# Article

# Cytokine-Related Genes Identified From the RIKEN Full-Length Mouse cDNA Data Set

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To identify novel cytokine-related genes, we searched the set of 60,770 annotated RIKEN mouse cDNA clones (FANTOM2 clones), using keywords such as cytokine itself or cytokine names (such as interferon, interleukin, epidermal growth factor, fibroblast growth factor, and transforming growth factor). This search produced 108 known cytokines and cytokine-related products such as cytokine receptors, cytokine-associated genes, or their products (enhancers, accessory proteins, cytokine-induced genes). We found 15 clusters of FANTOM2 clones that are candidates for novel cytokine-related genes. These encoded products with strong sequence similarity to guanylate-binding protein (GBP-5), interleukin-1 receptor-associated kinase 2 (IRAK-2), interleukin 20 receptor  $\alpha$  isoform 3, a member of the interferon-inducible proteins, one p27-like protein, and a hypothetical protein containing a Toll/Interleukin receptor domain. All four clones representing novel candidates of gene products from the family contain a novel highly conserved cross-species domain. Clones similar to growth factor-related products included transforming growth factor  $\beta$ -inducible early growth response protein 2 (TIEG-2), TGF $\beta$ -induced factor 2, integrin  $\beta$ -like 1, latent TGF-binding protein 4S, and FGF receptor 4B. We performed a detailed sequence analysis of the candidate novel genes to elucidate their likely functional properties.

[Supplemental material is available online at www.genome.org]

Cytokines are polypeptides secreted by immune cells that function as humoral regulators modulating functional activities of cells and tissues including cellular interactions and processes. Cytokine genes often encode functional variants through alternative splicing, resulting in proteins that may have slightly different biological activities (Ibelgaufts 1999). Cytokines are typically mutually unrelated, but some may be grouped in families.

Activated cytokine-producing cells often release many different cytokines, resulting in complex networks of interacting signals. Cytokines are crucial in immune cell activation, cell-to-cell communication, signal transduction, mitosis, cell survival, death, and transformation. In addition to their involvement in initiation and regulation of immune responses, cytokines are also important in embryogenesis, organ development, and neuronal processes (Ibelgaufts 1999). Clinical uses of cytokines include application in hematopoiesis (Smith 1990), tumor immunotherapies (Cao et al. 1998; Rosenberg 2001; Homey et al. 2002), bone marrow transplan-

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tation (Miyazaki and Kato 1999), gene therapy (Jeschke et al. 2001), and vaccine formulations (Knutson and Disis 2001).

The function of a particular cytokine can be assessed in vivo in transgenic or gene knock-out animals by studying the effects of an excess or lack of a particular cytokine gene product. It is of great importance for cytokine research, therefore, that the counterparts of human cytokine, cytokine receptors, and cytokine-related genes are identified in animals (e.g., the mouse) suitable for genetic manipulation. Because of the remarkable similarity between human and mouse genomes and their closely related biochemical, physiological, and pathological pathways, the mouse is an extremely useful animal model for immunological studies.

The FANTOM2 collaboration (Okazaki et al. 2002) involved functional annotation of a set of 60,770 RIKEN mouse cDNA clones (FANTOM2 clones). The FANTOM2 data set, the most complete picture of the mouse transcriptome to date, provides an ideal opportunity for identification of novel mouse genes, including cytokine-related genes. The annotation process included data from microarray expression, mapping and protein interactions, and sequence similarity search analyses. The annotation was performed by large-scale, computerized annotation combined with manual curation by human experts. We analyzed the FANTOM2 annotations, specifically looking for novel and existing mouse cytokines and

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cytokine-related genes. In this study, we focused on genes related to interferons (IFN), interleukins (IL), growth factors (epidermal growth factors [EGF], fibroblast growth factors [FGF], and transforming growth factors [TGF]), and small inducible cytokines (SIC). Some cytokines, such as tumor necrosis factors (TNF), belong to superfamilies that also include non-cytokines. Because this study was based on matching cytokine names from descriptions of database entries, cytokine families that belong to broader superfamilies were excluded from the analysis. Further study is required for a comprehensive analysis of all known cytokines and cytokine-related genes that have not been included in this work. The FANTOM2 clones contained more than 100 previously identified cytokine and cytokine-related genes. We also identified several candidates for novel mouse cytokine-related genes.

#### RESULTS

Of the 60,770 FANTOM2 clones, 256 had annotations that contained a cytokine name. A total of 222 clones had highsequence identity (>96% over 50% length of the reference sequence) to 108 known mouse genes, as listed in the major databases. Of these, clones represented 27 known interferonrelated genes (Table 1). The known mouse genes in the FANTOM2 set comprised 1 interferon (IFN- $\gamma$ ), 2 IFN receptors (IFN $\alpha\beta$  receptor, and IFN $\alpha\beta$  receptor 2), 20 inducible/ activatable genes, and 4 other IFN-related genes. Search of the protein translation of the GenBank database (Benson et al. 2002, as of September 2002) revealed that there were at least 15 nonredundant entries of various mouse IFNs, 10 IFN receptors, 190 IFN inducible or activatable proteins, and 60 IFN regulatory proteins.

A total of 77 clones represented 39 known interleukin-related genes (Table 2). The known mouse genes in the FANTOM2 set comprised 11 interleukins (IL) or chains thereof (IL-18, IL-1ε, IL-6, IL-7, IL-1F9, IL-16, IL-17B, IL-20, IL-23p19, and IL-25), 19 interleukin receptors (ILR-11, ILR-1-1, ILR-2a, ILR-2 $\beta$ , ILR-2 $\gamma$ , ILR-4 $\alpha$ , ILR-6 $\alpha$ , ILR-7, ILR-9, ILR-10 $\beta$ , 12β1, ILR-12β2, ILR-13α1, ILR-13α2, ILR-15α, ILR-17, ILR-18-1, ILR-21, and ILR-23αp19), and 9 other IL-related genes. Search of the protein translation of the GenBank database revealed that there are at least 40 nonredundant entries of mouse ILs (including variants and subunits), 50 IL receptors, and 80 other IL-related proteins. These results are in agreement with an earlier finding (Staudt and Brown 2000), which stated that only a half of human ILs, mainly those widely expressed, were found in the EST libraries. The ILs reliant on the activation of immune cells, namely IL-2, IL-3, IL-5, IL-9, IL-11, IL-12β, IL-14, and IL-17, were lacking in the EST libraries.

A total of 61 clones represented 23 known growth factor (EGF, FGF, and TGF)-related genes (Table 3). The known mouse genes in the FANTOM2 set comprised 5 growth factors (EGF, FGF 14, FGF 15, TGF $\beta$ , and TGF $\beta$  68 kD) and 18 other growth factor-related genes. Search of the protein translation of the GenBank database revealed that there are at least 140 nonredundant entries of various mouse growth factor (EGF, FGF, and TGF)-related genes (including variants and sub-units).

Finally, 30 clones represented 19 known small inducible cytokine-related genes in mouse (Table 4). Search of the protein translation of the GenBank database revealed that there are at least 38 nonredundant entries of various small inducible cytokines in mouse.

Clones representing 17 known mouse cytokine-related genes were derived from cDNA libraries from the non-obese diabetic (NOD) mouse. All of the remaining cytokine-related clones were derived from the C57BL/6 mouse strain. Six known genes were only found from the NOD mouse libraries. Clones representing a total of 11 known mouse genes were found in both C57BL/6 and NOD mouse cDNA libraries. The clones representing known cytokine-related genes were derived from a broad variety of RIKEN cDNA libraries. These libraries were generated from various tissues and developmental stages. Because this analysis focused on name matching from both the FANTOM2 data set and GenBank database, we can estimate that FANTOM2 clones cover ~20% of existing known cytokine-related genes. Cytokines induce cascades of cytokine networks as well as of regulatory molecules. Often it is unclear whether the induced molecule acts as a cytokine or has some other regulatory role. In this study, we consider all clones as cytokine related and did not discriminate between the two groups.

We have also identified 15 candidate novel cytokine or cytokine-related genes (Table 5), including 7 candidate novel mouse IFN-related genes (13 clones), 3 candidates for novel IL-related genes (5 clones), and 4 candidates for novel growth factor-related genes (3 TGF- and 1 EGF-related) (16 clones). All of the clones representing candidate novel genes were derived from the C57BL/6 mouse and showed  $\leq$ 96% identity over >50% length of the reference sequence.

#### Interferon-related Novel Gene Candidates

The clones 1110004C05, 4930507H06, A330075D07, and 1110036C17 represent four putative novel members of the membrane-associated family 1-8 of interferon-inducible proteins (Fig. 1). Sequences of these clones showed similarity to a variety of interferon-induced genes and proteins in human (Lewin et al. 1991), rat (Hayzer et al. 1992), cattle (Pru et al. 2001), and trout (SPTR Q8QFL3). Members of the 1-8 family of interferon (IFN)-inducible genes encode proteins thought to modulate cellular growth and adhesion wound healing, and inflammatory responses. It was hypothesized that they may function as membrane transport proteins (Pru et al. 2001). The 1-8 proteins participate in multimeric complexes involved in the transduction of homotypic and antiproliferative adhesion signals. Sequences of all four clones of novel candidate members of the 1-8 family contain a novel (not found in motif databases), motif DxxxWSxxxxxNxx-CLGxxAxxxSxKSRDRKMVGDxxGAxxxxTA that is conserved across species (fish, rat, mouse, and human). Furthermore, sequence of the clone 1110004C05 (Fig. 1) contains the motif YxxxxCLxI that conserves critical residues found in the active sites of human E2 ubiquitin conjugating enzymes (Pru et al. 2001). Sequences of the clones 4930507H06 and 1110036C17 contain similar motif FxxxxCLxI, whereas the remaining clone A330075D07 does not contain the corresponding motif. Each of the four candidate new members of the 1-8 family showed highest similarity to a distinct database entry and therefore each represents a distinct transcript (data not shown). The amino acid differences between the members of the 1-8 family at carboxy- and amino-termini may contribute to subtle functional differences or cellular localization.

Sequence of the clone 2310061N23 showed similarity to the gene for the IFN- $\alpha$ -inducible protein p27-h found in hamster cardiomyopathic tissue (Denovan-Wright et al. 2000) and human gene TLH29 (Q9H2X8) from liver tissue (Fig. 2). The

	Clone ID	Mouse strain	cDNA library	Gene name				
I	5830458K16	C57BL/6	Thymus	28-kD interferon $\alpha$ responsive protein				
	1810013G24	C57BL/6	Pancreas	(5830458K16RIK protein) homolog ( <i>Mus musculus</i> )				
	0710005K22	C57BL/6	Brain	Genes associated with retinoid-IFN-induced mortality 19				
	0710005K22	C57BL/6	Brain	Genes associated with realional invalidation induced mortality 15				
	2700054G14	C57BL/6	Embryo11					
			·					
	2410016H22	C57BL/6	ES	laterform ( and 0) meanter				
	A430075E22	C57BL/6	N0 Thymus	Interferon ( $\alpha$ and $\beta$ ) receptor				
	1100001K09	C57BL/6	18-day embryo	Interferon ( $\alpha$ and $\beta$ ) receptor 2				
	2510009P07	C57BL/6	Embryo 13-Liver					
	E430025C02	NOD	(NOD)N2 Thymic cells	Interferon-activated gene 202A (noncoding RNA)				
	A730048F03	C57BL/6	N7 Cerebellum	Interferon-activated gene 205				
'	A430077M01	C57BL/6	N0 Thymus	Interferon-inducible protein 203 (IFI-203) (interferon-inducible protein P203)				
	A530058P16	C57BL/6	Aorta and vein					
	A530093G11	C57BL/6	Aorta and vein					
	E430077M01	NOD	(NOD)N2 Thymic					
	9030603A05	C57BL/6	Colon	Interferon concensus sequence-binding protein				
	9830117K07	C57BL/6	Bone					
)	A830021G01	C57BL/6	N10 Cortex	Interferon-dependent positive-acting transcription factor 3 $\gamma$				
)	F830002I10	NOD	Activated spleen from NOD.Cz	Interferon $\gamma$				
·	F830032O20	NOD	Activated spleen from NOD.Cz					
	0610039K12	C57BL/6	Kidney	Interferon $\gamma$ induced GTPase				
	E430012H16	NOD	(NOD)N2 Thymic cells	Interieron y induced offase				
				laterform indusible metric 20				
	5730557A06	C57BL/6	Embryo 8	Interferon $\gamma$ inducible protein 30				
	2010205D02	C57BL/6	Small intestine	Interferon $\gamma$ inducing factor-binding protein				
	1110003K24	C57BL/6	18-day embryo					
	2310040F23	C57BL/6	Tongue					
	2310075112	C57BL/6	Tongue					
	2310047H18	C57BL/6	Tongue					
ł	0610011H23	C57BL/6	Kidney	Interferon inducible protein 1				
	9830146E22	C57BL/6	Bone	Interferon regulatory factor 2				
	E430001A19	NOD	(NOD)N2 Thymic					
,	F730005C13	C57BL/6	B6 derived	Interferon regulatory factor 4				
	F830028K14	NOD	Activated spleen	5 ,				
,	A130064P06	C57BL/6	N16 Thymus	Interferon regulatory factor 6				
	E230028105	C57BL/6	P2 oviduct					
;	0610039E12	C57BL/6	Kidney	Interferon regulatory factor 7				
	1110062H15	C57BL/6	18-day embryo	interior regulatory factory				
	A430014N21	C57BL/6	N0 Thymus					
	F830035B02	NOD	Activated spleen					
,				Interferen induced 25 kD protein homolog (ICD 25) homolog				
	2010008K16	C57BL/6	Small intestine	Interferon-induced 35-kD protein homolog (IFP 35) homolog ( <i>Mus musculus</i> )				
)	2010002M12	C57BL/6	Small intestine	Interferon-induced protein with tetratricopeptide repeats 1				
	F830004H17	NOD	Activated spleen					
	A130072L05	C57BL/6	N16 Thymus	Interferon-induced protein with tetratricopeptide repeats 2				
	A630085F01	C57BL/6	N3 Thymus					
2	5730412C13	C57BL/6	Embryo 8	Interferon-induced protein with tetratricopeptide repeats 3				
	5031412D17	C57BL/6	Ovary & Uterus (Preg. 11 day)	(IFIT-3) glucocorticoid-attenuated response gene 49				
	F830009I11	NOD	Activated spleen	interferon-inducible GTPase				
ļ	A430077M01	C57BL/6	N0 Thymus	Interferon-inducible protein 203 (Mus musculus)				
	0610030F02	C57BL/6	Kidney	Interferon-related developmental regulator 2				
	2900034/12	C57BL/6	Hippocampus	Interferon-stimulated protein (15 kD)				
,	1600023101	C57BL/6	Placenta	Interferon-stimulated protein (20 kD)				
	2010006M08	C57BL/6	Small intestine					
	2010009P14	C57BL/6	Small intestine					
	2010107M23	C57BL/6	Small intestine					

p27-h gene showed increased expression in cardiac and skeletal muscle tissue undergoing active necrosis. These sequences are similar (data not shown) to the human IFN- $\alpha$ -inducible protein p27 (GenBank NM\_005532) shown to have increased expression in breast carcinoma (Rasmussen et al. 1993). The p27 protein is relatively small (11 kD) and highly hydrophobic. The p27-like mRNA is present in multiple tissues treated with the TNF, and it was speculated that these genes may be linked to inflammation related to the necrotic

processes. However, the specific function of p27 has yet to be determined.

Clone A430075K09 represents a potential new member of a group of structurally related interferon-inducible proteins, the Ifi 200 cluster, that regulate biological activities of IFNs. The IFNs are immune response modulating cytokines that exhibit antiviral effects, induce expression of class I or class II major histocompatibility complex proteins, and help activate macrophages, natural killer cells, and neutrophils. Se-

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Clone ID	Mouse strain	cDNA library	Gene name
2210418104	C57BL/6	Stomach	Interleukin 1 family, member 5 (δ)
4632413N13	C57BL/6	Skin (Neonate 0 day)	
2310041K07	C57BL/6	Tongue	
2310063B08	C57BL/6	Tongue	
1110033G16	C57BL/6	18-day embryo	Interleukin 1 family, member 6 ( $\epsilon$ )
9330131B06	C57BL/6	Diencephalon	
9330171A08	C57BL/6	Diencephalon	
A330066C15	C57BL/6	Spinal cord	Interleukin 1 receptor accessory protein
B230304B05	C57BL/6	C.quadrigemina	
C630028F01	C57BL/6	Hippocampus	
F630041P17	NOD	NOD-derived	Interleukin 1 receptor antagonist
4632427H17	C57BL/6	Skin	
C130072K02	C57BL/6	E16 Head	Interleukin 1 receptor, type I
E230023C01	C57BL/6	P2 oviduct	Interleville 1 months and the difference
B230327C20	C57BL/6	C.quadrigemina	Interleukin 1 receptor-associated kinase
D830047P08	C57BL/6	N16 heart	Interlaukin 1 recenter like 1
9030010H19 A530048E20	C57BL/6 C57BL/6	Colon Aorta and vein	Interleukin 1 receptor-like 1
			Interleukin 10 receptor, $\beta$
1500012D04	C57BL/6	Cerebellum N0 Thymus	Interleukin 10-related T cell-derived inducible factor
A430092F05 A430056E01	C57BL/6 C57BL/6	NO Thymus NO Thymus	Interleukin 12 receptor, β 1 Interleukin 12 receptor, β 2
A430038E01 A530070009	C57BL/6	Aorta and vein	inteneukin iz receptor, p z
C430004G12	C57BL/6	E7 whole body	Interleukin 13 receptor, $\alpha$ 1
G430044106	C57BL/6	Mixed <sup>a</sup>	interiourin' i preceptory a l
F830008L10	NOD	Activated spleen	Interleukin 13 receptor, $\alpha$ 2
D630025K20	C57BL/6	N0 kidney	Interleukin 15
A530045E03	C57BL/6	Aorta and vein	
A530048H17	C57BL/6	Aorta and vein	Interleukin 15 receptor, $\alpha$ chain
A630048I20	C57BL/6	N3 Thymus	
A130095H14	C57BL/6	N16 Thymus	Interleukin 16
9830138F10	C57BL/6	Bone	Interleukin 17 receptor
A530085G20	C57BL/6	Aorta and vein	
1110006016	C57BL/6	18-day embryo	Interleukin 17B
E030029C19	C57BL/6	N0 lung	Interleukin 18 receptor 1
F630101C07	NOD	NOD-derived	
E030025K24	C57BL/6	N0 lung	Interleukin 18 receptor accessory protein
E430021B09	NOD	(NOD) N2 Thymic	Interleukin 2 receptor, $\alpha$ chain
A130010M06	C57BL/6	N16 Thymus	Interleulin 2 mars (
5430409005	C57BL/6	Head (Neonate 6 day)	Interleukin 2 receptor, β chain
9530010F16	C57BL/6	U.bladder	Interleukin 2 recenter
A130027N13	C57BL/6	N16 Thymus Evoball (adult)	Interleukin 2 receptor, $\gamma$ chain
7530415H19 A430056E07	C57BL/6 C57BL/6	Eyeball (adult)	Interleukin 20
		N0 Thymus	Interleukin 21 receptor
A830047G15 1110033L24	C57BL/6 C57BL/6	N10 Cortex 18-day embryo	Interleukin 23, α subunit p19 Interleukin 25
2210408F01	C57BL/6	Stomach	
2010016N09	C57BL/6	Small intestine	
5830455M19	C57BL/6	Thymus	
9330197K12	C57BL/6	Diencephalon	Interleukin 4 receptor, $\alpha$
A630083M23	C57BL/6	N3 Thymus	intenediant receptor, a
E430003K23	NOD	(NOD) N2 Thymic	
F830019G17	NOD	Activated spleen from	Interleukin 6
		NOD.Cz Idd3	
9530096101	C57BL/6	U.bladder	Interleukin 6 receptor, $\alpha$
B230372K20	C57BL/6	C.quadrigemina	
C230031H18	C57BL/6	N0 Cerebellum	
A430091K05	C57BL/6	N0 Thymus	Interleukin 7
A530098102	C57BL/6	Aorta and vein	
A630007F02	C57BL/6	N3 Thymus	
D430026C11	C57BL/6	E13 lung	
A430089D02	C57BL/6	N0 Thymus	Interleukin 7 receptor
A530022C19	C57BL/6	Aorta and vein	
A630041A17	C57BL/6	N3 Thymus	
A630076J23	C57BL/6	N3 Thymus	
E430023N04	NOD	(NOD) N2 Thymic	Interleukin 9 receptor
2700043J07	C57BL/6	Èmbryo 11	Interleukin enhancer-binding factor 2
3000002D13	C57BL/6	Embryo 12-head	5
6230405A16	C57BL/6	Head (embryo 11)	
6330441H14	C57BL/6	Medulla	
B130010B06	C57BL/6	Partenogenetic	

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(continued)

	Clone ID	Mouse strain	cDNA library	Gene name
5	5730447J21	C57BL/6	Embryo 8	Interleukin enhancer-binding factor 3
	C130034D23	C57BL/6	E16 Head	
	E430021F21	NOD	(NOD) N2 Thymic	
36	3200002B17	C57BL/6	Embryo (14+17)-Head	Interleukin-four induced gene 1
	E330015O10	C57BL/6	P2 ovary	-
37	E130317D06	C57BL/6	N0 eyeball	Nuclear factor, interleukin 3, regulated
8	C130076 06	C57BL/6	E16 Head	Mus musculus IL-1F9 (IL1F9)
39	1110054H05	C57BL/6	18-day embryo	Mus musculus similar to Interleukin enhancer-binding factor

<sup>a</sup>Mixed library (AL,AK,AD,AF,AG,AC,AE,AQ,AP,AH,AA,AO,AB,AI,AJ,AN,AM).

quence of the clone A430075K09 shows similarity (84.9%ID, 425 amino acids, 100% of the total length) to the Ifi 205, the new member of the Ifi 200 family (Table 5 and Fig. 4, supplemental research data; available online at www.genome.org). Little is known about the precise function of the Ifi 205. However, it contains the PAAD/DAPIN/Pyrin domain (Pfam 02758), which is present in other members of the Ifi 200 family, indicating a possible protein-protein interaction site. This domain was predicted to have the same six *a*-helicescontaining fold as the well-described Death Domain (Pfam 00531). The death domain is present in the TNFR-1 and FAS/ APO1 receptors involved in cell death (Hofmann and Tschopp 1995). The potential functions of our candidate novel Ifi 205 factor, therefore, is the inhibition of cell proliferation and the arrest of cell growth, possibly through modulation of the activity of several transcription factors (Gribaudo et al. 1999).

Sequence of the clone 5330409J06 showed a weak similarity (Table 5 and Fig. 5, supplemental research data) to the guanylate-binding protein 5-GBP5 (P26376), published only as a database entry (GenBank AF288815). To our knowledge, the function and structure of GBP5 has not been described or published. In humans, the members of the GBP family are the most abundant class of proteins induced by IFN- $\gamma$  (Prakash et al. 2000). GBPs have the ability to hydrolyze guanosine triphosphate (GTP) to both mono- (GMP) or di-phosphate (GDP) products (Schwemmle and Staeheli 1994) involved in cellular signal transduction. However, little is known about either the specific biological function of GBPs or their cellular localization (Vestal et al. 2000). Members of the GBP family were reported to have antiviral properties (Anderson et al. 1999), play a role in the macrophage activation processes (Han et al. 1998), and inhibit proliferation of endothelial cells (Guenzi et al. 2001). The clone 5330409J06 contains GBP (Pfam 002263) and GBP-C (Pfam 02841) domains. It is therefore a strong candidate for a novel gene-the mouse guanylate-binding protein mGBP-5.

#### Interleukin-Related Novel Gene Candidates

Sequence of the clone B430113A10 contains Toll/Interleukin receptor TIR domain structure and shows similarity (88% identity in 123 AA overlap) to the hypothetical 14.3-kD protein in long-tailed macaque (Table 5 and Fig. 6, supplemental research data). The TIR domain (Pfam 01582) is an intracellular-signaling domain found in MyD88, interleukin 1 receptor, and the Toll receptor. The Toll/Interleukin receptor domain is associated with a family of proteins that are involved in early responses to injury and infection (e.g., interleukin-1 receptor) (Slack et al. 2000). The Toll-like receptors in insects

and mammals mediate innate defensive responses against a variety of microbial products (Anderson 2000). The clone B430113A10 thus represents a hypothetical protein potentially involved in immune responses.

Sequence of the clone 4732448K15 is similar (69% identity in full-length sequence of 598 AA) to the human IL-1 receptor-associated kinase-2 (IRAK-2), a proximal mediator of IL-1 signaling (Table 5 and Fig. 7, supplemental research data). IRAK-2 binds to the IL-1 receptor and is involved in IL-1-dependent signaling (Muzio et al. 1997). The clone 4232448K15 contains a Death Domain (Pfam 00531) (Hofmann and Tschopp 1995) present in many apoptosis pathway proteins. It also contains a catalytic Protein Kinase domain (Hanks and Hunter 1995) that has a conserved catalytic core common in both serine/threonine and tyrosine protein kinases. It is therefore a strong candidate for defining a novel gene—the mouse interleukin-1 receptor-associated kinase 2—mIRAK-2.

Sequence of the clone E230031K19 is weakly similar (76.3% identity over 93.3% length) to the human interleukin 20 receptor  $\alpha$  isoform 3 (Table 5 and Fig. 8, supplemental research data). The conceptual translation (amino acids 1–292, PFAM 01108) contains a tissue factor motif (Muller et al. 1996) that is involved in blood coagulation. IL-20 was reported to be overexpressed in transgenic mice that have skin abnormalities similar to psoriatic lesions in humans (Dumoutier and Renauld 2002).

#### Growth Factor-Related Novel Gene Candidates

The FANTOM2 clone D430010015 is a representative of a cluster (TIEG2-like) that showed similarity to human TGF $\beta$ inducible early growth response protein 2 (TIEG2) (Table 5 and Fig. 9, supplemental research data). The conceptual translation of 9830142A17 showed a slightly higher similarity (75% identity) to the human, than to mouse TIEG2 (O89091) (74% identity). TIEG2 is ubiquitously expressed in human tissues, enriched in pancreas and muscle (Cook et al. 1998). Similarly, the TIEG2-like FANTOM2 clones were derived from eight different libraries. The TIEG2-like clone has a conserved zinc finger domain. TIEG2 is involved in cell growth, and its overexpression in hamster ovary cells inhibits cell proliferation (Cook et al. 1998).

Sequence of the clone 4921501K24 showed similarity— 93.7% identity over the full length—(Table 5 and Fig. 10, supplemental research data) to TGF- $\beta$ -induced factor 2 (TGIF2), a homeobox gene of the TALE superclass, thought to play a role in the development and progression of some ovarian tumors through a gene-amplification mechanism (Imoto et al. 2000). TGIF2 was shown to repress TGF- $\beta$ -responsive

	Clone ID	Mouse strain	cDNA library	Gene name				
1	D730020G24	C57BL/6	L2 mammary	Epidermal growth factor				
	D430013M15	C57BL/6	E13 lung	Epidemiai growaniaetoi				
		NOD	(NOD) N2 Thymic cells					
	E430016A08							
2	1300005M11	C57BL/6	Liver	Epidermal growth factor receptor (Epidermal growth factor				
	1300008123	C57BL/6	Liver	receptor isoform 3) homolog ( <i>Mus musculus</i> )				
	1300003K07	C57BL/6	Liver					
	C630023M09	C57BL/6	Hippocampus	Epidermal growth factor receptor pathway substrate 15				
ļ	9430061P05	C57BL/6	Upper body	Epidermal growth factor receptor pathway substrate 15, related sequence				
	9430070A01	C57BL/6	E12 Upper body	Epidermal growth factor receptor pathway substrate 15, related sequence				
5	9830147 04	C57BL/6	Bone	Epidermal growth factor receptor pathway substrate 15,				
,								
	9830168E22	C57BL/6	Bone	related sequence				
,	5230401N02	C57BL/6	Xiphoid	Epidermal growth factor-containing fibulin-like extracellular matrix protein 1				
8	0610011K11	C57BL/6	Kidney	Epidermal growth factor-containing fibulin-like extracellular matrix protein 2				
)	0610042A18 C57BL/6 Kidney			Eukaryotic translation initiation factor 3 subunit 2 (EIF-3 $\beta$ )				
	2410126D05	C57BL/6	ES	(EIF P36) (TGF- $\beta$ receptor interacting protein)				
	2700045C12	C57BL/6	Embryo 11	(TRIP-1) homolog ( <i>Mus musculus</i> )				
	4930503E24	C57BL/6	Testis					
				Eibroblast growth factor (acidia) intracellular hindian anatoin				
)	1190024F07	C57BL/6	18-day embryo	Fibroblast growth factor (acidic) intracellular-binding protein				
	2010004G08	C57BL/6	Small intestine					
	D930040K23	C57BL/6	E15 head					
	2810001F12	C57BL/6	Embryo 10 + Embryo 11					
	2010005N21	C57BL/6	Small intestine					
	3010027N18	C57BL/6	Embryo 12-Head					
	G430069F17	C57BL/6	Mixed <sup>a</sup>					
	5730551C10	C57BL/6	Embryo 8	Fibroblast growth factor 15				
			Embryo 9	Fibrobiast growth factor 15				
	D030003J18	C57BL/6	2					
	D230045H09	C57BL/6	E12 eyeball					
2	0610042J09	C57BL/6	Kidney	Fibroblast growth factor inducible 14				
	2810484E09	C57BL/6	Embryo 10 + Embryo 11	Fibroblast growth factor-regulated protein 2				
	1110020J11	C57BL/6	18-day embryo					
	9630030D05	C57BL/6	N16	Latent transforming growth factor $\beta$ -binding protein 1				
	E330034H12	C57BL/6	P2 ovary	55 1 51				
	C730041004	C57BL/6	SV40T antigen					
	9830146M04	C57BL/6	Bone					
	E230007P04	C57BL/6	P2 oviduct					
	1810044G20	C57BL/6	Pancreas	Latent transforming growth factor $\beta$ -binding protein 3				
	A230048E14	C57BL/6	Hypothalamus					
	B430105M07	C57BL/6	Adipose					
,	E130107B17	C57BL/6	N0 eyeball	MEGF 12 homolog (Mus musculus)				
,	E130317D06	C57BL/6	N0 eyeball	Nuclear factor, interleukin 3, regulated				
;	1300004005	C57BL/6	Liver	TGF $\beta$ -inducible early growth response				
	A730095A08	C57BL/6	N7 Cerebellum	,,,,				
	A430010P14	C57BL/6	N0 Thymus					
			<u>,</u>	Transforming growth factor () 1 induced to receive 1				
	1100001105	C57BL/6	18-day embryo	Transforming growth factor $\beta$ 1-induced transcript 1				
	A530093G23	C57BL/6	Aorta and vein					
	D430029G15	C57BL/6	E13 lung					
	E030027O16	C57BL/6	N0 lung					
	1110002A05	C57BL/6	E13 lung	Transforming growth factor $\beta$ -regulated gene 1				
	2610001N22	C57BL/6	Embryo 10					
	2610029J18	C57BL/6	Embryo 10					
	D230045E03	C57BL/6	E12 eyeball	Transforming growth factor, $\beta$ -induced, 68 kD				
			, , , , , , , , , , , , , , , , , , ,					
	1700105F20	C57BL/6	Testis	Transforming growth factor, $\beta$ receptor I				
	4930438119	C57BL/6	Testis					
	1700105F20	C57BL/6	Testis					
	5730530B02	C57BL/6	Embryo 8	Hypothetical 30.1-kD protein (TGF β inducible nuclear protei				
	6720481N02	C57BL/6	Extra testis (embryo 12)	TINP1) (hairy cell leukemia protein) (Homo sapiens)				
	2310004E21	C57BL/6	Tongue					
	2810027P06	C57BL/6	Embryo 10 + Embryo 11					
	3200001B21	C57BL/6	Embryo (14 + 17)-Head					
	0610010G24	C57BL/6	Kidney					
	0010010024		Embryo 8					

<sup>a</sup>Mixed library (AL,AK,AD,AF,AG,AC,AE,AQ,AP,AH,AA,AO,AB,AI,AJ,AN,AM).

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	Clone ID	Mouse strain	cDNA library	Gene name
1	1810065101	C57BL/6	Pancreas	Small inducible cytokine A1
2	2310074D07	C57BL/6	Tongue	Small inducible cytokine A11
	2310011L12	C57BL/6	Tongue	,
3	2700043D19	C57BL/6	Embryo 11	Small inducible cytokine A12
4	1600023B02	C57BL/6	Placenta	Small inducible cytokine A27
	1600002M16	C57BL/6	Placenta	,
5	4921506E08	C57BL/6	Testis	Small inducible cytokine A28
6	1010001B04	C57BL/6	Heart	Small inducible cytokine A5
7	0610030C20	C57BL/6	Kidney	Small inducible cytokine A6
	2010308M07	C57BL/6	Small intestine	,
8	8430433 09	C57BL/6	Lung (embryo 16)	Small inducible cytokine A7
9	1810063B20	C57BL/6	Pancreas	Small inducible cytokine A8, MCP-2
0	1810031C02	C57BL/6	Pancreas	Small inducible cytokine A17
1	A430095C13	C57BL/6	N0 Thymus	Small inducible cytokine A20
2	A930010L22	C57BL/6	Retina	Small inducible cytokine B, member 5
3	A630052H15	C57BL/6	N3 Thymus	Small inducible cytokine B (Cys-X-Cys), member 9
	A630095J17	C57BL/6	Thymus	
4	A430052M02	C57BL/6	NÓ Thymus	Small inducible cytokine B (Cys-X-Cys), member 11
	A430101A17	C57BL/6	N0 Thymus	
	C730001K22	C57BL/6	SV40T antigen transgenic mouse	
5	4631412M08	C57BL/6 (EST_a)	Skin (Neonate 0 day)	Small inducible cytokine B (Cys-X-Cys), member 13
6	1110012L01	C57BL/6 (EST_a)	18-day embryo	Small inducible cytokine B (Cys-X-Cys), member 14
	1110031L23	C57BL/6	18-day embryo	
	1200006123	C57BL/6	Lung	
	2810410F08	C57BL/6	Embryo 10 + Embryo 11	
	3300001J19	C57BL/6	Embryo 17-Head	
7	3110001P14	C57BL/6	Embryo 13-Head	Small inducible cytokine B, member 15
8	A430076J21	C57BL/6	N0 Thymus	C, member 1 (lymphotactin)
9	6430571D05	C57BL/6	Olfactory brain	Small inducible cytokine D, 1
	A430078 23	C57BL/6	N0 Thymus	

transcription through interaction with a TGIF-binding site in DNA (Melhuish et al. 2001).

Little is known about the integrin  $\beta$ -like 1 homolog (with EGF-like repeats), that showed similarity to the clone B930011D01 (Table 5 and Fig. 11, supplemental research data). The EGF-like domain is characteristic for extracellular domain of membrane-bound proteins (InterPro entry IPR000561). This product may possibly function as a cell-adhesion receptor.

The conceptual translation of clone E330012C08 showed similarity (Table 5 and Fig. 12, supplemental research data) to the human latent TGF-B-binding protein 4S, LTBP-4 (O75412). The LTPB-4 is a high molecular weight calciumbinding protein with EGF-like repeats. The cDNA of the LTBP-4 was found in several different forms generated by alternative splicing (Saharinen et al. 1998), consistent with the suspected frame shift positions found in the clone E330012C08. TGF-β proteins are multifunctional growth factors excreted as inactive precursors forming large protein complexes. These complexes include, among others, LTBPs which are believed to regulate both local tissue deposition of TGF-β and the related signaling (Sterner-Kock et al. 2002). The disruption of LTBP-4 in mice leads to ailments such as severe pulmonary emphysema, cardiomyopathy, and colorectal cancer (Sterner-Kock et al. 2002).

Finally, the conceptual translation of clone 9130025L13 showed similarity (Table 5 and Fig. 13, supplemental research data) to the human FGF receptor 4B—FGFR4b (Q9BXY2). FGF receptors are involved in multiple hormonal and proliferative processes of hormonal cells (Yu et al. 2002). A variant of the

FGF receptor 4, related to the FGFR4b was found to increase tumor cell motility (Bange et al. 2002).

#### DISCUSSION

The mouse is the most popular model organism for both basic and applied immunological studies, including the study of cytokines. Our analysis of the FANTOM2 data set focused on the subset of transcripts that showed similarity to the cytokine-related products, and which had cytokine names or abbreviations in the name or description fields of the corresponding database entries (the cytokine-name criterion). Our study is only partial because databases include cytokinerelated entries of genes and proteins that do not have an explicit cytokine name in the name or description fields. However, this study is representative because it focused on a well-defined subset of cytokine-related genes from which we can make an estimate of the coverage of cytokine-related genes in the FANTOM2 data set.

In comparison with database entries that conform to the cytokine-name criterion, we estimate that the ratio of cytokine-related products represented in the FANTOM2 data set is of the order of 20% of the total known cytokine-related products. This estimate is not surprising, because cytokines are inducible and often tissue-specific products. In addition, even when induced, the expression level of cytokines is often very low. The RIKEN libraries, although diverse, may not represent the optimal conditions required for induction of cytokines and their related genes. The regulatory processes involving cytokines are complex processes involving activation and suppression of multiple genes (Doly et al. 1998; Munshi et al. Brusic et al.

	Clone ID	DDBJ <sup>a</sup>	cDNA library	Similarity <sup>b</sup>	Similar gene name
1	1110004C05 <sup>c</sup>		18-day embryo	89.8, 100.0 (145 AA)	Interferon-inducible protein (Rattus norvegicus).
	1810060G19 1810061A10	AK003507	Pancreas Pancreas	89.1, 100.0 89.8, 100.0	Identical to 1110004C05RIK protein (Q9CQW9)
2	4930507H06 <sup>c</sup>		Testis	73.1, 75.9 (145 AA)	Interferon-inducible protein (Rattus norvegicus).
	4933438K12	AK076846	Testis	64.8, 89.0	Identical to 4933438K12RIK protein (Q9D3R8)
3	A330075D07 <sup>c</sup>	AK039626	Spinal cord	52.8, 81.8 (145 AA)	Interferon-induced transmembrane protein 2 (interferon-inducible protein 1-8D) ( <i>Homo sapien</i>
4	1110036C17 <sup>c</sup>	AK004121	18-day embryo	72.9, 78.1 (145 AA)	Homolog to interferon-inducible protein
5	2310061N23 <sup>c</sup>	AK010014	Tongue	78.0, 97.6 (84 AA)	<ul> <li>α-interferon inducible protein (fragment)</li> <li>(Mesocricetus auratus)</li> </ul>
6	A430075K09 <sup>c</sup>	AK040191	N0 Thymus	84.9, 100.0 (425 AA)	Interferon-activatable protein 205 (IFI-205), also
	9830003L02		Bone	77.3, 77.3	known as D3 protein ( <i>Mus musculus</i> ). Contains PAAD/DAPIN/ Pyrin domain (Pfam 02758) and t associated HIN-200/IF120x domain.
7	5330409J06 <sup>c</sup>		Pituitary gland	69.0, 93.6	IF-induced GUANYLATE-BINDING PROTEIN 5
	9530054H01	AK030414	U. bladder	65.1, 29.3	(Homo sapiens)
~	E030025M22	41/000070	N0 lung	69.0, 93.6 (551 AA)	The sector of the life of the sector of the sector
8	B430113A10 <sup>c</sup>	AK080873	Adipose	87.6, 100.0 (123 AA)	Hypothetical Toll/Interleukin receptor TIR domain structure containing protein
9	4732448K15 <sup>c</sup>		Skin (Neonate 10 day)	68.8, 100.0 (598 AA)	Interleukin-1 receptor-associated kinase 2
	9130227N12	AK028733		75.5, 24.2	(Homo sapiens)
~	6330415L08	44054015	Medulla oblongata	64.9, 100.0	Interlevilie 20 meanter informe 2 (11-ma
0	E230031K19 <sup>c</sup>	AKU54215	P2 oviduct	76.3, 93.3 (556 AA)	Interleukin 20 receptor $\alpha$ isoform 3 (Homo sapiens)
1	9830142A17		Bone	75.2, 99.8 (512 AA)	Transforming growth factor-β-inducible early grow
	C330038O18		ES cells	75.0, 99.8	response protein 2 (TGF $\beta$ -inducible early growth
	C730023J20		SV40T antigen transgenic	75.4, 99.8	response protein 2) (TIEG-2) (Kreppel-like factor
	D430010O15 <sup>c</sup> A230003P18	AK084913	E13 Hypothalamus	75.0, 99.8 75.2, 99.8	11) (Homo sapiens)
	D630011E14		N0 kidney	75.0, 99.8	
	E230024 15		P2 oviduct	75.2, 99.8	
	E430037H02		(NOD)N2 Thymic cells	75.1, 99.8	
2	4921501K224 <sup>c</sup>		Testis	93.7, 100.0 (237 AA)	DJ977B1.4.1 (isoform 1) (TGIF2) (TGFβ-induced factor
-	5730599009	AK076631	Embryo 8	89.9, 100.0	(TALE family homeobox)) homolog (Homo sapiens)
3	B930011D01 <sup>c</sup>	AK047014	N10 Cerebellum	88.2, 84.4 (153 AA)	Integrin, β-like 1 (with EGF-like repeat domains) homolog ( <i>Mus musculus</i> )
4	2310046A13 <sup>c</sup>		Tongue	88.3, 26.7	Latent transforming growth factor-\binding prote
	E330012C08	AK054297	P2 ovary	88.8, 96.5 (1505 AA)	4S homolog (Homo sapiens)
	A730031D03		N7 Cerebellum	95.4, 32.5	
5	9130025L13 <sup>c</sup>	AK033606	Cecum	77.1, 99.5 (423 AA)	FGF receptor 4B (Homo sapiens)
	9530005F04	/ 11033000	U. bladder	62.8, 99.5	

Thirty-one clones representing fourteen candidate new genes.

<sup>a</sup>DDBJ accession no. of the representative transcript.

<sup>b</sup>Similarity was defined by two parameters, first the percent identity and second the percent of the length of the reference sequence (similar-gene) having similarity to the FANTOM2 clone. The number in the parentheses in the SIMILARITY field shows the length of the alignment or the maximum length of multiple alignment. All clones were derived from the mouse strain C57BL/6, except for clone E430037H02, which was derived from NOD mouse.

<sup>c</sup>Representative transcripts.

1999; Vanden Berghe et al. 2000). This illustrates the need for creating more specific cDNA libraries that dynamically capture gene expression profiles for the study of the cellular processes involving cytokines.

It is not always possible, particularly in large-scale studies, to derive clear conclusions and explanations of the data. Our selection of the threshold that distinguishes novel from the known genes or gene candidates is arbitrary. The clone representing integrin  $\beta$ -like homolog (Fig. 11, supplemental research data) has only 10 amino acid differences to its reference sequence. Such clones may, therefore, represent variants or merely the inter-strain differences of the same transcript. Three of the clones representing the 1-8 family of IFNinducible proteins (Fig. 1) do not appear to be full length and may represent poor-quality sequence data. We should also be aware of the possible problems with the reference data. The Ifi

205 reference sequence (Fig. 4, supplemental research data) has a triple repeat of the sequence AGSTAQ, which is missing in the corresponding FANTOM2 clone. This may possibly represent a sequencing artifact in the reference sequence. Conversely, the FANTOM2 clone having sequence similarity to human latent TGF-β binding protein (Fig. 12, supplemental research data) contains P+R-rich regions that are not present in the reference sequence and may represent artifacts. Mapping FANTOM2 clones to the mouse genome sequence at Ensembl (Hubbard et al. 2002) indicated that some of the FANTOM2 clones represented clear hits, whereas others, (especially Ifi 200 clones) had multiple hits. The multiple hits results may be explained by possible artifacts or poor sequence data, but may also represent the complexity of transcriptional regulation of inducible genes warranting further investigation.

#### Cytokine-Related Genes in Mouse

	10	20	30	40	50	60	70	80
	1	1	Ĭ	I	T.	. 1	1	Ĭ.
SPTR   Q8QFL3	-MDQSPSYQPEFVP	MNGNKYMRLI	EDPHGAPKF-(	2HTVVLGLPQP	VVPQPR	DHIIWSLCS	LVYGNPLCLG	MLAVYF
SPTR Q9R175	MSHNSQAFLPA-NA	GLPPSYETI	KEEYGVTELGH	PNNSAVVRTT	VINMPREVS	VPDHVVWSLFN	TLFFNACCLG	FIAYAY
RI 1810061A10	MNHTSQAFITAASG	GQPPNYERI	KEEYEVAEMG/	APHGSASVRTT	VINMPREVS	VPDHVVWSLFN	TLFMNFCCLG	FIAYAY
RI 4930507H06		NQEMPI	KDQHEVVVMG	PHTSTSSTTT	IITMP-EIS	KPDYVVWSLFN	TLFMNFCCLG	FIAYAY
RI 1110036C17		EMPI	KEQQEVVVLGS	SPHISTSATAT	TINMP-EIS	TPDHVVWSLFN	TLFMNFCCLG	FVAYAY
RI A330075D07				APVPST	VVCINSDVI	QPDHITWSTFN	SVFMNGCCLG	FIAYIY
Conserved motif						Dx+xWSxxx	x++xNxxCLG	x+Axx+
	90	100	110	120	130	140		
	1	Î	Ī	Ï	l.	1		
SPTR Q8QFL3	SIKSRDRKMVGDLE	GARKHGKTAI	RCFNVVTLTLV	/ILGLLFLFII	YGFFIYNIS	HL	-	
SPTR Q9R175	SVKSRDRKMVGDVI	GAQAYASTA	KCLNISSLIFS	SVLMVIICIII	FSTTSAVVF	QSLSQRTPHSG	F	
RI 1810061A10	SVKSRDRKMVGDVT	GAQAYASTA	KCLNISTLVLS	SILMVVITIV-			-	
RI 4930507H06	SVKSRDRKMVGDMT	GAQAFASTA	RCLNISCLILS	SVVMVILFI			-	
RI 1110036C17	SVKSRDRKMVGDTT	the second se	The second s					
RI A330075D07	SVKSRDRKMVGDMT	GAQSHASTAI	KILNILALVIS	SLIFYIMLIVL			H	
Conserved motif	S+KSRDRKMVGDxx	GAXXXXXTA	kxxN+xxL					

**Figure 1** The alignment of conceptual translations of the four candidate novel mouse genes of the family 1-8 of IFN-inducible proteins with representative sequences from rainbow trout (Q8QFL3) and rat (Q9R175). A conserved motif characteristic for the members of interferon-induced 1-8 family of proteins in mice, rats, humans, cattle, and trout is shown with the alignment. Capital letters in the consensus motif stand for identical residues, + for conserved residues (aliphatic—I,V,L, or M; positively charged—K or R; aromatic—F,Y, or W), and x for any residue. The ubiquitin-conjugating enzyme E2 active site motif in 1810061A10 [25] and a similar motif in 4930507H06 are shaded. SPTR stands for SWISS-PROT/TrEMBL database [15], and RI stands for RIKEN clone.

Computational analysis has been applied previously to identification of novel cytokines from EST libraries, such as cardiotrophin-like cytokine (Shi et al. 1999). We have shown that this process can be enhanced by using the elimination strategy that involves multiple consecutive steps as follows: generation of multiple cDNA libraries, large-scale computational annotation, human curation, identification of relevant subsets of sequences, and individual clone analysis. This process involves a number of elimination steps-it is therefore critical that the loss of information is kept at a reasonable level. Too stringent criteria for elimination of clones from further analysis would lead to a small number of clones having a high-quality and high-confidence annotation. On the other hand, this would lead to the elimination of clones that represent results of complex transcriptional events, modifications, or variants. Too lenient criteria for the inclusion of clones for further analysis may lead to the proliferation of artifacts and errors into sequence data. To minimize the proliferation of errors, we excluded a number of clones from the analysis, including those that contained stop codons (data not shown).

We identified 15 candidates for novel cytokine-related genes. Because many cytokine-related genes do not contain cytokine names in their description, this should represent only a fraction of the novel cytokine-related candidate genes captured by the FANTOM2 library. Further strategies will involve making comprehensive lists of cytokine-related genes followed by comparison of these genes with the existing cDNA libraries and future cDNA libraries designed for capturing specific gene expression during the immune processes.

#### **METHODS**

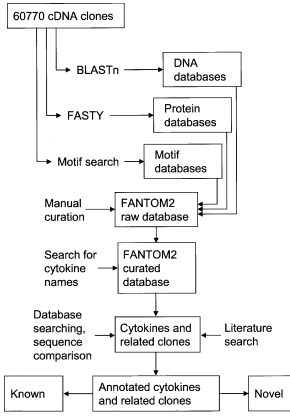
The FANTOM2 set of full-length mouse cDNA clones, referred to as clones in this text, contains 60,770 sequences with the total length of almost 120 million base pairs and more than 12 million conceptually translated amino acids. The FANTOM2 clones were functionally annotated using automatic computational annotation followed by expert human curation (Okazaki et al. 2002). The overall process of identification of novel candidate genes is shown in Fig. 3).

#### Sequence Analysis

The sequences of the cDNA clones were compared with the genetic, protein, motif and protein domain databases. The genetic databases used for the analysis of clones included GenBank (Benson et al. 2002), LocusLink (Wheeler et al. 2002), UniGene (Wheeler et al. 2002), Mouse Genome Database (MGD) (Blake et al. 2002), and TIGR databases (Quackenbush et al. 2001). The comparisons to the genetic databases were performed using the nucleotide to nucleotide sequence local alignment algorithm BLASTN (Altschul et al. 1990). The protein databases used for the analysis of FANTOM2 clones included SWISS-PROT and TrEMBL (Bairoch and Apweiler 2000), and PIR (Wu et al. 2002). The comparisons were performed using the program FASTY (Pearson et al. 1997) that compares a DNA sequence with protein sequences in a database. FASTY translates a DNA sequence in three frames and aligns the translations to each sequence in the protein database, permitting gaps and frame shifts. Finally, the FANTOM2 clones were compared with the SCOP (Lo Conte et al. 2002), Pfam (Bateman et al. 2002), and InterPro (Apweiler et al. 2001) databases for identification of potential protein domains and functional sites, and to the MDS (Maximum Den-

	10	20	30	40	50	60	70	80
	1	1	I	I	1	1	1	1
RI 2310061N23	GALAVAGA	APIALSAVGFTG#	GIAAGSIAAK	MMSAAAIANG	GGVAAGSLVA	TLQSAGVLGL	STSTNAILGA	AGA-AVGALL
Q9H2X8	AAVGGALAVGA	/PVVLSAMGFTG#	GIAASSIAAK	MMSAAAIANG	GGVSAGSLVA	TLQSVGAAGL	STSSNILLAS	SVGS-VLGACL
Q9QXH3	GAGAVVA	APVVLGAVGFTG	GIAAGSIAAN	MMSAAAIANG	GGVAAGSLVA	ILQSVGVAGL	STSSNIILGS	SVGAGSVGAAL
Consensus	GA AV	P+ L A+GFTGA	AGIAA SIAA	MMSAAAIANG	GGV AGSLVA	LQS G GL	STS+N +L	G+ +GA L

**Figure 2** Candidate novel gene represented by clone 2310061N23 similar to the fragments of IFN- $\alpha$  inducible proteins in human (Q9H2X8) and hamster (Q9QXH3). (+) Conserved replacements.



**Figure 3** The overall process of identification and functional annotation of cytokine-related FANTOM2 clones.

sity Subgraph) Motif Database (Kawaji et al. 2002) for the presence of potential protein motifs. Unless specified otherwise, the gene products references throughout the text correspond to SWISS-PROT and TrEMBL (SPTR) primary accession numbers.

# Functional Annotation and Identification of FANTOM2 Clones

Functional annotation of clones was performed in three steps as follows: data mining, manual annotation, and sequence analysis of the selected clones followed by literature text searching. First, the data-mining step involved a large-scale sequence comparison using BLASTN or FASTY against all of the selected databases. The results of these searches were accessible to the annotators via the FANTOM2 graphical interface, providing a convenient tool for the second step, the manual annotation. The most likely coding sequences (CDS) were determined by comparison of the results of four prediction programs (ProCrest, Decoder, rsCDS, and NCBI cds). Additional tools, such as ClustalW (Thompson et al. 1994) were available for facilitation of the functional annotation. The annotation results were then summarized and searched by keywords cytokine, interferon, IF, IFN, interleukin, IL, transforming growth factor, TGF, epidermal growth factor, EGF, fibroblast growth factor, and FGF. The clones that showed 50%-95% identity to the known cytokine-related genes were considered as candidate novel genes and were further analyzed using the programs BLAST or FASTA. The clone Á530093G11 was also included in analysis, because its conceptual translation showed 96% identity over the 50% of the reference sequence length. Finally, literature searches in the PubMed (Wheeler et al. 2002) were performed for identification of key publications. The possible functions of the candidate novel genes were elucidated from literature sources.

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