ELM: Enhanced lowest common ancestor based method for detecting a pathogenic virus from a large sequence dataset

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Supplementary data

This file contains Supplementary Figures S1 and S2 with legends.

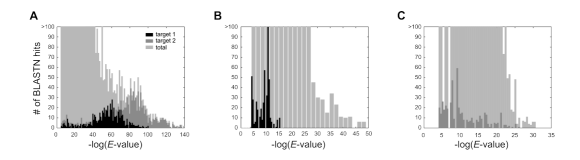


Figure S1. Distribution of *E*-values in BLASTN viruses for (A) rodent, (B) reptile and (C) simian samples.

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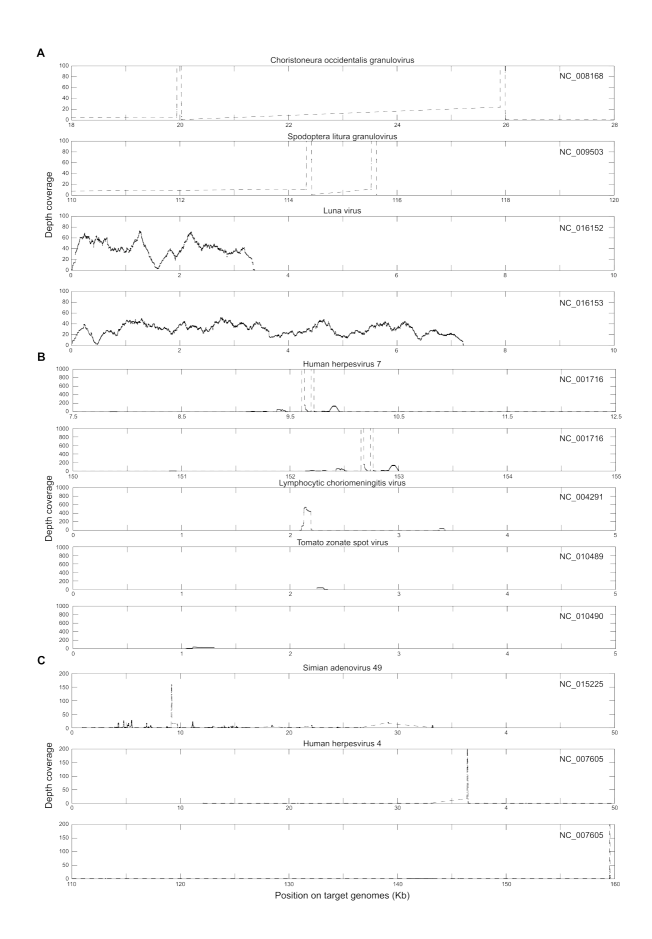


Figure S2. Read coverage on the position of target genomes for (A) rodent, (B) reptile and (C) simian samples.