

Recent segmental and gene duplications in the mouse genome

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Abstract

Background: The high quality of the mouse genome draft sequence and its associated annotations are an invaluable biological resource. Identifying recent duplications in the mouse genome, especially in regions containing genes, may highlight important events in recent murine evolution. In addition, detecting recent sequence duplications can reveal potentially problematic regions of the genome assembly. We use BLAST-based computational heuristics to identify large (≥ 5 kb) and recent ($\geq 90\%$ sequence identity) segmental duplications in the mouse genome sequence. Here we present a database of recently duplicated regions of the mouse genome found in the mouse genome sequencing consortium (MGSC) February 2002 and February 2003 assemblies.

Results: We determined that 33.6 Mb of 2,695 Mb (1.2%) of sequence from the February 2003 mouse genome sequence assembly is involved in recent segmental duplications, which is less than that observed in the human genome (around 3.5-5%). From this dataset, 8.9 Mb (26%) of the duplication content consisted of 'unmapped' chromosome sequence. Moreover, we suspect that an additional 18.5 Mb of sequence is involved in duplication artifacts arising from sequence misassignment errors in this genome assembly. By searching for genes that are located within these regions, we identified 675 genes that mapped to duplicated regions of the mouse genome. Sixteen of these genes appear to have been duplicated independently in the human genome. From our dataset we further characterized a 42 kb recent segmental duplication of *Mater*, a maternal-effect gene essential for embryogenesis in mice.

Conclusion: Our results provide an initial analysis of the recently duplicated sequence and gene content of the mouse genome. Many of these duplicated loci, as well as regions identified to be involved in potential sequence misassignment errors, will require further mapping and sequencing to achieve accuracy. A Genome Browser database was set up to display the identified duplication content presented in this work. This data will also be relevant to the growing number of investigators who use the draft genome sequence for experimental design and analysis.

Background

The evolutionary trajectory of duplicated genes has been an active area of investigation since gene duplication was first recognized as an important force in species evolution [1]. The availability of new sequence data and analyses has challenged the hypothesis suggesting most duplicated genes are destined to lose their function and become pseudogenes, with a few exceptions establishing new biological roles (reviewed in [2]). It is believed that the occurrence of gene duplication would result in relaxed selection of redundant copies permitting genes to evolve specialized sub-functions [3,4]. Moreover, nearly identical genomic regions provide important substrates for chromosomal rearrangements that permit rapid evolutionary changes to occur in a short period of time [5].

An estimated 3.5-5% of the human genome has undergone recent duplication [6-9], and these segmental duplications (also termed duplicons or low-copy repeats) are found to be hot spots, or predisposition sites, for the occurrence of non-allelic homologous recombination. This recombination can lead to genomic mutations such as deletion, duplication, inversion, or translocation, resulting in human disease [10]. Many mouse strains with chromosomal aberrations are known [11], and it remains to be seen whether segmental duplications have a role in any of these genomic mutations.

A high-quality draft genome sequence not only makes it possible to expand the known set of duplicate genes, but also reveals their genomic context. This genomic context contains the regulatory and structural elements responsible for gene expression that need to be interrogated for a better understanding of the mechanisms and consequences of gene duplication. Furthermore, an accurate and well-annotated mouse genome is an essential resource for many in the biomedical research community, especially those who use the sequence to design and interpret transgenic, mutagenesis, microarray, and proteomic studies.

Several lines of evidence show that the whole-genome shotgun (WGS) approach yielded a high-quality draft sequence that covers roughly 96% of the euchromatic genome excluding chromosome Y (a female C57BL/6J mouse was used in the sequencing project) [12]. The WGS sequence reads were assembled into sequence contigs using sequence-assembly programs to produce the February 2002 MGSCv3 working draft [12,13]. The newly released February 2003 assembly was a hybrid assembly comprising 705 megabases (Mb) of finished bacterial artificial chromosome (BAC) sequences incorporated into the MGSCv3 assembly.

We previously analyzed several versions of the human genome draft assemblies (NCBI Builds 28, 29, and 30) [9], and found substantial potential genome assembly errors in all builds, including approximately 40 Mb of sequence in Build 30. These assembly errors probably arose from difficulties in merging finished sequence or from incorrectly assigning

sequence contigs into the genome assembly. In such cases, completely identical or nearly identical sequences (due to allelic differences or sequencing errors) would be present at distinct regions in the genome sequence. These sequence mis-assignment errors would yield near-perfect duplication artifacts, detected as having extremely high sequence identities (exceeding 99.5% and over 5 kilobase (kb) in length), in genome assemblies. However, a small subset of such results could represent duplications that arose from very recent evolutionary events and will require further experimental analysis.

A number of web-based resources, specifically those provided by the National Center for Biotechnology Information (NCBI), Ensembl (at the European Bioinformatics Institute and Sanger Centre), and the University of California Santa Cruz (UCSC), make the genome sequence and associated annotations readily accessible. Because of the success of the mouse genome sequencing consortium (MSGC), investigators worldwide are utilizing the draft 'as is' in both medical and evolutionary studies. In this paper we show that even though the genome assembly is still in draft form, an initial analysis of the sequence can reveal novel genomic duplications and demarcate regions of the genome that require additional examination.

Results and discussion

We performed a search for all recent segmental duplications that were larger than 5 kb in size and showed greater than 90% sequence identity from both the February 2002 (numerical results for February 2002 assembly are presented at our web site [14]) and the February 2003 mouse genome sequence assemblies [15]. Our method was based on pairwise (mega-) BLAST2 [16] sequence comparisons between entire chromosome sequences. From our analysis of the February 2003 assembly, a total of 33.6 Mb (1.2%) of the genome sequence (2,695 Mb) was found to be involved in recent segmental duplications (Table 1) and 8.9 Mb of this sequence was unmapped data (found in the unmapped chromosome sequence). On the basis of the 20 mapped chromosomes, more than 712 distinct intrachromosomal segmental duplications, comprising 19.9 Mb of sequence (Figure 1), and 475 distinct interchromosomal duplications, comprising 7.1 Mb of sequence, were identified. We also found that 57% of the duplications were in tandem, which we defined as two related intrachromosomal duplicons located within 200 kb of one another.

Duplications can be found in all chromosomes analyzed, with chromosomes 6, 7, 17, and X having the highest, and chromosome 18 having the least, duplicated content (Table 1, Figure 1). Substantial amounts (8.9 Mb) of the duplicated content are found in the unmapped chromosome (ChrUn) sequence, suggesting that the correct chromosomal assignment of these segments remains a major assembly challenge. It is possible

that small subsets of these duplications are due to chimeric reads and other sequencing artifacts and thus should not be part of the finished genome sequence. On the other hand, these unmapped duplicated sequences represent true duplications that have been excluded from the assembly. One example of this occurs with a member of the mouse *Bcl2* family of apoptosis regulators, *Bcl2a1*. *Bcl2a1* contains four highly similar genes (> 97% identical at the nucleotide level) that have been mapped together on chromosome 9 of the C57BL/6 and 129SV genomes [17,18]. Currently, the *Bcl2a1* genes are not assembled on the mapped chromosome and are found in three distinct unmapped contigs. In the human genome only one copy of *BCL2A1* is found, although a recent, independent 8.5 kb tandem duplication containing the last exon of *BCL2A1* has occurred, forming a novel *BCL2A1*-related transcript (AF249277). An example of a region that has changed between assemblies is the *Amy2* locus. *Amy2* is known to vary in copy number between inbred strains of mice [19]. In the February 2002 assembly, only one copy of the *Amy2* gene resided on chromosome 3 in addition to a second copy found on a large 10 kb unmapped contig. In addition, partial high identity matches (> 95%) to four distinct unmapped contigs were found (note that these partial copies were not detected in our analysis as they are less than 5 kb long). In the February 2003 assembly, six *Amy2* genes exist, which is close to the five *Amy2*-like genes that were detected in the genome of strain A/J mice using quantitative densitometry of Southern blots [20]. It is, however, important to note that a gap, not bridged by a clone, still exists between the *Amy2* locus and the *Amy1* gene, and so the copy number in the C57BL/6J genome assembly may still vary.

We analyzed the distribution of segmental duplication content by sorting the duplications into six different sequence-similarity categories: 90-92%, 92-94%, 94-96%, 96-98%, 98-99.5%, and 99.5-100%, for both the February 2002 and 2003 assembly builds (Table 2). The amount of duplication content appears to be unevenly distributed across these categories, with a distinct rise in the 94-96% category. This might suggest recent duplicative events in the mouse genome have not occurred at a steady rate. However, it is unclear at this point how these results were affected by the draft status of the genome assemblies. Between the 2002 and the 2003 assembly builds we found that the amount of duplication content is nearly the same within each percent category except for the 99.5-100% category, which contained 4.8 Mb of sequence in 2002 and 18.5 Mb in 2003 (Table 2). Furthermore, we determined that the majority (88%) of the duplicated sequence in the 99.5-100% category occurred intrachromosomally, within

200 kb of each other. Using the assembly component tables (provided by UCSC [21]), which contain information about the underlying makeup of the February 2003 genome assembly (shotgun-assembled scaffolds and BAC sequences), we found that 215/216 (99.5%) of these duplications involved a BAC sequence. Hence, we suspect that the large increase in near-identical duplications could be the result of sequence misassignment errors arising from the inherent difficulty of merging finished BAC sequence with shotgun sequence contigs.

We previously observed that the human genome sequence assembled by Celera's WGS method [22] showed poor quality in regions with near-identical segmental duplications [23]. To assess the finishing status of duplicated regions in the WGS mouse genome assembly (February 2002 MGSCv3 assembly), we calculated the amount of unfinished sequence (regions with gaps or Ns) within the immediate neighborhood (20 kb) of each duplicon (the unmapped chromosome sequence was excluded from this analysis). We observed substantially higher amounts of unfinished sequence (number of Ns) in these regions. Whereas 8.0% of the assembly is comprised of Ns, regions harboring duplications contain an average of 12.2%. This average rises to 16.6% for duplications with more than 98% sequence identity (statistics can be obtained from our website [14]). This suggests that the WGS assembler had difficulty assembling regions containing recent sequence duplication and that these regions are good candidates for finishing using clone resources.

Using the NCBI Refseq and Ensembl mouse gene annotation, we identified 675 genes that mapped to duplicated regions of the mouse genome (a full list of genes can be obtained from our website [14]); 414 of these genes were found to be fully contained within a segmental duplication, thus representing the best candidates for whole-gene duplication. While it is likely that some of these duplicate copies have become pseudogenes, others may have evolved specialized functions [3]. Moreover, we sought to use the identified gene sequences, which were expressed sequence tags (ESTs) and/or cDNAs, as experimentally derived resources to help validate the genomic duplication content presented in this study. We aligned duplicated gene sequences to each genomic region using UCSC BLAT [17] and determined their percent identity matches. Unambiguous gene-to-genomic identity matches were established for all 128 gene pairs we examined. Each gene sequence was mapped to their respective genomic region with at least 99.1% identity (examples are shown in Table 3; a full table is available at [14]). We also examined the

Figure 1 (see following page)

Intrachromosomal segmental duplications identified in the mouse genome (chromosomes 1-X; results are based on the February 2003 assembly). Each line represents a duplicated module and connects a paralogous duplicon pair. Red, 99-100% sequence identity; purple, 96-98%; green, 93-95%; and blue, 90-92%. Correspondences to chromosome ideograms (obtained from Ensembl) are only crude. Graphics were produced using GenomePixelizer [34].

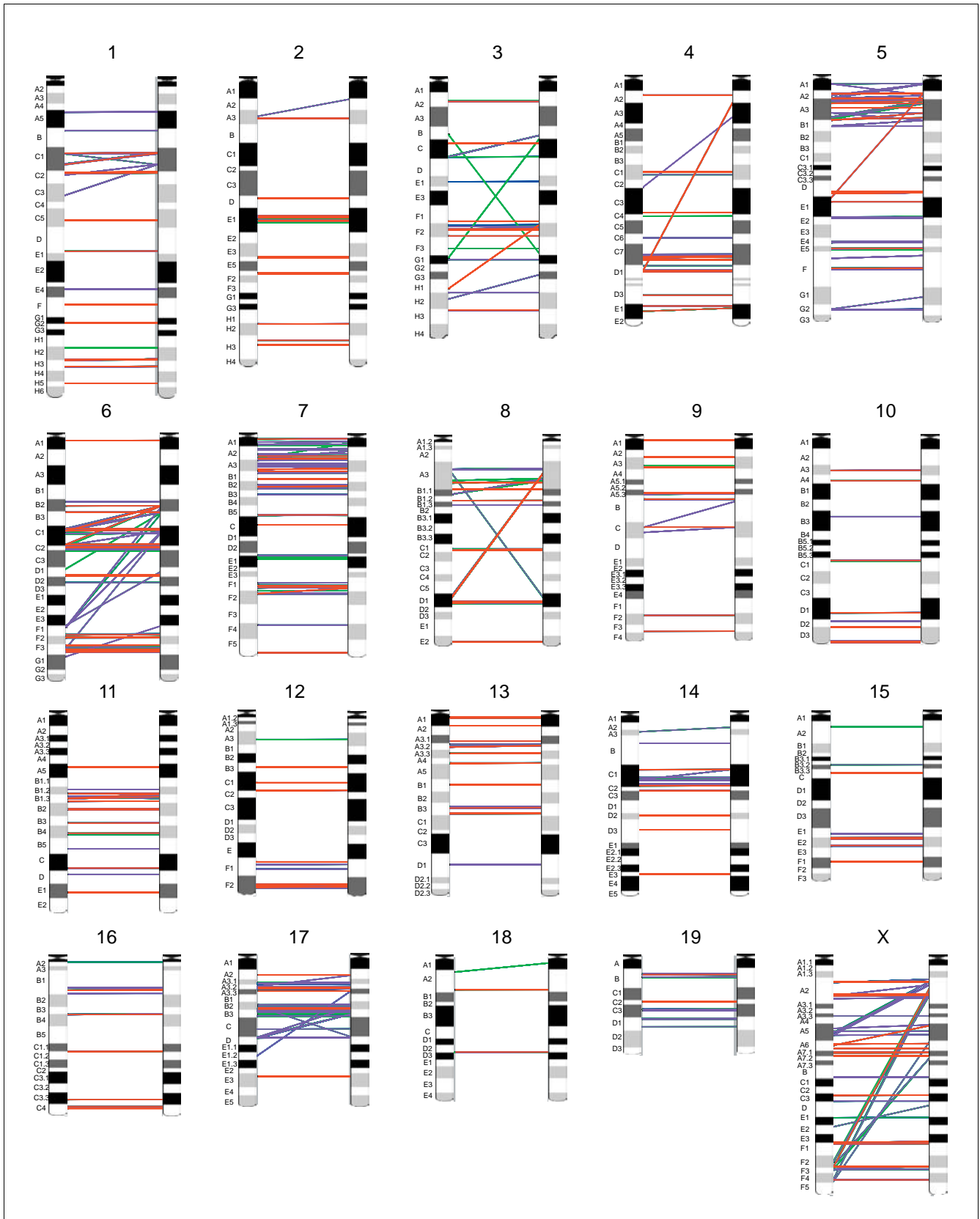


Figure 1 (see legend on previous page)

Table 1**Recent segmental duplication in the mouse genome**

Chromosome	Chromosome length	Intrachromosomal duplication	%	Interchromosomal duplication	%	Total	%
1	195,869,683	1,392,568	0.7	238,739	0.1	1,552,908	0.8
2	181,423,755	1,106,879	0.6	173,602	0.1	1,184,304	0.7
3	160,674,399	790,500	0.5	158,011	0.1	948,511	0.6
4	152,921,959	1,743,027	1.1	647,795	0.4	1,921,970	1.3
5	149,719,773	1,102,772	0.7	761,950	0.5	1,560,683	1.0
6	149,950,539	2,042,585	1.4	562,415	0.4	2,339,839	1.6
7	134,401,573	1,655,438	1.2	713,287	0.5	2,038,845	1.5
8	128,923,138	738,203	0.6	331,970	0.3	1,005,575	0.8
9	124,467,299	437,352	0.4	188,427	0.2	623,089	0.5
10	130,738,012	345,768	0.3	258,429	0.2	604,197	0.5
11	122,862,689	900,355	0.7	127,774	0.1	1,012,479	0.8
12	114,462,600	1,139,786	1.0	374,365	0.3	1,404,279	1.2
13	116,242,670	855,835	0.7	547,462	0.5	1,349,974	1.2
14	115,844,145	450,161	0.4	451,782	0.4	748,465	0.6
15	104,111,694	443,805	0.4	43,937	0.0	487,742	0.5
16	98,986,639	389,255	0.4	67,290	0.1	456,545	0.5
17	93,529,596	1,329,664	1.4	660,440	0.7	1,760,982	1.9
18	91,041,441	162,916	0.2	58,996	0.1	210,422	0.2
19	61,093,376	328,909	0.5	193,387	0.3	479,687	0.8
X	149,996,094	2,592,361	1.7	574,950	0.4	3,018,682	2.0
chrUn*	117,911,829	6,049,538	5.1	5,710,057	4.8	8,885,604	7.5
Total	2,695,172,903	25,997,677	1.0	12,845,065	0.5	33,594,782	1.2

The analysis is based on the February 2003 mouse genome assembly. *chrUn, unmapped chromosome sequence.

identified duplicated genes using their InterPro protein-domain classification present in 608 Ensembl genes to see whether specific kinds of genes or protein domains have been preferentially duplicated. We found that genes containing protein domains related to signal transduction (rhodopsin-like G-protein-coupled receptor superfamily), olfaction (olfactory receptors, vomeronasal receptors) immunity (immunoglobulin/MHC, serine protease), and drug metabolism (cytochrome P450) are significantly enriched (by at least threefold) (Table 4).

From this list of genes, we performed a detailed analysis of *Mater*, a maternal-effect gene of potential medical importance. *Mater* encodes an autoantigen in a mouse model for human autoimmune premature ovarian failure [24]. Knock-out studies have shown that it is essential for early embryonic development in mice [25]. *Mater* encodes a protein of 1,111 amino acids from a 3.5 kb transcript that spans 57 kb on mouse chromosome 7. A 42 kb segmental duplication involving two duplons (DUP1, where *Mater* is located; DUP2,

where a novel *Mater2* is located) are situated about 5 Mb apart and in an inverted orientation (Figure 2). DUP1 and DUP2 are on average 91.1% identical over the entire 42 kb genomic region, with a 96.6% average in the exonic regions. Furthermore, we identified an intron-less *Mater* pseudogene (*MaterP*), which shares 87% DNA sequence identity to *Mater*, at a location 10 Mb proximal to *Mater* (Figure 2; see Additional data files for a detailed comparative genomic analysis of the *Mater* locus). The mapping locations of these duplications have been confirmed by fluorescence *in situ* hybridization (FISH) (Figure 3). Thus, *Mater* serves as one example of a gene that has been knocked out in mice but for which there is a second, highly similar transcript whose biological role is not yet known.

In addition, we were interested in determining whether any of the 675 genes have undergone recent ($\geq 90\%$ sequence identity over ≥ 5 kb) and independent duplication in the human genome. Some of these genes could be recently evolving via the 'birth and death model of evolution' which has been used

Table 2**Comparison between genome assemblies**

Sequence identity level	February 2002 assembly*	February 2003 assembly
Duplication content (bp)		
90-92%	4,966,470	3,543,429
92-94%	15,685,840	13,981,642
94-96%	17,533,730	17,970,287
96-98%	11,539,392	11,731,958
98-99.5%	5,865,024	5,487,899
†Potential sequence misassignment error detected (bp)		
99.5-100%	4,832,594	18,456,096

The comparison is of duplication content by sequence identity and potential sequence misassignment errors between the February 2002 (MGSCv3) and February 2003 (a hybrid assembly of MGSCv3 with 705 Mb finished BAC sequence) genome assemblies. *Analysis of the duplication content for February 2002 assembly can be found at [14]. †Sequences detected to show extremely high percent identity duplications are likely to be genome assembly artifacts and were not included in the duplication content shown in Table 1.

to describe the evolution of the major histocompatibility complex (MHC) and immunoglobulin multigene families [26]. This model describes genes that are repeatedly created through duplication, with some genes becoming fixed while others are rendered nonfunctional by deleterious mutations [26].

We examined the 675 duplicated mouse genes using best reciprocal BLAST hits to identify their putative human orthologs. We subsequently analyzed regions containing these putative orthologs for recent sequence duplication in the human genome. Sixteen of the 675 genes were found to be involved in recent, independent gene duplication in mouse and human (see Table 5). Some of these regions containing whole-gene duplications are part of multigene families known to be evolving via duplication and are found in tandem duplicated arrays in both species (that is, the *Amy2*, *H2-Q1*, *Gst1*, and *Olf54* genes). An interesting example of a recent and apparently independent whole-gene duplication that occurred in mouse and human involves *Bmp8a* and a second intronic transcript *Oxct2*. Of the partial gene duplications, the recent duplication within the *Tnxb* gene and its human ortholog *TNXXB* (found at the MHC III locus of mouse chromosome 17 and human 6p21) is particularly intriguing. In humans, this locus consists of a tandem array of genes (*RP*, *C4*, *CYP21*, and *TNXXB* (RCCX)), which through gene duplication, can exist as mono-, di- and tri-modular forms in the caucasian population [27]. Recent studies have also shown the presence of a deletion haplotype in one individual, leading to a fusion of the *TNXXA/TNXXB* gene on one chromosome and a

duplication of *CYP21* on the other chromosome [28]. Furthermore, complex haplotypes of the complement genes (*C4A* and *C4B*) residing in the RCCX module have been characterized and postulated to have a role in individual susceptibility to infection and autoimmune disease [29]. A closer inspection of the genomic region surrounding this recent duplication in the mouse reveals that the C57BL/6J duplication encompasses homologous genes (*Tnxb*, *Slp* (a *C4* paralog), *Cyp21a1*, and *C4*). Similarly, in humans, this orthologous region of the mouse genome has been shown to undergo multiple recombination events, giving rise to a variety of haplotypes [30]. Overall, many of the genes that have recently experienced duplications in the mouse and human genomes are of biomedical and evolutionary interest. The complexity and polymorphic nature of these recent duplications underscores the need for, and the difficulty of, performing the detailed structural and functional analyses that will help discern their true genomic organization, evolutionary history, and biological implications.

Conclusions

Our current analysis of the presence and organization of recent segmental duplications in the mouse genome has identified recent gene-duplication events and potentially problematic regions of the mouse genome assembly. At a practical level, identifying regions with segmental duplication will be useful in highlighting the most dynamic regions of any mammalian genome assembly. For the genome-sequencing community, these potential misassemblies/putative duplications can become initial targets for clone-based finishing; and for the biologist, they can serve as sentinels for regions of the genome most likely to change in subsequent assemblies. Additional hierarchical shotgun sequencing effort [12] will undoubtedly be critical to finish the mouse genome sequence and reveal additional duplicated regions that are incomplete at the moment.

Many of the duplicated genes are of evolutionary and medical importance (that is, genes involved in immune defense, olfaction, and drug metabolism). Knowledge of these duplicated regions could be important for accurately mapping mutants derived from ethylnitrosourea (ENU) mutagenesis, designing targeting vectors for embryonic stem cell alterations, and validating putative single-nucleotide polymorphisms (SNPs) that may have arisen from recently duplicated sequences rather than allelic variants [31]. The ability to create large, sophisticated targeting vectors, by engineering BACs using homologous recombination in *Escherichia coli* [32], should prove very useful for designing *in vivo* experiments aimed at dissecting the function of recently duplicated genes. Knowledge of all recent duplications in mouse may also highlight regions subject to chromosomal rearrangement and polymorphism within and between species, and provide an opportunity to model the stability of such genomic architecture in a mammalian genome.

Table 3**Examples of recent mouse gene duplications**

Locus1*	Gene	Percent identity†	Annotation	Locus2*	Gene	Percent identity†	Annotation	Duplication % identity‡
1 F	NM_009888	99.6	Cfh (Complement component factor h)	1 F	M29010	99.0	Complement factor H-related protein mRNA	97.1
3 G1	NM_009669	100	Amy2 (Amylase 2, pancreatic)	3 G1	M11896	99.6	Pancreatic amylase B-1	97.6
5 E2	NM_053184	99.9	Ugt2a1 (UDP glycosyltransferase 2 A1)	5 E2	BF144793	99.6	cDNA clone IMAGE:4021939	95.7
5 E2	NM_009467	100	Ugt2b5(UDP-glucuronosyltransferase 2b5)	5 E2	NM_053215	100	RIKEN cDNA 0610033E06 gene	93.3
5 E4	NM_008620	99.9	Mpa2 (macrophage activation 2)	5 E4	BC007143	99.5	Similar to macrophage activation 2	90.6
5 G1	NM_029693	100	RIKEN cDNA 1700123K08	7 B2	NM_027702	100	RIKEN cDNA 4933421I07 gene	91.1
6 C1	NM_053238	100	V1rc8 (Vomer nasal I receptor, C8)	6 C1	NM_053239	99.7	V1rc9 (Vomer nasal I receptor, C9)	95.1
6 D1	NM_011467	99.9	Spr (sepiapterin reductase)	6 D1	BE862957	99.5	EST sequence	95.8
6 F1	AI505330	100	Similar to initiation factor eIF-4A1	6 F1	AI503670	99.8	Similar to initiation factor eIF-4A1	98.9
6 F2	NM_008646	99.9	Mug2 (Murinoglobulin 2)	6 F2	NM_008645	99.9	Mug1 (Murinoglobulin 1)	94.6
6 F3	NM_020257	99.8	Dcl1 (c-type lectin 1)	6 F3	NM_027562	99.9	4632413B12Rik (C-lectin related protein)	90.8
6 F3	NM_008463	99.7	Klra5 (Killer cell lectin-like receptor, A5)	6 F3	NM_008464	99.5	Klra6 (Killer cell lectin-like receptor, A6)	90.0
6 F3	NM_010649	99.8	Klra4 (Killer cell lectin-like receptor A4)	6 F3	NM_016659	99.8	Klra1 (Killer cell lectin-like receptor A1)	91.4
6 F3	NM_010737	99.8	Klrb1b (Killer cell lectin-like receptor 1b)	6 F3	NM_008527	99.9	Klrb1c (Killer cell lectin-like receptor 1c)	90.8
7 A2	NM_011860	100	Mater (Maternal effect gene)	7 A1	AK016782	100	Similar to Mater protein	96.6
7 B1	NM_032541	100	Hamp hepcidin antimicrobial peptide	7 B1	AK007975	99.8	Prohepcidin homolog	92.8
7 B2	NM_010115	99	Klk13 (Kallikrein 13)	7 B2	NM_008454	99.9	Klk16 (Kallikrein 16)	92.2
8 D1	LI1333	99.9	Carboxylesterase	8 D1	NM_144511	100	Es31	95.2
9 F4	NM_130864	99.6	Acaa acetyl-Coenzyme A acyltransferase	9 F4	BC019882	100	Similar to acetyl-CoA acyltransferase	96.6
10 B3	NM_013532	99.9	Gp49a (Glycoprotein 49A)	10 B3	NM_008147	100	Gp49b (glycoprotein 49B)	96.7
10 D2	NM_017372	100	Lyzs (Lysozyme)	10 D2	NM_013590	99.8	Lzp-s (P lysozyme structural)	95.3
11 A3.2	NM_172792	100	hypothetical protein 4932414J04	17 D	AK03001	100	Tyrosine protein kinase/ cysteine-rich region	94.0
11 B1.3	NM_011396	99.9	Slc22a5 (Solute carrier family 22)	11 B1.3	NM_019723	100	Slc22a9 (solute carrier family 22)	91.2
11 D	NM_021347	100	Gsdm (Gasdermin)	11 D	NM_029727	99.9	2200001G21Rik	94.2
12 F1	BC002065	99.6	Serine protease inhibitor 2-1	12 F1	BY761363	99.9	EST sequence	92.2
12 F1	NM_013772	100	Tcl1b3 (T-cell leukemia/lymphoma 1B, 3)	12 F1	NM_013776	100	Tcl1b5 (T-cell leukemia/lymphoma 1B, 5)	95.3
13 A1	NM_013778	99.5	Akr1c13 (Aldo-keto reductase 1, C13)	13 A1	NM_013777	99.5	Akr1c12 (Aldo-keto reductase 1, C12)	96.1
13 A3	NM_008864	99.2	Csh1 (chorionic somatomammotrophin 1)	13 A3.3	AK082929	100	Similar to placental lactogen 1	98.8
13 A4	NM_011456	100	Spi14 (Serine Protease Inhibitor 14)	13 A4	NM_011455	100	Spi13 (serine protease inhibitor 13)	95.2
13 D1	NM_010872	100	Birc1b (Neuronal apoptosis inhibitory 2)	13 D1	NM_008670	99.9	Birc1a (Neuronal apoptosis inhibitory 1)	90.0

Table 3 (Continued)**Examples of recent mouse gene duplications**

14 C1	NM_010373	99.7	Gzme (Granzyme E)	14 C1	NM_010372	99.9	Gzmd (Granzyme D)	94.7
14 C2	NM_172603	100	4933417L10Rik	14 C3	BE381578	100	EST sequence	94.0
15 E2	NM_007781	100	Csf2rb2 (Colony stimulating factor 2, β -2)	15 E2	NM_007780	99.8	Csf2rb1 (Colony stimulating factor 2, β -1)	95.0
15 E2	NM_010005	100	Cyp2d10 (Cytochrome P450, 2d10)	15 E2	NM_010006	100	Cyp2d9 (cytochrome P450, 2d9)	92.1
16 B1	NM_023125	100	Knk (Kininogen)	16 B1	BI330914	99.1	EST sequence	90.0
16 B3	M92418	99.8	MS2 (Cysteine proteinase inhibitor)	16 B3	BB654253	100	EST sequence	95.2
17 B2	NM_009780	99.9	C4 (Complement component 4)	17 B2	M21576	99.5	Slp (MHC sex-limited protein)	96.3
X A2	NM_008955	100	Pxl (Placenta specific homeobox 1)	X A2	NM_023894	100	Homeobox protein GPBOX	91.6

*Locations of duplicons by mouse chromosome banding; locus 1 and 2 represent a duplication pair. †Alignment percent identity between gene and genomic sequences showing correct matches. ‡% similarity: average DNA percent identity between paralogous gene/transcript sequences in locus 1 and 2 (duplicated pair)

Table 4**Protein domain enrichment found in recently duplicated mouse genes***

InterPro entry ID	Protein domain description	Number found in 608 duplicated genes	Number found in all 16,515 annotated genes in genome	Enrichment [†]
IPR000276	Rhodopsin-like GPCR superfamily	135	1229	3.0
IPR000725	Olfactory receptor	103	861	3.3
IPR003006	Immunoglobulin/major histocompatibility complex	46	372	3.4
IPR004072	Vomer nasal receptor, type 1	31	108	7.8
IPR001909	KRAB box	23	103	6.1
IPR001254	Serine protease, trypsin family	21	117	4.9
IPR002401	E-class P450, group I	20	61	8.9
IPR001128	Cytochrome P450	20	68	8.0
IPR007086	Zn-finger, C2H2 subtype	20	139	3.9
IPR001314	Chymotrypsin serine protease, family S1	19	108	4.8
IPR002403	E-class P450, group IV	17	56	8.2
IPR002397	B-class P450	13	29	11.9
IPR001304	C-type lectin	13	96	3.7
IPR000215	Serpin	12	48	6.8
IPR002402	E-class P450, group II	9	14	18.5
IPR006046	Glycoside hydrolase family 13	7	8	23.0
IPR006047	Alpha amylase, catalytic domain	7	10	19.2
IPR001400	Somatotropin hormone	7	32	6.1
IPR006048	Alpha amylase, C-terminal all-beta domain	6	7	24.7
IPR002018	Carboxylesterase, type B	6	13	12.3
IPR004073	Vomer nasal receptor, type 2	6	13	12.3
IPR001039	Major histocompatibility complex protein, class I	6	17	9.9
IPR001828	Extracellular ligand-binding receptor	6	29	5.5
IPR002213	UDP-glucuronosyl/UDP-glucosyl transferase	5	12	11.8
IPR002448	Odour-binding protein	4	9	13.2
IPR000068	Extracellular calcium-sensing receptor	4	10	11.0

*Only Ensembl gene annotation (608 genes) was used in this analysis. †All results shown are statistically significant with p -values $< 10^{-5}$ (χ^2 test).

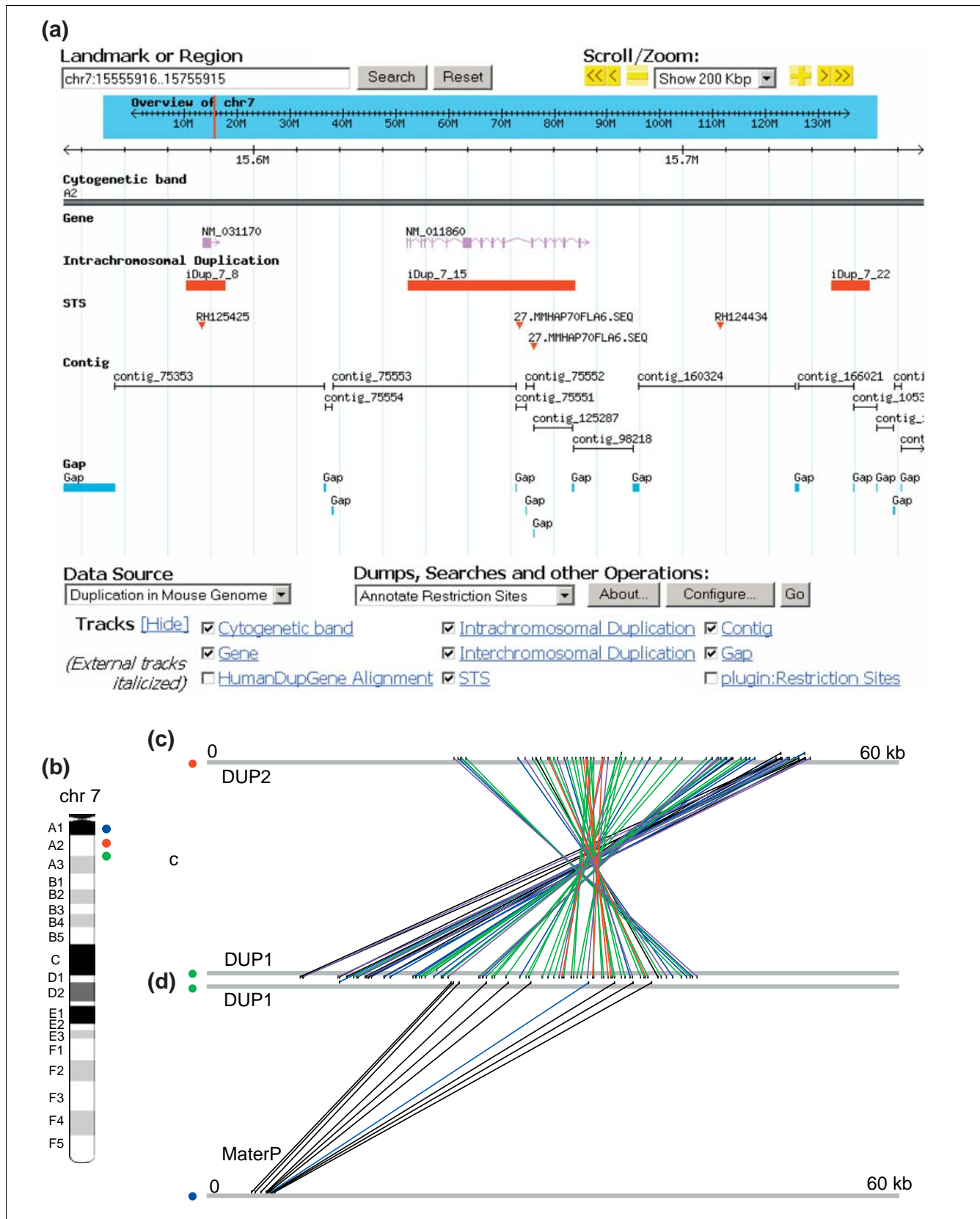


Figure 2 (see legend on next page)

Figure 2 (see previous page)

The genomic organization of the *Mater* duplication. **(a)** Location of the *Mater* duplication. A snapshot view of GMOD browser (details can be found at [14]). **(b)** Chromosomal view (mouse chromosome 7) of the three *Mater* duplication locations (DUP1, DUP2, *MaterP*). **(c)** Graphical view of the sequence similarity between DUP1 and DUP2 shown by GenomePixelizer. DUP2 is situated in an inverse orientation with respect to DUP1. Red, 99-100% sequence identity; purple, 96-98%; green, 93-95%; blue, 90-92%; black, 85-89%. **(d)** Graphical view of the sequence similarity between DUP1 and the *MaterP* region. As shown, *MaterP* is an intron-less, retrotransposed pseudogene. Blue, 90-92% sequence identity; black, 85-89%.

Materials and methods**Genome sequence and chromosome-wide BLAST**

We obtained the February 2002 (MGSCv3) and February 2003 mouse genome assemblies (lower-case repeat-masked sequences), as well as the assembly component tables, through the UCSC Human Genome Browser website [21]. For each assembly, detection of intrachromosomal segmental duplications involved comparing each of the 20 masked chromosome sequences (excluding the Y chromosome not targeted by the MGSC) and the masked unmapped chromosome sequence against itself by BLAST2 [16] (21 comparisons made). Interchromosomal analysis of segmental duplications involved pairwise comparisons between each of the 21 chromosomes (420 comparisons made). Analyses were repeated with the exclusion of the unmapped chromosome sequence to examine its contribution to the overall duplication content (results posted at [14]). All BLAST results were subsequently parsed to eliminate low-quality and fragmented alignments under the following criteria: BLAST results having $\geq 90\%$ sequence identity, \geq a length of 80 bp, and with expected value $\leq 10^{-30}$.

Parsing of BLAST results and duplication detection

Each BLAST report was sorted by chromosomal coordinates. All identical hits (same coordinate alignments), including suboptimal BLAST alignments recognized by multiple, overlapping alignments, as well as mirror hits (reverse coordinate alignments) from the BLAST results of the intrachromosomal set, were removed. Contiguous alignments separated by a distance of less than 3 kb, then 5 kb, and subsequently 9 kb, were joined stepwise into modules in order to traverse masked repetitive sequences and to overcome breaks in the BLAST alignments caused by insertions/deletions and sequence gaps. Such contiguous sequence-alignment modules represent sequence similarity between the subject and query chromosome sequence in question (at their respective positional coordinates) [9]. Potential sequence misassignment errors are results detected to have $> 99.5\%$ sequence identity with another region.

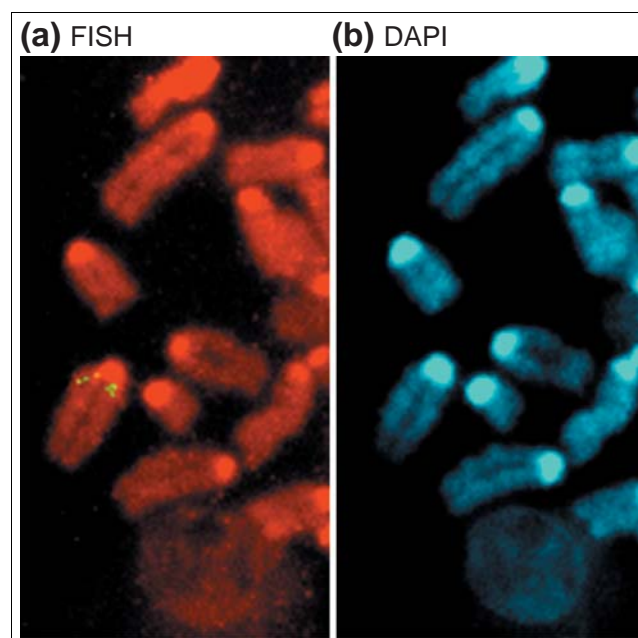
Online database for recent segmental duplications

We overlaid all duplication content and regions containing potential sequence misassignment errors onto the mouse genome sequence, which can be viewed using the interactive Generic Genome Browser [33] hosted at our website [14]. Results and analyses were presented for both the February

2002 and February 2003, each as a separate database. Results are also summarized in tables that include information on chromosomal coordinates, band locations, size of duplications, level of identity between duplicated copies, as well as genes mapped to these regions [14]. Graphical representation for intrachromosomal duplications was generated using the visualization tool GenomePixelizer (Figure 1) [34].

Identification of recent gene duplications

We obtained the NCBI Refseq gene annotation file (refGene.txt.gz) from the UCSC Downloads website [21] and the Ensembl gene annotation (Mus_musculus.cdna.fa.gz; Ensembl Known genes only) from the Ensembl website [35]. Genes that mapped to duplicated regions of the mouse genome were identified using their chromosome sequence coordinates. In total, 439 Refseq and 608 Ensembl annotated genes were found to be involved in duplicated regions;

**Figure 3**

FISH detection of *Mater* duplication. **(a)** Metaphase FISH showing three pairs of signals (yellow) detected on mouse chromosome 7 using BAC clone RP23-225F5 (detection frequency of 70%) mapping to duplicated *Mater* regions. **(b)** DAPI banding of the same partial mitotic figures for the identification of mouse chromosome 7. A control probe RP23-464L20 was mapped to a single location in the F2 region (data not shown).

Table 5**Genes that have undergone recent duplication in both the mouse and human genome***

Refseq†	Gene description
NM_007534	<i>B-cell leukemia/lymphoma 2 related protein A1b (Bcl2a1b)</i>
NM_007558	Bone morphogenetic protein 8a (Bmp8a)
NM_007812	Cytochrome P450, 2a5 (Cyp2a5)
NM_008181	<i>Glutathione S-transferase, alpha 1 (Ya) (Gsta1)</i>
NM_010390	<i>Histocompatibility 2, Q region locus 1 (H2Q1)</i>
NM_009467	UDP-glucuronosyltransferase 2 family, member 5 (Ugt2b5)
NM_009669	<i>Amylase 2, pancreatic (Amy2)</i>
NM_009888	Complement component factor h (Cfh)
NM_010856	Myosin heavy chain, cardiac muscle, adult (Myhca)
NM_013778	Aldo-keto reductase family I, member C13 (Akr1c13)
NM_022033	<i>3-Oxoacid CoA transferase 2 (Oxct2) (Imbedded in the Bmp8a gene)</i>
NM_026419	<i>Elastase 3B, pancreatic (Ela3b)</i>
NM_031176	Tenascin XB (Tnxb)
NM_130864	Acetyl-coenzyme A acyltransferase (peroxisomal 3-oxoacyl-Coenzyme athiolase) (Acaa)
NM_010997	<i>Olfactory receptor 54 (Olf54)</i>
NM_031170	Keratin complex 2, basic, gene 8 (Krt2-8)

*Six hundred and seventy-five duplicated mouse gene sequences were aligned to the June 2002 human genome assembly by BLAST (with an initial expected value cutoff of $<10^{-10}$). The best aligned human genes were subsequently used for reciprocal BLAST alignments (against the mouse genome sequence) to establish a putative orthologous relationship between the mouse and human gene pairs. Using results from our human genome duplication analysis [9,37], we examined regions of the human genome where the human genes are involved in recent segmental duplication. †Italics represents genes that are entirely within a duplication in the mouse genome.

together these two datasets made up 675 unique gene annotations (372 overlapped annotations). To establish gene-pair relationships between duplicated gene sequences, each of the 238 NCBI RefSeq genes that were found to be fully contained within a segmental duplication was searched against the UCSC and NCBI GenBank database for spliced ESTs, full-length cDNAs and additional annotated genes. A total of 128 gene pairs were established, most of which are likely to be novel gene paralogs previously unknown in the literature. In the analysis of protein-domain enrichment in duplicated genes, InterPro annotation for duplicated genes (608 Ensembl genes) as well as for the entire gene set (16,515) was obtained from Ensembl EnsMart [36]. We counted the number of times a protein-domain class is found in each gene set and tabulated our results (see Table 4). To examine the subset of genes that had undergone recent duplication in the human genome, each of the duplicated gene sequences was aligned to the June 2002 human genome assembly by BLAST (with an initial expected value cutoff of $<10^{-10}$). The best-aligned human genes were subsequently used for reciprocal

BLAST alignments (against the mouse genome sequence) to establish a putative orthologous relationship between the mouse and human gene pairs. Using results from our human genome duplication analysis [37], we examined regions of the human genome where the human genes were involved in recent segmental duplication.

Fluorescence in situ hybridization (FISH)

Mouse lymphocytes were isolated from the spleen and cultured at 37°C in RPMI 1640 medium supplemented with fetal calf serum, concanavalin A and lipopolysaccharide. After 44 hours, the cultured lymphocytes were treated with bromodeoxyuridine for an additional 14 hours. The synchronized cells were washed and recultured at 37°C for 4 hours in a minimal Eagle's medium with thymidine. Chromosome slides were made by conventional methods including hypotonic treatment, fixation and air-drying [38]. BAC probes RP23-225F5 (mapped to the *Mater* locus (DUP1) by BAC-end sequences) and RP23-464L20 (a control probe) were biotinylated respectively. Hybridization and detection were carried out according to [39]. FISH signals were observed under fluorescent microscopy using FITC and DAPI filters. Images were captured by CCD camera.

Additional data files

Further analysis of *Mater* duplication, including a figure showing a multiple percent identity plot of *Mater* versus *Mater2*, *MATER* (human), and *MaterP*, is available with the online version of this article (additional data file 1).

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