

# genomepy: Genes and Genomes made easy

This manuscript ([permalink](#)) was automatically generated from [vanheeringen-lab/genomepy\\_manuscript@787d60a](#) on January 10, 2022.

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# Abstract

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Analyzing functional genomics data, including ATAC-, ChIP- and RNA-sequencing, requires genomic data such as a genome assembly and gene annotations. These resources can generally be retrieved from multiple organizations, at multiple versions, and generated with varying methods. Meanwhile, most bioinformatic workflows and pipelines require the user to supply this genomic data manually, which can be a tedious and error-prone process.

Here we present genomepy, which can search, download, and preprocess the right genomic data for your analysis. Genomepy can search genomic data on GENCODE, Ensembl, UCSC and NCBI, and compare available gene annotations, to allow for an informed decision. The selected genome and gene annotation can be downloaded (from anywhere) and preprocessed with sensible, yet controllable, defaults. Additional supporting data can be automatically generated, such as aligner indexes, genome metadata and blacklists. These functionalities are available on command line interface, aimed at ease of use and integration in automated pipelines, with extended functionality on the Python application programming interface.

## Introduction

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Data analysis is increasingly important in biological research. Whether you are analyzing gene expression in two samples or protein binding motifs in genome atlases, you will need external information such as a reference genome or a gene annotation. For these types of data, there are three major providers: Ensembl [1], UCSC [2] and NCBI [3], and many model-system specific providers, such as GENCODE [4], ZFIN [5], FlyBase [6], WormBase [7], Xenbase [8] and more. Providers have different approaches to compiling genome assemblies and gene annotations, which effect formats, format compliance, naming, data quality, available versions and release cycle. These differences significantly impact compatibility with research [9], tools and (data based on) other genomic data.

You could try to find genomic data yourself, but there are many options with no clear metric for the “best” one. Ensembl, UCSC and NCBI each have FTP archives, web portals, and REST APIs, which you can use to search their individual databases. Alternatively, there are several tools that can be used to access some of these databases programmatically, such as ncbi-genome-download [10] and ucsc-genomes-downloader [11]. However, none of these can search, compare or download from all major genome providers data. Furthermore, downloading and processing genomic data manually can be tedious, error-prone, and poorly reproducible. Although the latter could be remedied by a data management tool, such as iGenomes [12], refGenie [13] or Go Get Data [14], data managers still require the user to supply new data manually.

We have developed genomepy to 1) find genomic data on major providers, 2) compare gene annotations, 3) select the genomic data best suited to your analysis and 4) provide a suite of functions to peruse and manipulate the data. Selected data can be downloaded from anywhere, and is processed automatically. To ensure reproducibility, data sources and processing steps are documented, and can be enhanced further by using a data manager. Genomic data can be loaded into genomepy, which utilizes and extends on packages including pyfaidx [15], pandas [16] and MyGene.info [17] to rapidly work with gene and genome sequences and metadata. Similarly, genomepy has been incorporated into other packages, such as pybedtools [18] and CellOracle [19]. Genomepy can be used on command line, and via its fully documented Python API, for a one-time analysis or integration in pipelines and workflow managers such as Nextflow [20], Galaxy [21] or Snakemake [22].

# Features of genomepy

The core functionalities of genomepy are to search, download and process genomes and gene annotations.

On first use, the `search` function queries the databases of GENCODE, Ensembl, UCSC and NCBI and caches the metadata on available assemblies (for up to 7 days). The input can be text, taxonomy identifiers or assembly accession identifiers. The input type is automatically recognized and used to find assemblies that 1) have the text in the genome names or various description fields, 2) (exactly) matches the taxonomy identifier or 3) (mostly) matches the assembly accession. The output of the function is a table with rows of metadata for each assembly found. This overview indicates whether a gene annotation can be downloaded for an assembly (or which of the four UCSC annotations) (see fig. 1a). The gene annotation(s) of an assembly can be inspected with the `annotation` function (fig. 1b).

```
$ genomepy search GRCh38
```

name	provider	accession	tax_id	annotation	species
other_info				n r e k	<- UCSC
options (see help)					
GRCh38	GENCODE	GCA_000001405.15	9606	✓	Homo sapiens
GENCODE annotation + UCSC genome					
GRCh38.p13	Ensembl	GCA_000001405.28	9606	✓	Homo sapiens
2014-01-Ensembl/2021-08					
hg38	UCSC	GCA_000001405.15	9606	✓ ✓ ✗ ✓	Homo sapiens
Dec. 2013 (GRCh38/hg38)					
GRCh38	NCBI	GCF_000001405.26	9606	✓	Homo sapiens
Genome Reference Consortium					

^  
Use name for genomepy install

```
$ genomepy annotation GRCh38.p13
```

```
12:00:00 | INFO | Ensembl
1      ensembl_havana  gene      1211340 1214153 .      -      .
      gene_id "ENSG00000186827"; gene_version "11"; gene_name "TNFRSF4";
      gene_source "ensembl_havana"; gene_biotype "protein_coding";
12:00:00 | INFO | NCBI
NC_000001.11      genomepy      transcript      11874 14409 .      +
      .      gene_id "DDX11L1"; transcript_id "NR_046018.2"; gene_name
      "DDX11L1";
```

```
$ genomepy install --annotation GRCh38.p13
$ ls -l ~/.local/share/genomes/GRCh38.p13
GRCh38.p13.annotation.bed
GRCh38.p13.annotation.gtf
GRCh38.p13.fa
GRCh38.p13.fa.fai
GRCh38.p13.fa.sizes
GRCh38.p13.gaps.bed
README.txt
assembly_report.txt
index/
```

An assembly name can be passed to the `install` function (fig. 1c). The genome FASTA file is downloaded with the desired sequence masking level [23,24] (softmasked by default). Reference assemblies often contain alternate sequences to reflect biological diversity. During sequence alignment however, similar reference sequences result in multiple alignment, leading to loss of data (as discussed in [25]). Therefore, genomepy downloads the primary assembly and filters out alternative sequences by default. Additional regex filters may be passed to either include or exclude contigs (chromosomes, scaffolds, etc.) by name. Once filtering is performed, genomepy generates a genome index using pyfaidx [15], and contig sizes and contig gap sizes are collected in separate files.

Gene annotations come in a variety of recognized formats (GFF3, GTF, BED12). The `install` function can download the most descriptive format to output the commonly used GTF and BED12 formats. Contig names of the genome and gene annotation sometimes mismatch, which makes them incompatible with tools such as splice-aware aligners. Therefore, genomepy will attempt to match the contig names of the gene annotations to those used in the genome FASTA.

The install function can be extended with postprocessing steps via plugins. The options can be inspected and toggled with the `plugin` function. Briefly, the blacklist plugin downloads blacklists by the Kundaje lab [26] for the supported genomes. Other plugins support the generation of aligner indexes, including DNA aligner indexes for Bowtie2 [27], BWA [28], GMAP [29] or Minimap2 [30], and splice-aware aligners such as STAR [31] and HISAT2 [32].

Assemblies not present on the major providers can be processed similarly by supplying the URLs or file paths to the `install` function. For data provenance and reproducibility, a README file is generated during the installation process with time, source files, processing steps, and filtered contigs.

The core features are available on both the command line interface and Python API. Additional features are available on the Python API, focussed around two classes. The `Genome` class can be used to extract exact or random sequences from the FASTA, filter the FASTA and list the contigs, contig sizes and contig gaps. The `Annotation` class can be used to browse and filter the BED12 or GTF files as pandas dataframes [16], map gene identifiers to other types using mygene.info [17], map chromosome names to naming schemes of other major providers, and create a dictionary of any two GTF columns or attribute fields (to easily convert gene identifiers to gene names for instance).

## Conclusion

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Obtaining suitable genomic data is a principal step in any genomics project. With genomepy, finding available assemblies becomes trivial. A genome, with the desired sequence masking, level of biological diversity, and contigs can be obtained with a single genomepy command. Gene annotations in GTF and BED12 format are commonly used, sometimes in combination with a genome, in analyses. The install command can download and prepare these annotations, and match contigs to the genome, with further options available in the Python API. Whatever install options you choose are logged, for reproducibly, allowing you to start your analysis with confidence.

## Code availability

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Genomepy can be installed using [Bioconda](#), [Pip](#) and [Docker](#). Code and documentation are available on [github](#) and [github-pages](#), respectively.

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