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GFScan: A Gene Family Search Tool at Genomic DNA Level

Zhenyu Xuan, W. Richard McCombie, and Michael Q. Zhang¹

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We have developed **GFScan** (**G**ene **F**amily **S**can), a tool that identifies members of a gene family by searching genomic DNA sequences with genomic DNA motifs (or matrices) that are representative of the family. We have tested **GFScan** on four human gene families including the neurotransmitter-gated ion-channels (NGIC) family, the carbonic anhydrases (CA) family, the Dbl homology (DH) domain family, and the ETS-domain family. All known members of these families with motifs mapped to sequenced genomic DNA regions were found, whereas some novel genomic locations were also found to match the motifs, which may indicate new members in these families. Compared with other methods, **GFScan** recognized all true positives with much fewer false positives. We also showed that motifs constructed based on human genes could be used to search the mouse genome to identify orthologous family members in mouse. This program is available at <http://www.cshl.org/mzhanglab/>.

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With the advances of several whole-genome sequencing projects, including human, mouse, *Drosophila*, and so on, more and more genomic DNA sequences have become available. These projects make it possible to analyze gene families in one species systematically. One of the well-known strategies for gene family analysis is to detect all the gene models first in one genome with some gene prediction methods, such as **Genscan** (Burge and Karlin 1997), **Genie** (Kulp et al. 1996), or **FGENES** (Solovyev and Salamov 1997); translate these genes into proteins; then try to find gene families at the protein level using similarity search or protein motif databases, such as **BLOCKS+** (Henikoff et al. 1999), **Pfam** (Bateman et al. 1999), **ProDom** (Corpet et al. 1999), **PRINTS** (Attwood et al. 1999), **PROSITE** (Hofmann et al. 1999), **IntroPro** (<http://www.ebi.ac.uk/interpro/>). Additionally, mRNAs can also be used to find gene family members by **BLAST** or **FASTA** searches (Pearson and Lipman 1988; Altschul et al. 1990, 1997). Recently, Henikoff (Henikoff and Henikoff 2000) had tried to use protein fragments in the **BLOCKS+** database to search the *Drosophila* genomic sequence using **BLAST**.

Our method seeks to find all members of a gene family by searching the whole genome with the representative genomic DNA motif of this family. Motif search at the protein level is a reliable method to find protein family members based on known proteins. However, protein motifs can only be used to search the known proteins, and some proteins remained undiscovered by existing experimental or theoretical methods. On the other hand, **TBLASTN**, a program of the **BLAST** package, can align protein sequences with genomic DNA sequence directly to find matched regions that may code new members of the gene family. However, as shown in the Results and Discussion sections, programs in the **BLAST** family are general sequence-alignment programs and find many false positives. To circumvent this problem, we developed **GFScan** (**G**ene **F**amily **S**can), which uses a representative DNA

motif of a gene family to search genomic DNA sequence directly to identify new members of the gene family. The representative genomic DNA motif is constructed based on protein motifs in **PROSITE** (release 16.0, updates up to September 2000) and the genomic structure of known members of the family. As more and more mRNA and protein sequences are submitted to the public databases, and as each genome becomes more complete, **GFScan** will be increasingly effective to find new members of a gene family.

RESULTS

GFScan was developed in C++ language. To show the usefulness of this program, we applied it to four gene families, searching for new members of the family in the whole human genome (Genome Sequencing Consortium 2001) in GoldenPath (April 2001 freeze and August 2001 freeze; <http://genome.ucsc.edu/>) and mouse genome in the Celera Genomics Company's database.

Neurotransmitter-Gated Ion-Channels (NGIC) Family

The human neurotransmitter-gated ion-channels family is a large family, whose members include GABA (γ -aminobutyric acid) A receptors, glycine receptors, acetylcholine receptors, and 5-hydroxytryptamine-3 receptor. All members of the family have a common protein motif, called **NEUROTR_ION_CHANNEL** in the **PROSITE** database (ID: PS00236). Using the known 37 human genes of this family in the public database and the protein motif in **PROSITE**, a 45-bp intronless genomic DNA motif was constructed. We also found that one family member, **CHRN1**, has an intron in the motif-matching genomic region, and the intron separates the 45-bp motif into two parts. An intron-containing genomic DNA motif was then constructed (see Methods). Both genomic DNA motifs were used to search the whole human genome. Of 37 known motif regions, 29 were found by **GFScan**. For the missed eight genes, all the genomic regions corresponding to the motifs fell into the gaps of the genome. Moreover, nine additional genomic regions were found. Three of them were

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Table 1. Results on Neurotransmitter-Gated Ion-Channels Family

No.	Chromosome	Strand	Motif position		Description
1	X	+	13920432	13920477	GLRA2
2	X	-	104174460	104174415	(Similar to mouse Glra1) ^{New}
3	X	-	152091851	152091806	GABRE
4	X	-	152454684	152454639	GABRA3
5	1	-	1478204	1478159	GABRD
6	2	-	178143495	178143450	CHRNA1
7	2	+	237873656	237873701	CHRNA2
8	2	+	237886519	237886564	CHRNA3
9	3	-	104298942	104298897	(Similar to Rat Gabrr3) ^{New}
10	4	-	50087519	50085574	(Similar to Rat Gabrg1) ^{New}
11	4	-	50330244	50330199	GABRA2
12	4	+	50782012	50782057	GABRA4
13	4	-	51222098	51222053	GABRB1
14	4	+	166964539	166964584	GLRB
15	4	-	184669299	184669254	GLRA3
16	5	-	174399338	174399293	GABRB2
17	5	+	174946361	174946406	GABRG2
18	5	+	183945827	183945872	GABRP
19	6	-	98310008	98309963	GABRR2
20	8	-	29298237	29298192	CHRNA2
21	8	+	45473102	45473147	CHRNA3
22	8	-	45498244	45498199	CHRNA6
23	10	-	58232334	58232289	(REPEAT region) ^{New}
24	10	+	96875579	96875624	(Similar to mouse Gabra3) ^{New}
25	11	-	2590955	2590910	CHRNA10
26	11	-	125455260	125455215	(HTR3A duplication) ^{New}
27	11	-	125490375	125490330	HTR3A
28	15	+	22459117	22459162	GABRB3
29	15	+	24261884	24261929	(Similar to Gallus Chrna8) ^{New}
30	15	+	24323276	24323321	(CHRNA7 duplication) ^{New}
31	15	+	26565917	26565962	CHRNA7
32	15	+	76709488	76709533	CHRNA5
33	15	-	76721764	76721719	CHRNA3
34	15	-	76752832	76752787	CHRNA4
35	17	-	5014125	5014080	CHRNA1
36	17	-	5277247	5277202	(CHRNA1 duplication) ^{New}
37	17	-	8070655	8070258	CHRNA1
38	20	-	63883819	63883774	CHRNA4

Missed known genes in this family: GABRA6 (NM_000811), GABRQ (NM_018558), CHRNA9 (NM_017581), GLRA1 (NM_000171), GABRA1 (NM_00806), GABRA5 (NM_000810), GABRR1 (NM_002042), CHRNA2 (NM_000748).

Table 2. Results on Carbonic Anhydrases (CA) Family

No.	Chromosome	Strand	Motif position		Description
1	1	-	230527424	230527143	(CA14 duplication) ^{New}
2	1	+	230570250	230570531	CA14
3	1	+	9063801	9065482	CA6
4	3	+	68787364	68790131	(PTPRG) ^{New}
5	4	+	138352740	138353600	(CA7 duplication) ^{New}
6	8	+	89796759	89803951	(Similar to mouse Car13) ^{New}
7	8	-	89874524	89871143	CA1
8	8	+	89982297	89984196	CA3
9	8	+	90013921	90014490	CA2
10	9	+	38809920	38810067	CA9
11	15	+	60482427	60485863	CA12
12	16	-	23050079	23047764	CA5
13	16	-	33403805	33401499	(CA5 duplication) ^{New}
14	16	-	33988707	33986402	(CA5 duplication) ^{New}
15	16	+	77218848	77219708	CA7
16	16	+	77672109	77672969	(CA7 duplication) ^{New}
17	17	-	55621766	55527814	LOC56934
18	17	+	64513194	64513368	CA4
19	19	-	57550194	57549926	CA11

Missed known genes in this family: CA8 (NM_004056), CASB (NM_007220).

duplications of the known genes. Among the remaining six novel genomic regions, one is located in the repeat region, and the other five were likely to be members of this gene family that are previously unidentified. Based on the human genome annotation in GoldenPath (<http://genome.ucsc.edu/>), these five regions were reported to be similar to mouse glycine receptor subunit $\alpha 1$, rat GABA A receptor subunit $\gamma 1$, rat $\rho 3$, mouse GABA A receptor subunit $\alpha 3$, and Gallus nicotinic acetylcholine subunit $\alpha 8$, respectively. With the exception of GABA A receptor $\alpha 3$, no mRNA or protein sequence has been known for the other four genes (see Table 1).

Carbonic Anhydrases (CA) Family

Human carbonic anhydrases (CA) are zinc metalloenzymes that catalyze the reversible hydration of carbon dioxide. There are 14 known members in the family. From the mRNAs of the known members, we first constructed a 57-bp cDNA motif based on the PROSITE protein motif (ID: PS00162). All of the genomic sequence regions corresponding to this cDNA motif contain one intron. The splice locations of the introns are identical among all members, but the lengths of the introns are different. We next constructed a genomic DNA motif from the cDNA motif incorporating information on the intron. By searching the whole human genome with the genomic DNA motif, 12 of 14 known genes were found, and the two genes that were missed had their motif-matching genomic region falling into the genomic gaps. Moreover, we found two additional genomic regions that match the motif: One was related to a non-CA family gene, *PTPRG* (protein tyrosine phosphatase, receptor type G) in Chromosome 3; the other was found in Chromosome 8, whose closest homologous gene was the mouse *Car13* gene. It is worth noticing that the human *CA13* gene has not been found before, and our finding may have shed light on this new member of the family (see Table 2).

Table 3. Results on DH-Domain Family

No.	Chromosome	Strand	Motif position		Description
1	X	-	141522542	141523440	DBL
2	9	-	137956245	137957015	VAV2
3	9	-	137956245	137964371	(VAV2 pseudo-site) ^{New}
4	13	+	117682791	117683814	DBS
5	17	-	631694	639380	ABR
6	19	+	28364356	28364681	(VAV duplication) ^{New}
7	19	-	7283797	7284122	VAV
8	21	-	29371499	29371577	TIAM
9	22	+	20261116	20264414	BCR

Missed known genes in this family: VAV3 (NM_006113) (found in chr1 125174019-125178912 in Goldenpath Aug 2001).

Dbl Homology (DH) Domain Family

The Dbl homology (DH) domain is responsible for the guanine nucleotide exchange factor (GEF) catalytic activity (Zhu et al. 2001). Eight human genes belong to this family, and some of these genes are oncogenes, including *DBL*, *Break Cluster Region (BCR)* oncogene, *VAV*, *VAV2*, and *VAV3*. The protein sequences of all eight members share the DH domain (PROSITE ID: PS00741). From their mRNA sequences, a 78-bp cDNA motif was constructed. In the genomic regions corresponding to the motif, no intron was found for one of the family members, *TIAM*; two introns were found for *ABR* and *BCR*; and one intron was found for the remaining five members of the family. Based on above information on gene structure, we next constructed three genomic DNA motifs of this domain from the cDNA motif. Searching the whole human genome with the genomic DNA motifs revealed nine genomic regions that significantly match the motifs. Among the nine

regions, seven contain known genes, one of the two new locations was the *VAV* gene's genomic DNA sequence duplication, and the other overlapped with the known *VAV2*'s motif region (see Table 3). *VAV3* was the only known member of the family that was missed by the search, and this is because the genomic region matching the motif region was not available in the April 2001 Goldenpath freeze (it was found in the August 2001 freeze).

ETS-Domain Family

The ETS-domain gene family includes a group of proteins that function as transcription factors under physiologic conditions and, if aberrantly expressed, can cause cellular transformation (Karim et al. 1990). These proteins share a conserved domain, the ETS domain, which is involved in DNA binding. From the mRNAs of the 19 known members and a protein motif in the PROSITE database (ID: PS00346), a 48-bp cDNA motif was constructed. Four of these 19 genes have one intron in their genomic regions matching the motif, and the splice location of the intron is the same. Therefore, we constructed an intron-containing genomic DNA motif, and it is used to search the human genome together with the cDNA motif. Twenty-six genomic regions were found to match the motifs, which include 18 of the 19 known genes. *ETV5*'s genomic DNA motif region was missed because the genomic DNA sequence around the motif-matching region was uncompleted. Out of the eight additional motif-matching regions, three were duplications of three known genes (i.e., *GABP*, *ETV6*, and *ERF*). The other five were related to unknown genes in human: one was in the *FEV* gene region, two were similar to mouse Ets-protein Spi-C (GenBank accession no. AF098863), and the last two were located in two genes predicted by *GenScan* and *Ensembl*. Both *FEV* and *Spi-C* are ETS-domain family members (Bemark et al. 1999). *FEV* was not listed in the PROSITE database because of the database-updating problem, and human *Spi-C* has not been found. Likely, these new motif-matching regions will provide experimental scientists with useful guidance to identify new members of the ETS-domain family in the human genome (see Table 4).

Table 4. Results on ETS-Domain Family

No.	Chromosome	Strand	Motif position		Description
1	X	+	46781345	46781393	ELK1
2	1	+	177575228	177576887	ETV3/PEP1
3	1	+	177611558	177611851	(<i>GenScan</i> predicted gene) ^{New}
4	1	-	229691250	229691298	ELK4
5	2	-	223797915	223797963	(<i>FEV</i> gene) ^{New}
6	6	-	40661831	40661879	TEL2
7	7	-	13211815	13211863	ETV1
8	7	+	63416487	63416535	(<i>GABP</i> duplication) ^{New}
9	11	-	142926355	142926403	ETS1
10	11	+	143364213	143364261	FLI1
11	11	-	34770889	34770937	(Similar to Mus. AF098863) ^{New}
12	11	-	48738245	48738293	SPI1
13	12	+	105487761	105487809	ELK3
14	12	+	110940578	110940626	(Similar to Mus. AF098863) ^{New}
15	12	+	110951587	110951635	(Similar to Mus. AF098863) ^{New}
16	12	+	13676162	13676210	ETV6
17	13	-	40208762	40210606	ELF1
18	17	-	45424547	45424595	ETV4
19	19	-	41606511	41606559	ETV2/ER71
20	19	-	50620314	50620753	(<i>Ensembl</i> predicted gene) ^{New}
21	19	+	51122030	51122466	(<i>ERF</i> duplication) ^{New}
22	19	-	51228203	51228639	ERF
23	19	+	62153841	62153889	SPIB
24	21	+	23995905	24000169	GABP
25	21	-	36613278	36613326	ERG
26	21	+	37052185	37052233	ETS2

Missed known genes in this family: *ETV5* (NM_004454).

Comparison with the BLAST Results

The other common method to search for new members of a gene family is to run the *BLAST* program against the whole genome using

Table 5. Comparison Results with BLAST

	GFScan	TBLASTN			BLASTN
		$E < E_m$	$E < 1e - 4$	$E < 10$	$E < 10$
A. NGIC Family ($E_m = 9e - 6$)^a					
Known member	37	37	37	37	37
Location found	38	45	48	59	33
Known location found	29	29	29	29	28
Potential candidates ^b	8	8	8	8	5
False positives ^c	1	8	11	22	0
Known location missed	8	8	8	8	9
B. CA Family ($E_m = 9e - 10$)					
Known member	14	14	14	14	14
Location found	19	19	23	38	16
Known location found	12	12	12	12	11
Potential candidates ^b	6	6	6	6	5
False positives ^c	1	1	5	20	0
Known location missed	2	2	2	2	3
C. DH-Domain Family ($E_m = 1c - 8$)					
Known member	8	8	8	8	8
Location found	9	11	16	44	5
Known location found	7	7	7	7	5
Potential candidates ^b	1	1	1	1	0
False positives ^c	1	3	8	36	0
Known location missed	1	1	1	1	3
D. ETS-Domain Family ($E_m = 1c - 10$)					
Known member	19	19	19	19	19
Location found	26	34	37	58	15
Known location found	18	18	18	18	14
Potential candidates ^b	8	8	8	8	1
False positives ^c	0	8	11	32	0
Known location missed	1	1	1	1	5

^a E_m : The minimum E -value used to find all known members by TBLASTN.

^bGenomic location that is not related to known members. The translated protein could match regular expression pattern of the gene family in the PROSITE database.

^cGenomic location where no gene family member locates (see detail in Methods).

known members' sequences as queries. We compared BLAST and GFScan on all four families. We searched the protein sequence of each known member of a given family in human genome using TBLASTN. We also used the motif region of the mRNA sequence of each known member to search the human genome using BLASTN. The results are listed in Table 5.

```

AF050105  GGCTCTGAACACCAGATCAACCATGAAGGCTTCTCCGCTGAG  GTGCGAGCTAATCCAC
AK004896  GGTTCAGACACACCGTCAATTTCAAGCCTTCCCATGGAG  CTCCACCTGATCCAC
NM_007607  GGTTCAGAGCAGATATTGATGGGAGACACTTTCGCATGGAG  ATGCACATCGTGCAC
NM_009799  GGCTCTGAGCACACCGTGGATGGAACCTAGATATTCCTGGAGAG  CTTCCACTTGTTCAC
NM_019513  GGCTCTGAGCACACCGTGGACAGTAAATGCTACCCAGCAGAG  CTTGCACCTTGGTACAT

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Motif

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GGBTCDGARCAYMNNVTBRRNNNNNNNNHHNNNNNNNGAR  VTBCABNTNRYNCAY
NM_011797  GAATCAGAGCACCAGATCAACAGTGAAGCCACGGCTGCGGGAG  CTCCACGTGGTTCAC
NM_030558  GGCTCCGAGCACAGCCTGGATGAGAAGCATggcTCTATGGAG  ATGCACATGGTCCAC
NM_024495  GGCTCAGAGCATgTGGTTAGACGGAGTGAGGTATGCTGCAGAG  CTGCATGTTGTCCAC
NM_007608  GGCTCAGAGCACCGCAGTGGACGGCCATACCTACCCAGCTGAG  CTTCCATCTgTTCatg
AF291660  GGCTCAGAGCACACAGTGGACGGCAAGTCCTCCCCAGCGAG  CTACATCTGTTTCAC
NM_009802  GGCTCTGAACACACCATTTGATGGGATCAGGgATATAATGGAG  GCTCACTTGTTTCAC

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Figure 1 The mRNA regions of 11 members of the mouse CA family identified by searching the Celera Mouse genome using the motif constructed based on CA family members in human. The first five lines were regions found by GFScan as they matched the human motif in the middle lines (R = A or G, Y = T or C, K = G or T, M = A or C, S = G or C, W = A or T, B = G or T or C, D = G or A or T, H = A or C or T, V = G or C or A, N = A or T or G or C). The bottom six lines show the other regions of the gene family that were missed by GFScan. The unmatched sites are in bold fonts. The underlined lower-case triplet represents the amino acid code that did not even match the protein motif of this family.

Table 5 indicates that GFScan had less false positives than TBLASTN (except for the CA family under a low E -value threshold, but the false positives of TBLASTN were increased when the E -value threshold was increased). In the BLASTN search, even with a very high E -value threshold (e.g., $E = 10$), some known genes were still not found, especially the ones whose motifs contain introns. For those genes, the match of the motif region to the genomic sequence is rather poor. Meanwhile, very few new genomic regions were found in this case. In short, compared with BLAST, GFScan offers both higher sensitivity and higher specificity, especially in intron-containing cases.

Mouse Genome Searching with Two Human DNA Motifs

We searched Celera's mouse genome using the motif constructed from human genes. For the neurotransmitter-gated ion-channels family, 23 of 24 known mouse members in the NCBI LocusLink Database (<http://www.ncbi.nlm.nih.gov/LocusLink/>) were found by GFScan. For the one that was missed (NM_017369: 1824–1868), the genomic DNA sequence of this gene was incomplete in the database. At the same time, 13 new motif-matching genomic locations

were found, which may correspond to 13 novel mouse members of this family.

The result was different for the CA family. For 13 known mouse CA members in the LocusLink Database, 11 had the genomic DNA sequence matches. Using GFScan and the motif constructed by human genes, we could only find five loci.

The reason for missing the other six was that the motif segments in these mouse genes are different from the motif in human genes (Fig. 1). Three of these six genes cannot even match the motif in human (NM_030558, mouse Car15; NM_009802, mouse Car6; NM_007608, mouse Car5a) at the protein level. However, two new genomic locations matching the human motif were still found, which may correspond to novel members in mouse.

In summary, GFScan is capable of identifying all the true members of a family with very few false positives and requiring no gene prediction. It performs especially well with intron-containing

Table 6. Genomic Locations with Highest Score Identified by Matrix Search

Chromosome	Motif location	Strand	Score	Index in pattern search
A. NGLC Family ($S_{\min} = 24.027^a$)				
20	63883774 63883819	-	29.4595	38
8	29298192 29298237	-	29.4595	20
X	104174415 104174460	-	29.0811	2
8	45473102 45473147	+	29.0541	21
X	13920432 13920477	+	28.8649	1
15	76752787 76752832	-	29.7568	34
17	8070258 8070655	-	28.5135	37
6	98309963 98310008	-	28.4324	19
17	5014080 5014125	-	28.4054	35
17	5277202 5277247	-	28.4054	36
2	237873656 237873701	+	28.3514	7
15	22459117 22459162	+	28.0811	28
1	1478159 1478204	-	28.0541	5
11	125455215 125455260	-	27.973	26
11	125490330 125490375	-	27.973	27
15	76721719 76721764	-	27.8919	33
4	184669254 184669299	-	27.6487	15
2	237886519 237886564	+	27.5946	8
15	76709488 76709533	+	27.5676	32
8	45498199 45498244	-	27.4324	22
X	152454639 152454684	-	27.4324	4
2	178143450 178143495	-	27.3514	6
15	24323276 24323321	+	27.1081	30
15	26565917 26565962	+	27.1081	31
5	183945827 183945872	+	27.0811	18
15	24261884 24261929	+	27.0541	29
5	174399293 174399338	-	26.7027	16
11	2590910 2590955	-	26.5946	25
4	51222053 51222098	-	26.5135	13
10	96875579 96875624	+	26.2432	24
4	50330199 50330244	-	26.2432	11
4	166964539 166964584	+	25.7027	14
5	174946361 174946406	+	25.6757	17
4	50782012 50782057	+	25.5135	12
3	104298897 104298942	-	25.1351	9
4	50085574 50085619	-	24.3243	10
X	152091806 152091851	-	24.027	3
B. CA Family ($S_{\min} = 32.6429$)				
16	77218848 77219708	+	39.2143	15
16	77672109 77672969	+	39.2143	16
4	138352740 138353600	+	39.2143	5
16	23047764 23050079	-	39.0714	12
16	33401499 33403805	-	39.0714	13
16	33986402 33988707	-	39.0714	14
8	89982297 90014490	+	38.8571	8
15	60482427 60485863	+	38.4286	11
9	38809920 38819255	+	37	10
8	89796759 89803951	+	36.9286	6
8	90013921 90014490	+	36.8572	9
17	64513194 64513368	+	36.4286	18
8	89871143 89874524	-	36.4286	7
1	9063801 9065482	+	35.8571	3
17	55604726 55621766	-	35.3571	17
3	68787364 68790131	+	34.5	4
19	57549926 57550194	-	34	19
1	230527143 230527424	-	32.6429	1
1	230570250 230570531	+	32.6429	2

^a S_{\min} is the minimum score of the motifs from known family members.

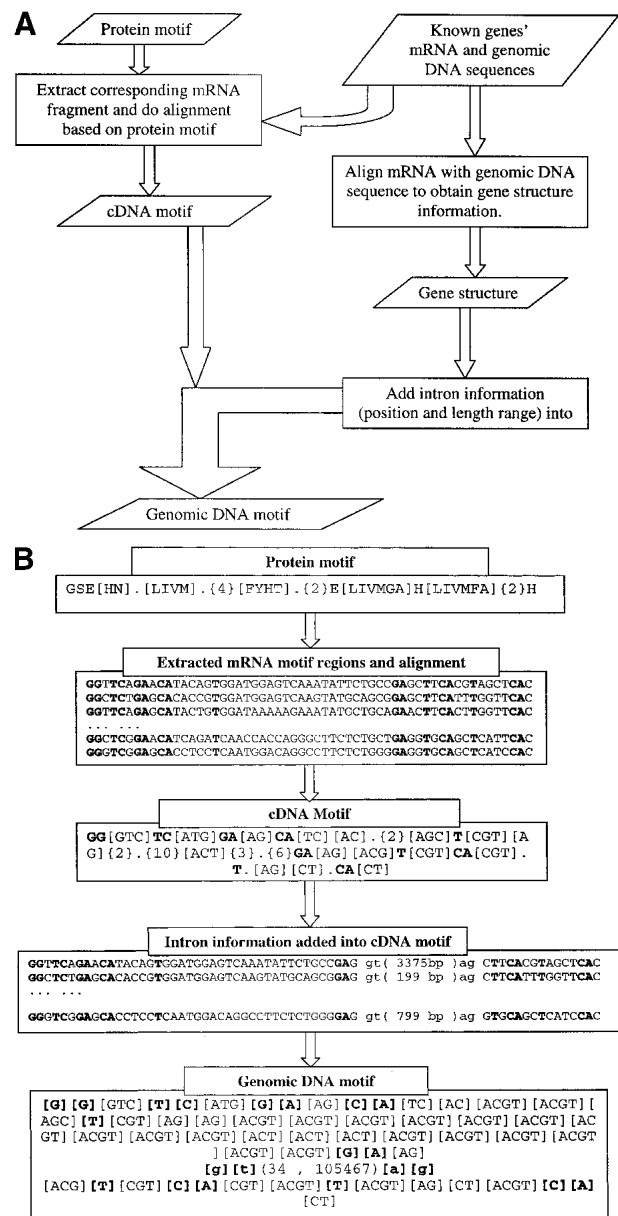


Figure 2 (A) Method to construct genomic DNA motif. Three steps were taken to construct the genomic DNA motif for a given family from known protein, mRNA, and genomic DNA sequences of the family. First, based on the locations of the protein motif in protein sequences, the corresponding mRNA regions were extracted and aligned to reveal the consensus pattern. Each site in the consensus pattern would include all nucleotides existing in the mRNAs at the site. Second, gene structures were obtained by aligning mRNA with genomic DNA sequences, and the intron information was collected. Third, the intron information was incorporated into the cDNA consensus pattern to generate the final genomic DNA motifs. (B) Motif construction example in the CA family. Conservative sites in DNA motifs are in bold font. Donors and acceptors of introns are in small letters. The number in the brackets in the DNA sequence alignments is the intron length in each gene. In the final Genomic DNA Motif, the two numbers separated by a comma in the parentheses (34 , 105467) are the minimum and maximum lengths of the intron in this position. Each pair of brackets in the DNA motif represents one site in the sequence, and the bases within each pair of brackets represent all possible nucleotides at that site.

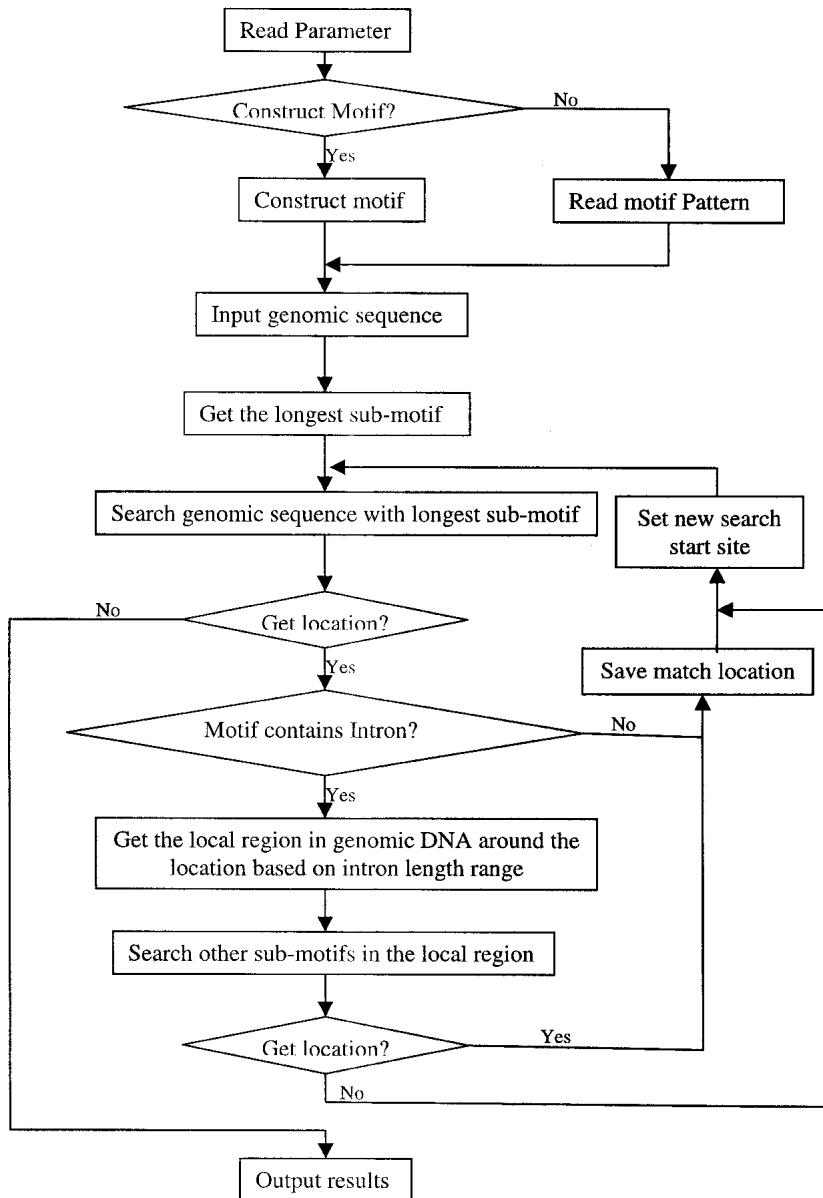


Figure 3 Flowchart of the motif search algorithm. Rectangle boxes represent steps, diamond boxes represent decision switches, and arrows show steps' order. For an intron-containing motif, the genomic DNA motif was separated into several submotifs, and the longest one was used to search the genome first. If a genomic region matches the longest submotif, this region is extracted based on intron information and the other submotifs would be only searched within this region.

motifs where most BLAST-based tools may fail. One should be cautioned when using GFScan for cross-species search, however, as the results may depend on the divergence among members of the family, as well as the evolutionary distance between the two species. By adding more mRNAs from different species or modifying a genomic motif to allow species-specific codon usages, further improvement on performance can be achieved. GFScan is implemented in a way that such customizations can be easily made (see Methods for more detail).

DISCUSSION

Same Species versus Cross-Species

As DNA sequences are usually less conserved than protein sequences in evolution, we recommend constructing motifs using known mRNAs in one species and then using the motif to search the genome of the same species. This will reduce false positives. For cross-species searches, this method sometimes worked well, as in neurotransmitter-gated ion-channels family; at other times it missed many true positives, as in the case of the CA family described above. As the program allows users to reconstruct motifs by adding more mRNAs from other species, it is easy to extend the search to the cross-species cases. One could also redefine the motif by relaxing on codon usage when searching related species or adding other conserved information into the motif.

Regular Expression Pattern Search and Weight Matrix Search

From the mRNA sequences and protein motifs of the known members of a given gene family, we could construct both a regular expression pattern and weight matrix for later searching. GFScan can use either of them to search the genomic DNA. Based on the matrix constructed, the scores of all known motif regions were calculated. When we chose the minimum score of the known motif regions as the threshold of matrix search to minimize false positives, we found that the genomic locations whose scores were higher than the threshold could all be found by a regular expression pattern search (Table 6), whereas the latter saved a lot of CPU time, because searching with regular expressions is almost 15–20 times faster than searching with matrices. However, because matrix search has higher sensitivity (at the expense of specificity and CPU time), the genomic locations missed by a regular expression pattern search may be recovered by a matrix search, especially in the cross-species cases.

Motifs

In the present program, the motif length is taken as a constant; in other words, all the motif regions in the family should have the same length. For those families whose pro-

tein motifs have variable lengths, it is difficult to construct the DNA motif, and allowing gaps in the motif can be very CPU-expensive. We will address these issues in future work.

Although *GFScan* constructs the genomic motif automatically, it also accepts user-defined motifs as its input. This makes *GFScan* a very flexible tool for gene family analysis at the genomic level. In conjunction with gene prediction tools, it can be used for gene finding and gene structure prediction as well.

METHODS

For a protein or a gene family, we collected protein, mRNA, and genomic DNA sequences of all known members, as well as the PROSITE entry. Using the protein motif in PROSITE, we extracted the protein motif fragments and their corresponding mRNA fragments. Based on the protein motif, these mRNA fragments were aligned, and the consensus pattern was created. Each site in the consensus pattern was determined from all the corresponding sites in the known mRNA sequences. In other words, each site in the protein motif was converted into three sites in the cDNA motif based on all existing codons in known mRNAs. Using *SIM4* (Florea et al. 1998) to align mRNAs with genomic DNAs, we find the potential intron position and its length range within the genomic regions that matches the motif regions. This intron information was incorporated into the cDNA motif as the genomic DNA motif of this family was constructed (see Fig. 2). For each genomic DNA motif, if there were introns inside, the motif was divided into several submotifs, and the longest submotif would be used first to find the potential match location, then the other submotifs were used to search the sequences around this location (see Fig. 3). Each genomic DNA region matching the motif would be translated into a protein sequence, and this protein fragment was tested by the protein motif to identify the false-positive results.

The weight matrix can be created while constructing the consensus regular expression pattern. In this algorithm, we simply used the nucleotide occupation frequencies at each site of the motif as the weights. For the intron-containing motif, we used the same strategy as we did in pattern search, namely, the longest submatrix was used first to find a candidate genomic location, and the local region around this location would be searched by the other submatrices.

We used protein sequences of all known members to search the human genome by *TBLASTN*, and we used the motif region of known members' mRNA sequences to search the human genome by *BLASTN*. As the exact number of the real members in a given gene family is unknown, we regarded the locations found by *GFScan* or *BLAST* false positives if the DNA fragment in these locations could not be translated into protein sequences without a stop codon, or the translated protein sequences did not match the motif pattern of the gene family. If the location is overlapped by one gene that is obviously not a member of the gene family by knowledge, the location would also be regarded as false positive. At the same time, those locations that do not code the known proteins listed in one PROSITE entry and are not false positive will be regarded as potential candidates. In *TBLASTN* search, only genomic DNA regions that could match the protein motif region partially or completely were considered as the locations of gene family members. The other genomic regions where the matches between genomic DNA sequence and protein sequence were outside of the motif were not considered. In *BLASTN* search, because the query sequences were so short that the significance of matches was low, only those genomic DNA match regions that could be aligned completely with the query sequence were regarded as the gene member's locations

to avoid many short, partial, and random matches. The Expect-value (*E*-value) was used as the threshold to filter the most significant match in *BLAST*. In our comparison, we chose different *E*-values as thresholds in *TBLASTN* searches and used the default setting in *BLASTN* (*E*-value < 10) searches. To compare the specificity with *GFScan* meaningfully, we chose the smallest *E*-value that could find all known gene members as the threshold for *TBLASTN*, then compared the new motif match locations number with that obtained from *GFScan*.

Availability

The program *GFScan* is available at <http://www.cshl.org/mzhanglab/>.

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REFERENCES

- Altschul, S.F., Gish, W., Miller, W., Myers, E.W., and Lipman, D.J. 1990. Basic local alignment search tool. *J. Mol. Biol.* **215**: 403–410.
- Altschul, S.F., Madden, T.L., Schaffer, A.A., Zhang, J., Zhang, Z., Miller, W., and Lipman, D.J. 1997. Gapped BLAST and PSI-BLAST: A new generation of protein database search programs. *Nucleic Acids Res.* **25**: 3389–3402.
- Attwood, T.K., Flower, D.R., Lewis, A.P., Mabey, J.E., Morgan, S.R., Scordis, P., Selley, J.N., and Wright, W. 1999. PRINTS prepares for the new millennium. *Nucleic Acids Res.* **27**: 220–225.
- Bateman, A., Birney, E., Durbin, R., Eddy, S.R., Finn, R.D., and Sonnhammer, E.L.L. 1999. Pfam 3.1: 1313 multiple alignments and profile HMMs match the majority of proteins. *Nucleic Acids Res.* **27**: 260–262.
- Bemark, M., Martensson, A., Liberg, D., and Leanderson, T. 1999. Spi-C, a novel Ets protein that is temporally regulated during B lymphocyte development. *J. Biol. Chem.* **274**: 10259–10267.
- Burge, C. and Karlin, S. 1997. Prediction of complete gene structure in human genomic DNA. *J. Mol. Biol.* **268**: 78–94.
- Corpet, F., Gouzy, J., and Kahn, D. 1999. Recent improvements of the ProDom database of protein domain families. *Nucleic Acids Res.* **27**: 263–267.
- Florea, L., Hartzell, G., Zhang, Z., Rubin, G.M., and Miller, W. 1998. A computer program for aligning a cDNA sequence with a genomic DNA sequence. *Genome Res.* **8**: 967–974.
- Genome Sequencing Consortium. 2001. Initial sequencing and analysis of human genome. *Nature* **409**: 860–921.
- Henikoff, J.G. and Henikoff, S. 2000. *Drosophila* genomic sequence annotation using the BLOCKS+ database. *Genome Res.* **10**: 543–546.
- Henikoff, S., Henikoff, J.G., and Pietrokovski, S. 1999. BLOCKS+: A non-redundant database of protein alignment blocks derived from multiple compilations. *Bioinformatics* **15**: 471–479.
- Hofmann, K., Bucher, P., Falquet, L., and Bairoch, A. 1999. The PROSITE database, its status in 1999. *Nucleic Acids Res.* **27**: 215–219.
- Karim, F.D., Urness, L.D., Thummel, C.S., Klemsz, M.J., McKercher, S.R., Celada, A., Van Beveren, C., Maki, R.A., Gunther, C.V., Nye, J.A., et al. 1990. The ETS-domain: A new DNA-binding motif that recognizes a purine-rich core DNA sequence. *Genes & Dev.* **4**: 1451–1453.
- Kulp, D., Haussler, D., Reese, M.G., and Eeckman, F.H. 1996. A generalized hidden Markov model for the recognition of human genes in DNA. In *Proceedings of the Fourth International Conference*

- on *Intelligent Systems for Molecular Biology* (eds. D. States et al.), pp. 134–142. AAAI Press, Menlo Park, CA.
- Pearson, W.R. and Lipman, D.J. 1988. Improved tools for biological sequence comparison. *Proc. Natl. Acad. Sci.* **85**: 2444–2448.
- Solovyev, V.V. and Salamov, A.A. 1997. The Gene-Finder computer tools for analysis of human and model organisms genome sequences. *Intell. Syst. Mol. Biol.* **5**: 294–302.
- Zhu, K.J., Debreceni, B., Bi, F., and Zheng, Y. 2001. Oligomerization of DH domain is essential for Dbl-induced transformation. *Mol. Cell. Biol.* **21**: 425–437.

WEB SITE REFERENCES

- <http://genome.ucsc.edu/>; GoldenPath.
<http://www.cshl.org/mzhanglab/>; GFScan program.
<http://www.ebi.ac.uk/interpro/>; IntroPro.
<http://www.ncbi.nlm.nih.gov/LocusLink/>; NCBI LocusLink Database.

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