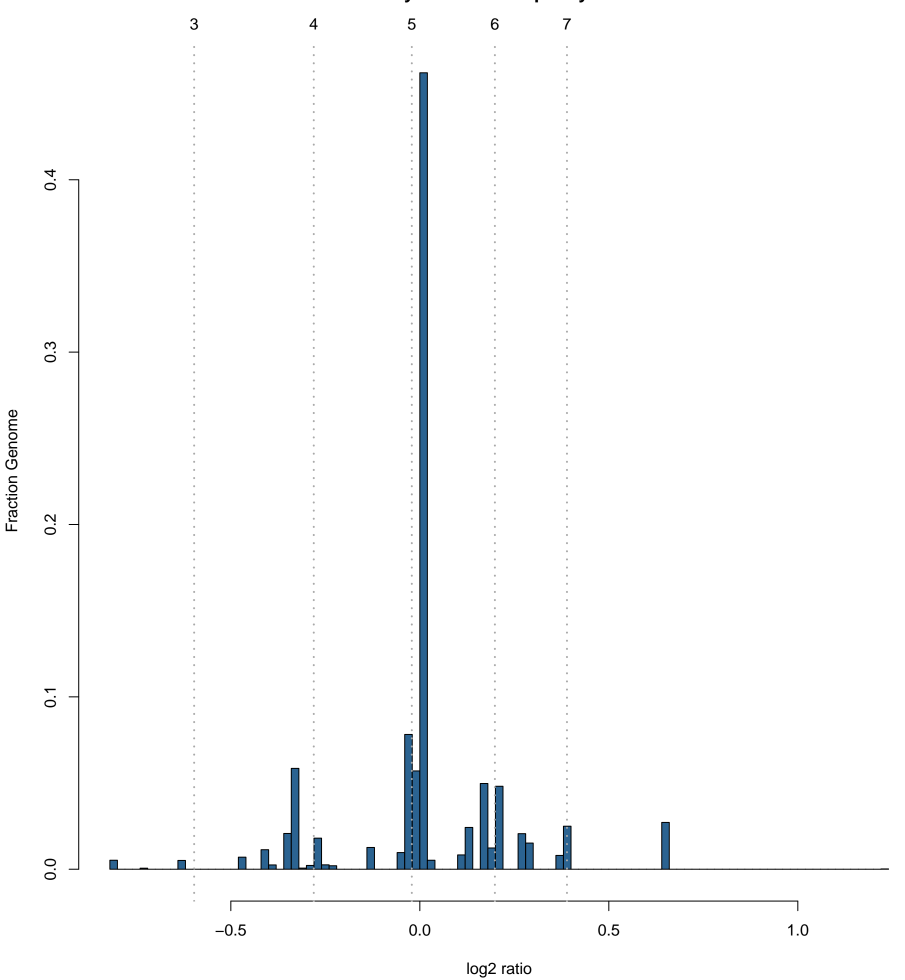
SCNA-fit log-likelihood: -4939.38

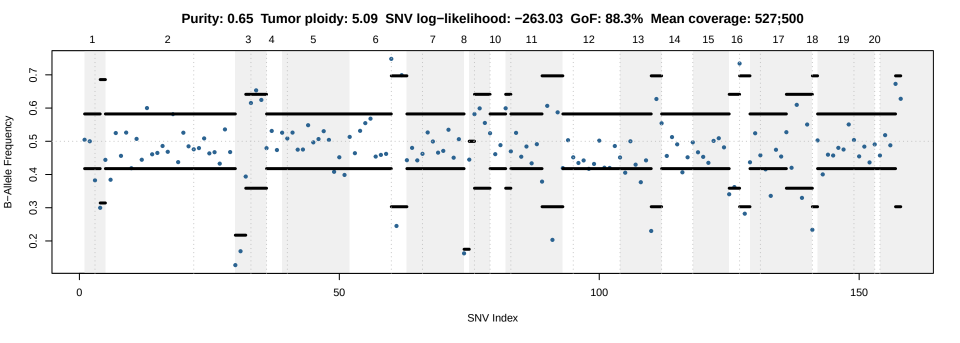




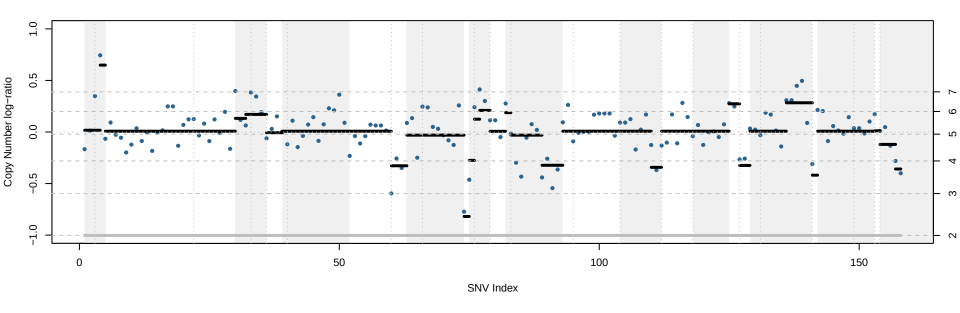


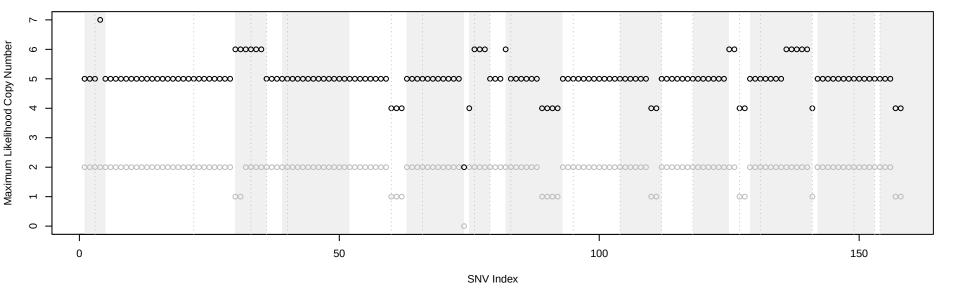
Purity: 0.65 Tumor ploidy: 5.09

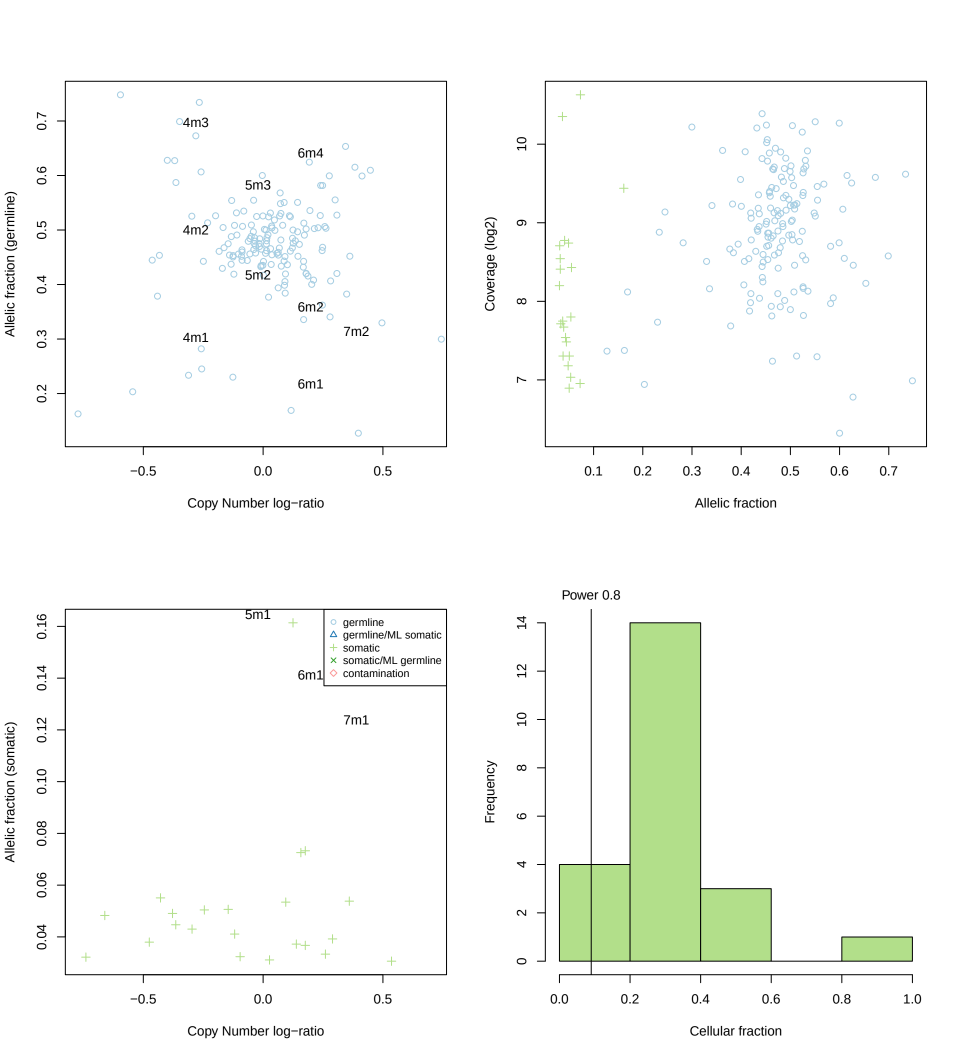




SCNA-fit log-likelihood: -4707.99

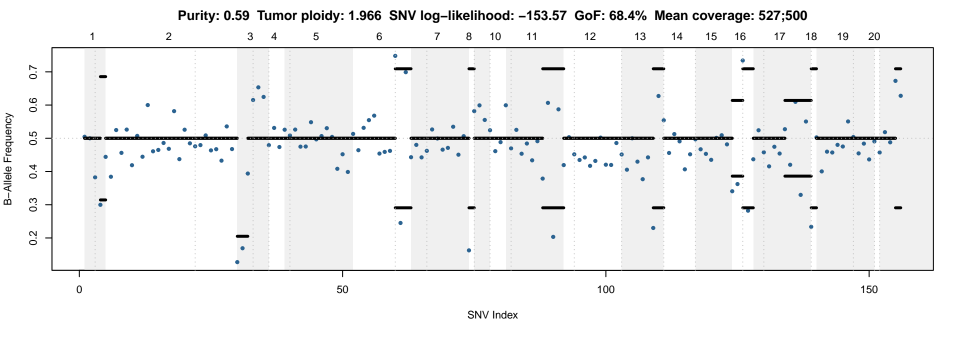






Purity: 0.59 Tumor ploidy: 1.966 3 2 6 0.3 Fraction Genome 0.2 0.1 0.0 -0.5 0.0 0.5 1.0

log2 ratio



SCNA-fit log-likelihood: -5082.08

