$_{\scriptscriptstyle 1}$ Abstract

Curated databases of species interactions are instrumental to exploring and understanding the spatial distribution of species and their biotic interactions. In the process of conducting such projects, data development and curation efforts may give rise to a data product with utility beyond the scope of the original work, but which becomes inaccessible over time. Data describing insect host-pathogen interactions are fairly rare, and should thus be preserved and curated with appropriate metadata. Here, we introduce the insectDisease R package, a mechanism for curating, updating, and distributing data from the Ecological Database of the World's Insect Pathogens, a database of insect host-pathogen associations, includ-10 ing attempted inoculations and infection outcomes for insect hosts and pathogens 11 (bacteria, fungi, nematodes, protozoans, and viruses). This dataset has been uti-12 lized for several projects since its inception, but without a well-defined, curated 13 and permanent repository, its existence and access have been limited to word-of-14 mouth connections. The current effort presented here aims to provide a means to 15 preserve, augment, and disseminate the database in a documented and versioned 16 format. This project is an example of the type of effort that will be necessary to 17 maintain valuable databases after the original funding disappears. 18

19 Running title: Ecological Database of the World's Insect Pathogens (EDWIP)

20

21 Introduction

There are a number of data sources documenting host-pathogen associations, especially for pathogens of mammals (Gibb et al., 2021, Patrick et al., 2017), birds (Bensch et al., 2009), and fish (Strona and Lafferty, 2012). Recent work from the Verena Consortium has developed a dynamically updated host-virus association database for all vertebrate hosts (VIRION) (Carlson et al., 2021), representing the largest collection of host-virus association data to date. These resources have been fundamental to our understanding of what determines pathogen host range, 28 pathogen species richness across a set of hosts, and overall host-pathogen network structure (e.g., Carlson et al. (2020), Dallas et al. (2018)). But while some host groups are well-studied, there are taxonomic gaps in our understanding of host-31 pathogen associations. Insect host-pathogen relationships have considerably less 32 open-source data available, despite their inherent importance to scientific studies and assessments of impacts to agricultural crops and spread of vector-borne disease, in addition to the sheer numerical dominance of insect species over other taxa (Stork et al., 2015). This is a clear knowledge gap.

Many of the existing species interaction databases have dedicated researchers, resources, and infrastructure to enable data deposition and curation in openly accessible formats. However, some data have not been as lucky, at no fault of the original data curators. These data run the risk of disappearing into a file drawer or on an external hard drive, potentially shared with a small number of researchers

- but not accessible to the scientific community at large. One data resource arguably close to this point of disappearance is the *Ecological Database of the World's Insect Pathogens* (EDWIP) (Onstad, 1997).
- The EDWIP data consist of experimental infections and field observations of
 the interactions between insect hosts and a number of bacterial, fungal, nematode, protozoan, and viral pathogens (Braxton et al., 2003). One particularly
 unique component of EDWIP is the existence of negative associations attempts
 to inoculate a host with a given pathogen that failed to infect for some host
 groups (Figure 1). Failed infections represent *true* absences or incompatibilities
 between a given host and pathogen. These data are incredibly useful to pathogen
 host range estimation and host-pathogen interaction modeling, but we rarely have
 data on these known non-interactions.
- Initially created in 1992, the data have been updated prior to 2000, but no clear semantic versioning was used. As such, it is unclear how long or how frequently this updating and curation continued, and thus, how many different versions of the data may be in existence presently. The database we present here, as the backbone of this R package, represents the most up-to-date version that we know of, though this may differ slightly from previous descriptions of the data (Braxton et al., 2003). Generally, we have attempted to preserve all of the original data in the original format.

Solution statement

To preserve these data in a format that is well-documented, openly accessible, versioned, and flexible for continued development, we created the insectDisease
R package. In doing so, we implicitly adhere to the FAIR (Findable, Accessible, Interoperable, Resuable) guidelines for managing data (Wilkinson et al.,
2016). By hosting the data openly on GitHub, and versioning releases of the data
with a permanent identifier (DOI), we ensure the longevity and versioned curation of this data resource. Finally, the incorporation of taxonomic data through
taxize (Chamberlain and Szöcs, 2013) ensures that host and pathogen taxonomic
names are updated periodically to accommodate for dynamic data or changing
taxonomies.

Data specification

Package structure Data products are broken down by pathogen group; nematodes (data(nematode)), viruses (data(viruses)), and non-viral pathogens,
which include protozoan, fungi, and bacteria (data(nvpassoc)). Data on negative associations is stored collectively instead of being delineated by pathogen
group (data(negative)), but information on pathogen group is provided within
each of these files, allowing for sorting of negative interactions based on the initial
pathogen groupings (Table 1). This data structure is inherited from the original
structure of the EDWIP data files, and code to process and join these different

data files is provided in the R package vignette.

Each of the pathogen groups differs slightly in the available ancillary data on experimental infections. For instance, nematode infections contain information on soil type and associated bacteria, virus infection data have information on viral dose, and non-viral pathogens (protozoans, fungi, and bacteria) have information on intermediate host species. We recommend the user explore these data and associated metadata from within R, as the metadata and data are neatly in the same place.

Data are also available on the insect host species themselves (e.g., data(hosts)).

These data contain some information on the Canadian province where the host is

found (ProvinceI column), what it eats (Food column), and what type of habitat

it is found in (Habitat column). Additionally, a column on host insect pest status

is present, offering the opportunity to explore study effort and pathogen specificity

dependent on the pest status of the insect host.

FAIR data The FAIR principles represent guidelines for making data more persistent, findable, and well-documented. Structuring the data as an R package ensures that metadata and data are packaged together, where R manual files contain column names and data descriptions for each data product (Findable). All code to take data from the raw data (data-raw folder) to the end product .RData and .csv files is contained in the versioned R data package, and integration with

Zenodo (https://doi.org/10.5281/zenodo.5821896) provides a DOI for each release (Accessible). Metadata are available in redundant forms, both from within 103 the R package as man files, and in the project README file such that installa-104 tion of the package (or navigation into the man folder) is not necessary. Apart 105 from providing data in these multiple formats, user access is aided by structuring 106 the data as a package in a very popular computing language among biologists 107 (and other folks too) and providing all code for data processing and serving in an 108 open and public-facing repository (Interoperable). Having all code and data in a streamlined, open, and versioned format, serving the data through an interactive 110 web portal, and publishing this software note collectively serve to promote the use 111 of this data resource (*Reusable*).

Metadata and package documentation Differences in features across the data on different pathogen types (e.g., ?nematodes relative to ?viruses) make 114 combining these data non-straightforward, without a degree of loss of information. 115 We provide some example code in the package vignette on how to go about 116 combining or linking the data across types, with the caveats of information loss, 117 and have standardized some key column names across the different data products. Further, we have documented each data resource using R package documentation, 119 allowing the metadata of each data product to be examined directly from R using 120 the help() function or the question mark notation (e.g., ?viruses). These same 121 metadata are also provided in the README file in the top-level of the GitHub 123 repository.

Data cleaning and taxonomic resolution We attempted to maintain as much of the original data structure from the raw data files provided by David Onstad, principal maintainer of the EDWIP data resource (Onstad, 1997). This includes files such as new_assoc, as this was likely a test file containing pathogen species such as "wormy thing", and newnema, a dataset identical to nematode. We document these idiosyncrasies in the metadata for each data product, providing a clear overview of the state of each data subproduct.

The first, and perhaps most important, novel augmentation, is the resolution of 131 host and pathogen taxonomic information. We achieved this by using the R pack-132 age taxize, specifically the NCBI taxonomic backbone (Chamberlain and Szöcs, 133 2013), making the data interoperable with existing data efforts by the Verena Consortium (e.g., VIRION; Carlson et al. (2021)). Cached versions of host and 135 pathogen taxonomic information are provided (data(hostTaxonomy) and 136 data(pathTaxonomy)), and the R code to generate these taxonomic backbones 137 and clean the data are provided in the package vignette. This taxonomic backbone 138 serves to both standardize host and pathogen nomenclature, while also correcting any taxonomic changes that have occurred in the past couple decades. This in-140 cludes the consideration of microsporidian parasites as fungi, not protozoans, a 141 change affecting a large set of records in the EDWIP data. All of the data within

the data and csv folders have already gone through these data cleaning steps.

However, these data may be dynamic, such that some form of continuous integration or updating of the host and pathogen taxonomy may be necessary. As such,

we provide a vignette which transparently shows the steps to clean and augment

the data resource, as well as reproduce figures from this manuscript. Finally, we

opt to store processed data in the csv folder, which contains all data files in .csv

format. This allows non-R users to access the csv-formatted data easily, and en
sures long-term stability of the data, as csv is a stable text file format. These data

are also provided as .rda files in the data folder.

Maintaining the data dynamically as described above allows users to access 152 the data programmatically or as versioned flatfiles (i.e., .csv files). However, for 153 users who do not wish to download the entire data resource, and simply want to 154 quickly query a static version of the database, there is also a standalone web user 155 interface (https://edwip.ecology.uga.edu/) that allows users to easily subset 156 and explore the data. The web interface serves arguably the most important 157 subset of the overall data (data files nematode, viruses, nvpassoc, negative, and 158 hosts). This interface allows users to quickly query based on host or parasite taxonomy as a dropdown list. This is perhaps more useful as a teaching tool or 160 for initial exploration of the data, while the programmatic interface and dynamic 161 data may be more useful for more rigorous analysis. This version of the EDWIP 162 data will also only be deployed with a single static copy of the data, such that users wanting to benefit from versioned and dynamic data will need to access the
data through the GitHub repository. Future efforts to integrate the web interface
and the existing dynamic data structure will be explored, but this is not currently
integrated.

Case study: covariance among pathogen groups in parasite species richness

Hosts that are infected by more pathogens of one type may also be more infected by 170 pathogens of another type, mediated by host life history traits, metabolic demands, 171 geographic distribution, and intensity of scientific study (Dallas and Becker, 2021). 172 We explore this in the EDWIP data by measuring the number of known positive associations of each of the pathogen groups for each insect host species, visualizing 174 the relationship between the number of pathogens per insect host as a correlation 175 matrix (Figure 2). We find very little evidence that pathogen groups have positive 176 covariance, which would be expected if host species traits or trait-based sampling 177 biases drove infection process across pathogen groups in the same manner. The 178 failure to detect strong positive relationships, and indeed some negative relation-179 ships appearing, could be a signal of the targeted nature of data collection, as 180 many insect host species were selected to study due to their potential as a crop 181 pest, and many pathogens were selected to study based on their potential use as biocontrol or perhaps for their ease of culture. 183

This potential sampling bias among insect host species would be evident if there 184 were a positive relationship between the number of positive interactions and the 185 number of negative interactions for a host species, as it would indicate that host 186 species with lots of known interactions also tended to appear in many studies and 187 have some negative interactions as well. We find evidence for a significantly neg-188 ative relationship based on a Spearman's rank correlation ($\rho =$ -0.1, p < 0.0001), 189 indicating no discernible influence of this relationship. This does not imply that 190 there is no sampling bias in the insect host species researchers opt to study, but that such bias was not so strong as to be clearly detected. 192

193 Concluding comments

While ecological data are growing in availability, size, accessibility, and stability, 194 there are still data resources that are aging in place, and should not be allowed 195 to fade out of existence. The EDWIP data provided to the authors were in a proprietary format ('Claris FileMaker Pro 5') that was already over 10 major 197 versions behind. With limited inter-version operability (e.g., .fmp5 files cannot 198 be opened in more recent versions of the software, or require multiple conversion 199 steps), these data seemed as if headed towards obsolescence. The insectDisease 200 package ensures that these data will be available to the broadest set of researchers, 201 be bound to relevant metadata, and be properly versioned. By hosting the data 202 openly, we welcome contributions from researchers interested in augmenting the 204 data or building off the existing resource.

205 Data accessibility

- ²⁰⁶ The insectDisease R package is currently available on GitHub
- (github.com/viralemergence/insectDisease), with '.csv' files in the csv directory
- 208 for long-term data stability. GitHub releases of the data ensure versioning is
- 209 maintained and all versions are accessible. At the time of this writing, the current
- version is 1.2.0 (available at
- ${\tt ^{211}} \quad https://github.com/viralemergence/insectDisease/releases/tag/1.2.0).$
- 212 Releases are given a DOI through integration with Zenodo (available at
- 213 https://doi.org/10.5281/zenodo.5821896).

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Tables

Table 1: Files associated with the EDWIP data resource. Metadata is stored in R package documentation, allowing the data and metadata to be intrinsically linked. For instance, users can use the help functionality from within R to see more information on data columns and unit (e.g., ?nematode).

filename	rows	columns	description
assocref	11005	16	references for some host-pathogen associa-
			tions
citation	1966	7	references but no host-pathogen association
			information
hosts	4392	21	insect host trait data
${\tt hostTaxonomy}$	4489	7	host taxonomic data updated with the
			getNCBI() function
negative	529	21	information on negative host-pathogen asso-
			ciations
nemaref	338	5	references from nematode pathogens
nematode	234	24	host-nematode interaction data
new_asso	19	25	likely a training document (perhaps do not
			use)
noassref	569	16	references for some host-pathogen associa-
			tions
nvpassoc	7164	23	non-viral pathogen infection data
pathogen	2041	9	pathogen trait data
pathTaxonomy	2282	7	pathogen taxonomic data updated with the
			getNCBI() function
viraref	2124	16	references from viral infections
viruses	1659	25	host-viral interaction data

Figures Figures

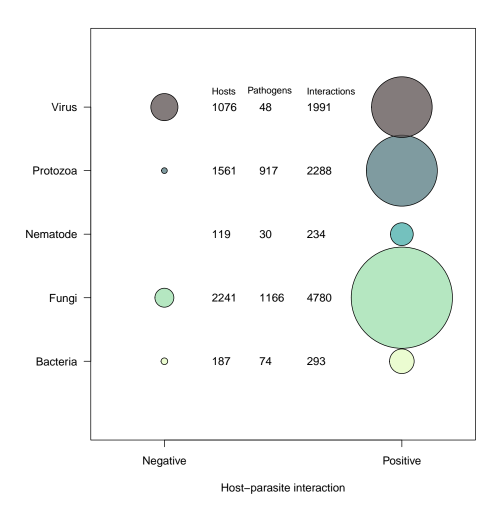


Figure 1: The number of known non-interactions (negative left panel) and known interactions (positive right panel) for the set of bacterial, fungal, nematode, protozoan, and viral pathogens (y-axis). Bubble size is proportional to the total number of interactions associated with that pathogen group and interaction type (i.e., negative or positive). Numeric columns correspond to the number of unique host species, pathogen species, and interactions for each pathogen group.

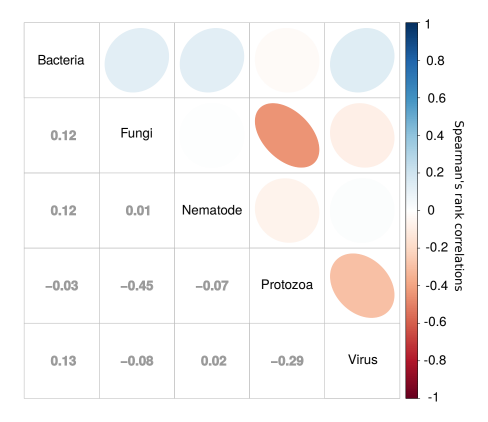


Figure 2: Correlations between each pathogen group in terms of pathogen richness of insect host species, where color corresponds to Spearman's rank correlation values (provided in the lower diagonal matrix). Fungal and protozoan pathogens were negatively related, as were viruses and protozoans. Understanding to what extent this is driven by sampling effects or insect host ecology is an outstanding research question that these could be used to begin addressing.