## Research

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# Genome-wide detection of segmental duplications and potential assembly errors in the human genome sequence Joseph Cheung<sup>\*</sup>, Xavier Estivill<sup>\*†</sup>, Razi Khaja<sup>\*</sup>, Jeffrey R MacDonald<sup>\*</sup>, Ken Lau<sup>\*</sup>, Lap-Chee Tsui<sup>\*‡§</sup> and Stephen W Scherer<sup>\*‡</sup>

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#### Abstract

**Background:** Previous studies have suggested that recent segmental duplications, which are often involved in chromosome rearrangements underlying genomic disease, account for some 5% of the human genome. We have developed rapid computational heuristics based on BLAST analysis to detect segmental duplications, as well as regions containing potential sequence misassignments in the human genome assemblies.

**Results:** Our analysis of the June 2002 public human genome assembly revealed that 107.4 of 3,043.1 megabases (Mb) (3.53%) of sequence contained segmental duplications, each with size equal or more than 5 kb and 90% identity. We have also detected that 38.9 Mb (1.28%) of sequence within this assembly is likely to be involved in sequence misassignment errors. Furthermore, we have identified a significant subset (199,965 of 2,327,473 or 8.6%) of single-nucleotide polymorphisms (SNPs) in the public databases that are not true SNPs but are potential paralogous sequence variants.

**Conclusion:** Using two distinct computational approaches, we have identified most of the sequences in the human genome that have undergone recent segmental duplications. Near-identical segmental duplications present a major challenge to the completion of the human genome sequence. Potential sequence misassignments detected in this study would require additional efforts to resolve.

#### Background

Segments of DNA with near-identical sequence (segmental duplications or duplicons) in the human genome can be hot spots or predisposition sites for the occurrence of non-allelic homologous recombination or unequal crossing-over leading to genomic mutations such as deletion [1], duplication [1], inversion [2] or translocation [3,4]. These structural

alterations, in turn, can cause dosage imbalance of genetic material or lead to the generation of new gene products resulting in diseases defined as genomic disorders [5].

Previous studies to identify segmental duplications in the human genome have analyzed older versions of the genome assembly, which contained higher amounts of unfinished sequence and incorrectly mapped regions, and have used different computational approaches all performed by the same group [6-8]. With the human genome sequence now nearing completion, we have examined its content for segmental duplications using two distinct computational methods. In the first, we utilized the rapid BLAST2 [9] algorithms that allow direct chromosomal-wide sequence comparisons to be made. All BLAST results reported in table formats can be subsequently grouped, parsed and analyzed for the detection of duplicated sequences. In addition, we have shown previously that there is a strong correlation between ambiguously mapped SNPs (ambSNPs), as well as the density of SNPs, and segmental duplications [10]. AmbSNPs are SNPs that were annotated to map to two locations on a particular chromosome in the NCBI dbSNP. A subset of these ambSNPs are not true SNPs but are likely to be computer-generated nucleotide mismatches from paralogous copies of duplicated sequences and should be more appropriately labeled as paralogous sequence variants (PSVs) [10]. Another subset is likely to be false ambSNPs of genomic sequences that have been misassigned in genome assemblies. Here, we report our analysis of all potential PSVs in the human genome and their correlation with segmental duplications as detected by our BLAST analysis. Furthermore, we provide a critical assessment on the three latest human genome assemblies from our analysis of sequence misassignments as identified from this study.

# Results and discussion

## Human genome segmental duplication content

On the basis of the June 2002 (NCBI Build 30) human genome assembly, a total of 107.4 Mb (3.53%) of the human genome content (3,043.1 Mb) were found to be involved in recent segmental duplications by our BLAST analysis criteria (Table 1). This content is composed of more than 1,530 distinct intrachromosomal segmental duplications (80.3 Mb or 2.64% of the total genome, Figure 1) and 1,637 distinct interchromosomal duplications (43.8 Mb or 1.44% of the total genome). In addition, 29% of all duplications are located in unfinished regions of the current genome assembly. Our results are shown using the Generic Genome Browser [11,12]. We have also found that 38% of the duplications (52.3 Mb) can be considered as tandem duplications - defined here as two related duplicons separated by less than 200 kb.

In this study, we only analyzed large (size  $\geq 5$  kb) and recent (sequence identity  $\geq 90\%$ ) duplications because we can achieve higher confidence and to prioritize those regions for their potential involvement in diseases. Previously, Bailey and colleagues [8] reported a total of 5.2% of the human genome involved in recent segmental duplications. The 1.6% discrepancy between our findings could be due to the difference in our detection criteria (size cutoff of 5 kb used in this study versus 1 kb used in Bailey *et al.* [8]). Moreover, we have identified 38.9 Mb of sequences (1.28% of the June 2002 genome assembly) likely to be artifactual duplications resulting from sequence misassignment errors present in the assembly. By comparing our results with those published previously [8], we found that 482/2579 clones that we identified to be involved in duplication were novel.

The molecular mechanism by which segmental duplications are created is still unclear at the moment. A recent report has suggested that Alu repeat clusters had a role as mediators of recurrent chromosomal rearrangements [13]. We have examined whether elevated amounts of repetitive elements could be found in duplicon junctions. We inspected all duplication borders from our results and calculated the occurrence of different repeat types within the 500 bp window outside each duplicon junction. The whole-genome average frequencies were determined by sampling random 500 bp windows across the genome (excluding gap regions). Overall, we found that there are significant enrichment (or relative fold increase) for the presence of small ribonucleoprotein RNA (srpRNA), satellite, long terminal (LTR) and SINE/Alu repeats (see Additional data file). In addition, our data also showed that for some chromosomes the amount of duplicated sequence is higher in the pericentromeric and subtelomeric regions of chromosomes (Figure 1), supporting the hypothesis that these repeat-dense regions have made an important contribution to the evolution of the human genome [14].

Regions containing recently occurring segmental duplications can harbor rapidly evolving hominoid-specific genes, as well as novel gene families that are unique to primates [15,16]. Using the National Center for Biotechnology Information RefSeq annotation, we identified 1,152 human genes that were mapped to duplicated regions. Of these, 475 genes were fully contained within duplicated regions and were best candidates for recent whole-gene duplication. We have carried out functional analysis of these 475 genes using the Gene Ontology Consortium database [17] and found that there is a significant increase in gene duplications for genes involved in immune defense (antibodies, blood-group antigens) and reproduction (pregnancy, sex differentiation) (see Additional data file).

# Sequence misassignment errors in the human genome sequence assembly

We were aware that *in silico* detection methods, such as the ones used in this study, would not allow us to distinguish completely true duplications from artifactual duplications arising from misassigned sequences, especially in cases where sequence identity between two detected duplications exceeded 99.5% over a substantial length (> 5 kb) in regions composed of draft sequences. Although a small proportion of such results (duplications with > 99.5% identity) might represent unfinished regions of the genome that contain true duplications that have arisen very recently in the evolution of the human genome (such as the large and nearly perfect

#### Table I

#### Segmental duplication content of the human genome

Chromo- some	Size (bp)	Intra- chromosomal duplication (bp)	% chromosome (previous)*	Inter- chromosomal duplication (bp)	% chromosome (previous)*	Total duplications (bp)	% chromosome (previous)*	Errors <sup>†</sup> (bp)	% chromosomes
I	246,874,334	5,278,549	2.1 (4.4)	2,854,898	1.2 (2.3)	7,056,274	2.9 (5.7)	4,369,406	1.8
2	240,681,600	4,917,160	2.0 (2.4)	3,298,723	1.4 (1.6)	6,892,585	2.9 (3.2)	2,311,522	1.0
3	194,908,136	2,128,493	1.1 (2.3)	1,654,201	0.8 (2.0)	3,146,570	1.6 (3.2)	3,979,610	2.0
4	192,019,378	2,599,650	1.4 (2.3)	2,164,382	1.1 (2.2)	4,061,432	2.1 (3.4)	2,482,740	1.3
5	180,966,400	3,519,480	1.9 (2.0)	1,464,945	0.8 (1.3)	4,530,406	2.5 (2.8)	2,297,998	1.3
6	170,309,517	2,358,252	1.4 (2.3)	743,875	0.4 (1.3)	2,877,392	1.7 (3.4)	569,918	0.3
7	157,432,793	8,636,434	5.5 (6.3)	2,614,326	1.7 (2.9)	10,139,669	6.4 (7.8)	205,130	0.1
8	143,874,322	2,318,984	1.6 (2.2)	1,125,241	0.8 (2.0)	2,612,280	1.8 (3.0)	3,956,756	2.8
9	132,438,756	7,248,232	5.5 (7.1)	4,801,871	3.6 (4.7)	8,341,767	6.3 (8.2)	1,589,734	1.2
10	134,416,750	5,279,301	3.9 (4.3)	1,375,341	1.0 (1.9)	6,334,458	4.7 (5.7)	1,250,157	0.9
П	137,442,545	3,622,080	2.6 (3.3)	1,670,412	1.2 (1.8)	4,363,619	3.2 (4.4)	2,028,875	1.5
12	131,300,572	1,894,547	1.4 (2.3)	971,490	0.7 (1.2)	2,816,187	2.1 (3.3)	3,383,730	2.6
13	113,446,104	918,255	0.8 (1.9)	1,202,102	1.1 (2.3)	1,855,806	1.6 (3.4)	146,198	0.1
14	104,324,908	531,219	0.5 (0.7)	820,880	0.8 (1.6)	1,335,177	1.3 (2.1)	13,814	0.0
15	99,217,355	4,593,233	4.6 (6.2)	2,344,618	2.4 (3.9)	5,634,201	5.7 (8.2)	1,739,894	1.8
16	81,671,585	4,917,218	6.0 (8.3)	2,228,116	2.7 (3.9)	6,012,178	7.4 (9.8)	2,113,843	2.6
17	80,052,782	4,775,137	6.0 (7.1)	646,968	0.8 (2.5)	5,274,195	6.6 (8.5)	2,145,614	2.7
18	77,516,809	525,636	0.7 (1.2)	700,654	0.9 (1.9)	1,226,290	1.6 (3.1)	1,443,775	1.9
19	60,013,307	2,700,984	4.5 (6.8)	704,757	1.2 (2.6)	3,156,687	5.3 (8.1)	335,190	0.6
20	62,842,997	592,441	0.9 (1.1)	873,152	1.4 (1.8)	1,052,248	1.7 (2.1)	147,940	0.2
21	44,626,493	481,879	1.1 (1.4)	1,303,776	2.9 (5.1)	1,504,333	3.4 (5.2)	0	0.0
22	47,748,585	1,741,766	3.6 (6.7)	1,374,363	2.9 (7.4)	2,770,386	5.8 (10.9)	0	0.0
Х	14,924,9818	2,625,206	1.8 (3.6)	2,927,714	2.0 (2.3)	5,518,712	3.7 (5.5)	2,185,046	1.5
Y	58,368,225	5,959,836	10.2 (28.4)	3,524,276	6.0 (25.0)	8,461,355	14.5 (40.7)	56,204	0.1
Un‡	1,391,854	179,709	12.9 (20.4)	378,110	27.2 (32.6)	407,013	29.2 (36.5)	116,923	8.4
Total 3	3,043,135,925	80,343,681	2.6 (3.8)	43,769,191	1.4 (2.6)	107,381,220	3.5 (5.2)	38,870,017	1.3

\*Previous data on segmental duplications distributed by chromosomes as reported in [8]. †Errors represent data that were detected as potential sequence misassignments. ‡Un, unmapped chromosome sequence.

palindromic repeats located in the AZFc region on chromosome Yq11.223 involved in male infertility [18]), we suspect that most of the duplications (> 99.5% identity and contain draft sequences) are in fact sequence misassignment errors in the genome assembly. An explanation for such errors would be when two identical sequences belonging to the same genomic location were misassigned to distinct regions in the genome assembly.

We have used the NCBI e-PCR [19] to evaluate our results (potential sequence misassignment errors) from the June 2002 human genome assembly. Using some of the largest interchromosomal misassignment errors detected in our study, we found that none of the STS markers located within these misassigned sequences maps to their incorrectly assigned chromosomes. For example, AC121339 is incorrectly mapped to 3q13.13 in the June 2002 genome assembly, as supported by a consensus number of chromosome X sequence-tagged site (STS) markers (Figure 2, Table 2).

From this genome assembly, we identified that a total of 38.9 Mb of sequences, representing 1.28% of the total sequence content, are involved in such potential errors (a full list of potentially misassigned sequences can be obtained from [12]) that would require additional effort and further sequencing to achieve resolutions. We also analyzed an



Figure I (see legend on the following page)

Query sequence:	AC121339,	1939461	bases
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#### from 3q13.13

Site (bases)	Marker	Chr.	Organism
75577665	DXS1407	Χ	Homo sapiens
2236322467	D11S3114		Homo sapiens
2556225796	G34118		Homo sapiens
3014030242	D11S3114		Homo sapiens
3841038570	DXS7950		Homo sapiens
5156251718	DXS1091		Homo sapiens
5163351843	DXS1090		Homo sapiens
5209552429	SHGC-12678	Χ	Homo sapiens
5329653457	RH38682		Homo sapiens
5844258621	DXS7470		Homo sapiens
5845158610	DXS1223	Χ	Homo sapiens
5846958587	AFM309yc1	Χ	Homo sapiens
9123191333	D11S3114		Homo sapiens
114549114649	DYF170S1		Homo sapiens
129101129428	SHGC-143878	Χ	Homo sapiens
133020133167	DXS7947		Homo sapiens
135574135719	D2S1401		Homo sapiens
138385138488	D11S3114		Homo sapiens
145671145808	DXS1497	Χ	Homo sapiens

#### Figure 2

An example of sequence misassignment error as indicated by e-PCR analysis. AC121339 is incorrectly mapped to 3q13.13 in the June 2002 human genome assembly as shown by a consensus number of chromosome X STS markers.

additional two previous versions of the human genome assemblies, December 2001 and April 2002, and our results showed that there has been a dramatic reduction in potential errors in the latest human genome assembly compared to the two previous genome assemblies (Table 3). Furthermore, we examined the distribution of the amount of duplications in five different categories on the basis of their level of sequence identity to each other (Table 3). We observed a large reduction in duplications that fall within the 98-100% category, supporting the fact that the genome assemblies continue to improve and correct errors made. In addition, our data showed that there have been major improvements for chromosomes 5, 6, 7, 13, 14 and 19 over the last three genome assemblies. And for chromosomes that had reached finished status, such as chromosomes 20, 21 and 22, the number of errors was negligible.

### Paralogous sequence variants in the human genome

We have previously shown a strong correlation between ambSNPs with segmental duplications [10]. AmbSNPs are SNPs that were annotated to map to two locations on a particular chromosome in the NCBI dbSNP. Here we show on a genome scale that ambSNPs most specifically correlate with intrachromosomal segmental duplications, suggesting they are paralogous sequence variants (PSVs) (Figure 1). These PSVs were perhaps mistakenly introduced into dbSNP by automated in silico-generated analysis, arising from nucleotide mismatches in paralogous copies of duplicated sequences. Overall, a surprisingly high proportion, 8.6% (199,965 of 2,327,473), of the refSNPs were annotated as ambSNPs from dbSNP (Build 108). A significant number of the ambSNPs (139,974 of 199,965 or 70.0%) are located within duplicated regions as identified by BLAST and should be regarded as PSVs (see Additional data file).

The non-identification by BLAST analysis of regions that contain ambSNPs could be due to one of three possibilities. First, the duplicated copy(s) could have been removed from the sequence assembly or the two have been conflated, that is, mistakenly thought to be the same sequence owing to their high sequence similarity. A second possibility is that the duplication is smaller than 5 kb and was excluded in our BLAST analysis. A third possibility is that a collection of ambSNPs could have been generated from misassigned sequences (identical sequences but misassigned to two difference locations in the genome) in older assembly builds due to sequencing errors or true SNPs (with high polymorphism rate) in the sequence. We also observed that the density of ambSNPs generally correlates with the size of the putative duplication, although this might be affected by the level of sequence identity between duplications. For example, two duplicated sequences sharing 98% sequence identity compared to 95% over the same length might contain fewer PSVs as the number of base-pair mismatches

Figure I (see figure on the previous page)

Intrachromosomal segmental duplications identified in the human genome. Three panels of results are displayed for each chromosome. Left, graphical views of the paralogous relationships between recent segmental duplications (graphics produced using GenomePixelizer [29,30]; each line represents a duplicated module; coloring scheme, red = 99% to 100% sequence identity, purple = 96% to 98%, green = 93% to 95%, and blue = 90% to 92%). Middle panel: segmental duplications as detected by BLAST analysis (size of duplication in kb plotted against the length of chromosome in Mb). Right panel: ambSNPs density plot (number of ambSNPs plotted against the length of chromosome in Mb). All analyses were done using the June 2002 human genome sequence assembly.

Table	2
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Examples of sequence misassignment errors								
Clone*	Location	Size of region involved (bp)	e-PCR results					
AC121339†	3q13.13	193,190	chrX					
AC016003	17q21.31	181,582	chr9					
AC119723	3q22.1	159,924	chr6					
AC093007	3q12.1	169,882	chr6					
AC110578	8p23.2	160,554	chr15					
AC108862	11p15.3	156,150	chr18					
AC113009	8q23.1	155,171	chrll					
AC104765	8q12.1	150,029	chr18					
AC105412	2p13.1	144,924	chr5					
AC092744	12p12.3	144,009	chr4					
AC099061	Ip21.3	140,516	chr15					
AC108735	3p24.3	136,005	chr I 6					
AC122689	3q23	120,057	chr12					
AC017027	l q32. l	116,265	chr5					
AC013530	3q26.1	99,768	chr8					
AC115093	11p15.4	98,715	chrl					
AC112921	Xp22.22	96,272	chr3					
AC108094	16q21	94,953	chr I 7					
AC079186	8q12.1	78,771	chr7					
AC024573	Unmapped	56,016	chr2					
AC115093	11p15.4	53,858	chrl					

\*A full list can be obtained from [12]. <sup>†</sup>See Figure 2 for e-PCR results supporting sequence misassignment.

would be fewer in the former. In addition, we observed that regions identified by our BLAST method but do not contain ambSNPs often correspond to artifactual duplications generated from assembly errors.

#### Duplicons related to genomic disorders

The size, orientation, and contents of segmental duplications are highly variable and most of them show great organizational complexity. This is perhaps due to successive transposition and rearrangement events leading to the creation of segmental duplications [14]. In many cases, a contiguous duplicon is organized into multiple modules with different orientations and sizes. For example, one of the largest segmental duplicons detected in this study was 359 kb in size at the Williams-Beuren locus on 7q11.23 [20,21]. In this case, the two duplicons are separated by 1.6 Mb of intervening sequence with the telomeric duplicon comprising several separate smaller modules as compared to the primary duplicon. The results presented in our study (provided in tables available at [12]) would also allow rapid identification of new duplicons that are potentially responsible for chromosome rearrangements and genomic disorders. For example, the

location of the duplicons on chromosomes 9q34/22q11 that have been suggested to mediate recombination leading to the Philadelphia chromosome [4] was identified in our analysis, as were other medically relevant chromosomal regions (Table 4) [22,23].

The characterization of most large segmental duplications is complicated by the fact that many of them (29% of all duplications) are only represented as draft sequences from the current genome assembly. Despite the fact that both BLAST and PSVs analyses allowed us to identify most segmental duplications involved in known genomic disorder mutations (Table 4), estimations of the size of rearranged regions were different from those previously reported [23]. In fact, with the exception of several small duplications and the segmental duplications on chromosome 22 [24], other regions containing duplications involved in genomic disorders were often erroneously assembled and misplaced. Furthermore, we have searched the Celera human genome C<sub>3</sub> (publicly released version [25]) and C4 (subscription-based version) sequence assemblies for large duplications found on chromosome 7. We observed that most of them were not represented in large scaffolds, but instead were located in their sequence gaps, or only partially found at ends of scaffolds leading into gaps (see Table 4) [26]. This suggests that the whole-genome assembly approach [25] alone might not be able to finish such duplicated regions in mammalian genomes.

#### Conclusions

We have used two different computational approaches to identify the locations of all recent segmental duplications in the current human genome draft sequence. The fidelity of the results reflects the quality of the assembly examined and the parameters used. In addition, our approach has detected numerous potential sequence misassignment errors in the current genome annotation, allowing rapid error detection in future sequence assemblies. The segmental duplication map of the human genome should serve as a guide for investigation of the role of duplications in genomic disorders, as well as their contributions to normal human genomic variability [2,3,27]. It is clear that genomic regions containing segmental duplications present a major challenge to the completion of the human genome sequence by April 2003. Focused efforts including targeted sequencing of allele-specific clones, high-resolution fluorescence in situ hybridization, and expert curation would be required to validate the actual (or proposed) organization of these complex regions as well as to complete the human genome reference sequence.

#### Materials and methods

#### Genome sequence and chromosome-wide BLAST

We obtained the December 2001, April 2002, and June 2002 (NCBI Build 28, 29 and 30 respectively) human genome assemblies through the University of California, Santa Cruz

#### Table 3

	December 2001				April 2002			June 2002		
	Length	Duplications	Errors	Length	Duplications	Errors	Length	Duplications	Errors	
Chrl	2,564	99	115	2,459	68	60	2,469	71	44	
Chr2	2,413	70	45	2,468	79	57	2,407	69	23	
Chr3	2,048	49	90	2,047	29	73	1,949	31	40	
Chr4	1,914	39	44	1,970	51	49	1,920	41	25	
Chr5	1,848	55	90	1896	55	112	1,810	45	23	
Chr6	1,783	58	56	1,828	43	153	1,703	29	6	
Chr7	1,638	130	48	1,605	119	27	1,574	101	2	
Chr8	1,457	35	66	1,484	33	43	1,439	26	40	
Chr9	1,330	83	38	1,291	75	27	1,324	83	16	
Chr10	1,421	74	51	1,385	72	39	1,344	63	13	
Chrll	1,414	51	84	1,341	43	36	1,374	44	20	
Chr12	1,396	30	83	1,342	24	32	1,313	28	34	
Chrl3	1,151	29	21	1,136	22	15	1,134	19	I	
Chr14	1,065	27	8	1,054	23	10	1,043	13	0	
Chr15	991	62	30	1,000	54	20	992	56	17	
Chr16	938	65	44	932	67	32	817	60	21	
Chr17	839	66	46	811	46	29	801	53	21	
Chr18	818	16	59	809	12	32	775	12	14	
Chr19	769	45	28	730	34	12	600	32	3	
Chr20	630	10	5	628	12	4	628	11	I	
Chr21	446	18	3	446	16	2	446	15	0	
Chr22	478	28	0	477	29	L	477	28	0	
ChrX	1,517	54	40	1,518	61	23	1,492	55	22	
ChrY	584	86	2	584	95	2	584	85	I.	
ChrUn	74	10	Ι	125	П	43	14	4	Ι	
Total	31,526	1,290	1,097	31,366	1,175	932	30,431	1,074	389	
% range*		Duplication	Error		Duplication	Error		Duplication	Error	
90-92%		135	0		137	0		117	0	
92-94%		334	0		334	0		311	0	
94-96%		391	0		382	0		367	0	
96-98%		451	0		444	0		418	0	
98-100%		884	1,097		724	932		665	389	

#### Comparison of duplications and potential sequence misassignment errors in genome assemblies

All numbers shown in the table are x 100 kb. \*Sequence similarity between duplication by five levels of percent identity.

Human Genome Browser [28]. All chromosome sequences were lower-case masked for highly repetitive elements by RepeatMasker (A.F.A. Smit and P. Green, unpublished). For each assembly build, each of the 25 masked chromosome sequences (including one unmapped chromosome sequence 'ChrUn') was compared against itself by chromosome-wide BLAST2 [9] to detect intrachromosomal segmental duplications (25 comparisons made), as well as pairwise comparisons to each of the other 24 chromosomes to detect interchromosomal segmental duplications (600 comparisons made). All BLAST results were subsequently parsed to eliminate low-quality and fragmented alignments under the

#### Table 4

# Segmental duplications involved in known genomic disorders and chromosome rearrangements identified by BLAST and ambSNP analyses

		First copy			Second copy(s)				
Disorders	Band	Start*	Size*	ambSNPs <sup>†</sup>	Start*	Size*	ambSNPs <sup>†</sup>	Identity	Celera C4‡
Gaucher disease	l q22	148108965	10,649	7	152776301	-10,479	10	95.19	S
Spinal muscular atrophy	5p14/5q13	21621854	79,183	1,032	69175603	-79,149	1,190	98.22	М
Williams-Beuren syndrome	7q11.23	70970126	359,416	380	72927299 73383317	,773 -227,260	56 355	99.60 99.20	P P
t(4;8)(p16;p23) Wolf-Hirschhorn syndrome	4p16/8p23	8769778	99,609	3†	7156209 7470072	-51,677 -82,189	18 387	95.65 95.81	P P
inv dup(8p) der(8)(8p23.1::p23.2-pter) del(8)(p23.1p23.2)	8p23.1	7084847	I 38,560	123	7756853 7651975	-126,769 54,807	229 463	99.16 96.93	M M
Prader-Willi syndrome and Angelman syndrome	5q  / 5q 3	19709020	75,325	102	19961243 20029574 20064937	34,902 41,965 74 780	55 83 65	98.70 98.79 99.01	P P P
Polycystic kidney disease I	16p13	2164789	38,034	136	16249164	24,076	243	98.32	P
Charcot-Marie-Tooth IA/Hereditary neuropathy with pressure palsies	7р 2/ 7р 2	14440158	23,599	272	15837032	23,585	286	98.42	Ρ
Smith-Magenis syndrome/ dup(17)(p11.2-p11.2)	17p12	18524425	152,700	547	20492073 25811482	-147,255 28,239	539 24	99.06 99.20	M M
Neurofibromatosis type I	17q11.2	28686414	63,356	163	28952984	-32,619	129	98.65	Р
DiGeorge syndrome and velocardiofacial syndrome	22q11.21	15662253	155,811	471	822 385  7742343  8 6437	155,996 9,740 -39,696	322 62 21	99.42 97.84 99.37	P P P
Chronic myeloid leukemia t(9;22)(pq34;q11)	9q34/ 22q11	123263651	36,956	NA	20552124	26,424	NA	91.81	S
Emery-Dreifuss muscular dystrophy	Xq28	147627873	11,030	2	147676529	11,034	2	99.61	S
Shwachman-Diamond syndrome	7q11.21	65091051	325,140	665	70647188	302,881	652	97.43	Р
Red green color blindness	Xq28	148439480	21,144	61	148476598	21,834	58	99.82	S
BRCA1 duplication	17q21	40983970	43,221	66	62252214	431,52	66	99.85	Р
Male infertility AZFc microdeletion region 2	Yq11.22 Yq11.22 Yq11.22	23322362 23908727 24794944	190,336 94,194 93,690	391§ 282§ 247§	23680552 24794944 27460935	-185,149 -93,690 -94,218	393§ 284§ 248§	99.88 99.92 99.93	P P P

This table represents a partial list of all known genomic disorders and chromosome rearrangements. \*Only the start coordinates (based on June 2002 assembly) for duplicons are shown. Results from BLAST analysis with chromosome coordinates and size of duplicon. For several genomic mutations (Williams-Beuren syndrome, Prader-Willi syndrome and Angelman syndromes, and DiGeorge syndrome) the duplicons shown are incomplete, most of which are composed of several duplication modules. The '-' sign indicates that the second duplicon is in the inverse orientation. <sup>†</sup>The number of ambSNPs (ambiguously mapped single-nucleotide polymorphisms) found within the genomic segment. NA, not applicable. The ambSNP analysis defines regions containing high densities of contiguous ambSNPs. For some of the segmental duplications involved in genomic disorders, the contiguous lengths of ambSNPs are much larger than those detected by BLAST. The specific sizes of the segmental duplications have to be resolved by detailed characterization of the different modules. <sup>‡</sup>Celera representation: S, both copies found in large (> 500 kb) sequence scaffolds; P, partially hit, single copy found, or less than perfect alignments; M, missing from large sequence scaffolds, hitting numerous fragments. <sup>§</sup>SNPs with multiple locations were used for evaluating the density of ambSNPs.

following criteria: BLAST results having  $\ge$  90% sequence identity,  $\ge$  80 bp in length, and with expected value  $\le$  10<sup>-30</sup>.

#### **BLAST** results parsing 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). This pairwise sequence comparison procedure serves as a rapid and robust way to detect duplication relationships. However, because of the use of masked sequences, our method would only yield a poor (on average 0.1-0.5 kb) resolution for the determination of the precise duplication alignment boundaries. Results were classified as either duplications or 'questionable' results based on sequencing status of the region and the percent sequence similarity between the detected alignments. Questionable duplications are results that fall within regions containing draft sequences with > 99.5 % detected sequence identity with another region. We consider these questionable duplications to be involved in potential sequence misassignment errors in the human genome assembly and would require further effort to achieve resolution.

#### Fine mapping of segmental duplications

Detailed information regarding segmental duplications as well as potential sequence misassignment errors identified by our analysis were presented using the Generic Genome Browser [11,12]. We have also summarized our results in table formats [12] that include information on size of duplications, chromosomal band locations, level of identity between duplicated copies, sequenced clones (accession numbers) and their sequencing status, as well as genes mapped to these regions. In addition, we have plotted the size of each intrachromosomal duplication (y-axis) against its chromosome position (x-axis) along each chromosome to indicate the intrachromosomal segmental duplication content of each chromosome (Figure 1) using the publicly available visualization tool GenomePixelizer [29,30]. Results generated from the detection of segmental duplications were subsequently converted into coordinate files as input for display using GenomePixelizer.

#### Paralogous sequence variants (PSV) density map

SNP mapping data from dbSNP were obtained through the NCBI ftp site [31]. Each chromosome SNP table, containing annotation regarding ambSNPs that have appeared twice in a particular chromosome, were extracted and sorted along with their corresponding chromosomal positions. The number of ambSNPs was tabulated along a 10-kb window to produce density plots of ambSNPs along the length of each chromosome (Figure 1).

### Additional data file

An additional data file with tables describing the relative frequency increase of repeat types at duplicon junctions and the number of ambiguousgly mapped SNPs within segmental duplications (number of PSVs), respectively, as well as a figure showing functional profiling of genes involved in recent whole-gene duplication vs human genome average is available with the online version of this article.

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