Review Article

Think Small: Zebrafish as a Model System of Human Pathology

J. R. Goldsmith¹ and Christian Jobin¹,²,³

¹ Department of Pharmacology, University of North Carolina, Chapel Hill, NC 27599, USA
² Department of Medicine, University of North Carolina, Chapel Hill, NC 27599, USA
³ Department of Microbiology and Immunology, University of North Carolina, Chapel Hill, NC 27599, USA

Correspondence should be addressed to Christian Jobin, job@med.unc.edu

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Although human pathologies have mostly been modeled using higher mammal systems such as mice, the lower vertebrate zebrafish has gained tremendous attention as a model system. The advantages of zebrafish over classical vertebrate models are multifactorial and include high genetic and organ system homology to humans, high fecundity, external fertilization, ease of genetic manipulation, and transparency through early adulthood that enables powerful imaging modalities. This paper focuses on four areas of human pathology that were developed and/or advanced significantly in zebrafish in the last decade. These areas are (1) wound healing/restitution, (2) gastrointestinal diseases, (3) microbe-host interactions, and (4) genetic diseases and drug screens. Important biological processes and pathologies explored include wound-healing responses, pancreatic cancer, inflammatory bowel diseases, nonalcoholic fatty liver disease, and mycobacterium infection. The utility of zebrafish in screening for novel genes important in various pathologies such as polycystic kidney disease is also discussed.

1. Introduction

Investigators have long utilized reductionist systems and animal models to mimic and study basic processes regulating cellular biology, organ function, and host homeostasis. While much work has been accomplished and continues to be undertaken in higher mammalian systems such as mice, rats, and rabbits, important discoveries have also been made using invertebrate systems such as C. elegans and Drosophila melanogaster. For example, RNA interference technology was first discovered in C. elegans [1], as was the initial caspase enzyme, caspase-1 (ced-3 in C. elegans) [2]. Similarly, it was in Drosophila melanogaster that the innate signaling Toll-like receptors (TLRs) were first identified via the discovery of the Toll gene, as was its linkage to the nuclear factor kappa-light-chain-enhancer of activated B cells (NF-κB) signaling cascade [3]. In the last ~15 years, there has been a growing appreciation for the vertebrate organism Danio rerio (zebrafish) as a tool to study human disease [4]. As opposed to C. elegans and Drosophila melanogaster, zebrafish is a vertebrate organism with physiological and anatomical characteristics of

its higher organism counterparts (see Figure 1 for a diagram of larval zebrafish anatomy), while maintaining the ease of use of a lower organism. The characteristics of zebrafish that make them desirable tools for the study of embryogenesis/development also make them useful for the study of human pathologies. Zebrafish have a fully mapped genome (http://www.sanger.ac.uk/Projects/D_rerio/), which has significant homology with the human genome, including noncoding regions (~60 per gene globally across all genes) [5] suggesting that numerous genes involved in human diseases could be matched in the zebrafish genome. Reverse (morpholino knock-down, Targeting Induced Local Lesions in Genome—TILLING [6]) and forward genetic (mutagenesis, transgenic) approaches are well established and commonly used to manipulate and characterize zebrafish gene function.

Zebrafish are highly fecund and breed rapidly; a pair of zebrafish produces over 100 embryos per clutch that are usable for larval experiments as early as 3 days post fertilization (dpf). These larvae are transparent through 7 dpf, and this can be extended to up to 9–14 dpf with the addition of the melanocyte inhibitor phenylthiourea. Moreover, the
recent generation of transparent adult zebrafish such as the Casper line adds new imaging possibilities [7]. The transparency of zebrafish, in conjunction with sophisticated utilization of fluorescent technology to mark signaling proteins or cellular entities, allows for powerful time-lapse imaging of biological and disease processes. Additionally, the vertebrate zebrafish has many features commonly found in mammals, including an innate immune system composed of neutrophils, NK cells, and monocyte/macrophages with functionality by 48 hours post fertilization (hpf) [8, 9] and an adaptive immune system that is fully functional at 4–6 weeks post fertilization [10]. The adaptive immune system is highly analogous to that of mammals, with T cells and B cells that have Rag-dependent V(D)J recombination (reviewed extensively in [9]). Finally, the zebrafish research community benefits from an up-to-date database of techniques, genetic strains, and other useful resources at http://zfin.org/.

In this paper, we focus our discussion on larval and adult zebrafish models that recapitulate human diseases, focusing on four separate branches of pathology: wound healing/restitution, gastrointestinal disease, microbe-host interactions, and genetic diseases and drug screens.

2. Wound Healing/Restitution

Wound healing represents a critical biological response of injured tissues and organs. Events causing epithelial injury and barrier breakdown initiate a biological response known as “restitution,” which is aimed at resealing the damaged region and reestablishing host homeostasis. This “wound healing” response involves migration of epithelial cells toward the injured regions as well as epithelial cell proliferation to replenish the cell pool. Understanding the cellular and molecular mechanisms regulating this response could have profound translational impact for patients suffering from chronic inflammation, ischemia/hypoxia, burn injury, and cancer. The powerful imaging modalities available to zebrafish researchers alongside their ease of genetic manipulation make this vertebrate system an ideal model for studying wound healing response to various injuries [11]. Additionally, the ability of zebrafish to regenerate both limbs and cardiac tissue [12] makes them a powerful animal model for understanding the molecular mechanisms involved in regenerative signaling.

The most popular zebrafish injury model is the larval tail wounding model, where a segment of the tail fin is resected. Using this injury model with transgenic zebrafish expressing EGFP under the transcriptional controlled of the neutrophil-specific myeloperoxidase (MPO) promoter—Tg(BACmpx:GFP)i114—investigators studied real-time neutrophil chemotaxis to the site of injury [13]. In this study, they observed retrograde chemotaxis of neutrophils toward the vasculature alongside where injury resolution was observed, suggesting for the first time that retrograde chemotaxis may play an important role in the resolution phase of the inflammatory response [13]. To determine the function of the reactive oxygen species H2O2 in the wound healing response, investigators injected mRNA encoding for the hydrogen peroxide sensor HyPer to one-cell stage developing zebrafish embryos [14]. Upon tail wounding of 3 dpf larvae, the HyPer sensor demonstrated increased H2O2 production along a gradient decreasing away from the wound site, which signaled leukocyte chemotaxis to the injured location [11]. Using pharmacological and morpholino blockade, the authors showed that generation of the H2O2 gradient is dependent on the activity of the dual oxidase (duox) gene. This HyPer reporter system has also been utilized in the study of neuronal regeneration, an important area of study for the treatment of many human diseases. H2O2 produced by injured keratinocytes was shown to induce somatosensory axonal regeneration in zebrafish. Similar to the tail fin wounding model, neuronal regeneration required activation of duox [15]. Moreover, H2O2 administration promoted axonal regeneration following neuronal injury, even without accompanying keratinocyte injury [15]. These results expand the understanding of posttraumatic nerve injury and subsequent loss of limb function in humans. The healing-enhancing properties of H2O2 have since been extended to studies in both rabbits [16] and horses [17] as well as one reported case study in a human patient [18].

Beyond cutaneous wounds, zebrafish have been used for the ability to regenerate cardiac tissue. Unlike mammals, which form scars and do not regenerate cardiac tissue following injury, zebrafish are able to fully regenerate their heart within 2 months after 20% ventricular resection [19]. This ability for zebrafish to overcome scar formation requires activation of Mps1, a mitotic checkpoint kinase [19]. Additionally, cardiac regeneration involves upregulation of wound healing genes and proliferative factors including apoeb, vegcf, and granulina (all of which have human analogs) and likely activation of platelet-derived growth factor B signaling [20]. Cardiac regeneration studies in zebrafish could produce novel paradigms and identify unique pathways/targets with profound translational impact for patients suffering from ischemic heart disease.

3. Gastrointestinal Diseases

The zebrafish gastrointestinal system is highly homologous to that of mammals, containing a liver, pancreas, gall bladder, and a linearly segmented intestinal track with absorptive and secretory functions [21, 22]. The intestinal epithelium displays proximal-distal functional specification and
Table 1: Zebrafish gastrointestinal models of pathology.

<table>
<thead>
<tr>
<th>Model</th>
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<td>2% ethanol to the water of 4 dpf zebrafish for 32 days</td>
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Abbreviations: GI: gastrointestinal; APC: adenomatous polyposis coli; HCV: hepatitis C virus; TNBS: 2,4,6-trinitrobenzene sulfonic acid; IBD: inflammatory bowel diseases; NAFLD: nonalcoholic fatty liver disease; ER: endoplasmic reticulum.

contains many of same epithelial cell lineages found in mammals including absorptive enterocytes, goblet cells, and enteroendocrine cells [21, 22]. Enterocytes have a basolateral nuclei and form tight junctions, apical microvilli, and an intestinal brush border [23]. In the last decade, numerous gastrointestinal pathologies have been modeled in the zebrafish, as summarized in Table 1.

Interestingly, a forward genetic mutagenesis screen (N-ethyl-N-nitrosourea (ENU), see section five) generated zebrafish mutants that develop spontaneous intestinal, pancreatic, and hepatic neoplasias [24]. These mutants were later found to be heterozygotic for a truncated form of APC, thereby leading to accumulation of nuclear β-catenin and increased expression of downstream genes such as cmyc and axin2 [24]. Adding the carcinogen 7, 12-dimethylbenz[a]anthracene to these mutant zebrafish increases the frequency of tumor development. Since germline-truncated APC mutations in both humans [34, 35] and mice [36] result in the spontaneous development of a large numbers of intestinal polyps, the APC zebrafish model could be useful in genetic, drug screening, and toxicology studies.

There are roughly 24,000 new cases of hepatocellular carcinoma (HCC) annually in the United States, resulting in over 18,000 deaths per year [37]. Worldwide, the disease incidence can be as much as 30-fold higher due to the increased prevalence of hepatitis infections [38]; globally estimated 564,000 new cases occur per year, accounting for 5.6% of all human cancers [38], and resulting in similar rates of mortality as seen in the USA [38]. Oncogenic profiling of liver tumors showed a significant overlap between human and zebrafish in 132 genes [25]. These genes included those involved in β-catenin and Ras-MAPK signaling pathways as well as genes implicated in cellular adhesion, apoptosis, and liver-specific metabolism found in both organisms [25]. HCC can be induced in zebrafish with the liver toxin thioacetamide, resulting in HCC-related pathology within 12 weeks of exposure. The timeline of thioacetamide-induced HCC could be significantly accelerated (6 versus 12 weeks) by using transgenic fish expressing the hepatitis C virus (HCV) core protein (HCP-transgenic fish) [26]. The increasing prevalence of HCC in humans has been attributed to the rise of HCV infection [39], in particular through the oncogenic action of the HCV core protein [40], making this model particularly relevant to human disease.

Pancreatic cancer represents the fourth most common cause of cancer-related mortality in the western world, likely due to limited diagnostic tools and inability to survey the disease. Exocrine pancreatic cells are responsible for more
than 95% of pancreatic cancer [37]. The cell population in the pancreas that gives rise to human exocrine pancreatic cancer remains unknown and investigators have turned to zebrafish models to investigate this important question. Transgenic zebrafish were generated using a bacterial artificial chromosome (BAC) approach that expressed EGFP-KRAS fusion protein under the control of the zebrafish pancreatic locus ptfla (Tg(ptfla:EGFP-Hsa.KRASG12V)jh7 [27]). Using these fish, investigators found that while normal pancreatic cell progenitors differentiated normally, KRAS-transgenic fish had blocked cellular differentiation, producing a pool of undifferentiated progenitors that progressed to invasive pancreatic cancer over a course of 3 to 9 months. These cancer cells also had increased expression of multiple genes.

The inflammatory phenotype was reduced by the administration of the gram-positive-specific antibiotic vancomycin, with human IBD [53]. In humans, IBD is the result of dysregulated interactions between a genetically susceptible host and their commensal gut microbiota [41–43]. IBD is primarily a disease of the Western world; it affects 1.4 million Americans and there is no cure for the disease [44]. Outside of North America, similar disease burdens are seen in Europe but are ∼3-fold lower in Asia and the Middle East, with the exception of Japan [45]. Pharmacological treatment centers around the use of anti-inflammatories and immunosuppressants, and surgical management of the disease is also possible though not always curative [46]. Numerous chemical and genetic approaches are currently used by the IBD research community to study this pathology. These models mimic various aspects of the disease, such as induced barrier function and impaired innate and adaptive immune responses [47, 48]. Among the chemical models, the hapten oxazolone has been used for studies of acute intestinal inflammation. This murine model consists of rectal administration of a single dose of oxazolone dissolved in ethanol, resulting in an acute colitis lasting up to 10 days, with peak inflammation at 2 days post administration in mice [49]. The model is characterized by a strong Th2-dependent immune response that can be abrogated by IL-4 neutralizing antibodies and has similar characteristics to human ulcerative colitis [48].

Investigators have tried to adapt the hapten oxazolone model of colitis to adult zebrafish. Brugman et al. showed intestinal epithelial damage and goblet cell depletion in oxazolone-treated fish within 5 hours of treatment, which lasts for up to 7 days [30]. This was accompanied by eosinophil infiltration into the intestine with increased il1b, tnf, and il10 gene production compared to untreated zebrafish. The inflammatory phenotype was reduced by the administration of the gram-positive-specific antibiotic vancomycin, suggesting a role for commensal bacteria in the inflammatory process of this model. The main constraint of this model is the need to use adult, nontransparent zebrafish which severely limits imaging capability. In addition, the need to rectally administer each individual zebrafish with oxazolone represents a technical challenge.

The hapten 2,4,6-trinitrobenzene sulfonic acid (TNBS) is another popular mammalian model of colitis characterized by a Th1-driven immune response [48] that is used for the study of both acute and chronic intestinal inflammation. In the acute model, TNBS is dissolved in ethanol and rectally administered in a single dose, while the chronic model can have multiple administrations weekly over a period of months [50]. The ethanol is responsible for disrupting the intestinal barrier while the TNBS serves to activate the immune system [48]. Two research groups have developed a larval model of colitis by exposing 3 dpf zebrafish for 3–5 days to TNBS in the fish media [28, 29]. Using the MPO reporter strain (Tg(BACmpx:GFP)i114) these investigators noticed neutrophil infiltration throughout the zebrafish [13], gut-specific increases in il1b expression, altered intestinal lipid metabolism, goblet cell hypertrophy, and intestinal shortening. This larval TNBS model presents several advantages, including a dose-dependent phenotype, a sensitivity to antibiotics treatment, and a response to anti-inflammatory agents (5-ASA and prednisolone) widely used to treat human IBD [29]. However, the intestinal histopathologies are poorly characterized in this model. In addition, the inflammatory component appears mostly to be nonintestine specific. Furthermore, no intestinal epithelial cell damage is observed histologically [29], suggesting a lack of significant intestinal injury. Finally, it is unclear how a hapten could cause an inflammatory reaction in larvae missing a functional adaptive immune system.

We have recently established a model of epithelial injury in zebrafish using the NSAID glafenine [51]. Administration of glafenine to 5 dpf zebrafish for 12 hours results in a dramatic increase in intestinal epithelial cell apoptosis. This intestinal-specific apoptotic response is mediated by induction of endoplasmic reticulum (ER) stress and is accompanied by attenuation of the unfolded protein response (UPR) coupled to an improper activation of downstream UPR mediators such atf6 and xbp1. Interestingly, loss of XBP1 in IECs results in the development of spontaneous enterocolitis in mice [52] and polymorphisms of this gene are associated with human IBD [53].

Chronic liver disease is responsible for over 25,000 deaths annually in the United States and represents the tenth leading cause of death, with a prevalence of over 5.5 million patients in 1998 [54] and that has since been estimated to affect up to 30% of the US population [55]. In the UK, the picture is grimmer, as it is the fifth leading cause of death [56]. Nonalcoholic fatty liver disease (NAFLD) is a highly prevalent form of severe chronic liver disease, affecting 30% of all adults in America [57]. It also has significant associated mortality, with a 5-year survival of 67% and 10-year survival of 59% [57]. Additionally, up to one-third of all Americans have some level of NAFLD [56]. Outside of the USA, where obesity is less prevalent, rates are also increasing. In Italy, which is traditionally considered low risk, an incidence of NAFLD of 20–25% of the population...
has been recently been reported [56]. In China and Japan, the disease incidence has been reported at 15% and 14% of the population, respectively [56]. There is no identifiable cause of NAFLD, but the pathology is linked to obesity, diabetes, and hyperlipidemia. Treatment involves managing these complex etiologies, and pharmacological therapies specific for NAFLD are not available for these patients [57]. Consequently, significant work has been invested to develop reliable zebrafish larvae models of liver pathology that could be easily utilized for drug targeting and screening.

The most characterized zebrafish model of NAFLD involved a forward genetic screen using viral insertion and screening for hepatomegaly. In the case of the NAFLD model, a 172 bp gene trap cassette was found to be inserted in the intron between exons 11 and 12 of the recently discovered zebrafish gene foie gras (foigr) [58], which results in a frame shift mutation, and generation of a stop codon. This mutation (foigr<sup>atf6</sup>) results in the development of steatosis (fatty liver disease) resembling human NAFLD, characterized by large lipid filled hepatocytes and cellular apoptosis in larvae as young as 5 dpf [58]. However, the exact function of this gene, which is highly conserved across species including humans, is not yet determined. Further studies with the foigr mutant have shown that the apoptosis observed increases ER stress and is regulated in part through the atf6 gene, which is highly conserved across species including humans [59]. Binge drinking leads to transient fatty liver disease [54], with liver pathology including fatty droplets, hepatocyte apoptosis, and upregulation of apoptotic genes such as bad, bax, p-38a, caspase-3 and 8, and jnk-1. Overall, the availability of NAFLD zebrafish models is burgeoning, and they are poised to be screened for pharmacological compounds that could for the first time effectively treat this disease.

Alcoholic fatty liver disease (ALD) is estimated to be involved in over 50% of all deaths due to liver failure secondary to liver cirrhosis, but accurate estimates for prevalence are unavailable [54]. Binge drinking leads to transient fatty liver disease, but chronic alcohol consumption can lead to fibrosis, cirrhosis, and steatohepatitis [59]. An ALD model has also been developed in zebrafish with administration of 2% ethanol to the water of 4 dpf zebrafish for 32 days. This regimen results in hepatomegaly and steatosis, alongside upregulation of hepatic cyp2e1, sod, and bip gene expression—indicating hepatotoxic metabolism of the ethanol [33]. Importantly, ethanol-induced steatosis was prevented by morpholino blockade of the sterol regulatory binding protein (Srebps). Because Srebps activation is important in chronic alcoholic liver disease [33] in humans, the zebrafish ALD model could be utilized to identify new therapeutic mechanisms and to screen for therapeutic agents.

4. Microbe-Host Interactions

The ability to generate germ-free and gnotobiotic zebrafish [60] has led to an increasing interest in understanding cellular and molecular mechanisms of microbial-host interactions using zebrafish. Microbial-host interactions have received significant interest in recent years and have been implicated in a wide variety of human diseases, including CRC [61], IBD [42, 43], obesity and diabetes [62], intestinal healing [63], irritable bowel syndrome [64], and inflammatory nociception [65]. Zebrafish are host to gram-positive and gram-negative bacteria, mycobacteria, protozoa, and viruses [66], also allowing for their use as a model for infectious diseases. In most cases, microorganisms associated with these pathologies are not intrinsically pathogenic but are rather part of the normal, commensal biota. The organ exposed to the highest amount of microorganisms in the human body is the gastrointestinal (GI) tract, especially the distal ileum and colon. Because of the high prevalence of bacteria and bacterial products in the GI tract, investigators have studied mechanisms controlling homeostasis in the face of such highly antigenic materials. The zebrafish GI tract also harbors the highest concentration of bacteria and as such represents an interesting model to study bacterial/host interactions.

Bacteria and their associated products such as DNA, RNA, and membrane structures (peptidoglycans, LPS—lipopolysaccharide) are typically detected by a series of innate sensors that are evolutionary conserved. A survey of the zebrafish genome has identified numerous innate sensors from the TLR and the Nod-like receptor family as well as their associated signaling pathways [67]. An almost complete set of TLRs (TLR1-5, 7-9, 21 [68]) exist in zebrafish, as do the intracellular sensors Nod1 and Nod2 [69]. Associated innate signaling proteins have also been identified in zebrafish such as myeloid differentiation primary response gene (MyD88) [68, 70], TIR domain-containing adaptor inducing IFN-β (TRIF) [71], and NF-κB [72, 73].

The most studied bacterial product is arguably the gram-negative-derived LPS. In mammals, LPS is detected by TLR4, in association with the coreceptors MD2 and CD-14 [74]. Once engaged, TLR4 recruits the adapter protein Myd88 or TRIF to propagate the signal to NF-κB [75] and the interferon responsive factor (IRF) 3, respectively [76]. This signaling cascade plays a critical role in host response to microbes and microbial products. As in humans, high doses of LPS are toxic to zebrafish [77]. Studies using commensal and germ-free zebrafish established that intestinal alkaline phosphatase (iap), known to detoxify the endotoxic lipid moiety A of LPS in mammals, is induced following microbiobial colonization [77]. Knockout of iap resulted in excessive intestinal neutrophil infiltration, a process involving functional myd88 and tnfr as determined by morpholino blockade [77].

Another important TLR system is TLR5, which detects flagellin and activates various signaling pathways including NF-κB [74]. Similarly to mammals, zebrafish detect flagellin (Salmonella derived in these experiments) to induce multiple matrix metalloproteinase genes as well as the inflammatory markers il1b, il8, ifn, and cxcl-C1c [78]. Morpholino blockade...
showed that flagellin-induced IIIb and metalloproteinase 9 (mmp9) were myd88 dependent whereas sfi and il1b were activated through another signaling system [78].

The functions of some of these TLR downstream signaling adaptors have also been studied in zebrafish. Both red and green transgenic Myd88 reporter strains such as Tg(myd88:EGFP)z163 and Tg(myd88-DsRED2)z164 have been developed [70]. These strains have been crossed to both macrophage (Tg(lyz:DsRED2)z50 [79]) and neutrophil (Tg(BACmpx:GFP)p114 [13]) reporter strains to monitor myd88 signaling. Studies with these dual-reporter fish demonstrated that MyD88 colocalized with both macrophages and neutrophils and that these myd88-expressing leukocytes migrated to wound sites and were involved in bacterial phagocytosis [70].

Another downstream mediator of LPS signaling, acting independently of MyD88, is TRIF. While zebrafish TRIF shares only 32% homology with the human protein, like in mammalian systems it can induce IFN and NF-κB luciferase-reporter responses as seen using an overexpression system in human HEK293T cells [71]. Interestingly TRIF-dependent gene induction through the TLR4 pathway was found to be nonfunctional in experiments with adult zebrafish gene induction through the TLR4 pathway was found to be nonfunctional in experiments with adult zebrafish [71]; however, this is to be expected given that zebrafish TLR4 paralogues are analogous to the human infectious agent M. marinum with 5 hpf embryos results in the formation of macrophage-driven granulomas within 5 dpf [90], a hallmark of TB pathology. In vivo imaging and macrophage depletion showed that granulomas, traditionally thought to be a host-protective mechanism, may in fact be a source of early TB tissue dissemination by passing M. marinum into uninfected macrophages during phagocytosis [91, 92]. Consequently, this system has provided novel insights into the pathological mechanisms of TB infection and its interactions with the host immune system and is poised to be used as a drug screen to identify novel anti-TB compounds.

Other pathogens have also been studied in zebrafish. These include S. aureus, S. pyogenes, and S. typhimurium. These and other infectious models in zebrafish have recently been reviewed in detail by Meeker and T rede [9]. Although very informative, these studies were not performed using a natural route of infection but rather by using direct delivery (injection) into embryos at various developmental stages. However, oral gavage technology in larvae would likely improve the physiological relevance of infectious models performed in zebrafish. Another limitation of zebrafish infection models is the difference in host temperature; zebrafish and their natural pathogens exist at a temperature of 28°C, while many human-relevant pathogens are only infectious at 37°C [9]. Finally, there is some evolutionary divergence in TLR signaling that must be taken into account when working with this organism. Specifically, in contrast to mammals, LPS is not sensed by zebrafish TLR4 and the sensor negatively regulates MyD88 signaling [80]. Despite these limitations, zebrafish remain a powerful tool for studying microbe-host interactions [67].

5. Genetic Diseases and Drug Screens

Because of their high fecundity, transparency, and ease of imaging, zebrafish are particularly well suited for genetic screening approaches. Forward and reverse genetic approaches can be undertaken to generate new zebrafish phenotypes and identify new genes of interest with potential relevance to human disease phenotypes. Zebrafish disease models can also be screened in a cost- and time-effective manner to discover disease-suppressing compounds. Using a whole organism is a particularly appealing aspect of zebrafish-based screens, since complex cell-cell and organ-organ interactions are kept intact.

There are two main approaches to forward-genetic screens in zebrafish that have been undertaken thus far. The
first involves exposing males to the mutagen ethylnitrosourea (ENU) and then screening for a phenotype [93, 94] shared by all mutants, such as renal cysts or heart failure. ENU is an alkylating agent that typically induces A→T base transversions. Zebrafish are relatively resistant to the ENU toxic side effects, allowing for higher dosages and thus increased rates of mutation [95]. An alternative mutagenesis approach employs random retroviral insertion [96, 97]. While this retroviral method is only one-ninth as efficient at generating a mutation as the ENU approach, it circumvents the need for positional cloning by tagging each insertion event, saving significant screening effort once a phenotype is found.

Duchenne muscular dystrophy (DMD) affects 1 in 3500 males and causes progressive muscle degeneration that can lead to death. DMD is the result of mutations in the sarcomeric protein dystrophin located on the X-chromosome [98]. During an ENU screen of zebrafish [94], a mutation referred to as sapje [99] was discovered. This mutation was later found to be located in the zebrafish Duchenne muscular dystrophy (dmd) gene and causes progressive muscular degeneration in zebrafish larvae [100]. Electron microscopy showed that muscular degeneration was the result of failure to form proper muscle attachments at the myotendinous junction. This pathological mechanism was hypothesized to occur in mammals but was lacking concrete evidence, making the zebrafish a novel system for studying the pathophysiology of this devastating disease.

The most common form of heritable polycystic kidney disease (PKD) is autosomal dominant PKD. This disease affects more than 1 in 1000 live births and approximately 10% of patients with PKD will require kidney transplant due to renal failure [101]. PKD has been linked to defects in primary cilia formation and function in humans [101]. This disease has also been successfully modeled in zebrafish by using retroviral insertion mutagenesis and screening for cystic kidneys in 5 dpf larvae [102]. The screen discovered 12 genes whose loss formed cysts in the glomerular-tubular region, including two already associated with human disease (vhnf1 and pkd2), demonstrating genetic and phenotypic homology between humans and zebrafish. Three of the 10 remaining genes are homologues of Chlamydomonas (a genus of green algae) genes, which encode components of intraflagellar transport particles involved in cilia formation [102], but have yet to be analyzed in human PKD patients. The other 7 genes discovered provide completely novel targets for studying the underlying genetics and mechanisms of PKD, as many of the genes that can result in PKD remain unidentified in humans [101].

Dilated cardiomyopathy (DCM) is the cause of at least half of the 5.8 million heart failure cases in the USA, typically resulting from prior heart damage due to a myocardial infarction or an infection [103]. Worldwide, DCM is the leading indicator for heart transplant [104], with incidences to be as high as 9.6 per 100,000 person-years in Europe [104] and China [105]. DCM results in poor heart function that eventually leads to death [103]. ENU zebrafish screens discovered numerous fish with cardiac abnormalities [93, 106, 107]. Two of these mutants, silh and pickwickm171, were found to have dilated hearts with thin walls and defective contractility [108, 109]. Further analysis showed that the silh (silent heart) mutants had a mutation in cardiac troponin T gene [108], while the pickwickm171 mutants had a mutation in the titin gene [109]. Since both of these genes are known to be important in human DCM, these mutants zebrafish represent interesting and complementary models to study this pathology.

All of the aforementioned models demonstrate that forward genetic screens in zebrafish can generate specific phenotypes that are highly homologous to human diseases. Further analysis of these mutants almost invariably leads to the identification of a gene with high homology and relevance to corresponding human disease.

Targeted, reverse genetic approaches are also quite successful in generating models of disease, especially if a homolog exists between the human gene(s) driving the disease and the zebrafish. One disease that has recently been modeled in zebrafish using a reverse-genetic approach is Hutchinson-Gilford progeria syndrome (HGPS), a rare (1 : 4,000,000 people) premature senescence syndrome that generally results from sporadic mutations that disrupts conversion of prelamin A to mature lamin A [110]. HGPS patients have markedly aberrant development characterized by lipodystrophy, osteolysis that is most pronounced in the cranium, and coronary dysfunction [110]. The mean life expectancy for patients with HGPS is less than 13 years. In zebrafish, investigators were able to use a combination of directed loss of function mutations and morpholino knockdowns of prelamin and progerin to generate zebrafish that were able to live to adulthood (though with shortened livespans). These fish had both phenotypic and molecular signs of early senescence, including lipodystrophy, aberrant muscular and craniofacial skeletal structure, increased cellular apoptosis, and cell-cycle arrest [111]. These phenotypic manifestations align remarkably well with those seen in human HGPS, demonstrating a proof of concept for reverse genetic approaches to developing zebrafish models of human diseases.

Some of the mutants generated during forward-genetic screens have been subsequently subjected to drug screening methodologies. The ability to rapidly and easily image large numbers of developing zebrafish makes them highly tractable as a preclinical system for drug screens [112]. A prototypical example of drug screening in mutant zebrafish discovered with the ENU forward-genetic approaches involves the gridlock (grl) mutant. The grl mutant lacks blood flow to the posterior trunk and tail due to a localized block in caudal blood flow at the base of the dorsal aorta [113]. This model has many characteristics similar to the human congenital disease coarctation of the aorta, making it of significant clinical interest. Coarctation of the aorta involves narrowing of the aorta that affects 1 in 2,500 live births [114]. It is the fifth most common form of congenital heart defect, and without surgical treatment the mean age of survival is only 31 years of age [114]. Surgical intervention can prevent early death, but significant morbidity, usually in the form of hypertension, can persist and results in decreased life expectancy [114]. Unfortunately, no clear etiology of the
disease exists, and the grl model in zebrafish is the best animal model currently in existence for the study of this disease [114].

The grl zebrafish has a hypomorphic mutation in the hey2 gene, a basic helix-loop-helix protein involved in aortic development [115]. Gene titration studies using morpholino technology demonstrated that dose-dependent loss of this gene results in ablation of progressively larger sections of the aorta and expansion of the contiguous region of vein [116]. A pharmacological screen, assessing restoration of blood flow and survival of larvae using bright-field microscopy, found that two structurally related compounds (GS3999 and GS4012, unknown targets) promoted activation of the vascular endothelial growth factor (VEGF) pathway and were able to rescue the phenotype [117]. Another unrelated screen found that phosphoinositide-3-kinase (PI3K) inhibitors were also able to rescue the phenotype by driving VEGF activation [118]. This included the uncharacterized flavone GS4898, a structurally similar compound to the PI3K inhibitor LY294002, and the PI3K inhibitor wortmannin. This work also demonstrated for the first time that ERK signaling in embryos drives aortic differentiation, while activation of PI3K signaling produces a venous fate. These findings have since been recapitulated in a murine system and have provided new insights into arterial morphogenesis [119] as well as generated some of the first insights into the underlying mechanisms behind coarctation of the aorta in humans.

A pair of robust screens has also been performed to evaluate compounds known to induce prolonged QT intervals in humans in the hopes of developing a preclinical cardiac toxicity screen. Cardiac QT elongation, which can cause the fatal heart arrhythmia known as Torsade de Pointes, is required to be evaluated in clinical drug trials [120] and thus these screens are of great clinical importance. Two such screens to evaluate bradycardia, atrioventricular blockade and arrhythmias in zebrafish have been performed thus far [121, 122]. In the first study 100 biologically active compounds were screened and 18 of the 23 that caused arrhythmias in humans were also positive in the zebrafish, while postscreen analysis showed that poor absorption explained 4 out of 5 of the false-negatives [122]. Additionally, the combination of erythromycin and cisapride was also positive during screening while each drug alone was not, recapitulating a known drug-drug interaction that causes arrhythmias in humans. Consequently, the zebrafish hold significant promise to be used as a screen to evaluate new drugs for their potential to generate this serious and often fatal side effect.

6. Overall Limitations

The use of zebrafish as a model organism for studying human disease is a relatively new and emerging field of research. Consequently, the number of available zebrafish strains and facilities that house zebrafish is much smaller than for well-developed model higher vertebrate organisms such as the mouse. Additionally, very few validated zebrafish reagents such as antibodies and cell lines are available to the research community. This significantly curtails the in-depth investigation of molecular and cellular details implicated in a given phenotypes.

Zebrafish also have numerous duplicate genes [123], which significantly complicate generation of knockout strains using either forward or reverse-genetic approaches. Forward-genetic approaches that disrupt one gene copy likely will not disrupt the second copy, and duplicate genes also make targeted knockout strategies more difficult as both copies must be deleted. Additionally, while the zebrafish genome has been fully sequenced, the annotation is still limited and more of a work in progress. Furthermore comparative genomic analyses between zebrafish and the human or murine genome counterparts have yet to be performed.

Zebrafish also have environmental conditions that differ substantially from humans. They must be raised in water with specific ionic concentrations and temperature (28°C). Consequently, some water-insoluble small molecules cannot be administered to zebrafish because carrier solvents (such as EtOH or DMSO) would reach toxic levels before solubility is achieved. Additionally, since drugs are administered directly to the fish media, bathing the entire fish in the test compound can result in nondesired toxic side effects. An example in our hands was the attempt to use dextran sodium sulfate (DSS) to induce colitis in larval zebrafish. At concentrations as low as 0.01% (v/v) the surfactant properties of the DSS choked the gills of the zebrafish, resulting in their death before an intestine-specific effect could be observed (Goldsmith and Jobin, unpublished observation). As discussed in a previous section, the development of an oral gavage approach could circumvent this important limitation.

7. Perspectives

Animal models have been used since the inception of medical research. For most models of human disease, the preferred and most utilized animal system of human disease is overwhelmingly the murine system. Despite the tremendous gain of knowledge provided by using murine experimental systems, the long gestation time (18–20 days) and sexual maturation rate (6–8 weeks) combined with high cost of housing and breeding represent significant limitations. Furthermore, experiments with mice are labor intensive and not particularly well suited for high-throughput screening. These limitations have spurred the need to develop other model organisms that could be used to provide initial gene or drug targets information before beginning investigations in more expensive systems.

Transparent, larval zebrafish models have the potential to fill this important niche in the study of human disease, enabling rapid, physiologically relevant in vivo screening. The transparent nature of zebrafish also allows for real-time imaging of pathogenesis, which has already provided key insights into the molecular mechanisms of metastasis [124–127] and tuberculosis dissemination [90–92]. The recent advent of the Casper fish promises to extend the imaging capacity beyond larvae to adult fish, permitting studies in fish with a functional adaptive immune system. The zebrafish has already demonstrated profound bench-to-bedside power, as evidenced by the rapid translation time
(~2-years) from the initial reports of the role of H$_2$O$_2$ in neutrophil chemotaxis during wound healing in zebrafish [11] to the first utilizations of such knowledge in human patients [18].

The zebrafish remains a relatively underdeveloped model organism with large amounts of untapped potential. As more is understood about the comparative genome, anatomy, and physiology of zebrafish to that of humans, the relevance and utility of this vertebrate model will only grow and provide a powerful complement to the murine system. Two decades of research has demonstrated the power and relevancy of the zebrafish in modeling human disease. Its unique properties make it an ideal *in vivo* system for initial use in the interrogation of a given pathology before translating the observations made in the model organism to more expensive murine systems.

### References


