# Ropensci fellowship application

#### Goals

Large datasets containing community taxonomic information are becoming increasingly common, but the hierarchical nature of taxonomic information makes manipulating these datasets difficult and many R packages made for this have similar, yet unique, data structures and functions. These differences make it difficult to use multiple packages on the same dataset and causes programmers to spend time creating and maintaining analogous solutions for the same low-level problems of data storage and manipulation, rather than focusing on novel functionality. These inefficiencies are likely to get worse as the cost of high-throughput sequences decreases, the interest in microbiome research increases, and the number of associated R packages increases. To address this problem, we are developing what we hope will become a standard in R for storing, manipulating, and plotting taxonomic information and any other application-specific data associated with it. Being awarded this fellowship will allow us to devote the time needed to bring this project to maturity and help integrate it into the community of R programmers working with taxonomic data. A common infrastructure will allow programmers to make focused, novel contributions without creating redundant functionality. A cohesive community of packages will encourage users to conduct analyses entirely within R, increasing the effectiveness of reproducibility tools like packrat and Rmarkdown.

We are working on developing the Ropensci package taxa, in collaboration with Sckott Chamberlain, to provide a set of all-purpose classes and associated manipulation functions for taxonomic data, modeled after the dplyr data-manipulation philosophy. In addition to classes that represent taxa, classifications, and taxonomic trees, the taxa packages has a class that stores any number of application-specific tables, lists, or vectors mapped to a taxonomic tree so that manipulations to the taxonomy are also applied to the corresponding data and visa-versa. Manipulation functions take into account the hierarchical relationships between taxa as well. For example, when filtering taxa or associated data, the supertaxa or subtaxa of taxa that meet some condition can be preserved or discarded. The taxa package also implements extremely flexible parsers to read data in from nearly any format containing taxonomic information.

Although the taxa package is already useful, to make it a solid foundation for the community to build on, we want to make it faster, more robust, and work to integrate it with enough other packages. By testing with very large data sets, we want to determine current limits and bottlenecks so that we can identify parts of code to refactor or replace with C++ to increase its speed. We also want to implement parallel processing to make better use of modern multi-core computers. Since we hope to make this a foundational package, we want to add extensive unit tests in addition to the many already implemented, to minimize bugs and their effect on dependent packages. Perhaps most importantly, we want to work with the maintainers of popular packages using taxonomic data to integrate taxa and jumpstart adoption. We have just started the process of integrating taxa into taxize, the primary package for downloading taxonomic data from public databases, and metacoder, our package for analysis and visualization of community taxonomic data. Once taxize and metacoder are refactored to use taxa, database searching/downloading and flexible visualization will be easy using the classes implemented by taxa, providing useful tools for a wide variety of research objectives and R packages. This useful, all-purpose functionality will incentivize others to adopt the taxa package as a standard.

The other related project we are working on is metacoder, an R package for visualization and analysis of community taxonomic data, with an emphasis on providing tools for microbiome research. metacoder introduced a way of visualizing statistics distributed throughout a hierarchy (e.g. a taxonomic tree) by quantitatively mapping statistics to the the color or size of nodes and edges in a tree. We call these "heat trees" and they have been enthusiastically received as an intuitive and information-dense alternative to stacked bar charts, which is the most common way taxon abundance in communities are currently plotted in publications. The heat\_tree function in metacoder is highly flexible, yet only requires a few arguments to produce high-quality figures in most cases, since raw statistics are automatically mapped to color/size and the size range of elements displayed are optimized for each graph to strike a balance between avoiding overlaps and maximizing the apparent differences. metacoder also provides a way to simulate PCR in R, which is important for selecting primers for metabarcoding research.

metacoder is currently being significantly refactored to use the classes provided by taxa and add more functionality specific to community taxonomic data analysis. We are considering splitting out the low-level plotting abilities of metacoder into its own package, since these are universally useful for any hierarchical data, not just community taxon abundance data from high-throughput sequencing. In that case, metacoder would focus exclusively on aiding analysis of community taxonomic data, generally from high-throughput sequencing. One particularly useful high-level visualization technique we want to facilitate is a matrix of unlabeled heat trees that show which taxa are differentially abundant between every pair-wise set of treatments accompanied by a larger, labeled tree that functions as a key. This can be used to show the results of thousands of statistical tests and the treatments/taxa they apply to in a graph that can fit in a publication. Currently, making these graphs is quite complicated, but we believe we can design a function that will make creating these graphs easy for users. We also want to implement pairs of parser and writer functions that handle the common formats used in metabarcoding research to make it easy for users to get their data into R. These parsers will return taxa objects, so they would be very useful for all packages that adopt the taxa package as a standard. They could also be used to convert between file formats, since a user could read from one format and write to another. The taxa classes are flexible enough that we should be able to losslessly read and write the same format, so this would provide a way to subset or otherwise modify large, complicated files, using the powerful manipulation functions provided by taxa.

Encouraging adoption of this framework by both users and developers is key to the success of this project. We hope to facilitate adoption by developers by offering to help maintainers adapt existing packages to taxa. Once a critical mass of packages have adopted taxa, we expect that it will naturally be adopted by new packages without our direct assistance. To attract users, we want to write a series of posts on major R blogs that cover specific uses of the framework and demonstrate its usefulness. We also plan to present this work at biological and computational conferences, including the UseR 2018 conference and the Ropensci 2018 unconf. We have already submitted a workshop proposal for the 2018 11th International Congress of Plant Pathology joint conference on reproducible analysis of microbiome data in R, which will focus on applying these tools. Finally, we hope to publish a paper on the taxa package in the F1000 open access journal. If this framework is adopted by the community has we hope, we would also want to publish an article describing the resources available for all-R analyses of taxonomic data using this framework.

## Expected outcomes and timeline

## Improvements to the taxa package

- Add missing dplyr analog functions (e.g. summarise, group\_by, etc)
- Add ability to use multiple cores to improve speed
- Port slowest parts of the code to C++ to improve speed
- Add exhaustive unit tests (test coverage of > 95%)

#### Improvements to metacoder

- Finish the transition to using taxa
- Add ability to plot multiple colors per node/edge automatically when multiple values per taxon are supplied. Nodes would be pie charts and edges would be stacked bar charts.
- Add function to easily create a heat tree matrix for pairwise comparison of treatments.
- For each major file format used in metabarcoding research, add a function to parse the file's contents into the classes provided by taxa and a corresponding writer to recreate the file format. We read from and written to the same format, the process should be lossless.
- Add automatic overlap avoidance for labels
- Add ability to highlight/delineate groups of taxa with shaded polygons
- Add support for plotting categorical information
- Add ability to automatically query, download and plot silhouettes representing taxa from the pylopics database in place of nodes.

- Add ability to plot user-supplied images in place of nodes.
- Add support for interactive plots.
- Increase support for simulated PCR and barcode gap analysis

### Outreach and community building

- Contact the maintainers of major packages using taxonomic information and offer to help adopt taxa.
- Finish refactoring of taxize to use taxa
- Present at the 2018 rOpenSci Unconf
- Present at the 2018 useR! conference
- Conduct a workshop and reproducibile microbiome analysis in R using these tool at the 2018 11th International Congress of Plant Pathology.
- Write a series of blogs for R-bloggers demonstrating the usefullness of the taxa/taxize/metacoder framework.
- Publish a journal aricle on taxa in F1000Research, or another open-access journal.
- Publish a journal aricle on reproducible microbiome analysis in R in an open-access journal.

### **Collaborators**

Niklaus Grünwald

## High level budget

#### 2018 useR! conference

- Airfair from Portland, OR to Brisbane Austrialia: \$1,400
- Per diem meals and incidentals Brisbane Austrialia:  $98 \times 4 \text{ days} = 392$
- Per diem lodging Brisbane Austrialia:  $$159 \times 4 \text{ days} = $636$
- Student registration fee: \$125

Estimated subtotal: \$2553

Salery