

# QTLViewer: an interactive webtool for genetic analysis in the Collaborative Cross and Diversity Outbred mouse populations

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## Abstract

The Collaborative Cross and the Diversity Outbred mouse populations are related multiparental populations, derived from the same 8 isogenic founder strains. They carry >50 M known genetic variants, which makes them ideal tools for mapping genetic loci that regulate phenotypes, including physiological and molecular traits. Mapping quantitative trait loci requires statistical and computational training, which can present a barrier to access for some researchers. The QTLViewer software allows users to graphically explore Collaborative Cross and Diversity Outbred quantitative trait locus mapping and related analyses performed through the R/qtl2 package. Additionally, the QTLViewer website serves as a repository for published Collaborative Cross and Diversity Outbred studies, increasing the accessibility of these genetic resources to the broader scientific community.

**Keywords:** multiparental populations; MPP; genome-wide association studies; GWAS; quantitative trait loci mapping; QTL mapping

## Introduction

Multiparental populations (MPPs; [de Koning and McIntyre 2017](#)) improve upon traditional experimental crosses by interbreeding more than 2 isogenic strains capturing greater genetic diversity. The Collaborative Cross (CC) and Diversity Outbred (DO) mouse populations are related MPPs that have been used to genetically dissect complex traits, including obesity ([Svenson et al. 2012](#)), bone strength ([Al-Barghouthi et al. 2021](#)), short-term memory ([Hsiao et al. 2020](#)), and benzene response ([French et al. 2015](#)). As recombinant populations, the CC and DO are well-suited for mapping quantitative trait loci (QTLs), which can be powerful in studies of -omic traits, including gene expression, protein abundance, and chromatin accessibility ([Aylor et al. 2011](#); [Chick et al. 2016](#); [Abu Toamih Atamni et al. 2018](#); [Keller et al. 2018](#); [Keele et al. 2020, 2021](#); [Takemon et al. 2021](#); [Gerdes Gyuricza et al. 2022](#)).

The CC and the DO were bred from the same 8 strains: A/J, C57BL/6J, 129S1/SvImJ, NOD/ShiLtJ, NZO/HILtJ, CAST/EiJ, PWK/PhJ, and WSB/EiJ ([Churchill et al. 2004, 2012](#); [Iraqi et al. 2012](#)). The use of classical and wild-derived founder strains drives high levels of genetic variability ([Yang et al. 2007, 2011](#)). The CC represents ~60 recombinant strains that are fully inbred (>99%) whereas DO mice are outbred, thus each mouse is genetically unique. Mapping resolution is finer for the CC and the DO compared to intercrosses or backcrosses due to additional generations of meiosis, which facilitates the identification of candidate genes ([Solberg Woods 2014](#)).

Mouse experiments support the collection of multiple types of data on the same individuals, including physiological traits and molecular assays (e.g. gene expression) across multiple tissues. Integrative approaches like mediation analysis can be used to delineate the relationships among traits with shared QTLs ([Chick et al. 2016](#); [Keele et al. 2020](#)). This series of analyses enable richer findings from CC and DO experiments but can also obstruct researchers without statistical and computational expertise.

The R/qtlcharts software ([Broman 2015](#)) provided a set of interactive data visualization tools for QTL analysis, built on the JavaScript library D3 ([Bostock et al. 2011](#)) and accessible from R ([R Core Team 2021](#)), but it does not provide a comprehensive or user-friendly graphical user interface. [Konganti et al. \(2018\)](#) developed the gQTL webtool, a graphical interface QTL mapping tool for CC data based on the R package DOQTL ([Gatti et al. 2014](#)). However, DOQTL is no longer maintained and instead, the R/qtl2 package ([Broman et al. 2019](#)) should be used. Here, we present the QTLViewer software (<https://qtlviewer.jax.org>), an interactive QTL mapping webtool built for both CC and DO that utilizes the R/qtl2 package to perform QTL mapping as well as variant association mapping based on the founder strain genotypes. QTLViewer can perform mediation analysis for QTL through -omic traits to identify candidate causal mediators of physiological trait QTL. The primary goal of QTLViewer is to allow users to interactively perform and visualize QTL mapping for publicly

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available CC and DO datasets, and so, QTLViewer represents a data repository that can be also downloaded for further investigation. This software will empower researchers to analyze and explore CC and DO data while facilitating access to relevant data and findings from others across the community.

## Methods

The QTLViewer is an interactive web application designed to interactively perform and visualize QTL mapping and related analyses in experimental CC or DO data. We make use of modern computing tools, such as application programming interfaces (APIs) and Docker containers, to make QTLViewer efficient, portable, and extendable.

## Implementation

The QTLViewer is comprised of 3 Docker (<https://www.docker.com/>—v18.09.3 last accessed 06/13/2022) images that are managed with Docker-Compose (<https://docs.docker.com/compose/>—v1.23.2 last accessed 06/13/2022). The user interface is contained in the churchilllab/qt2web container (v1.0.0), which is a Python (v3.6.9) web application that processes all requests made by the user. Each request is parsed and analyzed to determine which analyses need to be performed.

All computational requests are performed by the churchilllab/qt2rest container (v0.1.0). This container loads data upon startup and listens to Web API calls via RestRserve (<https://restrserve.org/>—v0.4.1 last accessed 06/13/2022). All data returned from these APIs are in JSON format. Since some requests require longer time to complete than others, we use another instance of the churchilllab/qt2web container to manage them. We utilize Celery (<https://docs.celeryproject.org/>—v4.4.7 last accessed 06/13/2022) with a Redis (<https://redis.io/>—v6.2 last accessed 06/13/2022) backend as a task queue.

The containers work together to handle all requests. Requests generated from the web page are submitted via AJAX (Asynchronous JavaScript) and processed by the churchilllab/qt2web container. A dialog box with a spinning wheel is displayed to the end-user to show that the request is being processed. The request is parsed to determine which APIs need to be called. There could be one to many API calls per request. Those API calls are bundled together and submitted to the task queue for processing. Once the task queue receives the request, a unique ID is generated and sent back to the web page. The web page parses the unique ID and begins polling the task queue until the task is complete or an error occurs. While the web page is polling the task queue for a status, the API calls are submitted to the churchilllab/qt2rest container for computational analyses. When all API calls are complete, the task queue bundles the data together and marks the job as complete. The background polling that has been happening on the web page see that the job is complete, removes the dialog box, and processes the JSON data to display the appropriate graphs and data.

A diagram illustrating the implementation process is shown in Fig. 1.

## Supporting tools

QTLViewer relies on a set of tools that facilitate the analyses of MPPs. Ensimpl (v 1.0.0) is a customized small version of Ensembl (<https://www.ensembl.org> last accessed 06/13/2022) that provides a web API to retrieve genomic information. Ensimpl extracts the necessary data from Ensembl and creates efficient SQLite (<https://www.sqlite.org/> last accessed 06/13/2022)

databases for querying purposes. The databases are separated by Ensembl release and species (mouse and human). The web API is written in Python utilizing FastAPI (<https://fastapi.tiangolo.com/> last accessed 06/13/2022) framework. The API functionality provides search, genomic location lookup, gene lookup, and gene history.

In addition to Ensimpl, QTLViewer also utilizes a Founder SNP Database (<https://churchilllab.jax.org/foundersnps> last accessed 06/13/2022). The Founder SNP database is a custom version of the Sanger SNP VCF files that have been processed via a Python script and stored in an SQLite (<https://www.sqlite.org/> last accessed 06/13/2022) database. The SNP genomic location and allele calls for the 8 founder strain alleles are stored. A custom Strain Distribution Pattern is stored for easy query and dissemination of data. Currently, GRCh38 ([https://www.ncbi.nlm.nih.gov/assembly/GCF\\_000001635.20/](https://www.ncbi.nlm.nih.gov/assembly/GCF_000001635.20/) last accessed 06/13/2022) release 1410 and 1505 are supported.

Data visualizations make use of the JavaScript library Highcharts (<https://highcharts.com> last accessed 06/13/2022) which is built on D3 (Bostock et al. 2011).

## Statistical methods

The QTLViewer carries out several types of analysis to map and characterize QTL. QTL mapping analysis is run using the R/qt2 package (Broman et al. 2019—v0.28). The QTLViewer maps QTL by testing for an additive locus effect (additive QTL) or a locus-by-factor interaction effect (interactive QTL). For the additive model, the following linear mixed model is used to test an additive locus effect at loci spanning the genome:

$$\text{trait}_i = \text{QTL}_{im} + \text{covariates}_i + \text{kinship}_i + \text{error}_i, \quad \text{Equation 1}$$

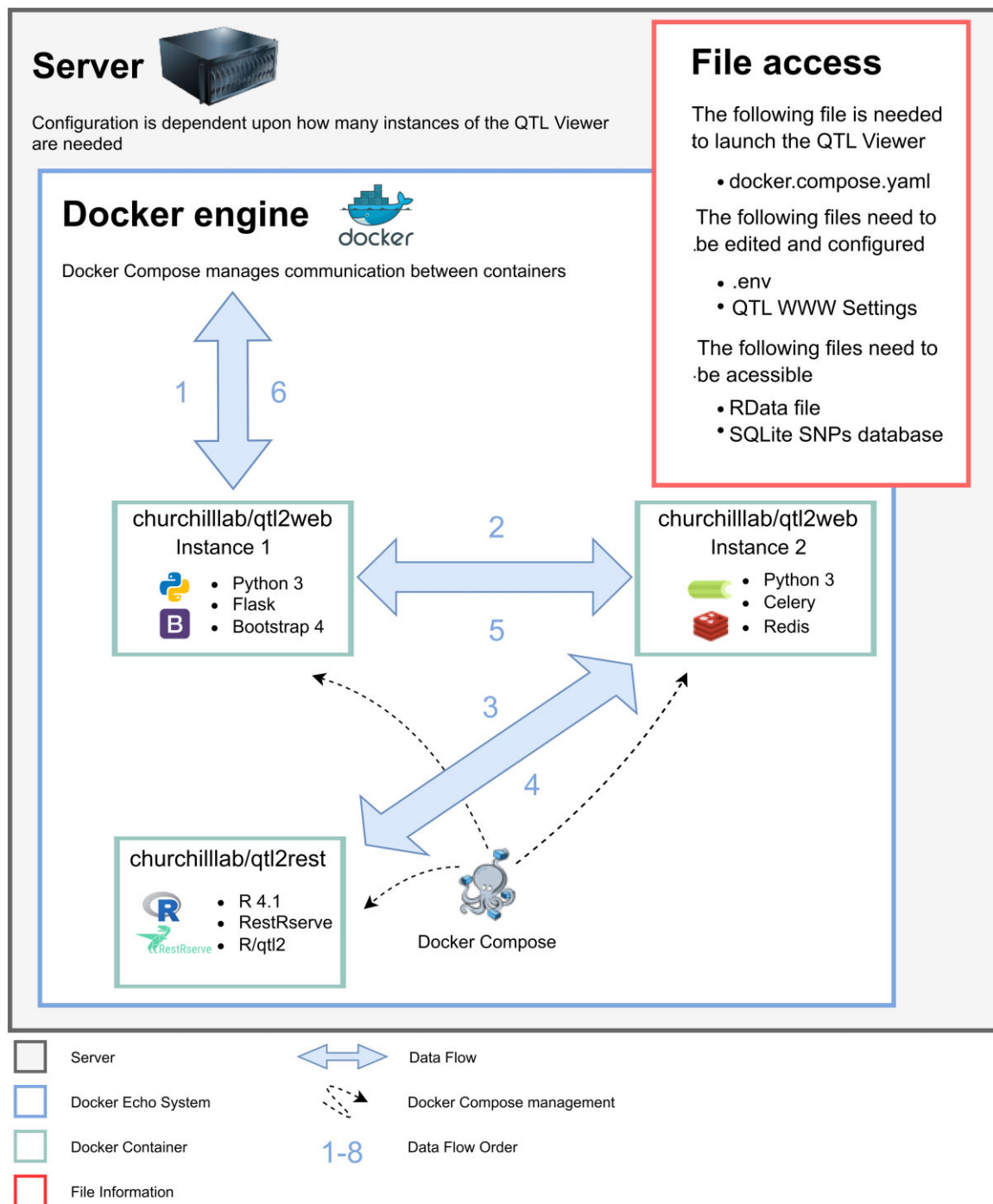
where  $\text{trait}_i$  is the phenotype value of mouse  $i$ ,  $\text{QTL}_{im}$  is the effect of locus  $m$  on mouse  $i$  being tested,  $\text{covariates}_i$  is the cumulative effect of all covariates on mouse  $i$ ,  $\text{kinship}_i$  is a random term that captures noise variation for mouse  $i$  due to population structure, and  $\text{error}_i$  is the independent random noise for mouse  $i$ . The structure of the kinship term is encoded in a genetic relationship matrix ( $K$ ) estimated from the genotypes. We use the “leave one chromosome out” (LOCO) approach in which the  $K$  used for each locus  $m$  fit by Equation (1) excludes all markers from the chromosome of locus  $m$ , which improves QTL mapping power (Wei and Xu 2016). The QTLViewer uses the same statistical model for mapping on all chromosomes including the X chromosome. The precomputed genotype probabilities take account of sex-specific differences (Broman et al. 2006).

For additive QTL, the  $\text{QTL}_{im}$  term represents allele dosages founder haplotypes at the locus. The QTLViewer will also plot the regression coefficients from the  $\text{QTL}_{im}$  term, i.e. the founder allele effects, when doing haplotype-based analysis. These effects can also be re-estimated as best linear unbiased predictors (BLUPs), which reduce the impact of rare alleles and can make signals clearer.

For interactive QTL, a similar model to Equation (1) is used to test a locus-by-factor interaction effect at loci across the genome:

$$\begin{aligned} \text{trait}_i = & \text{QTL}_{im} + \text{factor}_i + (\text{QTL}_m \times \text{factor})_{im} + \text{covariates}_i \\ & + \text{kinship}_i + \text{error}_i, \end{aligned} \quad \text{Equation 2}$$

where  $\text{factor}_i$  is a covariate of interest for mouse  $i$  that may have an interaction effect with genotype at the locus  $m$ ,  $(\text{QTL}_m \times \text{factor})_{im}$  is the QTL-by-factor interaction term at locus



**Fig. 1.** Diagram of QTLViewer data flow. (1) URL is parsed and, if not an API call, returns requested information (Flask is used as the web application framework and Bootstrap 4 as the interface framework). (2) If the requested URL is an API call, the tool either returns a cached version if found or proceeds to call the API (Celery is used as a distributed task queue). (3) R/RestRserve package routes the URL to the correct R method to be performed. (4) A compressed JSON object is returned and if the HTTP request creates several QTL API calls, Redis will store the intermediate result from each call until all of them are finished. (5) Results are cached, and data is returned to Flask/Python. (6) The request is complete. The headers are checked and, if the response needs to be compressed, Flask will compress the data and send to the end-user. The front-end user interface is now responsible for rendering the data.

$m$  being tested, and all other terms as defined before. Possible factors include sex and age.

The QTLViewer can also perform association mapping on biallelic variants using both additive and interactive models. Rather than encoding genetic effects based on doses of founder haplotypes, the  $QTL_{im}$  and  $(QTL_m \times \text{factor})_{im}$  terms in Equations (1) and (2) can be fit to doses of alleles of specific variants, imputed from

dense genotypes of the founder strains. Although variant association mapping sacrifices some of the information present in the founder haplotypes, it does enable researchers to potentially identify specific variants of interests and prioritize candidate genes.

Finally, the QTLViewer can perform mediation analysis for additive QTL to identify candidate causal mediators (Chick et al.

2016; Keller *et al.* 2018; Keele *et al.* 2021) if -omic data, such as gene expression, have been collected on the same mice or strains. Briefly, Equation (1) is reused (with the kinship term excluded for computational efficiency) and the QTL of interest retested, but now conditioning one-by-one on candidate mediators. A strong candidate mediator will localize near the QTL and strongly reduce the QTL LOD score.

## Results and discussion

The QTLViewer webtool can be accessed at <https://qtlviewer.jax.org/>. After clicking “Datasets” on the top right of the screen, users will find a list of available datasets from CC and DO mice with their corresponding references and links to the specific QTLViewer instances. Here, we describe how a user would get started in analyzing a publicly available CC or DO dataset using a QTLViewer.

### Step 1: Selecting a dataset

To start exploring QTLViewer data, users should click on the QTLViewer link corresponding to the project of interest. All QTLViewer projects contain at least 1 dataset such as a table of physiological traits, gene expression, or protein abundance data. Using the JAX Center for Aging Research DO data from the heart as an example (<https://churchilllab.jax.org/qtlviewer/JAC/DOHeart> last accessed 06/13/2022; Gerdes Gyuricza *et al.* 2022), we find transcript and protein data available. Users can switch datasets by clicking the selection box arrow next to the text “Current Data Set” (Fig. 2a). All datasets in a QTLViewer instance are linked to a common set of mice and genotypes.

### Step 2: Searching the dataset by key word

QTL mapping and additional analyses can be performed for any trait in the data. To conveniently specify traits of interests, users can search via key word in the *Search* text box and press *Enter*. The search algorithm is specialized for -omics data like gene expression, recognizing gene symbols, gene names, and Ensembl identifiers. All elements that match the search criteria will be displayed in a table below, but only elements that are in the dataset will be displayed in blue and clickable. For example, the genes *Ace* and *Ace2* are available in the transcriptome heart data, but not *Ace3* (Fig. 2b). This feature takes advantage of the Ensembl database and webservice, which currently hosts human and mouse gene information with backward compatible versioning through Ensembl (Yates *et al.* 2020).

### Step 3: Visualizing large-scale -omic QTLs (e.g. expression QTLs)

The QTL mapping features of the QTLViewer are flexible and tailored to different types of data. For -omics data, such as transcripts and proteins, an *LOD Peaks* tab is available (Fig. 2c), which shows the position of the QTL vs the genomic coordinate of the transcript or protein. When selecting this option, a genomic grid is displayed with LOD scores from each gene or protein and instead of visualizing single transcripts/proteins, users can visualize all the -omic QTLs at the same time. This is the easiest way to look for regions of the genome where many QTLs comap (genomic hotspots). Genomic hotspots are an indication that distal genetic variation regulates multiple genes/proteins that are potentially involved in common biological functions. The grid can be filtered based on an LOD threshold via a slider or manually setting the threshold in the *LOD Threshold* box. Factor-QTL interaction (e.g. sex-interactive QTL) LOD peaks can be viewed by

changing the *Plot type* select box. Additional information on specific QTLs can be interactively accessed by hovering and clicking on the points in the grid. The plot supports pan and zoom features.

### Step 4: Generating genome-wide LOD plots (i.e. genome scans) for individual traits

To visualize a genome-wide LOD plot for a specific trait, users should click on a point from the *LOD Peaks* grid or select a trait in the text search. Genome-wide LOD plots are useful to visualize peaks on the genome associated with the phenotype of interest. For example, the LOD plot for the gene *Sfi1* reveals a clear expression QTL (eQTL) on chromosome 13 at 65 Mb (Fig. 3a), which is considered a distal eQTL because *Sfi1* is located on chromosome 11. Information about each LOD score may be accessed by hovering. The *Plots* select box will switch between additive and factor-interactive QTL models. Clicking on a locus of interest (i.e. the peak LOD score) is used to drive additional analyses on the locus, which are visualized as plots in the lower panel. These analyses include estimation of founder allele effects, mediation of the locus effect through -omic traits, and SNP association mapping in the locus region.

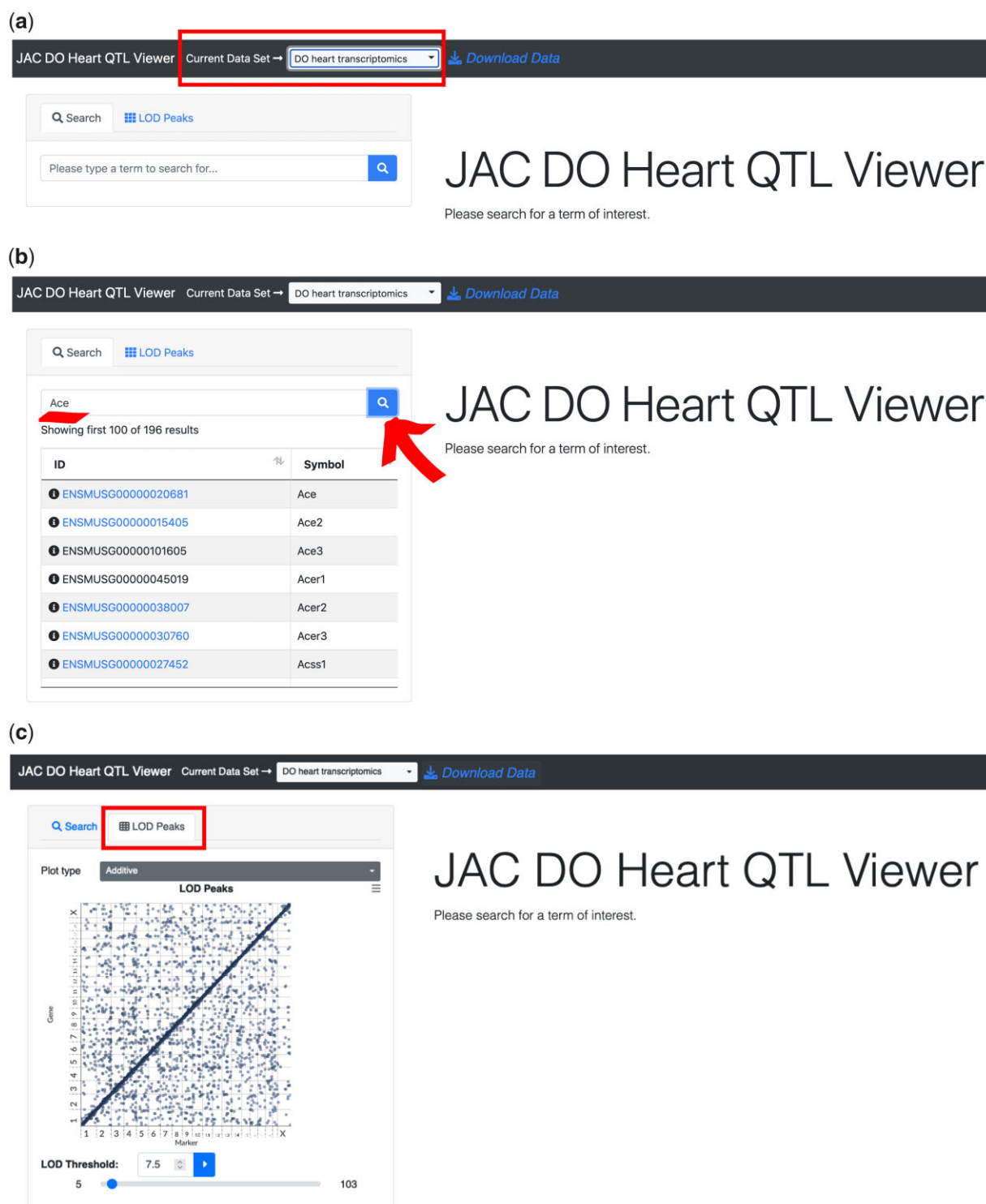
### Step 5: Generating founder allele effects plot

The estimation of founder allele effects is used to obtain the contributions of each founder haplotype to the QTL. To perform this analysis, users should click on a specific peak on the genome-wide LOD plot for a trait and, by default, the allele effects estimation will be the first analysis to be performed (Fig. 3b). The allele effects plot shows the estimated founder allele effects across the chromosome that contains the selected peak. The allele effects can be reported as fixed effects coefficients (default) or as constrained BLUPs by checking the bar on top of the effect plot. BLUP estimates are generally preferable because they shrink extreme effects from rarely observed alleles, but they are computationally more intensive to calculate. These effects can be used to distinguish which founder strains likely possess the causal genetic variants. They also highlight QTL that is multiallelic (Crouse *et al.* 2020)—a unique feature of MPPs. The allele effects of the distal eQTL on chromosome 13 for the gene *Sfi1* show a biallelic pattern driven by lower expression from the CAST/EiJ and C57BL/6J alleles compared to higher expression from the other alleles (Fig. 3b).

### Step 6: Performing mediation of QTLs through -omic traits

Mediation analysis can be used to identify candidate mediators of a QTL and thus reveals relationships between coregulated traits. For example, genetic variation local to a gene encoding a transcription factor can alter its expression levels, which then mediate changes in expression on its downstream targets. Another example of coregulated traits is a gene that possesses comapping eQTL for its transcript and pQTL for its protein, suggesting that its protein levels are transcriptionally regulated. Support for these relationships in the data can be assessed through mediation analysis. To generate mediation plots users should click on the *Mediation* tab adjacent to the *Effect* tab, and then click back on the locus of interest on the LOD plot (Fig. 4a). Doing so runs a mediation analysis which involves retesting a QTL effect at the locus of interest, iteratively conditioning on candidate mediators. Promising candidates will significantly reduce or drop the initial QTL LOD score and be encoded on the genome near the QTL. When performing a

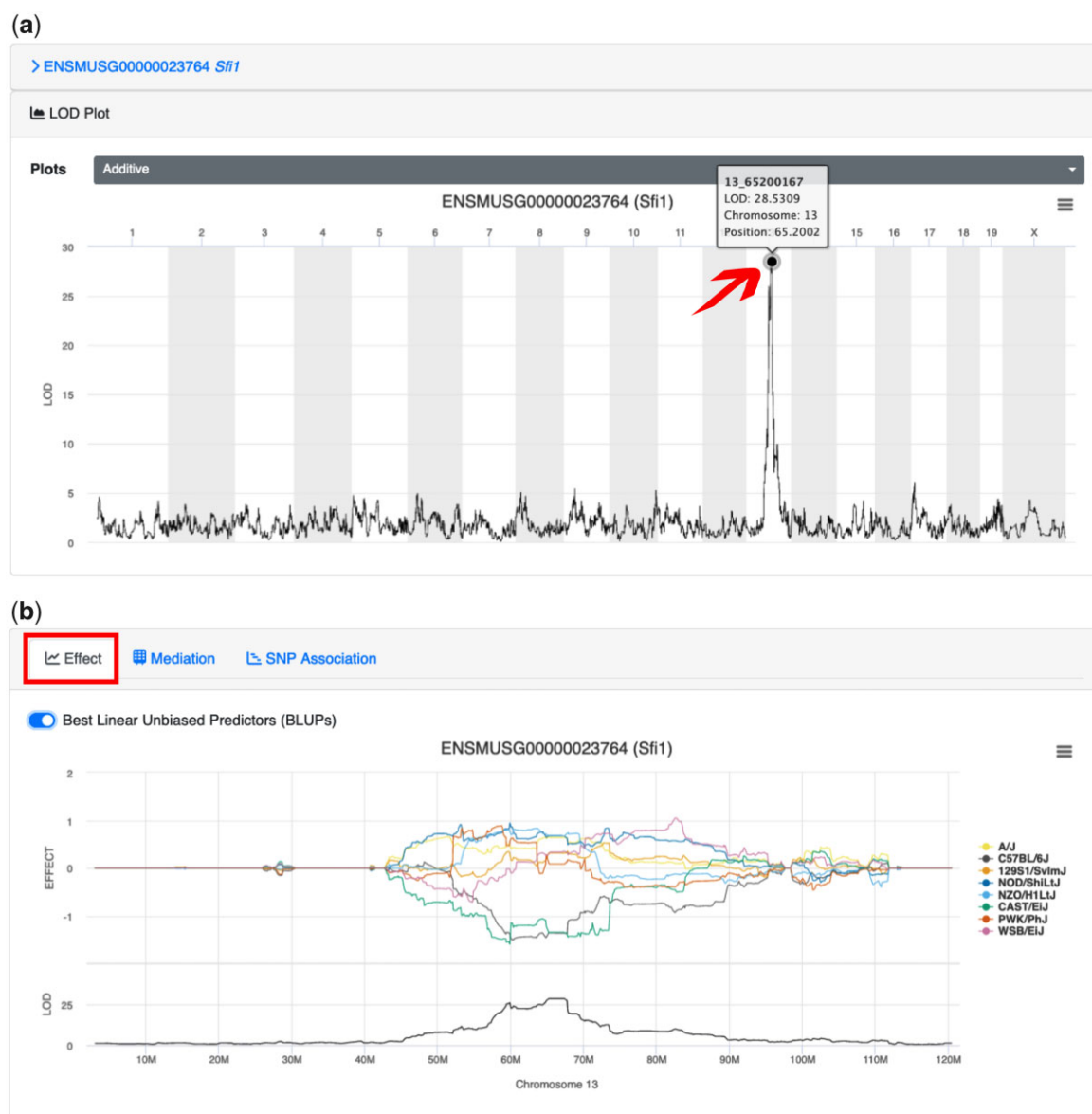




**Fig. 2.** Navigating datasets in the QTLViewer. The QTLViewer page for a project provides access to multiple datasets which can be explored by changing the option on “Current Data Set.” In the screenshot, the aging DO heart transcriptomics data is selected (a). Within this specific dataset, it is possible to search for specific traits. For example, in the transcriptome dataset, the genes *Ace* and *Ace2* are available, but not *Ace3* (b). Switching to the “Lod Peaks” mode, visualizes a genome-wide transcriptome map with marker IDs on the x-axis and gene position on the y-axis, which is a common plot type to summarize all the eQTLs in the data according to a specified LOD threshold (c).

mediation analysis of the gene *Sfi1* against the transcriptome heart data, we see an LOD drop on chromosome 13 (where the distal eQTL is located; Fig. 4a). The gene with the lowest LOD score on chromosome 13 is *Rsl1*, the candidate mediator. We also see an LOD drop on chromosome 11 at the position of the *Sfi1* gene (which corresponds to a local eQTL; Fig. 4a). This

“LOD-drop” method of mediation is a powerful tool to identify candidate mediators, but it is also susceptible to false-positive detection of linked independent effects (Chick et al. 2016; Crouse et al. 2020). Mediation can be performed against different datasets by changing the selection in the Mediate Against box.



**Fig. 3.** Genome-wide LOD and founder allele effects plots. Searching for specific traits or clicking on interesting QTLs on the transcriptome map reveals the genome-wide LOD plot with genome position across chromosomes on the x-axis and LOD scores on the y-axis (a). This plot reveals a strong eQTL on chromosome 13 at 65 Mb for the gene *Sfi1*, which is a distal eQTL because *Sfi1* is encoded on chromosome 11. By clicking on the “Effect” function and then clicking back on the locus of interest produces a founder allele effects plot at the locus (b). The allele effects plot has genomic position in Mb on the x-axis and the estimated allele effects (top) and LOD scores (bottom) on the y-axis. The peak on chromosome 13 for the gene *Sfi1* is driven by lower expression from the CAST/EiJ and C57BL/6J alleles compared to higher expression from the other 6 alleles.

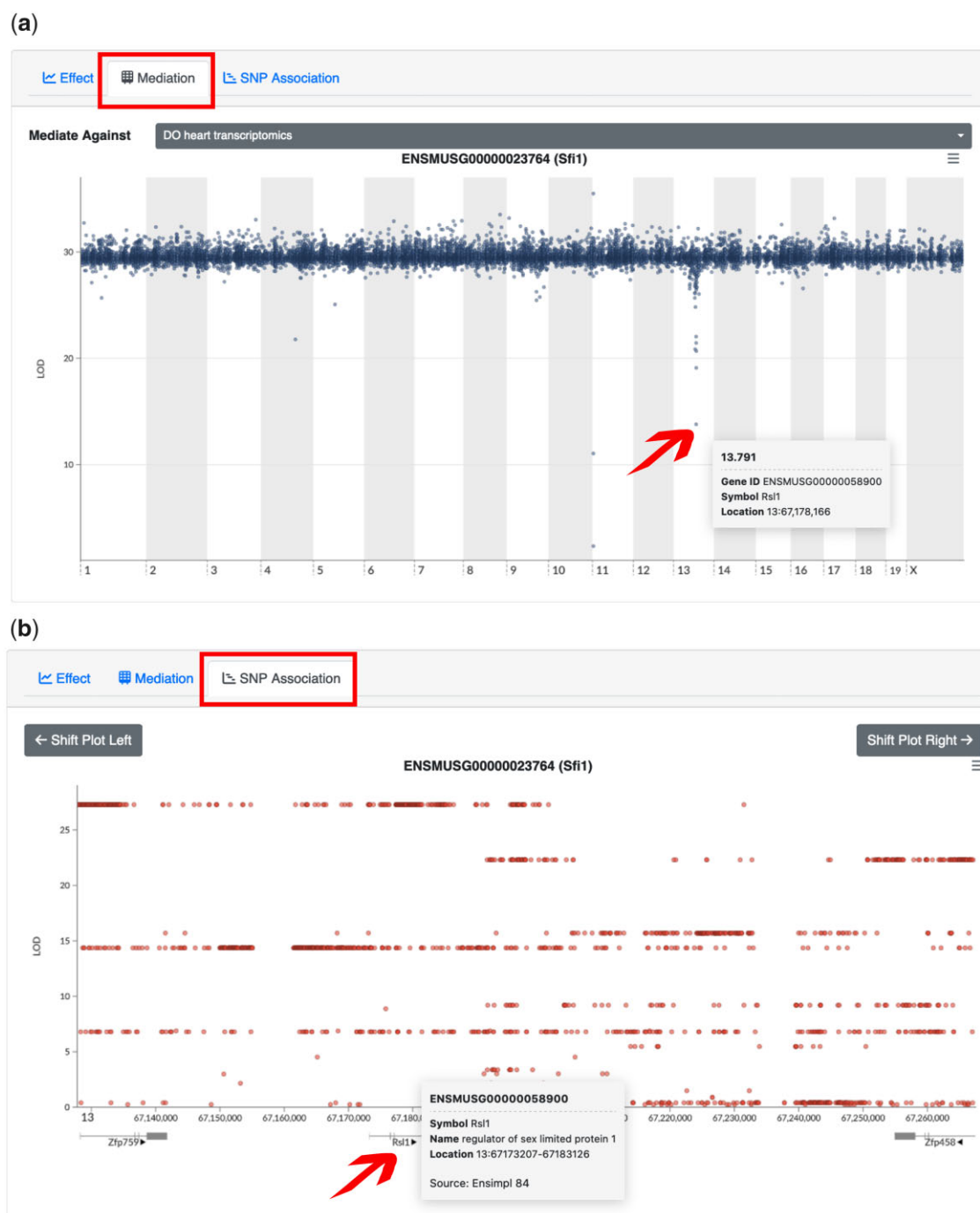
### Step 7: Performing SNP association scans

SNP (or variant) association mapping is useful to identify genetic variants of interest in the QTL region. To perform this analysis, users should click the SNP Association tab adjacent to the Mediation tab, and then click on a locus of interest on the genome-wide LOD plot (Fig. 4b). This will generate an SNP association plot, which shows LOD scores against the genomic position for all variants overlaid with annotated genes by their corresponding genomic positions on the bottom. The plot of the chromosome 13 shows multiple variants in strong linkage disequilibrium (LD) with each other (haplotype) and with high LOD scores at the position of the candidate mediator *Rsl1* (Fig. 4b). Users can pan and zoom in the SNP association plots. Hovering over a variant will display additional information such as variant consequences, and the allele distribution

pattern among the founder strains, which shows the magnitude and direction of the effect of each founder’s haplotypes on the trait.

### Step 8: Exploring the relationships among traits and covariates

In addition to QTL mapping results, the QTLViewer can be used to explore the relationship between a trait and a covariate of interests (e.g. sex). If users are interested in this type of analysis, they should scroll down to the bottom left corner of the screen, under the Profile Plot tab (Fig. 5a). The profile plot is generated automatically after searching or clicking in a trait of interest. If users wish to modify the plots based upon the covariates observed in the experiment, they should click on the Select your factors selection box, which will turn factors on or off. In addition,



**Fig. 4.** Mediation and SNP association plots. a) Mediation analysis can be performed on QTL of interest to identify candidate mediators as long as the QTL's trait and the mediators are observed for the same mice (for the DO specifically) or for the same strains. The mediation plot shows genomic position across chromosomes on the x-axis and conditional LOD scores on the y-axis. By mediating the distant eQTL of Sfi1 on chromosome 13 through the DO heart transcriptomics data, the gene Rsl1 on chromosome 13 at 67 Mb is identified as a candidate mediator, which matches the eQTL position. We also see candidate mediators on chromosome 11. The one with the lowest LOD score is the gene Sfi1 itself and the other is a gene model (*Gm11400*) that is likely on LD with Sfi1. b) An SNP association scan can be performed in the QTL region by clicking on the "SNP Association" button. The SNP association plot shows genomic position in bp on the x-axis and LOD scores on the y-axis. Annotated genes are overlaid below the x-axis. Variant association in the CC and DO reveals haplotypes shelves of variants in strong LD with each other. Additional information on variants or genes can be accessed by hovering the cursor over the dot or gene track.

users can assess the relationships among traits based on correlation. This analysis can be useful when, for example, checking the correlation between the trait of interest and its genetic mediator identified in the mediation analysis (Step 6). *Correlation* tab adjacent to the *Profile Plot* tab will show the correlation of the element of interest with any other element in the data (Fig. 5b). The tool

will display a scatter plot of the focus trait with another selected trait. To do that, users should choose a dataset in the *Select Correlation Dataset* box, and then select the element of interest from the table (Fig. 5c). The values of the elements can be adjusted by the covariates specified on the *Covariate Adjustment* box, and points can be colored by specific covariates determined



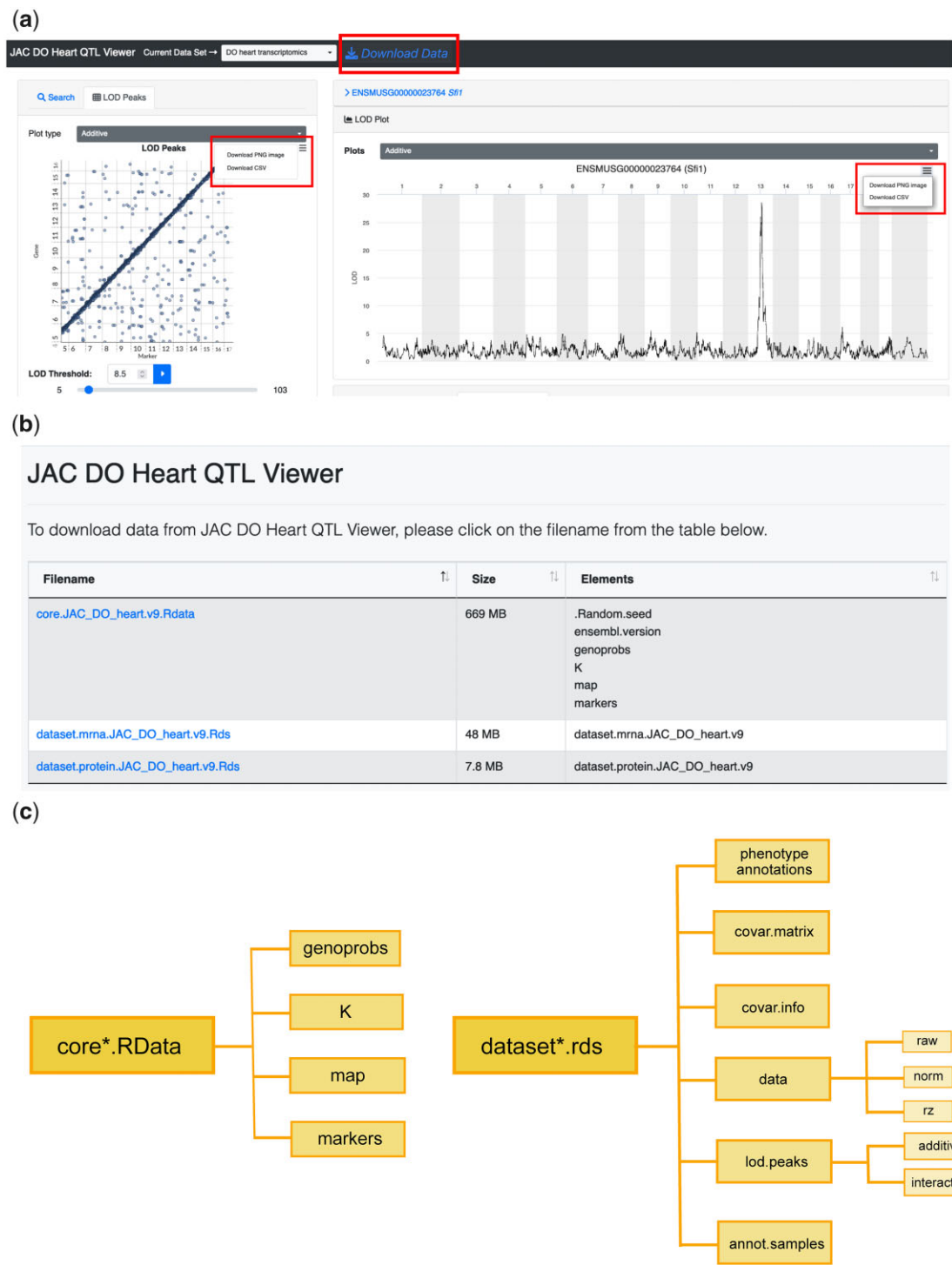
**Fig. 5.** Plots for relating traits and covariates. After searching for a specific trait or selecting one based on its QTL, QTLViewer can visualize the trait's profile according to covariates in the data with "Profile Plot." After clicking on the point corresponding to the gene *Sfi1* on the LOD peaks plots, QTLViewer outputs the normalized expression of the *Sfi1* (y-axis) categorized by sex (x-axis) as boxplots (a). Additionally, QTLViewer can display the correlation of a trait of interest with all the other elements of the data using the "Correlation" tab (b). The correlation data can also be downloaded locally by the user. All the elements on the correlation table are clickable. Clicking on "*Rsl1*" will generate a scatter plot between this gene and *Sfi1*, which illustrates the negative correlation between these 2 genes (c). *Sfi1* is negatively correlated with its mediator *Rsl1*, suggesting that *Rsl1* expression inhibits *Sfi1* expression.

on *Select a series to color*. We can see that *Sfi1* has a strong positive correlation with the gene *Gm114400* located on chromosome 11, which is likely in LD with *Sfi1* (Fig. 5c). The gene *Sfi1* also has a negative correlation with the candidate mediator *Rsl1* (Fig. 5c). The negative correlation between *Sfi1* and *Rsl1* makes sense given that *Rsl1* encodes a zinc-finger protein that functions as a negative regulator of gene expression (Krebs et al. 2003, 2014).

### Step 9: Downloading QTLViewer data objects

The QTLViewer contains multiple R data objects that supply all the necessary inputs for the analyses, including traits, covariates, and genotype probabilities. In addition to exploring the data interactively on the QTLViewer webpage, users can download the corresponding R data objects by clicking on *Download Data* on the top of the page (Fig. 6a). All plots and analyses generated above can be downloaded as figures or data tables by clicking on the top





**Fig. 6.** Figures and data download from QTLViewer. All plots generated by QTLViewer can be downloaded by clicking on the top right button in the application window (a). In addition to the figures, the processed data used as input to all analyses can be downloaded as R data files by clicking on “Download Data” (a). When using this option, users will be directed to a different webpage listing different components of the data for download (b). This includes a core RData object containing all information necessary for mapping, such as genotype probabilities, kinship matrix, and genomic map, and RDS files containing trait information, such as transcriptome (“dataset.mrna”) and proteome (“dataset.protein”) data. These RDS files are nested lists containing phenotype annotations, a matrix of covariates used for the QTL mapping (“covar.matrix”), information about the covariates (“covar.info”), trait data matrices, QTL mapping results (“lod.peaks”), and sample annotations (“annot.samples”) (c). With gene expression data, “dataset.mrna” is a list with different forms of data as matrices, including the raw counts (raw) the normalized data (norm), and inverse normal transformed data (rz) (c). The QTL mapping results “lod.peaks” is a list with QTL result tables from standard additive scans and potentially factor-interactive QTL scans (c).

right corner button on each plot (Fig. 6a). When clicking on Download Data, users will be redirected to a new page displaying all the downloadable RData files (Fig. 6b). There, users will find a

“core” RData file containing all the input needed for mapping, such as genotype probabilities and marker information. Users can also download the “dataset” remote desktop services (RDS)

files that contain the trait data, sample and assay metadata annotations, and a summary of the QTL mapping results (Fig. 6c). This functionality enables users to quickly gain access to processed data files to run further analyses.

## Building your own QTLViewer

The main goal of the QTLViewer is to allow users to interactively perform QTL mapping on publicly available data and to graphically explore results independent of their computational skills. However, if users wish to analyze their own data it is also possible to build a new QTLViewer object, which will require some coding experience. Instructions on how to set up a new QTLViewer can be found in the [Supplementary File](#).

## Future directions

The R/qtl2 software at the core of QTLViewer is very general and can accommodate a wide range of cross designs and data from model organisms other than the mouse. We plan to extend the QTLViewer to include mouse backcrosses, intercrosses, and 2-way recombinant inbred panels. Adapting QTLViewer to MPPs from species other than the mouse will require modification of settings in the configuration files and the gene-search features for -omics data, and creation of a variant database.

We are in the process of adding a new method for mediation analysis based on Bayesian model selection (Crouse et al. 2022) that can provide richer inference for mediators of interest. We continue to expand the QTLViewer for new types of genomic data, such as chromatin profiling data, which will be incorporated and layered onto the genome browser tracks.

We continue to add new datasets to the QTLViewer website, which already represents a powerful resource for the community. We welcome contributions from outside investigators and will assist in the process of preparing their data for the QTLViewer.

## Web resources

Users can access the QTLViewer webpage at <https://qtlviewer.jax.org/> (last accessed 06/13/2022). The QTLViewer software is version controlled and available from GitHub (<https://github.com/churchill-lab>; last accessed 06/13/2022).

The supporting tools Ensimpl and SNPDB are available online at <https://churchilllab.jax.org/ensimpl> (last accessed 06/13/2022) and <https://churchilllab.jax.org/foundersnps> (last accessed 06/13/2022), respectively. The Docker containers code can be found at <https://github.com/churchill-lab/qtl2web> (last accessed 06/13/2022), <https://github.com/churchill-lab/qtl2rest> (last accessed 06/13/2022), and <https://github.com/churchill-lab/ensimpl> (last accessed 06/13/2022). The Docker images are available at <https://hub.docker.com/r/churchilllab/qtl2web/tags> (last accessed 06/13/2022), <https://hub.docker.com/r/churchilllab/qtl2rest/tags> (last accessed 06/13/2022), and <https://hub.docker.com/r/churchilllab/ensimpl/tags> (last accessed 06/13/2022). The source code for QTL mapping analysis through R/qtl2 package can be found at <https://github.com/rqtl/qtl2> (last accessed 06/13/2022).

## Data availability

The aging heart data analyzed here to demonstrate the QTLViewer functions can be found online at <https://churchilllab.jax.org/qtlviewer/JAC/DOHeart> (last accessed 06/13/2022) and at figshare under accession number <https://doi.org/10.6084/m9.figshare.12378077>.

[Supplemental material](#) is available at G3 online.

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## Conflicts of interest

None declared.

## Literature cited

- Abu Toamih Atamni H, Nashef A, Iraqi FA. The Collaborative Cross mouse model for dissecting genetic susceptibility to infectious diseases. *Mamm Genome*. 2018;29(7–8):471–487.
- Al-Barghouthi BM, Mesner LD, Calabrese GM, Brooks D, Tommasini SM, Bouxsein ML, Horowitz MC, Rosen CJ, Nguyen K, Haddox S, et al. Systems genetics in diversity outbred mice inform BMD GWAS and identify determinants of bone strength. *Nat Commun*. 2021;12(1):3408.
- Aylor DL, Valdar W, Foulds-Mathes W, Buus RJ, Verdugo RA, Baric RS, Ferris MT, Frelinger JA, Heise M, Frieman MB, et al. Genetic analysis of complex traits in the emerging Collaborative Cross. *Genome Res*. 2011;21(8):1213–1222.
- Bostock M, Ogievetsky V, Heer J. D3 data-driven documents. *IEEE Trans Vis Comput Graph*. 2011;17(12):2301–2309.
- Broman KW, Gatti DM, Simecek P, Furlotte NA, Prins P, Sen S, Yandell BS, Churchill GA. R/qtl2: software for mapping quantitative trait loci with high-dimensional data and multiparent populations. *Genetics*. 2019;211(2):495–502.
- Broman KW, Sen S, Owens SE, Manichaikul A, Southard-Smith EM, Churchill GA. The X chromosome in quantitative trait locus mapping. *Genetics*. 2006;174(4):2151–2158.
- Broman KW. R/qtlcharts: interactive graphics for quantitative trait locus mapping. *Genetics*. 2015;199(2):359–361.
- Chick JM, Munger SC, Simecek P, Huttlin EL, Choi K, Daniel M. Defining the consequences of genetic variation on a proteome-wide scale. *Nature*. 2016;534(7608):500–5.
- Churchill GA, Airey DC, Allayee H, Angel JM, Attie AD, Beatty J, Beavis WD, Belknap JK, Bennett B, Berrettini W, et al.; Complex Trait Consortium. The Collaborative Cross, a community resource for the genetic analysis of complex traits. *Nat Genet*. 2004;36(11):1133–37.
- Churchill GA, Gatti DM, Munger SC, Svenson KL. The Diversity Outbred mouse population. *Mamm Genome*. 2012;23(9–10):713–718.
- Crouse WL, Keele GR, Gastonguay MS, Churchill GA, Valdar W. A Bayesian model selection approach to mediation analysis. *PLoS Genet*. 2022;18(5):e1010184.
- Crouse WL, Kelada SNP, Valdar W. Inferring the allelic series at QTL in multiparental populations. *Genetics*. 2020;216(4):957–983.
- de Koning D-J, McIntyre LM. Back to the future: multiparent populations provide the key to unlocking the genetic basis of complex traits. *Genetics*. 2017;7(6):1617–1618.
- French JE, Gatti DM, Morgan DL, Kissling GE, Shockley KR, Knudsen GA, Shepard KG, Price HC, King D, Witt KL, et al. Diversity outbred mice identify population-based exposure thresholds and genetic

- factors that influence benzene-induced genotoxicity. *Environ Health Perspect.* 2015;123(3):237–45.
- Gatti DM, Svenson KL, Shabalin A, Wu LY, Valdar WW, Simecek P, Goodwin N, Cheng R, Pomp D, Palmer A. Quantitative trait locus mapping methods for diversity outbred mice. *G3 (Bethesda)*. 2014;4(9):1623–1633.
- Gerdes Gyuricza I, Chick JM, Keele GR, Deighan AG, Munger SC, Korstanje RC, Gygi SP, Churchill GA. Genome-wide transcript and protein analysis highlights the role of protein homeostasis in the aging mouse heart. *Genome Res.* 2022;32(5):838–852.
- Hsiao K, Noble C, Pitman W, Yadav N, Kumar S, Keele GR, Terceros A, Kanke M, Conniff T, Cheleuitte-Nieves C, et al. A thalamic orphan receptor drives variability in short-term memory. *Cell*. 2020;183(2):522–536.e19.
- Iraqi FA, Mahajne M, Salaymah Y, Sandovski H, Tayem H, Vered K, Balmer L, Hall M, Manship G, Morahan G, et al. The genome architecture of the Collaborative Cross mouse genetic reference population. *Genetics*. 2012;190(2):389–401.
- Keele GR, Quach BC, Israel JW, Chappell GA, Lewis L, Safi A, Simon JM, Cotney P, Crawford GE, Valdar W, et al. Integrative QTL analysis of gene expression and chromatin accessibility identifies multi-tissue patterns of genetic regulation. *PLoS Genet.* 2020;16(1):e1008537.
- Keele GR, Zhang T, Pham DT, Vincent M, Bell TA, Hock P, Shaw GD, Paulo JA, Munger SC, Pardo-Manuel de Villena F, et al. Regulation of protein abundance in genetically diverse mouse populations. *Cell Genomics*. 2021;1(1):100003.
- Keller MP, Gatti DM, Schueler KL, Rabaglia ME, Stapleton DS, Simecek P, Vincent M, Allen S, Broman AT, Bacher R, et al. Genetic drivers of pancreatic islet function. *Genetics*. 2018;209(1):335–356.
- Konganti K, Ehrlich A, Ivan R, Threadgill DW. gQTL: a web application for QTL analysis using the Collaborative Cross mouse genetic reference population. *G3 (Bethesda)*. 2018;8:2559–2562.
- Krebs CJ, Larkins LK, Price R, Tullis KM, Miller RD, Robins DM. Regulator of sex-limitation (Rsl) encodes a pair of KRAB zinc-finger genes that control sexually dimorphic liver gene expression. *Genes Dev.* 2003;17(21):2664–2674.
- Krebs CJ, Zhang D, Yin L, Robins DM. The KRAB zinc finger protein RSL1 modulates sex-biased gene expression in liver and adipose tissue to maintain metabolic homeostasis. *Mol Cell Biol.* 2014;34(2):221–232.
- R Core Team. R: a language and environment for statistical computing. 2021 [accessed 2021 Dec 22]. <https://www.r-project.org/>.
- Solberg Woods LC. QTL mapping in outbred populations: successes and challenges. *Physiol Genomics*. 2014;46(3):81–90.
- Svenson KL, Gatti DM, Valdar W, Welsh CE, Cheng R, Chesler EJ, Palmer AA, McMillan L, Churchill GA. High-resolution genetic mapping using the mouse Diversity Outbred population. *Genetics*. 2012;190(2):437–447.
- Takemon Y, Chick JM, Gerdes Gyuricza I, Skelly DA, Devuyst O, Gygi SP, Churchill GA, Korstanje R. Proteomic and transcriptomic profiling reveal different aspects of aging in the kidney. *eLife*. 2021;10:e62585.
- Wei J, Xu S. A random-model approach to QTL Mapping in Multiparent Advanced Generation Intercross (MAGIC) populations. *Genetics*. 2016;202(2):471–486.
- Yang H, Bell TA, Churchill GA, Pardo-Manuel De Villena F. On the subspecific origin of the laboratory mouse. *Nat Genet.* 2007;39(9):1100–1107.
- Yang H, Wang JR, Didion JP, Buus RJ, Bell TA, Welsh CE, Bonhomme F, Yu AH-T, Nachman MW, Pialek J, et al. Subspecific origin and haplotype diversity in the laboratory mouse. *Nat Genet.* 2011;43(7):648–655.
- Yates AD, Achuthan P, Akanni W, Allen J, Allen J, Alvarez-Jarreta J, Amode MR, Armean IM, Azov AG, Bennett R, et al. Ensembl 2020. *Nucleic Acids Res.* 2020;48(D1):D682–D688.

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