Genome and assembly reports

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The genomes package reads genome or assembly reports from the NCBI genomes FTP. The main function reports lists files in the GENOME_REPORTS directory (or ASSEMBLY_REPORTS if assembly=TRUE) and uses the readr package to download the tables. Additional functions to download genome features and sequences in the genbank and refseq directories will be added soon (currently FTP paths in the prokaryotes.txt files are still missing).

```
R> reports()
R> proks <- reports("prokaryotes.txt")</pre>
```