



## NCBI News, February 2014

### Genome Workbench Update 2.7.15 released

*Wednesday, February 26, 2014*

Genome Workbench 2.7.15 has been released. The update includes several new features like Multiple Alignment View, Active Objects Inspector, and binary packages for Linux OpenSUSE 13.1. The [release notes](#) include more information on features, fixes and improvements.

### New CDD Release v.3.11 includes recomputed PSSMs and more

*Wednesday, February 19, 2014*

Conserved Domain Database (CDD) version 3.11 is now available with 596 new or updated NCBI-curated and 49,641 total domain models. The new version now contains the most recent Pfam release 27.

Updates to the Conserved Domain Database include:

- Position-specific score matrices (PSSMs) have been recomputed for many models in CDD, and frequency tables have been added to the PSSMs;
- The search databases distributed as part of this release can now be used with the more recent versions of RPS-BLAST (BLAST release 2.2.28 and up) using composition-based scoring. This abolishes the need to mask out compositionally biased regions in query sequences;
- Domain annotation displays in CD-Search, BATCH CD-Search, and other services now all use a uniform display style. A new display option in CD-Search and BATCH CD-Search provides “standard” results, in addition to “concise” and “full” results. “Standard” results will provide, for each region on the query sequence, the best0-scoring domain model (if any) from each of CDD’s database providers (Pfam, SMART, COG, TIGRFAMs, Protein Clusters, and the NCBI in-house curation project), but will suppress redundancy from within a single provider's results list.

You can access CDD at the [Conserved Domains homepage](#) and find updated content on the [CDD FTP site](#).

### GenBank has milestone 200th release

*Tuesday, February 18, 2014*

GenBank's 200th release is now available through NCBI's Entrez and BLAST services.

Release 200.0 (2/12/2014) has 171,123,749 non-WGS, non-CON records containing 157,943,793,171 base pairs of sequence data. In addition, there are 139,725,795 WGS records containing 591,378,698,544 base pairs of sequence data.

During the 64 days between the close dates for GenBank Releases 199.0 and 200.0, the non-WGS/non-CON portion of GenBank grew by 1,713,261,609 base pairs and by 1,792,342 sequence records. During that same period, 4,979,722 records were updated. An average of 105,813 non-WGS/non-CON records were added and/or updated per day. Between releases 199.9 and 200.0, the WGS component of GenBank grew by 34,614,377,046 base pairs and by 5,907,225 sequence records.

The total number of sequence data files increased by 34 with this release. The divisions are as follows:

- BCT: 4 new files, now a total of 118 files
- CON: 11 new files, now a total of 242 files
- GSS: 5 new files, now a total of 284 files
- INV: 2 new files, now a total of 38 files
- PAT: 5 new files, now a total of 204 files
- PLN: 2 new files, now a total of 67 files
- PRI 1 new file, now a total of 47 files
- TSA 3 new files, now a total of 150 files
- VRT: 1 new file, now a total of 32 files.

For downloading purposes, please keep in mind that the GenBank flatfiles are approximately 625 GB (sequence files only). The ASN.1 data are approximately 522 GB.

More information about GenBank Release 200.0 and coming changes are available in the [release notes](#).

## Human genome annotation release 106 now available

*Tuesday, February 11, 2014*

The human (*Homo sapiens*) genome annotation has recently been updated to [annotation release 106](#). The data is now available in the Nucleotide, Protein and Gene databases, is searchable using [BLAST](#), and can be downloaded from the [FTP site](#).

The annotated assemblies include [GRCh38](#) (GCF\_000001405.26), which was published in December 2013 and represents a major, chromosome-coordinate changing update to the reference assembly (for more information, visit the [GRC](#)). Annotation release 106 also includes a re-annotation of the assemblies [CHM1\\_1.1](#) (GCF\_000306695.2) and [HuRef](#) (GCF\_000002125.1).

Some highlights of the GRCh38 annotation results (as compared to [annotation release 105](#) of the previous assembly, GRCh37.p13) include:

1. A total of 29,399 genes are predicted (an increase of 5.6%)
2. A total of 69,826 protein-coding transcripts are annotated (an increase of 3.4%)
3. The number of CDSs annotated as partial decreased from 96 to 56
4. The number of curated RefSeq transcripts with an alignment split across scaffolds decreased from 30 to 5
5. The number of protein coding genes found only on alternate loci and/or novel patches increased from 28 to 64.

There are significant improvements to gene annotation on the new GRCh38 assembly. For example:

- [SRGAP2](#): Previously split across scaffolds with an inversion
- [DPP6](#): Previously split across scaffolds

- [EPPK1](#): Previously internally partial
- [DOC2B](#): Previously 3' partial

See what other annotation runs are in progress on the [Eukaryotic Genome Annotation Pipeline status page](#).

## NCBI releases Entrez Direct, the Entrez utilities on the UNIX command line

*Thursday, February 06, 2014*

NCBI has just released Entrez Direct, a new software suite that enables users to use the UNIX command line to directly access NCBI databases, as well as to parse and format the data to create customized downloads.

For over a decade, researchers have been able to access Entrez search, retrieval and linking functions through the web interface on the [NCBI site](#) or through [Entrez Utilities \(E-Utilities\)](#), the public API for Entrez. Entrez Direct now brings Entrez to the UNIX command line by offering a set of UNIX executables that call the E-utilities directly and provide a variety of post-processing functions. Using these functions, arbitrary fields from complex record formats can be parsed and converted to simple tables suitable for importing into spreadsheets or external databases.

Entrez Direct is available as a simple [FTP download](#) and has extensive [documentation](#) on the NCBI web site. The package itself consists of a master Perl script and several UNIX shell scripts that serve as an interface to the Perl script. Therefore, Entrez Direct will run on UNIX and Macintosh computers that have the Perl language installed, and under the [Cygwin](#) UNIX-emulation environment on Windows PCs.

The [Entrez Direct documentation](#) provides several examples of how these programs may be used, and we have provided three additional examples below. Links are provided to the commands and output files.

### 1. Retrieve a set of PubMed abstracts

### 2. Download a table of start and stop coordinates for all genes on human chromosome 17 from all available assemblies.

First, “`esearch`” retrieves all current genes on human chromosome 17, and these results are passed to “`esummary`”. The resulting XML document summaries are passed to “`xtract`”, which parses the assembly and gene coordinate details and writes them to an output [file](#). The first few lines of this file are shown below.

### 3. Download a table of start and stop coordinates for all genes on human chromosome 17 from the *current* reference assembly.

We will be posting additional Entrez Direct examples and explanations on the [NCBI Insights](#) blog and in the [Entrez Direct documentation](#), and will continue to announce updates on the [utilities-announce](#) list.

For more information:

- [Entrez Direct documentation](#)
- [E-utilities documentation](#)

## Sequence Viewer updated to version 3.1

*Tuesday, February 04, 2014*

```
esearch -db pubmed -query "thyroid peroxidase genetics" | efetch
-format abstract > example1.out
```

This [command](#) searches PubMed with the query “thyroid peroxidase genetics” and then downloads the retrieved records as abstracts, which are then written to a local file.

```
esearch -db gene -query "17[chr] AND human[orgn] AND alive[prop]" |
esummary | xtract -pattern DocumentSummary -element Id -block
LocationHistType -pfx "\n" -element
AnnotationRelease, AssemblyAccVer, ChrAccVer, ChrStart, ChrStop >
example2.out
```

This [command](#) demonstrates the parsing function “xtract” that is unique to Entrez Direct.

```
7157
105  GCF_000001405.25    NC_000017.10    7590867    7571719
105  GCF_000002125.1    AC_000149.1    7484282    7465168
105  GCF_000306695.2    NC_018928.2    7600002    7580865
104  GCF_000001405.22    NC_000017.10    7590867    7571719
...
1636
105  GCF_000001405.25    NC_000017.10    61554421    61575740
105  GCF_000002125.1    AC_000149.1    56922780    56944099
105  GCF_000306695.2    NC_018928.2    61618549    61639868
104  GCF_000001405.22    NC_000017.10    61554421    61575740
...
```

This [file](#) contains data segmented for each gene and begins with a line containing the Gene ID, followed by lines with these columns: annotation release, assembly accession.version, chromosome accession.version, start coordinate, stop coordinate.

```
esearch -db gene -query "17[chr] AND human[orgn] AND alive[prop]" |
esummary | xtract -pattern DocumentSummary -element Id -block
LocationHistType -match "AssemblyAccVer:GCF_000001405.25" -pfx "\n"
-element AnnotationRelease, ChrAccVer, ChrStart, ChrStop > example3.out
```

This [command](#) is an extension of Example 2 in that it uses the “-match” option to limit the output to data from the current reference assembly (GCF\_000001405.25).

[NCBI Sequence Viewer](#) provides a graphical view of sequences and color-coded annotations on regions of sequence stored in the Nucleotide and Protein databases. Sequence Viewer has recently been updated and now has improved compatibility with Internet Explorer for users embedding Sequence Viewer on their web sites and improved performance and response time.

A full list of new features, improvements and fixes is included in the [release notes](#).

```
7157
105  NC_000017.10  7590867  7571719
1636
105  NC_000017.10  61554421  61575740
6532
105  NC_000017.10  28562985  28521336
672
105  NC_000017.10  41277499  41196311
...
```

The output [file](#) has the same format as that for Example 2, except that column 2 (assembly accession.version) is omitted.