Genome project tables in the genomes package

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The number of genome sequencing projects submitted to public sequence databases is growing rapidly. In addition to the raw sequence data, the amount of associated metadata describing each project is also increasing. The genomes package collects genome project metadata from NCBI (http://www.ncbi.nlm.nih.gov) and provides tools to summarize, compare and plot the data in the R programming environment.

Genome tables are a defined class (*genomes*) and each table is a data frame where rows are genome projects and columns are the fields describing the associated metadata. At a minimum, the table should have a column listing the project name, status, and release date. A number of methods are available that operate on genome tables including print, summary, plot and update.

There are a number of ways to install this package. If you are running the most recent R version, you can use the biocLite command.

```
R> source("http://bioconductor.org/biocLite.R")
R> biocLite("genomes")
```

Since the format of online genome tables may change (and then update commands may fail), I would recommend downloading the development version for fixes in between the six month release cycle.

```
R> install.packages("genomes",
    repos="http://www.bioconductor.org/packages/devel/bioC")
```

Genome tables from the Genome Project database at NCBI include prokaryotic projects (lproks), eukaryotic projects (leuks), metagenomes (lenvs) and viruses (virus). The print methods displays the first few rows and columns of the table (either select less than seven rows or convert the object to a data.frame to print all columns). The summary function displays the download date, a count of projects by status, and a list of recent submissions. The plot method displays a cumulative plot of genomes by release date (Figure 1, use lines to add additional tables).

```
R> data(lproks)
R> lproks
```

A genomes data.frame with 5439 rows and 32 columns

```
status
       pid
                                                     name
     33011
                        Abiotrophia defectiva ATCC 49176
1
                                                             Assembly
2
     12997
                          Acaryochloris marina MBIC11017
                                                             Complete
3
     16707
                            Acaryochloris sp. CCMEE 5410 In Progress
4
     45843
                          Acetivibrio cellulolyticus CD2
                                                             Assembly
5
                            Acetobacter aceti NBRC 14818
     52649
                                                             Assembly
5439 34927 Zymomonas mobilis subsp. pomaceae ATCC 29192 In Progress
       released ...
1
     2009-03-17 ...
2
     2007-10-16 ...
3
           <NA> ...
4
     2010-08-11 ...
5
           <NA> ...
            . . . . . .
. . .
5439
           <NA> ...
R> summary(lproks)
$`Total genomes`
[1] 5439 genome projects on Mar 15, 2011
$`By status`
            Total
In Progress
             2133
Assembly
             1821
Complete
             1485
$`Recent submissions`
             NAME
  RELEASED
                                                                     STATUS
1 2011-03-10 Yersinia enterocolitica subsp. palearctica 105.5R(r) Complete
2 2011-03-09 Xanthomonas gardneri ATCC 19865
                                                                     Assembly
3 2011-03-09 Xanthomonas perforans 91-118
                                                                     Assembly
4 2011-03-09 Xanthomonas vesicatoria ATCC 35937
                                                                     Assembly
5 2011-03-08 Chlamydophila psittaci 6BC
                                                                     Complete
R> plot(lproks, log = "y", las = 1)
R> data(leuks)
R> data(lenvs)
R> lines(leuks, col = "red")
R> lines(lenvs, col = "green3")
```

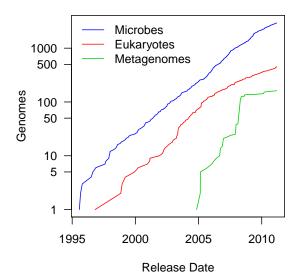


Figure 1: Cumulative plot of genome projects by release date at NCBI.

Most importantly, the update method downloads the latest version of the table from NCBI and displays a message listing the number of project IDs added and removed (not run).

R> update(lproks)

A number of additional functions assist in selecting, sorting and grouping genomes. The species and genus functions can be used to extract the species or genus from a scientific name. The table2 function formats and sorts a contingency table by counts.

R> spp <- species(lproks\$name) R> table2(spp)

	Total
Escherichia coli	338
Staphylococcus aureus	206
Salmonella enterica	140
Streptococcus pneumoniae	92
${\tt Mycobacterium\ tuberculosis}$	83
Enterococcus faecalis	73
Propionibacterium acnes	72
Acinetobacter baumannii	67
${\tt Staphylococcus}\ {\tt epidermidis}$	67
Streptococcus mutans	60

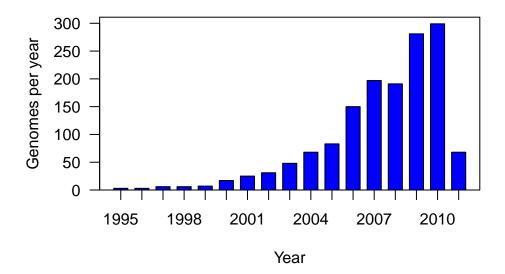


Figure 2: Number of complete microbial genomes released each year at NCBI

The month and year functions can be used to extract the month or year from the release date (Figure 2).

Because subsets of tables are often needed, the binary operator like allows pattern matching using wildcards. The plotby function can then be used to plot the release dates by status using labeled points, in this case to identify complete and draft sequences of Yersinia pestis (Figure 3).

```
R> yp <- subset(lproks, name %like% "Yersinia pestis*")
R> plotby(yp, labels = TRUE, cex = 0.5, lbty = "n")
```

A number of recent functions have been added that allow R users to run Entrez queries. For example, users can retrieve genome summaries or neighbors using a valid Entrez search query, list taxonomy names matching taxonomy ids, find the published dates of pubmed ids, or return the release dates given accession numbers. The full details about these functions and many others can be found in genomes help pages.

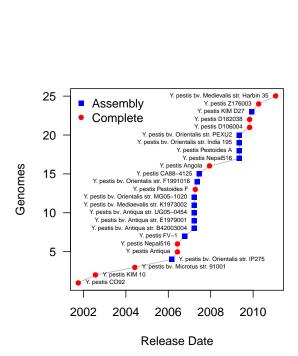


Figure 3: Cumulative plot of Yersinia pestis genomes by release date.