



RNA-seq–based mapping and candidate identification of mutations from forward genetic screens

Adam C. Miller, Nikolaus D. Obholzer, Arish N. Shah, et al.

Genome Res. 2013 23: 679-686 originally published online January 8, 2013

Access the most recent version at doi:[10.1101/gr.147322.112](https://doi.org/10.1101/gr.147322.112)

References This article cites 27 articles, 10 of which can be accessed free at:
<http://genome.cshlp.org/content/23/4/679.full.html#ref-list-1>

Creative Commons License This article is distributed exclusively by Cold Spring Harbor Laboratory Press for the first six months after the full-issue publication date (see <http://genome.cshlp.org/site/misc/terms.xhtml>). After six months, it is available under a Creative Commons License (Attribution-NonCommercial 3.0 Unported License), as described at <http://creativecommons.org/licenses/by-nc/3.0/>.

Email Alerting Service Receive free email alerts when new articles cite this article - sign up in the box at the top right corner of the article or [click here](#).



To subscribe to *Genome Research* go to:
<https://genome.cshlp.org/subscriptions>

Method

RNA-seq–based mapping and candidate identification of mutations from forward genetic screens

Adam C. Miller,^{1,3} Nikolaus D. Obholzer,² Arish N. Shah,¹ Sean G. Megason,² and Cecilia B. Moens^{1,3}

¹Division of Basic Sciences, Fred Hutchinson Cancer Research Center, Seattle, Washington 98109, USA; ²Department of Systems Biology, Harvard Medical School, Boston, Massachusetts 02115, USA

Forward genetic screens have elucidated molecular pathways required for innumerable aspects of life; however, identifying the causal mutations from such screens has long been the bottleneck in the process, particularly in vertebrates. We have developed an RNA-seq–based approach that identifies both the region of the genome linked to a mutation and candidate lesions that may be causal for the phenotype of interest. We show that our method successfully identifies zebrafish mutations that cause nonsense or missense changes to codons, alter transcript splicing, or alter gene expression levels. Furthermore, we develop an easily accessible bioinformatics pipeline allowing for implementation of all steps of the method. Overall, we show that RNA-seq is a fast, reliable, and cost-effective method to map and identify mutations that will greatly facilitate the power of forward genetics in vertebrate models.

[Supplemental material is available for this article.]

Forward genetic screens have illuminated how genes encode the information necessary for life (Crick et al. 1961; Brenner 1974; Nüsslein-Volhard and Wieschaus 1980; Meyerowitz and Pruitt 1985; Haffter et al. 1996; Nolan et al. 2000). However, the subsequent identification of the causal mutation has been the bottleneck in the forward genetics process. Indeed, only one-third of the mutants identified in the first large-scale forward screens undertaken in a vertebrate model (Haffter et al. 1996) have been cloned. This problem has been solved in invertebrate systems through the use of whole-genome sequencing (WGS) of mutant animals to identify candidate genes; the main advantage of both *Caenorhabditis elegans* and *Drosophila* is that the genomes are small, animals are isogenic, and chemically induced mutations are rare enough that the changes can be identified and quickly confirmed to be causative (Sarin et al. 2008; Blumenstiel et al. 2009). In the zebrafish, the genome is large (greater than 10 times larger than worm or fly), making it relatively expensive to use WGS. Additionally, the zebrafish genome is highly polymorphic, with each strain, and even each individual, carrying numerous polymorphisms. Thus sequencing a single animal is not sufficient to distinguish potential causative mutations from other polymorphisms. Here we describe an RNA-seq–based bulk segregant analysis (BSA) approach that allows for the inexpensive mapping and identification of candidate mutations from forward genetic screens. While we have used zebrafish as a model, this methodology is applicable to any model system with a sequenced genome.

BSA identifies regions of the genome that are linked to a causative mutation in a group of phenotypically mutant animals. This is accomplished by identifying regions of homozygosity within mutants at genetic markers found throughout the genome (Supplemental Fig. 1). BSA using PCR-based testing of infrequent microsatellite markers has been the standard for the initial map-

ping of zebrafish mutations (Geisler et al. 2007; Zhou and Zon 2011). The approach is both laborious and low-resolution and requires subsequent fine mapping using several hundreds to thousands of individual animals, each requiring multiple rounds of testing (Zhou and Zon 2011). This process is costly in terms of reagents and time. Next-generation sequencing (NGS) provides a means to identify the most abundant class of marker in the genome, single nucleotide polymorphisms (SNPs), in order to map mutations; additionally, in the same experiment, the data identify candidate mutations within the region of linkage that may be causal for the phenotype. Several WGS approaches have been successfully applied to the mapping and identification of candidate zebrafish mutations (Bowen et al. 2012; Leshchiner et al. 2012; Obholzer et al. 2012; Voz et al. 2012); however, because of the size of the zebrafish genome and the cost of WGS, these approaches have relied on relatively low coverage sequencing (two- to sevenfold). RNA-seq offers an alternative method to perform NGS mapping and presents several advantages over WGS: (1) It effectively reduces the representation of the genome, thereby decreasing the amount of sequencing needed to obtain high coverage and thus high-quality information; (2) the effect of candidate mutations on transcript splicing can be directly assessed in mutants; and (3) the effect of mutations on the expression levels of genes can be directly identified. Overall, the RNA-seq approach offers a number of advantages at reduced cost.

Here we have developed and validated *in vivo* methodology and an *in silico* bioinformatics pipeline using RNA-seq to map and identify mutations. The *in vivo* preparation is simple and straightforward, and the bioinformatics pipeline is constructed from existing open-source programs and custom scripts. We have validated our approach on several independent mutations, demonstrating the ability to map and identify mutations that are deleterious due to amino acid changes, altered splicing, or altered expression levels. Importantly, we developed a simple computational platform that allows data processing within one integrated application. Our methodology greatly increases the power of forward genetics approaches in model systems with large, polymorphic genomes.

³Corresponding authors
E-mail amiller@fhcrc.org
E-mail cmoens@fhcrc.org

Article published online before print. Article, supplemental material, and publication date are at <http://www.genome.org/cgi/doi/10.1101/gr.147322.112>.

Results

RNA-seq sample preparation and sequencing for BSA

RNA-seq-based and WGS-based BSA mapping and candidate identification are similar except that with RNA-seq the sample being sequenced is limited to the genes expressed at the time of RNA extraction. To test the applicability of RNA-seq to BSA mapping, we performed analysis on a number of known, independent, N-ethyl-N-nitrosourea (ENU)-induced mutations in the zebrafish: two nonsense mutations, *hoxb1b*^{b1219} (data not shown) and *nhs1b*^{f^{h131}} (Walsh et al. 2011); a mutation that causes splicing defects, *vangl2*^{m209} (Jessen et al. 2002); and a nonsense mutation causing nonsense-mediated decay, *egr2b*^{f^{h227}} (Monk et al. 2009). For each mutation, individual pairs of heterozygous mutants were crossed, and mutant progeny were selected based on their known phenotypes, just as would be done in the case of an unknown mutant. Separate pools with equal numbers (from eight to 80) (Table 1) of mutant (−/−) and wild-type (+/− or +/+) siblings were made (Supplemental Fig. 1). We reasoned that RNA extracted from embryos soon after the first appearance of the mutant phenotype would have the best chance of including transcripts carrying the causal mutation. We therefore performed RNA extractions on

mutant and sibling pools directly after scoring for each phenotype of interest (stages ranging from 2 to 5 days post-fertilization, depending on the phenotype; see Methods). Sequencing libraries were prepared using standard procedures. Briefly, total RNA was extracted, mRNAs were polyA selected and chemically fragmented, cDNA was prepared, and sequencing libraries were created for mutant and wild-type pools. Each pool was uniquely barcoded, allowing for multiplexing samples from several different mutants during sequencing. Sequencing was performed on an Illumina HiSeq 2000 machine with six libraries (three sibling/mutant pairs) per lane, resulting in an average of 43 million 50-bp paired-end reads per sample (Table1).

RNA-seq data processing and linkage mapping

Reads from each sibling/mutant pair were independently aligned to the zebrafish genome (Zv9.63) using TopHat/Bowtie, an intron and splice aware aligner (Supplemental Fig. 2; Trapnell et al. 2009). From the aligned data sets, we first identified SNPs within the wild-type sequence (using SAMtools mpileup and bcftools) (Li et al. 2009) that would serve as useful markers to test linkage within the mutant data (Supplemental Figs. 1C, 2). Within any genome, SNPs

Table 1. Characteristics of RNA-seq-based mapping experiments

Mutation type/effect	<i>hoxb1b</i>			<i>nhs1b</i>	<i>vangl2</i>	<i>egr2b</i>
	Nonsense	Nonsense	Nonsense	Nonsense	Splicing	Nonsense/NMD
No. of embryos in each pool	20	40	80	8	37	30
No. of 50-bp wild-type reads greater than Q30 (million)	48.5	41.5	35.4	47.9	56.9	38.6
No. of wild-type bases (Gb)	2.4	2.1	1.8	2.4	2.9	1.9
No. of 50-bp mutant reads greater than Q30 (million)	46.7	36.0	38.1	24.1	65.3	40.1
No. of mutant bases (Gb)	2.3	1.8	1.9	1.2	3.3	2.0
No. of SNPs wild-type greater than two reads ^a	630,994	564,915	506,331	508,872	483,282	374,887
No. of SNPs mutant greater than two reads ^a	636,480	523,281	545,295	333,315	553,698	398,334
Mapping						
No. of markers (greater than 25-fold/25%) ^b	40,203	33,987	27,945	53,993	50,301	24,852
Linked region (Mb) ^c	11.6	7.9	6.5	40.8	2.3	6.6
Candidate identification						
No. of 50-bp mutant reads greater than Q30 (thousand) ^d	424	245	205	4,989	120	106
Average coverage of genes ^{d,e}	32	33	40	29	12	26
Mode coverage of genes ^{d,e}	2	3	4	1	2	1
% Genes covered at greater than fourfold ^d	71	79	84	52	41	44
% Genes covered two- to fourfold ^d	26	19	14	34	58	27
% Genes covered less than twofold ^d	3	2	2	14	1	29
No. of homozygous SNPs greater than two reads ^{a,d}	3,851	2,175	1,922	6,167	839	1,637
No. of SNPs left after filtering ^{d,f}	662	317	283	719	126	232
No. of SNPs affecting coding ^{d,g}	46	18	19	129	29	58
No. of Nonsense ^{a,d,g}	1 ⁱ	1 ⁱ	1 ⁱ	2 ⁱ	0	0
No. of Missense ^{a,d,g}	6	1	1	20	2	4
No. of Isoforms altered ^{d,h}	1	0	0	nd	1 ⁱ	0
No. of expression levels altered ^{d,i}	0	0	0	3	0	1 ⁱ

(NMD) Nonsense mediated decay; (SNP) single nucleotide polymorphisms; (nd) not determined; (Q30) quality score with an accuracy of 99.9%.

^aAt least one forward alternative and one reverse alternative read.

^bGreater than 25-fold coverage at SNP with >25% heterozygosity.

^cRegion defined as having an average marker frequency within 1% of peak marker frequency.

^dValues from mutant RNA-seq data within the linked region.

^eEach gene's coverage is the average depth of reads found across all exons.

^fKnown wild-type SNPs removed from further consideration.

^gPredicted by Variant Effect Predictor from SNPs.

^hAssessed manually using Integrative Genomics Viewer.

ⁱKnown lesion included.

^jPredicted by Cuffdiff from aligned reads, greater than twofold change.

are most likely to be present in intergenic/intronic regions that are not represented in RNA-seq data. However, we find an average of ~500,000 SNPs within each of our transcriptome pools (Table 1), most of these residing within UTRs. In contrast to the relatively low coverage of WGS, the highly expressed genes within the transcriptome allow for the identification of high-quality SNP markers directly in the parental background under investigation; these markers provide highly reliable mapping information. We therefore identified SNPs within each wild-type sibling pool covered by at least 25 reads, of which at least 25% of the calls represented an alternative allele (using the custom R script RNAmappe.R) (Supplemental Fig. 2). This resulted in an average of 40,000 high-confidence markers per experiment (Table 1) that were then used to interrogate the mutant RNA-seq data for regions of the genome linked to the mutation of interest (Supplemental Fig. 1).

In bulk RNA extracted from a pool of many animals with the same mutant phenotype, the mutation underlying the phenotype of interest, as well as linked regions of the genome, will be homozygous. In contrast, due to recombination during meiosis and independent chromosome assortment, regions unlinked to the mutation (both on the same and independent chromosomes) will be heterozygous (light and dark gray bars in Supplemental Fig. 1B). Thus the SNP marker frequency at and near the mutation will be 1 (all alleles are the same), and this frequency will gradually decline with increasing genetic distance from the mutation (Supplemental Fig. 1B–D). We therefore calculated the allele frequency within the mutant RNA-seq data at the positions identified as high-quality markers and then used a sliding window of 50 neighboring SNPs (average window size of 1.9 Mb, average step size of 37.5 kb; see Methods) to average this frequency and plot it against chromosome position. This allowed for the identification of regions of linkage (Supplemental Fig. 1D; Fig. 1).

Validation of RNA-seq-based mutation mapping

We tested our RNA-seq-based mapping strategy on four known, independent mutations (*hoxb1b*^{b1219}, *nhs11b*^{fh131}, *vangl2*^{m209}, *egr2b*^{fh227}). We found that in each case the peak of highest allele frequency was centered on the known mutation (Fig. 1). For each experiment, the average allele frequency on the linked chromosome steadily rose until reaching its highest frequency surrounding the known locus (Fig. 1B). In most experiments, the average allele frequency reached greater than 0.98 (homozygous = 1) surrounding the known locus (Fig. 1). In contrast, the highest frequency on unlinked chromosomes never exceeded 0.89, and the average was ~0.65 (the average frequency of unlinked SNPs is higher than 0.5 because some SNPs are heterozygous in both parents, giving an allele frequency of 0.5, while others are homozygous in one of the two parents, giving an allele frequency of 0.75) (Fig. 1A). In one case, *egr2b*^{fh227}, the highest peak frequency in the genome was 0.93, yet this peak surrounded the *egr2b* locus (Fig. 1B). This lower peak allele frequency is likely due to the missorting of wild-type animals into the mutant pool. We examined this missorting idea directly by computationally adding reads from the wild-type pool into the mutant data and then performing the mapping experiment. We used the *hoxb1b*^{b1219} experiment because it contained a clear region of homozygosity. The method still provided a single mapping peak even after being “contaminated” with up to 30% of wild-type reads within the mutant pool (Supplemental Fig. 3). Thus even some limited missorting of wild-type individuals into the mutant pool can be tolerated.

The size of the linked region from a mapping experiment is expected to decrease with an increase in the number of mutant animals pooled due to an increased likelihood of recombination between the causative mutation and nearby SNP markers. We tested this prediction by sequencing pools of 20, 40, and 80

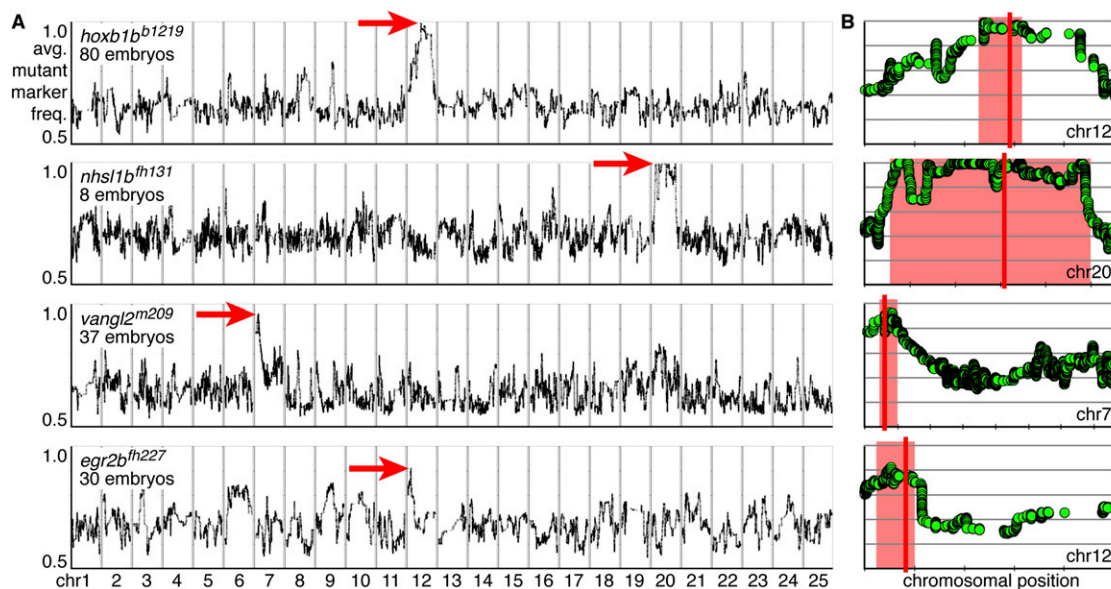


Figure 1. RNA-seq-based mapping identifies single peaks of linkage centered on the known mutations in all experiments. (A,B) Rows represent individual experiments and are labeled by genotype and the number of mutant embryos used for mapping. (A) Genome-wide mapping data. The average frequency of mutant markers (black marks) is plotted against genomic position. In each case, a single region emerges with an allele frequency near one (red arrow). Each chromosome is separated by vertical lines and labeled at the bottom. (B) Detail of the chromosome containing the linked interval for a given experiment (row). The average frequency of mutant markers (green discs) is plotted against chromosomal position. A red box marks each region of linkage, and a red line marks the position of the known mutation. Each tick mark on the x-axis represents 10 Mb. Each y-axis is the same as in A, first row.

hoxb1b^{b1219} mutant embryos and found that, as expected, increasing the number of embryos decreased the size of linkage: 11.6, 7.9, and 6.5 Mb, respectively (linkage was defined by the “leftmost” and “rightmost” position with an average mutant allele frequency within 1% of the peak frequency). For *hoxb1b^{b1219}*, increasing from 20 to 40 embryos decreased the linked region by 32%, while from 40 to 80, there was an 18% decrease (Fig. 2; Table1). At the *vangl2^{m209}* and *egr2b^{fh227}* loci, given the number of input embryos, we observed smaller homozygous intervals than would be predicted by the *hoxb1b^{b1219}* experiments (37 and 30 embryos, 2.3 and 6.6 Mb, respectively) (Fig. 2); this likely reflects the nonhomogeneous rate of recombination across the zebrafish genome (Bowen et al. 2012). Since in some cases it is difficult to acquire high numbers of mutant embryos, we also tested whether mutations could be mapped using a small number of mutant embryos. For *nhs1b^{fh131}*, we used pools of eight mutant and eight wild-type embryos with the mapping experiment resulting in a single, large region of linkage (~40 Mb) surrounding the known locus (Fig. 1). So while very few embryos can be used to accurately map mutations, pooling more mutants is advisable to minimize

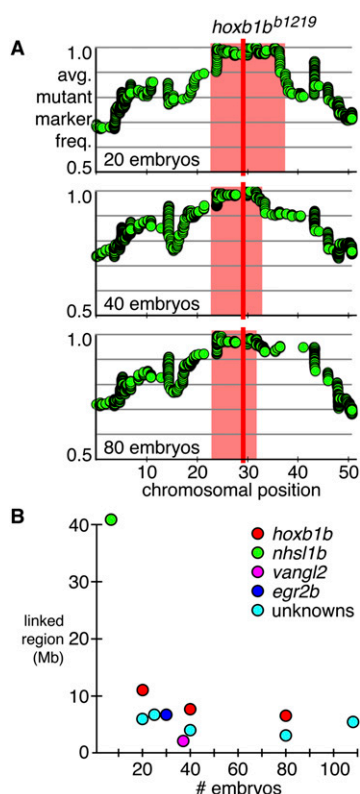


Figure 2. Increasing the number of embryos in an RNA-seq-based mapping experiment decreases the linkage size of the mapped region. (A) Detail of chromosome 12 containing the linked interval for each *hoxb1b^{b1219}* mapping experiment. Each row is labeled with the number of embryos used in the experiment. The average frequency of mutant markers (green discs) is plotted against chromosomal position. A red box marks each region of linkage, and a red line marks the position of the *hoxb1b* gene; linkage was defined as the region between the “leftmost” and “rightmost” positions within 1% of homozygosity. Each y-axis is the same as in the first row. (B) Comparison of linked regions to the number of embryos used in each RNA-seq-based mapping experiment. The *hoxb1b^{b1219}* experiments are labeled in red; *nhs1b^{fh131}*, in green; *vangl2^{m209}*, in magenta; and *egr2b^{fh227}*, in blue; and unknown mutations mapped using this method, in cyan. Increasing the number of embryos decreases the linked region with diminishing returns.

the size of the linked region. However, we found that increasing the number of embryos beyond 40 led to diminishing returns in reducing the mapped region size (Fig. 2B).

While each mapping experiment identified a single region of linkage that was centered on the known mutation, as previously described (Leshchiner et al. 2012), we found unexpected deflections from homozygosity within several linked regions; these are likely due to misplaced contigs in the current assembly of the zebrafish genome, which place unlinked SNPs that have an allele frequency less than 1.0 into the region of homozygosity. Additionally, while our mapping produced single mapping peaks, it is possible to identify regions of homozygosity due to shared lineage instead of due to linkage to the mutation, particularly in inbred backgrounds (Bowen et al. 2012); thus mutations are often outcrossed to mapping strains. Our mutants were maintained in a variety of backgrounds (see Methods), but we note that the *hoxb1b^{b1219}* and *egr2b^{fh227}* alleles were generated in a *AB background, were maintained through outcrosses to the *AB background, and were in the F3 generation post-mutagenesis; in species with high intrastrain polymorphism like the zebrafish, it is therefore possible to use RNA-seq to map mutations from forward screens without outcrossing to mapping strains, although outcrossing does provide a higher frequency of high-quality markers (Supplemental Fig. 4). Overall, the RNA-seq mapping strategy provided robust mapping of mutations to correct regions of the genome.

Identification of candidate deleterious SNP mutations

The most powerful aspect of WGS-based mapping is that it has the potential to directly identify causal mutations within the homozygous interval. After identifying a region of linkage, we revisited the mutant RNA-seq data and extracted all SNPs within the region (using the custom R script RNAidentify.R) (Supplemental Fig. 2). A concern in using RNA-seq data is that it may not sequence the mutant transcript of interest given that only genes expressed at the time of RNA extraction are captured. We found that, on average, 62% of genes within our homozygous intervals are covered by greater than four sequencing reads, 32% are covered by two to four reads at each nucleotide (average mode = 2.17), while the remaining 8% of genes are covered at levels below twofold (Table1). Thus most genes are sequenced at levels that allow for the identification of candidate mutations. Furthermore, since we isolate RNA at the time the mutant phenotype first emerges, it is likely that the transcript carrying the lesion of interest will be detected. In support of this idea, we found either the altered transcript or the effect of the mutation on transcript levels directly in the RNA-seq data for each of the known mutations (Fig. 3, see below).

Within the linked region of each mapping experiment, we used the RNA-seq data to analyze the number of SNP changes that could represent mutations of interest if mapping unknown mutants. We found that, on average, 275 alternative SNPs became homozygous per Mb of linkage (Table1). We first removed from further analysis SNPs that are known to exist in wild-type zebrafish strains. These wild-type SNPs were compiled (using the custom R script VCFmerge.R) from independent RNA-seq analysis of wild-type strains from our own facility, WGS projects (Bowen et al. 2012; Obholzer et al. 2012), and from the standard public databases (dbSNP, Ensembl). This filtering removed 86% of SNPs, leaving, on average, 40 SNPs per Mb of linkage (Table1). We then assessed whether these remaining SNPs caused nonsynonymous

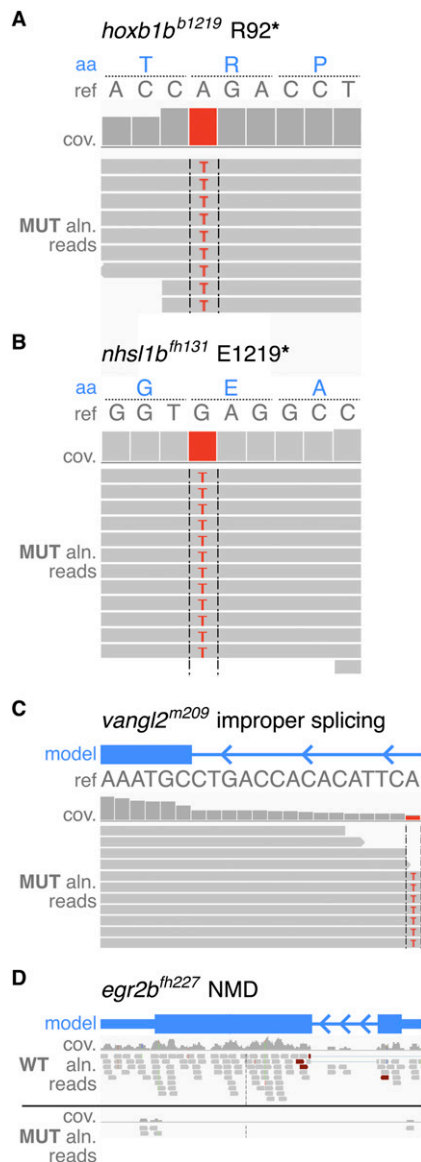


Figure 3. RNA-seq-based mapping identifies candidate mutations creating nonsense and missense changes, affecting splicing, and affecting gene expression. Reads are shown aligned to each known lesion site. Aligned reads are shown as gray boxes; differences from reference (ref) are highlighted by colored letters. (aa) Amino acid; (cov) coverage; (aln) aligned. (A–C) RNA-seq data from the mutant pool identified the known A-to-T transversion in *hoXB1b*^{b1219} (A), the G-to-T transversion in *nhsl1b*^{fh131} (B), both creating stop codons, and the creation of a splice acceptor introducing 15 bp of intronic sequence in the *vangl2*^{m209} mutation (C). (D) The down-regulation of *egr2b* (via NMD of the *egr2b*^{fh227} nonsense mutation) is evident in a comparison of the wild-type and mutant aligned reads (identified as significantly down-regulated by 25-fold via Cufflinks, $q = 0.00011423$).

changes using the Ensembl tool Variant Effect Predictor (McLaren et al. 2010) and visually confirmed each of these using the Integrative Genomics Viewer (IGV) (Thorvaldsdóttir et al. 2012). Each *hoXB1b*^{b1219} mapping experiment (80, 40, and 20 embryos) contained exactly one nonsense change in the linked interval, which was the known lesion (Fig. 3A; Table1). There was one missense change in each 80- and 40-embryo *hoXB1b*^{b1123} linked region. The

larger linkage region in the *hoXB1b*^{b1219} 20-embryo pool included five additional missense mutations (Table1). In the case of *nhsl1b*^{fh131}, which mapped to an interval of ~40 Mb due to the small mutant pool size of eight mutant embryos, there were only two nonsense mutations detected, one of which was the known lesion, and 20 missense mutations (Fig. 3B, Table1); thus even when using a limited number of embryos, the RNA-seq approach provides a manageable number of SNP candidates that might be causative. In the *vangl2*^{m209} and *egr2b*^{fh227} intervals, there were zero nonsense mutations and two and four missense mutations detected, respectively. This methodology allows for the identification of a very small number of high priority nonsense and missense candidates underlying a phenotype of interest (Table1).

Identification of candidate mutations that affect splicing

Many mutations from forward genetic screens alter the splicing of transcripts either by abolishing endogenous splice donor or acceptor sites or by creating new ones, as is the case of *vangl2*^{m209}. While WGS approaches could detect such mutations as homozygous SNPs within the linked interval, the effect of such changes can be difficult to predict, particularly in the case of the creation of a new splice acceptor or donor or in the case of unannotated exons. We analyzed how many splicing variants were identified within the mapped regions by using IGV to visually assess the transcripts with defects in splicing. Within the linked intervals, we identified very few alterations to splicing patterns: In the 80- and 40-embryo *hoXB1b*^{b1123} and *egr2b*^{fh227} experiments, there were zero splicing alterations. In the 20-embryo *hoXB1b*^{b1123} and *vangl2*^{m209} pools, there was one alteration to the splicing of a transcript, with the *vangl2*^{m209} change being the known lesion (the splice alteration identified in the *hoXB1b*^{b1123} 20-embryo pool is outside the region of linkage obtained from the 80- and 40-embryo experiments) (Fig. 3C; Table1). The ability to directly identify and assess the consequences of splice-altering mutations is thus a benefit of the RNA-seq-based approach.

Identification of candidate mutations affecting gene expression levels

A mutation can alter the level of expression of genes by (1) creating a nonsense change that results in the elimination of mutant transcripts by nonsense-mediated decay (NMD), as is the case of *egr2b*^{fh227}; (2) affecting transcription by disrupting regulatory elements; or (3) secondarily altering the expression of downstream target genes that could be in the homozygous interval. Although WGS data can detect mutations that disrupt gene expression, it does not include information about the effects of these mutations, making them hard to recognize as causal. We used Cufflinks (Trapnell et al. 2012) and the custom R script RNAeffecto.R (Supplemental Fig. 1) to identify the genes within linked regions whose expression levels are different in the wild-type and mutant pools. We found that only two of the linked regions contained significant expression level changes of greater than twofold: In the large, ~40 Mb, *nhsl1b*^{fh131} region of linkage, there were three genes affected, and in the *egr2b*^{fh227} mutant pool, there was one gene affected, *egr2b* itself (Fig. 3D; Table1). Differentiating between the different possible causes of down-regulation is challenging; however, in the case of NMD the nonsense transcript might be captured and sequenced at low frequency. Indeed, in the *egr2b*^{fh227} case, there was one read in the wild-type pool carrying the nonsense change (data not shown). In the case of regulatory mutations, sequencing of

genomic DNA surrounding the candidate gene would allow for the identification of mutations in conserved enhancer regions that could be responsible for down-regulation. The direct identification of genes with altered expression levels is another powerful benefit of the RNA-seq-based approach.

Overall our bioinformatics pipeline identified a limited number of high priority candidate mutations within the linked region of each experiment (Table 1) and, in each case, identified the known lesion. To facilitate the usefulness of the technique, we have developed an integrated, bioinformatics pipeline called RNAmapper running on the Galaxy platform (Giardine et al. 2005; Blankenberg et al. 2010; Goecks et al. 2010). This package integrates all of the tools used in the pipeline and can be downloaded and used locally or run in the cloud by creating an Amazon Machine Image. RNAmapper and its documentation can be found at www.RNAmapper.org and make the RNA-seq-based mapping approach accessible.

Discussion

Together our results show that RNA-seq-based mapping and candidate gene identification is a powerful approach allowing for the rapid and inexpensive identification of mutations from forward genetic screens. Within this article, for validation purposes, we applied the technology to only known mutations. We have also used this approach to map several unknown mutants to unique regions of the genome (Supplemental Fig. 4) and to identify a limited number of candidate lesions for each (including nonsense, missense, and splice altering lesions) (data not shown). After mapping of unknown mutations and candidate identification, regardless of linkage size and candidate number, further experiments are necessary to link a potential lesion to the phenotype definitively (e.g., demonstrating linkage of the candidate lesion to the phenotype in a large number of individuals, phenocopy via candidate knockdown/removal, and/or rescue via exogenous candidate expression). However, the short list of candidates generated using RNA-seq is likely to yield results quickly. While we have focused on zebrafish, the use of RNA-seq-based mapping is applicable to other systems. A similar approach was recently used to map a mutation in maize (Liu et al. 2012); thus the technique is applicable broadly to many genetic systems with a sequenced genome. There are several other vertebrate model systems that are commonly used to perform forward genetic screens, and their genomes are of a similar size to that of the zebrafish (zebrafish, ~1.5 Gb; *Xenopus tropicalis*, ~1.4 Gb; *Mus musculus* and *Rattus norvegicus*, ~2.7 Gb), suggesting the RNA-seq-based mapping described here can be used to identify candidate mutations in these organisms. For the larger, less polymorphic genome of the mouse, a map cross will be essential to ensure sufficient markers are identified for mapping the mutation. Additionally, because both mouse and rat have larger genomes, but have ~3000 fewer genes than zebrafish, the mapping resolution will be reduced; however, such a reduction would still provide a relatively small linked region of mapping and a small number of candidate mutations. Our RNA-seq-based approach is simple, using common laboratory procedures and free bioinformatics programs packaged into the RNAmapper program (<http://www.RNAmapper.org>).

While it is encouraging that we found the known lesion in each of our test cases, a consideration when using RNA-seq for positional cloning is that the mutant transcript of interest could be missed due to low or zero expression. We found that within our mapped intervals, from 16% to 56% of genes had low levels of

expression (less than fourfold) that would make it difficult to identify candidate mutations (these numbers were highly variable in the different linkage regions) (Table 1). To mitigate this concern in the case of an unknown mutation, we suggest extracting RNA at a timepoint as close to the first emergence of the mutant phenotype as possible, as this increases the likelihood that the transcript carrying the causal lesion will be expressed. While such early selection of the phenotype might lead to increased misphenotyping and thereby inclusion of wild-type embryos in the mutant pool, we have found our method to be surprisingly robust against such contamination (Supplemental Fig. 3). Alternatively, RNA could be extracted from a number of different developmental stages or, in the case of mapping an adult phenotype, a number of different tissues. This will increase the breadth of transcripts captured and the likelihood of sequencing the mutant transcript itself. In the worst-case scenario—where the transcript is missed—our RNA-seq approach will still provide a mapping interval due to linked SNPs from neighboring transcripts becoming homozygous. Additionally, the sequencing of transcribed genes within the interval will allow many (44%–84% of genes have greater than fourfold coverage) (Table 1) to be ruled out as candidates, narrowing the search to a limited number of genes. While the possibility of missing the causal mutation using an RNA-seq-based approach remains, care in experimental setup is likely to make this concern minimal, and the mapping will identify a region of linkage with a small list of candidates to validate in subsequent experiments.

Recently developed WGS-based BSA approaches effectively map mutation, and identify 10-fold greater SNPs in each experiment than our RNA-seq-based approach (Bowen et al. 2012; Obholzer et al. 2012). However, we find that our RNA-seq methodology maps mutations to intervals of similar size compared with WGS methods (Supplemental Fig. 5), and both approaches are able to identify nonsense and missense mutations. The RNA-seq-based approach offers three main advantages: First, sequencing the transcriptome allows for the visualization of annotated and unannotated intron/exon boundaries, allowing for the direct identification of mutations affecting splicing. WGS approaches may identify changes that alter known splice acceptor/donors but would fail to directly detect mutations affecting nonannotated isoforms or creating new splice acceptor/donors. Second, sequencing of the mutant and wild-type transcriptomes allows for the identification of candidate genes whose expression is affected by regulatory mutations. While WGS would detect the mutations themselves, other cosegregating, noncoding polymorphisms could mask the identity of the causal lesion. By providing a direct comparison of expression levels, RNA-seq identifies the effects of such mutations, but in the case of noncoding regulatory mutations, it will not detect the mutation itself. In this case, further targeted genomic sequencing would be necessary to identify the causative mutation (Gupta et al. 2010); however, the expression data would focus the search for causal noncoding mutations to those surrounding the candidate whose expression was affected. Here we compared only a single mutant to a single wild-type transcriptome; additional biological replicates would increase the significance of any expression differences between mutant and wild-type pools. Finally, RNA-seq comes at a significantly reduced cost compared with WGS approaches. Currently, WGS approaches require one to two lanes on an Illumina HiSeq 2000 for each mutant (Bowen et al. 2012; Leshchiner et al. 2012; Voz et al. 2012). In contrast, we have multiplexed six samples (three mutant/wild-type pairs) in a single lane. The RNA-seq approach thus incurs one-third to one-sixth the expense of equivalent WGS approaches.

Additionally, we found that computationally decreasing the number of reads by half (to ~20 million 50-bp reads per sample) still allowed for mapping to a small region but came at the cost of reducing the number of reads at the lesion site (Supplemental Fig. 6; Supplemental Table 1). Thus doubling the number of samples multiplexed would further decrease the cost of RNA-seq-based mapping, but these savings would come at the expense of identifying candidate mutations. As sequencing costs fall, it will become feasible to use both WGS and RNA-seq approaches, which would confirm and complement one another powerfully. Currently, RNA-seq offers many advantages at reduced cost.

Methods

The *hoxb1b*¹²¹⁹, *nhs11b*^{fh131}, and *egr2b*^{fh227} mutations were generated in the *AB strain and maintained in either a *AB (*hoxb1b*¹²¹⁹, *egr2b*^{fh227}) or a *AB/Tu background (*nhs11b*^{fh131}). The *vangl*^{m209} was generated in the Tu strain and maintained in a *AB background. *hoxb1b*¹²¹⁹ and *egr2b*^{fh227} embryos were collected in the F3 generation, while *nhs11b*^{fh131} and *vangl*^{m209} were outcrossed for greater than five generations. A single-pair of heterozygous carriers were crossed for each mutation, and embryos were collected and sorted, based on morphological phenotypes, into mutant and wild-type pools: *hoxb1b*¹²¹⁹ mutants were identified based on reduced otic vesicle size and lack of hindbrain segmentation (data not shown), *egr2b*^{fh227} mutants based on lack of hindbrain segmentation (data not shown), *vangl*^{m209} for shortened anterior/posterior axis (Jessen et al. 2002), and *nhs11b*^{fh131} for defective motor neuron migration using the *Tg(isl1:GFP)rw0* line (Walsh et al. 2011). Some phenotypes can only be detected after fixation and subsequent processing that destroys RNA; in such cases, the portion of the animal necessary for phenotype identification can be fixed and screened, while the rest of the animal can be saved for RNA extraction (this was successful in our hands in the case of mutants 3–5 in Supplemental Fig. 4).

Total RNA was extracted from each pool separately using a standard acid guanidinium thiocyanate and phenol chloroform extraction (TRIzol, Invitrogen). RNA was tested for quality using a spectrophotometer and an Agilent 2100 Bioanalyzer; RNA was only accepted if it was uncontaminated with phenol or guanidinium thiocyanate and the RNA Integrity Number (RIN) was greater than 9.0. Approximately 1.0 µg of total RNA was then polyA selected and chemically fragmented to ~200 bp, and cDNA was created using random hexamer primers. Library preparation followed the TruSeq Illumina protocol with each individual library receiving a unique Illumina barcode, allowing for their identification after multiplexed sequencing. RNA-seq was performed on an Illumina HiSeq 2000 machine with six libraries multiplexed per lane using 50-bp paired-end reads. This resulted in an average of 250 million reads per lane, with an average of 43 million reads per sample.

Raw reads were aligned to the zebrafish genome (Zv9.63) using TopHat/Bowtie, an intron and splice aware aligner (Trapnell et al. 2009). SNPs were identified using the SAMtools mpileup and bcftools variant caller (Li et al. 2009), requiring the map and nucleotide quality to be greater than 30 (i.e., the probability of a read being mismapped is one in 1000 effectively removing any repetitive sequence) and, importantly, allowing for anomalous pairs to be mapped—these “anomalous” pairs being reads spanning large introns. For the purposes of mapping, SNPs were further filtered for quality based on expression level (at least 25-fold) and for high alternative allele frequency (at least 25%) using the custom R script RNAmapper.R. RNAmapper.R then

assessed the mutant allele frequency at the positions of the high-quality wild-type SNP markers and averaged these frequencies using a sliding window of 50 neighboring markers with a step size of one SNP. Markers are spaced, on average, every 37.5 kb, but because SNPs identified are within coding regions, the distance between markers is variable based on the locations of genes within the genome. The average allele frequency was then plotted across the genome, and linkage was identified by analyzing the genome-wide mapping data for the region of highest average frequency.

Once a region of linkage was identified, the custom R script, RNAidentifie.R, was used to extract all SNPs within this region. These include any SNPs that are detected regardless of coverage level, providing the broadest list possible of potentially causal mutations but requiring the user to determine the quality and coverage level required for further characterization. RNAidentifie.R then filtered the mutant SNPs against independently identified wild-type SNPs to remove SNPs that existed in the background before the mutagenesis. A custom R script, VCFmerge.R, was written to combine VCF formatted SNPs from multiple sources, including RNA-seq data from our in-house wild-type strains, recent WGS projects (Bowen et al. 2012; Obholzer et al. 2012), and standard community sites (dbSNP, Ensembl). Linked SNPs remaining after filtering were then assessed for consequences to proteins using Ensembl's Variant Effect Predictor (VEP) (McLaren et al. 2010) or snpEff (Cingolani et al. 2012). The custom R script VEPsorte.R was used to sort and prioritize SNP candidates from VEP. The Cufflinks package was used to assess differences in expression between the wild-type and mutant pools (Trapnell et al. 2012). The custom R script RNAeffecto.R was used to extract and identify genes with significant expression level changes within the linked region. IGV (Thorvaldsdóttir et al. 2012) was used to assess splice changes at intron/exon boundaries and also to visually assess each potential candidate mutation. All custom scripts were written in R and are available for download (<http://www.RNAmapper.org>) with an open-source BSD license.

To generate a user-friendly mapping platform, we developed RNAmapper based on Galaxy (<http://galaxy.psu.edu>) and created a downloadable package that can be run on a powerful desktop workstation. We also created an Amazon Machine Image to allow users to instantiate their own RNAmapper server on the Amazon Elastic Compute Cloud. We packaged RNAmapper and all associated required programs and reference data into a single bundle using VirtualBox (<https://www.virtualbox.org/>). Alternatively, all programs and custom scripts listed in Supplemental Figure 2 can be run from the unix/linux command line. The source code and virtual machines are free to download at www.RNAmapper.org.

Data access

All RNA-seq data have been submitted to the NCBI Sequence Read Archive (SRA) (<http://www.ncbi.nlm.nih.gov/sra>) under the BioProject accession PRJNA172016, and each sequencing file can be found under the accession numbers SRS352960, SRS352996, SRS352997, SRS352998, SRS353000, SRS353001, SRS353003, SRS353004, SRS353006, SRS353007, SRS353008, SRS353009. All scripts, source code, and programs developed here can be found at www.RNAmapper.org.

Acknowledgments

We thank John Morgan and Anja Tjaden for help in RNA extraction; the Fred Hutchinson Cancer Research Center's Genomic Resource Center—particularly Jeff Delrow, Andy Marty, and Alyssa Dawson—for help with experiment design, sequencing library

preparation, and sequencing; and Ryan Basom for help in data processing. Funding was provided by the National Institutes of Health, R01HD037909 and R01HG002995 to C.B.M., the NRSA fellowship F32NS074839 to A.C.M., and NIDCD R21DC012097 to S.G.M.

Author contributions: A.C.M. and C.B.M. performed the in vivo experiments and sample preparation. A.C.M., A.N.S., and N.D.O. wrote the custom R scripts. N.D.O. created the Galaxy implementation of the bioinformatics pipeline. A.C.M. and C.B.M. wrote the manuscript. All authors edited the manuscript.

References

- Blankenberg D, Von Kuster G, Coraor N, Ananda G, Lazarus R, Mangan M, Nekrutenko A, Taylor J. 2010. Galaxy: A web-based genome analysis tool for experimentalists. *Curr Protoc Mol Biol* **89**: 19.10.1–19.10.21.
- Blumenstiel JP, Noll AC, Griffiths JA, Perera AG, Walton KN, Gilliland WD, Hawley RS, Staehling-Hampton K. 2009. Identification of EMS-induced mutations in *Drosophila melanogaster* by whole-genome sequencing. *Genetics* **182**: 25–32.
- Bowen ME, Henke K, Siegfried KR, Warman ML, Harris MP. 2012. Efficient mapping and cloning of mutations in zebrafish by low coverage whole genome sequencing. *Genetics* **190**: 1017–1024.
- Brenner S. 1974. The genetics of *Caenorhabditis elegans*. *Genetics* **77**: 71–94.
- Cingolani P, Platts A, Wang LL, Coon M, Nguyen T, Wang L, Land SJ, Lu X, Ruden DM. 2012. A program for annotating and predicting the effects of single nucleotide polymorphisms, SnpEff: SNPs in the genome of *Drosophila melanogaster* strain *w¹¹¹⁸*; *iso-2*; *iso-3*. *Fly* **6**: 80–92.
- Crick FH, Barnett L, Brenner S, Watts-Tobin RJ. 1961. General nature of the genetic code for proteins. *Nature* **192**: 1227–1232.
- Geisler R, Rauch G, Geiger-Rudolph S, Albrecht A, van Bebber F, Berger A, Busch-Nentwich E, Dahm R, Dekens MPS, Dooley C, et al. 2007. Large-scale mapping of mutations affecting zebrafish development. *BMC Genomics* **10**: 865–867.
- Giardine B, Riemer C, Hardison RC, Burhans R, Elnitski L, Shah P, Zhang Y, Blankenberg D, Albert I, Taylor J, et al. 2005. Galaxy: A platform for interactive large-scale genome analysis. *Genome Res* **15**: 1451–1455.
- Goecks J, Nekrutenko A, Taylor J, Galaxy Team. 2010. Galaxy: A comprehensive approach for supporting accessible, reproducible, and transparent computational research in the life sciences. *Genome Biol* **11**: R86.
- Gupta T, Marlow FL, Ferriola D, Mackiewicz K, Dapprich J, Monos D, Mullins MC. 2010. Microtubule actin crosslinking factor 1 regulates the Balbiani body and animal-vegetal polarity of the zebrafish oocyte. *PLoS Genet* **6**: e1001073.
- Haffter P, Granato M, Brand M, Mullins MC, Hammerschmidt M, Kane DA, Odenthal J, van Eeden JM, Jiang Y, Heisenberg C, et al. 1996. The identification of genes with unique and essential functions in the development of the zebrafish, *Danio rerio*. *Development* **123**: 1–36.
- Jessen JR, Topczewski J, Bingham S, Sepich DS, Marlow F, Chandrasekhar A, Solnica-Krezel L. 2002. Zebrafish trilobite identifies new roles for Strabismus in gastrulation and neuronal movements. *Nat Cell Biol* **4**: 610–615.
- Leshchiner I, Alexa K, Kelsey P, Adzhubei I, Austin CA, Cooney JD, Anderson H, King MJ, Stottmann R, Ha S, et al. 2012. Mutation mapping and identification by whole genome sequencing. *Genome Res* **22**: 1541–1548.
- Li H, Handsaker B, Wysoker A, Fennell T, Ruan J, Homer N, Marth G, Abecasis G, Durbin R, 1000 Genome Project Data Processing Subgroup. 2009. The sequence alignment/map (SAM) format and SAMtools. *Bioinformatics* **25**: 2078–2079.
- Liu S, Yeh CT, Tang HM, Nettleton D, Schnable PS. 2012. Gene mapping via bulked segregant RNA-Seq (BSR-Seq). *PLoS ONE* **7**: e36406.
- McLaren W, Pritchard B, Rios D, Chen Y, Flicek P, Cunningham F. 2010. Deriving the consequences of genomic variants with the Ensembl API and SNP Effect Predictor. *BMC Bioinformatics* **26**: 2069–2070.
- Meyerowitz EM, Pruitt RE. 1985. *Arabidopsis thaliana* and plant molecular genetics. *Science* **229**: 1214–1218.
- Monk KR, Naylor SG, Glenn TD, Mercurio S, Perlin JR, Dominguez C, Moens CB, Talbot WS. 2009. A G protein-coupled receptor is essential for Schwann cells to initiate myelination. *Science* **325**: 1402–1405.
- Nolan PM, Peters J, Strivens M, Rogers D, Hagan J, Spurr N, Gray IC, Vizor L, Brooker D, Whitehill E, et al. 2000. A systematic, genome-wide, phenotype-driven mutagenesis programme for gene function studies in the mouse. *Nat Genet* **25**: 440–443.
- Nüsslein-Volhard C, Wieschaus E. 1980. Mutations affecting segment number and polarity in *Drosophila*. *Nature* **287**: 795–801.
- Obholzer N, Swinburne IA, Schwab E, Nechiporuk AV, Nicolson T, Megason SG. 2012. Rapid positional cloning of zebrafish mutations by linkage and homozygosity mapping using whole-genome sequencing. *Development* **139**: 4280–4290.
- Sarin S, Prabhu S, O'Meara MM, Pe'er I, Hobert O. 2008. *Caenorhabditis elegans* mutant allele identification by whole-genome sequencing. *Nat Methods* **5**: 865–867.
- Thorvaldsdóttir H, Robinson JT, Mesirov JP. 2012. Integrative Genomics Viewer (IGV): High-performance genomics data visualization and exploration. *Brief Bioinform* doi: 10.1093/bib/bbs017.
- Trapnell C, Pachter L, Salzberg SL. 2009. TopHat: Discovering splice junctions with RNA-Seq. *Bioinformatics* **25**: 1105–1111.
- Trapnell C, Roberts A, Goff L, Pertea G, Kim D, Kelley DR, Pimentel H, Salzberg SL, Rinn JL, Pachter L. 2012. Differential gene and transcript expression analysis of RNA-seq experiments with TopHat and cufflinks. *Nat Protoc* **7**: 562–578.
- Voz ML, Coppieters W, Manfroid I, Baudhuin A, Von Berg V, Charlier C, Meyer D, Driever W, Martial JA, Peers B. 2012. Fast homozygosity mapping and identification of a zebrafish ENU-induced mutation by whole-genome sequencing. *PLoS ONE* **7**: e34671.
- Walsh GS, Grant PK, Morgan JA, Moens CB. 2011. Planar polarity pathway and Nance-Horan syndrome-like 1b have essential cell-autonomous functions in neuronal migration. *Development* **138**: 3033–3042.
- Zhou Y, Zon LI. 2011. The zon laboratory guide to positional cloning in zebrafish. *Methods Cell Biol* **104**: 287–309.

Received August 3, 2012; accepted in revised form December 18, 2012.