## Species occurrence data (spocc), version 0.08

#### Introduction

The rOpenSci projects aims to provide programmatic access to scientific data repositories on the web. A vast majority of the packages in our current suite retrieve some form of biodiversity or taxonomic data. Since several of these datasets have been georeferenced, it provides numerous opportunities for visualizing species distributions, building species distribution maps, and for using it analyses such as species distribution models. In an effort to streamline access to these data, we have developed a package called Spocc, which provides a unified API to all the biodiversity sources that we provide. The obvious advantage is that a user can interact with a common API and not worry about the nuances in syntax that differ between packages. As more data sources come online, users can access even more data without significant changes to their code. However, it is important to note that spocc will never replicate the full functionality that exists within specific packages. Therefore users with a strong interest in one of the specific data sources listed below would benefit from familiarising themselves with the inner working of the appropriate packages.

#### **Data Sources**

spoce currently interfaces with five major biodiversity repositories. Many of these packages have been part of the rOpenSci su

- 1. Global Biodiversity Information Facility (rgbif)
  - GBIF is a government funded open data repository with several partner organizations with the express goal of providing access to data on Earth's biodiversity. The data are made available by a network of member nodes, coordinating information from various participant organizations and government agencies.
- 2. Berkeley Ecoengine (ecoengine)
  - The ecoengine is an open API built by the Berkeley Initiative for Global Change Biology. The repository provides access to over 3 million specimens from various Berkeley natural history museums. These data span more than a century and provide access to georeferenced specimens, species checklists, photographs, vegetation surveys and resurveys and a variety of measurements from environmental sensors located at reserves across University of California's natural reserve system.
- 3. iNaturalist (rinat) iNaturalist provides access to crowd sourced citizen science data on species observations.
- 4. VertNet (rvertnet) Similar to rgbif, ecoengine, and rbison (see below), VertNet provides access to more than 80 million vertebrate records spanning a large number of institutions and museums primarly covering four major disciplines (mammology, herpetology, ornithology, and icthyology). Note that we don't currenlty support vertnet data in this package, but we should soon
- 5. Biodiversity Information Serving Our Nation (rbison)
  Built by the US Geological Survey's core science analytic team, BISON is a portal that provides access to species occurrence data from several participating institutions.
- 6. eBird (rebird)
  - ebird is a database developed and maintained by the Cornell Lab of Ornithology and the National Audubon Society. It provides real-time access to checklist data, data on bird abundance and distribution, and community reports from birders.

**Notes:** It's important to keep in mind that several data providers interface with many of the above mentioned repositories. This means that occurrence data obtained from BISON may be duplicates of data that are also available through GBIF. We do not have a way to resolve these duplicates or overlaps at this time but it is an issue we are hoping to resolve in future versions of the package.

#### Data retrieval

The most significant function in spoce is the occ (short for occurrence) function. occ takes a query, often a species name, and searches across all data sources specified in the from argument. For example, one can search for all occurrences of Sharp-shinned Hawks (Accipiter striatus) from the GBIF database with the following R call.

```
library(spocc)
df <- occ(query = "Accipiter striatus", from = "gbif")
df

## Summary of results - occurrences found for:
## gbif : 25 records across 1 species
## bison : 0 records across 1 species
## inat : 0 records across 1 species
## ebird : 0 records across 1 species
## ecoengine : 0 records across 1 species
head(df$gbif$data[[1]])</pre>
```

```
##
                             key longitude latitude prov
                                             4.724 gbif
## 1 Accipiter striatus 768992325
                                 -76.10
                                  -97.32
## 2 Accipiter striatus 773408845
                                            32.821 gbif
## 3 Accipiter striatus 773414146
                                 -122.27
                                            37.771 gbif
## 4 Accipiter striatus 859267562
                                  -108.34
                                            36.732 gbif
## 5 Accipiter striatus 859267548
                                   -108.34
                                            36.732 gbif
## 6 Accipiter striatus 859267717
                                   -108.34
                                            36.732 gbif
```

The data returned are part of a S3 class called occdat. This class has slots for the five data sources described above. One can easily switch the source by changing the from parameter in the function call above.

Within each data source is the set of species queried. In the above example, we only asked for occurrence data for one species, but we could have asked for any number. Let's say we asked for data for two species: *Accipiter striatus*, and *Pinus contorta*. Then the structure of the response would be

If you only request data from gbif, like from='gbif', then the other four source slots are prsent in the response object, but have no data.

You can quickly get just the data by indexing to the data element, like

## head(df\$gbif\$data\$Accipiter\_striatus)

```
##
                              key longitude latitude prov
## 1 Accipiter striatus 768992325
                                     -76.10
                                               4.724 gbif
## 2 Accipiter striatus 773408845
                                     -97.32
                                              32.821 gbif
## 3 Accipiter striatus 773414146
                                    -122.27
                                              37.771 gbif
## 4 Accipiter striatus 859267562
                                    -108.34
                                               36.732 gbif
## 5 Accipiter striatus 859267548
                                    -108.34
                                               36.732 gbif
## 6 Accipiter striatus 859267717
                                    -108.34
                                               36.732 gbif
```

When you get data from multiple providers, the fields returned are slightly different, e.g.:

```
df <- occ(query = "Accipiter striatus", from = c("gbif", "ecoengine"))
head(df$gbif$data$Accipiter_striatus)</pre>
```

```
##
                              key longitude latitude prov
                   name
## 1 Accipiter striatus 768992325
                                      -76.10
                                                4.724 gbif
                                      -97.32
## 2 Accipiter striatus 773408845
                                               32.821 gbif
## 3 Accipiter striatus 773414146
                                     -122.27
                                               37.771 gbif
## 4 Accipiter striatus 859267562
                                     -108.34
                                               36.732 gbif
## 5 Accipiter striatus 859267548
                                     -108.34
                                               36.732 gbif
## 6 Accipiter striatus 859267717
                                     -108.34
                                               36.732 gbif
```

#### head(df\$ecoengine\$data\$Accipiter\_striatus)

```
url
## 1 http://ecoengine.berkeley.edu/api/observations/LACM%3ABirds%3A5893/
## 2 http://ecoengine.berkeley.edu/api/observations/LACM%3ABirds%3A5883/
## 3 http://ecoengine.berkeley.edu/api/observations/LACM%3ABirds%3A5884/
## 4 http://ecoengine.berkeley.edu/api/observations/LACM%3ABirds%3A5885/
## 5 http://ecoengine.berkeley.edu/api/observations/LACM%3ABirds%3A5886/
## 6 http://ecoengine.berkeley.edu/api/observations/LACM%3ABirds%3A5887/
##
     observation_type
                                          name
                                                     country state_province
## 1
             specimen Accipiter striatus velox United States
                                                                  Minnesota
## 2
             specimen Accipiter striatus velox United States
                                                                  California
## 3
             specimen Accipiter striatus velox United States
                                                                 California
             specimen Accipiter striatus velox United States
## 4
                                                                  Minnesota
## 5
             specimen Accipiter striatus velox United States
                                                                  California
## 6
             specimen Accipiter striatus velox United States
                                                                    Illinois
                  end_date
##
     begin_date
                                                                  source
## 1 1894-09-15 1894-09-15 http://ecoengine.berkeley.edu/api/sources/2/
## 2 1899-04-11 1899-04-11 http://ecoengine.berkeley.edu/api/sources/2/
## 3 1914-11-11 1914-11-11 http://ecoengine.berkeley.edu/api/sources/2/
## 4 1893-09-23 1893-09-23 http://ecoengine.berkeley.edu/api/sources/2/
## 5 1899-04-17 1899-04-17 http://ecoengine.berkeley.edu/api/sources/2/
## 6 1908-04-07 1908-04-07 http://ecoengine.berkeley.edu/api/sources/2/
    remote_resource geojson.type longitude latitude
                                                          prov
```

```
## 1
                             Point
                                       -92.11
                                                  46.78 ecoengine
## 2
                                      -118.13
                                                  34.14 ecoengine
                             Point
                                                  33.63 ecoengine
## 3
                             Point
                                      -116.15
## 4
                                       -92.11
                                                  46.78 ecoengine
                             Point.
## 5
                             Point
                                      -118.13
                                                  34.14 ecoengine
## 6
                                       -87.79
                                                  41.85 ecoengine
                             Point
```

We provide a function occ2df that pulls out a few key columns needed for making maps:

# head(occ2df(df))

```
##
                   name longitude latitude prov
## 1 Accipiter striatus
                           -76.10
                                     4.724 gbif
## 2 Accipiter striatus
                           -97.32
                                    32.821 gbif
## 3 Accipiter striatus
                          -122.27
                                    37.771 gbif
## 4 Accipiter striatus
                          -108.34
                                    36.732 gbif
## 5 Accipiter striatus
                          -108.34
                                    36.732 gbif
## 6 Accipiter striatus
                                    36.732 gbif
                          -108.34
```

#### Fix names

One problem you often run in to is that there can be various names for the same taxon in any one source. For example:

```
df <- occ(query = "Pinus contorta", from = c("gbif", "inat"), limit = 50)
head(df$gbif$data$Pinus_contorta[, 1:2])</pre>
```

```
## name key
## 1 Pinus contorta Douglas ex Loudon 856965570
## 2 Pinus contorta Douglas ex Loudon 856964858
## 3 Pinus contorta 773428981
## 4 Pinus contorta 866502616
## 5 Pinus contorta 866517786
## 6 Pinus contorta 856022134
```

#### head(df\$inat\$data\$Pinus\_contorta[, 1:2])

```
## name Datetime
## 1 Pinus contorta contorta 2014-01-17 00:00:00 +0000
## 2 Pinus contorta 2013-12-23 18:55:03 +0000
## 3 Pinus contorta 2013-12-23 19:02:55 +0000
## 4 Elaphocordyceps capitata 2013-12-11 22:05:22 +0000
## 5 Pinus contorta murrayana 2013-10-01 00:00:00 +0000
## 6 Pinus contorta 2013-11-30 20:17:00 +0000
```

This is fine, but when trying to make a map in which points are colored for each taxon, you can have many colors for a single taxon, where instead one color per taxon is more appropriate. There is a function in spocc called fixnames, which has a few options in which you can take the shortest names (usually just the plain binomials like *Homo sapiens*), or the original name queried, or a vector of names supplied by the user.

```
df <- fixnames(df, how = "shortest")</pre>
head(df$gbif$data$Pinus_contorta[, 1:2])
##
               name
                          key
## 1 Pinus contorta 856965570
## 2 Pinus contorta 856964858
## 3 Pinus contorta 773428981
## 4 Pinus contorta 866502616
## 5 Pinus contorta 866517786
## 6 Pinus contorta 856022134
head(df$inat$data$Pinus_contorta[, 1:2])
##
               name
                                      Datetime
## 1 Pinus contorta 2014-01-17 00:00:00 +0000
## 2 Pinus contorta 2013-12-23 18:55:03 +0000
## 3 Pinus contorta 2013-12-23 19:02:55 +0000
## 4 Pinus contorta 2013-12-11 22:05:22 +0000
## 5 Pinus contorta 2013-10-01 00:00:00 +0000
## 6 Pinus contorta 2013-11-30 20:17:00 +0000
df_comb <- occ2df(df)</pre>
head(df_comb)
##
               name longitude latitude prov
## 1 Pinus contorta
                       -3.630
                                  55.73 gbif
## 2 Pinus contorta
                       -3.644
                                  55.72 gbif
## 3 Pinus contorta
                     -120.040
                                  38.87 gbif
## 4 Pinus contorta
                     -122.271
                                  47.78 gbif
## 5 Pinus contorta
                     -123.803
                                  39.29 gbif
## 6 Pinus contorta
                       20.350
                                  63.71 gbif
tail(df_comb)
##
                 name longitude latitude prov
                          47.03
                                   -124.2 inat
## 95
      Pinus contorta
                           47.03
                                   -124.2 inat
## 96 Pinus contorta
## 97
       Pinus contorta
                             NA
                                       NA inat
## 98
       Pinus contorta
                           44.77
                                   -116.3 inat
## 99 Pinus contorta
                          47.03
                                   -124.2 inat
## 100 Pinus contorta
                          47.03
                                   -124.2 inat
```

#### Visualization routines

#### Interactive maps

Leaflet.js

Leaflet JS is an open source mapping library that can leverage various layers from multiple sources. Using the leafletR library, it's possible to generate a local geoJSON file and a html file of species distribution maps. The folder can easily be moved to a web server and served widely without any additional coding.

It's also possible to render similar maps iwth Mapbox by committing just the geoJSON file to GitHub or posting it as a gist. All the remaining fields will become part of a table inside a tooltip, providing a extremely quick and easy way to serve up interactive maps. This is especially useful when a user does not have access to web hosting.

Here is an example of making a leaflet map:

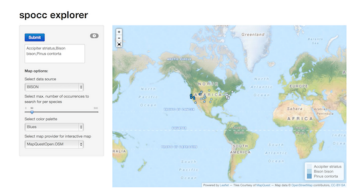
```
spp <- c("Danaus plexippus", "Accipiter striatus", "Pinus contorta")
dat <- occ(query = spp, from = "gbif", gbifopts = list(georeferenced = TRUE))
data <- occ2df(dat)
mapleaflet(data = data, dest = ".")</pre>
```



## $Shiny\ map$

We've created an interactive map using RStudio's Shiny, via the mapshiny function. This function uses the package rCharts, which you'll need to install if you want to use the mapshiny function. Instead of passing in occurrence data to the function, the function creates an interactive app in your browser (or the pop up RStudio window), in which you can enter speciess names to search for, choose data source, set limit on number of occurrences, choose the background map, and the color palette for the map symbols. You can make this map yourself if you like by looking at two files: system.file("shiny/ui.r", package = "spocc") and system.file("shiny/server.r", package = "spocc").

### mapshiny()



#### Geojson map as a Github gist

We've created an interactive map using RStudio's Shiny, via the mapgist function. This function uses the package rCharts, which you'll need to install if you want to use the mapgist function. You have to

have a Github account to use this function. Github accounts are free though, and great for versioning and collaborating on code or papers. When you run the mapgist function it will ask for your Github username and password. You can alternatively store those in your .Rprofile file by adding entries for username (options(github.username = 'username')) and password (options(github.password = 'password')).

```
spp <- c("Danaus plexippus", "Accipiter striatus", "Pinus contorta")
dat <- occ(query = spp, from = "gbif", gbifopts = list(georeferenced = TRUE))
dat <- fixnames(dat)
dat <- occ2df(dat)
mapgist(data = dat, color = c("#976AAE", "#6B944D", "#BD5945"))</pre>
```

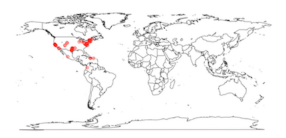


## Static maps

base plots

Base plots, or the built in plotting facility in R accessed via plot(), is quite fast, but not super easy to use, but are good for a quick glance at some data.

```
spnames <- c("Accipiter striatus", "Setophaga caerulescens", "Spinus tristis")
out <- occ(query = spnames, from = "gbif", gbifopts = list(georeferenced = TRUE))
plot(out, cex = 1, pch = 10)</pre>
```



ggplot2

ggplot2 is a powerful package for making visualizations in R. Read more about it here. We created a simple wrapper function mapggplot to make a ggplot2 map from occurrence data using the ggmap package, which is built on top of ggplot2. Here's an example:

```
ecoengine_data <- occ(query = "Lynx rufus californicus", from = "ecoengine")
mapggplot(ecoengine_data)</pre>
```



## Upcoming features

- As soon as we have an updated rvertnet package, we'll add the ability to query VertNet data from spoce.
- We're helping on a new package rMaps to make interactive maps using various Javascript mapping libraries, which will give access to a variety of awesome interactive maps. We will integrate rMaps once it's on CRAN.