
BioMercator

Release 5.0

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QUICK START

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INTRODUCTION

BioMercator V4 is a genetic map compilation and QTL meta-analysis software providing tools to integrate these data with genome structural and functional annotation. It offers wizards to run analyses and display input and output maps through a user friendly graphical user interface (GUI). In addition to map compilation and meta-analysis methods available in previous BioMercator version, this new release includes algorithms from the MetaQTL package (<http://www.bioinformatics.org/mqtl/wiki/>).

The GUI has evolved, and now offers new ways of zooming. A cascading zoom allows to progressively enlarge a region of a map while conserving alongside the whole chromosome representation. It is now possible to use the zoom from the whole map representation to focus on zones of interest of several chromosomes. Image can be exported in various format including SVG for subsequent image editing in third-party software.

BioMercator V4 also implements a new file loading wizard capable of reading an unlimited number of files and automatically recognize their format.

The possibility of using MetaQTL's package algorithms directly inside BioMercator V4 came with a file format evolution ; to use fully those algorithms, more data will be required than in the previous BioMercator version.

The user will be able to export maps and analyses in a format compliant for submission to the URGI databases (GnpIS).

1.1 How to install BioMercator

BioMercator is a Java program; all you need is Java (v1.6 or above) installed on your machine. To install java see <http://www.java.com/>.

On Windows: Double-click on the BioMercatorV4.jar to launch the program

On other OS: Open a terminal and execute the command line :

```
java -jar BioMercatorV4.jar
```

Memory management : On any OS, If the software doesn't launch or exits with a memory error (when working with large genome annotation dataset), you should open a console (terminal) and type the following line :

```
java -jar -Xmx512m BioMercatorV4.jar
```

1.2 Tutorial

This tutorial will guide you through new functionalities and analyses in BioMercatorV4; data files used are situated in the “Tutorial_files” folder.

When launching the software for the first time, the workspace is totally empty.

1.2.1 Data Loading

First, we'll load the genetic maps. The wizard is accessible through the “File” menu and “Genetic data loading” sub menu. Choose “New project” and name it “data”.

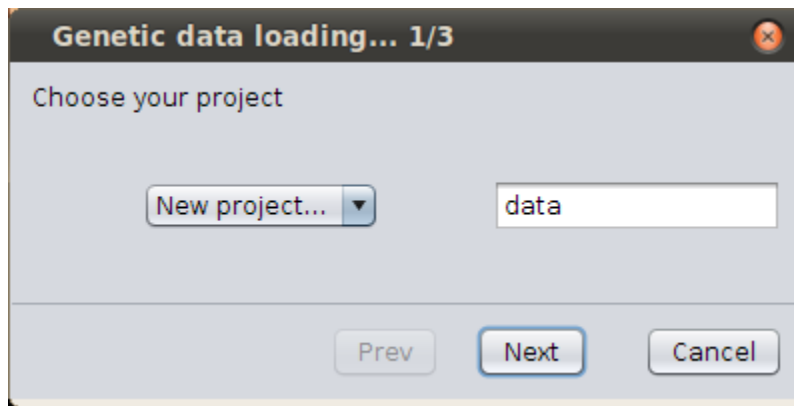


Fig. 1: *Create new project*

The next step is the files choosing; browse your files and select all files inside the “tutorial/Genetic Files” folder. (hold the “shift” or “control” button to select all files, or press the combined keys “Control” and “a” (replace “control” by the “apple” key on mac OS))

The next step shows a summary on all loaded files.

Once the wizard is finished, the new maps will be displayed in the explorer on the left side, inside the chosen project. You can delete a single map, click right on the map Laurence and then click on supress button.

Chromosomes display

For displaying a whole map, click on the map in the explorer, move the mouse in the drawing panel and release the mouse button. The whole map should appear with all chromosomes and linkage groups. You can drag and drop a chromosome or linkage group the same way. Drag the chromosomes ‘1’ from ‘Cardinal_2001’, ‘Groh_1998’ and ‘Mechin_2001_inv1’ maps to see the dynamic comparison. You see now the common markers linked with blue and red lines; red lines show the inverted markers. Now drag the ‘Mechin_2001_inv1’ from the right side to the left side of ‘Groh_1998’ by moving it (holding the mouse left button). We see lots of inversion; right click on the ‘Mechin_2001_inv1’ and select “Reverse linkage group”; the whole linkage group is now in the right order. To remove an element from the display panel, you have 2 ways : * use the middle click of the mouse on the element to remove * right click on the element, then select “remove from view” in the contextual menu

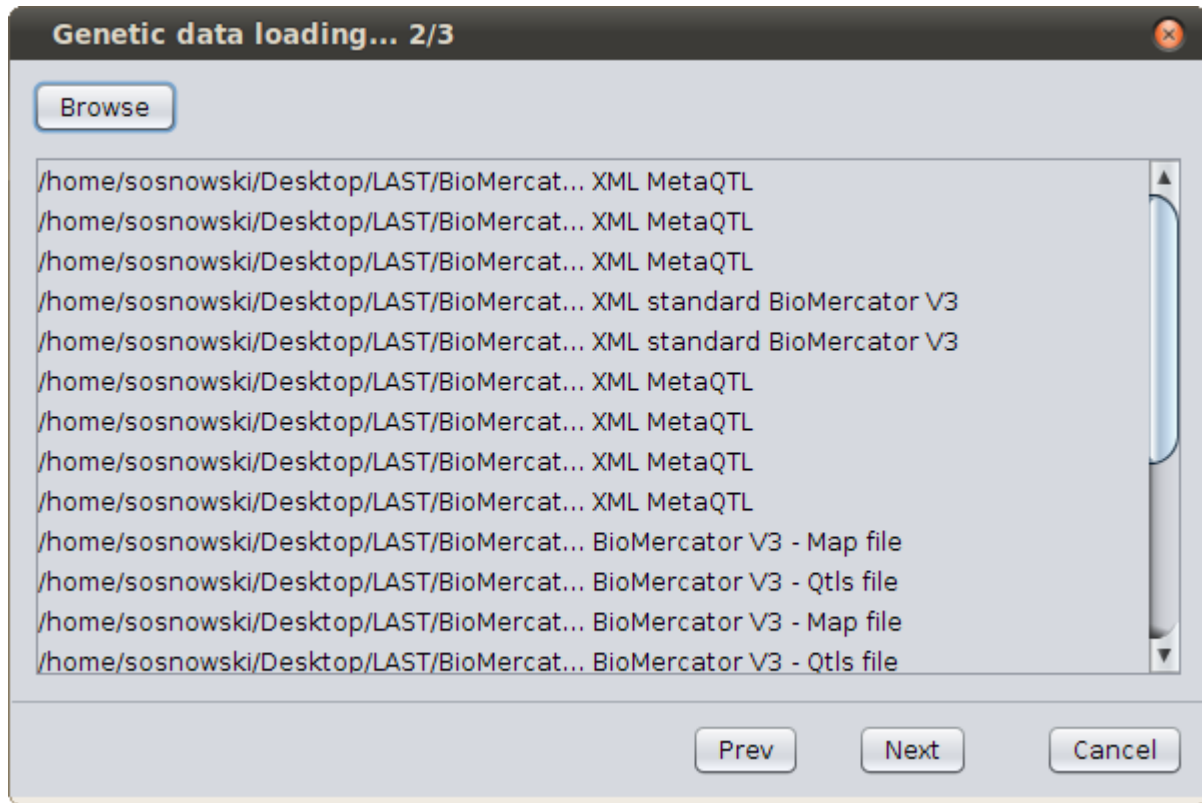


Fig. 2: Genetic maps loaded

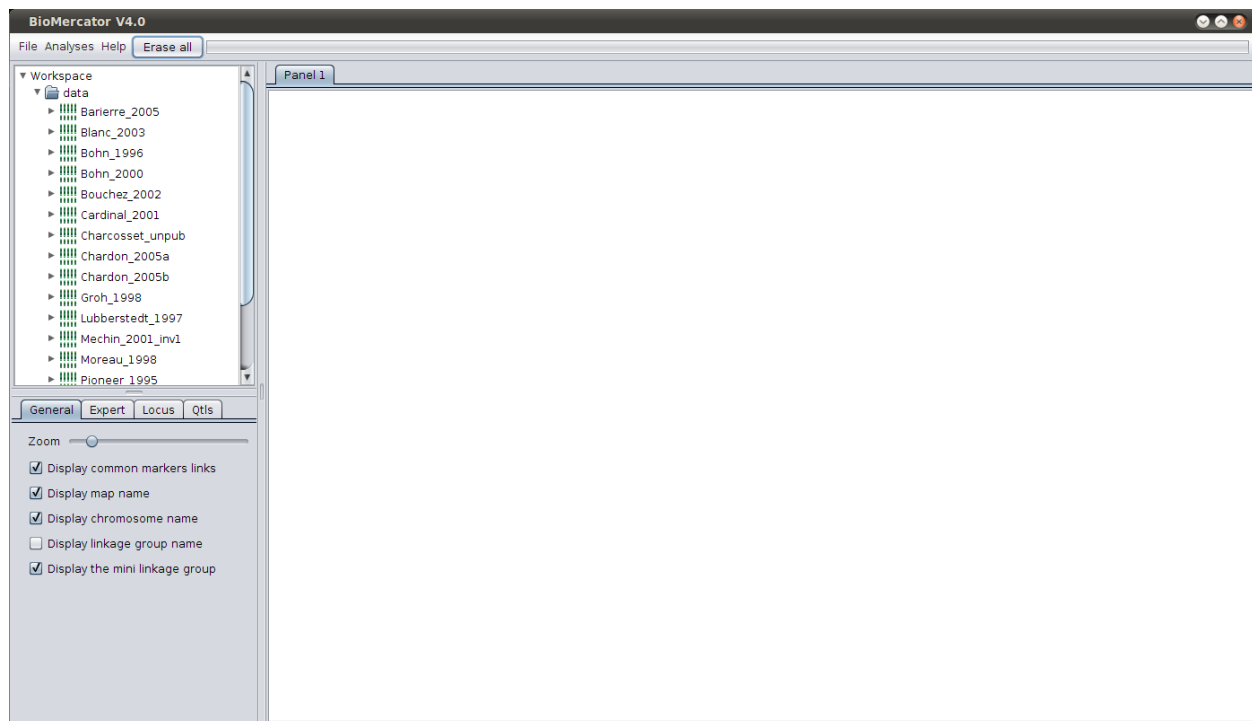


Fig. 3: Maps explorer on the left

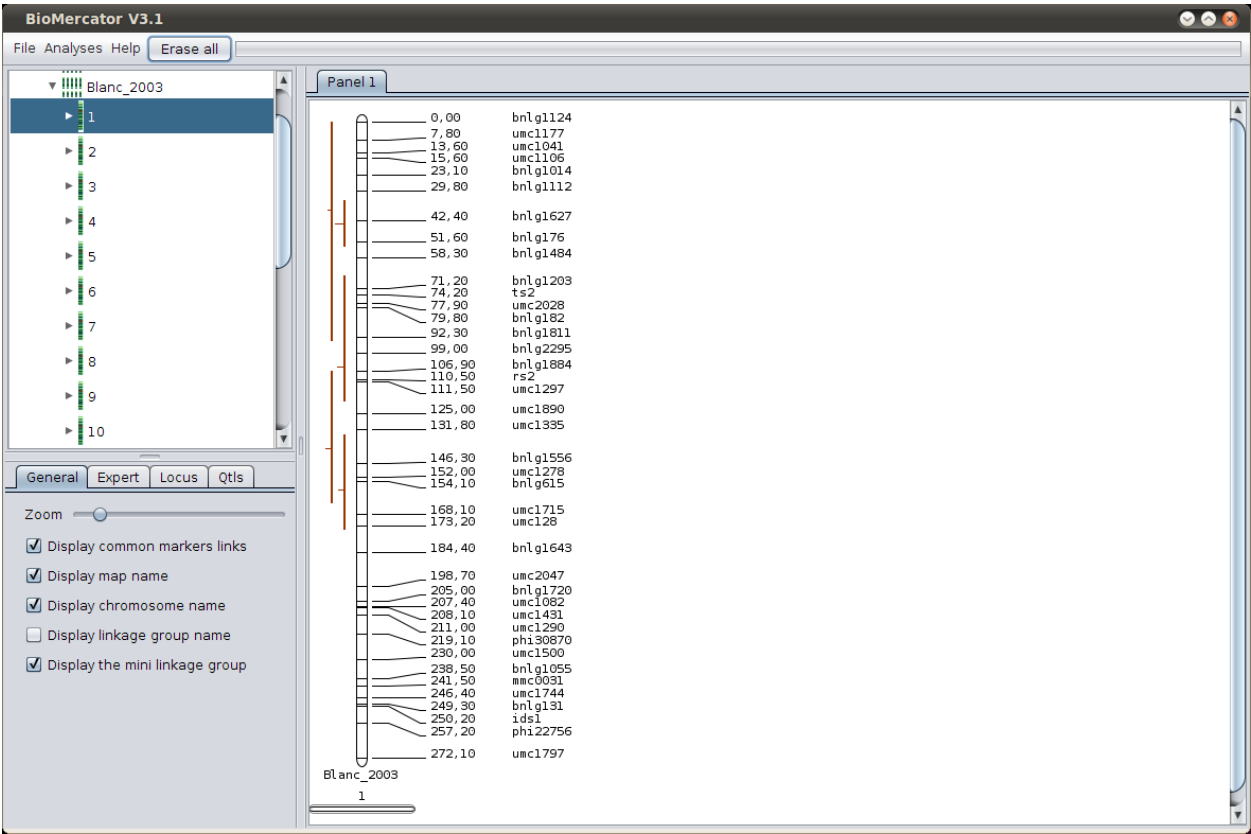


Fig. 4: Chromosome display

Individual zoom

Drag several chromosomes into the display panel; now, put the mouse inside a chromosome, and use the mouse wheel; you will zoom inside the chromosome, keeping other chromosome un-zoomed. (You can also right click on the chromosome, and choose “Zoom in/out”. To keep an idea of your position in the chromosome, a mini-chromosome is displayed below, with a scroller indication your viewing window. (Try this functionality a bit later on chromosomes with more markers on them.)

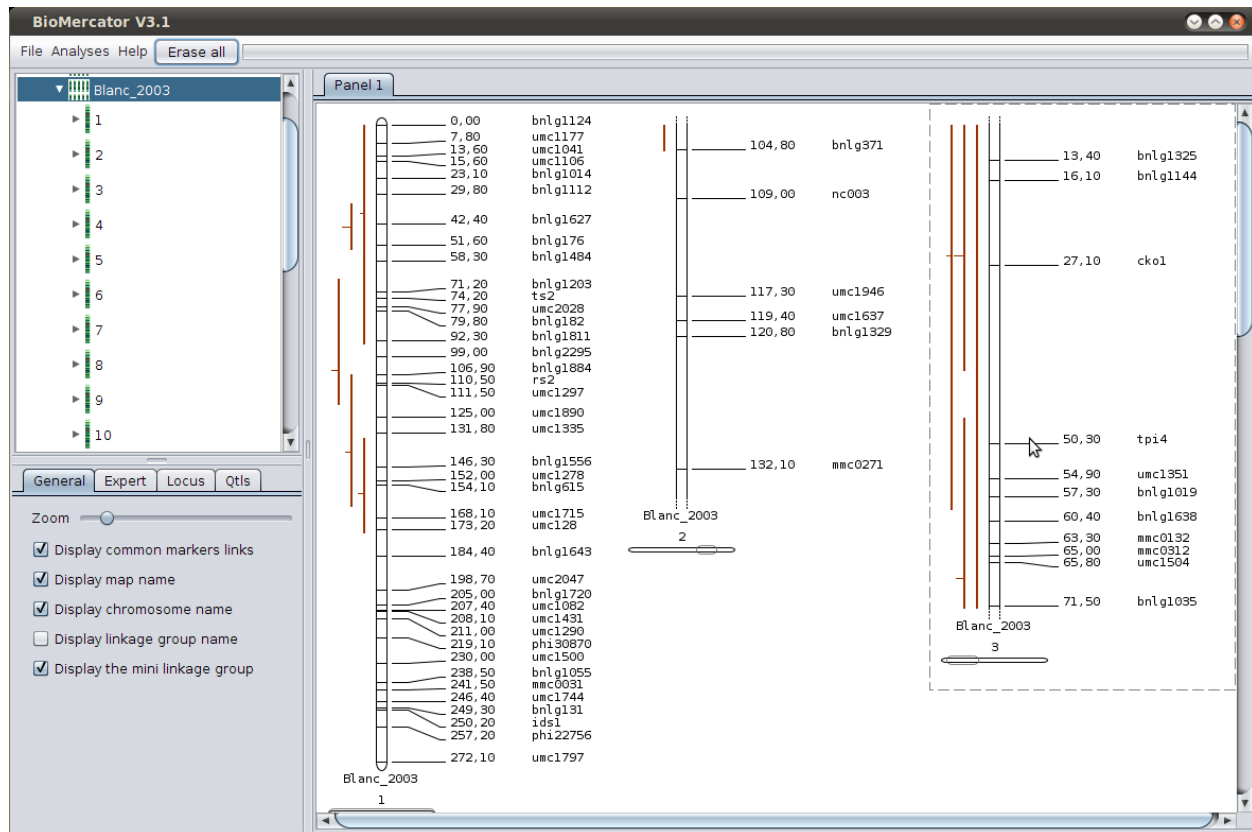


Fig. 5: *Individual zoom*

Cascading zoom

Clear the drawing panel and drag the chromosome 8 from ‘Chardon_2005a’. Clicking on it will launch the cascading view; a scroller appears and can be move/resized with your mouse for exploring the chromosome with more details.

Fig6 here

You can click on the zoomed chromosome for more details.

The zoom allows to display more information on the markers; (some markers are hidden by default when there are too many; this option can be changed in the “Expert” tab, below the maps explorer).

Next step

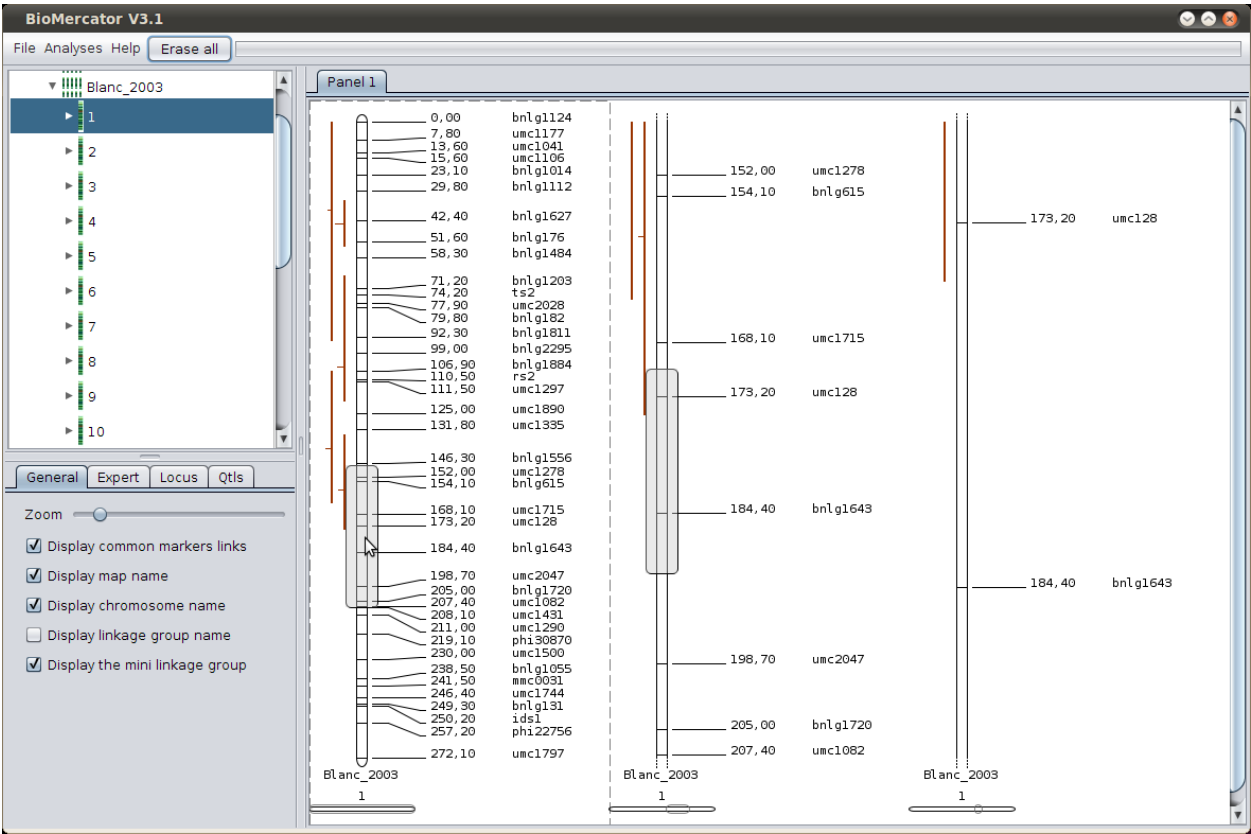


Fig. 6: Cascading zoom

1.2.2 Genetic maps analyses

Previous BioMercator version allows you to directly use command line analyses from the metaQTL toolbox. The standard workflow is the following: * Verify connectivity between input maps * Create a consensus map * Launch a meta-analysis on all QTLs from the consensus map

Connectivity (InfoMap)

This method creates text files about connectivity (where dynamic comparison is only used in a visual way), in order to determine if the can be used for creating a consensus map.

- Click on “Analysis”, “Statistics”, “InfoMap”
- Create a project “Statistics” and validate
- Select all maps from the “Data” project
- Set the result name as “connectivity”
- Click ‘Next’ for launching the analysis

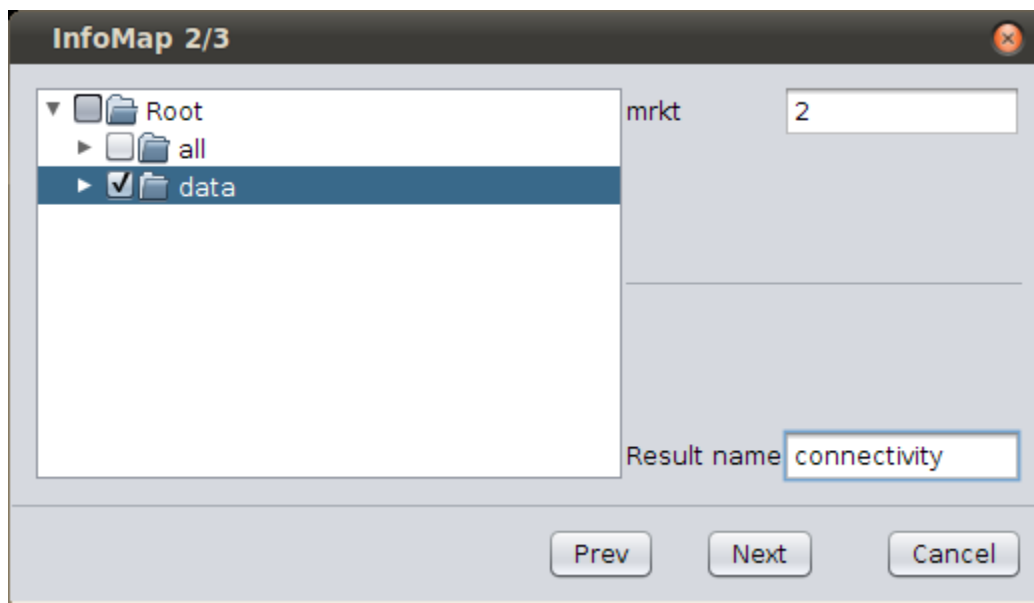


Fig. 7: Select dataset

Once the analysis done, you'll see 2 created files in the Statistics folder; they contain informations about markers and their connectivity.

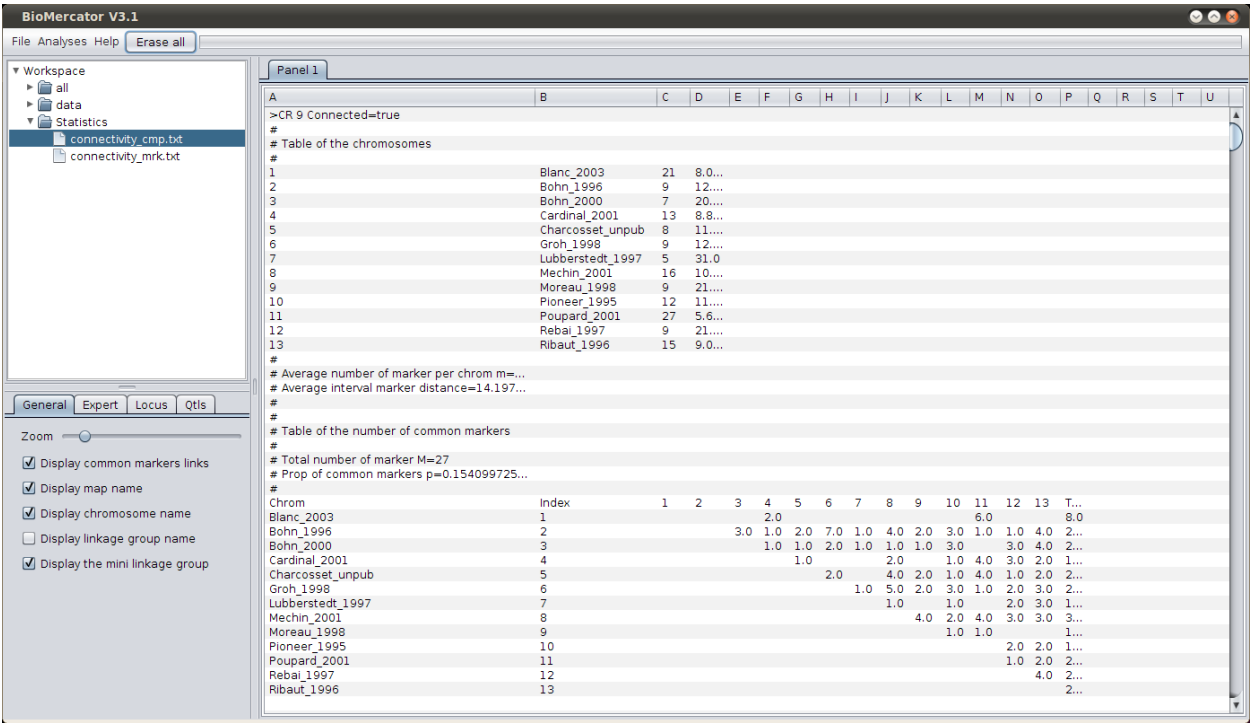


Fig. 8: Connectivity files (1/2)

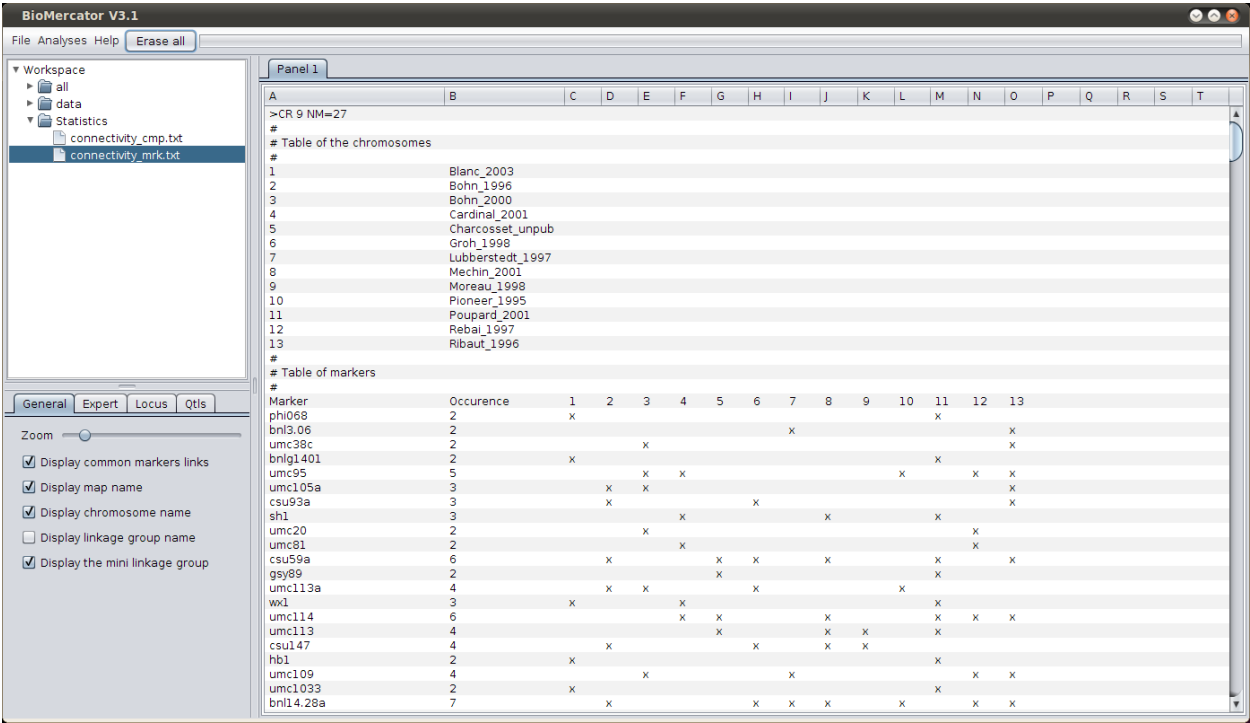


Fig. 9: Connectivity files (2/2)

Consensus map creation (ConsMap)

This analysis is done in single one step, avoiding the iteration part. * Click on “Analysis”, “Map compilations”, “MetaQTL Cons” * Create a project “Consensus” and validate * Select all maps from the “data” project * Set the resulting map name to “pre_consensus” * Launch the analysis

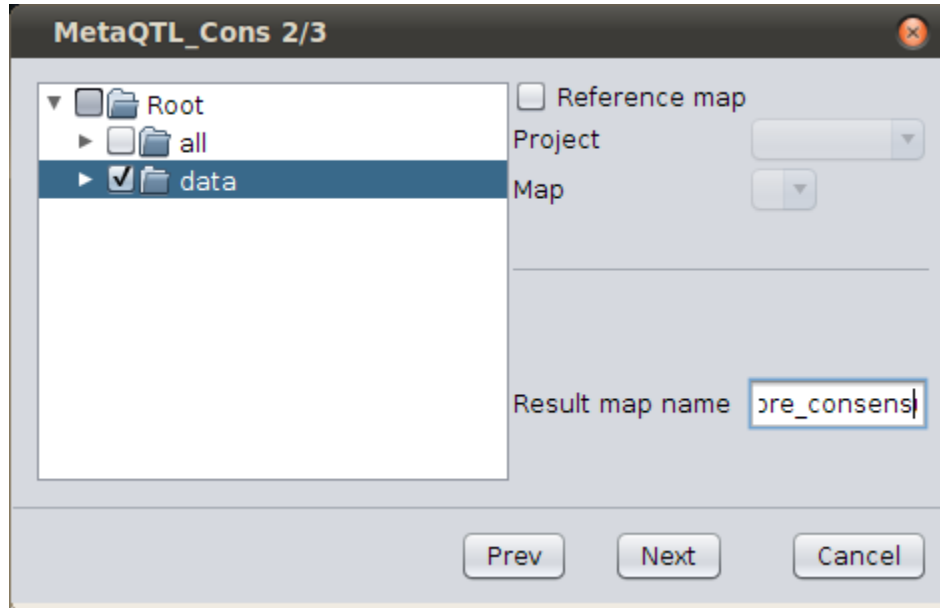


Fig. 10: Select dataset

When the analysis finishes, an information dialog is shown; the analysis should be a success, and the resulting consensus map should be visible in the explorer. Drag it to display it; you notice that no QTL is present; the next analysis is needed.

QTLs projection (QTLProj)

Once the consensus map is created, use this analysis in order to project QTLs from all the maps used. * Click on “Analysis”, “Map compilations”, “MetaQTL QTLProj” * Choose the “Consensus” project and validate * In the next window, in the left explorer, select all maps from the “data” project * In the right explorer, select the “pre_consensus” map from the “Consensus” project * Set the resulting map name to “consensus” * Launch the analysis

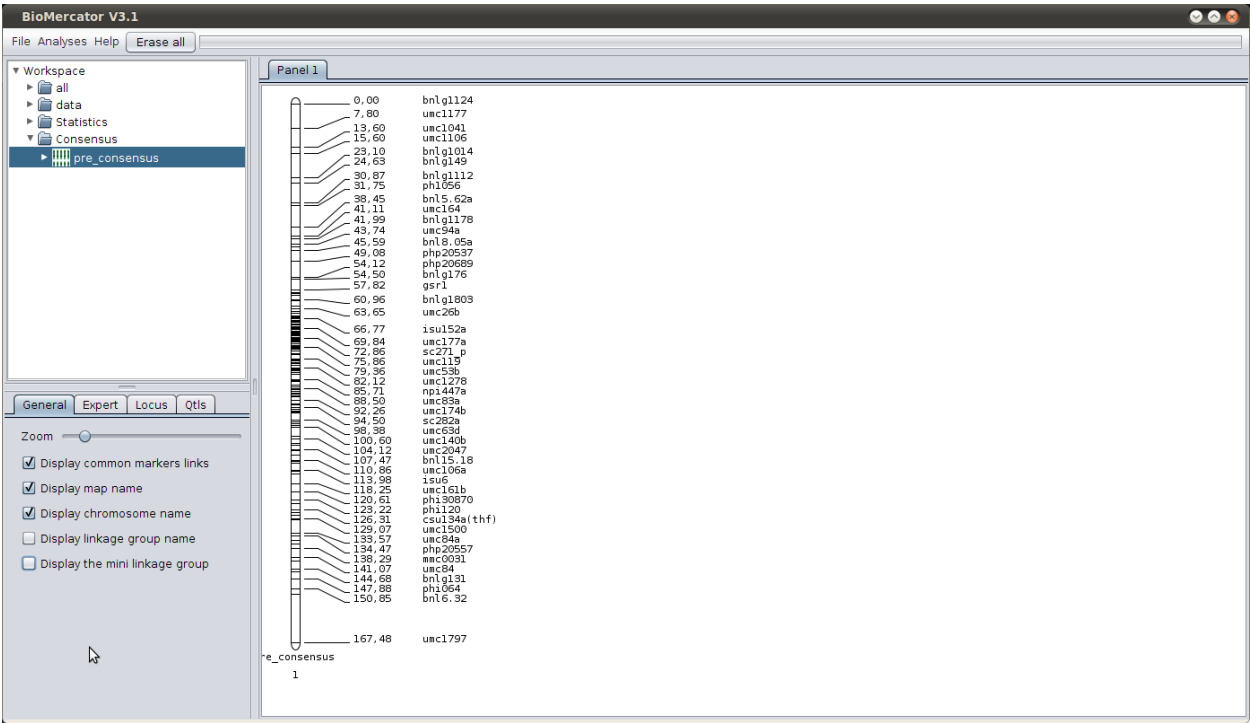


Fig. 11: Display consensus map

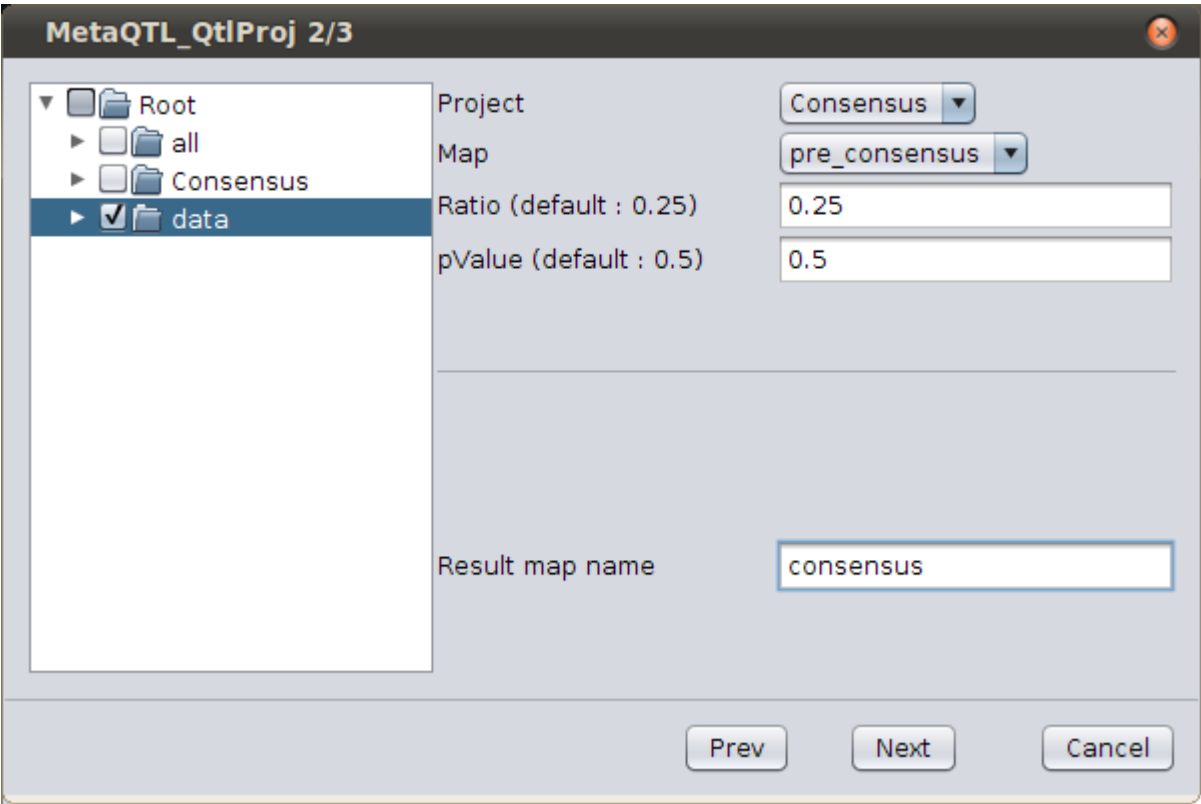
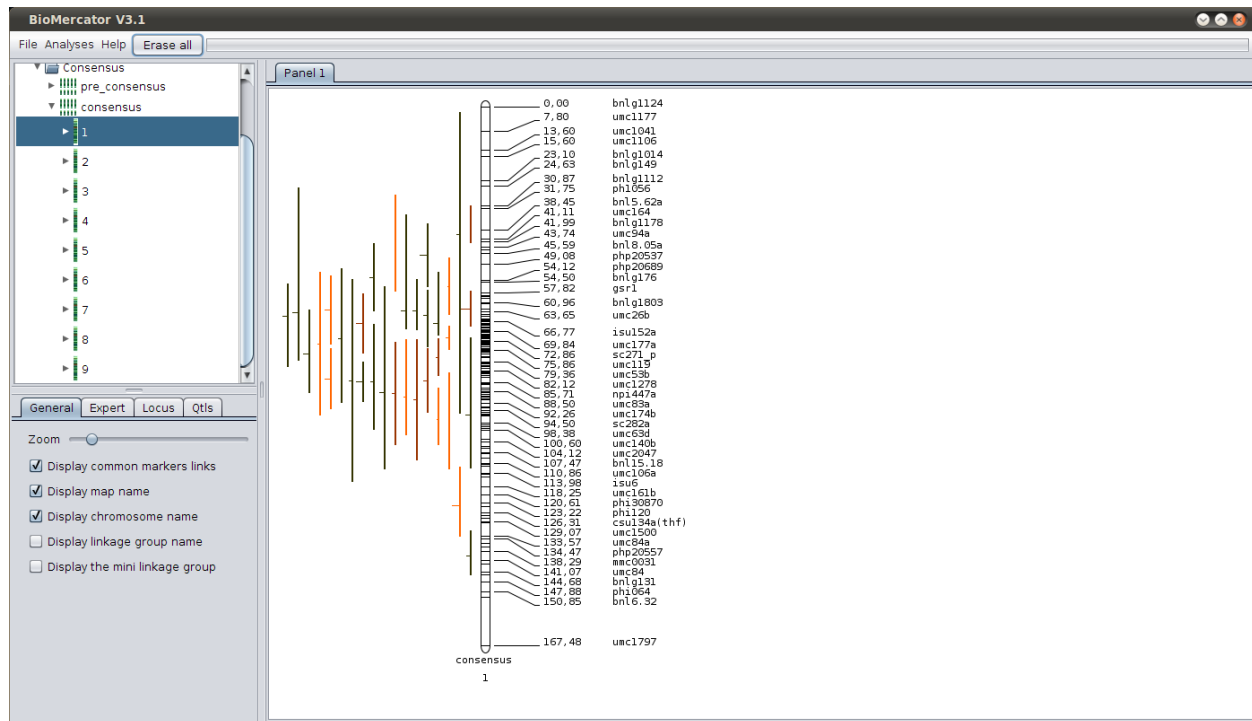


Fig. 12: Select dataset

Fig. 13: *QTLs projection*

QTLs Meta-analysis (QTLClust)

This meta-analysis is a two steps analysis; the first one calculates and estimates the best models (ie number of meta QTLs), so you can choose the one to show in the second.

First step

- Click on 'Analyses', 'QTL Meta analyses', 'MetaQTL Meta analysis 1/2'
- Choose 'meta_v1' for the meta analysis' name
- Select the 'Consensus' project
- Select the 'consensus' map
- Select the '1' chromosome
- Select the '1' linkage group
- Choose to regroup the traits into a single meta trait named 'FT' (for Flowering Time)
- Click 'Next' to launch the analysis

Once the analysis is done, browse the explorer down to the created meta analysis situated inside the previously selected linkage group (project 'Consensus', map 'consensus', chromosome '1', linkage group '1'). You'll see 3 created files; drag the one named 'meta_v1_model.txt'; it corresponds to the most probable model given different criterions.(for more explications, please refer to the 'MetaQTL' software documentation)

MetaQTL_QTLClust 1/2

Meta-analyse

meta_v1

Project

Consensus

Map

consensus

Chromosome

1

Linkage group

1

QTLs choice

kMax

10

ci mode

1

ci miss

1

emrs

50

emeps

1.e-8

☐ Don't regroup traits

☒ Regroup all traits

☐ Use an ontology file

Meta-trait

FT

Choose an ontology file...

Prev

Next

Cancel

Fig. 14: *QTLs meta-analysis parameters*

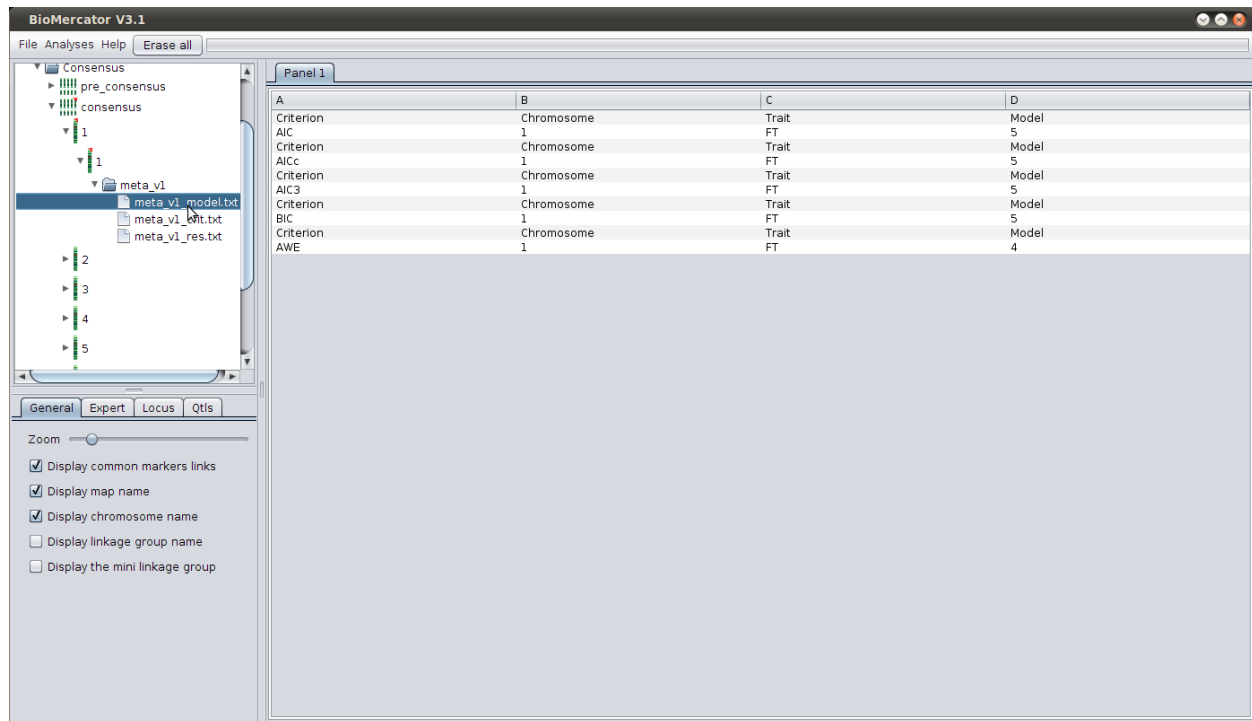


Fig. 15: First step analysis

Second step

- Click on 'Analyses', 'QTL Meta analyses', 'MetaQTL Meta analysis 2/2'
- Select the 'Consensus' project
- Select the 'consensus' map
- Select the '1' chromosome
- Select the '1' linkage group
- Choose 'meta_v1' for the meta analysis' name
- Choose the meta trait named 'FT'
- Click 'Next' to launch the analysis
- Set '5' for the 'best' parameter

Once the analysis is done, browse the explorer down to the created meta analysis situated inside the previously selected linkage group (project 'Consensus', map 'consensus', chromosome '1', linkage group '1'). Inside the folder, you'll see a linkage group named '1', drag it to see the meta-QTLs along with the QTLs and their percentage of belonging.

Next step

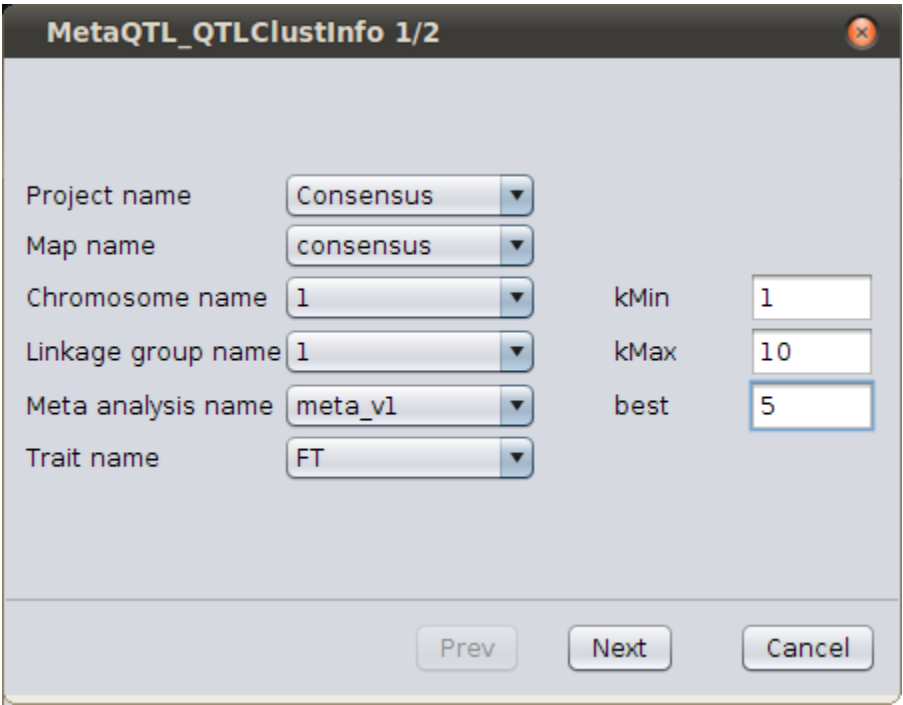


Fig. 16: Second step parameters

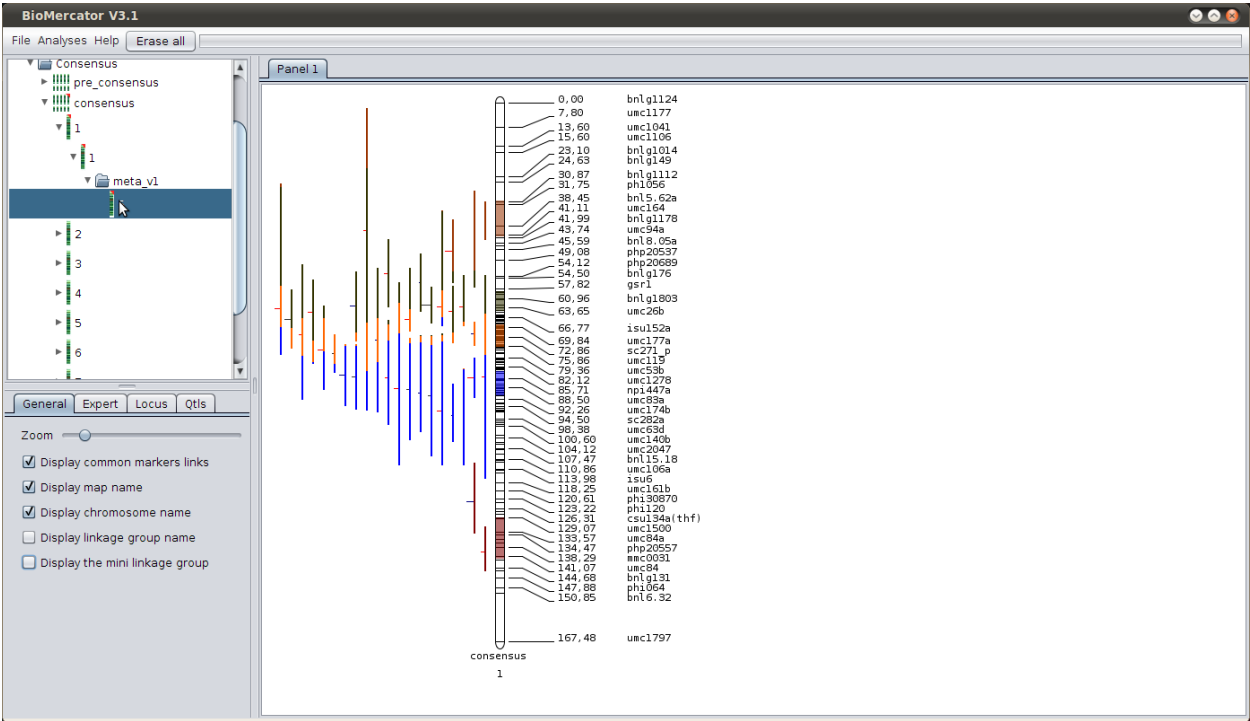


Fig. 17: Display meta-analysis results

1.2.3 Genome viewer

BioMercator integrates structural and functional annotation; a genome version corresponds to a structural annotation, a functional annotation and anchors between the genetic map and the sequence.

Loading a new genome version

You'll need to download 2 files corresponding to the annotation (too heavy to be included in the BioMercator package). You'll find them in the "tutorial/Physic Files" folder. * The structural annotation in maize sequence ZmB73_5b_FGS.gff.gz * The functional annotation ZmB73_5a_xref.txt.gz Extract each file into the 'tutorial/Physic Files' folder.

Click in the menu: 'File/New genome version'; Choose a name for the genome version ('maize_1') Click on browse for the structural annotation, and choose the 'ZmB73_5b_FGS.gff' file

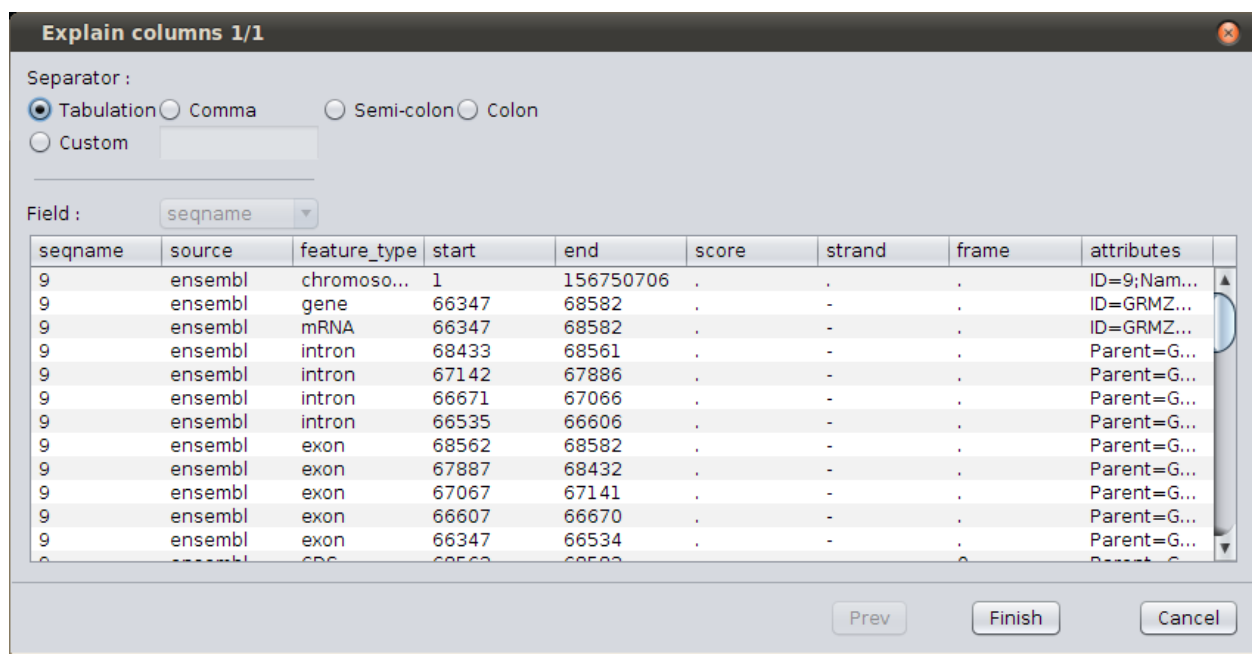


Fig. 18: Browse GFF file

The wizard show you the first lines in the file; it is set by default to use a GFF3 file, so you don't need to change anything; just click on Finish.

Now, click on browse for the functional annotation, and choose the 'xref.txt' file. As before, just click finish as the columns are in the right order. Click on browse for the anchors and choose the 'anchors.csv' file.

As you can see, the wizard fails to display correctly the file; here, the file separator should be "space-comma-space"; select the 'Custom' separator and type ", " (in words : "space" "comma" "space"). The display should now be better.

As you can see, the columns titles don't corresponds to the files columns. Click on the column and use the combo box to change their title until they are similar to the following screenshot.

Carefully verify that the columns are the same, and then click on 'Finish' At last, click on browse for the database links and choose the 'dblinks.txt' file. Choose tabulation as separator, and then click on 'Finish'.

You should wait for the files to be loaded until you get the following dialog; then, just click 'Finish' to save the genome version.

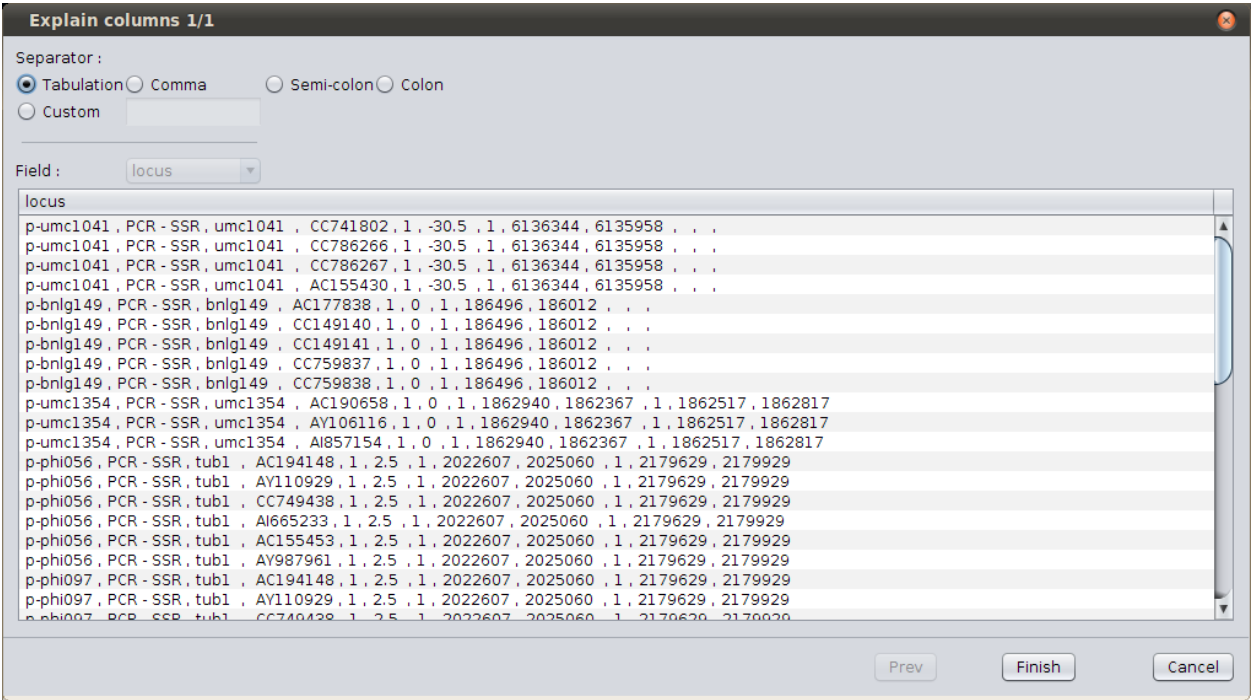


Fig. 19: Browse anchor file

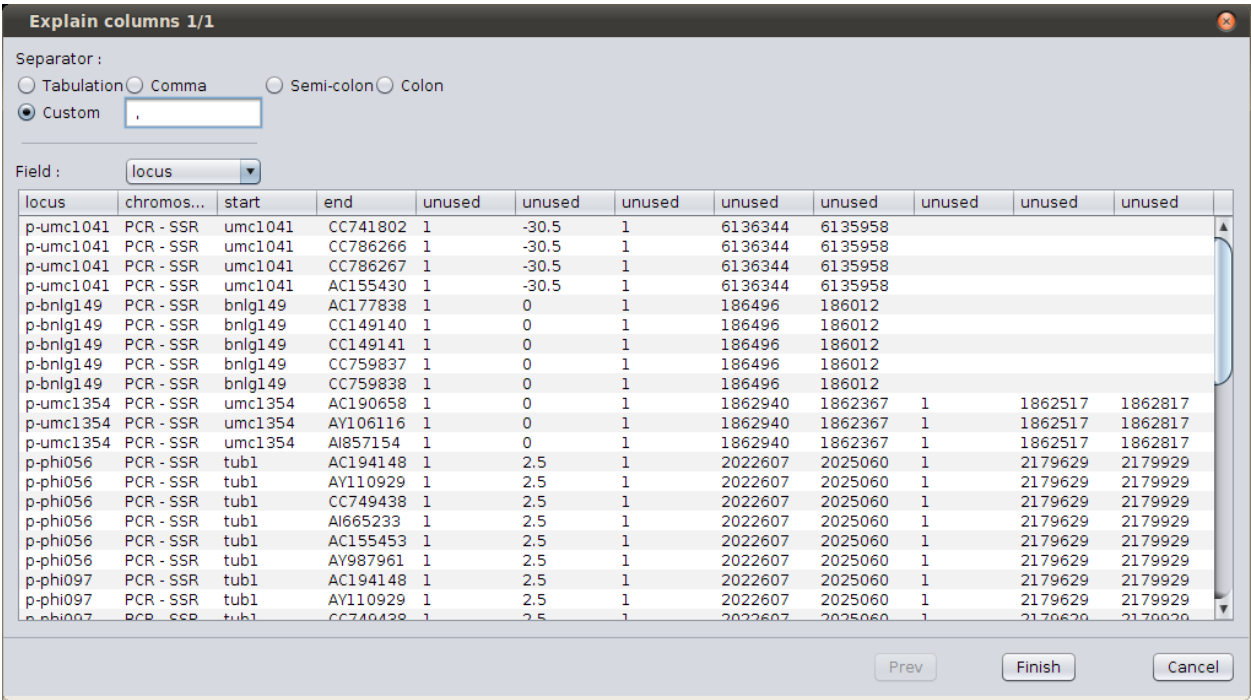


Fig. 20: Define custom separator

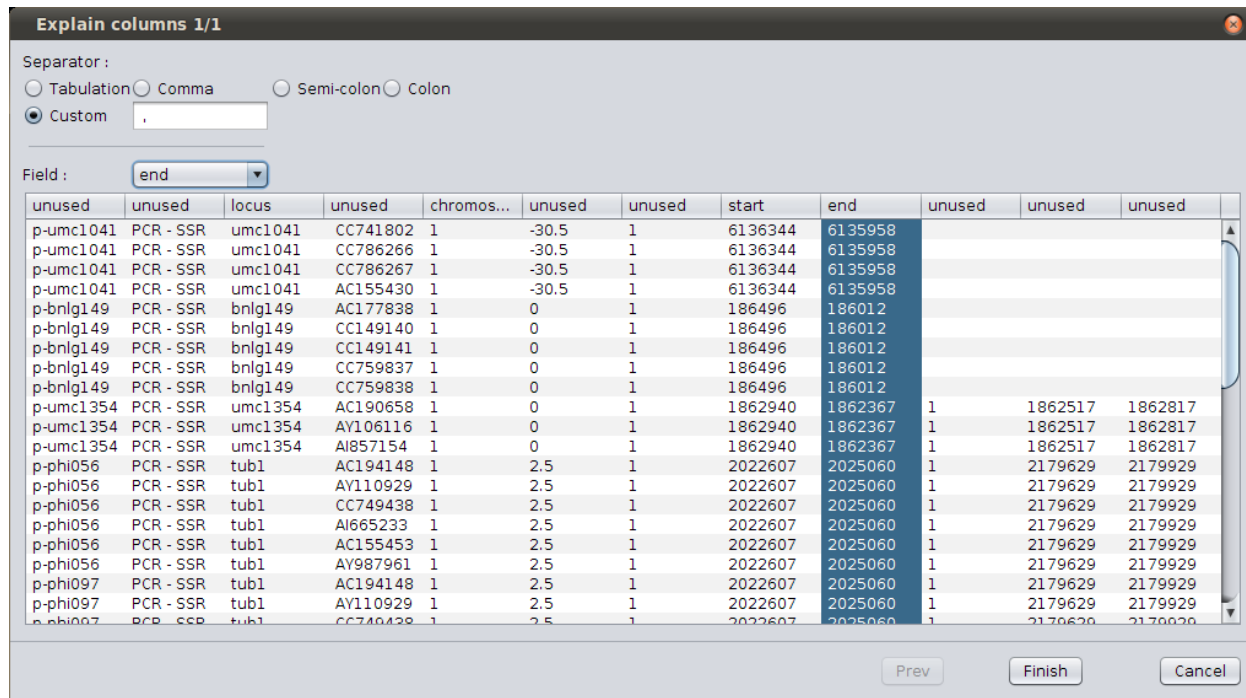


Fig. 21: Change columns order



Fig. 22: Progress state of file loading

Display a new genome version

We're now going to display the new genome version; in the tab 'Genome Version' select 'maize_1'; a progress bar should appear; wait until it's completely loaded. Drag the linkage group from the explorer: Consensus/consensus/1/1/meta_v1 and unselect the QTLs display under the QTL tab (for a clearer view). Drag the scroller on the genetic map, and you'll see it move along the sequence (vertical line in the middle), as well as the genes under the selected area (on the right side). As the ratio between cM and bp isn't always the same, look at the scroller's size on the sequence as you move the other one on the genetic map. (you can also resize the scoller or move the other one). You can also move the mouse over a gene. A menu appears with links to URGI, gramene or maizeGDB databases. You can click on one of them, it will open a browser with the page corresponding to the selected gene.

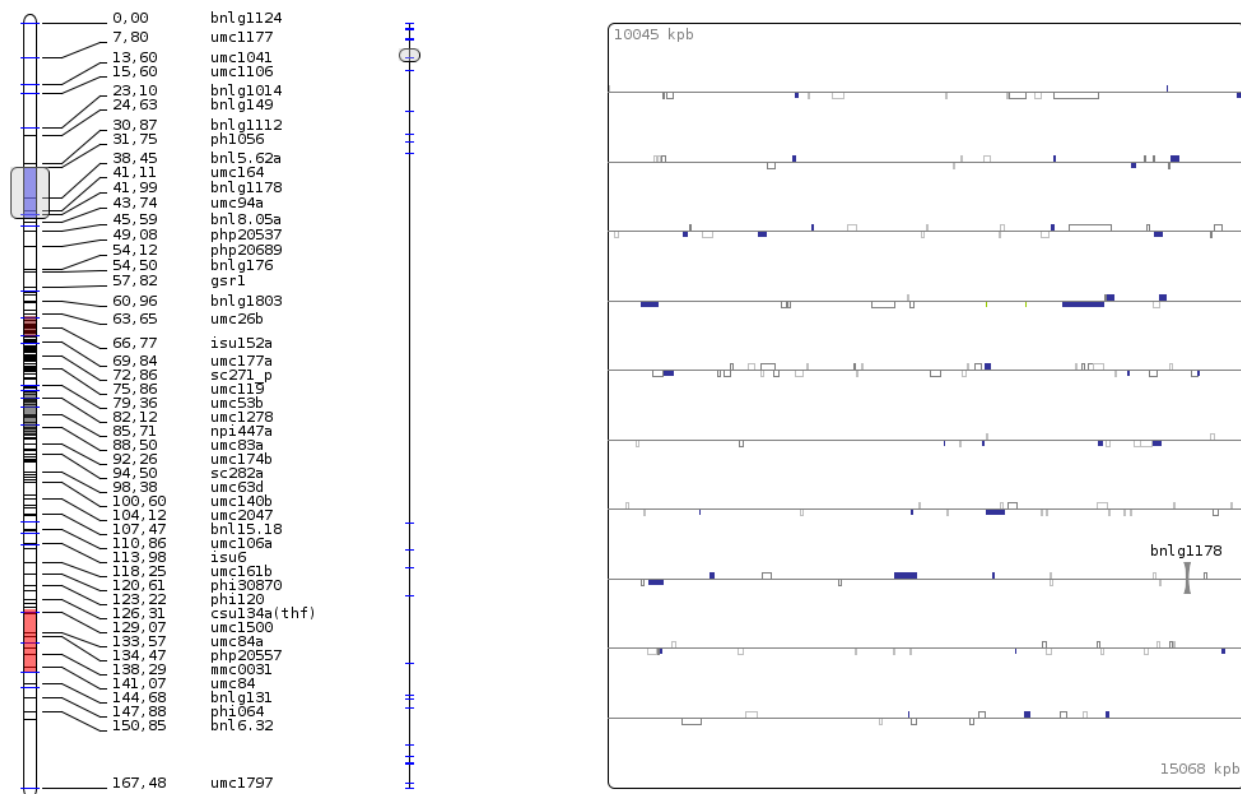


Fig. 23: Display genome view

Click then on the first meta-QTL. The Cursor is positioned automatically around it. Now, in the 'View' menu, choose 'Show the GO view'. A panel appears at the bottom. The colors shown in the chart represent the Gene Ontology terms, and are used for the genes color; clicking on a chart section will enlighten the corresponding genes, and clicking a second time (not a double-click) will make you go deeper in the GO hierarchy.

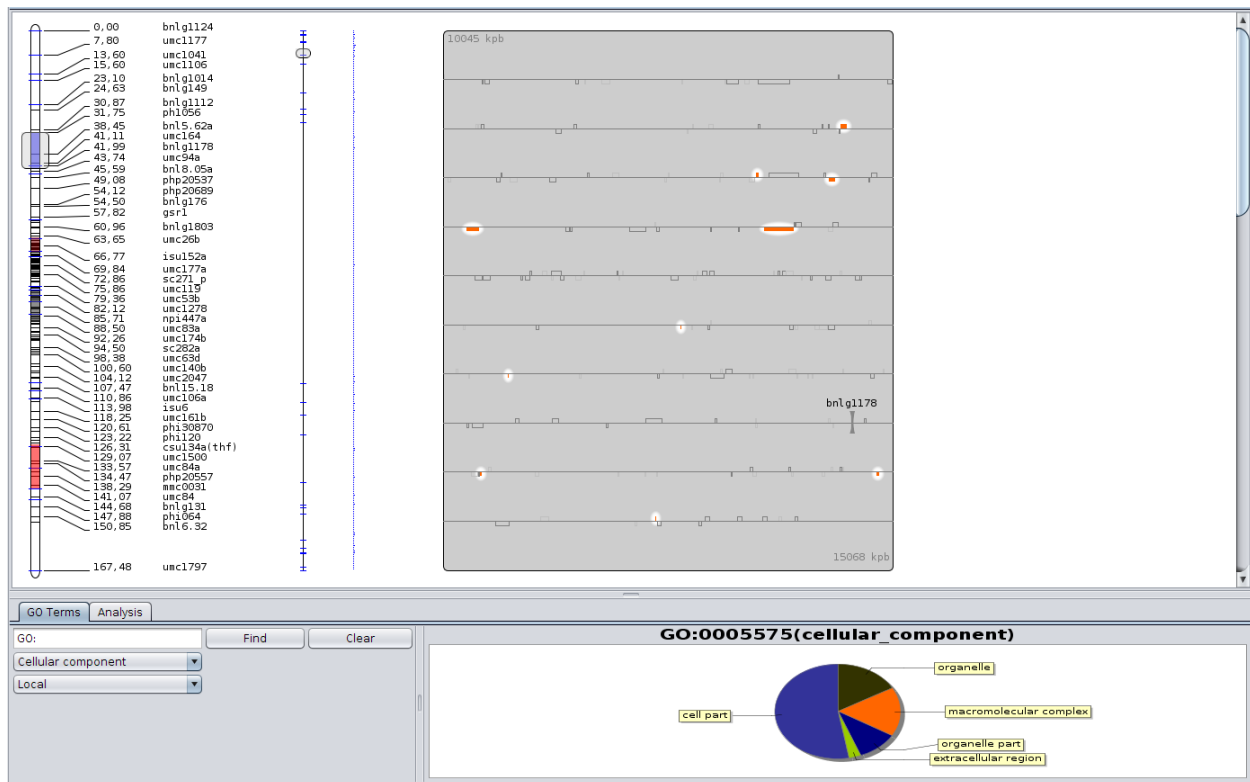


Fig. 24: Display Gene Ontology piechart

Analyzing GO terms over-representation

Another tool in BioMercator is the GO representation analysis; select the Analysis tab in the new panel, check the scroller is situated around the first metaQTL (click on the metaQTL if not) and just launch the analysis. The analysis gives you a list of GO terms over and under represented in the selected area (here, the meta-QTL) compared to the whole linkage group. The columns gives you more information about the term. Clicking on it will enlighten the genes with the corresponding annotation. Other analysis parameters are available, you could decide to analyze the representation along all intervals of a trait compared to the whole genome for instance.

Finally you can export genes into GFF3 file; Right click on the genes window, and select 'Export to GFF'.

You can export all features or only genes. You can choose the window of export (just the scroller's area, the whole chromosome or the whole genome). And you can choose to export only genes with the corresponding GO functional annotation. Browse to select the path of your new file, and click next. You'll see in the created GFF the genes with all the structural annotation, but with also the GO annotation. The QTLs and meta-QTLs will be present in the GFF file.

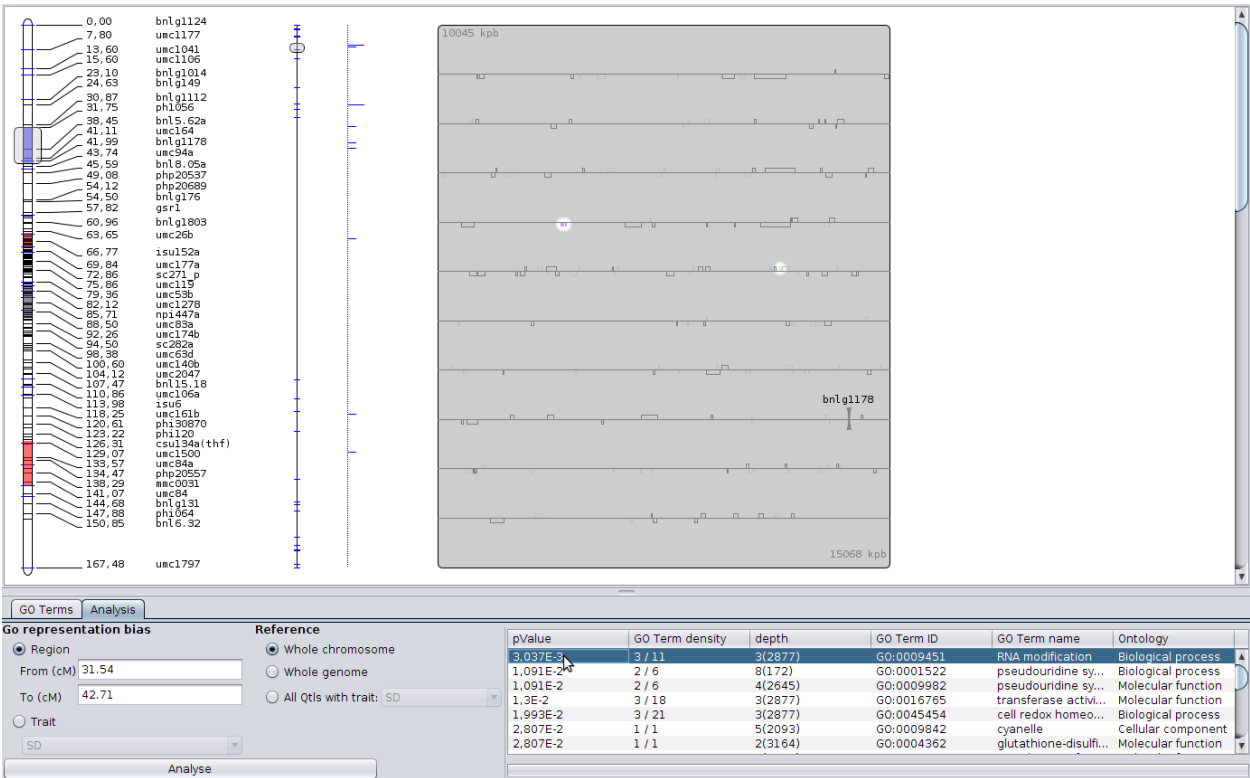


Fig. 25: Emphasis annotated genes

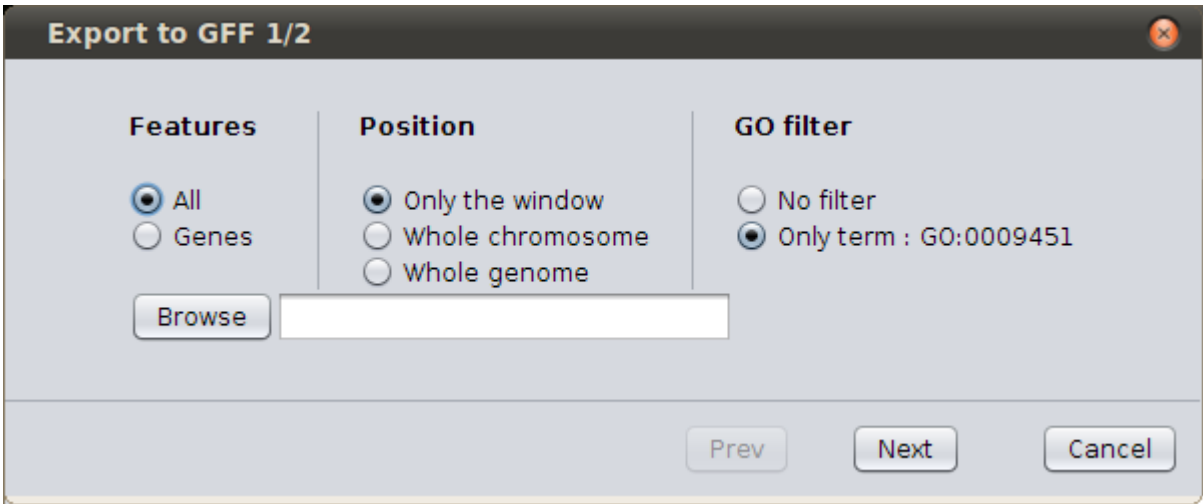


Fig. 26: GFF export form

1.3 Load data

1.3.1 Genetic Maps files

1.3.2 Physical Maps files

Structural annotation file

Functional annotation file

Anchor file

Working without genetic map

1.3.3 GWAS data files

1.4 Export data

1.4.1 Export a genetic map

1.4.2 Export genome region as a GFF file

1.4.3 Export a screenshot

1.5 Analyses

1.5.1 Statistical Analyses

Connectivity

Connectivity image

1.5.2 Map Compilation

Iterative map projection

Consensus map compilation

QTL projection

1.5.3 QTL meta-analysis

Gerber & Goffinet method

Veyrieras method

1.6 Display data

1.6.1 Genetic maps

1.6.2 Genome browser

1.6.3 GWAS data