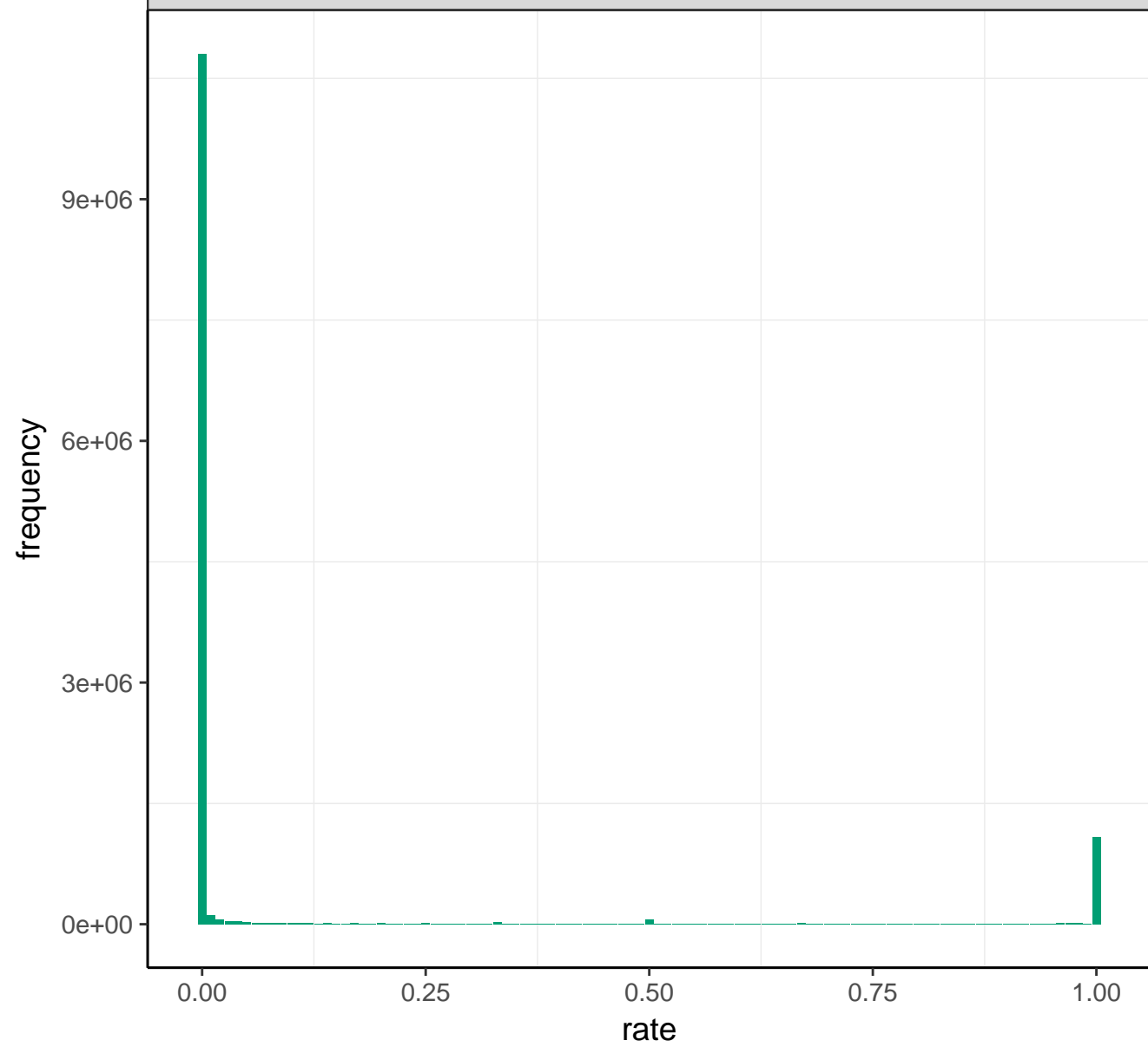
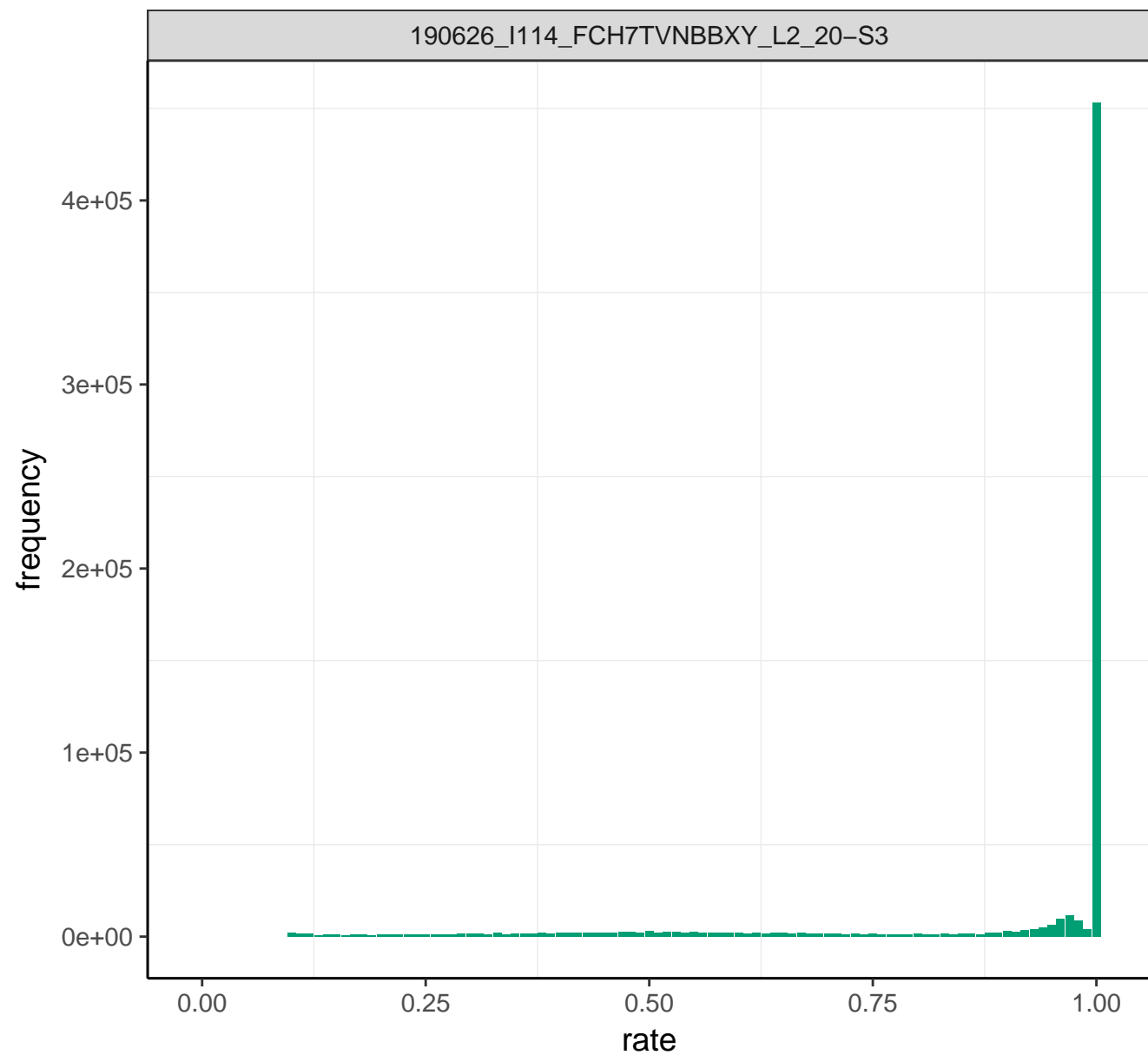


methylation rates at all positions

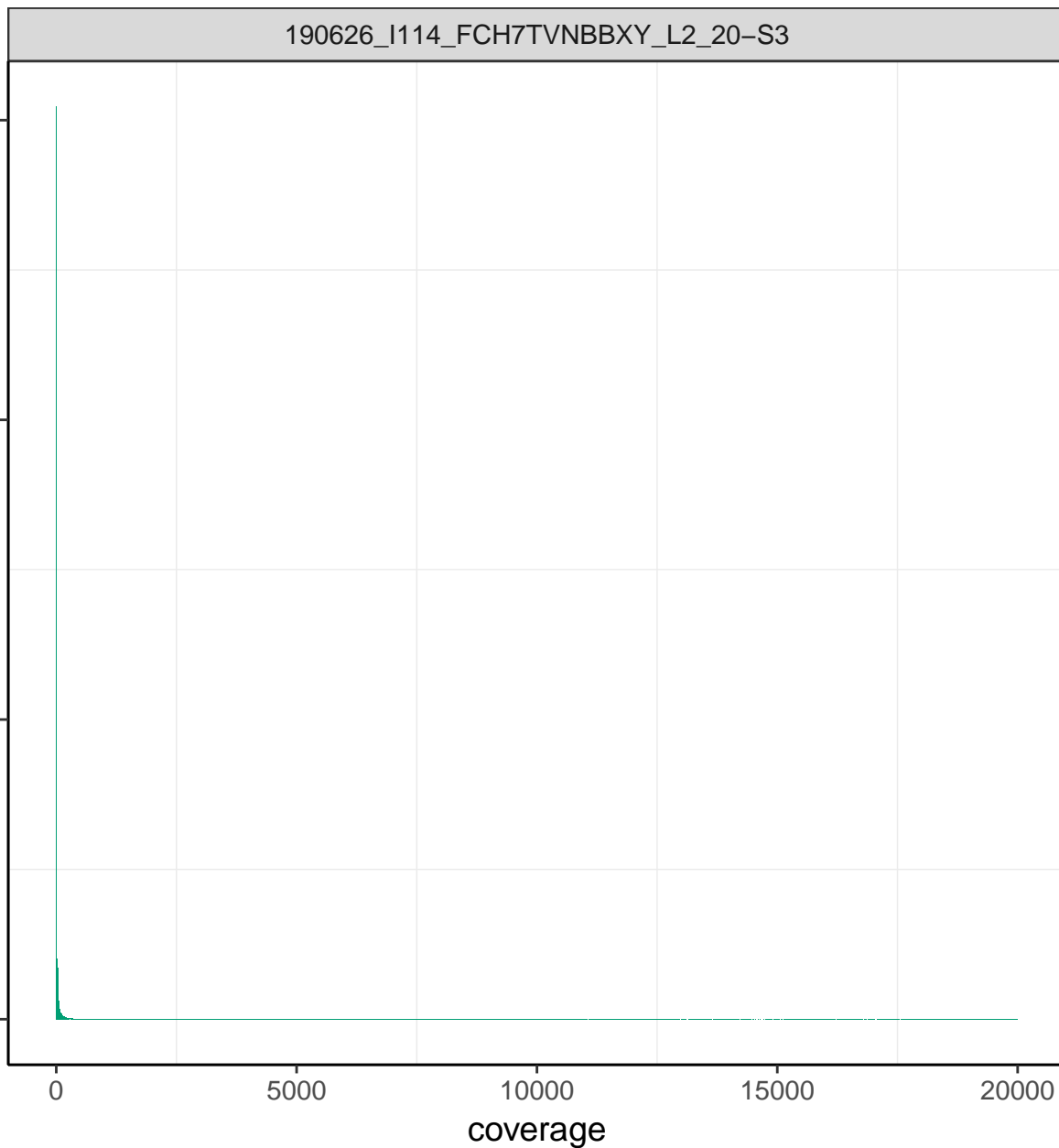
190626_I114_FCH7TVNBBXY_L2_20-S3



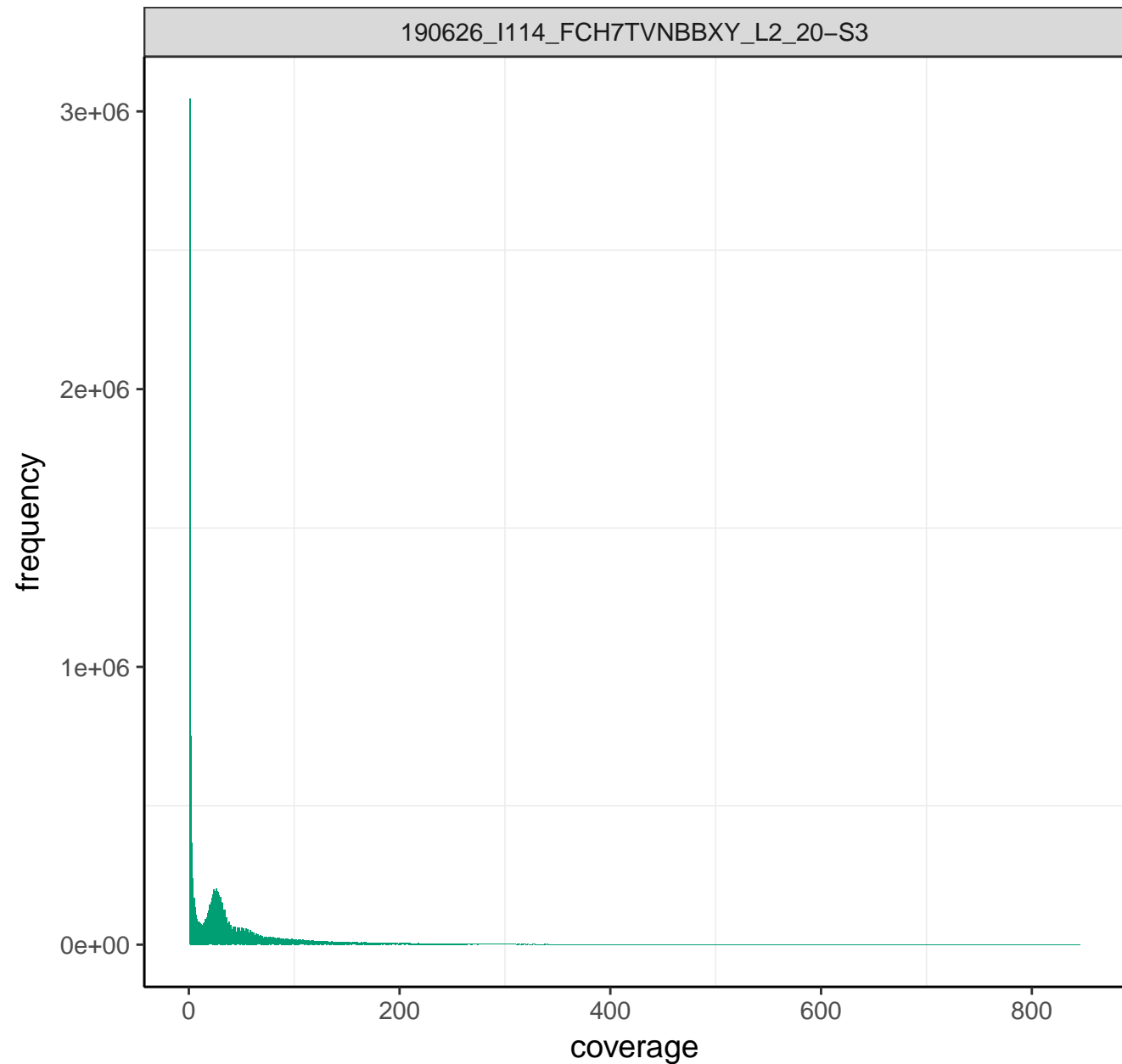
methylation rates at filtered positions
context:CG, min MDP:10, max MDP:100, min MR:0.1



coverages at all positions



coverages at all positions, in 1–846 (90%–quantile)



coverages at filtered positions

context:CG, min MDP:10, max MDP:100, min MR:0.1

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