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Recent Segmental Duplications in the Working Draft Assembly of the Brown Norway Rat

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We assessed the content, structure, and distribution of segmental duplications ($\geq 90\%$ sequence identity, ≥ 5 kb length) within the published version of the *Rattus norvegicus* genome assembly (v.3.1). The overall fraction of duplicated sequence within the rat assembly (2.92%) is greater than that of the mouse (1%–1.2%) but significantly less than that of human (~5%). Duplications were nonuniformly distributed, occurring predominantly as tandem and tightly clustered intrachromosomal duplications. Regions containing extensive interchromosomal duplications were observed, particularly within subtelomeric and pericentromeric regions. We identified 41 discrete genomic regions greater than 1 Mb in size, termed “duplication blocks.” These appear to have been the target of extensive duplication over millions of years of evolution. Gene content within duplicated regions (~1%) was lower than expected based on the genome representation. Interestingly, sequence contigs lacking chromosome assignment (“the unplaced chromosome”) showed a marked enrichment for segmental duplication (45% of 75.2 Mb), indicating that segmental duplications have been problematic for sequence and assembly of the rat genome. Further targeted efforts are required to resolve the organization and complexity of these regions.

Segmental duplications have long been recognized as important mediators of both gene and genome evolution (Muller 1936; Ohno 1970). From the genic perspective, such duplications often encode protein products which, although not essential for viability of the organism, are important for the adaptation of the species to specific ecological niches (Duda and Palumbi 1999). Among mammalian species, commonly duplicated genes include those associated with the recognition of environmental molecules and include genes associated with innate immunity, drug detoxification, olfaction, and sperm competition. From the perspective of genome structure, lineage-specific segmental duplications or large repeats often delineate regions of recurrent evolutionary lability (Eichler and Sankoff 2003). Recent comparative sequencing efforts among closely related eukaryotes, for example, shows that highly homologous repetitive sequence frequently associate with the breakpoints of large-scale chromosomal rearrangement (Dehal et al. 2001; Kellis et al. 2003). Understanding the nature and pattern of segmental duplications provides fundamental insight into functional redundancy, adaptive evolution, and the structural dynamics of chromosomal evolution.

One of the surprising findings from the analysis of the Human Genome Project data was the relevant abundance of large blocks of sequence with a high degree of sequence identity (Bailey et al. 2001; International Human Genome Sequencing Consortium [IHGSC] 2001). A variety of computational and experimental methods (Bailey et al. 2001, 2002; Cheung et al. 2001) now estimate 5%–6% of the human as duplicated (≥ 1 kb and $\geq 90\%$). Compared to other sequenced organisms such as fly and worm, the human genome is enriched for recent segmental duplications, particularly interspersed duplications (Bailey et al. 2002). Such comparisons, however, typically assess duplication content with lower-bound estimates of length. For example, these cross-species comparisons rarely characterize duplications less than 500 bp in length. This may introduce an ascertainment

bias, particularly among invertebrates, whose genomes can be orders of magnitude smaller compared to human. Larger genomes may simply harbor larger segmental duplications. The purported “unique” properties of the human genome can only be assessed by detailed comparison with other mammalian genomes where genome sizes are equivalent. With the whole-genome shotgun sequence assembly of the rat genome, we can now assess the nature and pattern of segmental duplication of a third mammalian genome (Rat Genome Sequencing Project Consortium [RGSPC] 2004; Waterston et al. 2002).

We present a preliminary, genome-wide analysis of the segmental duplication content of the rat (*Rattus norvegicus*). Any assessment of segmental duplication content is highly dependent on the methodology and quality of the sequence assembly (Bailey et al. 2001; Eichler 2001; Cheung et al. 2003a). Discrimination between highly paralogous copies and allelic regions that have not been properly assembled requires an estimate of the levels of both allelic variation and sequencing error. For most regions, paralogous sequences are more divergent than allelic copies. Another consideration is the method of genome assembly. Assembly algorithms based on sequence overlap from working-draft BAC clones were shown to overestimate the frequency of segmental duplication, due to a failure to properly merge allelic overlaps (Bailey et al. 2001; IHGSC 2001). Alternatively, assembly strictly from whole-genome shotgun sequence reads tends to over-collapse and therefore underrepresent such regions due to the recruitment of both paralogous and allelic sequence reads (Eichler 1998; Bailey et al. 2002; Estivill et al. 2002). Interestingly, the assembly algorithm of the rat genome represents a hybrid of whole-genome- and clone-ordered-based approaches. The ability of this approach to resolve segmental duplications has not been tested previously. In light of these inherent difficulties associated with the assembly of highly similar duplications, the analysis should be considered a first approximation of the recent duplication properties of the rat genome. Such initial analyses, however, are essential in providing a more accurate and robust “final” version of the rat genome as well as insight into genome-assembly approaches. The results of the present study have been made publicly available through the UCSC genome browser as

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Table 1. Length vs. Number of Pairwise Alignments (Rat v. 3.1)

Seed length (bp)	# Pairwise
>250	1,283,258
>1000	532,720
>5000	45,835
>10,000	4798
>20,000	171

Seed length determined after masking of common rat repeats (Repeatmasker June 2003 version).

well as through our own local database (<http://ratparalogy.cwru.edu>), providing a resource for the rat sequencing and genetics community.

RESULTS

We initially examined the entire draft genome of the rat using a previously described BLAST-based whole-genome sequence comparison method (see Methods; Bailey et al. 2001). Assembled draft sequence of any genome may be operationally divided into two categories: sequence which can be mapped to a chromosome, and that which cannot. In the case of the rat genome, 75 Mb of assembled sequence was ambiguous in its placement. Because segmental duplications are particularly enriched in this category, we separately considered this category throughout our

analysis. In order to detect segmental duplications specific to the rat lineage, we examined all duplications that showed <10% sequence divergence. Based on sequence divergence between mouse and rat (0.175–0.195; RGSPC 2004), such regions likely represent either lineage-specific duplications or large-scale gene conversion events. During the initial phases of this analysis, we discovered an overabundance of pairwise alignments <5 kb in size (Table 1). Their high-copy number, relatively small size, well defined borders, and their highly interspersed nature both within and between chromosomes suggested contamination by high-copy repeats, despite the removal of rodent retroelements using the latest curated version of Repeatmasker (June 2003). Such contamination could be due to either incomplete masking of unknown repeat elements or transduction of flanking sequence (Goodier et al. 2000; Pickeral et al. 2000). As our goal was to identify genomic sequence that arose as a consequence of duplication (not retrotransposition or retroposon-induced transduction), we raised our threshold for seeding alignments to 5 kb—the effective insertion length of most retroelements is <5 kb in length, whereas most transduced sequences are less than 1 kb in length. For comparisons we considered additional alignment length thresholds (5, 10, 20 and 50 kb; Fig. 1) which were certain to exclude all transposable elements, including full-length retroviral repeats (Table 2).

Sequence Properties of Rat Segmental Duplications

We calculate a total of 2.92% (82.8 Mb/2835 Mb) of the rat genome as duplicated ($\geq 90\%$ sequence identity, ≥ 5 kb; Figs. 1, 2).

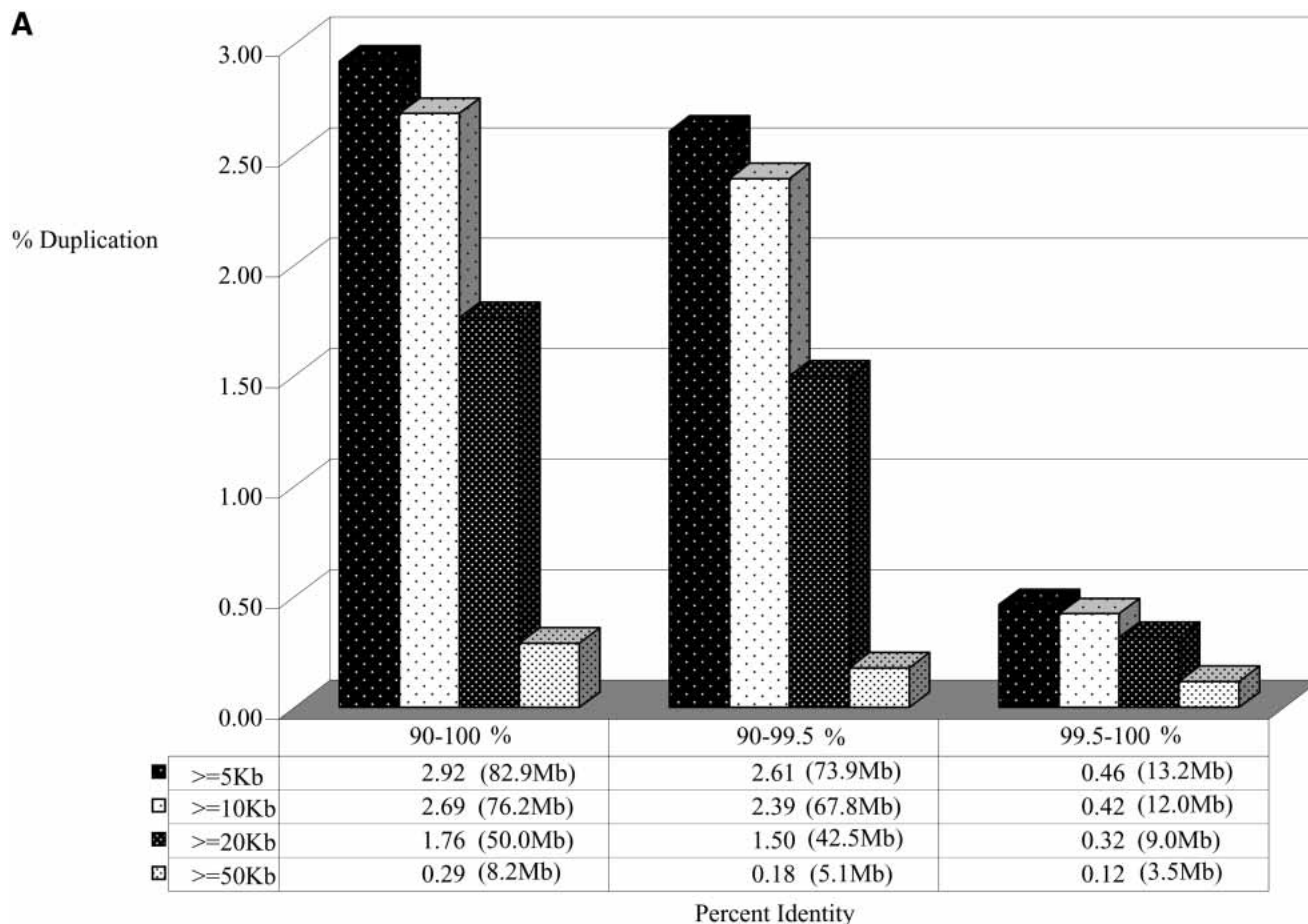


Figure 1 (Continued on next page)

These correspond to 43,597 pairwise alignments and represent 3237 distinct regions of the rat genome (Table 2). Pairwise alignments may be redundant in nature, as the same sequence may be duplicated to multiple locations in the genome. Therefore, the number of distinct, nonoverlapping regions is substantially fewer. Figure 1 depicts the duplication content of the rat genome as a function of the length of alignment and the degree of sequence identity. As described above, we included and excluded the unplaced sequence contigs to show the disproportionate representation of duplicated sequence in this category. Based on our analysis of the entire genome, the median length of alignments (9749 bp) is not significantly different between interchromosomal and intrachromosomal duplications. The largest alignment detected is 104 kb. The average degree of sequence identity among all alignments is 94.4%. Interestingly, when we considered the percent nucleotide sequence identity for segmental duplications as a function of the number of aligned base pairs, we observed a distinct bimodal distribution (Fig. 2B). Two peaks were observed corresponding to 95.5% and 92.5% sequence identity (0.045 substitutions per site and 0.075 substitutions per site). This bimodal distribution was consistently observed whether unmapped genomic sequence was excluded or included in the analysis.

Estimates of segmental duplication from the human genome working draft sequence assembly initially overestimated the fraction of duplicated bases (10.8% of the genome). This was

the result of a failure to merge allelic overlaps during the genome assembly process. Subsequent analysis of finished genome sequence showed that such alignments showed an extraordinary degree of sequence identity consistent with missed allelic overlaps (Bailey et al. 2001). To eliminate such potential artifacts in the rat genome assembly, we separately considered all alignments where the degree of sequence identity is less than 99.5%. We derive a conservative estimate of the duplication content of the rat genome to be 2.61% (73.9 Mb/ 2835.2 Mb, 2928 distinct genomic regions; Fig. 1). It is unlikely, therefore, that the majority of rat segmental duplications identified in this study arise as a consequence of a failure to merge overlaps during assembly.

Rat segmental duplications show a bias toward intrachromosomal alignments (68.1 Mb) compared to interchromosomal duplications (48.2 Mb; Figs. 2, 3, Table 2). Interestingly, the number of intrachromosomal and interchromosomal pairwise alignments differs more dramatically. By this measure, intrachromosomal duplications are three times more frequent than interchromosomal duplications (32,527 intrachromosomal alignments vs. 11,070 interchromosomal alignments; Table 2). It should be noted, however, that a significant fraction of the rat genome sequence (115.2 Mb) has not been assigned to a chromosome (unplaced chromosome), nor has it been assigned specifically within a chromosomal region (random chromosome bins). The above calculations treat the unmapped sequence as a separate chromosome when classifying duplications as inter- or intrachro-

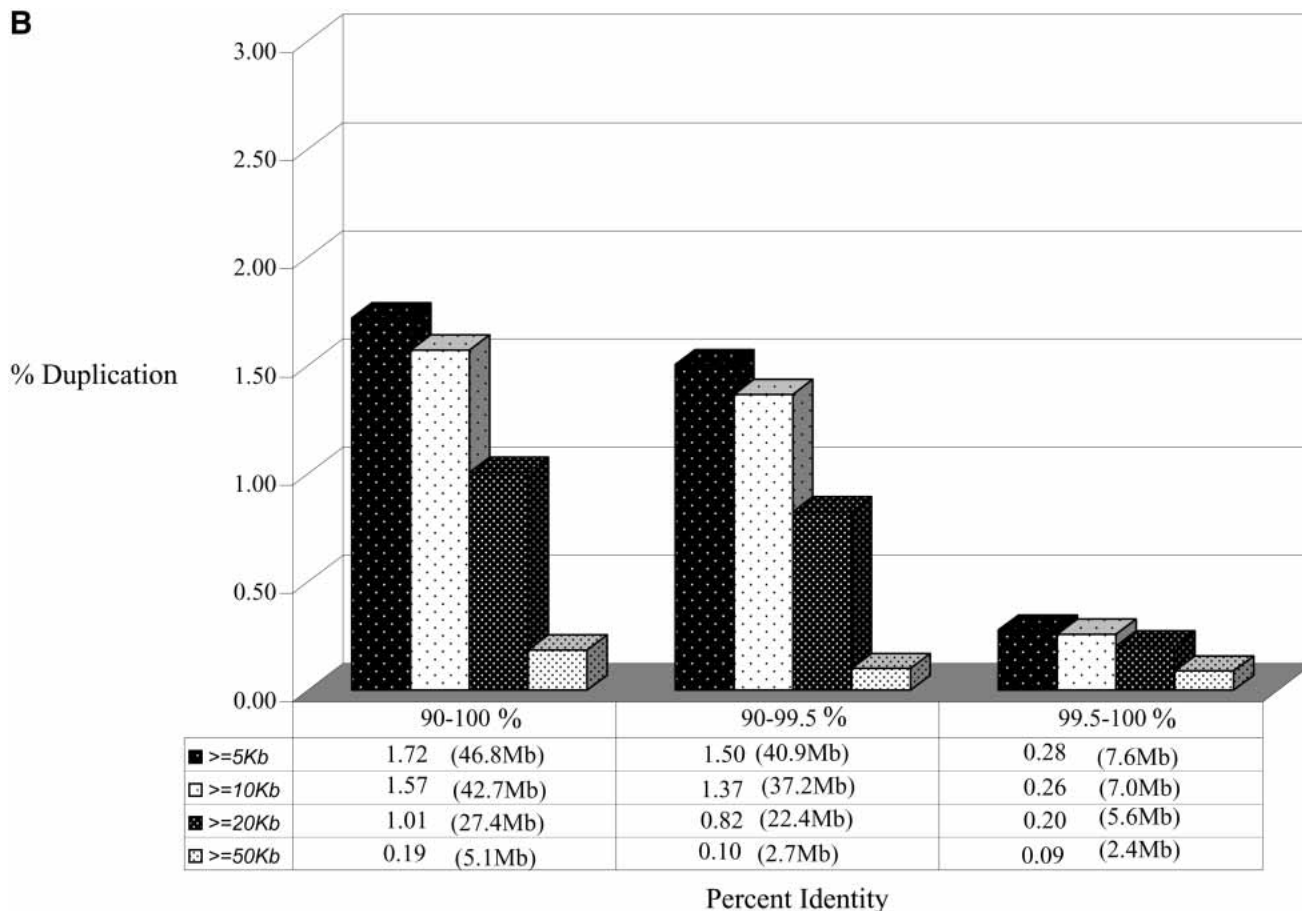


Figure 1 Duplicated fraction in the rat genome. The figure depicts the proportion of the genome that shows duplication (A) when all genomic sequence was compared, and (B) for the rat genome excluding random, unassigned sequence contigs. Various lengths and % identity thresholds are shown. A very small portion of the rat genome shows segmental duplications with $\geq 99.5\%$ sequence identity. This suggests that the majority of segmental duplications are bona fide and are not the result of missed allelic overlaps during genome assembly.

Table 2. Rat Segmental Duplication Sequence Alignment Statistics

Size (kb)	Number of alignments			Average sequence identity			Average length		
	Inter	Intra	All	Inter	Intra	All	Inter	Intra	All
5	568	1265	1833	0.940	0.952	0.948	5631	5630	5631
6	1255	3769	5024	0.941	0.947	0.945	6528	6529	6529
7	1532	3968	5500	0.936	0.944	0.942	7496	7495	7495
8	1329	4501	5830	0.934	0.942	0.940	8506	8472	8480
9	1145	3487	4632	0.938	0.943	0.942	9533	9463	9481
10–19	4367	13550	17917	0.944	0.946	0.945	13376	13280	13303
20–29	673	1571	2244	0.950	0.953	0.952	23894	23601	23689
30–39	134	277	411	0.956	0.960	0.959	33183	33858	33638
40–49	33	70	103	0.965	0.964	0.964	44493	44211	44301
50+	34	69	103	0.979	0.967	0.971	59546	56850	57740
Total	11070	32527	43597	0.941	0.946	0.944	11520	11253	11321

Alignments were binned into groups based on 1-kb increments (i.e., 5, 6 kb, etc) and 10 kb increments (i.e., 10–19.9 kb), the absolute number of alignments, average sequence identity, and average length for interchromosomal, intrachromosomal, and all alignments are shown after seed alignments were joined (see Methods). Consequently, the total number of joined alignments is less than the number of seed alignments (Table 1).

mosomal. We estimate that 45% (36.1 Mb/82.8 Mb) of the duplications are mapped to these intractable regions of the rat genome. Their map locations are ambiguous, and intra/interchromosomal distribution is technically unknown. If we exclude these two categories of sequence, a total of 1911 (46.7 Mb, 1.72% of the genome) regions of duplication are identified which have been unambiguously mapped within the rat genome. Again, a stronger preference for intrachromosomal duplications (38.8 Mb) was observed compared to interchromosomal duplications (17.7 Mb). With few exceptions, most intrachromosomal duplications are organized as clusters of tandem or inverted duplications within close proximity. Using these conservative criteria, ~21% of the duplicated bases (8.8 Mb) were part of interchromosomal and intrachromosomal duplication alignments.

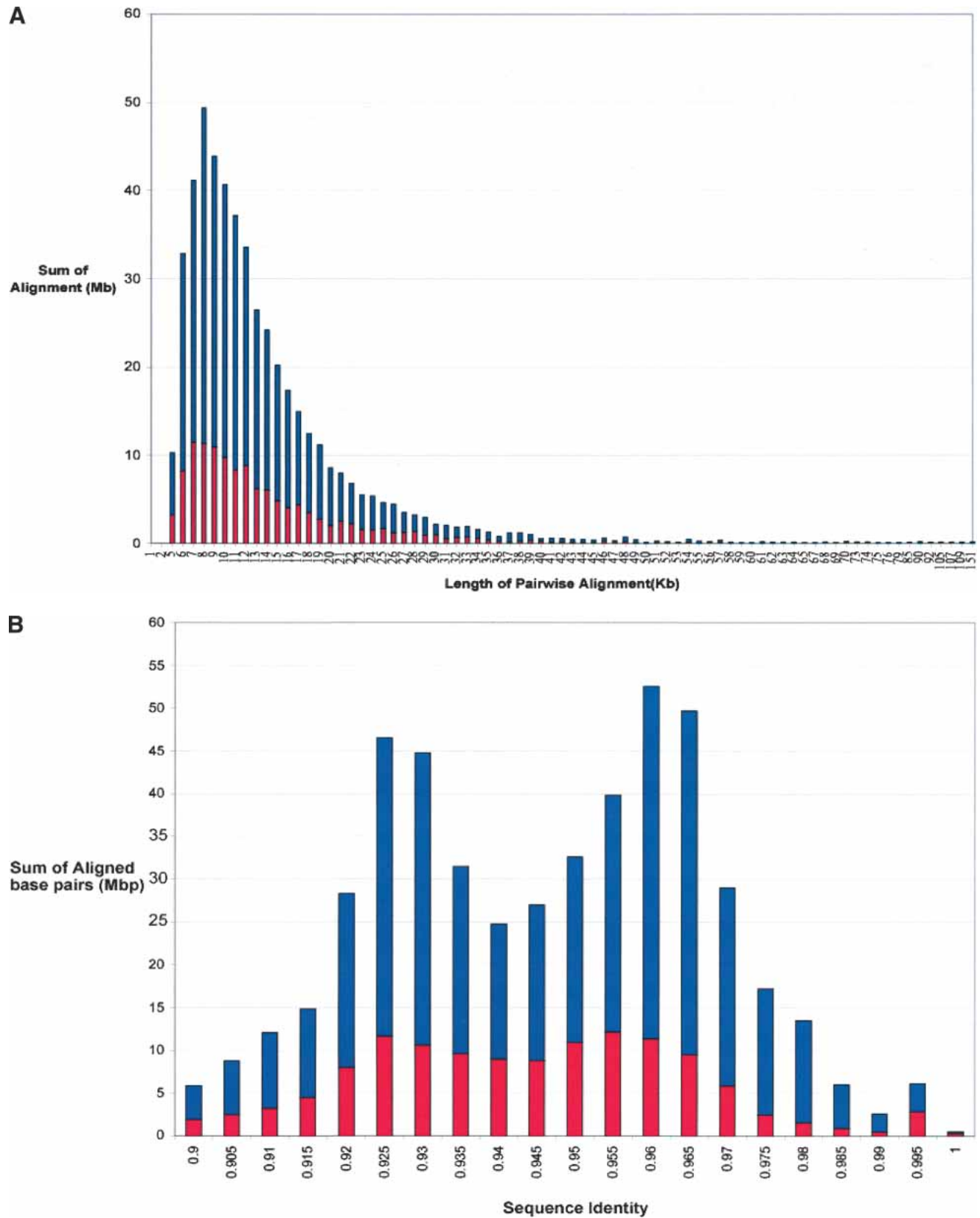
As a final analysis of the sequence properties of rat segmental duplications, we compared the repeat content of duplicated sequence, flanking sequence and the whole genome (Table 3, Methods). Unlike human segmental duplications, which are enriched for SINE repeats (Bailey et al. 2003), no SINE enrichment (nor any other retroelement) was associated with rat segmental duplications. The working draft nature of the rat genome sequence prevents a detailed analysis of the sequence structure at the transition regions between unique and duplicated sequence. Nevertheless, two clear patterns emerge regarding repeat content. Although the common repeat content of most duplications appears to be reduced, SINE content shows the greatest reduction compared to the genome average (1.97% vs. 7.1%). This gradually increases to the genome average as sequences flanking the duplications are considered (Table 3). An opposite trend is observed with respect to centromeric satellite repeat sequences. Rat segmental duplications show a fourfold enrichment for satellite repeat content compared to the genome average. When individual repeat subfamilies are considered, satellite repeat classes 91ES8_RN and RNSAT1 show the greatest enrichment (10-fold and sevenfold, respectively). This association is most pronounced among blocks of interchromosomal duplication (see below).

Organization of Rat Segmental Duplications

The recent segmental duplications of the rat genome are distributed in a nonrandom fashion at two different levels. First, duplication content varies significantly among different chromosomes. Chromosomes 12, 7, 15, and 1 show the greatest enrichment for segmental duplication (Fig. 4A) with twofold the duplication content of the genome average (excluding unplaced

sequence contigs). Most of this effect is due to an increase in intrachromosomal duplication content localized as specific clusters. During the analysis of segmental duplications, large tracts were identified which were populated by a high density of segmental duplications. These tracts, termed “duplication blocks” (Bailey et al. 2001) ranged from 500 kb to as large as 3 Mb in size (Table 4), were generally gene-poor, and were characterized by assembly inconsistencies. A total of 41 discrete duplication blocks were identified which exceeded 1 Mb in length (Table 4). Typical block structures for chromosomes 1 and 7 are depicted (Fig. 4B). Analysis of the pairwise alignments underlying these block structures showed considerable variation in sequence identity (90%–99% identity), often within the same block. Two types of duplication block structures were distinguished: chromosome-specific blocks which consisted largely of interspersed segmental duplications (Table 5), and clustered interchromosomal pairwise alignments with considerable range in sequence identity. Interestingly, within a specific duplication block, multiple pairwise alignments among specific subsets of chromosomes could be identified (Table 5).

In humans (Eichler et al. 1996; Jackson et al. 1999; Horvath et al. 2000) and to a lesser known extent in mouse (Thomas et al. 2003), segmental duplications show particular biases for pericentromeric and subtelomeric regions of the genome. Based on the current rat genome assembly, regions of segmental duplications (100 kb–1.5 Mb in size) were observed for 13 of the 40 possible most distal sequence contigs, suggesting a subtelomeric preponderance. Most of these subtelomeric blocks showed complex patterns of interchromosomal duplication among specific subsets of rat chromosomes. Characterization of a pericentromeric bias for segmental duplication is more difficult to determine, because the location of rat centromeres are generally not as well identified as mouse and human. We attempted to approximate the centromere position within the rat genome assembly using two independent methods (RGSPC 2004). The first approach mapped the most proximal STS/gene marker to the p and q arm of each rat chromosome by FISH, and considered the interval between these markers within the assembly as a possible centromere location. The second approach identified dense clusters of classic rat satellite repeats (particularly SAT1_RN and ISAT_RN) within the assembly. Six rat chromosomes showed a correlation by these two different methods and allowed a likely assignment of the centromere region. Of these four chromosomes, large blocks of interchromosomal segmental duplication were identified ranging in size from 300 kb–3 Mb. Once again, analysis of underlying pair-



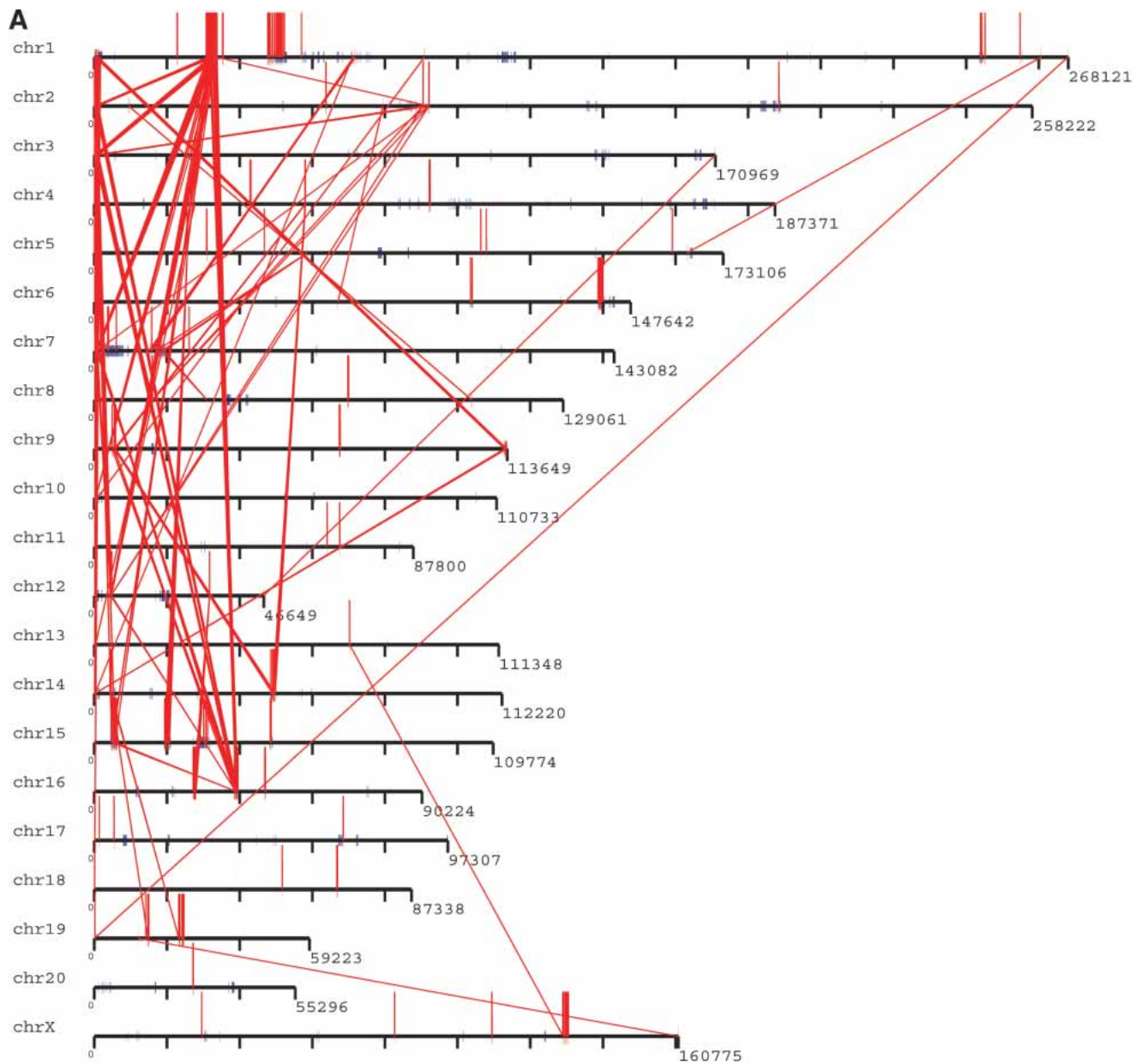


Figure 3 (Continued on next page)

wise alignments identified sequence homology among specific sets of rat chromosomes (Table 5). A dearth of RefSeq genes or spliced rat mRNA within these regions was noted.

Gene Analysis of Recent Rat Segmental Duplications

We considered the genomic duplication content of all RefSeq mRNA aligned to the rat genome. To eliminate potential false positives, we limited our analysis to duplications showing $\geq 1\%$ sequence divergence, well below the polymorphic level of variation for this inbred strain. The duplications therefore likely represent bona fide recent gene duplication or gene conversion events within the rat lineage. A total of 45/4250 rat RefSeq genes were identified that were embedded within the segmental duplications detected by whole-genome analysis comparison (Table

6). Even though interchromosomal duplications constitute \sim one-third of all pairwise alignments and 40% of all duplicated bases, genes are largely biased to intrachromosomal duplications (41/45 or 91%). Of these, almost all pairwise were < 1 Mb apart, indicating that most “functional” duplicates within the rat genome are tandem gene clusters, as opposed to widely interspersed duplications. Indeed, in our analysis of the RefSeq genes alone, 19/45 genes belonged to known clusters of tandem gene families. Due to the limited number of characterized RefSeq genes, we broadened our analysis to consider known rat mRNA which possessed two or more exons. Although a few putative novel gene families were identified (e.g., α -latrotoxin G-coupled protein receptor, low-voltage activated calcium channel gene family, and a dynein-like protein subfamily; Supplemental Table 1), most mRNA corresponded to additional members of previously characterized genes (RGSPC 2004).

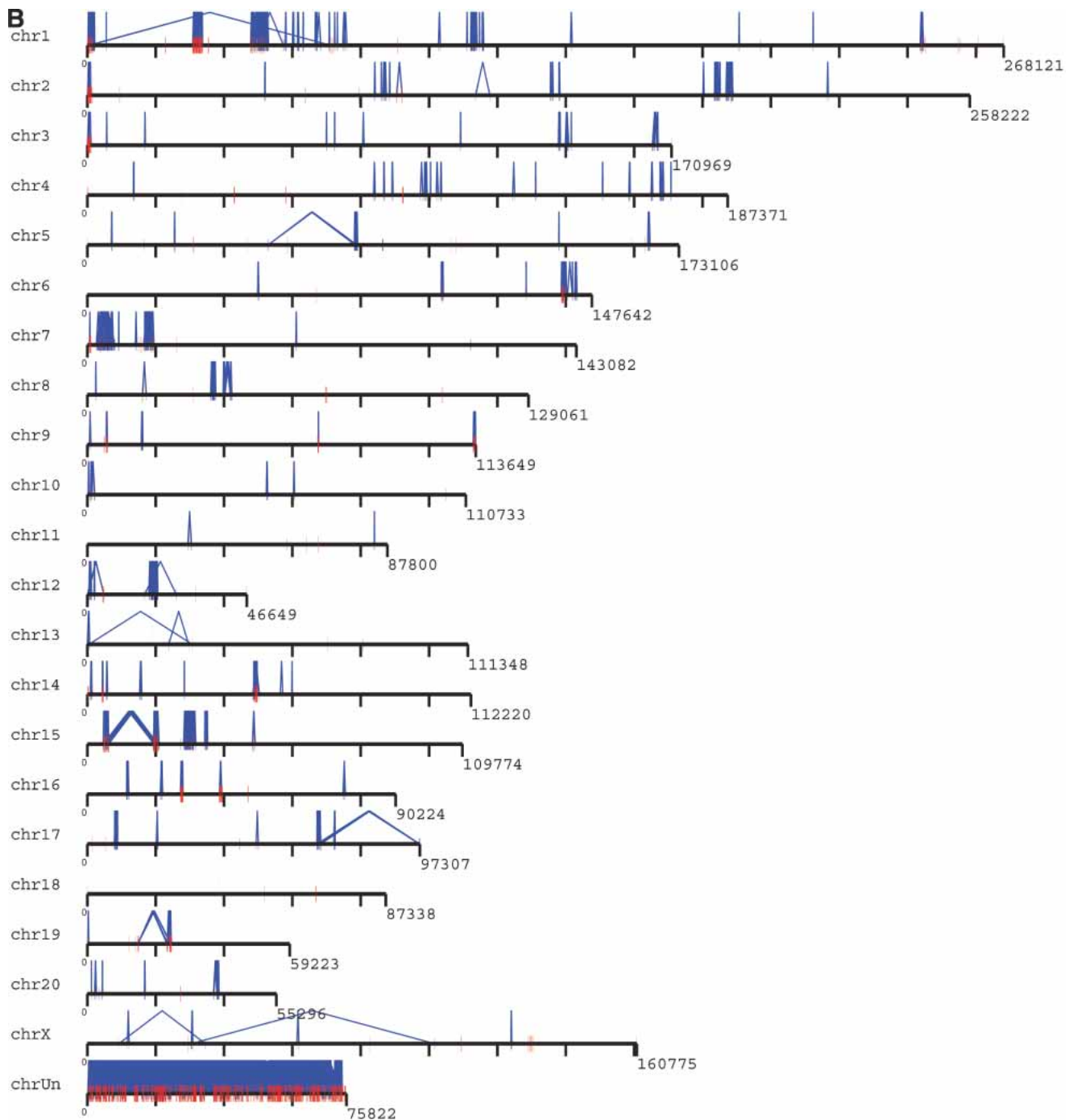


Figure 3 Distribution of segmental duplications ($\geq 90\%$ and ≥ 10 kb) in the rat genome. The pattern of (A) interchromosomal duplications (red) and (B) intrachromosomal duplications (blue) are depicted for all duplications $\geq 90\%$ sequence identity and ≥ 10 kb in length. For clarity, interchromosomal distribution patterns with the random, unassigned sequence contigs (chrUn) are not shown for (A). For more detail, including % identity and pairwise relationships of all duplications and alignments, see <http://ratparalogy.cwru.edu>.

The genes identified in our analysis fall into three categories. These include genes associated with foreign compound detoxification (cytochrome P450 and carboxylesterase genes), environmental signal recognition (α -2 globulin and pheromone receptors), and innate immune response (rat serine protease inhibitors, natural killer cell receptors, T-cell receptor, major histocompatibility locus, and immunoglobulin variable heavy chain locus, etc.; RGSPC 2004). Despite the abundance of

rat segmental duplications on the “unknown” chromosome, only eight duplicate genes with two or more exons are identified within this 45 Mb of duplicated sequence. This included caveolin-2 (AF439788), a vacuolar protein sorting homolog (U35244), two copies of a carboxylesterase E gene (D00362), and various gene fragments/orphans of immunoglobulin γ and ϵ variable chain, T-cell receptor, and cytochrome P450 genes. It is likely that these sequences represent displaced members of tandem

Table 3. Repeat Properties of Rat Genome, Duplications, and Flanking Regions

Repeat	Duplications		Duplicated blocks		20-kb flanks		Genome		Enrichment in duplication content
		%		%		%		%	
DNA	237468	0.29	321957	0.24	77076	0.33	20671634	0.81	0.358
LINE	13919243	16.79	25958082	19.01	5722640	24.23	579171737	22.57	0.744
SINE	1958333	2.36	2691829	1.97	724501	3.07	181352249	7.07	0.334
LTR	5993558	7.23	9605961	7.04	1524692	6.46	218127658	8.5	0.851
Satellite	526306	0.63	1151170	0.84	105150	0.45	4160764	0.16	3.938
Simple	957019	1.15	1324043	0.97	293020	1.24	6040369	2.34	0.491
Low complexity	268705	0.32	382987	0.28	80465	0.34	16757951	0.65	0.492
Total repeat	24152896	29.1	41894101	30.69	8614159	36.47	1094133363	42.65	0.682
Total bp analyzed	82887797		136512088		23609256		2565547630		

The repeat contents of four regions of the rat genome were compared: duplicated regions as detected by whole-genome analysis comparison; duplicated blocks where pairwise alignments within 100 kb were merged; 20-kb flanking regions immediately flanking the clustered duplications and the genome average. Enrichment was defined as the repeat content of duplicated sequence divided by the repeat content of unique sequence.

gene clusters which proved difficult to integrate into the genome assembly due to the high degree of sequence identity.

DISCUSSION

We present a preliminary analysis of recent segmental duplication content of the rat genome. In order to avoid some of the difficulties and artifacts associated with detection and characterization of low-copy repeat sequence (Bailey et al. 2001), we

implemented several precautions during our in silico analysis of the draft sequence. First, to avoid overestimating segmental duplication content due to common repeats, we purposefully set our alignment length criteria to exclude uncharacterized retroelements considering thresholds at both 5 and 10 kb. Second, we considered separately the proportion of duplications with near perfect sequence identity (Fig. 1; Table 1). Initial analyses of the Human Genome Project overestimated the amount of segmental

A

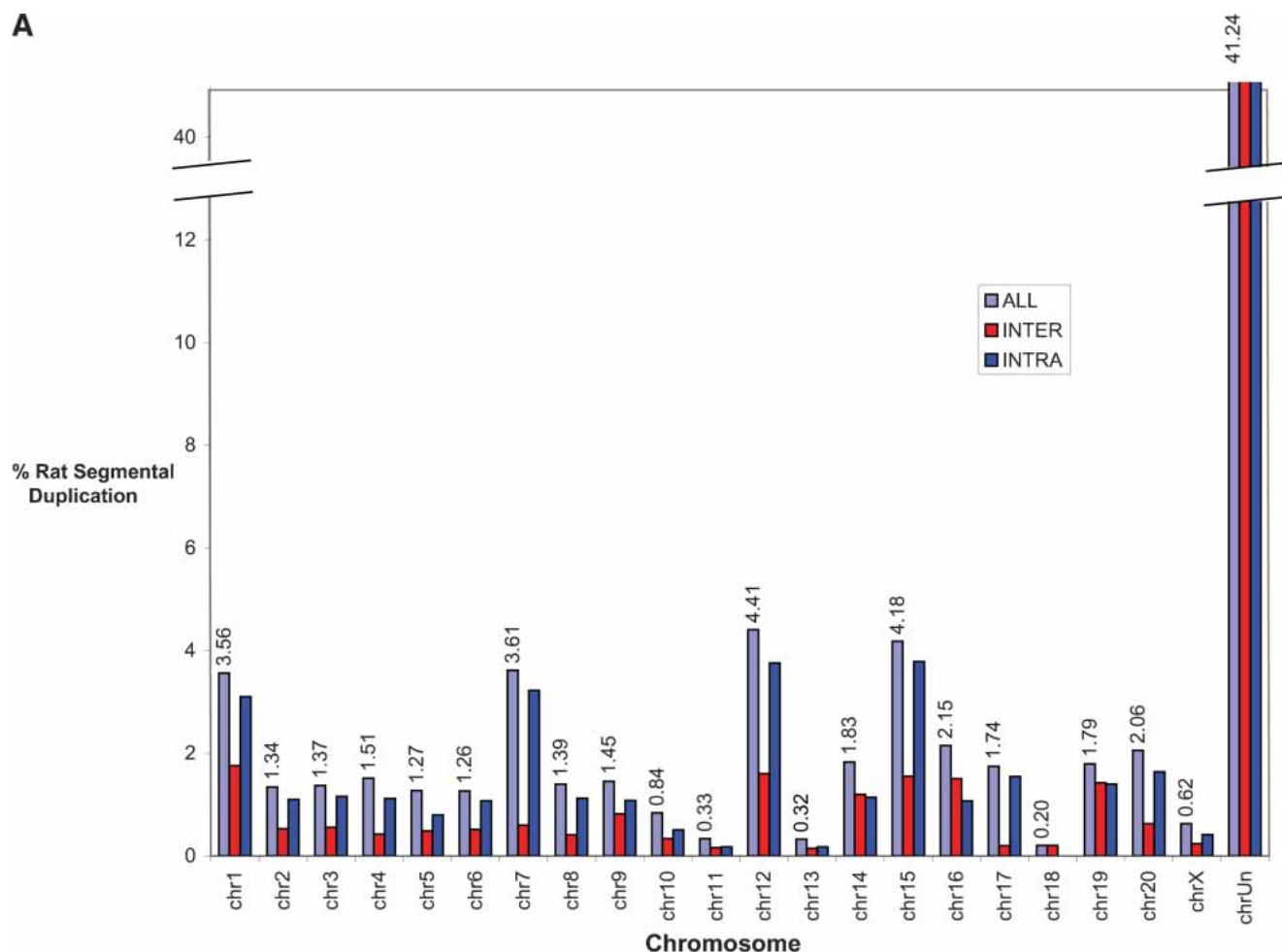


Figure 4 (Continued on next page)

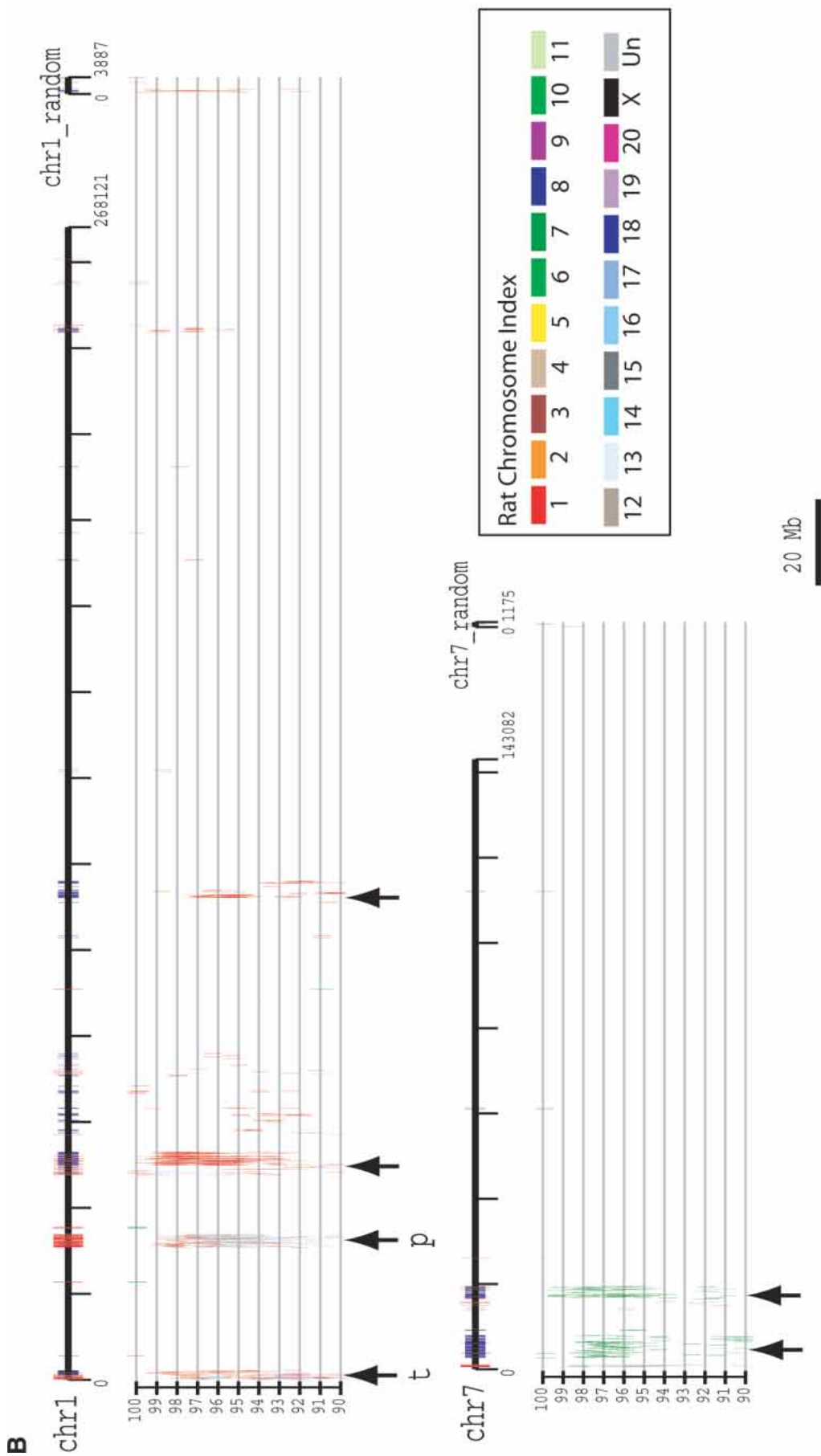


Figure 4 (A) Segmental duplication content per chromosome. The relative proportion of intrachromosomal and interchromosomal duplications for each chromosome is shown. The above calculations treat the unmapped sequence as a separate chromosome when classifying duplications as inter- or intrachromosomal. Forty-five percent of the unplaced chromosome is made up almost entirely of duplicated sequence. (B) Duplication blocks. Rat segmental duplications clustered into larger regions ranging from 100 to 3000 kb in length. We termed these structures “duplication blocks.” Examples of duplication blocks on chromosomes 1 and 7 are presented (arrows) with the underlying degree of sequence identity for each pairwise depicted below the graph. Chromosome 1, green; chromosome 7, red. A subtelomeric (t) and pericentromeric (p) block are indicated. The regions of the rat genome are typified by low gene density (RefSeq/EST/mRNA), a high frequency of gaps within the assembly, and an excess of pairwise alignments.

Table 4. Block Structure of Rat Segmental Duplications

Block Size	Assigned	Unknown	Total
≥2 Mb	12	5	17
≥1 Mb	19	22	41
≥500 kb	25	49	76

Unknown refers to unplaced sequence either random or “unknown” chromosome.

duplication as much as threefold due to a failure to merge allelic overlap during the clone-ordered assembly process (IHGSC 2001). Subsequent analyses showed that such artifactual duplications were readily distinguished by an unusually high degree of sequence identity consistent with allelic levels of variation ($\geq 99.8\%$). Our analysis of the rat genome indicates that the vast majority ($\geq 91\%$) of the pairwise alignments show sequence identity $< 99.5\%$ —far below the estimated levels of sequence variation due to error and/or allelic variation. In fact, the individual rat used for genome sequencing was highly inbred, with virtually no allelic variation—the product of more than 50 brother–sister matings (Methods). The fact that the majority of duplication alignments were $< 99.5\%$ identical suggests relatively few false positives during this analysis.

There are some limitations of this analysis that should be noted. Regions of extremely high sequence identity may have been collapsed during assembly. Thus, the relative small fraction

of the genome that shows duplication $\geq 99.5\%$ (Fig. 1) may be an underestimate. The fact that the α -2-globulin cluster shows only five duplicated genes as opposed to the estimated 15–20 copies (McFadyen et al. 1999; McFadyen and Locke 2000) may be a consequence of such an effect. Although many of the expected rat gene duplications and highly homologous gene families (i.e., carboxylesterases, α -2-globulin, cytochrome P450 genes, serine protease inhibitor, T-cell receptor, MHC genes, etc.; Atchison and Adesnik 1986; Pages et al. 1990; Yan et al. 1995; McFadyen et al. 1999; Ioannidu et al. 2001; Oldfield et al. 2001; Rolstad et al. 2001) were validated during our analysis, not all were detected. For example, the pancreatic type ribonuclease I represents a single-copy gene within most mammalian lineages that has expanded specifically within the genus *Rattus* (Dubois et al. 2002). It was not detected as a duplicated gene within rat genome assembly v. 3.1 by our criteria. A more detailed analysis of the gene showed that segmental duplications were indeed present, but the effective length of these alignments was less than 5 kb (below our detection threshold). Surprisingly, duplications of this gene were detected within a ≥ 5 -kb pairwise alignment within a previous rat genome assembly (v.2.1; <http://ratparalogy.gene.cwru.edu>). The presence of sequence gaps, changes in sequence contig orientation, and our length threshold prevented its detection within the newer assembly. It is clear that duplications have been problematic during sequence and assembly. The analysis of the unplaced chromosome and random chromosomal sequence provides the best testament to this effect. The “unplaced” chromosome showed a marked enrichment for blocks of segmental duplication, with almost half (36.1/82.8 Mb) of the duplications

Table 5. Largest Blocks of Segmental Duplication in the Rat Genome

Chromosome	v.3.1begin	v.3.1end	Block size	Homology	Content
chr15	28153750	31785691	3631941	chr15	T-cell receptor
chr1	49778386	53005805	3227419	chr1	noRefseq, noRatmRNA, noRatEst, but homology to non-rat mRNA
chr1	30806167	33768308	2962141	chr1, 3, 7, 12, 16	noRefseq, noRatmRNA, limited RatEst
chr7	4210045	6569239	2359194	chr7	noRefseq, noRatmRNA, noRatEst, but homology to non-rat mRNA
chr1	5500	2196674	2191174	chr1, 7, 9, 14	noRefseq, noRatmRNA, limitedRatEst, subtelomeric homology
chr14	48526685	50326618	1799933	chr14, 9	noRefseq, noRatmRNA, RatEST
chr15	19229668	20983782	1754114	chr15, 3, 12	prostaglandin D2 receptor
chr15	4606290	6304612	1698322	chr15	MIC2 like 1/Rhombex40
chr7	2608728	4103847	1495119	chr7	noRefseq, noRatmRNA, noRatEst, but homology to human mRNA
chr19	23253330	24673455	1420125	chr14, 19	noRefseq, noRatmRNA, noRatEst, but homology to non-rat mRNA
chr12	18092439	19509542	1417103	chr12	noRefseq, noRatmRNA, limited RatEst
chr6	138660326	140060338	1400012	chr6	Immunoglobulin heavy chain variable region (IGHV)
chr17	67155883	68547346	1391463	chr17	noRefseq, noRatmRNA, RatEst
chr2	14691	1387166	1372475	chr1, 2, 3, 7, 16	noRefseq, limited RatmRNA and RatEST, subtelomeric
chr7	18156845	19521121	1364276	chr7(prim), 8, 9	noRefseq, noRatmRNA, RatEST
chr7	16446057	17651248	1205191	chr7	noRefseq, noRatmRNA, RatEST (conserved)
chr3	20974	1139288	1118314	chr1, 2, 3, 7, 12, 15, 16	noRefseq, noRatmRNA, RatEST
chr17	7851444	8937241	1085797	chr17	Cathepsin M ^a
chr8	36551927	37612862	1060935	chr8	ATPase inhibitor
chr1	112988982	114038449	1049467	chr1	noRefseq, noRatmRNA, RatEST
chr5	78092980	79128821	1035841	chr5	alpha 2 μ globulin PGCL3 ^b Zfp37
chr9	4870998	5902959	1031961	chr9, 14, 16	noRefseq, noRatmRNA, RatEST

Blocks were defined as clusters of segmental duplication with < 100 kb of intervening sequence between duplons. The largest 22 blocks which were assigned to a chromosome are shown. For a complete listing of all blocks, see <http://ratparalogy.gene.cwru.edu>. Begin and end coordinates within build 3.1, block size, and homologous regions are shown. Content was based on assigned Refseq, Rat mRNA, and ESTs within intron/exon structure within the UCSC browser. ^aBest sequence match of Cathepsin M is on chromosome 5. ^bEstimated 20 copies, but only five copies can be distinguished within the assembly.

Table 6. Genes Within Rat Segmental Duplications

Accession	Gene name	Gene product	Chrom	txStart	txEnd	Exon count	Exons hit	Gene size	# Dupbp
NM_181693	Adam28	A disintegrin and metalloprotease domain 28	chr15	49036202	49100982	22	12	2357	1200
NM_023103	Mug1	Alpha(1)-inhibitor 3, variant I	chr4	158528836	158582575	36	21	4656	2829
NM_147214	LOC259246	Alpha-2 μ globulin PGCL1	chr5	78712436	78715858	7	7	878	878
NM_147212	LOC259244	Alpha-2 μ globulin PGCL3	chr5	78094054	78171960	7	7	807	807
NM_147215	LOC259247	Alpha-2 μ globulin PGCL4	chr5	78093991	78097430	7	7	1010	1010
NM_147213	LOC259245	Alpha-2 μ globulin PGCL5	chr5	78686553	78689873	7	7	733	733
NM_012718	Andpro	Androgen regulated 20 kDa protein	chr3	137991232	137997597	4	4	828	828
NM_032072	Appbpl	APP-binding protein 1	chr19	382555	408667	20	7	1780	692
NM_012915	Atpi	ATPase inhibitor	chr8	36902394	36905032	3	3	415	415
NM_022281	Abccl	ATP-binding cassette, sub-family C (CFTR/MRP)	chr10	452238	575705	31	13	4998	2360
NM_031565	Ces1	Carboxylesterase 1	chr19	14892725	14929187	14	11	1936	1389
NM_133295	Ces3	Carboxylesterase 3	chr19	14933410	14971894	14	14	1892	1892
NM_144743	LOC246252	Carboxylesterase isoenzyme gene	chr19	37685	44736	12	12	1872	1872
NM_181378	Ctsm	Cathepsin M	chr17	8931467	8936876	8	8	1355	1355
NM_031561	Cd36	CD36 antigen	chr4	13472462	13554416	13	12	2447	2285
NM_153313	CYP2D1	Cytochrome P450 2D1	chr7	120803930	120808335	9	9	1632	1632
NM_017158	Cyp2c39	Cytochrome P450, 2c39	chr1	243799046	244827001	9	3	1591	754
NM_017158	Cyp2c39	Cytochrome P450, 2c39	chr1	243935780	244719024	12	10	1737	1518
NM_173304	Cyp2d5	Cytochrome P450CMF1b	chr7	120794663	120799170	9	9	1599	1599
NM_022849	Dmbt1	Deleted in malignant brain tumors 1	chr1	190539537	190620410	33	7	4360	1635
NM_138902	Loc192264	Eosinophil cationic protein	chr15	27287836	27288707	2	2	713	713
NM_053689	Crfg	G protein-binding protein CRFG	chr17	72111731	72131468	17	11	1927	1267
NM_181440	Grp-Ca	Glutamine/glutamic acid-rich protein GRP-Ca	chr4	170757736	170828910	5	5	876	876
NM_138517	Gzmb	Granzyme B	chr15	35211149	35214166	5	5	1035	1035
NM_019261	Klrc2	Killer cell lectin-like receptor subfamily C	chr4	167201553	167212693	7	6	1309	751
NM_133421	Lkap	Limkain b1	chr10	826802	872214	28	27	7645	7519
NM_152848	Ly49i2	Ly49 inhibitory receptor 2	chr4	168584327	168608948	7	7	1522	1522
NM_153726	Ly49s3	Ly-49 stimulatory receptor 3	chr4	168268533	168454309	7	4	1401	932
NM_173291	Ly49	Lymphocyte antigen 49 complex	chr4	168135002	168354511	9	9	1145	1145
NM_134459	Mic211	MIC2 like 1	chr15	5673663	5719158	11	4	4177	359
NM_022247	Pdcl	Phosducin-like protein	chr3	17003800	17012826	4	1	2303	1825
NM_022241	Ptgdr2	Prostaglandin D2 receptor	chr15	19345554	19352929	2	2	1317	1317
NM_080770	Psbp1	Prostatic steroid-binding protein 1	chr1	212320755	212323535	3	3	518	518
NM_173315	LOC286981	Putative pheromone receptor (Go-VN2)	chr1	71238505	71258976	6	2	3572	512
NM_173318	LOC286984	Putative pheromone receptor (Go-VN4)	chr1	61650206	61748606	8	8	3650	3650
NM_173320	LOC286986	Putative pheromone receptor Go-VN13C	chr18	34192	61266	6	5	3346	3111
NM_173113	VN1	Putative pheromone receptor VN1	chr4	124479450	124720930	2	1	1378	251
NM_173298	VN2	Putative pheromone receptor VN2	chr4	124479445	124487113	3	3	1663	1663
NM_012646	RT1-N1	RT1 class 1b gene, H2-TL-like, grc region	chr20	278581	2789031	8	8	1186	1186
NM_176076	S100RVP	S100 calcium-binding protein	chr2	183684144	183687494	4	4	831	831
NM_012657	Spin2b	Serine protease inhibitor 2b	chr6	128383141	128390546	5	5	1669	1669
NM_031664	Slc28a2	Solute carrier family 2, member 2	chr3	109256103	109276805	17	17	2644	2644
NM_053752	Suclg1	Succinate-CoA ligase, GDP-forming, alpha	chr8	36133697	36137263	4	4	476	476
NM_133547	Sult1c2	Sulfotransferase family, cytosolic, 1C, member	chr9	1030810	1160809	7	4	2432	2029
NM_058209	Zfp37	Zinc finger protein 37	chr5	78484370	79157374	6	3	2492	2167

^aOnly genes within segmental duplications where the alignments were between 90%–99% identical are shown. A total of 63 genes were detected within duplications 90–100% identical. The number of exons (exons hit) and genic bases within the duplicated region (Dupbp) are indicated. Gene size is the sum of exon lengths from the rat genome assembly.

assigned to this category. Further targeted efforts are required to resolve the true location, organization, and complexity of these regions.

Despite these methodological and assembly limitations, some important trends regarding rat segmental duplications emerged during our study. The overall content of highly homologous duplications as determined by the sequence assembly is greater within the rat (2.92%) than the mouse (1%–1.2%; Cheung et al. 2003b). Both are significantly reduced for segmental duplications compared to human (4.78%) for similar length thresholds (>5 kb; Bailey et al. 2002). The threefold difference between rat and mouse is surprising and may reflect biological differences or differences in the strategy for genome assembly. The mouse genome assembly strategy depended almost solely upon whole-genome shotgun (WGS) assembly, which has been predicted to overcollapse segmental duplication (Eichler 1998, 2001; Waterston et al. 2002). In contrast, the rat genome was assembled using a hybrid strategy, termed “BAC-enrichment.” The BAC-enrichment hybrid strategy entailed low-pass sequencing of 20,987 individual rat BAC clones, followed by an enrichment phase where individual WGS reads were mapped to specific BAC projects based on sequence overlap (RGSPC 2004). In such a scenario, paralogous regions within BACs would compete to optimally place WGS mate pairs and in so doing prevent overlap of duplicated regions.

Based on the current assembly, recent duplications are distributed in a nonuniform fashion across the genome. In addition to chromosome differences, we identified 41 duplication blocks (Fig. 3B) over 1 Mb in size. The extreme variation in sequence identity underlying the pairwise alignments (<http://ratparalogy.gene.cwru.edu>) within these blocks suggests that these areas have been the target of recurrent duplication over millions of years of evolution. The majority of duplications are organized as clusters of tandem or inverted intrachromosomal duplications. A similar bias toward clustered duplications was observed in the mouse genome assembly (Cheung et al. 2003b). Regions of extensive interchromosomal duplication were observed, particularly near the subtelomeric and pericentromeric regions. In the absence of detailed mapping information regarding the precise positions of centromeres and telomeres, it is difficult to assess these properties for all rat chromosomes. Our preliminary analyses of the rat genome, however, clearly shows a pericentromeric and subtelomeric bias for segmental duplications, suggesting that these may be general properties of mammalian chromosomal architecture. An analysis of the evolutionary genetic distance of all segmental duplications as a function of the sum of aligned base pairs (43,597 alignments) showed a bimodal distribution, particularly for intrachromosomal segmental duplications. Two peaks were observed, at 0.045 substitutions per site and 0.075 substitutions per site. Assuming that the rat and mouse lineages diverged 16–23 million years ago (Springer et al. 2003) and a neutral sequence divergence range of 0.173–0.195 years (RGSPC 2004), this bimodal distribution may correspond to bursts of segmental duplication that occurred approximately four and eight million years ago, respectively.

An analysis of the RefSeq genes (Methods) showed that segmental duplications are generally gene-poor based on their genomic representation (~1.3% vs. 2.9%). Of the 63 genes that were identified within duplicated sequence, 33 were part of alignments which contained a complete complement of exons. Most of the duplications that contained genes were part of intrachromosomal alignments. A similar effect was observed when assigned rat mRNA was considered. This suggests that regions containing interchromosomal duplications are conspicuously transcriptionally silent. Our analysis was designed to recover genes that had emerged specifically within the rat lineage, because

aligned genomic sequence between rat and mouse shows on average 0.175–0.195 substitutions per site and our study was limited to alignments showing less than 0.10 substitutions/site. Many of the rat duplication gene clusters recovered during this analysis (natural-killer cell receptor, serine protease inhibitor, carboxylesterase, cytochrome P450 gene families, etc.; Atchison and Adesnik 1986; Pages et al. 1990; Yan et al. 1995; McFadyen et al. 1999; Ioannidu et al. 2001; Oldfield et al. 2001; Rolstad et al. 2001) were also detected during an analysis of “recent” segmental duplication within the mouse lineage (Cheung et al. 2003b). The fact that such tandem duplications with extensive sequence identity exist within both lineages argues for active gene conversion (Atchison and Adesnik 1986) to maintain such homologous structures within each species.

Since the original analyses of working draft sequences of human and mouse (IHGSC 2001; Waterston et al. 2002), global studies of segmental duplication content have become an effective measure to assess one aspect of the quality of whole-genome sequence assemblies (Bailey et al. 2001, 2002; Cheung et al. 2001). Regions of recent segmental duplication remain one of the greatest challenges to finishing a genome sequence. Within the “finished” human genome assembly, for example, there is a striking correspondence between the position of sequence “gaps” among finished chromosomes and regions of large highly homologous duplications. Such areas have proven problematic for both clone-based methods and whole-genome shotgun sequence (Bailey et al. 2001, 2002; Cheung et al. 2001). In general, it is well recognized that the greater the proportion of large, highly homologous repeats, the more difficult a genome is to finish. Among certain genomes such as the human and mouse, high-quality ordered and oriented finished genome sequence is the stated goal. Concomitantly, it is expected that the structure and organization of such regions will ultimately be resolved—albeit with considerable effort and expenditure (Eichler 2001). Among other genomes such as the rat, finished genome sequence is not the stated goal. An initial assessment of segmental duplication content therefore provides an important level of annotation for the user of genome sequence information in the design and interpretation of experiments. Moreover, we argue that these initial analyses precisely delineate potential regions where whole-genome shotgun or a BAC-enrichment strategy will provide insufficient information for the biologist. In this study for example, we have identified <100 regions where the segmental duplications and bona fide gene families intersect. These regions include gene families important in drug detoxification, chemotaxis, and immunity. The content and structure of these regions will be pivotal to the full realization of the rat as a physiological model of pharmacology and complex genetic diseases (Jacob and Kwitek 2002). We therefore propose that such highly duplicated, generic regions be uncoupled from WGS sequencing strategies and be targeted for high-quality BAC-based finishing. The analysis presented here should provide a framework for the prioritization of such regions.

METHODS

Genome Resources

All reported analyses were performed on the June 2003 rat genome assembly (version 3.1). A complete segmental duplication analysis was also performed on an earlier assembly (version 2.1). The results of both analyses including pairwise sequence alignment locations, statistics, and gene content are available at <http://ratparalogy.gene.cwru.edu>. Segmental duplication analyses for version 3.1 have been added as a segmental duplication browser track as part of the UCSC browser (<http://genome.ucsc.edu>). Both rat genome assemblies were constructed using the

BAC-enrichment strategy, which represents a hybrid between whole-genome shotgun sequence and clone-ordered approaches (see <http://www.hgsc.bcm.tmc.edu/> for details). Genome sequences used in this study were derived from an inbred strain (BN/SsNHsd/MCW) of the brown Norway rat (*Rattus norvegicus*). The original inbred founder pair Harlan Sprague Dawley showed limited allelic variation; 6 of ~4338 microsatellite loci (<http://rgd.mcw.edu/>). Brother–sister matings were performed for an additional 13 and 14 generations. A mother–daughter pair were the source of the whole-genome shotgun sequence library and the large-insert BAC library (CHORI-230).

Rat Segmental Duplications Detection

To analyze rat segmental duplications, we applied a BLAST-based whole-genome assembly comparison (Bailey et al. 2001). This BLAST-based method was designed to detect highly similar ($\geq 90\%$ identity) lineage-specific segmental duplications (≥ 1 kb) after extracting common repeat sequences. We applied this method to the rat but detected an excess of smaller putative segmental duplications (Table 1) after using an updated Repeat-masker library database (June 2003). Upon inspection, many of the shorter alignments corresponded to incompletely masked high-copy repeats (LTR elements) or composite repeat elements (LTR/LINE hybrids). Because our detection algorithm extends seeding alignments into adjacent high-copy repeats, partially masked repeats will be lengthened to include the entire element. To circumvent the high-copy repeat overabundance, we selected a higher length threshold (≥ 5000 bp of seeding sequence). At this threshold, most uncharacterized transposable element alignments were eliminated. These seeding alignments were then trimmed to better define their end points, and optimal global alignments were performed to generate accurate alignment statistics. Alignments were then joined for gaps up to 10 kb in size. To avoid the potential of larger transposable elements as well as composite repeats, we considered various length thresholds (5, 10, and 20 kb). Sequence alignment statistics were calculated from optimal global alignments as described (Bailey et al. 2001), and paralogous sequence relationships were generated using Parasight graphical visualization software (J. Bailey, unpubl.).

Block-Size Delineation

We clustered duplications into larger blocks by examining the proximity of flanking sequences. A “weld” was performed if another pairwise alignment was identified within 100 kb from the coordinates of a pairwise alignment (Table 3). Gaps were not included in this calculation. Clustering proceeded in both directions from the seed pairwise alignment until a unique region (no duplications) of at least 100 kb was encountered per each cluster. (Table 3). Analysis of flanking sequences was performed based on these “weld” coordinates.

Gene Analysis

Gene content of rat segmental duplications was assessed using two differences sources of data: LocusLink RefSeq gene annotations and rat mRNAs in GenBank. All mRNAs were aligned using BLAT as described (Kent et al. 2002), and intersections between segmental duplication coordinates and exon positions were compared using MySQL queries of the UCSC browser database. During our analysis, a total of 63 RefSeq genes (from a genome total of 4532) and 945 rat mRNAs (from a genome total of 11,560) were identified that had been assigned to duplicated regions. Of these, 716 mRNAs were identified that did not overlap with RefSeq gene coordinates. In addition, 61/63 of the RefSeq genes contained two or more exons.

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- <http://rgd.mcw.edu/>; Rat Genome Database at Medical College of Wisconsin.

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