

# The Syntenic Relationship of the Zebrafish and Human Genomes

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The zebrafish is an important vertebrate model for the mutational analysis of genes effecting developmental processes. Understanding the relationship between zebrafish genes and mutations with those of humans will require understanding the syntenic correspondence between the zebrafish and human genomes. High throughput gene and EST mapping projects in zebrafish are now facilitating this goal. Map positions for 523 zebrafish genes and ESTs with predicted human orthologs reveal extensive contiguous blocks of synteny between the zebrafish and human genomes. Eighty percent of genes and ESTs analyzed belong to conserved synteny groups (two or more genes linked in both zebrafish and human) and 56% of all genes analyzed fall in I18 homology segments (uninterrupted segments containing two or more contiguous genes or ESTs with conserved map order between the zebrafish and human genomes). This work now provides a syntenic relationship to the human genome for the majority of the zebrafish genome.

Zebrafish is an important model system for analysis of vertebrate development (Kimmel 1989; Driever et al. 1996) and an emerging model system for human disease (Zon 1999). Understanding the relationship between the zebrafish and human genomes will help identify roles for human genes from zebrafish mutations, and help identify zebrafish models for genes identified by human disease (Brownlie et al. 1998). Hundreds of zebrafish genes and thousands of zebrafish ESTs have been identified that provide the basis for comparing the relationship between the human and zebrafish genomes. These can be compared with human genes to identify orthologs. Subsequent mapping can be used to define the extent of conservation between zebrafish and human genomes. Earlier reports identify map locations for 124 zebrafish genes with mapped human orthologs (Postlethwait et al. 1998; Gates et al. 1999). Analysis of this mapping data revealed many instances of conserved synteny, whereby two or more genes that are found on the same chromosome in zebrafish are also found on the same chromosome in humans. In some cases, members of such syntenic groups were contiguous with one another and had conserved map order suggesting no large-scale rearrangements between zebrafish and human genomes in these regions (we call these homology segments). Nevertheless, not enough genes were analyzed to give a global picture of the extent of conserved synteny

between zebrafish and human genomes. We have increased the number of analyzed genes and ESTs to 523, allowing a more complete analysis of the syntenic relationship between human and zebrafish genomes.

## RESULTS

We used 523 mapped zebrafish genes and ESTs with mapped human orthologs to compare the syntenic relationship of the zebrafish and human genomes. These included 25 genes and 228 ESTs mapped in this study on the LN54 zebrafish radiation hybrid panel (Hukriede et al. 1999) in addition to 270 genes and ESTs with previously reported map positions (Johnson et al. 1996; Postlethwaite et al. 1998; Gates et al. 1999; Geisler et al. 1999; Hukriede et al. 1999). Related gene clusters (such as *hox* clusters, *dlx* gene pairs, the major histocompatibility complex, or hemoglobin loci) are represented as single genes in our analysis to prevent an overestimate of the extent of conserved synteny. Orthology was determined by WU-BLAST analysis (W. Gish, unpubl.; <http://BLAST.wustl.edu>), selecting for highly significant matches (maximum WU-BLASTN probability of  $e^{-20}$ , see Materials and Methods). Genes and ESTs positioned with other mapping panels were integrated onto our map with respect to markers shared between each panel (Johnson et al. 1996; Postlethwaite et al. 1998; Gates et al. 1999; Geisler et al. 1999; Hukriede et al. 1999). Approximately 400 additional mapped genes and ESTs were excluded from this analysis because they had no obvious human or mouse ortholog, or map positions of human orthologs were unknown (data not shown). A small subset of ESTs and genes had multiple possible orthologs, which pre-

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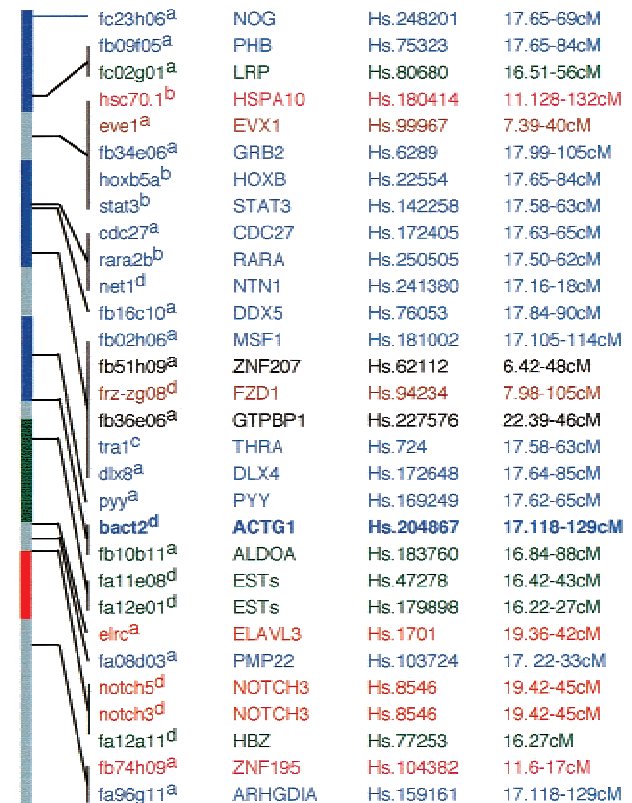
Article and publication are at [www.genome.org/cgi/doi/10.1101/gr.144700](http://www.genome.org/cgi/doi/10.1101/gr.144700).

vented unambiguous orthology assignments (see below).

An example of the extent of syntenic correspondence of zebrafish and human genomes is shown in Figure 1. Of the 29 LG3 genes and ESTs with mapped human orthologs, 27 (93%) belong to five conserved synteny groups, corresponding to human chromosomes Hsa7, Hsa11, Hsa16, Hsa17, and Hsa19. The 14 genes of the LG3-Hsa17 conserved synteny group (excluding *bact2* for this analysis; see below) are separated into four uninterrupted segments of conserved map order (fc23h06–fb09f05, fb34e06–*net1*, *rara2*–fb02h06, and *dlx8*–*pyy*) that likely represent homologous segments conserved intact, or nearly intact, between human and zebrafish. An additional two ESTs, fa08d03 and fa96g11 from the LG3–Hsa17 conserved synteny group (that BLAST analyses suggest identify zebrafish orthologs to human *PMP22* and *ARHGDI*A genes) are not contiguous with other genes from the conserved synteny group. However, their membership in the LG3–Hsa17 conserved synteny group adds support to the predicted orthology, and suggests that these ESTs may nucleate additional zebrafish–human homology segments as more genes are analyzed. By similar logic, the other four conserved synteny groups represented on LG3 may identify an additional nine multiple- or single-gene homology segments, increasing the number of homology segments on LG3 to 15. Two ESTs on LG3, fb51h09 and fb36e06, are not identified as members of defined conserved synteny groups and thus lack independent support for the existence of additional homology segments (see below for possible alternatives). We refer to this class of mapped gene as singletons.

Genome-wide, 421 of 523 mapped genes and ESTs were in 113 conserved synteny groups, averaging 4.5 groups (range 2–7) per zebrafish chromosome (Table 1). As observed above for LG3, genes and ESTs in conserved synteny groups fall into two classes: one class of uninterrupted segments of two or more genes and ESTs with conserved gene order in zebrafish and human that likely represent homology segments conserved intact, or nearly intact, between human and zebrafish; and a second class of single genes and ESTs that belong to conserved synteny groups, but are otherwise isolated from members of their conserved synteny group. Thus, we found 292 genes and ESTs (56% of total) in the first class arranged in 118 multiple-gene homology segments and a further 129 genes and ESTs in the second class separated from other members of their conserved synteny group (presumably by intrachromosomal rearrangements). The fact that this second class of genes are part of conserved synteny groups tends to support their predicted orthology, thus providing evidence for additional homology segments and therefore raising the number of likely zebrafish–human homol-

LG 3



**Figure 1** Syntenic relationship between zebrafish linkage group 3 and the human genome. Vertical staff shows map of zebrafish LG3 derived from genes and ESTs (column 1) typed on the LN54 Radiation Hybrid panel 1, or genes and ESTs typed on other panels integrated onto the LN54 map with respect to SSLP markers typed in common. Because gene and EST marker order cannot always be precisely determined when typed on different panels, we show them in high-confidence bins with respect to position of framework markers of the LN54 panel (Hukriede et al. 1999). Order within confidence bins is not established and we have inferred minimal chromosomal rearrangements for our analysis. Superscripts indicate sources of mapping data: (<sup>a</sup>) the LN54 zebrafish RH panel (Hukriede et al. 1999; this study), (<sup>b</sup>) the MOP meiotic panel (Johnson et al. 1996; Postlethwait et al. 1998), (<sup>c</sup>) the GAT meiotic panel (Gates et al. 1999), or (<sup>d</sup>) the Goodfellow zebrafish RH panel (Geisler et al. 1999). Orthologous human genes (column 2), UniGene reference sequence (<http://www.ncbi.nlm.nih.gov/UniGene>) (column 3), and Gene Map 98 (Deloukas et al. 1998) position (column 4) are shown to right. Conserved synteny groups are as shown as follows: blue, Hsa17; green, Hsa16; light red, Hsa7; dark red, Hsa19; pink, Hsa11; and singletons, black. Contiguous regions with two or more genes from the same conserved synteny group are shaded the corresponding color on the map staff (left). Bold type shows gene (*bact2*) where determination of orthology was assisted by syntenic relationships. See <http://zfish.wustl.edu>, or supplemental information at the Genome Research web site (<http://www.genome.org>) for maps showing other zebrafish-to-human or human-to-zebrafish relationships.

ogy segments to 247 (118 + 129). The remaining 102 mapped genes and ESTs (19% of total) that are not currently in conserved synteny groups (thus, singletons, see Figure 2), may reflect the existence of addi-

**Table 1.** Zebrafish–Human Conserved Synteny

Zebrafish linkage group	Human chromosome
1	1, 2, 4, 13, 14
2	1, 2, 3, 7, 8, 9, 19
3	7, 11, 16, 17, 19
4	3, 7, 11, 12
5	5, 9, 11, 14, 17, 19, X
6	2, 12, 13, 19
7	7, 11, 16, 19
8	1, 3, 4, 5, 7, 8, X
9	2, 11, 21, X
10	3, 4, 11, 21
11	1, 3, 8, 12, 17
12	2, 10, 17, 22
13	4, 6, 10, 19
14	5, 11, X
15	3, 11, 17
16	3, 6, 8, 17, 19
17	2, 4, 14, 20
18	11, 15, 19, 22
19	1, 3, 6, 7
20	2, 4, 6, 20
21	5, 6, 9, 10, 11
22	1, 2, 7, 12, 19
23	1, 3, 6, 7, 12, X
24	8, 10
25	5, 11, 15, 22

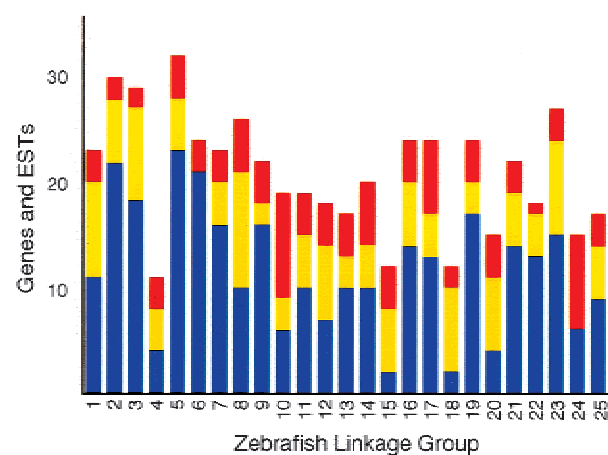
Human chromosomes (*right*) with two or more orthologous genes or ESTs mapped on corresponding zebrafish linkage groups (*left*).

tional conserved synteny groups and homology segments, or instead may reflect errors in determining orthology, errors in mapping, yet unidentified genes in the human (or mouse) data set, or instances where the corresponding orthologous gene has been lost from the human lineage. Putting these possibilities aside and assuming a Poisson distribution of genes and ESTs in synteny groups and singletons suggests the existence of a further 69 synteny groups not yet identified by mapped genes (data not shown). Therefore, the 247 homology segments supported by syntenic relationships provides a lower limit for the number of such segments but there may be upwards of 418 (247 + 102 + 69) homology segments defining the relationship between the zebrafish and human genomes (DeBry and Seldin 1996).

Previous analyses have suggested that a genome-wide duplication may have occurred in the teleost lineage since its divergence from the tetrapod lineage (Amores et al. 1998; Postlethwaite et al. 1998; Wittbrodt et al. 1998; Gates et al. 1999). Consistent with the notion of genome-wide duplication, we find 38 examples where two or more mapped, unlinked zebrafish genes share a single mammalian ortholog (Table 2). These are distributed on 20 of the 25 ze-

brafish linkage groups, and 14 of 23 human chromosomes. A further seven pairs of tightly linked zebrafish genes also share a single human ortholog, suggesting that in some cases, tandem duplications may also have played a role in generating extra zebrafish genes. However, paralogous gene pairs are not the rule for the described zebrafish genes. Analysis of ESTs from 12 ribosomal protein genes, an abundantly expressed class of genes that has been sufficiently sampled to draw inferences about gene number, revealed only two with duplicate expressed genes (S. Johnson, unpubl.), raising the possibility that if the entire genome were additionally duplicated, most of the duplicate copies have been lost or inactivated.

The described syntenic relationship between the zebrafish and human genomes can be used as a tool for predicting human orthologs for zebrafish genes and ESTs. We found 32 zebrafish genes or ESTs where multiple human homologs were suggested by WU-BLAST analysis. For 20 of these genes (61%), the syntenic relationships revealed by the foregoing analysis allowed us to predict the human orthologs (Table 3). For example, our WU-BLAST analysis failed to distinguish between human *ACTB* (on Hsa1), *ACTC* (on Hsa15), and *ACTG1* (on Hsa17) as the most likely ortholog for zebrafish *bact2* (Kelly and Reversade 1997). The map position for *bact2* on LG3 (Geisler et al. 1999) near *Pyy* (on Hsa17; Lundell et al. 1997) argues that *bact2* is the zebrafish ortholog for *ACTG1*, rather than *ACTB* or *ACTC*. Similarly, WU-BLAST analysis fails to unambigu-



**Figure 2** Distribution of genes and ESTs in synteny groups. Bars indicate the distribution of zebrafish genes and ESTs according to class of synteny relationship (Y-axis) for each linkage group (X-axis). Number of genes and ESTs from homology segments with two or more contiguous members where gene order is conserved between zebrafish and human are shown in blue. Additional genes and ESTs in conserved synteny groups but not in contiguous sets are shown in yellow. Genes and ESTs that are not part of conserved synteny groups (singletons) are depicted in red. Together these three classes account for all the mapped genes and ESTs with orthologs predicted unambiguously by WU-BLAST analysis (see Methods).

<b>Table 2. Human Genes with Two or More Zebrafish Orthologs</b>					
<b>Human gene</b>	<b>Reference (NCVI unigene)</b>	<b>Human map position</b>	<b>Zebrafish ortholog</b>	<b>Reference (NCBI gi)</b>	<b>Zebrafish map position</b>
HES5	no ref	1.49-52cM <sup>a</sup>	her2	1279391	8.472cR <sup>c</sup>
HFH2	Hs.166188	1.95-102cM	her4	1279395	23.99cR <sup>c</sup>
SOX11	Hs.32964	2.0-32cM	fk d8	2982352	8.299cR <sup>d</sup>
			fk d6	2982348	6.273cR <sup>b</sup>
RARA	Hs.173205	2.51-54cM	sox11a	NA	17.234cR <sup>d</sup>
			sox11b	NA	20.499cR <sup>d</sup>
SIX3	Hs.227277	2.73-88cM	rara2a	704369	12.125cR <sup>c</sup>
			rara2b	215025	3.161cR <sup>d</sup>
EN1	Hs.227277	2.127-134cM	six6	3047418	12.188cR <sup>f</sup>
			six3	304716	13.278cR <sup>f</sup>
DLX2	Hs.419	2.182-188cM	eng4	4322043	1.59cR <sup>d</sup>
			eng1	62515	9.9cR <sup>d</sup>
IHH	Hs.69351	2.200-215cM	dlx5	1620515	1.179cR <sup>c</sup>
			dlx2	460126	9.131cR <sup>c</sup>
FZD5	Hs.152251	2.211-218	ehh	1616584	6.115cR <sup>d</sup>
			hha	NA	9.140cR <sup>d</sup>
FZD7	Hs.173859	2.200-206cM	fz8a	4164470	24.133cR <sup>f</sup>
			frz-zg06	1245193	2.438cR <sup>f</sup>
GATA2	Hs.760	3.142-146cM	frz-zg07	1245195	9.170cR <sup>f</sup>
			<b>fb38g02</b>		<b>6.115cR<sup>b</sup></b>
ATP1B3	Hs.76941	3.157-158cM	frz-zg13	1245207	6.129cR <sup>f</sup>
			gata1	1132418	11.230cR <sup>d</sup>
EPHA5	Hs.31092	4.68-78cM	gata2	1132420	11.390cR <sup>c</sup>
			atp1b	974773	2.150cR <sup>f</sup>
NPY1R	Hs.169266	4.157-169cM	fb13c07		15.57cR <sup>a</sup>
			fb82e05		24.301cR <sup>b</sup>
EFNA5	Hs.37142	5.108-116cM	<b>rtk7</b>	<b>3005904</b>	<b>24.301cR<sup>b</sup></b>
			zya	3098345	17.79cR <sup>d</sup>
CSX	Hs.54473	5.161-163cM	zyb	2739140	8.563cR <sup>d</sup>
			zyc	3098347	10.385cR <sup>d</sup>
<b>MSX2</b>	<b>Hs.89404</b>	<b>5.185-196cM</b>	al1	1834430	8.10cR <sup>c</sup>
			ephra5	2462952	21.129cR <sup>b</sup>
ISL1	Hs.505	5.54-61cM	nkx2.7	1518150	8.505cR <sup>e</sup>
			nkx2.5	1518148	14.341cR <sup>d</sup>
AHR	Hs.170087	7.24-35cM	<b>msxe</b>	<b>1399516</b>	<b>14.27cR<sup>c</sup></b>
			<b>msxa</b>	<b>608508</b>	<b>14.464cR<sup>d</sup></b>
EVX1	Hs.99967	7.38-42cM	<b>msxd</b>	<b>62544</b>	<b>21.211cR<sup>c</sup></b>
			islet1	497897	5.143cR <sup>c</sup>
HOXA	N/A	7.39-40cM	islet2	1037165	25.406cR <sup>c</sup>
			islet3	1037167	25.406cR <sup>f</sup>
EN2	Hs.134989	7.167-175cM	ahr2	4321818	22.88cR <sup>f</sup>
			ahr	2764987	16.196cR <sup>b</sup>
SHH	Hs.121539	7.181-184cM	eve1	475049	3.113cR <sup>c</sup>
			evx1	no ref.	16.175cR <sup>d</sup>
SLUG	Hs.93005	8.57-68cM	hoxa13b	4322052	16.175cR <sup>d</sup>
			hoxa4a	4322059	19.170cR <sup>c</sup>
NOTCH1	Hs.121539	7.181-184cM	eng2	62517	7.158cR <sup>c</sup>
			eng3	62521	2.343cR <sup>c</sup>
RXRA	Hs.20084	9.143-166cM	shh	5714439	7.158cR <sup>c</sup>
			twhh	1171139	2.346cR <sup>d</sup>
FTH1	Hs.62954	11.16-23cM	sna2	841423	23.41cR <sup>d</sup>
			sna1	468620	11.284cR <sup>c</sup>
WNT11	Hs.108219	11.80-84cM	notch1b	2569967	5.267cR <sup>f</sup>
			notch1	433866	21.75cR <sup>f</sup>
HSPA10	Hs.180414	11.128-132cM	rxrg	1046288	5.222cR <sup>f</sup>
			rxra	1046294	2.309cR <sup>c</sup>
SPON1	Hs.5378	11.24-25cM	fb06g09		7.45cR <sup>b</sup>
			fb01e08		24.144cR <sup>b</sup>
HSPA10	Hs.180414	11.128-132cM	wnt11	3169686	5.125cR <sup>e</sup>
			wnt11r	NA	10.306cR <sup>d</sup>
SPON1	Hs.5378	11.24-25cM	hsc70.1	1408566	3.113cR <sup>d</sup>
			fb01g06		10.304cR <sup>b</sup>
SPON1	Hs.5378	11.24-25cM	fspdin2	2529226	25.70cR <sup>f</sup>
			mindin1	2529220	14.379cR <sup>f</sup>
			mindin2	2529222	14.341cR <sup>f</sup>

**Table 2.** (Continued)

Human gene	Reference (NCVI unigene)	Human map position	Zebrafish ortholog	Reference (NCBI gi)	Zebrafish map position
HOXC	N/A	12.70-72cM	hoxc5a	414104	23.324cR <sup>c</sup>
ASCL1	Hs.1619	12.106-113cM	hoxc13b	4322091	11.459cR <sup>d</sup>
			zasha	540237	4.149cR <sup>c</sup>
			zashb	540239	7.177cR <sup>c</sup>
OTX2	II.5015	14.0-1cM	otx2	540243	17.304cR <sup>b</sup>
			<b>otx3</b>	<b>633134</b>	<b>1.381cR<sup>c</sup></b>
RTN1	Hs.99947	14.54-58cM	deltab	2772824	5.125cR <sup>d</sup>
			dla	2809388	1.395cR <sup>d</sup>
HOXB	N/A	17.62-69cM	hoxb4a	341108	3.113cR <sup>f</sup>
			hoxb1b	1127809	12.188cR <sup>c</sup>
LHX1	Hs.157449	17.58-63cM	lim1	577524	15.189cR <sup>d</sup>
			lim6	2155288	5.171cR <sup>d</sup>
NOTCH3	Hs.8546	19.42-45cM	notch3	3153196	3.430cR <sup>f</sup>
			notch5	2569969	3.430cR <sup>f</sup>
PR65	Hs.173902	19.59-98cM	fa02h04		5.171cR <sup>f</sup>
			fb38a08		15.138cR <sup>b</sup>
CKM	Hs.118843	19.59-98cM	fa28d05		5.125cR <sup>f</sup>
			fc14g11		13.183cR <sup>b</sup>
MYRL2	Hs.9615	19.59-98cM	fa93e09		7.284cR <sup>b</sup>
			fa97a12		2.340cR <sup>b</sup>
BMP2	Hs.73853	20.18-27cM	bmp2	2804174	20.678cR <sup>c</sup>
			bmp2a	2149147	17.43cR <sup>d</sup>
SNAP25	Hs.84389	20.27-37cM	snap25a	3703097	20.459cR <sup>c</sup>
			snap25b	3703099	17.79cR <sup>c</sup>
L1CAM	Hs.1757	X.188-198cM	nadl1.1	1065713	23.22cR <sup>c</sup>
			nadl1.2	1065715	23.163cR <sup>c</sup>

Orthologs predicted with aid of syntenic correspondence (see Table 3) are shown in bold.

<sup>a</sup> Position for human gene is inferred from map position of orthologous mouse gene and the mouse-human syntenic relationship (DeBry and Seldin 1996).

<sup>b</sup> Genes and ESTs mapped in this study.

<sup>c</sup> Hukriede et al. 1999.

<sup>d</sup> Postlethwait et al. 1998, Amores et al. 1998.

<sup>e</sup> Gates et al. 1999.

<sup>f</sup> Geissler et al. 1999.

ously establish the orthologous relationship between zebrafish *msxa*, *msxb*, *msxc*, *msxd*, and *msxe* genes (Eker et al. 1997) and the human *MSX1* and *MSX2*, and mouse *Msx3* (human *MSX3* has not yet been identified) genes. Because the regions of the zebrafish linkage groups in which *msxa* (LG14), *msxd* (LG21) and *msxe* (LG14) reside are syntenic to or map near syntenic regions to the region on human chromosome 5 that contains *MSX2*, syntenic comparison suggests that the zebrafish *msxa*, *msxd*, and *msxe* genes are orthologous to human *MSX2*. Likewise, synteny analysis suggests that the zebrafish *msxb* gene (LG1) is orthologous to *MSX1* (Hsa4) and zebrafish *msxc* is orthologous to mouse *Msx3*. These and other zebrafish-human orthology relationships predicted by synteny are shown in Table 3.

## DISCUSSION

Increasing the number of mapped zebrafish genes and ESTs with likely human (or in a few cases, mouse) orthologs to 523 has revealed extensive conserved synteny between the zebrafish and human genomes. We find 80% of genes and ESTs in this analysis fall in con-

served synteny groups, averaging 3.7 genes/syntenic group. A previous analysis of 124 zebrafish genes and ESTs identified only 64% (79/124) in conserved synteny groups, averaging 2.8 genes/group (Gates et al. 1999). Presumably, as more and more zebrafish genes and ESTs are mapped, the fraction that fall in synteny groups will continue to increase, and may approach 100%. Similarly, Gates et al. (1999) identified 28 synteny groups between zebrafish and human, and our analysis increases this number to 113. The existence of yet unidentified synteny groups is suggested by the 102 genes and ESTs in the singleton class. Singletons may reflect errors in mapping or in orthology determination, or may instead nucleate additional synteny groups as additional genes are mapped. Using the singleton class for Poisson analysis (and assuming no error) predicts a further 69 synteny groups as yet undiscovered. This allows us to predict an upper limit for synteny groups between zebrafish and human of 284 (113 + 102 + 69).

The finding that most zebrafish genes in this study are in conserved synteny groups with human genes

**Table 3.** Predicting Orthology Using Synteny Relationship

Zebrafish gene	Reference NCBI gi	Zebrafish map position	Human synteny predictions <sup>a</sup>	Possible human orthologues	Reference NCBI unigene	Human map position
bact	3044209	1.59cR <sup>b</sup>	1, 2	<b>ACTB</b> ACTG1 ACTC	<b>Hs.180952</b> Hs.204867 Hs.118127	<b>1.49-82cM</b> 17.118-129cM 15.25-32cM
bact2	2822455	3.304cR <sup>g</sup>	16, 17	<b>ACTG1</b> ACTB ACTC	<b>Hs.204867</b> Hs.180952 Hs.118127	<b>17.118-129cM</b> 1.49-82cM 15.25-32cM
brn1.2	222975	6.218cR <sup>d</sup>	1, 2, 9, 17	<b>POU3F1</b> POU3F2 POU3F3 POU3F4	<b>Hs.1837</b> Hs.182505 Hs.248158 Hs.2229	<b>1.49-82cM</b> 6.91-96cM 3.80-100 X.97-105cM
elrd	608548	8.108cR <sup>d</sup>	1	<b>ELAVL4</b> ELAVL2	<b>Hs.75236</b> Hs.3198	<b>1.49-82cM</b> 9.57-93cM
frz-zg01	1245183	15.272cR <sup>g</sup>	2, 3, 11, 17	<b>FZD4</b> FZD9	<b>II.8322</b> Hs.158335	<b>11.84-100cM</b> 7.84-91cM
glr	3378595	14.433cR <sup>b</sup>	5, 11, 12	<b>GLRA1</b> GLRA3 GLRA2	<b>Hs.121490</b> Hs.167742 Hs.2700	<b>5.153-158cM</b> 4.170cM X.0-42cM
groucho1	2104717	7.119cR <sup>b</sup>	11, 15, 16	<b>TLE3</b> TLE1 TLE4 TLE2	<b>Hs.167086</b> Hs.28935 Hs.83958 Hs.173063	<b>15.70-71cM</b> 9 9.77.7-82.3cM 19.0.0-31.9cM
hha	N/A	9.140cR <sup>e</sup>	2	<b>IHH</b> SHH	<b>Hs.69351</b> Hs.121539	<b>2.200-215cM</b> 7.181-184cM
ldb4	3078004	13.278cR <sup>g</sup>	2, 6, 10	<b>LDB1</b> LDB2	<b>Hs.26002</b> Hs.4980	<b>10.114-131cM</b> 4.0-32cM
msxa	608508	14.464cR <sup>e</sup>	5	<b>MSX2</b> MSX1 MSX3	<b>Hs.89404</b> Hs.194 Mm.4816	<b>5.185-199cM</b> 4.4-28cM 10.170-182cM <sup>c</sup>
msxb	608510	1.381cR <sup>b</sup>	4, 13, 14	<b>MSX1</b> MSX2 MSX3	<b>Hs.194</b> Hs.89404 Mm.4816	<b>4.4-28cM</b> 5.185-196cM 10.170-182cM <sup>c</sup>
msxc	399912	13.312cR <sup>d</sup>	6, 10	<b>MSX3</b> MSX1 MSX2 MSX3	<b>Mm.4816</b> Hs.194 Hs.89404 Mm.4816	<b>10.170-182cM<sup>c</sup></b> 4.4-28cM 5.185-196cM 10.170-182cM <sup>c</sup>
msxd	62544	21.211cR <sup>d</sup>	5, 7, 10	<b>MSX2</b> MSX2 MSX3	<b>Hs.89404</b> Hs.89404 Mm.4816	<b>5.185-196cM</b> 5.185-196cM 10.170-182cM <sup>c</sup>
msxe	1399516	14.27cR <sup>d</sup>	5, 6, 8, 22	<b>MSX2</b> MSX1 MSX3	<b>Hs.89404</b> Hs.194 Mm.4816	<b>5.185-196cM</b> 4.4-28cM 10.170-182cM <sup>c</sup>
otx3	633134	1.381cR <sup>d</sup>	4, 7, 14	<b>OTX2</b> OTX1	<b>II.5015</b> II.5013	<b>14.0-1cM</b> 2.84-88cM
plasticin	1881763	11.390cR <sup>f</sup>	3, 12, 17	<b>PRPH</b> VIM	<b>Hs.37044</b> Hs.2064	<b>12.53-70cM</b> 10.40-44cM
rtk7	3005904	24.301cR <sup>b</sup>	4, 8	<b>EPHA5</b> EHK-1 EPHNA4 EPHA7 EPHA3	<b>Hs.31092</b> Hs.194771 Hs.739641 Hs.73962 Hs.123642	<b>4.67.7-77.9cM</b> N/A N/A 6.101-104cM 3.111-113cM
zef1	4099173	14.534cR <sup>d</sup>	4, 5, 12, X	<b>ELF4</b> ELF1	<b>Hs.151139</b> Hs.154365	<b>X.150-184cM</b> 13.37-46cM
fb38g02		6.115cR <sup>b</sup>	2, 19	<b>FZD7</b> FZD2	<b>Hs.173859</b> Hs.81217	<b>2.200-212cM</b> 17.74-75cM
fb18b11		24.388cR <sup>b</sup>	1, 8	<b>UBE2V2</b> UBE2V1 FZD10	<b>Hs.79300</b> Hs.75875 Hs.31664	<b>8.66-67cM</b> 20.74-75cM 12.160-169cM

Human genes in bold are orthologues predicted by syntenic correspondence.

<sup>a</sup> Corresponding human synteny group or groups for zebrafish genes in same mapping bin or flanking positions to zebrafish gene in column 1.

<sup>b</sup> Genes and ESTs mapped in this study.

<sup>c</sup> Corresponding human map position inferred from human-mouse syntenic relationship and mouse gene position.

<sup>d</sup> Hukriede et al. 1999.

<sup>e</sup> Postlethwait et al. 1998.

<sup>f</sup> Gates et al. 1999.

<sup>g</sup> Geissler et al. 1999.

raises the possibility that significant portions of the zebrafish genome are uninterrupted by rearrangements since the teleost–tetrapod divergence. Indeed, we find that 292 of the genes and ESTs analyzed in this study define 118 homology segments (uninterrupted segments with conserved map order) covering ~56% of the zebrafish genome (assuming random marker distribution). Taking into account the  $1.7 \times 10^9$  bp size of the haploid zebrafish genome (Hinegardner 1968), we suggest an average size of  $8.1 \times 10^6$  bp/homology segment identified in this study. This analysis suggests that zebrafish workers wishing to positionally clone zebrafish mutant genes can profitably use the syntenic comparison between zebrafish and human to identify candidates from the nearly complete human genome sequence.

Comparative biology often utilizes functional analysis of orthologous gene pairs, yet gene orthology is not always solvable by sequence comparison. For instance, members of multigene families may be too similar for BLAST or phylogenetic methods to unambiguously distinguish orthologous pairs of genes. One alternative to sequence-based orthology determination is a synteny-based approach. Such an approach first requires an understanding of the syntenic relationship between species compared. We suggest that the extensive correspondence between the human and zebrafish genomes revealed by this analysis can be used in predicting orthologous gene relationships. Of 32 zebrafish genes or ESTs whose human ortholog could not be unambiguously identified by BLAST analysis (data not shown), we suggest a human ortholog for 20 of these based on the syntenic correspondence of the zebrafish and human genomes (Table 3). Examples of such predictions include members of the zebrafish *msx* gene family. BLAST analysis fails to confidently predict the orthology relationships between the zebrafish *msxa*, *msxb*, *msxc*, *msxd*, or *msxe* genes and the human *MSX1* and *MSX2* and mouse *MSX3* genes. Phylogenetic analysis (data not shown), suggests that zebrafish *msxb* and *msxc* are orthologous to mouse *Msx3* (the human ortholog has not been identified), and zebrafish *msxe* is orthologous to human *MSX1*. We can use synteny as an alternative predictor of orthology, which suggests that *msxa*, *msxd*, and *msxe* are orthologous to *MSX2*; zebrafish *msxb* is orthologous to *MSX1*; and zebrafish *msxc* is orthologous to mouse *MSX3*. The addition of more genes to the zebrafish genetic map may further resolve this issue.

Recent observations suggest a whole genome duplication occurred in the teleost lineage since its divergence from the tetrapod lineage (Amores et al. 1998; Postlethwaite et al. 1998; Wittbrodt et al. 1998; Gates et al. 1999). Consistent with this notion are the 38 examples where two or more mapped, unlinked zebrafish genes share a single mammalian ortholog, dis-

tributed among 20 of the 25 zebrafish chromosomes. The alternative hypothesis, that the duplications observed may have accrued individually, rather than in a single, whole-genome event, cannot yet be excluded. Indeed, instances of three zebrafish orthologs for a single human gene may argue for some role of regional duplication in generating duplicate copies of zebrafish genes. For instance, two of the three *ISL1* orthologs, *islet2* and *islet3*, map to a similar location on LG 25 (Geissler et al. 1999; Hukriede et al. 1999), and thus may have arisen by a tandem duplication. Identifying the syntenic relationship between the entire zebrafish and human genome may help resolve this issue.

A full understanding of the role of human genes in development and physiology will require models where gene function can be examined readily. Forward mutant screens in zebrafish are performed routinely, resulting in sizable collections of mutations causing a variety of developmental and physiological defects (e.g., Driever et al. 1996; Haffter et al. 1996; Henion et al. 1996). Molecular analysis of these mutations is beginning to reveal their utility as models for human disease (Zon 1999). Furthermore, the zebrafish is being established as a genetic and physiological model for vertebrate-specific processes such as organogenesis (Zhong et al. 2000). Knowledge of the relationship between the zebrafish and human genomes will provide the link to compare zebrafish genes and mutations with their orthologous human genes and diseases.

## METHODS

### RH Mapping and Map Construction

RH mapping was performed as described (Hukriede et al. 1999) on the LN54 zebrafish RH panel. Briefly, STS primers for genes were designed from 3' ends of gene sequences obtained from GenBank (<http://www.ncbi.nlm.nih.gov>), or for representative 3' EST reads preselected for highly significant WU-BLASTX matches to the nonredundant protein database (<http://zfish.wustl.edu>). Primer sequences were designed using OSP (Hillier and Green 1991), (see <http://zfish.wustl.edu> for primer sequences). Each marker was positioned relative to the LN54 framework (Hukriede et al. 1999) using the RHMAPPER radiation hybrid mapping program (<http://waldo.wi.mit.edu/ftp/distribution/software/rhmapper/>) by web submission of the RH vector to <http://mgchd1.nichd.nih.gov:8000/zfrh/beta.cgi>, and placed accordingly in the bin following the framework marker, using the position of the framework marker to denote their position on the map.

### Orthology Prediction

Each mapped zebrafish EST or gene was subjected to extensive WU-BLASTX and WU-BLASTN (filter = seg,  $E = 1e^{-10}$ ) (W. Gish, unpubl.; <http://blast.wustl.edu>) analysis against the comprehensive GenBank EST database, release 113 (<http://ncbi.nlm.nih.gov>) as well as the nonredundant protein and nucleotide database. The reports were postprocessed to recover the top matching hits from zebrafish, and the top EST, protein, and nucleotide hits from human sequences. All align-

ments were assessed manually, using a BLASTN cutoff at a maximum p value of  $e^{-20}$  (the vast majority of predicted ortholog showed matches with p values  $< e^{-40}$ ). Zebrafish-human sequence pairs identified as putative orthologs by BLASTN similarity were likewise confirmed by BLASTX similarity. When available, we determined the UniGene reference sequence (<http://www.ncbi.nlm.nih.gov/UniGene/>) representing the human ortholog and acquired its Gene Map 98 map location (Deloukas et al. 1998; <http://www.ncbi.nlm.nih.gov/genemap98>). In some cases human mapping data was obtained from Online Mendelian Inheritance in Man (OMIM) (<http://www.ncbi.nlm.nih.gov/Omim>). All zebrafish-human orthologous pair BLASTN/BLASTX results, GenBank accession numbers, GenBank records, human reference numbers, and map positions are available at <http://www.zfish.wustl.edu>.

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