The Gene Expression Database for Mouse Development (GXD): Putting Developmental Expression Information at Your Fingertips

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Because molecular mechanisms of development are extraordinarily complex, the understanding of these processes requires the integration of pertinent research data. Using the Gene Expression Database for Mouse Development (GXD) as an example, we illustrate the progress made toward this goal, and discuss relevant issues that apply to developmental databases and developmental research in general. Since its first release in 1998, GXD has served the scientific community by integrating multiple types of expression data from publications and electronic submissions and by making these data freely and widely available. Focusing on endogenous gene expression in wild-type and mutant mice and covering data from RNA in situ hybridization, in situ reporter (knock-in), immunohistochemistry, reverse transcriptase-polymerase chain reaction, Northern blot, and Western blot experiments, the database has grown tremendously over the years in terms of data content and search utilities. Currently, GXD includes over 1.4 million annotated expression results and over 260,000 images. All these data and images are readily accessible to many types of database searches. Here we describe the data and search tools of GXD; explain how to use the database most effectively; discuss how we acquire, curate, and integrate developmental expression information; and describe how the research community can help in this process. Developmental Dynamics 243:1176–1186, 2014. © 2014 The Authors Developmental Dynamics published by Wiley Periodicals, Inc. on behalf of American Association of Anatomists.

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Introduction

Gene expression data provide crucial insights into the molecular mechanisms of development, differentiation, and disease. However, the data are voluminous, complex, and heterogeneous. They are generated by many different laboratories and scattered through thousands of publications. Without the help of centralized databases, it is impossible to keep abreast of all this information, let alone to access and search these data in a cohesive and integrated way.

The Gene Expression Database for Mouse Development (GXD) was one of the first databases to address these critical issues (Ringwald et al., 1994, 1999). As a mammalian model system, the mouse is heavily used for developmental research. Tissues from all developmental stages and from different mouse strains and mutants are subject to detailed expression studies. For many years, the GXD project has been curating mouse developmental expression data from the published literature, as well as acquiring data through direct submissions and collaboration with efforts that generate pertinent expression data at a large-scale. For example, GXD has incorporated the in situ hybridization data from the EurExpress (Diez-Roux et al., 2011), GenePaint (Visel et al., 2004), GUDMAP (Genitourinary Molecular Anatomy Project; Harding et al., 2011), and BGEM (Brain Gene Expression Map; Magdaleno et al., 2006) projects. GXD integrates data from all these different sources and, as a major component of the Mouse Genome Informatics (MGI) resource (www.informatics.jax.org), combines the expression information with genetic, functional, and phenotypic data. Therefore, these expression data are readily accessible to many types of database searches (Finger et al., 2011; Smith et al., 2014; Blake et al., 2014).

Here, we describe the current status of GXD with a particular emphasis on its search utilities. Furthermore, we illustrate issues of data curation and integration that apply to developmental research in general.
**Concepts, Scope, and Expression Data Content**

GXD covers all developmental stages and all organ systems and comprises expression data from wild-type and mutant mice. The main focus is on endogenous gene expression data during development. As data accumulate, GXD aims to provide increasingly complete information about which RNA and protein products are made from a given gene, where and when these products are expressed, and how their expression varies in different mouse strains and mutants. Because there is no single assay type that can provide answers to all these questions, GXD is designed as a system that can integrate different types of expression data.

At this point, GXD captures data from RNA in situ hybridization, in situ reporter (knock-in), immunohistochemistry, reverse transcriptase-polymerase chain reaction (RT-PCR), Northern blot, and Western blot experiments.

Expression patterns (i.e., the time and space of gene expression) are described in a standardized way by using an extensive anatomical ontology that has been developed in collaboration with the eMouseAtlas (EMAP) project (Bard et al., 1998; Hayamizu et al., 2013). The ontology is structured hierarchically, allowing the integrated description of expression patterns from experiments with differing spatial resolution, as well as enabling searches that include anatomical structures and their substructures (described in detail below).
As illustrated in Figure 1, each database record describes the results obtained for each specimen, including the level and pattern of expression for each anatomical structure examined, as well as the molecular probe and the experimental conditions used. Images of the original expression data accompany the annotations whenever possible. By capturing these elemental data, different types of expression data can be represented and integrated in a robust manner. Genes and mutant alleles are recorded using official nomenclature, and all data are associated with a reference. Genes, probes, alleles, anatomical structures, and references are key points of data integration that tie genetic, genomic, expression, functional, and phenotypic data closely together, enabling search capabilities unique to GXD/MGI.

GXD currently includes detailed expression records for >13,800 genes. The data come from >67,000 expression assays and include >1.4 million expression result annotations and >260,000 images of primary expression data. Eighty-two percent of the data are from RNA in situ hybridization studies and 10% from RT-PCR experiments, reflecting the detailed spatial resolution and sensitivity required in developmental expression studies. In addition to data from different strains of wild-type mice, GXD currently includes expression data from >1,900 mouse mutants.
In the following we illustrate different ways to effectively search these data.

**Searching for Expression Data**

**The GXD Home Page**

One way to access the expression data for a given gene is to use the Quick Search, accessible from all MGI pages to find the corresponding gene detail page and then follow the links in the “Expression” section (see Fig. 2). However, GXD’s search tools allow you to do much more than look up expression data gene by gene. To take full advantage of GXD, we recommend the GXD Home Page as a starting point: http://www.informatics.jax.org/expression.shtml.

This page gives access to GXD’s search forms and provides more information about GXD, including user help, news announcements, and instructions for submitting data to GXD. We now describe the use of three of GXD’s search forms in more detail.

**Gene Expression Data Query**

The Gene Expression Data Query form (http://www.informatics.jax.org/gxd) is the most versatile and powerful search form for detailed expression data in GXD. It offers two different query utilities by means of the Standard Search and Differential Expression Search tabs.

The Standard Search tab (shown in Fig. 3) allows investigators to ask both broad and very specific questions pertinent to their...
research interests and quickly obtain matching expression results. Researchers can find expression data for specific genes, or for sets of genes as defined by their biological function or by association with annotated mouse phenotypes or human diseases. They can search for expression data for specific anatomical structures and/or specific developmental stages. Searches can be limited to expression data from wild-type mice or to expression data from mice that have been mutated in specific genes. Assay type(s) can also be chosen. Furthermore, using combinations of the query parameters described above makes it possible to formulate complex queries. For example, one could search for genes associated with DiGeorge Syndrome expressed in heart; for genes involved in left/right asymmetry that are expressed in the primitive streak; or for genes involved in signaling pathways that are expressed in the eye of Pax6 mutant mice.

The Differential Expression Search tab allows querying for genes that are expressed in one anatomical structure but not in another and/or at some developmental stages but not others. For example, one can search for genes that are expressed in the epithalamus but not in the hypothalamus; or for genes that are expressed at the morula stage (Theiler stage 3) but not at the blastocyst stages (Theiler stages 4 and 5). These searches will return a list of genes whose expression has been shown to be absent (not detected), as well as genes whose expression has not been analyzed or recorded in the database for the specified structures and stages. The two cases can be distinguished by filtering the results summaries for instances where expression was not detected (see below).

The Standard Search and Differential Expression Search return a results page with tabbed summaries for assay results, assays, genes, and images that match the search parameters. The assay results tab (upper) is displayed by default. It lists the gene studied, the assay type used, the anatomical system, age and tissue examined, indicates whether expression was detected, provides a link to the corresponding images, lists the mutant alleles of the specimen (if applicable), and provides the reference from which the data were derived. Links in the Result Details and Images columns lead to detailed expression records, such as the one shown in Figure 1. Arrowheads in column headers indicate that the column is sortable (one set is circled). The assay results tab (as well as the genes tab) allows for the export of results in text and spreadsheet formats (buttons in table header). The images tab (lower) shows all the images that match the search criteria, together with the gene(s) examined in that image and the assay type used and provides a link to the corresponding part of the detailed expression record. The expression summaries can be refined by using the "click to modify search" button or by using the filter options provided on the summary page. The content of all four tabbed summaries will change accordingly.
de-selected to interactively revise and refine the data summaries. Most of the data columns of the summaries are sortable. Furthermore, data from the Assay Results and Genes summaries can be downloaded in text or spreadsheet formats allowing upload into other applications.

Each row in the assays, assay results, and image summaries includes links to detailed expression records. Figure 1 shows, as an example, an entry for an immunohistochemistry experiment, illustrating the detail in which data in GXD are annotated. These Assay Detail pages display image panes together with their annotations. Each image pane is linked to the corresponding image detail page that shows the publication figure, which is often multi-paned, thus presenting the image pane in context. Image detail pages might also include links to external resources. For example, detail pages for the EurExpress and GenePaint images link back to the corresponding records at these sites where utilities such as serial-section browsers and high-resolution images are available. Some image detail pages also
have links to the Edinburgh Mouse Atlas and Gene Expression Database (EMAGE). GXD makes all its RNA in situ and immunohistochemistry images, together with their annotations, available to EMAGE (Richardson et al., 2014), so that the expression patterns can be mapped and queried spatially. If in situ images have been spatially mapped, GXD provides links from image detail pages to the corresponding mapped images in EMAGE. However, only a subset of the wild-type expression images can be spatially mapped (the 3D atlas is based on wild-type embryos). The standardized text-based description of expression patterns used by GXD is essential as it allows the representation and integration of all types of expression data from wild-type and mutant mice, as well as the further integration with other data that relate to anatomy, such as mouse phenotype and human disease data.

In short, the Gene Expression Data Query form enables researchers to quickly find specific sets of expression data. Query summaries can be interactively refined and lead to the detailed expression records.

Mouse Developmental Anatomy Browser

As discussed above, GXD and EMAP have developed an extensive ontology for mouse developmental anatomy to describe the time and space of gene expression in a standardized way. This enables intuitive and comprehensive expression searches at variable anatomical resolution. The ontology is structured hierarchically. Each term can have multiple parents, allowing the anatomy to be represented and searched from different perspectives. For example, “brain” is represented as part of the “nervous system” and as part of the “head”. Searches for expression in the “nervous system” or in the “head” will both return expression data for the brain.

The newest version of Mouse Developmental Anatomy comprises all 28 Theiler stages, including the embryonic (TS 1–26), newborn (TS 27), and postnatal mouse (TS 28). There is one “abstract” (nonstage specific) representation of the mouse anatomy that lists the anatomical structures for all Theiler stages, as well as the stage range during which each structure is present.
Acquiring and Curating Data

Data are acquired from the literature, by means of electronic data submissions, and through collaborations with projects that generate pertinent expression data at a large-scale. All data are reviewed by curators and annotated in standardized ways by making extensive use of controlled vocabularies and ontologies as illustrated in Figure 1. This is a prerequisite for the data integration and search capabilities that GXD provides.

GXD is the only effort that curates mouse developmental expression data from the literature in a systematic way. The first step in this process is populating the Gene Expression Literature Index as described above. The next step is to annotate the details of the expression data from these articles, including supplemental data, as illustrated in Figure 1. Papers are prioritized for detailed annotation based on the information in the Gene Expression Literature Index. Prioritization criteria include: genes for which there is no detailed data in the database; genes associated with human diseases; genes which, based on the number of publications, are underrepresented in the detailed portion of the database; publications that include a large amount of expression data; and publications that characterize the developmental expression pattern of a given gene in detail (often the first publications for that gene).

Large-scale electronic data sets are reviewed by a combination of computational and manual checks to make sure that the probe-to-gene assignments are up-to-date and that data entries are complete. Nomenclature issues, data ambiguities, and questions that might arise during the mapping of submitted data to ontologies are resolved together with the data provider before the data are loaded into GXD.

All data, including those from large-scale data sets, gain significant value when integrated with the other data in GXD and the larger MGI resource. All data can be explored together and searches can be done that are unavailable elsewhere. Furthermore, GXD ensures that data and data connections are maintained and kept up-to-date after large-scale projects have ended and gene models or the gene names used in the literature have changed.

Authors base their conclusions on much more primary data than will appear in a publication, they are specialists in their fields, and they have detailed knowledge about specific experimental parameters and potential pitfalls that is not available to curators. For these reasons, GXD curators rely on the text of the manuscript, or of data submissions, to derive annotations of expression patterns. These annotations are standardized by using terms from the anatomical ontology (which is expanded as required). However, curators do not interpret images themselves, thereby trying to derive additional expression results that have not been asserted in the study. For the reasons discussed above, interpretation of images by curators to infer expression or absence of expression would be problematic and error-prone. Instead, GXD displays standardized text annotations together with the corresponding images so users can see the original expression data (see Fig. 1). We have obtained permissions to include images from all major developmental journals, as well as from many others. Currently, over 70% of the result annotations are accompanied by images. Also in cases where GXD does not have the permission to include images, we annotate the data based on the information in the manuscript and provide a reference to the corresponding figure panes.

It is thus important to note that the detail of expression annotations in GXD is determined by the details provided in the text of published articles and electronic data submissions. Detailed descriptions of expression results will lead to more fine-grained data.

Gene Expression Literature Query

GXD provides researchers with an effective way to search the mouse embryonic expression literature. Our curators survey journals to find all publications that contain the types of data that GXD collects. As a first annotation step, they index all these publications with regard to the genes that have been studied, the expression assay types used, and the ages analyzed. These annotations are then combined with bibliographic information from PubMed to generate the Gene Expression Literature Index. The index is complete and up-to-date from 1990 onward for all major journals (~150). Currently, the index covers >21,800 references reporting expression data for >15,100 genes. An average of 1,200 papers are added to the index per year. All this information is available by means of the Gene Expression Literature Query form (http://www.informatics.jax.org/gxdlit). The query form allows researchers to quickly find publications that report specific sets of expression data (see Fig. 6). These searches are more effective and complete than PubMed searches because the index uses standard nomenclature for genes, assay types, and ages and because the annotations are based on the entire article, including supplemental data.
and complete annotations. Furthermore, a clear correspondence in the publication between stated expression results, probes, specimens, and images facilitates data annotation. Concise, unambiguous descriptions of expression patterns result in robust annotations. Issues of completeness, consistency, data identity, and clarity apply not only to GXD but to all databases that curate data from the published scientific literature. This includes other developmental organism databases such as GEISHA (Antin et al., 2014), Xenbase (James-Zorn et al., 2013), ZFIN (Howe et al., 2013), FlyBase (St. Pierre et al., 2014), and Wormbase (Harris et al., 2014). Publications that present data, results, and interpretations clearly and identify genes, strains, reagents, and methods adequately can significantly facilitate data curation and dissemination, thus greatly benefiting the researchers who generated the data initially as well as improving data access to the research community at large.

The Case for Electronic Data Submission

Both journal publications and databases are essential for the research enterprise, and they fulfill complementary roles. Journal publications excel in the narrative descriptions of novel discoveries and discussions of data implications and future applications. However, they usually include only part of the data pertinent to a study due to space constraints or because authors report only the results most relevant to the papers’ narrative. Expression (or non-expression) in anatomical systems peripheral to the narrative is often not reported. Furthermore, journal publications cannot and do not provide a framework for data integration and regular updates, with the result that the data cannot be searched adequately. Databases, on the other hand, do not have space constraints or an incentive to focus on specific scientific narratives; and they excel at the handling, maintenance, and integration of data, and in making them accessible to searches. Currently, data generated from conventional (non largescale) laboratories are primarily reported in free-text descriptions in journal publications. Unless database curators extract data from these publications and bring them into formats proper for integration and searching, these data will remain poorly integrated into our scientific knowledge base. This is clearly not a good solution for making research data accessible.

An obvious and effective solution to address this problem is to combine journal publications with electronic data submissions. Following the example of sequence and array expression data, the types of developmental expression data discussed here could be submitted to pertinent public databases in conjunction with publications. Submitters would receive accession numbers that can be cited in the publications. Without causing an undue burden on submitters, they could, for example, provide an expanded legend for each image that includes more complete descriptions of the expression patterns seen. Thus, electronic submission can include more primary data and results can be described in greater detail and in more standardized ways. While such submissions would still require review and annotation by database curators, it would significantly facilitate the acquisition, integration, and dissemination of data. Having the data widely accessible to many types of searches would benefit everyone: the journal (because all the data would be tied to the publication reference), the submitter (because their data would be more accessible), and the scientific community as a whole (because of the increased database content and utility).

GXD accepts electronic data submissions for the types of data it collects; see the “Send us your data” tab at the bottom of the GXD Home Page for instructions (http://www.informatics.jax.org/gxd/gxd_submission_guidelines.shtml). Other developmental databases, such as FlyBase, GEISHA, ZFIN, and Xenbase, are welcoming electronic data submissions as well. The value of all these community resources is proportional to the amount and the quality of data they contain, and it is time for the research community to realize the potential of electronic data submissions.

Future Directions

GXD will continue to acquire data from the literature and electronic data submissions and to improve its search and display capabilities. Later this year, we plan to add interactive matrix views of expression data. Tissue-by-developmental stage matrices will provide high-level overviews of the spatio-temporal expression patterns of genes. Tissue-by-gene matrices will enable a comparison of expression patterns. Both types of matrices can be expanded (and collapsed) along the tissue axis, based on the hierarchical organization of the anatomy. Thus, these matrix views will provide users with intuitive high-level summaries of expression results from where they can drill down to more detail.

GXD and the larger MGI resource are also planning to implement additional links to other developmental organism databases. In the shorter term, gene-based links will enable users to look up and compare expression data (and other types of data) for orthologous genes. As described above, we are also working on establishing cross-references between anatomical ontologies to enable, in the longer term, an anatomy-based comparison of developmental gene expression patterns.

User Support and Outreach

User Support and GXD curatorial staff actively seek to provide presentations, demonstrations and training sessions on GXD and the larger MGI resource at many meetings. Upon request user support personnel also provide on-site visits and training workshops, as well as remote interactive sessions. GXD is continually looking for ways to improve the user experience and, thus, direct user interactions, surveys and collaborations provide valuable feedback on the project from biologists, biomedical researchers, ontology developers and computational biologists who use our data. Furthermore, the GXD Advisory Board provides critical input and guidance for the project.

Database Access and Contact Information

The GXD home page can be accessed directly at http://www.informatics.jax.org/expression.shtml; by means of the MGI home page (http://www.informatics.jax.org/) by following the topic “Gene Expression Database (GXD)”; or from any other page within MGI by clicking the “Expression” tab of the navigation bar. For Web-based access to GXD and MGI, we recommend Firefox, Chrome, or Safari. Online help is available by means of the FAQs on the GXD home page and by clicking on the question marks in the upper right corner of most GXD pages. User Support personnel can be contacted by means of email to mgi-help@jax.org or by means of the “Contact Us” link in the navigation bar. This article focuses on Web-based access to the GXD database.
proper. Additional computational tools to explore GXD’s expression data are available, such as the GXD BioMart (accessible by means of the GXD home page) and MouseMine (www.mouse-mine.org). MouseMine, in particular, offers advanced iterative search and filtering capabilities.

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