The Zebrafish Model Organism Database: new support for human disease models, mutation details, gene expression phenotypes and searching


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ABSTRACT

The Zebrafish Model Organism Database (ZFIN; http://zfin.org) is the central resource for zebrafish (Danio rerio) genetic, genomic, phenotypic and developmental data. ZFIN curators provide expert manual curation and integration of comprehensive data involving zebrafish genes, mutants, transgenic constructs and lines, phenotypes, genotypes, gene expressions, morpholinos, TALENs, CRISPRs, antibodies, anatomical structures, models of human disease and publications. We integrate curated, directly submitted, and collaboratively generated data, making these available to zebrafish research community. Among the vertebrate model organisms, zebrafish are superbly suited for rapid generation of sequence-targeted mutant lines, characterization of phenotypes including gene expression patterns, and generation of human disease models. The recent rapid adoption of zebrafish as human disease models is making management of these data particularly important to both the research and clinical communities. Here, we describe recent enhancements to ZFIN including use of a new experimental conditions ontology, ‘Fish’ records in the ZFIN database, support for gene expression phenotypes, models of human disease, mutation details at the DNA, RNA and protein levels, and updates to the ZFIN single box search.

INTRODUCTION

The Zebrafish Model Organism Database (ZFIN; http://zfin.org) is the community resource for genetic and genomic data involving zebrafish (Danio rerio). Data in ZFIN includes mutants, gene expression, phenotypes, knockdown reagents, antibodies, transgenic constructs, and reporter lines. These data are integrated from direct data submissions from the research community and expert curation of published literature by biocurators. ZFIN aims to provide the best support possible for current needs of the scientific research community. In this paper, we describe new features added to ZFIN in the past two years. These include use of a new experimental conditions ontology, curation of gene expression phenotypes, zebrafish models of human disease and molecular details of mutants, as well as the addition of a Reporter Lines category and a figure gallery to the single box search interface. The release schedule for these new projects, descriptive metrics for the new data types, and descriptive metrics for usage and data content of the ZFIN resource in general can be found in the Supplementary Tables S1–S4.

THE ‘FISH’ CONCEPT AND EXPERIMENTAL CONDITIONS

Gene expression patterns, phenotypes, and models of human disease are all functions of the combined effects of the genetic constitution of an organism (genotype and genetic background), gene knockdown reagents, and the experimental conditions under which observations are made. At ZFIN, we have long been curating gene expression and phenotype from the zebrafish literature to a combination of a genotype plus the experimental conditions (1–3). Historically, ZFIN included the sequence targeting reagents (STRs) morpholinos, TALENs, and CRISPRs in experimental conditions when they were used as transient genetic modifiers (i.e. not used as germline mutagens). In order to obtain a better distinction between the genetic modifiers of a fish and the non-genetic experimental conditions, the STRs were moved out of the experimental conditions and included along with the genotype in a new database ob-
bject we call a ‘Fish’ (Figure 1). The Fish data model allows grouping of the transient and non-transient genetic modifiers, allowing researchers to more easily understand the number and variety of affected genes for which gene expression and phenotype observations are made. Fish records get unique and persistent identifiers in ZFIN in the form ZDB-FISH-########-###### (see example in Figure 1).

All gene expression and phenotype annotations, and models of human disease are currently annotated to a Fish record associated with specific experimental conditions. Information about Fish records (genotype, background, and STRs when applicable) can be found on ZFIN Fish record pages and in downloads files. Data from different model organism databases is currently annotated to records having varying degrees of specificity with regard to the genotype involved. In some cases annotations are made directly to allele records and in other cases annotations are made to more complex records like genotypes or ‘Fish’. This presents a challenge when trying to work with or combine these data. The ‘Fish’ database object aligns with ‘effective genotype’ (GENO:00005252) term from the GENO ontology, which was developed by Monarch Initiative to combine data that may be annotated to different levels of granularity in the genotype/allele hierarchy.

Experimental conditions are the conditions under which experiments are performed and data are gathered. They can range from ‘Standard’, as defined in the Zebrafish Book (6) or ‘Generic Control’ (non-standard control treatments such as a vehicle DMSO injection) to any sort of manipulation of the environment of the fish (as in exposure to a chemical or a non-standard light/dark cycle) or physical manipulation of the fish itself (as in amputation). Historically, experimental conditions in ZFIN were limited to a small list of frequently used conditions. Details regarding the conditions, such as the structure amputated or the chemical used, were recorded in a text note which lacked the necessary structure for effective searching. To address those limitations, ZFIN experimental conditions are now annotated using a new ontology called the Zebrafish Experimental Conditions Ontology (ZECO). ZECO provides a more expressive representation of experimental conditions used in zebrafish research while providing an ontological framework to support searching, data reporting, and interoperability of ZFIN annotations using experimental conditions. The full scope and use of the ZECO ontology will be described in another publication.

**GENE EXPRESSION AS A PHENOTYPE**

Gene expression patterns in mutants frequently differ greatly from patterns in wild-type animals in spatial extent, level (increased or decreased), or location. These differences illuminate genetic interactions and pathways. In some cases, particularly during early development, a change in an expression pattern may be the only notable phenotype. Until recently, phenotype data and gene expression data were entirely separate in ZFIN. The ZFIN database could store expression annotations such as ‘gene \( xyz \) is expressed in the kidney in \( abc \) mutant fish’ and phenotype annotations such as ‘the kidney is enlarged in \( abc \) mutant fish’. However, there was no way to record that the gene expression pattern of gene \( xyz \) in the kidney of \( abc \) mutant fish differed from the wild-type pattern. Database support for expression patterns as phenotypes (EaP) now allows these important phenotypes to be curated and searched. Between January and September 2016 ZFIN curators added 3087 EaP annotations from 253 publications. During that period, 59% of the 429 publications curated for gene expression and 20% of all 1281 expertly curated publications contained at least one EaP annotation (Supplementary Table S2). This underscores the importance of these expression phenotype annotations in the more general phenotype annotation landscape.

**Expression phenotype format**

Gene expression phenotypes follow the same ‘Entity:Quality:Tag’ (EQT) format as other phenotypes in ZFIN (1,7). EQT annotations are typically composed of an ‘Entity’ term commonly from the Zebrafish Anatomy Ontology (AO) (8) or the Gene Ontology (GO) (9,10) combined with a Quality term from the Phenotypic Quality ontology (PATO) (11) and a Tag (‘normal’, ‘abnormal’, or ‘ameliorated’). The Entity in an EaP annotation combines an anatomical structure with either a gene or an antibody. An antibody is used in cases of antibody labeling data that is not connected with expression of a single gene, e.g. labeling by antibodies that recognize carbohydrate epitopes or multiple members of a gene family. Table I describes two phenotypes in the heart, one standard anatomical phenotype and one EaP. The standard anatomical phenotype could be stated in words as ‘the heart has abnormally increased volume’. The EaP could be stated as ‘\( /f2 \) expression in the heart is abnormally mislocalized.’

There are two possible EaP annotation tags—abnormal and ameliorated. The ameliorated tag is used to describe an expression pattern observed in a mutant that is significantly or completely rescued by an additional mutation or reagent. The EaP Quality is chosen from a small subset of terms from the PATO ontology (Table 2). After extensive discussion within ZFIN and with members of the zebrafish research community, it was concluded that the vast majority of user needs could be met by using only seven PATO quality terms for EaPs having the ‘abnormal’ tag, and three PATO quality terms for EaPs having the ‘ameliorated’ tag. The use of a small collection of PATO quality terms made the addition of EaP curation to the ZFIN curation interface simpler, keeping the day-to-day process of EaP curation quick and efficient.

**Logical representation of EaP data**

EaP annotations come in two forms: gene and antibody. Which form is used depends upon whether the expression pattern is for a gene or for an antibody staining pattern that
Figure 1. The Fish concept in ZFIN. Information found at the top of a Fish page in ZFIN. Fish records include the genotype (Green; Genotype), the genetic background (Yellow, G.B.), and the sequence targeting reagents (Red, STRs) when applicable. Fish IDs have the form ZDB-FISH-#####-#####. Fish ZDB-FISH-150901-11301 (https://zfin.org/ZDB-FISH-150901-11301) includes a homozygous \textit{hand2^c99/c99} gene on the AB genetic background plus three morpholinos targeting the \textit{dlx3b}, \textit{dlx5a} and \textit{dlx4b} genes.

Table 1. Howe et al. NAR database issue 2016

<table>
<thead>
<tr>
<th>Annotation Type</th>
<th>Entity</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anatomical phenotype</td>
<td>Heart</td>
<td>Increased volume</td>
</tr>
<tr>
<td>Expression phenotype</td>
<td>Heart \textit{lft}2 expression</td>
<td>Mislocated</td>
</tr>
</tbody>
</table>

Table 2. Howe et al. NAR database issue 2016

<table>
<thead>
<tr>
<th>EaP Qualities with ‘Abnormal’</th>
<th>Tag corresponding PATO ID</th>
</tr>
</thead>
<tbody>
<tr>
<td>Increased amount</td>
<td>PATO:0000470</td>
</tr>
<tr>
<td>Decreased amount</td>
<td>PATO:0001970</td>
</tr>
<tr>
<td>Absent</td>
<td>PATO:0000462</td>
</tr>
<tr>
<td>Mislocalized</td>
<td>PATO:0000628</td>
</tr>
<tr>
<td>Spatial pattern</td>
<td>PATO:000060</td>
</tr>
<tr>
<td>Increased distribution</td>
<td>PATO:0001671</td>
</tr>
<tr>
<td>Decreased distribution</td>
<td>PATO:0001672</td>
</tr>
</tbody>
</table>

EaP Qualities with ‘Ameliorated’ Tag

<table>
<thead>
<tr>
<th>Tag corresponding PATO ID</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amount</td>
</tr>
<tr>
<td>Position</td>
</tr>
<tr>
<td>Spatial pattern</td>
</tr>
</tbody>
</table>

is not clearly attributable to a single gene. In each case the phenotype is manifest in the expression of a gene or epitope in an anatomical location.

- **Gene Form**: \(<\text{PATO}>\ <\text{RO:inheres_in}>\text{some (gene and \text{RO:expressed_in} some \text{ZFA})}\)
- **Antibody Form**: \(<\text{PATO}>\ <\text{RO:inheres_in}>\text{some (\text{SO:epitope} and \text{RO:expressed_in} some \text{ZFA})}\)

\(<\text{PATO}>\) denotes a term from our short list of EaP PATO qualities. \(<\text{ZFA}>\) denotes a term from the zebrafish anatomical ontology which can also optionally be post-composed with a term from the GO Cellular Component ontology. \(<\text{RO:expressed_in}>\) is the term ‘expressed in’ (RO:0002206) from the Relations Ontology (RO) (12). \(<\text{SO:epitope}>\) is the term ‘epitope’ (SO:0001018) from the Sequence Ontology (SO) (13). In both cases, the phenotype inheres in something being expressed in an anatomical structure.

Curation of expression phenotypes

Curators add EaP data to ZFIN quickly using a simple extension of the existing ‘building block’ style gene expression curation interface. In the gene expression section of the ZFIN curation interface, ‘Experiments’ are created that combine Gene, Antibody (optional), Assay, Fish, and Experimental Conditions. Combining Experiments with a developmental stage and Figure number makes ‘Expressions’.

The ‘Anatomy pile’ is a list of anatomy terms that is built by curators as they work on a particular publication. Associating one or more ‘expressions’ with one or more anatomical structures from the ‘Anatomy pile’ produces gene expression annotations (Figure 2). This approach maximizes re-use, so that experiments and anatomy terms are only entered once, and then they are combined as needed for different figures.

EaP terms are represented on the Anatomy Pile along with non-EaP anatomy terms, so the pile might contain ‘posterior lateral plate mesoderm’ and also ‘posterior lateral plate mesoderm: decreased amount’. Adding ‘posterior lateral plate mesoderm’ to an Expression creates a gene expression entry in the database. Adding ‘posterior lateral plate mesoderm: decreased amount’ creates both gene expression and gene expression phenotype annotations.

Displaying and searching EaPs

EaP phenotypes are shown along with non-EaP phenotypes in the Phenotype section of Fish and Figure pages. A variety of different phenotypes, including EaP as well as non-EaP phenotypes can be seen in the ‘Phenotype Details’ section of a typical Figure page (Figure 3).

Searching for EaP annotations is done in the same manner as searching for expression or phenotype annotations. For example, an EaP annotation representing ‘fgf8a expression is mislocalized to the retina’ can be found in the fgf8a gene expression data as well as the retina phenotype data.
Figure 2. A portion of the ZFIN gene expression curation interface. (A) portion of the ZFIN gene expression curation interface showing EaP and non-EaP expression annotations. (A) Two Expression rows, the first holding an ordinary gene expression annotation in a wild-type fish, the second row holding an EaP annotation in a mutant fish. (B) The Anatomy Pile section of the ZFIN curation interface showing the anatomical structure lateral plate mesoderm in standard format as well as expression phenotype format for lateral plate mesoderm - decreased amount. Image captured on 18 August 2016.

Figure 3. The Phenotype Details section of the ZFIN figure page. https://zfin.org/ZDB-FIG-151214-52#expDetail shows examples of EaP and non-EaP phenotypes displayed in the Phenotype column of the Phenotype Details section for the figure. Image captured on 18 August 2016.

EaP has been integrated into ZFIN’s new single box search page. (https://zfin.org/search)

A new facet—“Misexpressed Gene”, aids EaP discovery in five different data categories: Gene/Transcript, Phenotype, Fish, Mutation/Tg and Figure.

Clicking on a gene symbol in the ‘Misexpressed Gene’ facet will filter the result set to only include results that have EaP data for that gene. Another way to access EaP data is to use ‘misexpression’ in the search string. Table 3 includes examples of four common kinds of searches using the Misexpressed Gene facet or the ‘misexpression’ key word.

EaP data downloads

Data in ZFIN are available from the downloads page (http://zfin.org/downloads), where the ‘Phenotype Data’ section has two new download files to support EaP annotations:


HUMAN DISEASE

The zebrafish has become a well-established vertebrate model organism. The optical clarity, accessibility and rapid development of the zebrafish embryo allow organ systems and tissues to be closely observed as they form. There are close parallels between the genomes of zebrafish and humans. Sequencing of the zebrafish genome has revealed that 71% of human genes have at least one zebrafish ortholog while 69% of zebrafish genes have at least one human ortholog. In addition, 82% of human disease genes in OMIM have a zebrafish ortholog (14). A variety of powerful tools for genome manipulation are now available, facilitating the creation of zebrafish mutants that model specific human diseases. This section will highlight recent enhancements at ZFIN that allow users to access and navigate human disease information and zebrafish disease models for use in translational research.

A link to human disease information in ZFIN is provided through two different methods, one automated, one curated. An automated pipeline links OMIM diseases associated with human genes with the orthologous zebrafish genes using curated orthology. Orthology curation is done
Table 3. Howe et al. NAR database issue 2016

<table>
<thead>
<tr>
<th>Search</th>
<th>Category</th>
<th>Filter settings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Which Fish have perturbed gene expression in the heart?</td>
<td>Fish</td>
<td>Affected Anatomy: heart</td>
</tr>
<tr>
<td>Which phenotypes include \textit{arg1} misexpression?</td>
<td>Phenotype</td>
<td>Misexpressed Gene: \textit{arg1}</td>
</tr>
<tr>
<td>Which figures show \textit{lef1} misexpression?</td>
<td>Figure</td>
<td>Misexpressed Gene: \textit{lef1}</td>
</tr>
<tr>
<td>Which phenotypes include misexpression of members of the \textit{bmp} gene family?</td>
<td>Phenotype</td>
<td>No Filters set Query string = ‘bmp misexpression’</td>
</tr>
</tbody>
</table>

Figure 4. Human disease information in ZFIN. (A) The human disease section from the \textit{atxn2} gene page showing OMIM phenotypes associated with the human ortholog of zebrafish \textit{atxn2}. (B) The Disease page for Parkinson’s disease showing details from the DO including disease name, synonyms, definition, cross references to other resources and term relationships within DO. The zebrafish models section shows models of this disease generated in zebrafish with chemical treatments or genetic alterations. Image captured on 24 August 2016.
using the most recent human and zebrafish genome assemblies available at Ensembl at the time of each analysis (15). Expert curation of publications link zebrafish models of human disease, human disease, and the relevant publications utilizing the Disease Ontology (DO) (16). Each of these methods is described in more detail below. Phenotype based comparison approaches for human disease is under consideration (17). A tool that has been developed by the Monarch Initiative presents a grid detailing similarities between human and zebrafish phenotypes (18). ZFIN is evaluating integration of this widget on the disease term pages to provide users with a different option for linking data to human disease.

**Gene-disease association via OMIM**

Gene-to-disease associations are brought into ZFIN through a data exchange with NCBI. This association is established through an automated pipeline that relates OMIM disease IDs with orthologous zebrafish genes at ZFIN via the NCBI human Gene ID. Zebrafish genes that are orthologous to human genes implicated in disease display OMIM disease data on the ZFIN gene pages, providing users with a snapshot of diseases for which the zebrafish gene could provide a model (Figure 4A). Human genes and orthologous zebrafish genes are also made available on the ZFIN human disease page (Figure 4B). It is important to note that this gene-to-disease relationship is derived solely through orthology determinations between zebrafish and human genes and are not reviewed by curators at ZFIN.

**Disease annotation through literature curation**

Expert manual curation of zebrafish models of human disease is another way in which ZFIN supports human disease model discovery. Zebrafish disease models are annotated in ZFIN through a combination of Fish, experimental conditions and terms from the Disease Ontology. The Fish could include a wild-type strain, or a mutant or transgenic line. The experimental conditions refer to non-standard treatment conditions such as temperature, diet or chemical treatment applied to the fish to produce a phenotype. If an experimental treatment was not used on the fish, ‘standard’ or ‘generic control’ is used as the experimental condition. The DO is used to make an annotation to the human disease and represents diseases in an ontological hierarchy, with cross references to disease concepts from OMIM, SNOMed, NCI Thesaurus and MeSH (16). The data structure that ZFIN uses to record disease models can be summarized as: ‘Fish in specific ‘experimental conditions’ is a model of ‘disease X’ as described in ‘Citation N’. A summary of disease models in this format can be seen on the Parkinson’s disease page shown in Figure 4B. In addition, terms from the DO are used to establish a relationship between a human disease and a journal article about the disease, regardless of whether a zebrafish model is created.

Every annotation linking a zebrafish model of human disease to a disease term is supported by an evidence code...
Figure 6. Mutation details as seen on the genomic feature page and the gene page. Mutation details are displayed on the genomic feature and gene pages in ZFIN. (A) Snap shot of the $b1127$ point mutation feature page. The mutation details section (bracket) shows the DNA/cDNA change with the corresponding location within the gene, and the consequences at the transcript and protein level. (B) Snap shot of the $fam20b$ gene page. The mutations available for this gene are displayed with details regarding their localization within the gene, and the transcript consequences of the mutation (bracket). Image captured on 18 August 2016.

Figure 7. An example search in the Reporter Lines Category in ZFIN single box search interface. In this example, the Reporter Line Category was selected and values were set in the Expression Anatomy, Stage, and Reporter Gene filters to answer the question 'Are there any reporter lines that express mCherry in Meckel's cartilage during the larval stage?'. Image captured on 24 August 2016.
Figure 8. The single box search figure gallery. The figure gallery provides a tiled view of images associated with results of a single box search. Here, the Expression category has been searched for ‘heart’ and the ‘show images’ button was clicked to get to the figure gallery view (lower). Clicking on any figure produces a magnified view of that image (red arrow). The magnified view includes the figure caption and curated data associated with that figure. Left and Right arrows provide continued navigation through the image set in the magnified view. The particular image clicked here is Figure 1 from Lou et al. (23).

Images captured on 24 August 2016.

from the Evidence and Conclusion Ontology (ECO) (19). The evidence code ‘Traceable Author Statement (TAS)’ is used when authors make an explicit assertion about a Fish plus experimental condition as a model for a specific disease. The ‘Inferred by Curator (IC)’ evidence code is used if authors do not make an explicit assertion about the disease model but instead report phenotypes similar to the human patient pathologies, and the intent of their publication is to elucidate portions of the disease mechanism based on results obtained in zebrafish.

As of 10 October 2016 there are 372 curated zebrafish disease models and annotations using 357 DO terms. 16% of curated publications between 17 June and 10 October 2016 included a disease model (Supplementary Table S2).

**Searching for disease models**

Human disease information in ZFIN can be queried by entering a disease name or gene of interest into the search box on the home page. The results can be refined further by selecting a data category on the left of the search results or by selecting a ‘related data’ link in the body of one of the search results (Figure 5).
Disease term search results link to the Disease term page, which provides more details on the Human disease including definitions, synonyms and cross-references in other databases (Figure 4B). The relationship of the disease in the DO to other terms is also provided. The zebrafish models section of the disease page contains details of the genetic background of the fish, experimental conditions and the citation referring to the use of this model. Following either the Fish or Citations link in the models section provides access to phenotype information for each individual model. A list of established disease models and the ability to search for them will be very useful to researchers and/or clinicians studying a particular disease.

**MUTATION DETAILS**

Information about nucleotide or peptide changes for alleles has previously been provided in a free text note on genomic feature (allele) pages. Annotation and display of these data have now been updated to provide better reporting and searching of mutation details. The genomic feature page now displays detailed information about the DNA/cDNA change, as well as associated transcript and peptide changes for alleles (Figure 6A). These data originate from publications, direct submissions to the nomenclature coordinator, or via data loads. As of October 2016, ZFIN includes DNA, transcript, or protein mutation details for 83%, 82% and 3% of applicable feature records respectively (Supplementary Table S2).

Various types of DNA/cDNA changes are described based on the type of mutation that has occurred. Nucleotide substitutions, like Adenine to Thymine, are reported for point mutations while the numbers of base pairs inserted and/or deleted are reported for mutations that are described as insertion, small deletion, or indel. The position of the nucleotide alteration within the DNA/cDNA and the associated reference sequence as well as the corresponding location within the gene (such as exon, intron, splice acceptor site, promoter) are reported when these details are provided. Complex mutations, where there are multiple substitutions, are reported in notes.

Nucleotide changes can have multiple effects on the resulting mutant transcript producing missense or splicing variants that can encode a frameshift or premature stop for example. These changes can be described by multiple transcript consequences. Terms from the SO (20) are used to annotate these sequence variants. A single mutation can have different effects on multiple transcripts originating from a single gene. ZFIN does not currently report the effect of mutations on each specific transcript.

Ultimately, nucleotide changes and transcript alterations can result in modifications at the protein level. The ‘Protein consequence’ section of the mutation details includes these changes such as protein truncation, amino acid insertion, or amino acid deletion. Amino acid sequence modification, such as amino acid change (e.g. Cys > Arg, Tyr > Stop) and the number of amino acids inserted or deleted are also reported, as well as the position of this modification and its sequence of reference when available. Information that cannot be effectively annotated into a particular section (for example protein domains affected by the mutation) remains in the notes.

Currently mutation details are only recorded for point mutations, small deletions, insertions, indels, and transgenic insertions where the insertion site in a gene is known. It is important to note that mutation details as described above are rarely reported in their entirety. In addition to being displayed on the genomic feature page, mutation consequences for a particular allele are also displayed on the associated gene page, where a detailed view of known mutations for a single gene is provided (Figure 6B). Transcript consequences are also displayed in single box search results in the Mutants and Transgenics category, and can be used to filter the search results.

**UPDATES TO SINGLE BOX SEARCH**

The ZFIN single box search was updated to a Solr-backed faceted interface in October 2014 (15). Since that initial release, there have been numerous small updates and several major additions. Here we describe two major new features that were recently added: The Reporter Genes category and the Figure Gallery.

**Reporter lines category**

Reporter lines have become indispensable tools for a wide array of zebrafish research projects. However, locating a reporter line that meets specific criteria for the reporter gene, the promoter driving the reporter expression, or the anatomical location and developmental time at which the reporter is expressed can be challenging. To help address this need, the ZFIN single box search now has a ‘Reporter Lines’ category. This Category is limited to lines that express a reporter gene, such as EGFP or mCherry, and excludes lines those lines with any mutant alleles, sequence targeting reagents, or experimental treatments. Four parameters are provided to further refine reporter line searches; reporter gene, anatomical structure, transgene regulatory region, and reporter expression developmental stage. Together, these filters readily support inquiries such as ‘Are there any reporter lines that express mCherry in Meckel’s cartilage during the larval stage?’ ZFIN currently includes two such lines: hu5910Tg and nu15Tg (Figure 7). Queries involving a specific transgene regulatory region are also supported. For example, ‘Are there any reporter lines that use the regulatory region of the agr2 gene to drive expression of any reporter in the anterior crista?’ ZFIN currently includes one reporter line meeting those criteria: as11Tg. The search results provide links to all the expression data for each reporter line as well as direct access to associated figures. The new figure gallery (discussed below) provides another way to rapidly review images of reporter gene expression.

**Figure gallery**

Quick and easy access to images is a favorite feature for ZFIN users. In response to user input, a new figure gallery has been added to the single box search interface. All single box search results that return images now include a button to ‘Browse Images’. The image browsing view is a familiar
web paradigm for surveying sets of images in a tiled format (Figure 8). Clicking on a figure in the tiled format displays a magnified view that includes curated figure summary data. Left and right arrows allow navigation through the image set in the magnified view format.

DISCUSSION

ZFIN aims to provide the best support possible for the zebrafish research community and for the broader biomedical community having an interest in the data, tools, and resources that zebrafish research generates. Expanded support for curated zebrafish models of human disease will be especially useful to biomedical researchers.

A great strength of zebrafish as a research tool is their transparency during early development, allowing the detailed observation of gene expression patterns in wild-type and mutant fish embryos and larvae. New features have been added to ZFIN to support curation, search and display of phenotypic expression patterns.

As mutagenesis techniques have evolved, zebrafish researchers have asked ZFIN to provide more details about mutations and their consequences. In response to this, more details about the nature of mutations and their molecular consequences is now provided in a structured format. These new data are curated from literature and added via data loads. Results of mutation searches can be filtered by the transcript consequences. These data will provide an additional way for researchers to evaluate the utility of a specific mutant for their research.

Searching for data is a critical component of any on-line data resource. A new ‘Reporter Line’ Category has been added to the ZFIN single box search to streamline and simplify searches for reporter lines with particular attributes of interest. Images are an important part of many searches. The new Figure Gallery feature brings simple image browsing to the single box search. These new additions to ZFIN will help members of the research community find data of interest and generate new hypotheses.

The ZFIN team strives to have as complete a set of data as possible. Community input is always welcomed to help supplement the data in ZFIN. New information regarding disease models, gene expression phenotypes, and mutation details are particularly relevant now that support has been added for those new features. To make a contribution, contact us at zfinadmin@zfin.org or use the ‘Your Input Welcome’ button found on the top right of every page in ZFIN.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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REFERENCES


