MakeHub User Guide

Author and Contact Information

Katharina J. Hoff, University of Greifswald, Institute for Mathematics and Computer Science, Bioinformatics Group (katharina.hoff@uni-greifswald.de)

Contents

- What is MakeHub?
- Installation
 - Quick start
 - Dependencies
 - MakeHub
- Data preparation
- Running MakeHub
 - Creating a new hub
 - Adding tracks to existing hub
 - Options explained
- Example data
- Output of MakeHub
- How to use MakeHub output with UCSC Genome Browser
- Bug reporting
- Citing MakeHub
- License

What is MakeHub?

MakeHub is a command line tool for the fully automatic generation of of track data hubs ¹ for visualizing genomes with the UCSC genome browser ². Track data hubs are data structures that contain all required information about a genome for visualizing with the UCSC genome browser.

Assembly hubs need to be hosted on a publicly available webspace (that might be user/password protected) for usage with the UCSC genome browser.

MakeHub is implemented in Python3 and automatically executes tools provided by UCSC for generation of assembly hubs (http://hgdownload.soe.ucsc.edu/admin/exe (http://hgdownload.soe.ucsc.edu/admin/exe)) on Linux and MacOS X x86_64 computers. For visualization of RNA-Seq alignment data from BAM files, MakeHub uses Samtools 3]. If installed, the AUGUSTUS 4 tool bam2wig is used to speed up BAM to wig format conversion (https://github.com/Gaius-Augustus/Augustus/Augustus/Augustus)), which is otherwise performed without bam2wig.

MakeHub can either be used to create entirely new assembly hubs, or it can be used to add tracks to hubs that

were previously created by MakeHub.

For display by the UCSC Genome Browser, assembly hubs need to be hosted on a publicly accessible web server.

Installation

Quick Start

MakeHub is a Python3 script for Linux or MacOS X with x86-64 architecture. It requires Python3, Biopython, gzip, sort and - in the case that BAM files are provided - samtools, and optionally the AUGUSTUS tool bam2 hints

Many users who create the input data for MakeHub, e.g. with BRAKER ⁵, have the required dependencies already installed on their system and my thus skip ahead to section Running MakeHub. In case of doubt, read the following sections about installation of Dependencies and MakeHub installation.

Dependencies

In the following, we give instructions on where dependencies can be obtained, and how they may be installed on Ubuntu Linux.

Python3 is available from https://www.python.org/downloads/ (https://www.python.org/downloads/), or as package for many Unix systems.

For example, on Ubuntu, install Python3 with:

sudo apt install python3

We recommend to use pip for installing further python modules. pip is available at https://pypi.org/project/pip/ (https://pypi.org/project/pip/). It is also available as package for many Unix systems.

For example, on ubuntu, install pip with:

sudo apt install python3-pip

Further, MakeHub uses Biopython (e.g. for parsing a genome file in order to determine which parts of the genome have been masked for repeats). Install biopython with pip as follows:

pip3 install biopython

- bedToBigBed
- genePredCheck
- faToTwoBit
- gtfToGenePred
- hgGcPercent
- ixIxx

- twoBitInfo
- wigToBigWig
- genePredToBed

You may download these binaries and make them available in your \$PATH. However, if you skip installing these tools, they will be downloaded during MakeHub execution, automatically.

MakeHub uses Samtools for BAM file sorting and conversion. Samtools is avilable at https://github.com/samtools/ (https://github.com/samtools/) . It is also avilable as package with many linux distributions.

For example, on ubuntu, install samtools with:

sudo apt install samtools

MakeHub has been tested with Samtools 1.8-20-g4ff8062. It might not be fully downward compatible with older versions.

MakeHub uses gzip for compressing wig files that were created from BAM files. gzip is available at https://ftp.gnu.org/gnu/gzip/ (https://ftp.gnu.org/gnu/gzip/). It often installed by default on Unix systems. If not, it is usually available as a package.

If missing, on Ubuntu, install with:

p
 sudo apt install gzip

MakeHub uses Unix sort. sort should be installed by default on all Unix systems.

MakeHub can use the AUGUSTUS tool bam2wig, if that tool is available in the \$PATH. bam2wig is available as part of AUGUSTUS at https://github.com/Gaius-Augustus/Augustus (https://github.com/Gaius-Augustus/Augustus). Please follow the compilation instructions in Augustus/auxprogs/bam2wig/README.txt in case the default make command fails.

MakeHub

MakeHub is a python3 script named make_hub.py. It does not require a particular installation procedure after download.

It can be executed either with

python3 make_hub.py

If you add make_hub.py to your \$PATH (i.e. by adding the location of make_hub.py at the bottom of your ~/.bashrc file similar to PATH=/path/to/MakeHub: \$PATH, followed by loading the ~/.bashrc file in case you did not re-open a new bash session with source ~/.bashrc)and make it executable (i.e. with chmod u+x make_hub.py), it can be executed with

make_hub.py

from any location on your computer.

Data Preparation

MakeHub accepts files in the following formats:

- genome file in FASTA format (simple FASTA headers without whitespaces or special characters); if the file is softmasked, a track with repeat information will automatically be generated. Note that the FASTA headers must be consistent with BAM-, hints- and gene prediction files.
- BAM file(s) with RNA-Seq to genome alignments
- gene prediction file(s) in GTF-format, e.g. from BRAKER
- AUGUSTUS hints files in BRAKER-specific GFF hints format
- Gene prediction files in GFF3-format from MAKER ⁶ and Gemoma ⁷

Running MakeHub

MakeHub can be used either to create new assembly hubs, or to add tracks to assembly hubs that had previously been created.

Creating a new hub

The essential arguments for creating a new assembly hub are:

- FMAIL, --email EMAIL Contact e-mail adress for assembly hub. This e-mail adress will be displayed on all HTML pages that describe this hub and its tracks. Providing an e-mail adress is a requirement for UCSC assembly hubs, e.g. described at http://genomewiki.ucsc.edu/index.php/Assembly_Hubs (http://genomewiki.ucsc.edu/index.php/Assembly_Hubs) and http://genomewiki.ucsc.edu/index.php/Public_Hub_Guidelines#Track_description_page_recommendations (http://genomewiki.ucsc.edu/index.php/Public_Hub_Guidelines#Track_description_page_recommendations).
- -g GENOME, --genome GENOME Genome file in FASTA format. If the file contains softmasked repeats, a repeat masking track with softmasking information will automatically be generated.
- -1 SHORT_LABEL, --short_label SHORT_LABEL Short label (without whitespaces and special characters) for identifying assembly hub, will also be used as directory name for hub, e.g. --short_label fly

At the point in time of assembly hub creation, we strongly recommend the additional usage of

• -L LONG_LABEL, --long_label LONG_LABEL Long label for hub, e.g. english organism name, if it contains whitespaces, pass it with quotation marks: ---long_label "fruit fly"

You may at the point of time of creating a hub already supply information about all gene prediction and evidence tracks that you would like to see in your final hub. Please have a look at the section Options Explained for information about possible tracks. The section also describes how to add latin species name and assembly version.

Usage example 1:

```
make_hub.py -l hmi1 -L "Rodent tapeworm" -g data/genome.fa -e \
katharina.hoff@uni-greifswald.de
```

The resulting hub is trivial, as it only displays very basic information about the genome, such as the GC-content, restriction enzyme sites and repeat masking segments.

If you want to visualize the result, connect the following hub with the UCSC genome browser (see section How to use MakeHub output with UCSC Genome Browser): http://augustus.uni-greifswald.de/bioinf/makehub/examples/hmi1/hub.txt (http://augustus.uni-

greifswald.de/bioinf/makehub/examples/hmi1/hub.txt)

Usage example 2:

```
make_hub.py -1 hmi2 -L "Rodent tapeworm" -g data/genome.fa -e \
katharina.hoff@uni-greifswald.de -a data/annot.gtf -b data/rnaseq.bam \
-d
```

In comparison to the first example, the resulting hub has a track with reference annotation genes, and a track with coverage information from RNA-Seq data, and it displays the native BAM-file (-d).

If you want to visualize the result, connect the following hub with the UCSC genome browser (see section How to use MakeHub output with UCSC Genome Browser): http://augustus.uni-greifswald.de/bioinf/makehub/examples/hmi2/hub.txt (http://augustus.uni-greifswald.de/bioinf/makehub/examples/hmi2/hub.txt)

Usage example 4:

```
make_hub.py -1 hmi4 -L "Rodent tapeworm" -g data/genome.fa -e \
    katharina.hoff@uni-greifswald.de -a data/annot.gtf -b data/rnaseq.bam \
    -d -X data -M data/maker.gff -E data/gemoma.gff \
    -N "Hymenolepsis microstoma" -V GCA_000469805.2
```

In comparison to the first two examples, the resulting hub has a large number of evidence and gene prediction tracks from BRAKER, MAKER and Gemoma.

If you want to visualize the result, connect the following hub with the UCSC genome browser (see section How to use MakeHub output with UCSC Genome Browser): http://augustus.uni-greifswald.de/bioinf/makehub/examples/hmi4/hub.txt (http://augustus.uni-greifswald.de/bioinf/makehub/examples/hmi4/hub.txt)

Adding tracks to existing hub

If a hub already exists, you may add tracks to this existing hub using the option -A, --add_track. The minimal required arguments

- besides giving the approriate information that you would like to add are:
- -A, --add_track
- -e EMAIL, --email EMAIL Contact e-mail address for assembly hub.
- -1 SHORT_LABEL, --short_label SHORT_LABEL Short label (without whitespaces and special characters) for identifying assembly hub.
- -A, --add_track Add track(s) to existing hub

Usage example 3:

First, we create a novel track hub hmi3 that is identical to Usage example 2:

```
make_hub.py -1 hmi3 -L "Rodent tapeworm" -g data/genome.fa -e \
    katharina.hoff@uni-greifswald.de -a data/annot.gtf -b data/rnaseq.bam \
    -d
```

Subsequently, we add a number of tracks:

The resulting hub has many gene prediction tracks from the BRAKER output directory data, and from the MAKER output file data/maker.gff.

Let's add one more track (only for the sake of demonstration, this track could have been included in the previous example, or course, or at the point of time of track generation):

If you want to visualize the result, connect the following hub with the UCSC genome browser (see section How to use MakeHub output with UCSC Genome Browser): http://augustus.uni-greifswald.de/bioinf/makehub/examples/hmi3/hub.txt (http://augustus.uni-greifswald.de/bioinf/makehub/examples/hmi3/hub.txt)

Options explained

In the following, we explain all options of make_hub.py

- -h, --help Print help message and exit.
- -p, --printUsageExamples Print usage examples for make_hub.py to command line (for demonstration).
- -e EMAIL, --email EMAIL Contact e-mail address for assembly hub. This is a requirement for all publicly listed assembly hubs. It is obligatory for make_hub.py.
- -g GENOME, --genome GENOME Genome file in FASTA format. If the file is softmasked for repeats, a repeat masking track will automatically be generated, unless the option:
- -n, --no_repeats Disable repeat track generation from softmasked genome sequence is activated (this may save runtime, particularly for large genomes).
- -L LONG_LABEL, --long_label LONG_LABEL Long label for hub, e.g. english organism name, if it contains whitespaces, pass it with quotation marks: ---long_label "fruit fly"
- -1 SHORT_LABEL, --short_label SHORT_LABEL Short label (without whitespaces and special characters)
 for identifying assembly hub. The short label will also be used as assembly version, unless the following
 option is specified:
- -V ASSEMBLY_VERSION, --assembly_version ASSEMBLY_VERSION Assembly version, e.g. "BDGP R4/dm3". This argument must be provided if the hub is supposed to be added to the public UCSC list.
- -N LATIN_NAME, --latin_name LATIN_NAME Latin species name, e.g. "Drosophila melanogaster". This argument must be provided if the hub is supposed to be added to the public UCSC list.
- -s SAMTOOLS_PATH, --SAMTOOLS_PATH SAMTOOLS_PATH Path to samtools executable. By default, make_hub.py will search for a samtools executable in your \$PATH. On some systems, e.g. high performance compute clusters, it may be more conventient to specify the path to samtools with this option while calling make_hub.py
- -B BAM2WIG_PATH, --BAM2WIG_PATH BAM2WIG_PATH Path to bam2wig executable. bam2wig from AUGUSTUS auxprogs is not required for converting a BAM to a WIG file with make_hub.py. It may be a little faster than the built-in conversion function, though. By default, make_hub.py will search for a bam2wig executable in your \$PATH. On some systems, e.g. high performance compute clusters, it may be more conventient to specify the path to bam2wig with this option while calling make_hub.py
- -b BAM [BAM ...], --bam BAM [BAM ...] BAM file(s) space separated with RNA-Seq information, will be displayed as BigWig coverage track.
- -d, --display_bam_as_bam Display BAM file(s) as bam tracks (in addition to BigWig coverage tracks)
- -c CORES, --cores CORES Number of cores for samtools sort processes that are used for producing BAM

tracks. Usage of more than one core may significantly speed up track generation.

- -a ANNOT, --annot ANNOT GTF file with reference annotation (may be particularly interesting to visualize in case of re-annotation of genomes).
- -X BRAKER_OUT_DIR, --braker_out_dir BRAKER_OUT_DIR BRAKER output directory with GTF files. If this option is specified, the following options are set, automatically, using the files in BRAKER_OUT_DIR (if these files exist):
 - -i HINTS, --hints HINTS
 - -t TRAINGENES, --traingenes TRAINGENES
 - -m GENEMARK, --genemark GENEMARK
 - -w AUG_AB_INITIO, --aug_ab_initio AUG_AB_INITIO
 - -x AUG_HINTS, --aug_hints AUG_HINTS
 - -y AUG_AB_INITIO_UTR, --aug_ab_initio_utr AUG_AB_INITIO_UTR
 - -z AUG_HINTS_UTR, --aug_hints_utr AUG_HINTS_UTR
- -i HINTS, --hints HINTS GFF file with BRAKER hints (AUGUSTUS-specific GFF format of BRAKER).
- -t TRAINGENES, --traingenes TRAINGENES GTF file with training genes.
- -m GENEMARK, --genemark GENEMARK GTF file with GeneMark predictions.
- -w AUG_AB_INITIO, --aug_ab_initio AUG_AB_INITIO GTF file with ab initio AUGUSTUS predictions
- -x AUG_HINTS, --aug_hints AUG_HINTS GTF file with AUGUSTUS predictions with hints
- -y AUG_AB_INITIO_UTR, --aug_ab_initio_utr AUG_AB_INITIO_UTR GTF file with ab initio AUGUSTUS predictions with UTRs
- -z AUG_HINTS_UTR, --aug_hints_utr AUG_HINTS_UTR GTF file with AUGUSTUS predictions with hints with UTRs
- -M MAKER_GFF, --maker_gff MAKER_GFF MAKER2 output file in GFF3 format. This file could be the result of a gff3_merge -d *_master_datastore_index.log command.
- -E GEMOMA_FILTERED_PREDICTIONS, --gemoma_filtered_predictions GEMOMA_FILTERED_PREDICTIONS GFF3 output file of Gemoma (filtered_predictions.gff)
- -G GENE_TRACK [GENE_TRACK ...], --gene_track GENE_TRACK [GENE_TRACK ...] Gene track with user specified label, argument must be formatted as follows for adding a single track: --gene_track file.gtf tracklabel
- -A, --add_track Add track(s) to existing hub
- -o OUTDIR, --outdir OUTDIR Output directory to write hub to (default is the current working directory). This directory must be writable.
- -r, --no_tmp_rm Do not delete temporary files (e.g. for debugging purposes).
- -v VERBOSITY, --verbosity VERBOSITY If INT VERBOSITY > 0, verbose logging output is produced (e.g. for debugging purposes).

Example data

Example data is located in the directory data/. It consists of the following files:

- genome.fa: sequence LN902858_1 of *Hymenolepis microstoma*, assembly version GCA_000469805.2 from GenBank.
- rnaseq. fa: RNA-Seq reads of library ERR337976 that mapped to sequence LN902858_1 with Hisat2.
- annot.gtf: NCBI reference annotation of scaffold LN902858_1.
- augustus.ab_initio.gtf: AUGUSTUS ab inito gene predictions from a BRAKER run (run was performed on the complete genome, predictions corresponding to LN902858_1 were extracted) with Hisat2 alignments from RNA-Seq library ERR337976.
- augustus.hints.gtf: AUGUSTUS gene predictions with hints from a BRAKER run (run was performed on the complete genome, predictions corresponding to LN902858_1 were extracted) with Hisat2 alignments from RNA-Seq library ERR337976.
- GeneMark-ET/genemark.gtf: GeneMark-ES/ET predictions from a BRAKER run (run was performed on the complete genome, predictions corresponding to LN902858_1 were extracted) with Hisat2 alignments from RNA-Seq library ERR337976.
- hintsfile.gff: Hints from a BRAKER run (run was performed on the complete genome, hints corresponding to LN902858_1 were extracted) with Hisat2 alignments from RNA-Seq library ERR337976.
- gemoma.gff: Gemoma predictions from a Gemoma run with Hisat2 alignments from RNA-Seq library ERR337976 and proteins of *Echinococcus multilocularis*. (Run was performed on the complete genome, predictions corresponding to LN902858_1 were extracted)
- maker.gff: MAKER2 predictions from a run with BRAKER gene models as model_gff, Cufflinks assembly
 of Hisat2 alignments of RNA-Seq library ERR337976, a custom repeat library for RepeatMasker,
 AUGUSTUS with BRAKER-trained parameters, BUSCO predictions as evidence, and GeneMark-ES/ET
 predictions with BRAKER-trained parameters.

Output of MakeHub

make_hub.py produces a directory that is called identical to the argument for option --short_label/-1. Let's assume the short label had been species.

species contains the following files:

- hub.txt this file contains basic information about the assembly hub, for example, the short and long labels, a reference to genomes.txt, and contact information.
- genomes.txt this file contains references to the configuration files trackDb.txt and groups.txt, as well as for example a default browsing location.
- aboutHub.html this file should contain a meaningful description of your assembly hub. Please edit this file, manually.

Furthermore, species contains another directory species in which the hub configuration files trackDb.txt and groups.txt, as well as all files that are required for browsing tracks, reside. The number of files may differ depending on how many tracks have actually been created.

Importantly, species also contains *.html files for all tracks. These files should be edited, manually, to contain meaningful information!

How to use MakeHub output with UCSC Genome

Browser

Copy the complete hub folder (e.g. species) to a publicly accessible web server.

Go to https://genome.ucsc.edu/index.html (https://genome.ucsc.edu/index.html), click on My Data -> Track Hubs -> My Hubs and add the link to your publicly available hub.txt file into the URL window. Subsequently, click on Add Hub.

Bug reporting

Before reporting bugs, please check that you are using the most recent versions of MakeHub. Also, check the open and closed issues on github at https://github.com/Gaius-Augustus/MakeHub/issues (https://github.com/Gaius-Augustus/MakeHub/issues) for possible solutions to your problem.

Reporting bugs on github

If you found a bug, please open an issue at https://github.com/Gaius-Augustus/MakeHub/issues (https://github.com/Gaius-Augustus/MakeHub/issues) (or contact katharina.hoff@uni-greifswald.de).

Information worth mentioning in your bug report:

make_hub.py prints information about separate steps on STDOUT. Please let us know at which step and with what error message make_hub.py caused problems.

Citing MakeHub

Hoff KJ, "MakeHub: Fully automated generation of UCSC Genome Browser Assembly Hubs." *bioRxiv*: doi: https://doi.org/10.1101/550145 (https://doi.org/10.1101/550145)

License

All source code is under GNU public license 3.0 (see https://www.gnu.org/licenses/gpl-3.o.de.html (https://www.gnu.org/licenses/gpl-3.o.de.html)).

References

[1] Raney BJ, Dreszer TR, Barber GP, Clawson H, Fujita PA, Wang T, Nguyen N, Paten B, Zweig AS, Karolchik D, Kent WJ. 2014. "Track Data Hubs." *Bioinformatics* 1;30(7):1003−5. ↔

[2] Kent WJ, Sugnet CW, Furey TS, Roskin KM, Pringle TH, Zahler AM, Haussler D. 2002. "UCSC Genome Browser." *Genome Res.* 12(6):996−1006. ↔

[3] Li H, Handsaker B, Wysoker A, Fennell T, Ruan J, Homer N, Marth G, Abecasis G, Durbin R. 2009. "The sequence alignment/map format and SAMtools." *Bioinformatics* 26(16):2078–2079. ↔

[4] Stanke M, Diekhans M, Baertsch R, Haussler D. 2008. "Using native and syntenically mapped cDNA

alignments to improve de novo gene finding." Bioinformatics 24(5):637-644.€

- [5] Hoff KJ, Lange S, Lomsadze A, Borodovsky M, Stanke M. 2015. "BRAKER1: unsupervised RNA-Seq-based genome annotation with GeneMark-ET and AUGUSTUS." *Bioinformatics* 32(5), 767-769. ↔
- [6] Holt C, Yandell M. 2011. "MAKER2: an annotation pipeline and genome-database management tool for second-generation genome projects." *BMC Bioinformatics* 12(1), 491. ↔
- [7] Keilwagen J, Hartung F, Paulini M, Twardziok SO, Grau J. 2018. "Combining RNA-seq data and homology-based gene prediction for plants, animals and fungi." *BMC Bioinformatics* 19(1), 189. ←