How many insect viruses are there?

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The core philosophy

This is a back-of-the-envelope estimate of the global diversity of insect-infective viruses. It uses an approach developed by Carlson *et al.* (2019) *Nature Ecology and Evolution*, which was used at the time to estimate global viral diversity in mammals. That approach estimates total symbiont diversity by (1) generating a bipartite scaling curve from a host-symbiont association dataset, and extrapolating to a higher total number of hosts; then (2) using a more complete metagenomic dataset on one specific host species to correct for undersampling in the association data. For example, in that study, there are two species - a bat and a monkey - that have been fully inventoried. After using the HP3 dataset to estimate total mammal virus diversity, that number gets multiplied by 1 / (the proportion of monkey/bat viruses in HP3 out of all of their known viruses).

We're going to try the same thing here. On Twitter, Eddie Holmes said that *Drosophila melanogaster* probably has the best-sampled insect virome. We're going to go to GenBank and use that to grab every named *D. melanogaster* virus. That'll be our "complete" species. Then, we're going to use the insectPathogen package that Tad shared to generate our edgelist of insects and viruses, and use the codependent package I made in 2019 to estimate insect virus diversity. We'll use an estimate of ~6 million insects on Earth, based on:

Larsen, Brendan B., et al. "Inordinate fondness multiplied and redistributed: the number of species on earth and the new pie of life." The Quarterly Review of Biology 92.3 (2017): 229-265.

Some install

If you need these packages, you can install both from Github. However, because insectDisease is still private, you'll need to clone it and install locally.

You will also need to unzip the file called GenBank_as_Edgelist.zip before you go any further (yes, the file is too big for GitHub, no, I don't feel like solving that problem a responsible way in the first version of this).

Some loading

```
library(codependent)
library(InsectDisease)
library(tidyverse)
```

How many viruses does *Drosophila melanogaster* have?

```
setwd("~/Github/drosophily")
gb <- read_csv('GenBank_as_Edgelist.csv')</pre>
##
## -- Column specification -----
## cols(
    Host = col_character(),
    Species = col_character()
##
## )
gb %>%
  filter(Host == 'Drosophila melanogaster') %>%
  select(Host, Species) %>%
  unique() -> dm
dm %>%
 nrow()
## [1] 30
dm %>% pull(Species)
   [1] "Esparto virus"
##
##
   [2] "Tomelloso virus"
## [3] "Drosophila melanogaster sigmavirus"
## [4] "Kallithea virus"
## [5] "Drosophila melanogaster totivirus SW-2009a"
## [6] "Drosophila A virus"
## [7] "Nora virus"
## [8] "Galbut virus"
## [9] "Chaq virus"
## [10] "Vera virus"
## [11] "Chaq-like virus"
## [12] "Vesanto virus"
## [13] "Drosophila-associated nudivirus"
## [14] "Drosophila-associated filamentous virus"
## [15] "Yalta virus"
## [16] "Drosophila C virus"
## [17] "Mauternbach virus"
## [18] "La Jolla virus"
## [19] "Dansoman virus"
## [20] "Motts Mill virus"
```

```
## [21] "Craigies Hill virus"
## [22] "Newfield virus"
## [23] "Torrey Pines virus"
## [24] "Viltain virus"
## [25] "Drosophila virus JTM-2015"
## [26] "Drosophila reovirus"
## [27] "Thika virus"
## [28] "Flock House virus"
## [29] "Drosophila melanogaster birnavirus SW-2009a"
## [30] "Drosophila melanogaster tetravirus SW-2009a"
```

There are about thirty viruses in the NCBI taxonomy that infect D.m.

Bringing in the GenBank data

Let's grab the insect virus data, and check *D. melanogaster*:

```
id <- InsectDisease::viruses

id %>%
    filter(Host == 'Drosophila melanogaster') %>%
    pull(Virus)

## [1] Drosophila sigma virus Drosophila C virus Retrovirus (RTV)
## [4] reovirus-like particle
## 267 Levels: ... Wiseana cervinata IV
```

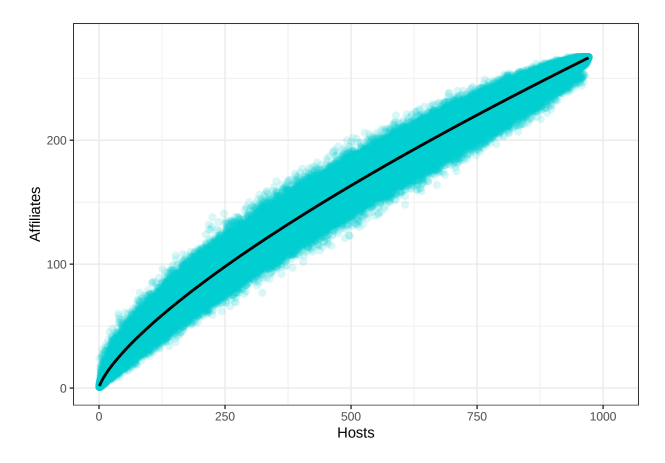
Drosophila C virus and sigma virus are named, both in GenBank. Retrovirus (RTV) and reovirus-like particle are not; reovirus is probably equivalent to Drosophila reovirus, though. So let's say there are 3 viruses here compared to 30 in GenBank - that means we'll use a multiplication factor of 10.

Building the edgelist and curve

Let's get a unique edgelist of hosts and vectors, and try building the scaling curve:

```
id %>%
  select(Host, Virus) %>%
  unique() -> assn

b <- binera(assn, iter = 100, plots = TRUE)</pre>
```



b

##

```
## Nonlinear regression model
##
     model: n.par ~ b * n.host^z
##
      data: cu
##
        b
## 1.6724 0.7373
##
   residual sum-of-squares: 8929546
## Number of iterations to convergence: 8
## Achieved convergence tolerance: 1.782e-09
Now let's try extrapolating:
c <- copredict(assn, iter = 100, n.indep = 6000000)</pre>
## [[1]]
##
     mean.b lowerCI upperCI
## 166897.6 163670.4 170182.9
##
## [[2]]
##
           b
## 1.6675970 0.7377172
```

```
## [[3]]
## 2.5 % 97.5 %
## b 1.6580443 1.6771497
## z 0.7368342 0.7386002
```

And finally, let's multiply by that correction factor:

```
c[[1]][1]*10

## mean.b
```

If there are about 6 million insect species on Earth, there should be proportionally at least ~ 1.66 million viruses in insects.

Frequently asked questions

Q: Carroll *et al.* (2018) *Science* famously estimated there are 1.67 million viruses in mammals and waterfowl, which is a number you corrected in the 2019 *Nature Ecology and Evolution* paper down to 40-60,000.

A: Yes, I was there. That's not a question, really.

Q: That 1.67 million viruses number is absurdly, weirdly close to 1.66 million insect viruses.

A: Yeah, it is.

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Q: Does that mean anything?

A: Absolutely not, but it's going to make for a ridiculous sentence in the abstract.