Use of zebrafish models to investigate rare human disease

Kathryn Isabel Adamson,1 Eamonn Sheridan,2 Andrew James Grierson1,3

ABSTRACT

Rare diseases are collectively common and often extremely debilitating. Following the emergence of next-generation sequencing (NGS) technologies, the variants underpinning rare genetic disorders are being unearthed at an accelerating rate. However, many rare conditions lack effective treatments due to their poorly understood pathophysiology. There is therefore a growing demand for the development of novel experimental models of rare genetic diseases, so that potentially causative variants can be validated, pathogenic mechanisms can be investigated and therapeutic targets can be identified. Animal models of rare diseases need to be genetically and physiologically similar to humans, and well-suited to large-scale experimental manipulation, considering the vast number of novel variants that are being identified through NGS. The zebrafish has emerged as a popular model system for investigating these variants, combining conserved vertebrate characteristics with a capacity for large-scale phenotypic and therapeutic screening. In this review, we aim to highlight the unique advantages of the zebrafish over other in vivo model systems for the large-scale study of rare genetic variants. We will also consider the generation of zebrafish disease models from a practical standpoint, by discussing how genome editing technologies, particularly the recently developed clustered regularly interspaced repeat (CRISPR)/CRISPR-associated protein 9 system, can be used to model rare pathogenic variants in zebrafish. Finally, we will review examples in the literature where zebrafish models have played a pivotal role in confirming variant causality and revealing the underlying mechanisms of rare diseases, often with wider implications for our understanding of human biology.

THE IMPORTANCE OF RESEARCHING RARE GENETIC DISEASES

Rare genetic diseases commonly manifest as chronically debilitating illnesses, which often dramatically diminish life expectancy and quality of life.1 Although such diseases are individually rare, defined as having a prevalence lower than 1 in 2000,2 the term ‘rare disease’ has been ascribed to over 6000 distinct disorders.3 Collectively, these conditions affect around 30 million people in the European Union alone,4 thus posing a tremendous societal burden.5 Approximately 80% of rare diseases have a genetic aetiology.6 The advent of next-generation sequencing (NGS) technologies has made it possible to rapidly and cost-effectively sequence large regions of DNA in an unbiased manner, which has been especially advantageous for diagnosing rare genetic diseases. This has triggered rapid progress in the identification of their causative mutations in recent years,7 and several collaborative large-scale initiatives, comprising national and international networks of clinicians and researchers, have now been established with the aim of accelerating rare disease gene discovery by NGS (table 1).

Despite these advances, treatments for most rare diseases remain scarce.8 For ‘ultrarare’ diseases, in which candidate variants have only been identified in a handful of individuals within a single pedigree, validation of pathogenicity can be challenging. Additionally, because translational research efforts have historically been more focused on common disorders,2 the underlying mechanisms contributing to rare diseases often remain poorly understood, even where the genetic aetiology has been confirmed.

Considering the growing number of candidate rare disease variants being identified through NGS, there is an increasing need for research into these diseases. Such research will promote insight into their pathogenic mechanisms, helping to accelerate therapeutic development and alleviate the collective burden created by rare illnesses. Moreover, research to reveal the causes of rare conditions can often illuminate fundamental pathways governing human biology, which may advance our understanding of more common disorders.8 For example, the rare metabolic condition, Gaucher’s disease, is caused by homozygous mutations in GBA, resulting in deficiency of the β-glucocerebrosidase protein.9 Interestingly, some heterozygous GBA variants confer risk for the comparatively common neurodegenerative disorder, Parkinson’s disease.10 This genetic link between two clinically disparate diseases has revealed a role for β-glucocerebrosidase deficiency in some forms of Parkinson’s disease.11 Because of this, clinical trials of ambroxol, a pharmaceutical compound that restores the trafficking and enzymatic activity of β-glucocerebrosidase,12 are currently underway for Parkinson’s disease (NCT02941822; NCT02914366).

MODELLING RARE GENETIC DISEASES

Progress in the study of rare diseases necessitates the development of experimental models in which candidate variants can be validated and disease mechanisms can be explored. Information about the functionality of variants can often be gleaned using in vitro or cellular approaches, or through analysis of patient tissue samples.13–16 However, these approaches are insufficient to demonstrate pathogenicity at the whole organism level. For
many rare diseases, in vivo models are therefore required to confirm causality, especially for neurological disorders where pathologically relevant patient tissue is often inaccessible, or for conditions where interactions between multiple cell types or organ systems are required for a disease phenotype to manifest. A caveat to this is that some mutations may have species-specific effects, so the presence or absence of a phenotype in a model organism must be interpreted carefully. For example, several murine cystic fibrosis models fail to accurately recapitulate key respiratory features of the human phenotype, despite harbouring the pathogenic deletion mutation present in most patients.20 21

Besides the capacity to validate candidate genes, in vivo models of rare disease offer further opportunities to dissect disease mechanisms within biologically relevant systems, and to explore the phenotypic consequences of therapeutic intervention. It is important that this can be done at scale, considering initiatives such as the 100 000 Genomes Project, which aim to reveal the genetic causes of rare diseases in thousands of patients.22

In vivo models for rare genetic disease must be genetically tractable, with a fully sequenced genome that shares substantial homology with the human genome and an amenability to genetic manipulation on a whole organism scale. Historically, the mouse (Mus musculus) has been considered a pre- eminent model organism for human genetic disease, but mice would be impractical for validating the large number of candidate variants that are likely to be identified through large-scale rare disease programmes, due to their high husbandry costs and small litter sizes. Invertebrates, such as Drosophila melanogaster and Caenorhabditis elegans, are also commonly used as models for human genetic diseases. These are better-suited to large-scale analyses, but their evolutionary distance from humans and resulting physical differences can make them a less relevant system in which to investigate the physiological consequences of disease-linked mutations and therapeutic interventions.

**ZEBRAFISH AS AN EMERGING MODEL ORGANISM FOR RARE GENETIC DISEASES**

In recent years, the zebrafish (Danio rerio) has become an attractive model organism for translational research. Zebrafish uniquely combine many of the genetic and physiological advantages of mammalian models with the high-throughput capabilities and experimental manipulability of invertebrate models. Their growing popularity is reflected by a continued increase in the use of zebrafish in biomedical research publications (figure 1).

![Figure 1](http://dan.corlan.net/medline-trend.html)

**Figure 1** The use of zebrafish in biomedical research articles has been rising since the early 1990s. The line graph shows the number of publications indexed by PubMed under the term ‘zebrafish’ each year between 1990 and 2016. Raw data obtained from Medline Trend (http://dan.corlan.net/medline-trend.html).

**Genetic and physiological conservation**

The zebrafish genome possesses considerable homology with the human genome, with orthologues having been identified for approximately 70% of human genes. Zebrafish are also highly genetically tractable, and tools for generating genetically modified zebrafish models continue to be developed and optimised. Anatomically and physiologically, the zebrafish is more distantly removed from humans than the mouse. Consequently, it can be more challenging to model genetic diseases affecting structures that are absent in fish, such as the lungs. Nonetheless, the straightforward vertebrate architecture of the zebrafish enables the simplified study of disease in numerous organ systems and structures that are common to both zebrafish and humans. For example, the zebrafish has successfully been used to model genetic diseases affecting the human cardiovascular, nervous, visual, renal and muscular systems, among others.

**Study of genes essential for mammalian placental development**

As a non-placental vertebrate, the zebrafish allows us to study phenotypes linked to genes that are essential for mammalian

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**Table 1** National and international collaborative networks aiming to accelerate research into rare genetic diseases

<table>
<thead>
<tr>
<th>Programme/network name</th>
<th>Locations</th>
<th>Active years</th>
<th>Aims and specific disease interests</th>
<th>Key publications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Finding the Genetic Basis of Learning Disability study</td>
<td>Based at the Cambridge Institute for Medical Research (UK)</td>
<td>2001–present</td>
<td>Identifying genetic causes of X linked intellectual disability</td>
<td>87</td>
</tr>
<tr>
<td>Deciphering Developmental Disorders study</td>
<td>Collaborations across 24 Regional Genetics Services (UK and Ireland) and Wellcome Trust Sanger Institute (UK)</td>
<td>2011–2016</td>
<td>Identifying genetic causes of developmental disorders</td>
<td>88–90</td>
</tr>
<tr>
<td>Finding of Rare Disease Genes Canada Consortium</td>
<td>Network of 21 genetics centres and 3 science and technology innovation centres across Canada, with ad hoc international collaborations across 17 additional countries</td>
<td>2011–2013</td>
<td>Identifying genes associated with rare monogenic diseases with paediatric onset</td>
<td>89</td>
</tr>
<tr>
<td>International Rare Diseases Research Consortium</td>
<td>International network of researchers and organisations with an interest in rare disease research</td>
<td>2011–present</td>
<td>Contributing to the development of novel rare disease therapies and the means to uncover the genetic causes of rare diseases</td>
<td>92</td>
</tr>
<tr>
<td>Undiagnosed Diseases Network</td>
<td>Network of clinical and scientific sites across the USA, coordinated by Harvard Medical School</td>
<td>2014–present</td>
<td>Accelerating identification of genetic causes of rare diseases, with use of Drosophila and zebrafish models to validate candidate genes</td>
<td>93–94</td>
</tr>
</tbody>
</table>
Investigation of developmental processes

The zebrafish additionally offers unparalleled opportunities for the investigation of fundamental developmental processes. Three-quarters of rare diseases arise during childhood and 30% of rare disease patients do not survive past their fifth birthday, so perturbed developmental processes are likely to be implicated in the pathogenesis of many rare conditions. Zebrafish embryos develop externally and rapidly from the one-cell stage, and morphogenesis of most major organ systems is complete within 48 hours of fertilisation. These characteristics, combined with their optical transparency as embryos, and the array of reporter lines and imaging techniques available for detailed visualisation of developmental and physiological processes, make zebrafish immensely useful for exploring the effects of rare disease-linked mutations on vertebrate development.

Large-scale phenotypic screening

Zebrafish are also amenable to large-scale analyses. As adults, their small size and inexpensive husbandry mean that they can be housed in large numbers, but their advantages in large-scale applications are especially apparent at embryonic and larval stages. A pair of adult zebrafish can produce hundreds of eggs in a single clutch, and the resulting embryos and larvae are sufficiently small that they can be housed in 96-well or 384-well plates, allowing phenotypic analysis using high-throughput microscopy and behavioural analysis systems. Soluble compounds can also be dissolved directly into the medium in which the embryos are housed, providing a convenient platform for rapid and efficient screening of small molecule phenotype modifiers once relevant pathophysiological mechanisms have been identified. This positions the zebrafish as an unrivalled tool combining the scalability of in vitro and cell-based assays with a multidimensional capacity to explore disease mechanisms, phenotypes and therapeutic strategies within a living vertebrate.

GENERATING ZEBRAFISH MODELS OF RARE GENETIC DISEASES

Over time, various approaches have been used to model human genetic diseases in zebrafish. Summarised in table 2, these encompass strategies for generating stable mutant lines by random mutagenesis or targeted gene editing, as well as methods for transient interrogation of the effects of altered gene expression. A limitation of this approach is that the base-pair of interest must be located within an optimal window of proximity to a protospacer adjacent motif (PAM). However, efforts are currently underway to overcome this through the development of Cas9

Generation of stable zebrafish disease models

Random mutagenesis

Traditionally, genetic manipulation of zebrafish has focused on random mutagenesis using chemical mutagens such as N-ethyl-N-nitrosourea, or retroviral-mediated insertional methods. Mutations in genes of interest can then be identified, and stable mutant lines can subsequently be generated. This has taken place on a large scale and huge libraries of stable mutants for numerous zebrafish genes are now available. However, mutants are not available for every gene and this method also does not enable specific disease-associated mutations to be modelled, limiting its value for testing the pathogenicity of individual candidate variants.

Targeted gene-editing

More recently, powerful methods involving the use of engineered nucleases, including zinc finger nucleases (ZFNs), transcription activation-like effector nucleases (TALENs) and the clustered regularly interspaced repeat (CRISPR) system have been employed to generate stable models of human disease, enabling targeted mutations to be created in specific zebrafish orthologues of interest. Both ZFNs and TALENs require generation of a tailored protein component for each target locus, which can be a costly and labourious process, making these systems less compatible with large-scale applications. In contrast, the CRISPR/Cas9 system relies on recognition of the target site by a custom guide RNA (gRNA) molecule, and simply requires the design of a single oligonucleotide for each target site.

Using CRISPR/Cas9 technology to model rare genetic diseases in zebrafish

The CRISPR/Cas9 system has therefore become the tool of choice for generating stable models of human genetic diseases in recent years. A mechanistic overview of the classical CRISPR/Cas9 system in the context of disease modelling is shown in figure 2. Most successes from CRISPR-mediated gene modification in zebrafish have arisen from the generation of frameshift null alleles through non-homologous end-joining (NHEJ)-mediated repair of CRISPR-induced DNA breaks. This is useful for generating models of human disease occurring due to LOF alleles and work to further improve the mutagenesis efficiency of this method in zebrafish is ongoing.

It may be desirable to introduce specific disease-associated mutations, particularly where complete LOF alleles are lethal or a variant is predicted to act through a gain-of-function mechanism. Knock-in of specific mutations and exogenous DNA sequences has been achieved in zebrafish through both homology-independent and homology-directed repair (HDR), but this approach is currently less efficient than generation of LOF alleles.

Many genetic diseases are caused by missense mutations, which generally arise from single base-pair substitutions. Novel CRISPR/Cas9-derived gene editing tools, known as base editors, have recently been developed, allowing targeted deaminase-mediated conversion of a single base-pair of interest to another without requiring DNA cleavage. Base editors are currently being assessed for their potential to facilitate modelling of point mutations in various systems, including zebrafish. A limitation of this approach is that the base-pair of interest must be located within an optimal window of proximity to a protospacer adjacent motif (PAM). However, efforts are currently underway to overcome this through the development of Cas9
Methods

Table 2  Summary of methods for generating stable and transient zebrafish models of genetic human diseases

<table>
<thead>
<tr>
<th>Technique</th>
<th>Mechanism</th>
<th>Required components</th>
<th>Advantages for modelling rare genetic diseases in zebrafish</th>
<th>Limitations for modelling rare genetic diseases in zebrafish</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stable models</td>
<td></td>
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</tr>
<tr>
<td>ENU-mediated mutagenesis</td>
<td>Random mutagenesis</td>
<td>Database of mutants that have already been generated by ENU-mediated random mutagenesis</td>
<td>If LOF mutants for genes of interest are available, this abrogates need to generate a new model</td>
<td>Mutants for genes of interest are not always available. Outcrossing is often required to generate a stable model.</td>
</tr>
<tr>
<td>Retroviral-mediated insertional mutagenesis</td>
<td>Random mutagenesis</td>
<td>Database of mutants that have already been generated by retroviral-mediated insertional mutagenesis</td>
<td>If LOF mutants for genes of interest are available, this abrogates need to generate a new model</td>
<td>Mutants for genes of interest are not always available. Outcrossing is often required to generate a stable model.</td>
</tr>
<tr>
<td>ZFNs+NHEJ-mediated repair</td>
<td>Creates double-stranded DNA break at target site, resulting in repair by NHEJ</td>
<td>Multiple DNA-binding zinc finger peptides (which each recognise 3bp of target DNA) fused to FokI nuclease domain</td>
<td>Enables targeted frameshift mutations to be introduced in candidate genes of interest</td>
<td>Tailored protein component needs to be generated for each genomic target. Outcrossing is often required to generate a stable model.</td>
</tr>
<tr>
<td>TALENs+NHEJ-mediated repair</td>
<td>Creates double-stranded DNA break at target site, resulting in repair by NHEJ</td>
<td>Customisable DNA-binding domain (peptide-based) fused to FokI nuclease domain</td>
<td>Enables targeted frameshift mutations to be introduced in candidate genes of interest</td>
<td>Tailored protein component needs to be generated for each genomic target. Outcrossing is often required to generate a stable model.</td>
</tr>
<tr>
<td>CRISPR/Cas9+NHEJ-mediated repair</td>
<td>Creates double-stranded DNA break at target site, resulting in repair by NHEJ</td>
<td>Specific 20 nt guide RNA complementary to target site+Cas9 endonuclease</td>
<td>Enables targeted frameshift mutations to be introduced in candidate genes of interest</td>
<td>gRNAs can easily be designed for different targets. LOF models can be efficiently generated through NHEJ-mediated repair</td>
</tr>
<tr>
<td>ZFNs, TALENs or CRISPR/Cas9+HDR-mediated repair</td>
<td>Creates double-stranded DNA break at target site. Simultaneous addition of DNA repair template results in HDR and incorporation of specific sequences or mutations of interest</td>
<td>Zinc finger peptides or customisable peptide-based DNA-binding domain fused to FokI nuclease domain, or specific – 20 nt guide RNA complementary to target site+Cas9 endonuclease (+DNA repair template containing sequence of interest)</td>
<td>Allows knock-in of specific mutations of interest (most commonly via CRISPR/Cas9)</td>
<td>Currently less efficient than NHEJ-mediated LOF model generation. Outcrossing is often required to generate a stable model.</td>
</tr>
<tr>
<td>CRISPR/Cas9-mediated base-editing</td>
<td>Deaminates cytidine or adenine bases at genomic target site, resulting in single base-pair substitutions</td>
<td>Specific – 20 nt guide RNA complementary to target site+catalytically inactive Cas9 (dCas9), fused to cytidine or adenine deaminase enzyme</td>
<td>Allows introduction of disease-relevant missense mutations arising due to C-T or A-G base substitutions</td>
<td>Some base-pair substitutions cannot be modelled using this approach. Evidence of efficacy in zebrafish is limited. Specificity for target site needs to be established.</td>
</tr>
<tr>
<td>Transient models</td>
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<tr>
<td>Morpholino knockdown</td>
<td>Blocks mRNA translation or splicing (post-translational)</td>
<td>Synthetic 25 bp oligonucleotide</td>
<td>Allows for rapid examination of LOF phenotypes. Could be used to rapidly obtain evidence to support causality of LOF candidate variants</td>
<td>Effects are short-lasting. May be associated with significant off-target effects. Cannot be used to model gain-of-function or patient-specific mutations.</td>
</tr>
<tr>
<td>CRISPR/dCas9 (CRISPR interference)</td>
<td>Blocks transcription (and can be coupled to transcriptional activators or repressors to further control gene dosage)</td>
<td>Specific – 20 nt gRNA complementary to target site+dCas9 (which can be fused to a transcriptional activator or repressor)</td>
<td>Can be used to model both gain and LOF phenotypes. The potential to be used on a large scale</td>
<td>Currently not widely used in zebrafish. Specificity for target site needs to be established.</td>
</tr>
</tbody>
</table>

 Cas9, CRISPR-associated protein 9; CRISPR, clustered regularly interspaced repeats; dCas9, catalytically inactive Cas9; ENU, N-ethyl-N-nitosourea; HDR, homology-directed repair; LOF, loss-of-function; NHEJ, non-homologous end joining; TALENs, transcription activation-like effector nucleases; ZFNs, zinc finger nucleases.

Methods for transiently modelling genetic disease in zebrafish

Transient genetic manipulation of zebrafish may also be useful where complete LOF of a gene of interest is undesirable. This has traditionally been accomplished through morpholino-mediated knockdown of zebrafish orthologues of interest; however, concerns have been raised about the on-target specificity of morpholinos.70 CRISPR interference (CRISPRi), another variation on the CRISPR/Cas9 system, can be used to reversibly silence expression of a target gene at the transcriptional level.71 The CRISPRi machinery can also be coupled to transcriptional repressors or activators to alter gene dosage, allowing LOF or gain-of-function variants to be reversibly mimicked.72 Most examples of the use of this technology have come from cellular experiments,73 74 although CRISPRi has successfully been used to silence gene expression in zebrafish.75 As CRISPRi is developed further, it is expected to offer additional possibilities for modelling human genetic diseases in zebrafish.75

Using zebrafish to model rare genetic diseases

The benefits of the zebrafish as a model for rare genetic disease are illustrated by several examples in the literature. Here, we
Methods

Overview of clustered regularly interspaced repeat (CRISPR)/CRISPR-associated protein 9 (Cas9)-mediated genome editing. A 20-nucleotide guide RNA (gRNA) directs Cas9 endonuclease to a chosen genomic target site (also known as the protospacer) immediately upstream of a protospacer-adjacent motif (PAM). On binding to the target site, Cas9 cleaves the genomic DNA to create a double-stranded DNA break 3–4 bp upstream of the PAM. Subsequent DNA repair via the non-homologous end joining (NHEJ) or homology-directed repair (HDR) pathways can be exploited to generate disease models through the creation of knockout alleles (to investigate loss-of-function of a gene of interest), or knock-in alleles (to study the consequences of a specific disease-associated mutation). For HDR approaches, a tailored repair template containing the variant of interest is also required.

review examples where zebrafish models of rare diseases have been instrumental in confirming the pathogenicity of candidate variants, for elucidating disease mechanisms, and in the functional annotation of rare disease genes.

Confirming causality of candidate rare disease-associated variants

The value of zebrafish models in supporting the causality of candidate rare disease variants is exemplified by the identification of several novel genes linked to Hirschsprung disease. Hirschsprung disease is a rare disorder characterised by aberrant development of enteric neurons, resulting in an absence of innervation in parts of the colon. Whole exome sequencing (WES) analysis of 24 trios, comprised of patients with Hirschsprung disease and their parents, led to the discovery of several novel LOF and missense variants in genes with no prior links to either enteric nervous system development or Hirschsprung disease pathogenesis.18 Given an absence of clues that would substantiate the pathogenicity of these variants in patients, the authors investigated the effects of LOF of zebrafish orthologues for 12 of these candidate genes on enteric nervous system development or Hirschsprung disease pathogenesis.18 Given an absence of clues that would substantiate the pathogenicity of these variants in patients, the authors investigated the effects of LOF of zebrafish orthologues for 12 of these candidate genes on enteric nervous system development or Hirschsprung disease pathogenesis.18 Given an absence of clues that would substantiate the pathogenicity of these variants in patients, the authors investigated the effects of LOF of 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Increasing understanding of rare disease pathophysiology

Zebrafish models have also successfully been used to reveal pathogenic mechanisms in rare genetic diseases. Diamond-Blackfan anaemia (DBA) is a rare disorder affecting red blood cell production, which often arises due to mutations in ribosomal proteins. Mutations in RPS19, encoding the ribosomal protein S19, account for most DBA cases. To clarify the contribution of ribosomal protein deficiency to the phenotype observed in patients with DBA, TALENs were used to generate a knockout model for the zebrafish orthologue, rps19.76 This model mimics pathogenic features of DBA, including impaired haematopoiesis and p53 activation. The authors also demonstrated that globin protein production is impaired in rps19-null mutants, with negligible effect observed at the mRNA level. This was replicated in an additional zebrafish model of DBA, harbouring a null mutation in rpl11. By generating a transgenic zebrafish reporter line exhibiting erythroid-specific mCherry expression, they determined that the defective globin production was likely a result of faulty protein production in these cells rather than a pathogenic effect specific to globin genes.76

Previously, zebrafish rps19 morphants were treated with L-leucine on the basis of its known stimulatory effects on protein synthesis.77 This improved their anaemic phenotype through activation of the mTOR pathway.76 Correspondingly, L-leucine treatment of rps19 and rpl11 mutants ameliorated globin protein production with partial phenotypic rescue, suggesting

Figure 2  Overview of clustered regularly interspaced repeat (CRISPR)/CRISPR-associated protein 9 (Cas9)-mediated genome editing. A 20-nucleotide guide RNA (gRNA) directs Cas9 endonuclease to a chosen genomic target site (also known as the protospacer) immediately upstream of a protospacer-adjacent motif (PAM). On binding to the target site, Cas9 cleaves the genomic DNA to create a double-stranded DNA break 3–4 bp upstream of the PAM. Subsequent DNA repair via the non-homologous end joining (NHEJ) or homology-directed repair (HDR) pathways can be exploited to generate disease models through the creation of knockout alleles (to investigate loss-of-function of a gene of interest), or knock-in alleles (to study the consequences of a specific disease-associated mutation). For HDR approaches, a tailored repair template containing the variant of interest is also required.
that L-leucine may activate translation in erythroid cells. Here, zebrafish models have progressed understanding of DBA pathophysiology, and present an in vivo system that could be used to further investigate pathogenesis and screen for possible therapeutic modifiers of the disease phenotype.

**Functional annotation of rare disease-associated genes**

As well as their specific advantages in relation to the study of rare diseases, zebrafish models of rare genetic conditions can also contribute novel insights into the biological pathways and processes in which rare disease genes are implicated. These findings have value in a translational context and may ultimately improve our understanding of more common disorders.

Identification of a novel pathway involved in human bone formation

Members of the vacuolar ATPase (V-ATPase) protein family form multicomponent complexes that are responsible for regulating intracellular and extracellular pH through acidification. This acidification process is especially important in osteoclasts, where it critically regulates proper bone resorption. Accordingly, mice deficient for the V-ATPase gene \( \text{Atp6i} \) exhibit osteopetrosis (increased bone density), which is underpinned by impaired bone resorption due to deficient extracellular acidification by osteoclasts. Through the NIH Undiagnosed Diseases Programme, a rare heterozygous mutation in the \( \text{Atp6v1h} \) gene (encoding another member of the complex) was found to segregate with a phenotype of osteoporosis (reduced bone density) in three generations of a single pedigree. The human disease-associated mutant \( \text{Atp6v1h} \) protein was shown to be less stable than the wild-type protein when expressed in HEK cells, suggesting that \( \text{Atp6v1h} \) haploinsufficiency may contribute to pathogenesis; this was unexpected given the contrasting phenotypes of the patients and \( \text{Atp6i} \)-null mice.

A zebrafish model harbouring a CRISPR-induced null mutation in \( \text{atp6v1b} \) was developed to investigate the consequences of loss of \( \text{Atp6v1h} \) function on bone homeostasis. At 6 days post-fertilisation, bone mineralisation was reduced in homozygous null mutant embryos compared with wild-type zebrafish and this was not rescued by injection of mRNA harbouring the pathogenic mutation from patients with osteoporosis. Heterozygous mutants exhibited no obvious skeletal defects until adulthood, at which point decreased bone density, volume and surface area became apparent, along with a marked reduction in calcification of the vertebral centrum. Thus, this model recapitulates the osteoporotic phenotype observed in patients with the \( \text{Atp6v1h} \) mutation and supports a role for \( \text{Atp6v1h} \) deficiency in the disease process.

The authors additionally explored the pathways linking \( \text{atp6v1b} \) deficiency to aberrant bone homeostasis in their zebrafish model. They discovered that \( \text{atp6v1b} \) deficiency results in elevated transcript levels of the osteoclast marker, matrix metalloproteinase-9 (\( \text{Mmp9} \)), and the authors investigated whether \( \text{atp6v1b} \) deficiency also led to abnormal expression of the osteoclast marker, \( \text{Southpaw} \) (\( \text{Spaw} \)), indicating that the looping defects are linked to aberrant control of left-right asymmetry. This supports the causality of the \( \text{LOF Mmp21} \) mutation in patients with heterotaxia and demonstrates involvement of MMPs in the establishment of organism polarity.

Further cell-based investigation of the connection between \( \text{Mmp21} \) and control of laterality revealed that \( \text{Mmp21} \) likely acts as a negative regulator of the Notch signalling pathway, which is required for establishing left-right asymmetry. Several target genes of Notch signalling were upregulated in the zebrafish \( \text{mmp21} \) morphants, emphasising the contribution of this pathway to \( \text{Mmp21} \)-mediated control of laterality.

Here, a zebrafish model of a rare genetic disease has advanced our understanding of both the underlying pathophysiology of this rare instance of heterotaxia and of the fundamental pathways involved in establishing vertebrate polarity. Specifically, the zebrafish has exposed a previously undiscovered role for MMPs in this process.

**CONCLUSION**

Recent advances in DNA sequencing technology and the introduction of large-scale rare disease gene discovery programmes mean that a growing number of candidate disease-associated variants are being rapidly identified for many rare disorders. Combined with the challenging nature of confirming pathogenicity of candidate variants for rare diseases affecting small numbers of patients, this has resulted in a bottleneck in the validation of these variants and elucidation of underlying disease mechanisms. This is a critical hurdle to overcome, as a better understanding of these rare conditions will improve quality of life for rare disease patients by facilitating therapeutic development and will also advance our knowledge of human biology and the mechanisms of more prevalent diseases through the functional annotation of rare disease genes.

Progress in the identification and validation of rare disease variants is of particular relevance to individuals born to consanguineous parents—a population whose risk of congenital anomalies is doubled. These abnormalities frequently arise from autosomal recessively inherited variants, and NGS-based approaches are increasingly revealing the rare genetic mutations responsible for diseases in these families. Such conditions...
often present great obstacles with regard to validation of variant pathogenicity; in many cases they are extremely rare, with mutations only reported in a single pedigree.

Here we present the zebrafish as an ideal in vivo model for addressing this vast pool of candidate rare disease genes. With a highly conserved vertebrate genome and a capacity for large-scale genetic manipulation, the zebrafish is well-positioned as a tool for modelling a substantial proportion of rare genetic variants and the scope for modelling these mutations in zebrafish is likely to expand with continued technological advances in the field of CRISPR/Cas9-mediated genome-editing. Indeed, large-scale initiatives, such as the NIH Undiagnosed Diseases Network (table 1), have now been developed with the aim of using zebrafish to assist in the validation of rare disease variants to uncover the genetic causes of previously undiagnosed rare disorders. Moreover, zebrafish can be used in large-scale screening of phenotypes associated with many conserved vertebrate organ systems and structures and they have a capacity for high-throughput testing of therapeutic compounds that is often impractical in other vertebrate model systems. This unique combination of attributes renders the zebrafish unparalleled by other classical model systems in its potential for advancing our understanding of rare genetic diseases.

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