Recent advances in high-throughput gene targeting and conditional mutagenesis are creating new and powerful resources to study the in vivo function of mammalian genes using the mouse as an experimental model. Mutant ES cells and mice are being generated at a rapid rate to study the molecular and phenotypic consequences of genetic mutations, and to correlate these study results with human disease conditions. Likewise, classical genetics approaches to identify mutations in the mouse genome that cause specific phenotypes have become more effective. Here, we describe methods to quickly obtain information on what mutant ES cells and mice are available, including recombinase driver lines for the generation of conditional mutants. Further, we describe means to access genetic and phenotypic data that identify mouse models for specific human diseases.

1. Introduction

Closely related to human and readily accessible to detailed genetic, molecular, and phenotypic analysis, the mouse serves as a premier animal model in biomedical research. The classical forward genetics approach starts with a mouse phenotype that resembles a human disease and determines the mutations that cause the phenotype. Reverse genetics creates specific mutations, characterizes the resulting phenotypes and correlates them with human disease conditions. Both approaches have become much more effective in recent years because of the availability of complete genome sequences and due to the development of high-throughput targeting methods, many of which are described in this issue. Complementary in nature, both approaches deliver, at a rapid rate, mutant ES cells and/or mice as a starting point for further investigations aimed to understand the molecular mechanisms that lead from a genotype to a phenotype. This article describes methods to readily access up-to-date information on what mutant ES cells and mice are available, as well as information about mouse phenotypes and their correlation with human disease conditions. Methods are framed as use cases that start with specific questions.

2. Description of methods

2.1. Use Case 1: Are there mutant ES cells or mice available for my gene of interest? What mutations does the gene carry in these mutants? How can I obtain specific mutant ES cells or mice?

There are currently several large-scale programs underway to mutate all protein-coding genes in the mouse using gene trapping and gene targeting technologies: the Knockout Mouse Project (KOMP; USA) [1]; the European Conditional Mouse Mutagenesis Program (EUCOMM) [2]; the North American Conditional Mouse Mutagenesis Project (NorCOMM; Canada); and the high-throughput gene trapping effort by the Texas A&M Institute for Genomic Medicine (TIGM; USA) [3]. In order to pursue their work in a collaborative and coordinated manner, these groups have formed the International Knockout Mouse Consortium (IKMC) [4,5]. The IKMC web portal [www.knockoutmouse.org] is the central public web site for IKMC data, giving researchers world-wide access to up-to-date information on IKMC knockout vectors, ES cells and mice, and links to repositories from which these products can be ordered [6]. The IKMC database also provides data and tools to the IKMC for selecting and prioritizing genes. For example, it indicates to the IKMC for what genes mutant ES cells or mice have already been generated by other researchers. Because all this external information and links to pertinent resources are also readily available through the web site, the IKMC web portal is an excellent central starting point for the queries listed above. In the following, we describe a workflow for performing these queries: Go to www.knockoutmouse.org.
Enter the gene symbol(s) or gene ID(s) for your gene(s) of interest into the Search box and start the search.

Search results are listed as a summary with a record (row) for each gene that matches your query (Fig. 1 top). Each record includes (i) the gene symbol; (ii) a high-level summary of IKMC knockout attempts that states which programs are working on the gene, provides the status of the most advanced targeting effort per program with a link to more details, and indicates the availability of targeting vectors, mutant ES cells and mice, with links to the repositories that distribute the respective IKMC products; and (iii)
other resources that report on mutant ES cells or mice for the gene of interest, with the number of, and links to, corresponding entries at the respective sites.

If you are interested in IKMC products, follow the ‘Details’ link in the ‘IKMC Knockout Attemps’ column. This brings you to a page that lists all IKMC knockout attempts for your gene of interest (Fig. 1 bottom). One program might try different targeting vectors to mutate a gene, resulting in several targeting projects. The production pipeline status for each project is displayed, and the availability of products is indicated via Order links that lead to the respective repositories. As soon as targeting vectors have been generated for a given project, an ‘Allele Details’ link is displayed to the right of the progress bar. Clicking on this link opens a graphical display that illustrates the salient molecular features of the targeted allele. On the top left side of this newly opened display is another link ‘View this project’ that leads to more comprehensive information about the mice, ES cells and targeting vector generated by a specific project. Order links guiding users to the respective repositories are provided on this page as well.

The search path just described leads to information about gene targeting products (vectors, ES cells and mice) from KOMP, EU-COMM and NorCOMM, and gene trap ES cell lines from TIGM. Information about gene trap ES cell lines from EU-COMM and NorCOMM will be fully integrated into the IKMC web portal in the near future. Then this data will be searchable and displayed in the same way as the other IKMC products described above.

Currently, if you are interested in EU-COMM and NorCOMM gene traps, or in non-IKMC products, follow the links provided in the ‘Other Resources’ section of the initial query summary to find the entries for your gene in these databases (Fig. 1 top). The International Mouse Strain Resource (IMSR) [7] is a searchable database of inbred and mutant mouse strains and stocks available worldwide and provides access to the respective holder sites, including all the members of FIMRE, the Federation of International Mouse Resources. The International Gene Trap Consortium (IGTC) database [8] records all gene trap lines available from the consortium and provides links to corresponding holder sites. The Mouse Genome Informatics (MGI) database [9,10] represents all mutants for a given gene that have been reported in the published literature, by the IKMC, and by the IMSR. If pertinent mutant products are recorded in MGI, the ‘Other Resources’ section provides two links, one to targeted mutations in MGI, one to other mutations in MGI. The latter category includes gene trap ES cell lines from EU-COMM and NorCOMM. All mutant records in MGI include a reference to the original publication. ES cells or mice that are not available through the IKMC or the IMSR might be obtained by contacting the authors.

IKMC data are also accessible through MGI and the IMSR; IKMC gene trap data are also available through the IGTC. However, the IKMC web portal provides the most up-to-date and the most detailed molecular information about its products, reports on work in progress, and maintains up-to-date links to pertinent external resources. Therefore, the search strategy presented here offers effective means to find, globally, mutant ES cells and mice together with detailed information about targeting vectors and mutant alleles.

2.2. Use Case 2: Are recombinase driver lines available under promoters of my choice? Are there recombinase driver lines that show activity in tissues of my choice?

A large number of targeted and gene trapped alleles produced by the IKMC, and an increasing number of targeted alleles generated by the biomedical research community, feature recombinase recognition sites that allow the generation of conditional mutants [2,11–13]. To take advantage of these conditional alleles, mouse lines that carry recombinases under specific promoters/enhancers are being developed [14]. Time and tissue specificity of these recombinase lines is characterized, either by examining recombinase expression patterns, or, more accurately, by using reporter strains to determine the conditional allele pattern generated by the recombinase [15]. Pertinent data about transgenic and knock-in recombinase lines are available through the recently established Cre-Portal (www.creportal.org). The site lets you search for recombinase driver lines that display recombinase activity in specific tissues, or for mouse lines with recombinases under specific promoters/enhancers. As part of the larger MGI system, the portal provides the recombinase expression and reporter data, as well as phenotypic data obtained in conditional mutants. Further it indicates which mouse lines are available through the IMSR and provides links to corresponding entries in the IMSR database. The following query example illustrates how to find information about recombinase driver lines with specific promoters (see also Fig. 2):

Go to www.creportal.org.

In the ‘Access Data’ section use the ‘Search for allele by promoter/driver specificity’. Select the promoter/driver of your choice from the selection list and click the ‘Go’ button.

The query result summary lists all the mouse lines/alleles that match your query. The ‘Recombinase Data’ column indicates whether recombinase activity data are available and in how many anatomic systems activity has been measured. The ‘Find Mice’ column indicates whether mouse lines are available and provides links to the IMSR. The ‘Refs’ column gives the number of publications that report on the allele/recombinase line, as one potential measure for the utility of the line, and links to the complete list of references.

To look up recombinase activity data for a specific allele, expand the corresponding information in the ‘Recombinase Data’ column by clicking on the triangle to the left of the phrase ‘Detected in x systems’. Then follow the link for a specific anatomic system to look up pertinent data, including the specific anatomic structures in which recombinase activity was reported, pertinent image data, as well as additional assay and genetic information.

To look up all recombinase and phenotype data for a given allele, on the query results summary, in the ‘Allele Symbol’ column, follow the link ‘phenotype data’.

The search and navigation path presented here is just one way to address this specific use case. For more information about the utility of the Cre Portal, you can consult the FAQs provided at the site and explore the database and its content in more detail.

2.3. Use Case 3: Are there mouse models available for a given human disease?

As pointed out above, mouse models are generated by forward and reverse genetic approaches. A key to both strategies is the phenotypic characterization of mouse mutants and the correlation of these data with human disease conditions. The MGI database is a primary resource for this type of information. MGI provides various means to search for genes and mouse mutants based on phenotypic and human disease information. Pertinent search forms and FAQs are available at www.informatics.jax.org/phenotypes.shtml. Here we illustrate one example: finding mouse models based on human disease terms.

In the Access Data Section, follow the link ‘Human Disease (OMIM) Browser’.

OMIM [16] disease terms are listed in alphabetical order and the availability of mouse models is indicated, in parentheses, to the right of terms. Search or browse for your disease term of interest and click on the corresponding term (Fig. 3 top left).

Fig. 3 (top right) shows the entry for Angelman Syndrome; AS (OMIM ID: 105830). As indicated on this page, human disease characteristics are associated with mutations in the human gene...
UBE3A and with mutations in the mouse orthologous gene Ube3a. However, mouse mutants for the Gabrb3 and Snrpn genes also show disease characteristics for Angelman Syndrome although OMIM data currently do not associate the disease with the orthologous human genes. Conversely, there are three additional human genes associated with Angelman Syndrome in OMIM: ANCR, CDKL5, and MECP2. The mouse ortholog for ANCR is not yet identified, and mouse mutants in Cdkl5 and Mecp2 do not show the disease characteristics. This clearly illustrates that whether a mouse mutant is a good model for a human disease conditions cannot be inferred simply by orthology. The detailed curation of mouse disease model information performed by MGI is needed to enable database queries such as the one described here.

The page illustrated by the top right of Fig. 3 serves as an entry point to much more detailed information about the respective mouse models. By following the links in the ‘Mouse Models’ section, you can look up the detailed annotations for each allele, including information about the nature of the mutation and the phenotypes displayed on different genetic backgrounds. If MGI has recorded expression information for the respective mutant, links to these data are provided as well. To see the full detail of the phenotype information on the ‘Allele Detail’ pages, use the ‘show all annotated terms’ or ‘show all phenotypic details’ functions, or expand specific sections by clicking on the triangles.

Allele Detail pages in MGI can contain a lot of information just for one specific allele. Moreover, there can be many alleles, and thus many Allele Detail pages for a given gene. All the information that MGI holds for a given gene can be accessed from the respective Gene Detail page, including information about chromosomal location, mammalian orthologs, sequences, mutant alleles and phenotypes, Gene Ontology (GO) classifications, and expression data. Many links to external resources are also provided.
3. Concluding remarks

There are many methods to study mammalian gene function. It is beyond the scope of this article to describe methods for accessing all the databases that hold pertinent information. Instead we have focused on databases that deal with recent developments holding great promise for understanding the in vivo function of mammalian genes and their role in human health and disease: high-throughput gene targeting and trapping, tools to create conditional mouse mutants, and the correlation of mouse phenotype and human disease information. The methods described here access central data hubs that provide extensive, highly-curated, and

Fig. 3. Finding mouse models for specific human diseases. Top left: Part of the page that shows the search results for ‘Angelman Syndrome’, upon using the ‘Human Disease (OMIM) Browser’ at MGI. Top right: ‘Human Disease and Mouse Model Detail’ page for Angelman Syndrome. Based on available phenotypic data, orthologous mouse and human genes are categorized according to their association with human disease characteristics. Mouse models for the respective categories are listed at the bottom part of the page. Each mouse model record specifies the exact allele composition and the genetic background under which the phenotype was observed, and points to the corresponding publication. Each mouse model record links to much more detailed allele and phenotypic data (red arrow). Bottom: ‘Allele Detail’ page for $Ube3a^{tm1Alb}$. This page provides detailed molecular and phenotypic information for each allele. The sections ‘Phenotype summary’ and ‘Phenotypic data by genotype’ can be expanded by using the ‘show all’ functions or by clicking on the triangles.
up-to-date information on these subject matters, and feature connections to many additional databases. There are other methods to query the resources described here, and the specific ways to perform the example queries illustrated in this article might change as the databases evolve. However, once one is familiar with these resources, it will be easy to exploit them more fully and to adapt to changes. All three web sites described here provide up-to-date online help and FAQ pages for further guidance.

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