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Correction: Protein embeddings reveal a continuous molecular landscape of host adaptation in waterfowl parvoviruses

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A Correction on

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The Data Availability statement was erroneously given as “The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.”

The correct Data Availability statement is “All data and code used in this study are available at <https://github.com/ShaoNihui/gpv-hostadaptation-embeddings.git>. The repository contains the input datasets (alignments, embeddings, distance matrices, structural input files), all custom analysis scripts, and the Conda environment file (environment.yml) specifying the software versions used, and is provided for community use and reference.”

The original version has been updated.

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