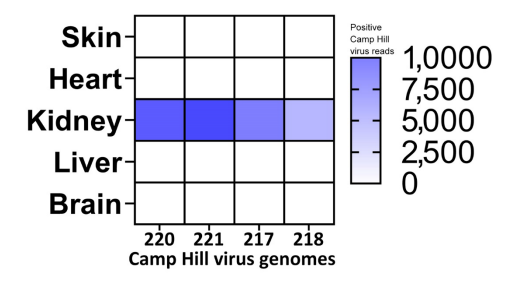
Article DOI: http://doi.org/10.3201/eid3102.241155

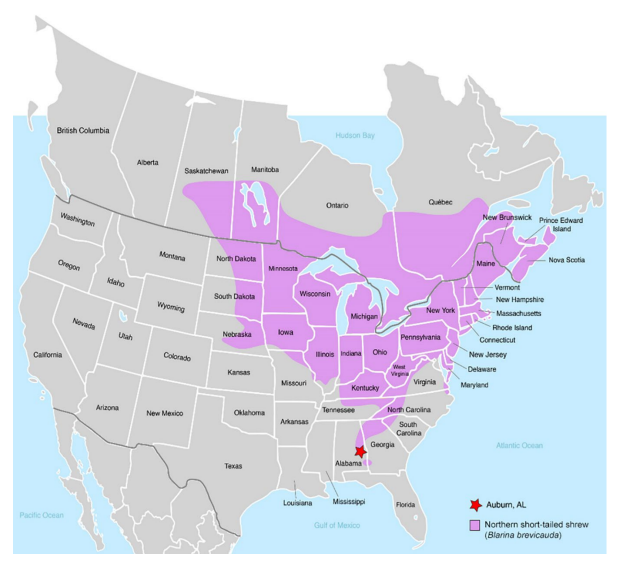
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Henipavirus in Northern Short-Tailed Shrew, Alabama, USA

Appendix



Appendix Figure 1. Tissue tropism of Camp Hill virus across various tissues from *Blarina brevicauda* (genomes 220, 221, 217, 218; Genbank ID PQ140948-PQ140951). Abundance data was derived from trimmed RNA-seq reads mapped to the Camp Hill virus genome using Bowtie2 (https://bowtie-bio.sourceforge.net/bowtie2), illustrating viral distribution across different tissue types.



Appendix Figure 2. *Blarina brevicauda* incidence data in North America and Camp Hill virus-positive samples from Auburn, Alabama, USA. Incidence data adapted from *Histoire naturelle des mammifères du Canada, 2016* (Naughton et al, 2016;

https://www.editionsmichelquintin.ca/produit/1234-histoire-naturelle-des-mammiferes-ducanada.html).