Bioinformatics tools and database resources for systems genetics analysis in mice—a short review and an evaluation of future needs

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Submitted: 28th February 2011; Received (in revised form): 11th April 2011

Abstract

During a meeting of the SYSGENET working group 'Bioinformatics,' currently available software tools and databases for systems genetics in mice were reviewed and the needs for future developments discussed. The group evaluated interoperability and performed initial feasibility studies. To aid future compatibility of software and exchange of already developed software modules, a strong recommendation was made by the group to integrate HAPPY and R/qtl analysis toolboxes, GeneNetwork and XGAP database platforms, and TIQS and xQTL processing platforms. R should be used as the principal computer language for QTL data analysis in all platforms and a 'cloud' should be used for software dissemination to the community. Furthermore, the working group recommended that all data models and software source code should be made visible in public repositories to allow a coordinated effort on the use of common data structures and file formats.

Keywords: QTL mapping; database; mouse; systems genetics

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INTRODUCTION
The study of complex traits in genetic reference populations (GRP) of mice and rats enables the unravelling of molecular interaction networks and the gain of insights into the underlying biological mechanisms. This knowledge has also considerably contributed to a better understanding of the genetic basis of common diseases in humans [1–9].

A wide variety of bioinformatics resources and tools exist which are available for researchers to dissect complex traits through the analysis of genotype–phenotype linkage/associations. However, different institutions and research groups developed these independently and some of the resources are not widely known, several are partially redundant and most of them are not easily interoperable.

In a recent workshop in Groningen, Netherlands, the Bioinformatics Working Group (WG) of the SYSGENET consortium [10] addressed some of these IT issues. SYSGENET represents a network of scientists in Europe who use the mouse as an experimental model system to identify complex genetic factors influencing phenotypic traits that are relevant for human diseases. The Bioinformatics WG is focusing on reviewing the availability and interoperability of existing tools and resources for complex trait analyses and to make suggestions for future developments and needs. Here, we report about the results of the Groningen workshop.

RESULTS AND DISCUSSION
We examined available user interfaces (front-end) and facilities (back-end) for large scale computing which will be needed to analyse the next generation of mouse genetics experiments (Figure 1). We also explored the tools available for analysis, databases, software dissemination and code sharing. For each, we investigated needs for interoperability and made recommendations for the next steps.

HAPPY and R/qtl—analysis tools
Mouse genetics researchers rely on the availability of suitable statistical tools that accommodate the wide range of experimental designs now available. However, many resources and tools are constantly being developed by research groups (http://www.stat.wisc.edu/~yandell/statgen/software/). But it would be very desirably if these tools could be combined. Therefore, the WG explored what is required to enable such integration by examining two tools: HAPPY and R/qtl.

HAPPY [11] is designed to analyse complex crosses of model organisms from multiple founders, e.g. mouse heterogeneous stocks (HS) or the Collaborative Cross [12–21]. It includes the ability to infer founder ancestry between genotyped markers and functions for single QTL mapping (including a Bayesian method) and two-locus epistasis (genetic interactions).

R/qtl [22, 23] is a versatile, stand-alone software package with a large number of functions and plotting tools for a wide range of analyses for standard model organism crosses, e.g. F2, backcrosses and intercrosses, and an increasing functionality for the more complex crosses analysed by HAPPY. These QTL mapping packages are both freely available and largely complement each other in functionality.
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including genetic markers, genotype microarrays,
and Genome Wide Association Studies (GWAS),
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builds on the MOLGENIS software toolkit [25]
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allowing stored data to be extracted from XGAP,
for annotation. It is highly integrated with R/qtl,
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used for Genome Wide Linkage Studies (GWLS)
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including genetic markers, genotype microarrays,
gene expression microarrays, Nuclear Magnetic
Resonance and mass spectrometry, in both outbred
populations (including humans), inbred model or-
ganisms (including mouse, rat, Arabidopsis and
Caenorhabditis elegans) and commercial crops (e.g. brassica). XGAP is actively maintained and used in vari-
ous EU projects and is responsive to requests from
users for additional functionalities.

GeneNetwork (GN, www.genenetwork.org) is
an open-source web service for systems genetics
that stores, analyses and displays classic phenotypes
as well as large gene expression data sets with
matched genomic data for several species (human,
mouse, rat, fly, barley and Arabidopsis). GN exploits
the familiar Linux/Apache/Python/MySQL archi-
tecture (www.genenetwork.org). GN is intended
primarily for biologist researchers and does not re-
quire training in bioinformatics. All genotype data
are built into the system and calculations and displays
are entirely via the web. GN includes extensive
multivariate statistical tools from R and custom
QTL mapping modules (QTL Reaper) used for clas-
sical and expression QTL studies. GN is able to rap-
idly compute QTL maps for up to 100 traits,
simultaneously, using the Cluster Mapping function
using Haley–Knott regression code, but does not
store the full QTL vector. GN does store the peak
QTL locations and displays summarized full tran-
scriptome results using the GenomeGraph tool.
GN is also used to study the correlation structure
among large numbers of traits, and to perform data
mining in genomic regions containing candidates for
quantitative trait genes. The GN system can handle
standard mapping panels but is not yet compatible
with complex multi-generation crosses or human
linkage studies. GN is now mirrored at nine research
institutions in Europe, the USA, Australia and Asia.

To allow bioinformatics laboratories easy access to
the many data elements stored in GN and incorpo-
rate them into their R analyses, data extraction tools
and/or converters between XGAP and GN would
be of great value. In this way, GN may benefit from
the large data capabilities, programmers interfaces
and flexibility of XGAP. Vice versa, XGAP users
would benefit from GN as a way to make results
generated in XGAP easy to explore for a broader
public. Feasibility studies were successfully per-
formed at the Bioinformatics WG meeting in
Groningen. Data could be exported as text or
Excel files from both GN and XGAP. A number of
data sets from GeneNetwork, including BXD
genotypes, hippocampus expression studies, marker

XGAP and GeneNetwork
The scale of experimental data requires database in-
frastucture to track, trace and explore all experimen-
tal data and annotations involved.

XGAP [24] is flexible, open-source software for
storing and accessing large scale xQTL data, includ-
ing user interfaces to manage, search and conduct
quality control checks on data, import/export tools
to up/download data to Excel, and interfaces to the
R language to connect to statistical analysis and
graphics capabilities and interfaces to web services
for annotation. It is highly integrated with R/qtl,
allowing stored data to be extracted from XGAP,
analysed in R/qtl and results (like high-throughput
eQTL profiles) stored back into XGAP. XGAP
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data sets from GeneNetwork, including BXD
genotypes, hippocampus expression studies, marker
and probe annotations, were imported into XGAP, and via XGAP successfully transferred into R/qtl allowing whole phenome and genome analysis.

In the future, the Bioinformatics WG of SYSGENET will collaborate closely with the developers of both systems and data providers to integrate more codes, share best practices, merge systems where suitable, and develop new ideas to bring the demands of biomedical research and statistical genetics together. The WG has a particular interest to integrate back-end open source mapping code such as HAPPY and R/qtl and focus on the interpretation of concordant and conflicting findings in order to improve methods. The WG will also host a workshop between GN and XGAP/SYSGENET developers to further foster their integration. Similarly, the developers of R/qtl will be included in future activities of the WG.

**TIQS and xQTL—processing infrastructures**

The scale of analysis requires automated computational protocols to analyse all expression, proteomics and metabolomics QTLs on marker maps of ever increasing density. These should include web access tools for both experts and non-experts in sophisticated statistics analysis and high performance computing.

The interactive QTL System (TIQS) (http://eqtl.berlios.de) is a web application that guides its users through the analysis steps needed. It maximizes the distribution of computational effort (supporting traditional clusters, cloud-based systems and the grid) and presents the results in two ways. The user is guided through input forms to specify queries to the database and gets the results displayed in tabular form. That approach can also be taken by external tools to query the system if those do not access the database directly. Secondly, all major functionality and graphical displays in R/qtl are wrapped by web forms and bidirectionally linked to the tabular displays. That approach can also be taken by external tools to query the system if those do not access the database directly. Secondly, all major functionality and graphical displays in R/qtl are wrapped by web forms and bidirectionally linked to the tabular displays. The focus of current developments is on comparisons of multiple independent xQTL data sets and the adoption of the above concepts to HAPPY for the study of advanced intercross lines.

xQTL workbench is also a web application to ease large scale QTL processing that builds on XGAP (http://www.xgap.org). As per trait QTL analysis methods are perfectly tailored for running on a cluster, the xQTL workbench distributes analysis in a highly transparent way and the analysis time will be reduced by a factor of approximately the number of cluster nodes assigned to the task. At the heart of the xQTL system is a flexible model that allows bioinformaticians to perform new analyses via R scripts and input and output data sets via the web user interface. Based on this model, biologist researchers obtain an auto-generated user interface that allows the running of pre-defined analyses in a user friendly way. A job manager shows the status of the current (and past) jobs, resubmits failed individual subjobs and allows browsing of the history of previous analysis. The current version of xQTL workbench can be linked to standard compute cluster software and includes R/qtl analysis, providing various QTL mapping methods like the new Multiple QTL Mapping [23].

TIQS and xQTL workbench allow the running of analyses on whole assays of (molecular) phenotypes as a batch. This enables genetical genomics studies without waiting times. TIQS is particularly strong in using a ‘cloud’ for large scale computing while xQTL uses ‘pbs’ based traditional clusters and is more developed for data management and definition of new analyses, so the desire is to work together. Both systems use R as the back-end language for data analysis in all platforms, which will enable transfer of analysis protocols between experiments and institutes. Both xQTL and TIQS have a number of analysis tools and functions integrated into the framework and these tools are described using a standard format, or ‘model’, which includes input parameters, output parameters and scripts to actually run them. The WG recommended harmonizing the ‘models’ used presently in light not only of TIQS and xQTL workbench but also of established formats from broader toolkits like the Galaxy (http://galaxy.psu.edu/) format tool.xml and the Taverna (http://www.taverna.org.uk) format scufl.xml as first steps towards integration.

**Git and SVN—software code sharing and versioning**

Access and sharing of software code of all aforementioned tools is essential to avoid duplication of efforts, promote interoperability and to really collaborate within the mouse genetics bioinformatics community. Only then does it really become clear what is available and how there can be data and software flow between projects.

HAPPY has versions in C and R code and beta versions under development, and each change is not...
necessarily released individually as an executable because of the additional effort needed. Other bioinformaticians may want to do this additional work but can only do so if they have access to the code. Similar arguments hold for R/qtl, GN, XGAP and TIQS.

An example tool to facilitate such sharing and collaboration on code level is Git: Git allows users to follow updates (versions) of software packages and easily provide feedback. During the WG workshop therefore the most recent stable R version of HAPPY (version 2.1) was placed in Git, along with beta versions and some development code. Git will help to control versions for the authors and facilitate compatibility with R/qtl, although most users will generally find the last stable version (Git master branch) the most useful and user-friendly.

Presently, R/qtl, HAPPY and TIQS are in Git (www.github.com/kbroman/qtl, www.github.com/cdurrant/happy, http://eqtl.berlios.de). An alternative to Git is Subversion (SVN). GN code was developed using Subversion with Python as its primary language. GN’s code and a shell (partial) database are available on Source Forge (www.sourceforge.org/projects/genenetwork). The mapping and permutation code used by GN for real time analysis is based on fast Haley–Knott approximations and is available on Source Forge (http://qtlreaper.sourceforge.net/). xQTL and XGAP code were also developed in Subversion using Java as its primary language and R for its analysis interfaces. XGAP code and binaries are all available as open source at http://www.xgap.org and http://www.molgenis.org. From a developer’s perspective both Git and SVN are equally well-suited for publishing code and may be used in parallel.

The Bioinformatics WG of SYSGENET clearly recommended that all projects should be using source code management software, preferably with a web presentation. The open source spirit of mutual collaboration is of increasing importance, but also increasingly difficult with the specialization of software. The open source paradigm discussed in this article is mainly founded on scientific principles to allow local modifications of a protocol. Nonetheless, all inter-project aims of this work group outlined in this article demand free access to the source code. Those contributions, however, shall be non-interruptive to the main development, and inter-exchangeable immediately without the demand of manual involvement of the main developers.

**Debian packaging of BioLib—software dissemination**

Local installation of publicly available software tools is often challenging in the light-specific software requirements. Moreover, even if the software is successfully installed, use of different programming languages may represent a barrier for successful inter-operation between different packages.

We explored BioLib and Debian Med as methods to ease software dissemination and installation. The goal of the BioLib project is to make existing C/C++ libraries and modules, such as R/qtl and EMBASS, available to users of Java, Perl, Python and Ruby. This would greatly reduce the limitations for software interoperation and in particular makes existing libraries accessible to mouse systems genetics programmers. The Debian Med project is an effort to package biomedical software together with Debian into one complete and easy-to-install Linux system, offering a portal to help developers (http://www.debian.org/devel/debian-med/) and with a large existing user base on Debian and Ubuntu [26]. It also easily migrates to openBSD, Apple Macintosh OSX and by means of virtualization also to Windows. Moreover, Debian packaging by Debian Med volunteers allows easy distribution of software packages to large numbers of users via clusters, local servers and cloud computing which is essential in light of the processing needs described above.

Feasibility studies with simple examples such as packaging and running PAML (Phylogenetic Analysis through Maximum Likelihood) in the Cloud were successfully performed at the WG meeting. An SVN repository was created for storing packaging information. The WG recommended that software for QTL analysis should be packaged as part of DebianMed distribution. However, before we can fully benefit from BioLib, more conceptual work on packaging will be needed in the future, since BioLib is quite complex and highly integrated with other packages which render maintenance of BioLib packages difficult.

**Cloud computing—transparent access to software and hardware resources**

With increasing experimental scale, it will not be sufficient to have analysis software as simple downloads, because the researcher will also need sizable
compute and storage power [27]. Thus, researchers will need easy access to ‘software as a service’ such as ‘cloud computing’.

The term ‘cloud’ is used in many different ways. It basically means to have software available on demand, on suitable compute and storage hardware, with the user being charged for the time that it is being used. Following the concept of the leading cloud provider Amazon, it refers in practice to use of a virtual Linux machine, i.e., a virtualized compute server with Linux pre-installed, hosted within some large compute infrastructure, with complete freedom to be customized by the user although not for free [28]. Many commercial, national and local compute centres are also increasingly providing computer server capacity in cloud fashion. This makes these virtual machines an easy method to distribute software without the need for users to install software. The collaboration in SYSGENET includes the preparation of one hard drive image (virtual copy of a compute hard drive) that can be started by any collaborator as a virtual machine on the ‘cloud’, with data that can be shared between different collaborators’ instances. Thus, the infrastructure is specified jointly but for the actual computation every partner can be private with their data. This approach grants enormous computational power to everyone with minimal preparation—one that shared image is finalized.

The setup of such a compute cloud is not trivial but several initiatives are underway to ease this process. At the workshop, the Bioinformatics WG tested the packaging system provided by the Debian Med initiative (described above) as a method to create a cloud and we consider this as a seed for the image to then be publicly shared. Instructions to create such images were prepared for the workshop and made available at http://wiki.debian.org/DebianMed/LiveCD. Using this method the cloud infrastructure can be transferred to local computer clusters if desired. Every participant has access to the server and can grant access to collaborators without having to pay the hosting fees. When complete, this server image can be ported to Amazon to be reused by others.

CONCLUSIONS
Several feasibility studies were performed during the meeting of the Bioinformatics WG of the SYSGENET network and have already produced results with direct implication to the research community.

At the meeting, data were successfully transferred between HAPPY, R/qtl, GeneNetwork, TIQS, xQTL and XGAP, and subsequent analysis of GeneNetwork data were performed with R/qtl using XGAP and R/qtl ‘cross’ as exchange formats. This allows researchers to benefit from complementary features available between these tools and resources. Debian packaging of BioLib was also successfully tested for some simple examples. Significant improvements have been made or are planned to further increase the interoperability of a number of existing packages, with the final goal of providing the systems genetics community with an integrated suite of state-of-the-art programs and bioinformatics platforms to analyse complex trait data.

A series of recommendations were made:

(i) Specific data converters and tools will be required to transfer data from one package to another and convert code into different languages.

(ii) R should be used as the back-end language for data analysis in all platforms, to aid future compatibility of software and exchange of already developed analysis modules.

(iii) Source code should be made visible in systems like Git and Subversion, to enable review of the back-end code and allow a coordinated effort on the use of common data structures and file formats across packages.

(iv) Suitable packaging systems should be used to enable software sharing.

(v) Data(base) models for raw data, annotations and the reporting of computing protocols should be shared to aim for interoperability and enable sharing of ‘all information’ associated with QTL experiments.

(vi) New tool and resource developers should try to build/extend on existing data structures like those in R/qtl and XGAP file format so these new tools can be used in combination with existing solutions.

As the next step, the Bioinformatics WG of SYSGENET will bring all tools and resources described above into the cloud, to allow communication of advancements of the workflows between experts and dissemination of tools to the research community. The cloud technology also allows
OUTLOOK
In the future, further considerations should include integration of mapping analysis tools with genomic annotation and sequence data. Another goal should be the generalization of analysis frameworks to accommodate not only mouse and human data but also data from model species with different genetic pedigrees, such as pets (e.g. dogs), farm animals (e.g. horses, cattle, pigs) and plants (e.g. *brassica*, maize). While the frameworks described here have the ability to absorb new analysis methods, consideration should be given to allow high flexibility. It will be very difficult to anticipate totally new demands and requirements, or even new developments which could provide a better alternative than databases for data and R for analysis. XGAP and TiQS are frameworks that incorporate appropriate mechanisms to support new demands and developments [29].

**Key Points**
- Current software tools and databases for systems genetics in mice were reviewed.
- The interoperability between databases and tools was evaluated.
- Recommendations were made to use R as the common language for QTL analysis and to deposit software source codes in public repositories to better integrate tools and data between research groups.

Acknowledgements
We like to thank Rob Williams for critical comments and his contributions to the descriptions of GN. C.D. and M.A.S. wrote the manuscript and communicated with the other authors who contributed various parts to the final manuscript. R.C.J. and K.S. organized and developed the concepts for the workshop and wrote the manuscript.

**FUNDING**
SYSGENET is funded through the COST framework, an intergovernmental framework for European Cooperation in Science and Technology (http://www.cost.eu/about_cost) and is coordinated by Klaus Schughart (more details at: http://www.helmholtz-hzi.de/sysgenet/).

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