BMC Genomics



Research article Open Access

Cynomolgus monkey testicular cDNAs for discovery of novel human genes in the human genome sequence

Naoki Osada*^{1,2}, Munetomo Hida³, Jun Kusuda¹, Reiko Tanuma¹, Makoto Hirata¹, Yumiko Suto², Momoki Hirai², Keiji Terao⁴, Sumio Sugano³ and Katsuyuki Hashimoto¹

Address: ¹Division of Genetic Resources, National Institute of Infectious Diseases, 1-23-1 Toyama-cho, Shinjuku-ku, 162-8640, Japan, ²Laboratory of human evolution, Depertment of Integrated Biosciences, Graduate School of Frontier Sciences, The University of Tokyo, 5-1-5, Kashiwanoha, Kashiwa-shi, Chiba, 277-8562, Japan, ³Department of Genome Structure Analysis, Institute of Medical Science, The University of Tokyo, 4-6-1 Shirokanedai, Minato-ku, Tokyo 108-8639, Japan and ⁴Tsukuba Primate Center For Medical Science, National Institute of Infectious Diseases, Hachimandai-1, Tsukuba-shi, Ibaraki 305-0843, Japan

Email: Naoki Osada* - osada@nih.go.jp; Munetomo Hida - ss77233@ims.u-tokyo.ac.jp; Jun Kusuda - jkusuda@nih.go.jp; Reiko Tanuma - rtanuma@nih.go.jp; Makoto Hirata - mhirata@nih.go.jp; Yumiko Suto - suto@k.u-tokyo.ac.jp; Momoki Hirai - mhirai@k.u-tokyo.ac.jp; Keiji Terao - terao@nih.go.jp; Sumio Sugano - ssugano@ims.u-tokyo.ac.jp; Katsuyuki Hashimoto - khashi@nih.go.jp

* Corresponding author

Published: 23 December 2002 BMC Genomics 2002. 3:36 Received: 9 October 2002 Accepted: 23 December 2002

This article is available from: http://www.biomedcentral.com/1471-2164/3/36

© 2002 Osada et al; licensee BioMed Central Ltd. This is an Open Access article: verbatim copying and redistribution of this article are permitted in all media for any purpose, provided this notice is preserved along with the article's original URL.

Abstract

Background: In order to contribute to the establishment of a complete map of transcribed regions of the human genome, we constructed a testicular cDNA library for the cynomolgus monkey, and attempted to find novel transcripts for identification of their human homologues.

Result: The full-insert sequences of 512 cDNA clones were determined. Ultimately we found 302 non-redundant cDNAs carrying open reading frames of 300 bp-length or longer. Among them, 89 cDNAs were found not to be annotated previously in the Ensembl human database. After searching against the Ensembl mouse database, we also found 69 putative coding sequences have no homologous cDNAs in the annotated human and mouse genome sequences in Ensembl.

We subsequently designed a DNA microarray including 396 non-redundant cDNAs (with and without open reading frames) to examine the expression of the full-sequenced genes. With the testicular probe and a mixture of probes of 10 other tissues, 316 of 332 effective spots showed intense hybridized signals and 75 cDNAs were shown to be expressed very highly in the cynomolgus monkey testis, but not ubiquitously.

Conclusions: In this report, we determined 302 full-insert sequences of cynomolgus monkey cDNAs with enough length of open reading frames to discover novel transcripts as human homologues. Among 302 cDNA sequences, human homologues of 89 cDNAs have not been predicted in the annotated human genome sequence in the Ensembl. Additionally, we identified 75 dominantly expressed genes in testis among the full-sequenced clones by using a DNA microarray. Our cDNA clones and analytical results will be valuable resources for future functional genomic studies.

Background

Progress in genome biology has revealed the complete genome sequences of many non-mammalian species, such as yeast, nematodes, and the fruit fly. In addition, the much larger and more complicated genome sequences of the mouse and the human will soon be made completely available. However, decoding the genome sequences, especially the human sequence will be a long process. In order to achieve a comprehensive understanding of how an organism is established by its genome sequence, we must identify the structure, function, and interaction of as many genes as possible. First, we should accumulate and compile many types of evidence from computational and empirical data. The immediate challenge is establishing a complete map of transcribed regions in the human genome. Current comprehensive studies predicting proteincoding genes from the human genome [1,2] mainly employ three sources of information: empirical evidence provided by expressed sequence tags (ESTs) and cDNAs, nucleotide and protein sequence similarity to those of known genes, and statistical probability calculated by computer algorithms (ab initio prediction). All of these sources more or less lead to false-positive or false-negative types of errors. EST and cDNA sequences usually contain sequences that are not actually transcribed in vivo, i.e. artifacts arising from splicing intermediates, genomic DNA contamination, and transcription from nongenic regions [3,4]. Moreover, rarely expressed genes that may represent only a small portion of all transcripts cannot be easily represented in cDNA libraries. Predictions based on nucleotide and protein sequence similarities to those of other gene families and organisms might misassign pseudo genes, and cannot identify evolutionarily diverged genes that have no sequence similarity to known genes. Ab initio prediction works well for some organisms, such as yeast, nematodes, and the fruit fly. However, the human genome makes ab initio prediction of protein-coding genes difficult because it generally consists of small exons separated by long introns. Ultimately, in order to make a complete catalog of human genes, it will be necessary to gather undiscovered evidence from experiments and discard spurious evidence.

Our strategy for finding novel genes is to perform cDNA analysis using an organism closely related to humans, the cynomolgus monkey (*Macaca fascicularis*). In previous studies, we accumulated a number of 5'-end sequences of many clones derived from the oligo-capped cDNA libraries of the brain with high mRNA complexity, and determined approximately 1,500 full insert sequences of the clones whose 5'-end sequences showed no significant similarity to sequences in the public databases [5,6]. This method allowed us to identify many novel transcripts in the human genome sequence. Using fresh cynomolgus monkey tissues makes it possible to isolate rarely ex-

pressed genes, because mRNAs are so fragile that considerable portions of them degenerate during the usual construction of a cDNA library for humans. As an advantage of using cynomolgus monkey, evolutionary inspection can also provide information on gene function. If there are genes that evolved rapidly after the divergence of humans and cynomolgus monkeys, the function of the proteins and parts of the proteins might be important for human evolution. Moreover, biomedical interest in nonhuman primate genomes has been increased rapidly [7], especially in macaques, which also have been a material as transgenic primates [8], and thus genomic analysis of macaques will be important after the completion of human genome sequencing. In this study, we analyzed the cDNA library of the cynomolgus monkey testis. Analysis of testicular cDNA libraries has high potential for finding novel genes [9,10], because the testis is an organ in which transcripts have high complexity and where important biological processes, such as cell differentiation and meiosis, occur. The genes expressed in the testis are also important for medical, evolutionary, and developmental research. It is ironic that one of the most attractive tissues for biology expresses a number of undiscovered genes. We anticipated that analysis of the testicular cDNA library would lead to the discovery of novel genes that would facilitate post-genomic studies to attempt to unravel the complex genomes of higher organisms. Further, we conducted an expression analysis of our full-sequenced cD-NAs with cDNA microarray. DNA microarrays are a versatile tool for evaluating gene expression and sequence variation [11]. We used a cDNA microarray, to determine whether our putative genes were actually transcribed in cynomolgus monkey tissues and whether they were expressed dominantly in the cynomolgus monkey testis.

Results

We constructed the cDNA library derived from the cynomolgus monkey testis (library name: QtsA) by the oligo-capping method. The 5'-ends of 10,426 clones isolated from the library were sequenced and yielded 5,381 clusters of sequences (the redundancy rate was 1.94). To classify these cynomolgus monkey cDNAs and find their human homologues, we performed the BLAST search [12] to human RefSeq databases [13]. The 5'-end sequences of 6151 clones were found to have high similarity to 2321 human RefSeq genes with a cut-off value of e-60. The results showed that most of frequently occurring genes in cDNA library, QtsA were those specifically expressed in testis and sperm. Breakdown of the numerically represented genes is shown in Table 1. The clones whose 5'-end sequences had no homologies to the sequences in the nr and EST databases in the Genbank and had the possibility of being a certain length of ORF were subjected to full-insert sequencing. The entire sequences of 512 clones were determined as a result, but the total number of non-re-

Table 1: The list of 10 most frequently represented genes in the library QtsA.

Accession	No. of clones	Gene name (Gene symbol)
NM_002762	318	Protamine 2 (PRM2)
NM_004645	121	Coilin (COIL)
NM_004362	108	Calmegin (CLGN)
NM_005425	105	Transition protein 2 (TNP2)
NM_004396	95	DEAD/H box polypeptide 5 (DDX5)
NM_017769	83	Hypothetical protein FLJ20333 (FLJ20333)
NM_030941	80	Exonuclease NEF-sp (LOC81691)
NM_001402	77	Eukaryotic translation elongation factor alpha (EEFTAT)
NM_021009	76	Ubiquitin C (ÚBC)
NM_004724	63	ZW10 (Drosohplila) homolog (ZW10)

dundant transcripts was smaller because we could not completely exclude the 5'-truncated clones of the same transcripts at the stage of 5'-end sequences. Further, we might obtain some alternatively spliced transcripts from the same gene. In such cases, we used the longest transcripts in this study. Ultimately, we obtained a total of 394 non-redundant full insert cDNA sequences (Figure 1). After masking the common repetitive elements in the Repbase Update database [14], we assigned all cDNA sequences to the human genome draft sequence by using the BLAST program. With 85% or greater sequence identity and 50% or greater overlap of cDNA sequence length as criteria, 12 clones were deduced to be chimeric because they could be divided into two regions, of which DNA sequences showed homology to sequences of different human chromosomes. Sequences of 317 cDNAs had only one homologous region in the human genome sequences, while 18 cDNA sequences had homology to more than two human chromosomal regions. The remaining 47 had no homologous sequences in the human genome based on the above criteria. The average nucleotide length of all full-sequenced clones was 2016 bp. Of the 382 non-chimeric sequences, 302 carried a putative CDS (coding sequence) longer than or equal to 300 bp. In order to determine how many human homologues of our full-sequenced cDNAs have been annotated from the human genome sequences, a search was made for 302 putative CDSs to the Ensembl human database (Release 5.28.1) [15], which comprised 29,076 putative transcribed sequences classified as 'known' and 'novel' genes (BLAST cut-off value was e^{-60} , and the coverage was $\geq 50\%$ of each putative CDS length). Genes classified as 'known' in Ensembl are more reliable and have valid cDNA and/or evolutionary evidence, whereas 'novel' genes lack credible sources of expression and are sometimes supported by only ab initio methods and ESTs. As a result, 124 and 89 putative CDSs had human homologous sequences in the known category and novel category, respectively. The oth-

er 89 putative CDS had no homologous sequences in Emsembl human database based on these criteria. We also searched 302 cDNA sequences against the Ensembl mouse database (Release 7.3b.2), in which 28,097 putative transcribed sequences were annotated, (cut-off: e-30, coverage: ≥ 50% of ORF length), resulting that 74 and 67 cDNAs had homologous mouse sequences predicted as Ensembl 'known' and 'novel' genes, respectively. Finally, 69 putative CDSs have no homologous sequences in the annotated mammalian genome sequences in Ensembl. The putative functions of 302 hypothetical proteins were predicted by searching against the InterPro database [16]. A list of their name and other information on the 302 cD-NAs is provided in the supplementary table (additional file 1). We then constructed the putative human transcribed sequences corresponding to the 302 cynomolgus monkey cDNA sequences carrying enough length of ORFs by using the human genome draft sequences (see Materials and methods). The result showed that 117 putative human transcribed sequences, including 55 'known' and 48 'novel' genes in Ensembl had almost identical genomic structure to those of cynomolgus monkeys. We tested how many exons of 48 'novel' and 12 'unidentified' putative transcribed sequences can be predicted by the ab initio program, GENSCAN [17]. In total, 240 (53%) and 79 (17%) of 455 exons were correctly and partially predicted by GENSCAN, respectively, however, 136 exons (30%) were unpredictable. The list of putative human transcribed sequences is presented in Table 2 and their sequences are provided in the supplementary data (additional file 2), but the sequences have not been registered in the public database because they were not actually sequenced. We also compared the nucleotide and protein sequence similarity of 117 putative transcribed sequences between humans and cynomolgus monkeys. Amino acid sequence identity, nucleotide sequence identity for CDS, synonymous substitution per synonymous site, and non-

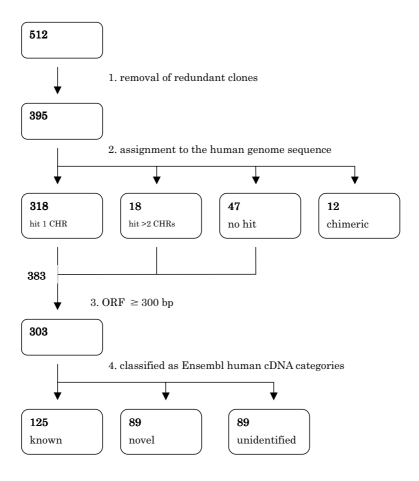


Figure I
Flow of full-sequencing analysis of unidentified clones. 1) The 512 full-sequenced clones were reduced to 394 because slightly different 5'-end sequences could be derived from the same transcripts. 2) 394 non-redundant clones were assigned to the human genome sequence. 3) 302 of 383 non-chimeric clones carried ORFs longer than 300 bp. 4) 302 putative genes were classified as 'known' and 'novel' categories of Ensembl human cDNA sequences. *CHR: Chromosome

synonymous substitution per non-synonymous site are presented in Table 2.

In order to investigate the expression pattern of the testicular full-sequenced cDNAs, we designed a DNA microarray containing approximately 400 spots of cDNA, full-sequenced samples and controls. Fifty clones carrying common repetitive elements and 12 clones deduced to be chimeric were excluded from further analysis, although they were spotted on the slides. Ultimately, 332 spots were used for quantification of gene expression. First, we investigated whether the putative genes were transcribed in a ubiquitous manner or had a tissue-specific pattern of

expression especially in the testis. RNA pools from the testis of the cynomolgus monkey and the mixture of equal amounts of RNA from 10 other cynomolgus monkey tissues (brain, heart, skin, liver, spleen, renal, pancreas, stomach, small intestine, and large intestine) were independently labeled and co-hybridized to the DNA microarray. When the signal intensity of the testicular probe is greater than that of the mixed probe, the gene was concluded to be over-expressed in the testis, or to be transcribed in the testis and a few other tissues, but not ubiquitously. When the intensity of both signals was equal, the gene was concluded to be expressed in a ubiquitous manner. When the signal intensity of the testicular

Table 2: The list of 117 putative human transcribed sequences.

Referenced macaca cDNA ^a	Ensembl status ^b	length	CDS length: startend (bp)	aa identity (%)	nt identity of CDS (%)	Ka ^c	Ks ^d
QtsA-10152	novel	1789	413AA: 421283	96.1	97.6	0.019	0.039
QtsA-10154	known	2010	502AA: 3771885	98.2	98.3	0.009	0.032
QtsA-10162	novel	2444	718AA: 722228	96.5	97.6	0.017	0.040
QtsA-10245	known	2598	752AA: 2982556	94.4	96.1	0.027	0.078
QtsA-10439	novel	2566	538AA: 2711887	88.6	93.7	0.059	0.076
QtsA-10472	known	2159	418AA: 761332	95.5	96.7	0.025	0.062
QtsA-10491	known	2439	346AA: 13222362	98.0	98.6	0.009	0.027
QtsA-10636	novel	2627	440AA: 571379	100.0	100.0	0.000	0.000
QtsA-10679	known	2415	523AA: 7262297	96.0	96.8	0.021	0.061
QtsA-10739	novel	1880	231AA: 141836	93.4	97.0	0.034	0.022
QtsA-10833	known	2234	673AA: 892110	92.2	95.2	0.037	0.077
QtsA-10891	novel	2049	343AA: 21033	93.2	95.2	0.038	0.084
QtsA-10947	known	2132	540AA: 1121734	93.2	94.4	0.033	0.126
QtsA-10963	known	1924	462AA: 4331821	98.9	98.6	0.005	0.039
QtsA-11068	unidentified	2299	594AA: 4052189	91.8	94.9	0.042	0.080
QtsA-11127	known	2084	550AA: 541706	89.8	95.5	0.053	0.034
QtsA-11181	known	3414	566AA: 841784	99.8	98.9	0.001	0.036
QtsA-11319	unidentified	1559	104AA: 168482	100.0	100.0	0.000	0.000
QtsA-11379	known	2805	690AA: 1062178	98.8	98.2	0.005	0.050
QtsA-11535	known	2116	474AA: 3041728	98.3	98.0	0.008	0.057
QtsA-11567	unidentified	1376	376AA: 2001330	96.0	97.7	0.020	0.034
QtsA-11570	unidentified	2437	117AA: 19022255	90.6	95.5	0.046	0.042
QtsA-11661	novel	2228	588AA: 2271993	97.6	98.0	0.013	0.038
QtsA-11670	unidentified	1785	325AA: 4131390	99.7	99.2	0.002	0.028
QtsA-11842	known	2173	225AA: 142819	100.0	100.0	0.000	0.000
QtsA-12007	novel	2316	724AA: 282202	96.6	97.1	0.017	0.053
QtsA-12095	novel	710	231AA: 16711	94.4	96.8	0.030	0.039
QtsA-12142	known	1731	404AA: 3951609	94.1	96.3	0.027	0.060
QtsA-12155	known	1305	329AA: 2521241	95.5	97.7	0.024	0.034
QtsA-12190	unidentified	1962	600AA: 211823	94.0	96.6	0.030	0.044
QtsA-12219	known	2480	793AA: 182399	100.0	100.0	0.000	0.000

Table 2: The list of 117 putative human transcribed sequences. (Continued)

Qish 12324 Nove 270 700Ak 97.1 97.9 0.013 0.036 0.035 0.045 0.045 0.068 0.045 0.068 0.	ubic 2. The list	or patative na		bea sequences (Cont.	nace,			
QtzA-12352 novel 2177 695AA: 91.2178 93.3 95.9 0.034 0.068 QtzA-12457 novel 245 689AA: 105.2174 94.9 96.5 0.026 0.059 QtzA-12579 novel 1499 491AA: 81.483 93.1 94.9 0.034 0.103 QtzA-12757 known 1280 278AA: 97.6 98.2 0.012 0.038 QtzA-12767 novel 2100 627AA: 98.6 98.0 0.007 0.060 QtzA-12769 known 1530 335AA: 92.2 95.8 0.038 0.052 QtzA-12850 known 2825 52.0 97.5 97.7 0.011 0.053 QtzA-12850 known 2826 2808 873AA: 92.2 95.8 0.038 0.052 QtzA-13252 novel 2229 669AA: 69.2 97.7 96.9 0.021 0.059 QtzA-13272 novel 1833 207AA: 98.1 98.1 96.6 0.010 0.033 QtzA-13327 <	QtsA-12282	novel	2270		97.1	97.9	0.013	0.036
Picka-12457 novel 2405 689AAk 94.9 96.5 0.026 0.059	QtsA-12354	known	2082	674AA: 52029	84.4		0.095	
Clizk-12579 novel 1499 491A-8. B.1483 93.1 94.9 0.034 0.103	QtsA-12362	novel	2177		93.3	95.9	0.034	0.068
QtsA-12649 novel 2114 634AAK: 33.1973 97.6 98.2 0.012 0.088 QtsA-12757 known 1280 278AAK: 98.6 97.7 0.008 0.055 QtsA-12767 novel 2100 622AAK: 98.6 98.0 0.007 0.060 QtsA-12769 known 1530 395AAK: 92.2 95.8 0.038 0.052 QtsA-12850 known 2825 854AAK: 97.5 97.7 0.011 0.053 QtsA-13222 known 2806 2806 873AK: 97.7 95.0 0.038 0.085 QtsA-13252 novel 229 669AAK: 95.7 95.7 96.9 0.021 0.059 QtsA-13272 novel 1833 207AAK: 98.1 98.6 0.010 0.033 QtsA-13343 novel 1960 131AAK: 99.6 96.2 0.041 0.026 QtsA-13406 known 1855 266AAK: 98.9 98.9 98.9 0.005 0.022 QtsA-13400 known	QtsA-12457	novel	2405		94.9	96.5	0.026	0.059
Cital-1275 Rinown 1280 278AA: 98.6 97.7 0.008 0.055	QtsA-12579	novel	1499	491AA: 81483	93.1	94.9	0.034	0.103
Care	QtsA-12649	novel	2114		97.6	98.2	0.012	0.038
QtsA-12767 novel 2100 622AA: 98.6 98.0 0.007 0.080 QtsA-12769 known 1510 395AA: 92.2 95.8 0.038 0.052 QtsA-12850 known 2825 854AA: 97.5 97.7 0.011 0.053 QtsA-13222 known 2806 2806 873AA: 92.7 95.0 0.038 0.085 QtsA-13252 novel 2229 669AA: 95.7 96.9 0.021 0.059 QtsA-13272 novel 1833 207AA: 98.1 98.6 0.010 0.033 QtsA-13343 novel 1960 131AA: 91.6 96.2 0.041 0.026 QtsA-13430 movel 1761 438AA: 91.6 98.9 98.9 0.005 0.022 QtsA-13406 known 1855 266AA: 92.0 96.2 0.039 0.040 QtsA-13452 known 1718 428AA: 97.7 98.1 0.011 0.038 QtsA-13450 known 1492 427AA: 95.0 97.2	QtsA-12757	known	1280	278AA:	98.6	97.7	0.008	0.055
T5.1262	QtsA-12767	novel	2100		98.6	98.0	0.007	0.060
Carta-1322	QtsA-12769	known	1530		92.2	95.8	0.038	0.052
Cash 13252 novel 2229 669An. 95.7 96.9 0.021 0.059 65.2074 95.7 96.9 0.021 0.059 65.2074 95.7 96.9 0.021 0.033	QtsA-12850	known	2825		97.5	97.7	0.011	0.053
Care 1833 1207AA: 1814 1815 1814 1815	QtsA-13222	known	2806		92.7	95.0	0.038	0.085
Tell	QtsA-13252	novel	2229		95.7	96.9	0.021	0.059
T71.566	QtsA-13272	novel	1833		98.1	98.6	0.010	0.033
QtsA-13406 known 1855 266AA: 26AA: 2	QtsA-13343	novel	1960		91.6	96.2	0.041	0.026
QtsA-13432 known 1718 428AA: 428A	QtsA-13392	unidentified	1761		98.9	98.9	0.005	0.022
Cach	QtsA-13406	known	1855		92.0	96.2	0.039	0.040
QtsA-13672 novel 1824 363AA: 734.1825 95.3 96.9 0.023 0.046 QtsA-13918 known 1730 537AA: 734.1825 97.2 98.1 0.012 0.047 QtsA-13925 novel 1678 515AA: 92.0 95.4 0.041 0.061 QtsA-14022 novel 1653 517AA: 96.3 96.8 0.020 0.064 QtsA-14166 novel 1121 293AA: 177.1058 99.7 0.052 0.060 QtsA-14351 novel 2938 839AA: 91.8 96.0 0.017 0.056 QtsA-14618 known 1273 363AA: 91.8 96.0 0.041 0.046 QtsA-14618 known 1273 363AA: 94.5 97.0 0.027 0.037 QtsA-14668 known 996 150AA: 405.857 97.0 0.027 0.037 QtsA-14766 known 2049 528AA: 96.8 97.2 0.016 0.060 QtsA-14816 unidentified 2515	QtsA-13432	known	1718		97.7	98.1	0.011	0.038
QtsA-13918 known 1730 537AA: 120.1733 97.2 98.1 0.012 0.047 QtsA-13925 novel 1678 \$15AA: 92.0 95.4 0.041 0.061 QtsA-14022 novel 1653 \$15AA: 102.1655 96.3 96.8 0.020 0.064 QtsA-14166 novel 1121 293AA: 89.5 94.7 0.052 0.060 QtsA-14245 known 1784 \$31AA: \$5.1600 96.6 97.2 0.017 0.056 QtsA-14351 novel 2938 839AA: 91.8 96.0 0.041 0.046 QtsA-14618 known 1273 363AA: 91.8 97.0 0.027 0.037 QtsA-14653 known 96 150AA: 100.0 98.7 0.000 0.049 QtsA-14766 known 2049 528AA: 96.8 97.2 0.016 0.060 QtsA-14752 known 2049 328AA: 91.8 96.8 97.2 0.016 0.060 QtsA-14816 unident	QtsA-13460	known	1492		95.0	97.2	0.023	0.035
Canal Cana	QtsA-13672	novel	1824		95.3	96.9	0.023	0.046
QtsA-14022 novel 1653 517AA: 1021655 96.3 96.8 0.020 0.064 QtsA-14166 novel 1121 293AA: 1771058 89.5 94.7 0.052 0.060 QtsA-14245 known 1784 531AA: 5.1600 96.6 97.2 0.017 0.056 QtsA-14351 novel 2938 839AA: 2257244 91.8 96.0 0.041 0.046 QtsA-14618 known 1273 363AA: 571148 94.5 97.0 0.027 0.037 QtsA-14653 known 996 150AA: 405.857 100.0 98.7 0.000 0.049 QtsA-14746 known 2049 528AA: 171603 96.8 97.2 0.016 0.060 QtsA-14752 known 904 235AA: 184891 97.5 97.0 0.012 0.078 QtsA-14816 unidentified 2515 134AA: 242646 98.4 99.2 0.008 0.013 QtsA-14970 known <td< td=""><td>QtsA-13918</td><td>known</td><td>1730</td><td>1201733</td><td>97.2</td><td>98.1</td><td>0.012</td><td>0.047</td></td<>	QtsA-13918	known	1730	1201733	97.2	98.1	0.012	0.047
Cart	QtsA-13925	novel	1678		92.0	95.4	0.041	0.061
1771058 QtsA-14245 known 1784 531AA: 51600 96.6 97.2 0.017 0.056 QtsA-14351 novel 2938 839AA: 91.8 96.0 0.041 0.046 QtsA-14618 known 1273 363AA: 94.5 97.0 0.027 0.037 QtsA-14653 known 996 150AA: 100.0 98.7 0.000 0.049 QtsA-14746 known 2049 528AA: 96.8 97.2 0.016 0.060 QtsA-14752 known 904 235AA: 97.5 97.0 0.012 0.078 QtsA-14816 unidentified 2515 134AA: 98.4 99.2 0.008 0.013 QtsA-14824 known 2965 891AA: 95.7 97.5 0.020 0.036 QtsA-14970 known 1282 168AA: 100.0 99.2 0.000 0.017 QtsA-15013 novel 2349 303AA: <td< td=""><td>QtsA-14022</td><td>novel</td><td>1653</td><td></td><td>96.3</td><td>96.8</td><td>0.020</td><td>0.064</td></td<>	QtsA-14022	novel	1653		96.3	96.8	0.020	0.064
QtsA-14351 novel 2938 839AA: 2252744 91.8 96.0 0.041 0.046 QtsA-14618 known 1273 363AA: 57148 94.5 97.0 0.027 0.037 QtsA-14653 known 996 150AA: 405857 100.0 98.7 0.000 0.049 QtsA-14746 known 2049 528AA: 96.8 97.2 0.016 0.060 QtsA-14752 known 904 235AA: 97.5 97.0 0.012 0.078 QtsA-14816 unidentified 2515 134AA: 98.4 99.2 0.008 0.013 QtsA-14824 known 2965 891AA: 95.7 97.5 0.020 0.036 QtsA-14970 known 1282 168AA: 6391145 100.0 99.2 0.000 0.017 QtsA-15013 novel 2349 303AA: 3641275 90.1 95.3 0.045 0.047 QtsA-15139 novel 2487 740AA: 96.4 96.4 96.9 0.017 0.061	QtsA-14166	novel	1121		89.5	94.7	0.052	0.060
2252744 QtsA-14618 known 1273 363AA: 571148 94.5 97.0 0.027 0.037 QtsA-14653 known 996 150AA: 405857 100.0 98.7 0.000 0.049 QtsA-14746 known 2049 528AA: 71.1603 96.8 97.2 0.016 0.060 QtsA-14752 known 904 235AA: 97.5 97.0 0.012 0.078 QtsA-14816 unidentified 2515 134AA: 98.4 99.2 0.008 0.013 QtsA-14824 known 2965 891AA: 95.7 97.5 0.020 0.036 QtsA-14970 known 1282 168AA: 6391145 100.0 99.2 0.000 0.017 QtsA-15013 novel 2349 303AA: 303AA: 90.1 95.3 0.045 0.047 QtsA-15139 novel 2487 740AA: 96.4 96.4 96.9 0.017 0.061	QtsA-14245	known	1784	531AA: 51600	96.6	97.2	0.017	0.056
57.1148 QtsA-14653 known 996 150AA: 405.857 100.0 98.7 0.000 0.049 QtsA-14746 known 2049 528AA: 71.1603 96.8 97.2 0.016 0.060 QtsA-14752 known 904 235AA: 81.891 97.5 97.0 0.012 0.078 QtsA-14816 unidentified 2515 134AA: 98.4 99.2 0.008 0.013 QtsA-14824 known 2965 891AA: 95.7 97.5 0.020 0.036 QtsA-14970 known 1282 168AA: 100.0 99.2 0.000 0.017 QtsA-15013 novel 2349 303AA: 90.1 95.3 0.045 0.047 QtsA-15139 novel 2487 740AA: 96.4 96.4 96.9 0.017 0.061								
405.857 QtsA-14746 known 2049 528AA: 96.8 97.2 0.016 0.060 171603 QtsA-14752 known 904 235AA: 97.5 97.0 0.012 0.078 184891 QtsA-14816 unidentified 2515 134AA: 98.4 99.2 0.008 0.013 242646 QtsA-14824 known 2965 891AA: 1512826 1512826 95.7 97.5 0.020 0.036 1512826 QtsA-14970 known 1282 168AA: 6391145 0.00 99.2 0.000 0.017 0.0017 0.047 0.047 0.041 QtsA-15013 novel 2349 303AA: 3641275 0.047 0.061 90.1 95.3 0.045 0.017 0.061 QtsA-15139 novel 2487 740AA: 96.4 96.9 0.017 0.017 0.061	QtsA-14618	known	1273		94.5	97.0	0.027	0.037
Tr. 1603 QtsA-14752 known 904 235AA: 97.5 97.0 0.012 0.078	QtsA-14653	known	996		100.0	98.7	0.000	0.049
184891 QtsA-14816 unidentified 2515 134AA: 242646 98.4 99.2 0.008 0.013 QtsA-14824 known 2965 891AA: 1511.2826 95.7 97.5 0.020 0.036 QtsA-14970 known 1282 168AA: 6391145 100.0 99.2 0.000 0.017 QtsA-15013 novel 2349 303AA: 3641275 90.1 95.3 0.045 0.047 QtsA-15139 novel 2487 740AA: 96.4 96.9 0.017 0.061	QtsA-14746	known			96.8	97.2	0.016	
242646 QtsA-14824 known 2965 891AA: 95.7 97.5 97.5 0.020 0.036 1512826 QtsA-14970 known 1282 168AA: 100.0 99.2 0.000 0.017 6391145 0.000 0.017 0.017 QtsA-15013 novel 2349 303AA: 3641275 0.047 3641275 90.1 95.3 0.045 0.047 0.061 QtsA-15139 novel 2487 740AA: 96.4 96.9 0.017 0.061	QtsA-14752	known	904		97.5	97.0	0.012	0.078
Cartest Cart	QtsA-14816	unidentified	2515		98.4	99.2	0.008	0.013
G39.1145 QtsA-15013 novel 2349 303AA: 90.1 95.3 0.045 0.047 3641275 QtsA-15139 novel 2487 740AA: 96.4 96.9 0.017 0.061	QtsA-14824	known	2965		95.7	97.5	0.020	0.036
3641275 QtsA-15139 novel 2487 740AA: 96.4 96.9 0.017 0.061	QtsA-14970	known	1282		100.0	99.2	0.000	0.017
		novel		3641275				
	QtsA-15139	novel	2487	740AA: 752297	96.4	96.9	0.017	0.061

Table 2: The list of 117 putative human transcribed sequences. (Continued)

rubic 2. The list	or its paracive in		sea sequences. (Cont.	ucc,			
QtsA-15186	known	2181	588AA: 971863	93.1	96.4	0.034	0.044
QtsA-15224	novel	2089	290AA: 3361208	96.1	96.8	0.021	0.074
QtsA-15268	novel	1808	396AA: 2031393	96.2	96.4	0.018	0.092
QtsA-15315	known	1284	344AA: 2171251	88.8	94.0	0.054	0.073
QtsA-15384	known	2169	565AA: 2131910	95.9	96.9	0.019	0.062
QtsA-15676	novel	2153	581AA: 1841929	92.1	95.8	0.039	0.068
QtsA-15696	novel	1856	563AA: 1441835	93.1	96.5	0.034	0.046
QtsA-15812	novel	2293	576AA: 4912221	96.4	97.2	0.019	0.052
QtsA-15844	known	2569	653AA: 1742135	95.6	96.1	0.023	0.086
QtsA-15875	known	2327	654AA: 3572321	96.6	97.7	0.017	0.038
QtsA-16005	known	1987	518AA: 4331989	100.0	99.4	0.000	0.021
QtsA-16015	known	2389	671AA: 3452360	98.1	97.7	0.009	0.054
QtsA-16028	known	1624	447AA: 231366	99.8	97.7	0.001	0.082
QtsA-16077	known	2307	571AA: 5762291	99.7	98.7	0.002	0.047
QtsA-16107	known	2039	432AA: 3011599	100.0	98.8	0.000	0.034
QtsA-16118	known	1396	415AA: 571304	97.6	96.7	0.012	0.096
QtsA-16284	novel	1199	342AA: 311059	96.5	96.8	0.017	0.071
QtsA-16373	known	2085	433AA: 6191920	99.5	98.5	0.002	0.048
QtsA-16429	known	1783	413AA: 421283	96.1	97.6	0.019	0.039
QtsA-16453	novel	1757	185AA: 7931350	87.3	93.5	0.066	0.065
QtsA-16496	novel	1599	448AA: 721418	95.5	96.1	0.023	0.081
QtsA-16602	unidentified	2482	263AA: 3151106	96.6	97.1	0.015	0.079
QtsA-16622	novel	1364	313AA: 2911232	95.1	96.9	0.024	0.045
QtsA-16678	known	2325	688AA: 912157	98.5	96.5	0.008	0.096
QtsA-16765	novel	2501	415AA: 8622109	98.1	97.4	0.010	0.068
QtsA-16837	known	3268	586AA: 8732633	99.3	98.5	0.004	0.041
QtsA-17449	known	1858	506AA: 2621782	90.9	95.4	0.044	0.053
QtsA-17495	novel	1026	261AA: 62847	96.2	97.3	0.018	0.068
QtsA-17616	known	2471	617AA: 4352288	98.4	98.2	0.008	0.044
QtsA-18070	novel	1997	585AA: 1341891	97.8	97.7	0.009	0.054
QtsA-18134	known	1832	309AA: 5921521	99.7	98.2	0.002	0.053
QtsA-18363	novel	1807	315AA: 6381585	95.9	97.5	0.020	0.040
QtsA-18372	novel	972	128AA: 337723	97.7	97.4	0.011	0.069

Table 2: The list of 117 putative human transcribed sequences. (Continued)

OtsA-18427	known	2198	565AA:	99.3	98.9	0.003	0.033
20,110127	Mom	2.70	4162113	,,,,	,	0.005	0.000
QtsA-18831	unidentified	2133	555AA: 471714	92.1	95.1	0.041	0.082
QtsA-18885	known	3250	642AA: 3142242	96.6	95.8	0.017	0.102
QtsA-19023	novel	2072	500AA: 841586	91.0	95.6	0.043	0.047
QtsA-19036	novel	955	214AA: 313957	100.0	99.5	0.000	0.014
QtsA-19380	unidentified	2158	412AA: 6251863	98.1	97.3	0.009	0.071
QtsA-19788	novel	1080	295AA: 1161003	98.6	98.3	0.006	0.040
QtsA-19856	known	2055	352AA: 4971555	98.9	98.4	0.005	0.039
QtsA-19961	known	1025	283AA: 62913	100.0	98.0	0.000	0.069
QtsA-20273	novel	1783	420AA: 791341	92.7	96.1	0.039	0.029
QtsA-20302	known	2889	882AA: 872735	94.6	97.2	0.026	0.042
QtsA-20424	unidentified	2056	505AA: 1471664	99.2	98.4	0.005	0.041
QtsA-20433	novel	1981	559AA: 731752	94.8	96.5	0.027	0.057
QtsA-20664	known	2396	616AA: 2312081	97.1	96.2	0.015	0.095
QtsA-20987	known	3090	561AA: 6362321	97.9	97.7	0.011	0.053
QtsA-21536	novel	1409	350AA: 1341186	92.3	95.7	0.042	0.052
QtsA-21565	novel	1810	367AA: 2761379	94.2	95.6	0.028	0.093
QtsA-21583	novel	2640	761AA: 2602545	90.4	95.0	0.046	0.060
QtsA-21585	known	2252	202AA: 38646	91.8	94.5	0.045	0.085

a) Cynomolgus monkey cDNA sequence that was used to deduce putative human transcribed sequence. b) Classification of human transcribed sequence in the Ensembl human database. c) Synonymous substitution rate per synonymous site between human and cynomolgus monkey genes. d) Non-synonymous substitution rate per non-synonymous site between human and cynomolgus monkey genes.

probe was lower than that of the mixed probe, the gene was concluded to be mainly transcribed in non-testicular tissues. We calculated the ratio of the testicular probe intensity to the mixed probe intensity and the ratio was normalized by using the beta-actin cDNA spot. A total of 316 (95%) of the 332 effective spots showed an intense and reproducible signal with either the testicular RNA probe or the mixed RNA probes or both. The signals of 75 spots were four fold or more intense with the testicular probe, and human homologues of the 15 genes among 75 cD-NAs had been registered in the RefSeq database (Table 3). Eight of the 15 RefSeq genes were reported to be expressed exclusively or dominantly in the human testis in the literature and the databases: TSGA10, expressed during spermatogenesis [18]; ACTL7B, an intronless gene strongly expressed in the testis and weakly expressed in the prostate [19]; SOX30, Sry-related transcriptional factor specifically expressed in the testis [20]; and five NYD-SP genes,

functionally anonymous but highly expressed in the testis in other DNA microarray experiments [21]. The other seven genes had ORFs of hypothetical proteins and were deduced from only the cDNA sequence evidence. Four of the cDNA clones were derived from human testis, and the other three cDNAs were from brain, placenta, or teratocarcinoma (Table 3). The results indicated that the remaining 60 clones that have no human RefSeq homologues are expressed exclusively or dominantly in the cynomolgus monkey testis.

Discussion

In this study we analyzed a cDNA library derived from a cynomolgus monkey testis. Although most of the human genome sequence has been determined, many unidentified genes remain, and a complete catalog of protein-coding genes is desired. Sequence similarity search of our full-sequenced cDNAs to the human draft genome sequence

Table 3: The list of genes that were highly expressed in a testis and had human RefSeq homologues

Macaca clone	Human RefSeq	Description	Ratio ^a	Expression (Reference)
QtsA-10833	NM_032559	kinesin protein (LOC84643)	8.7	derived from testis
QtsA-13647	NM_025244	testis specific, 10 (TSGA10)	8.6	testis specific [18]
QtsA-16118	NM_006686	actin-like 7B (ACTL7B)	8.5	testis and prostate [19]
QtsA-14409	NM_018418	hypothetical protein (HSD-3.1)	7.8	derived from testis
QtsA-13567	NM_033122	testis development protein NYD-SP26 (NYD-SP26)	7.5	testis
QtsA-14035	NM_033123	testis-development related NYD-SP27 (NYD-SP27)	7.2	testis
QtsA-11842	NM_032130	hypothetical protein DKFZp434J0113 (DKFZP434J0113)	6.9	derived from testis
QtsA-15256	NM_032126	hypothetical protein DKFZp564J047 (DKFZP564J047)	6.6	derived from brain
QtsA-14560	NM_032599	testes development-related NYD-SP18 (NYD-SP18)	6.6	testis
QtsA-12850	NM_019038	hypothetical protein (FLJ11045)	6.4	derived from placenta
QtsA-15384	NM_030672	hypothetical protein FLJ10312 (FLJ10312)	5.1	derived from teratocarcinoma
QtsA-10245	NM_007017	SRY (sex determining regionY)-box 30 (SOX30)	5.0	testis specific [20]
QtsA-18012	NM_032596	testes development-related NYD-SP22 (NYD-SP22)	5.0	testis
QtsA-14618	NM_032598	testes development-related NYD-SP20 (NYD-SP20)	4.7	testis
QtsA-19865	NM_033364	AATI-alpha (AATI) kinesin-like 6 (mitotic centromere-associated kinesin)	4.5	derived from testis
QtsA-16015	NM 006845	(KNSL6)	4 .1	thymus and testis [21]

a) The ratio of signal intensity of testicular probe to mixed probe.

resulted in the assignment of 347 cDNA sequences to at least one human chromosome, indicating that most genes in the cynomolgus monkey have homologous regions in the human genome. The primary objective of this analysis was to find genes that have not been experimentally identified in the human genome. Among the 302 cDNAs carrying enough length of ORFs (= 300 bp), we succeeded in identifying 89 putative genes that have no counterparts in the Ensembl 29,076-gene set. Another 89 genes that had highly similar sequences to Ensembl 'novel' genes were discovered in our full-sequenced cDNAs. The latter 89 genes strongly support the existence of predicted 'novel' cDNA sequences, which are relatively less accurate.

Many genes expressed in the testis cause male infertility in humans [22]. Since it is estimated that up to 11% of all genes in the fruit fly might lead to male sterility [23], in view of the complexity of the human genome, at least 4000 genes might be responsible for male infertility in humans and there must be many as yet unidentified genes that are related to male fertility. Functional analysis of 75 genes found to be highly expressed in the cynomolgus monkey testis may contribute such a medical interest about male infertility. A DNA microarray analysis is an appropriate method not only of annotating the pattern of expression of our full-length cDNAs, but of demonstrat-

ing that our strategy for finding novel gene works well. In the first set of the DNA microarray experiment, among the 199 genes that displayed two fold or more higher expression with the testicular probes than with the mixed probes, 67 (34%) were classified as the Ensembl 'known' genes, whereas among the 45 genes that showed ubiquitous pattern of expression (signal intensities within 1.5 fold of each other with both probes), 23 (51%) were classified as Ensembl 'known' genes. This finding indicated that the probability that transcripts overexpressed in testis are derived from unidentified novel genes is significantly higher than that of ubiquitous transcripts (p = 0.028: Fisher's exact test).

Evolutionary inspection is also important, especially for gene analysis of the testis, because genetic diversity in the male reproductive system is quite large, even among closely related species. Many reproductive proteins have evolved rapidly at the molecular level [24,25]. We compared 117 sequences of cynomolgus monkey cDNA and the corresponding human genome sequences described above, and use of the cDNA microarray revealed that 79 of the 117 cDNAs were overexpressed in testis (= 2.0 fold in testis) and 15 were ubiquitously expressed (within 1.5 fold of each other). We estimated the sequence divergence of putative coding sequences between humans and cy-

nomolgus monkeys and found that the average non-synonymous nucleotide divergence of testis-dominantly expressed genes (0.024) was significantly greater than that of ubiquitously expressed genes (0.012; p value < 0.01), whereas divergence in synonymous sites were not different significantly (testis-dominant genes: 0.54, ubiquitous genes: 0.51). This finding is also highly consistent with a report that the proteins of genes expressed in a tissue-specific manner evolve an average of twice as fast as those that are ubiquitously expressed [26].

Although a number of full and partial sequences of human genes have been deposited in the public databases, many of the genes in the human genome have not yet to be discovered experimentally. Most of the undiscovered genes may be expressed very seldom or their expression may be restricted to certain tissues and developmental stages. The complete human genome will be available in 2003, and a search of the entire genome for novel genes by oligonucleotide-based microarray analysis is designed; i.e. an attempt to predict all candidate human genes from the human genome and experimentally confirm the transcript status of the predicted regions as well as the entire region by using a oligo-nucleotide-based microarray [27,28]. However, it is difficult to overcome the problem of rarely or temporarily expressed genes for practical reasons. The transcriptional and genomic approaches will compensate for each other's blind spots, and many tissues, developmental stages, and other organisms should become experimental subjects for finding undiscovered genes to complete the human gene catalog.

Materials and Methods cDNA library from cynomolgus monkey testis

A 15-year-old male cynomolgus monkey was used as the source of the testis, and a 1-year-old and 21-year-old female cynomolgus monkeys were used for other RNA samples. The monkeys were cared for and handled according to guidelines established by the Institutional Animal Care and Use Committee of the National Institute of Infectious Diseases (NIID) of Japan and the standard operating procedures for monkeys at the Tsukuba Primate Center, NIID, Tsukuba, Ibaraki, Japan. Tissues were excised in accordance with all guidelines in the Laboratory Biosafety Manual, World Health Organization, and were carried out at the P3 facility for monkeys of the Tsukuba Primate Center, NIID. Immediately after collection, the tissues were frozen with liquid nitrogen and used for RNA extraction. Oligocapped cDNA libraries were constructed according to the method described previously [29,30].

DNA sequencing

The 5'-end sequences of the clones were sequenced using ABI 3700 sequencer (Applied Biosystems), and categorized using DYNACLUST (DYNACOM), based on a BLAST

search against the GenBank database. The entire sequences of clones were determined by the primer walking method. Cycle sequencing was performed with an ABI PRISM BigDye Terminator Sequencing kit (Applied Biosystems) according to the manufacturer's instructions.

Computational analyses

The Sim4 program was used to align each cynomolgus monkey cDNA sequence with the human genome sequence [31]. Whenever Sim4 failed to align cynomolgus monkey cDNA sequence with human genome DNA sequence, comparison by BLAST program was executed, and the alignment was corrected manually. In the intron sequences, GT at the 5' splice site and AG at the 3' site (GT-AG pattern), and the GC-AG pattern were regarded as conserved splice sites, and corresponding human genome regions were concatenated to construct a hypothetical human transcribed sequence. 117 Cynomolgus monkey cDNA sequences and the putative human transcribed sequences were aligned by using the ClustalW program [32]. Synonymous substitution per synonymous site and nonsynonymous substitution per non-synonymous site were estimated by the method of Li [33].

cDNA microarray

An aliquot of the same DNA preparation used in the 5'end-sequencing reactions provided material for the PCRs. Inserts were amplified by PCR using 5'-CTTCT-GCTCTAAAAGCTGCG-3' as a forward primer and 5'-CGACCTGCAGCTCGAGCACA-3' as a reverse primer, in a volume of 100 µl. Successful amplification was confirmed by agarose gel electrophoresis. When the first PCR failed to amplify enough products, the first PCR products were amplified again. Four hundred cDNA clones were amplified and samples of approximately 300 µg/ml DNA in 2 × Solution-T reagent (Takara Bio) were printed on duplicate glass-slides with a GMS 417 arrayer (Genetic MicroSystems). The testicular RNA was obtained from only one 15-year-old male cynomolgus monkey, and the other RNA was a mixture of RNA obtained from 10 tissues (brain, heart, skin, liver, spleen, renal, pancreas, stomach, small intestine, and large intestine) of two cynomolgus monkeys, a 1-year-old female and a 21-year-old female. RNA was isolated with Trizol (Life Technologies) and purified with Oligo-Tex (Takara Bio). Both 0.7 µg mRNA probes were labeled with Cy3- and Cy5- dioxynucleotide (Pharmacia) and co-hybridized to DNA spots. The amount of RNA from each tissue was 0.07 µg in the mixed RNA probe. After the hybridization and washing procedure, slides were scanned with ScanArray (GSI Inc.). Several experiments were conducted, and the duplicated spots on the slides, where the most intense signals were obtained, were processed to measure the transcriptional status. When the relative intensity of Cy3/Cy5 signals of duplicated spots differed more than 1.5 times compared

to that of the corresponding spots in duplicate, the spots were not processed for the subsequent analyses. Finally, the ratio of signal intensities of Cy3 (the testicular probe) and Cy5 (the mixed probe) was obtained from average value of duplicated spots and normalized by dividing by the ratio of the beta-action spots as a control.

List of abbreviations

EST: expressed sequence tag.

CDS: coding sequence.

ORF: open reading frame.

Authors' contributions

NO was involved in design of the study, construction of cDNA library, in silico analysis, expression analysis with DNA microarray and preparation of the manuscript. M. Hida and SS performed construction of cDNA libraries and analysis of 5'-end sequence analysis. JK participated in the design and implementation of the study, contributed to writing and revising the manuscript. RT and M. Hirata participated in the sequencing of cDNAs and in-silico analyis of cDNA sequences. YS and M. Hirai participated in the design and implementation of the study on microarray, and obtained funding for the study. KT contributed to obtaining the tissues for cDNA libraries and total RNA from cynomolgus monkeys. KH was involved in the design and implementation of the study, writing and editing the manuscript and obtained funding for the study.

All authors read and approved the final manuscript.

Additional material

Additional file 1

Click here for file [http://www.biomedcentral.com/content/supplementary/1471-2164-3-36-S1.htm]

Additional file 2

Click here for file [http://www.biomedcentral.com/content/supplementary/1471-2164-3-36-S2.txt]

Acknowledgements

This study was supported in part by the Health Science Research grant for the Human Genome Program from the Ministry of Health, Labor and Welfare of Japan.

References

- International Human Genome Sequencing Consortium: Initial sequencing and analysis of the human genome. Nature 2001. 409:860-921
- Venter JC, Adams MD, Myers EW, Li PW, Mural RJ, Sutton GG, Smith HO, Yandell M, Evans CA and Holt RA The sequence of the human genome. Science 2001, 291:1304-1351
- Adams MD, Kerlavage AR, Fleischmann RD, Fuldner RA, Bult CJ, Lee NH, Kirkness EF, Weinstock KG, Gocayne JD and White O Initial assessment of human gene diversity and expression patterns based upon 83 million nucleotides of cDNA sequence. Nature 1995, 377(Supplement):3-17
- Hillier LD, Lennon G, Becker M, Bonaldo MF, Chiapelli B, Chissoe S, Dietrich N, DuBuque T, Favello A and Gish W Generation and analysis of 280,000 human expressed sequence tags. Genome Res 1996, 6:807-828
- Osada N, Hida M, Kusuda J, Tanuma R, Iseki K, Hirata M, Suto Y, Hirai M, Terao K and Suzuki Y Assignment of 118 novel cDNAs of cynomolgus monkey brain to human chromosomes. Gene 2001, 275:31-37
- Osada N, Hida M, Kusuda J, Tanuma R, Hirata M, Hirai M, Terao K, Suzuki Y, Sugano S and Hashimoto K Prediction of unidentified human genes on the basis of sequence similarity to novel cD-NAs from cynomolgus monkey brain. Genome Biol 2002, 3:research0006.1-0006.5
- Eichler EE and Dejong PJ Biomedical applications and studies of molecular evolution: A proporsal for a primate genomic library resource. Genome Res 2002, 12:673-678
- Chan AWS, Chong KY, Martinovich C, Simerly C and Schatten G Transgenic monkeys produced by retroviral gene transfer into mature oocytes. Science 2001, 291:309-312
- Andrews J, Bouffard CG, Cheadle C, Lu Jining, Becker KG and Oliver B Gene discovery using computational and microarray analysis of transcription in the Drosophila melanogaster tesis. Genome Res 2000, 10:2030-2043
- Wiemann S, Weil B, Wellenreuther R, Gassenhuber J, Glassl S, Ansorge W, Bocher M, Blocker H, Bauersachs S and Blum H Toward a catalog of human genes and proteins: sequencing and analysis of 500 novel complete protein coding human cDNAs. Genome Res 2001, 11:422-435
- Nooedewier MO and Warren PV Gene expression microarrays and the integration of biological knowledge.: Trends Biotecnol 2001, 19:412-415
- Altschul SF, Madden TL, Schaffer AA, Zhang J, Zhang Z, Miller W and Lipman DJ Gapped BLAST and PSI-BLAST: a new generation of protein database search programs. Nuc Acids Res 1997, 25:3389-3402
- Pruitt KD and Maglott DR RefSeq and LocusLink: NCBI genecentered resources. Nucleic Acids Res 2001, 29:137-140
- Jurka J Repbase Update: a database and an electronic journal of repetitive elements. Trends Genet 2000, 9:418-420
- Hubbard T, Barker D, Birney E, Cameron G, Chen Y, Clark L, Cox T, Cuff J, Curwen V and Down T The Ensembl genome database project. Nuc Acid Res 2002, 30:38-41
- Apweiler R, Attwood TK, Bairoch A, Bateman A, Birney E, Biswas M, Bucher P, Cerutti L, Corpet F and Croning MD The InterPro database, an integrated documentation resource for protein families, domains and functional sites. Nucleic Acids Res 2001, 29:37-40
- Burge C and Karlin S Prediction of complete gene structures in human genomic DNA. J Mol Biol 1997, 268:78-94
- Modarressi MH, Cameron J, Taylor KE and Wolfe J Identification and characterisation of a novel gene, TSGA10, expressed in testis. Gene 2001, 262:249-255
- Chadwick BP, Mull J, Helbling LA, Gill S, Leyne M, Robbins CM, Pinkett HW, Makalowska I, Maayan C and Blumenfeld A Cloning, mapping, and expression of two novel actin genes, actin-like-7A (ACTL7A) and actin-like-7B (ACTL7B), from the familial dystonomia candidate region on 9q31. Genomics 1999, 58:302-209
- Osaki E, Nishina Y, Inazawa J, Copeland NG, Gilbert DJ, Jenkins NA, Ohsugi M, Tezuka T, Yoshida M and Semba K Identification of a novel Sry-related gene and its germ cell-specific expression. Nuc Acid Res 1999, 27:2503-2510
- 21. Jiahao S, Zuomin ZH and Jianmin L Preparation of human testicular cDNA microarray and initial research of gene expres-

- sion library related to spermatogenesis. In Epithelial Cell Biology-A primer (Edited by: Chan HS) Beijing 2000, 274-277
- Okabe M, Ikawa M and Ashkenas J Male infertility and the genetics of spermatogenesis. Am J Hum Genet 1998, 62:1274-1281
- Hackstein JHP, Hochstenbach R and Pearson PL Towards an understanding of the genetics of human make infertility: lessons from flies. Trends Genet 2000, 16:565-572
- Wyckoff GJ, Wang W and Wu Cl Rapid evolution of male reproductive genes in the decent of man. Nature 2000, 403:304-309
- Swanson WJ, Clark AG, Waldrip-Dail HM, Wolfner MF and Aquadro CF Evolutionary EST analysis identifies rapidly evolving make reproductive proteins in Drosophila. Proc Natl Acad Sci USA 2001, 98:7375-7379
- Duret L and Mouchiroud D Determinants of substitution rates in mammalian genes: expression pattern affects selection intensity but not mutation rate. Mol Biol Evol 2000, 17:68-74
 Shoemaker DD, Schadt EE, Armour CD, He YD, Garrett-Engele P,
- Shoemaker DD, Schadt EE, Armour CD, He YD, Garrett-Engele P, McDonagh PD, Loerch PM, Leonardson A, Lum PY and Cavet G Experimental annotation of the human genome using microarray technology. Nature 2001, 409:922-927
- Kapranov P, Cawley SE, Drenkow J, Bekiranov S, Strausberg RL, Fodor SPA and Gingeras TA Large-Scale Transcriptional Activity in Chromosomes 21 and 22. Science 2002, 296:916-919
- Suzuki Y, Ishihara D, Sasaki M, Nakagawa H, Hata H, Tsunoda T, Watanabe M, Komatsu T, Ota T and Isogai T Statistical analysis of the 5' untranslated region of human mRNA using "Oligo-Capped" cDNA libraries. Genomics 2000, 64:286-297
- Hida M, Suzuki Y, Sugano S, Hashimoto K, Terao K, Hayasaka I and Hirai M Construction and preliminary characterization of full length enriched cDNA libraries for nonhuman primates. Primate Res 2000, 16:95-110
- Florea L, Hartzell G, Zhang Z, Rubin GM and Miller W A computer program for aligning a cDNA sequence with a genomic DNA sequence. Genome Res 1998, 8:967-974
- Thompson JD, Higgins DG and Gibson TJ CLUSTAL W: improving the sensitivity of progressive multiple sequence alignment through sequence weighting, position-specific gap penalties and weight matrix choice. Nucleic Acids Res 1994, 22:4673-4680
- 33. Li WH Unbiased estimation of the rates of synonymous and nonsynonymous substitution. J Mol Evol 1993, 36:96-99

Publish with **Bio Med Central** and every scientist can read your work free of charge

"BioMed Central will be the most significant development for disseminating the results of biomedical research in our lifetime."

Sir Paul Nurse, Cancer Research UK

Your research papers will be:

- available free of charge to the entire biomedical community
- peer reviewed and published immediately upon acceptance
- cited in PubMed and archived on PubMed Central
- yours you keep the copyright

Submit your manuscript here: http://www.biomedcentral.com/info/publishing_adv.asp

