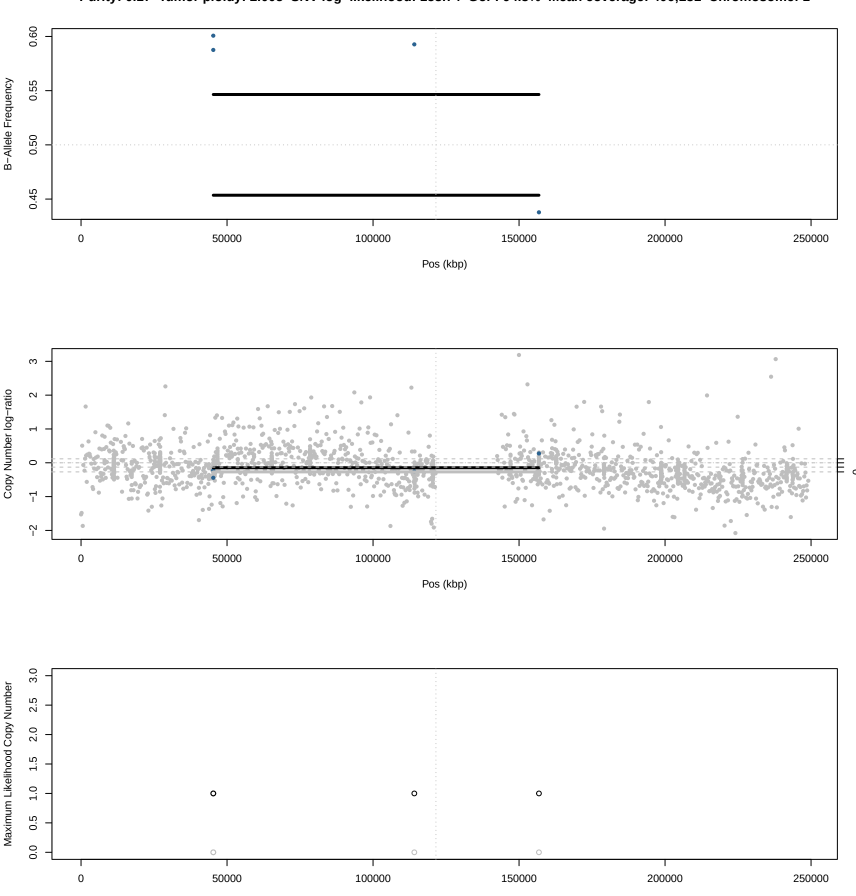
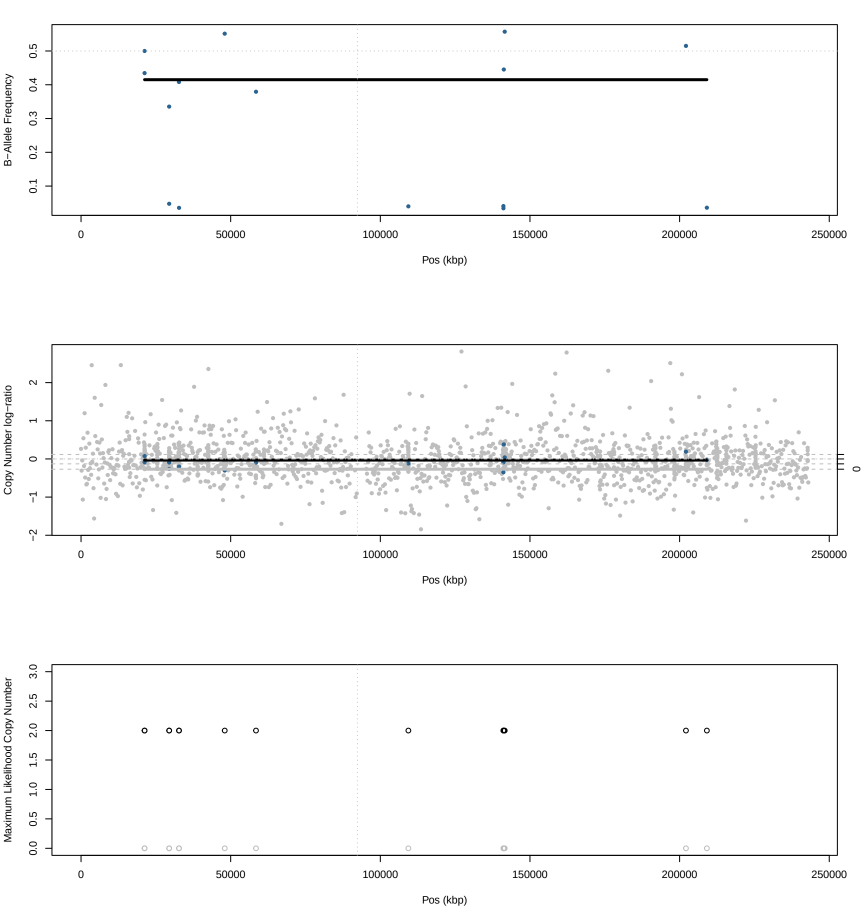
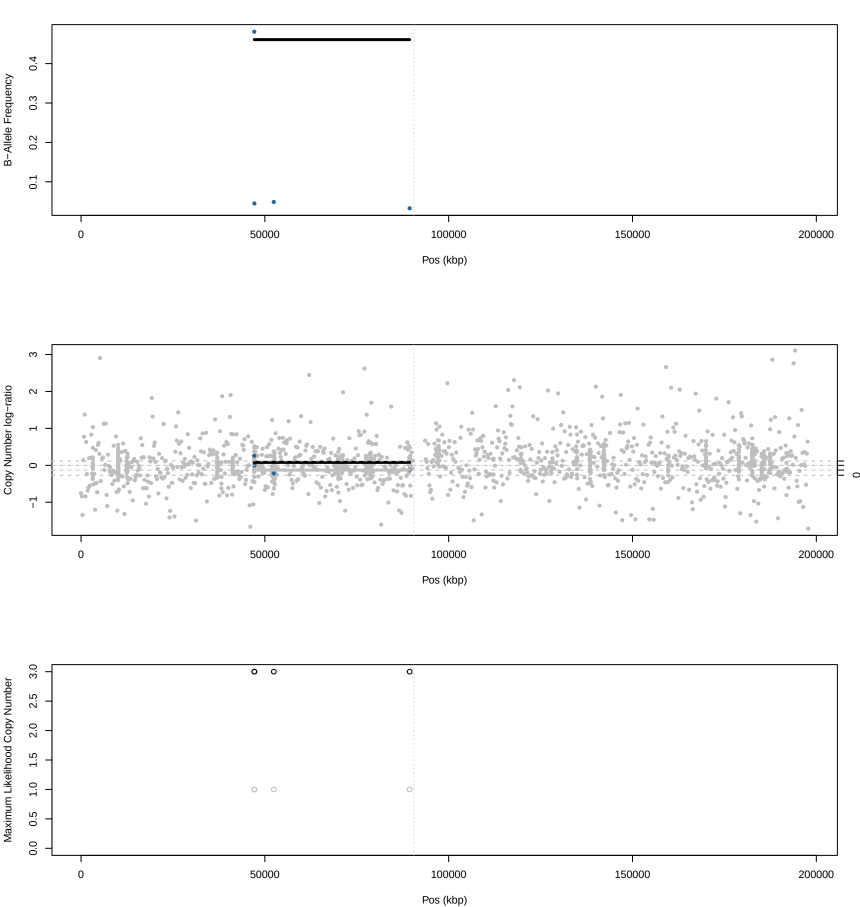
Purity: 0.17 Tumor ploidy: 2.005 SNV log-likelihood: 158.74 GoF: 94.3% Mean coverage: 406;281 Chromosome: 1



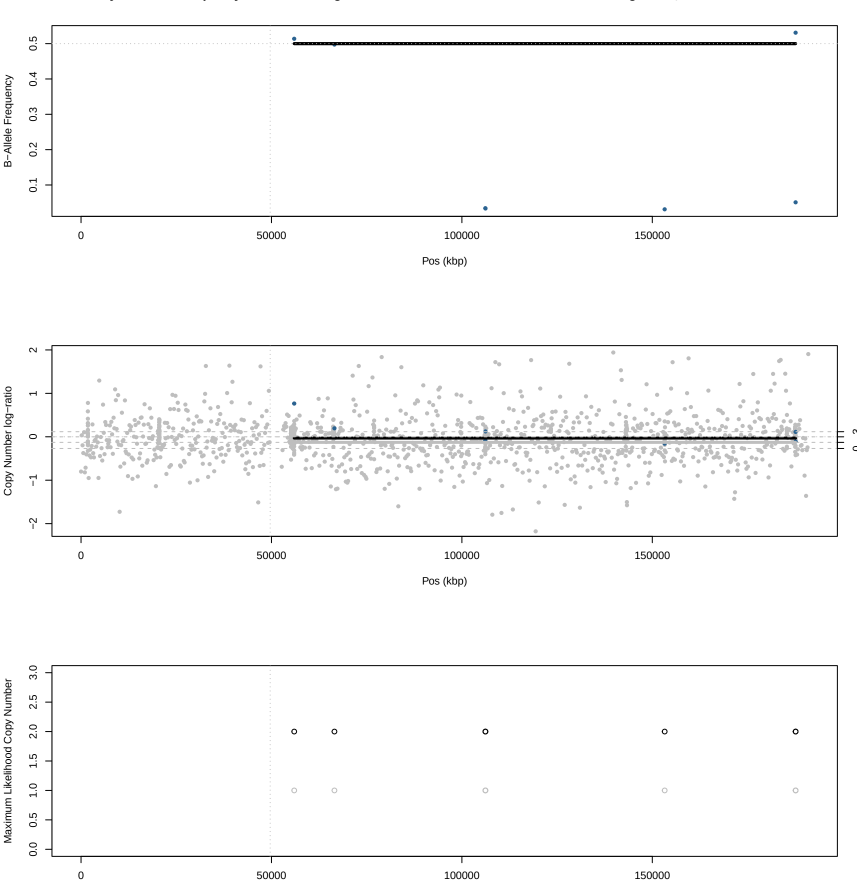
Purity: 0.17 Tumor ploidy: 2.005 SNV log-likelihood: 158.74 GoF: 94.3% Mean coverage: 406;281 Chromosome: 2



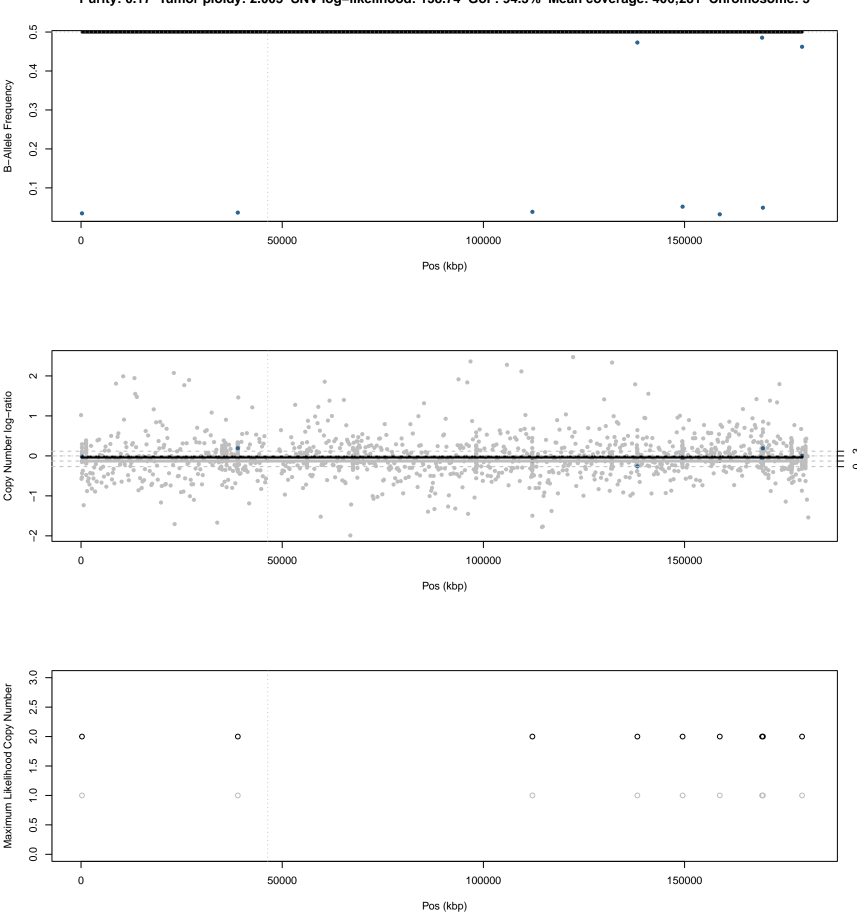
Purity: 0.17 Tumor ploidy: 2.005 SNV log-likelihood: 158.74 GoF: 94.3% Mean coverage: 406;281 Chromosome: 3



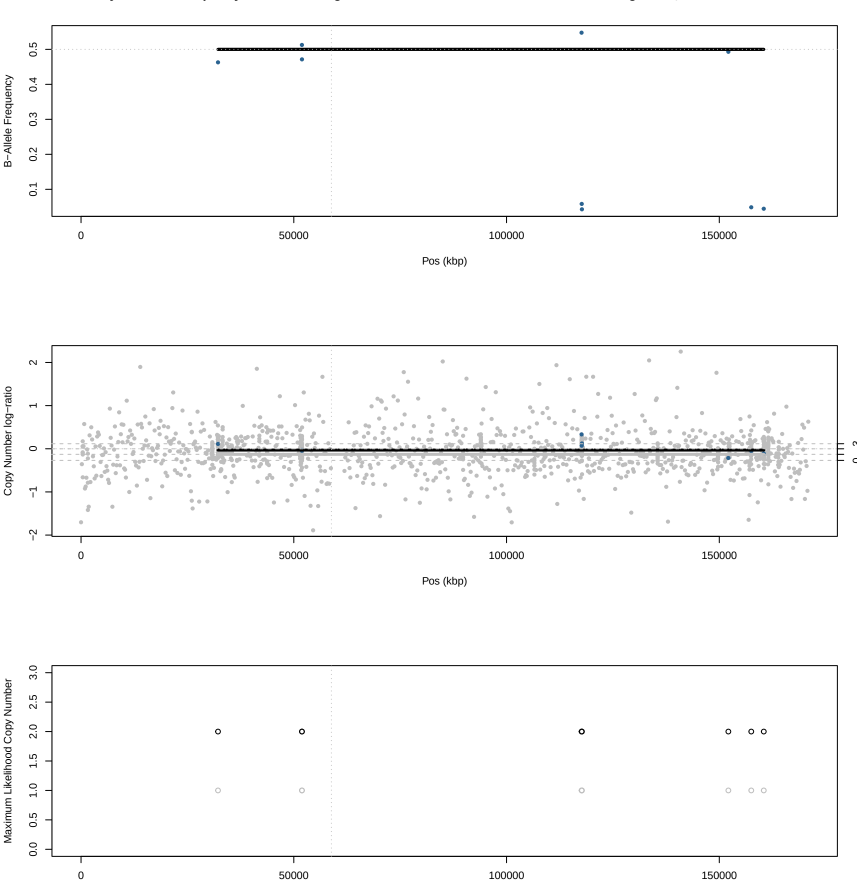
Purity: 0.17 Tumor ploidy: 2.005 SNV log-likelihood: 158.74 GoF: 94.3% Mean coverage: 406;281 Chromosome: 4



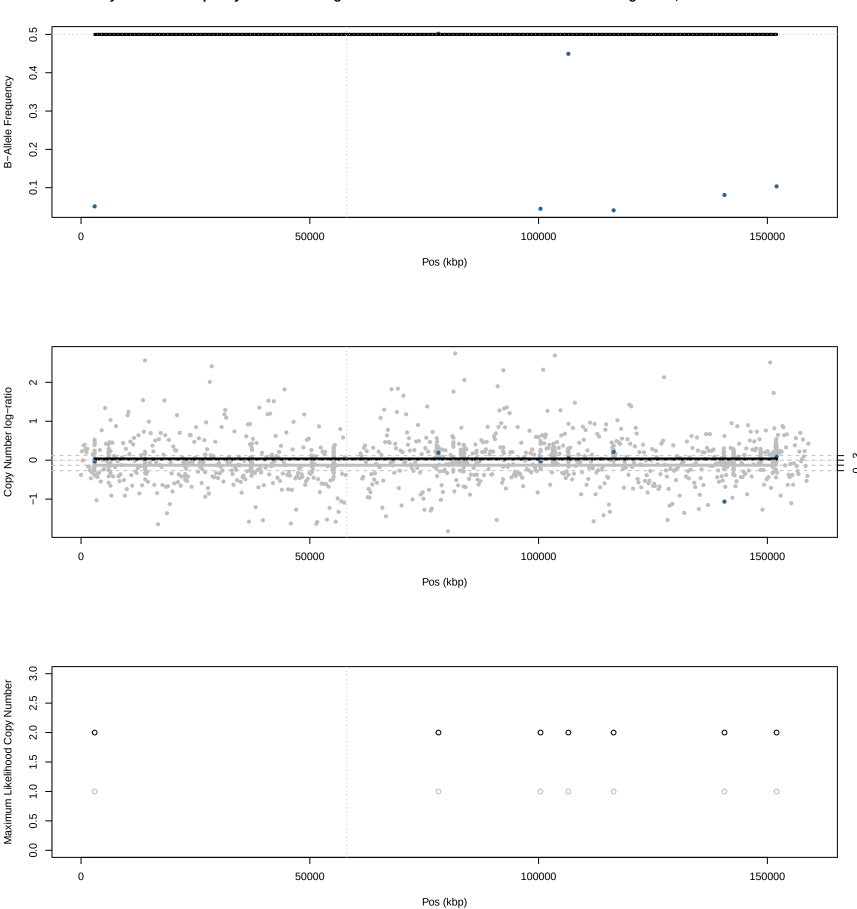
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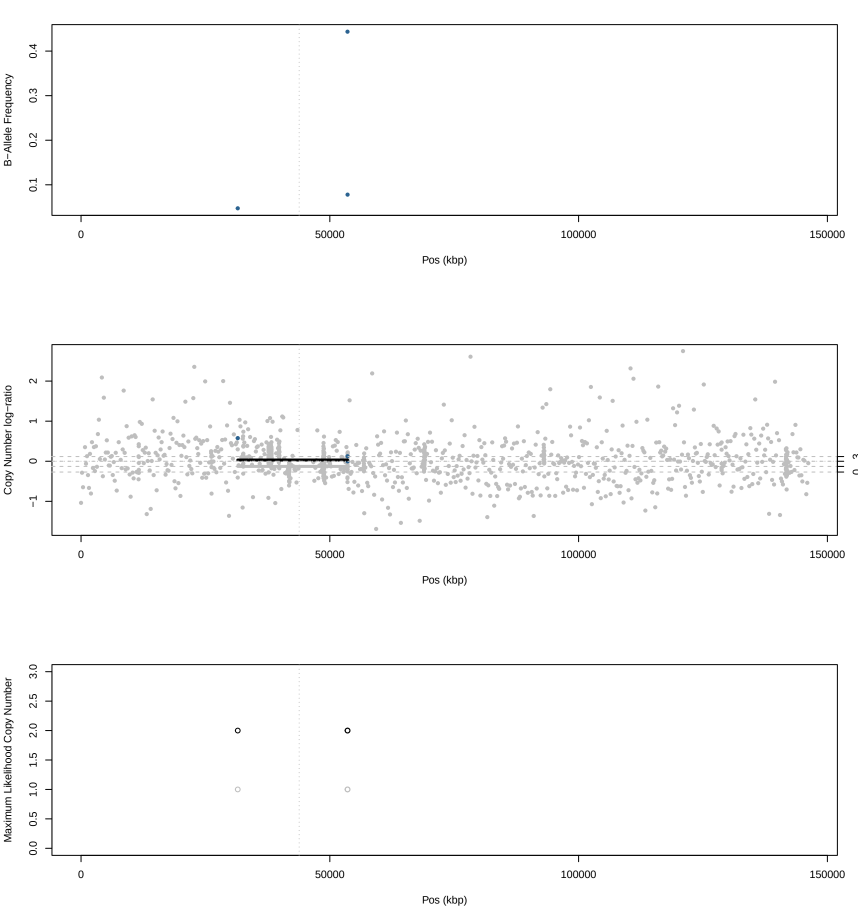
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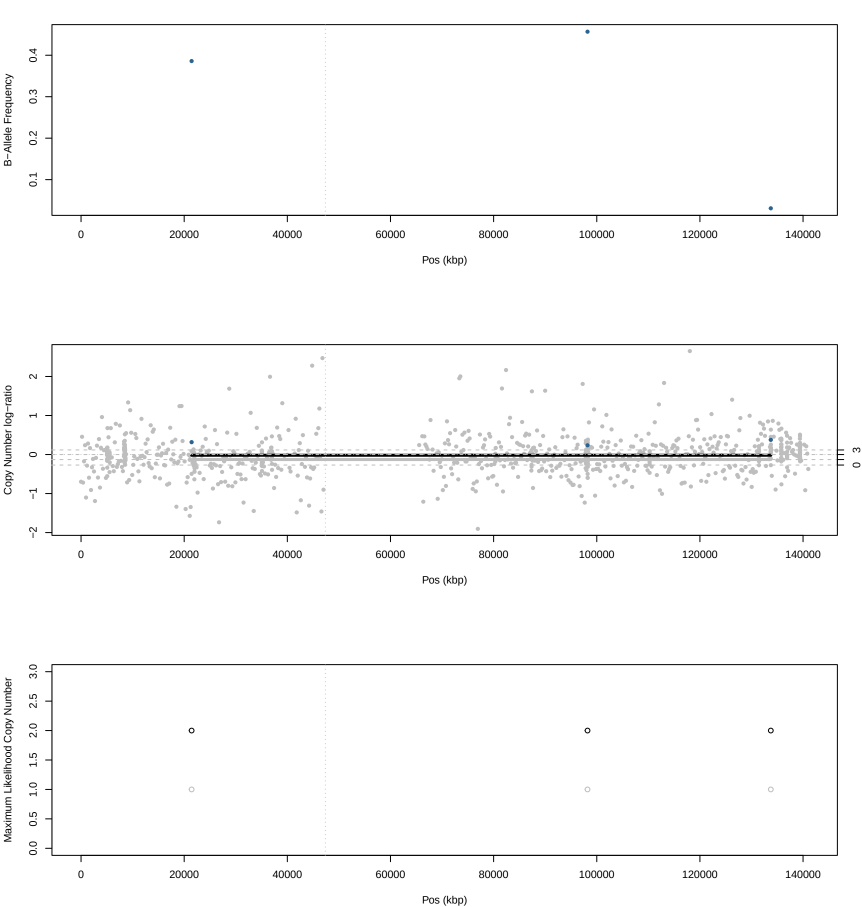
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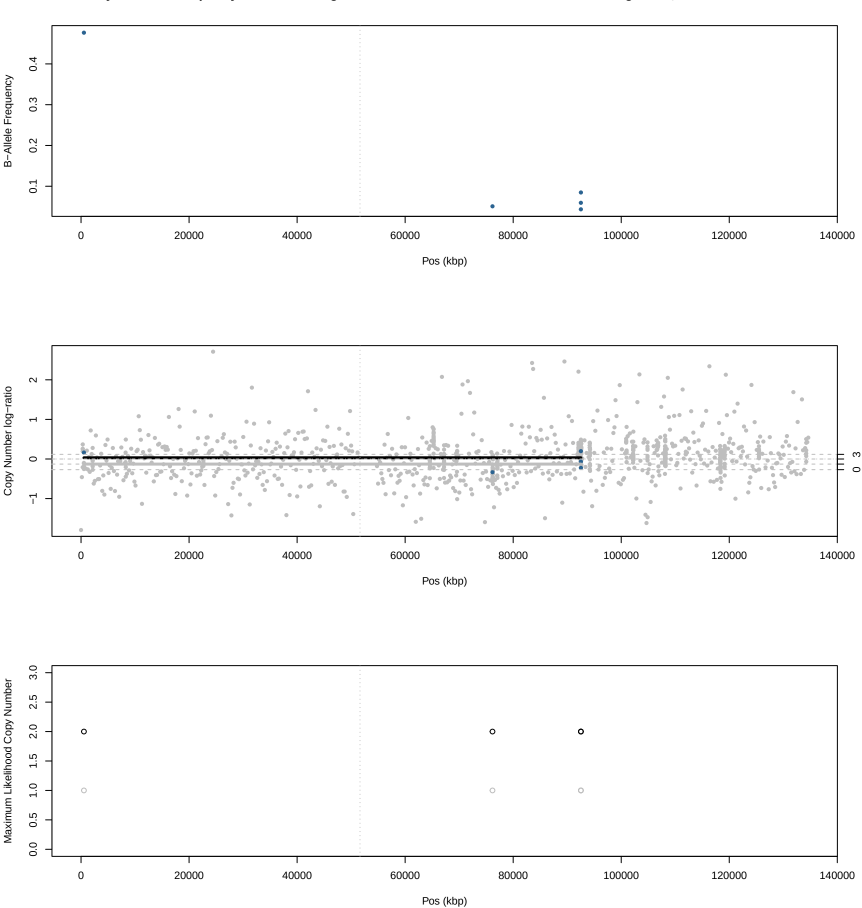
Purity: 0.17 Tumor ploidy: 2.005 SNV log-likelihood: 158.74 GoF: 94.3% Mean coverage: 406;281 Chromosome: 8



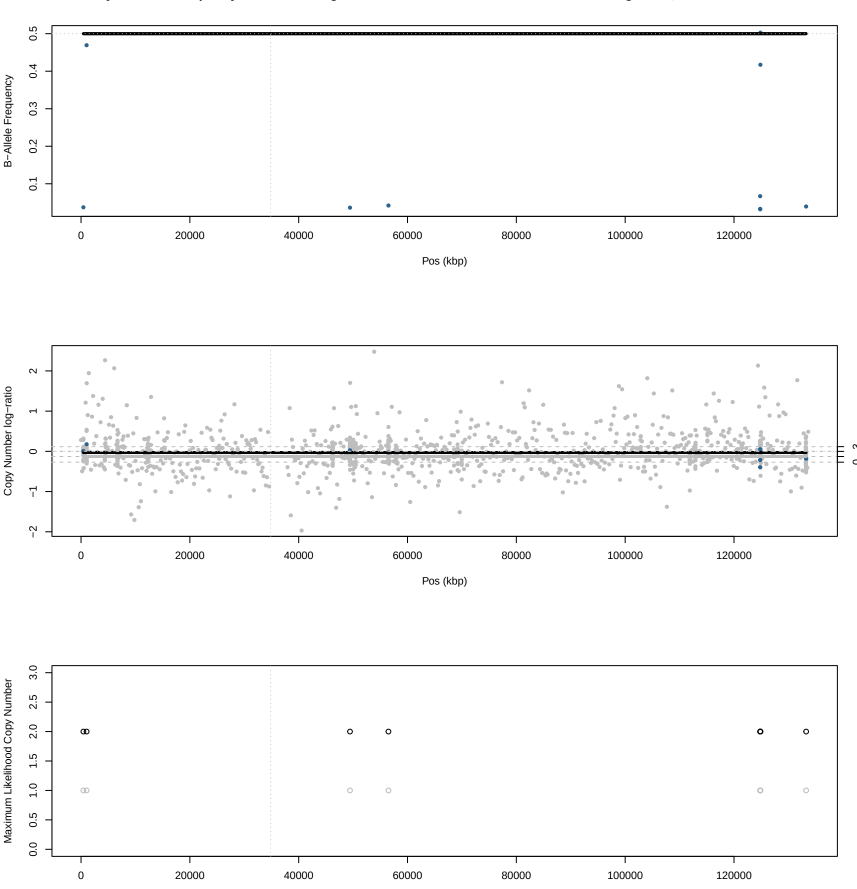
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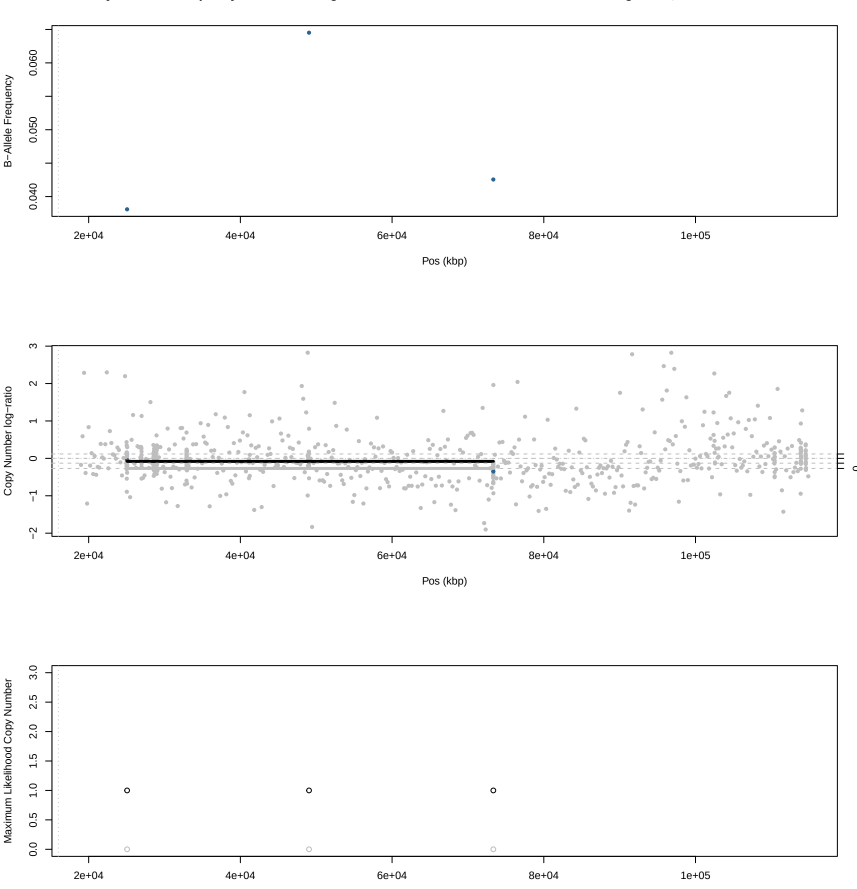
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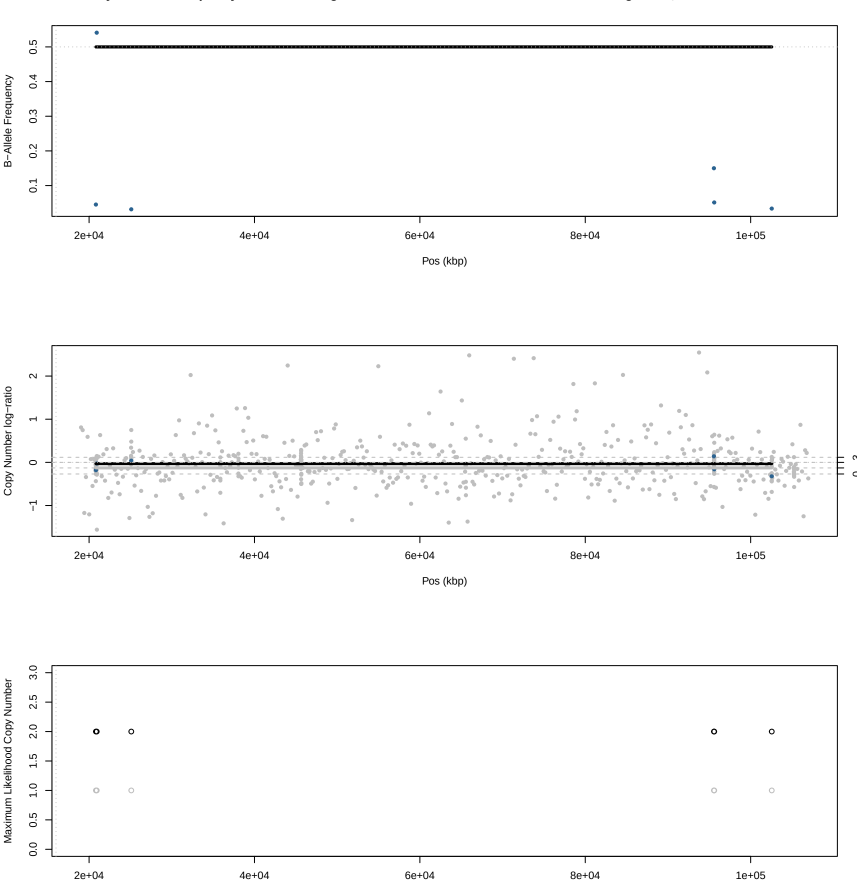
Purity: 0.17 Tumor ploidy: 2.005 SNV log-likelihood: 158.74 GoF: 94.3% Mean coverage: 406;281 Chromosome: 12



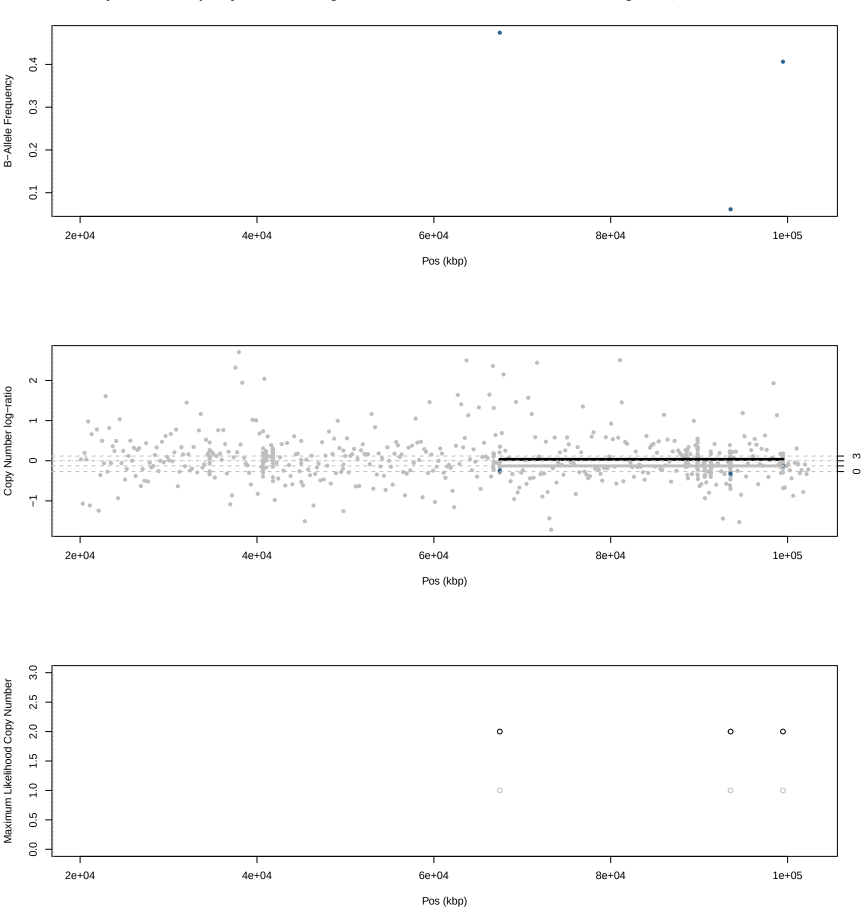
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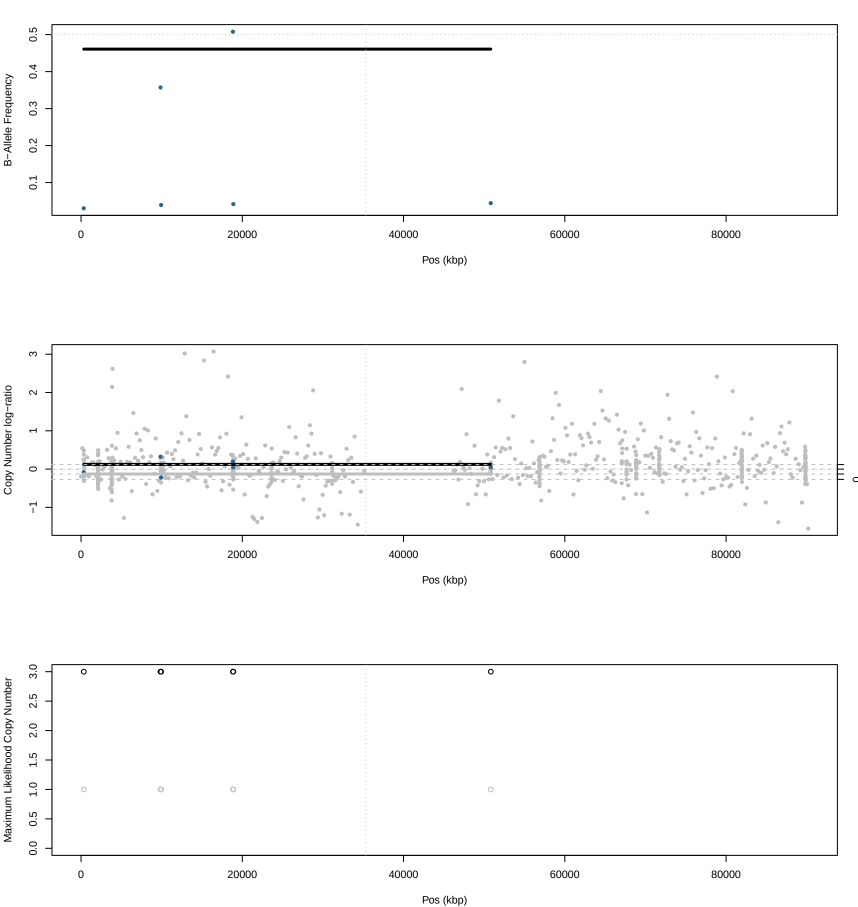
Purity: 0.17 Tumor ploidy: 2.005 SNV log-likelihood: 158.74 GoF: 94.3% Mean coverage: 406;281 Chromosome: 14



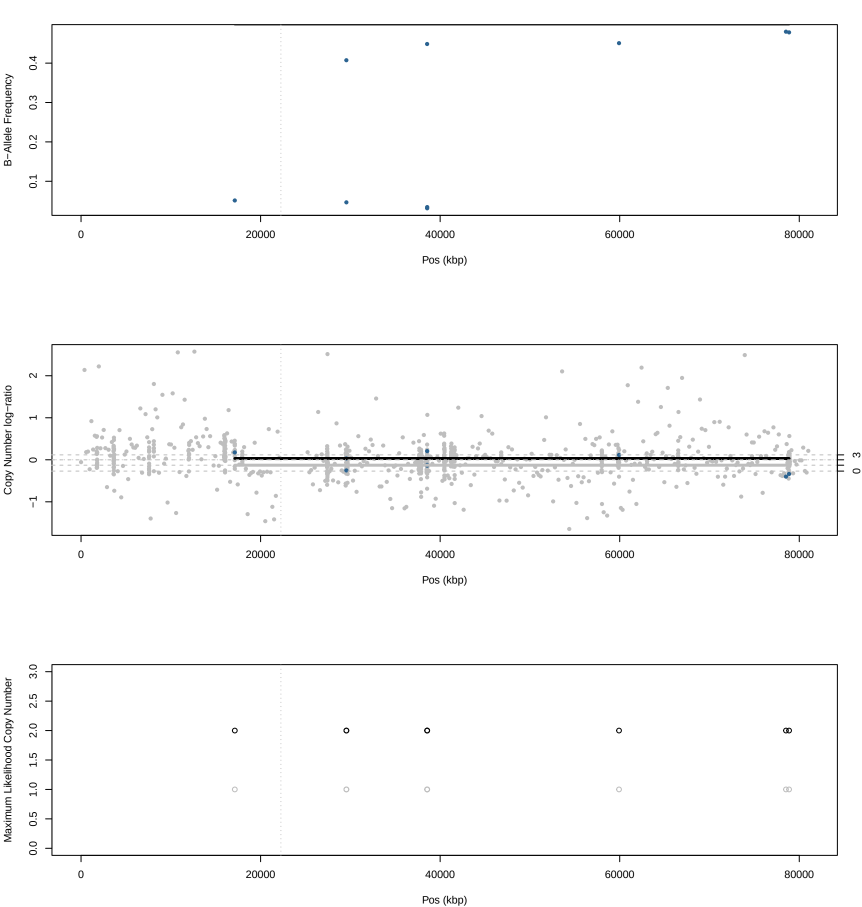
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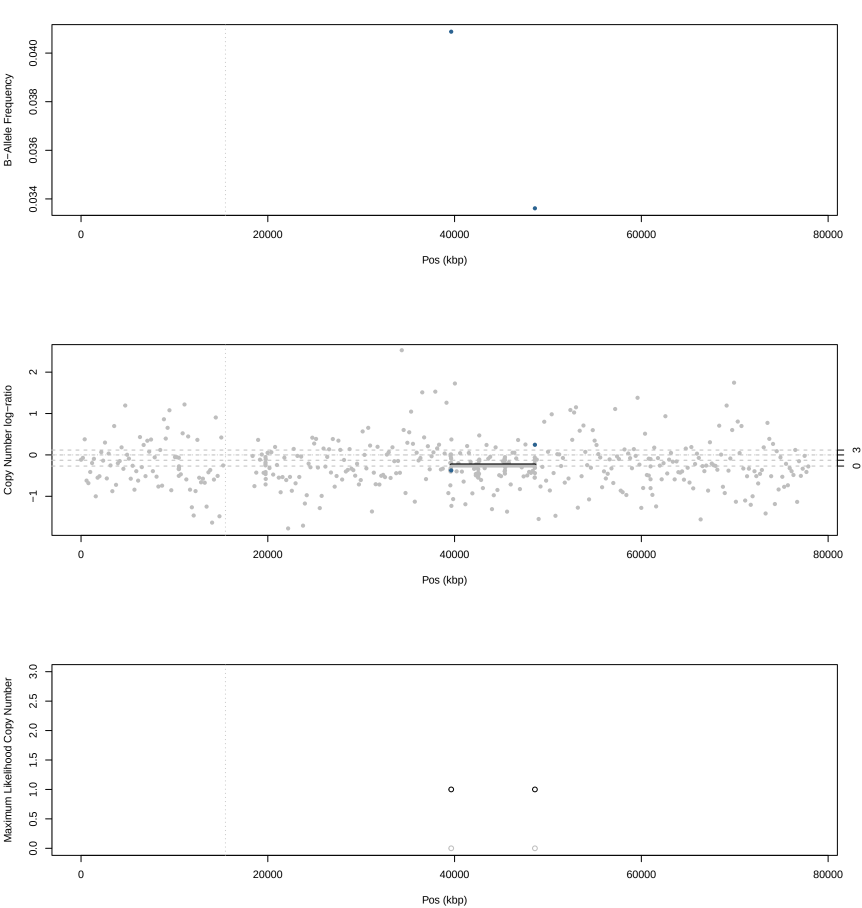
Purity: 0.17 Tumor ploidy: 2.005 SNV log-likelihood: 158.74 GoF: 94.3% Mean coverage: 406;281 Chromosome: 16



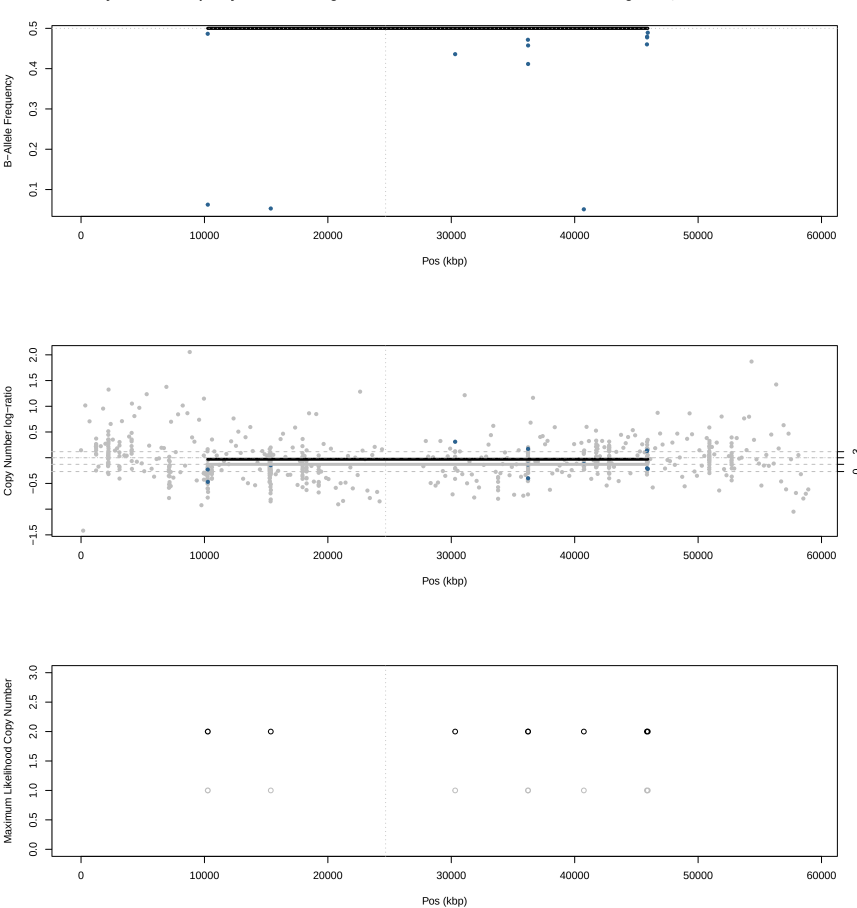
Purity: 0.17 Tumor ploidy: 2.005 SNV log-likelihood: 158.74 GoF: 94.3% Mean coverage: 406;281 Chromosome: 17



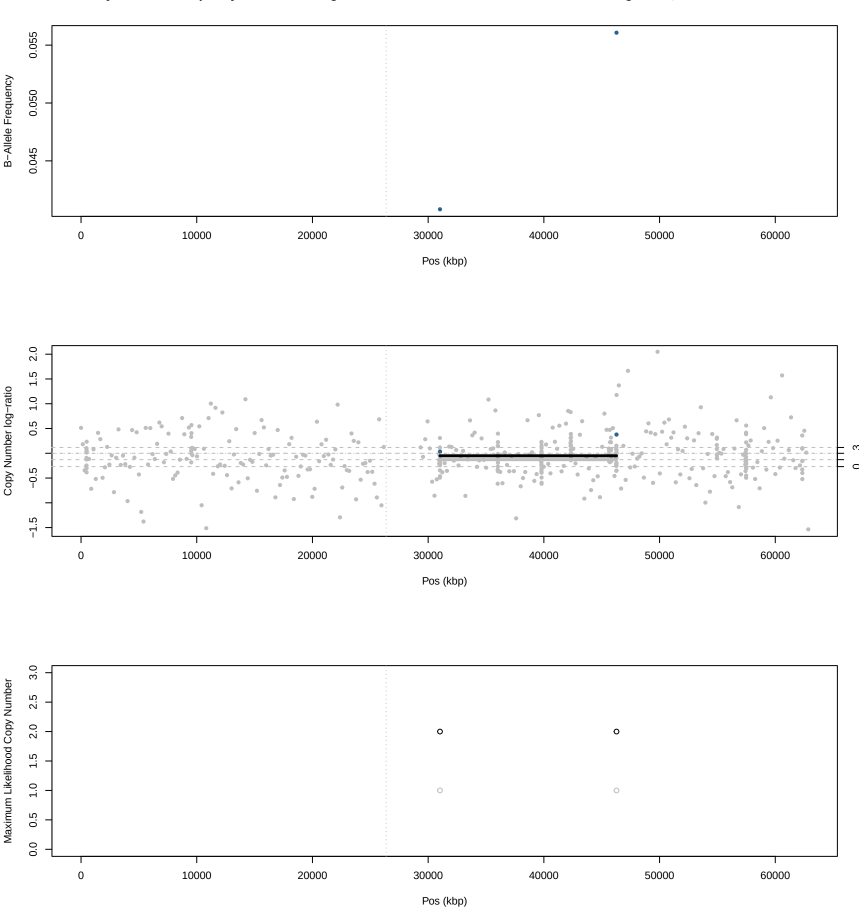
Purity: 0.17 Tumor ploidy: 2.005 SNV log-likelihood: 158.74 GoF: 94.3% Mean coverage: 406;281 Chromosome: 18



Purity: 0.17 Tumor ploidy: 2.005 SNV log-likelihood: 158.74 GoF: 94.3% Mean coverage: 406;281 Chromosome: 19



Purity: 0.17 Tumor ploidy: 2.005 SNV log-likelihood: 158.74 GoF: 94.3% Mean coverage: 406;281 Chromosome: 20



Purity: 0.17 Tumor ploidy: 2.005 SNV log-likelihood: 158.74 GoF: 94.3% Mean coverage: 406;281 Chromosome: 21

