

View a bacterial genome in IGB

Introduction

This tutorial describes how to import sequence and annotations into IGB when your genome of interest is not available from an IGB QuickLoad or Distributed Annotation site.

In this tutorial, we will demonstrate importing a bacterial genome (E. coli) using files downloaded from NCBI. Overall, to view custom genome and annotations in IGB properly a Synonym File needs to be created.

Get genome data from NCBI

First visit NCBI to retrieve the sequence data and annotations.

- Go to the NCBI GenBank record for E. coli K-12 substr. MG1655: http://www.ncbi.nlm.nih.gov/nucore/NC_000913.3

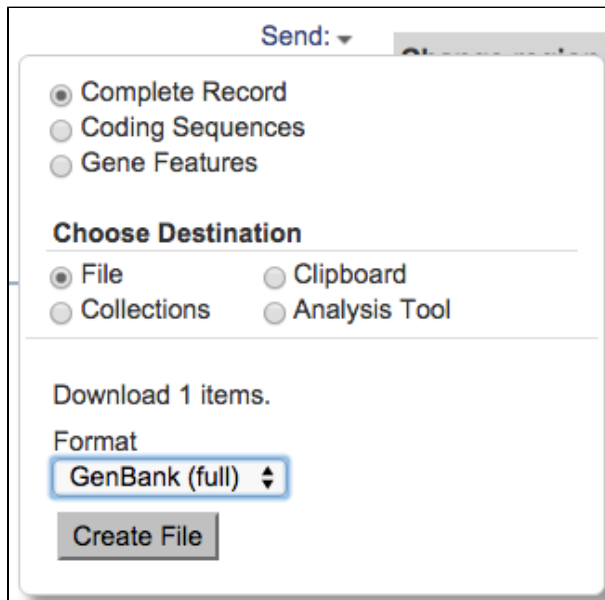
Download annotations file

- Click **Send** menu (on the right side of the record)
- Select **Complete Record, File, GenBank(Full)**, and **Create File**

Save the file to your computer; by default, it will be named "sequence.gb".



If you change the name, be sure to use the ".gb" file extension so that IGB can recognize the file and the file format.



Download sequence file

- Click on the **FASTA** link (on the left side of the record)
- Click **Send** menu
- Select, **Complete Record, File, FASTA**, and **Create File**

Save the file to your computer; by default, it will be named "sequence.fasta".

Make sure that the extensions for the FASTA file is either '.fasta' or '.fa'. If the extension is '.txt', you can safely change it to '.fa'.

NCBI Resources How To My NCBI Sign In

Nucleotide Nucleotide Search

Display Settings: FASTA Send: Show options shown

Escherichia coli str. K-12 substr. MG1655 chromosome, complete genome

NCBI Reference Sequence: NC_000913.2

GenBank Graphics

>gi|49175990|ref|NC_000913.2| Escherichia coli str. K-12 substr. MG1655 chromosome, complete genome

AGCTTTTCATTTCTGACGCAACGGGCAATATGCTCTGTGTGGATTAAAAAAGAGTGTCTGATAGCAGC
TTCTGAACCTGTTACCTGCGCTGAGTAAATTTAAATTTTATTGACTTAGTGCACATAAATACTTTAACCAA
TATAGGCATAGCGCAGACAGAGATAAAATTTACAGAGTACACAACATCCATGAAACGCATTAGCACCACC
ATTACCACCACCATCACCATTACCACAGGTAACGGTGGGGCTGACGGGTACAGGAAACACAGAAAAAG
CCCGCACCTGACAGTGGGGCTTTTTTTTCGACCAAGGTAACGAGGTAACAACCATGCGAGTGTGAA
GTTCCGGGTACATCAGTGGCAAAATGCAGAACGTTTTTCGCGTGTTCGGATATTCTGGAAGCAATGCC
AGGCAGGGGAGGTGGCCACCGTCTCTGCCCCGCCAAAATCACCACCACTGGTGGCGATGATTG
AAAAAACCATTAGCGGCCAGGATGCTTTACCAATATCAGCGATGCCGAACGTATTTTGGCGAACTTTT
GACGGGACTCGCCGCCCGCCAGCGGGTTCCTGCGGCAATTGAAAACCTTCGTCGATCAGGAATTT
GCCCAATAAACATGTCCTGCATGGCATTAGTTTGTGGGGCAGTGCCGGATAGCATCAACGCTGCGC
TGATTTCGCGTGGCGAGAAAATGTCGATGCCATTATGGCGGCGTATTAGAAGCGCGCGGTCAACAGT
TACTGTTATCGATCCGTCGAAAACTGCTGGCAGTGGGCAATTACCTCGAATCTACCGTCGATATTGCT

Choose Destination

Complete Record
Coding Sequences
Gene Features

File Clipboard
Collections Analysis Tool

Download 1 items.

Format
FASTA

Create File

REBASE enzyme EcoKMcra
[REBASE - The Restriction Enzy...]
REBASE enzyme EcoKMcra
[REBASE - The Restriction Enzy...]
REBASE enzyme M.EcoKII

Create Synonyms File

When IGB shows the sequence.gb file, it uses the 'LOCUS' name from the sequence.gb. To show the sequence.fasta file, IGB uses the 'sequence name' that follows the >. Note, when the labels for each file type (FASTA sequence files, annotation files, .bam/alignment files, etc.) for the exact same chromosome/sequence are all different IGB will treat each one as a separate chromosome; they will not be visualized together.

To overcome this, we enter 'synonyms'; IGB already has some internal synonyms, for example "1" and "chr1" are equivalent. You will need to know the sequence names of each of these files; if you are not sure, a quick way is to drag all of the files into IGB at the same time.

Determine 'sequence names' in sequence and annotation files:

1. Drag into IGB the sequence.fasta and sequence.gb
2. In **Current Genome Panel** look under **Sequences**
 - a. Note the sequences.fasta is called **gi|556503834|ref|NC_000913.3|** and the sequence.gb is labelled **NC_000913**

NC_000913 (Unknown Genome 1) - Integrated Genome Browser 6.7.1

NC_000913 : 0 - 4,639,675

NC_000913
4639675 bp
DNA
circular BCT
11-JAN-2012 (+/-)

sequence.fasta (+/-)

Coordinates

0 1,000,000 2,000,000 3,000,000 4,000,000

2,319,837

Data Access Selection Info Search Sliced View Graph Adjuster External View Plug-ins

Data Sources and Data Sets - Configure

Local Files (Local Files)

sequence.fasta
sequence(1).gb

Data Management Table

	FG	BG	2 Track	Load Mode	Data Set/File...	Track Name (Dou...
				Region	sequence(1).gb	NC_000913...
				Region	sequence.fasta	sequence.fasta

Load All Sequence Load Sequence In View Load Data

Species

Unknown Species 1

Genome Version

Unknown Genome 1

(0) Sequence(s) Length

Sequence(s)	Length
gi 49175990 ref NC_000913.2	4,639,675
NC_000913	4,639,675

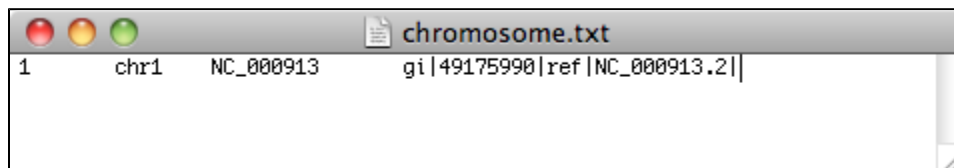
genome

94M of 2039M

Create synonym file

Now that we know the headers of the files we will create a tab-delimited 'personal synonym' file:

1. Open a Text Editor
2. On the first line, type **1** then press tab. Type **chr1** then press tab
 - a. 1 and chr1 are standard names for the first chromosome, and many files use these as their headers. Typically, we always include these two options in a synonym file
3. Type in the header of the sequence.gb file, **NC_000913**, then press tab
4. Type in the header of the sequence.fasta file, **gi|556503834|ref|NC_000913.3|**
5. **Save** this file with the name **chromosome.txt**

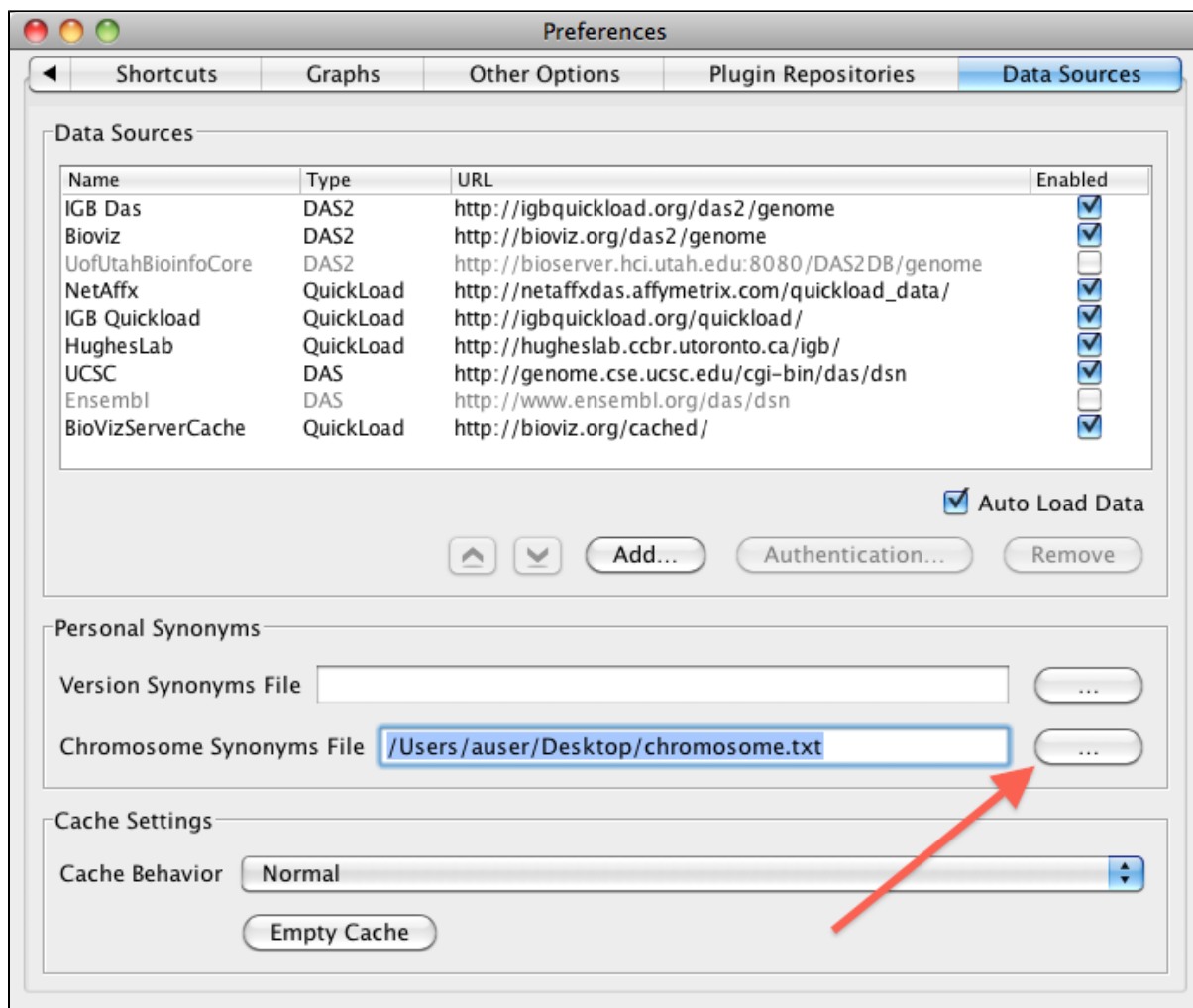


If you are making a synonym file for a multi-chromosomal organism, then make a new line in the file for each chromosome, and just add all of the 'names' associated with it (make sure that there is a 'tab' between each name!). If you include a file that has a new name, open the chromosome.txt, and add the name to the proper line

Import Synonym File into IGB

In IGB:

1. Select **Configure** (under Data Access panel) to open the Preferences Menu
2. Select **Data Sources** Tab
3. Next to **Chromosome Synonyms File** browse for the chromosome.txt file
4. Close the Preferences Menu
5. Restart IGB



Visualizing the Reference Sequence and Models

When you open the new instance of IGB, your synonyms will be loaded for you. At this point, we will open the files, sequences.fasta and sequences.gb, so you can begin analysis of your own data.

To open the sequence and annotation files:

1. **File > Open Reference Sequence...**
2. Browse for the 'sequence.fasta' file.
3. **File > Open File...**
4. Browse for the 'sequence.gb' file.
5. Click the **Load Data** button. **NOTE:** you need to click the **Load Data** button and NOT the **Load Sequence** button to visualize your sequence file!

Both the sequence and the models will load. You will be zoomed out, so the sequence will appear as a grey bar; as you zoom in the colors and nucleotides will become visible. At this point, you can begin viewing the gene models and sequence. Keep in mind that if the name of the chromosome(s) is different in **your** files, you will have to add the name to the chromosome.txt and then reopen IGB so it can load in the new information.