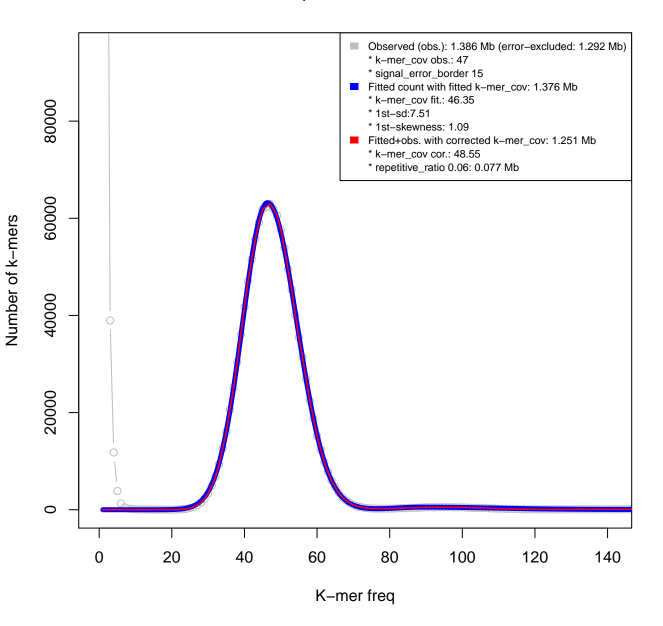
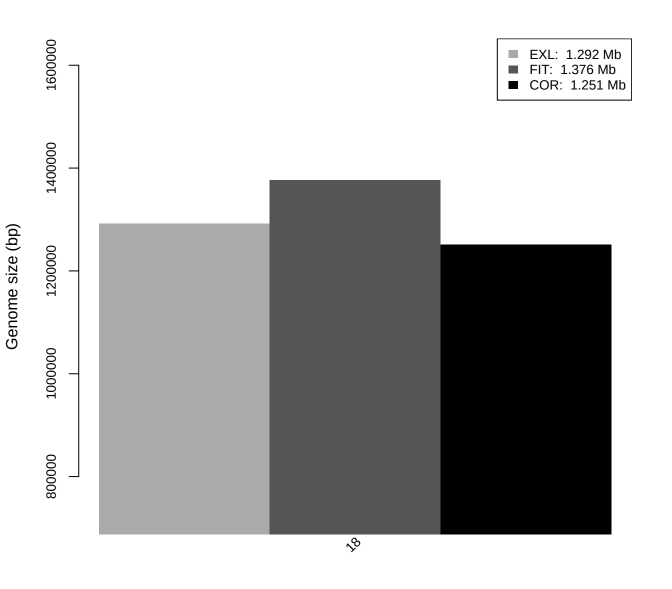
Sample sim2.histo k= 18



Genome size estimation by error-excluding, fitting, and correcting



Size k