



Research Article

# SAND, a new protein family: from nucleic acid to protein structure and function prediction

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## Abstract

As a result of genome, EST and cDNA sequencing projects, there are huge numbers of predicted and/or partially characterised protein sequences compared with a relatively small number of proteins with experimentally determined function and structure. Thus, there is a considerable attention focused on the accurate prediction of gene function and structure from sequence by using bioinformatics. In the course of our analysis of genomic sequence from *Fugu rubripes*, we identified a novel gene, *SAND*, with significant sequence identity to hypothetical proteins predicted in *Saccharomyces cerevisiae*, *Schizosaccharomyces pombe*, *Caenorhabditis elegans*, a *Drosophila melanogaster* gene, and mouse and human cDNAs. Here we identify a further *SAND* homologue in human and *Arabidopsis thaliana* by use of standard computational tools. We describe the genomic organisation of *SAND* in these evolutionarily divergent species and identify sequence homologues from EST database searches confirming the expression of *SAND* in over 20 different eukaryotes. We confirm the expression of two different *SAND* paralogues in mammals and determine expression of one *SAND* in other vertebrates and eukaryotes. Furthermore, we predict structural properties of *SAND*, and characterise conserved sequence motifs in this protein family. Copyright © 2001 John Wiley & Sons, Ltd.

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## Introduction

Many prokaryotic genomes have now been fully sequenced, as well as several eukaryotic genomes including: *Saccharomyces cerevisiae* (Goffeau *et al.*, 1996; *The yeast genome directory*, 1997), *Caenorhabditis elegans* (The *C.elegans* sequencing consortium, 1998), *Drosophila melanogaster* (Adams *et al.*, 1999) and that of the first plant genome *Arabidopsis thaliana* (Lin *et al.*, 1999; Mayer *et al.*, 1999; Salanoubat *et al.*, 2000; Tabata *et al.*, 2000; Theologis *et al.*, 2000). The first complete vertebrate genomic sequence, that of man, is available but at this time is not fully assembled (Venter *et al.* 2001; International human genome sequencing consortium, 2001). Many gaps in the sequence remain to be filled and only two chromosomes are in a finished state (Dunham *et al.*, 1999; Hattori *et al.*,

2000). It is now becoming apparent that the annotation of these genomic sequences relies heavily on gene prediction programs, of which the best only fare reasonably. Continuous sequence similarity searches of large genomic sequences against an increasing pool of cDNAs and ESTs is proving to be costly. A feature of the analysis of these genomes is the number of unknown proteins that they are predicted to encode. For example, in the relatively small genome of *C. elegans*, 60% of predicted genes encoded proteins of unknown function (The *C. elegans* sequencing consortium, 1998). In this study, we have identified a novel gene in the model vertebrate *Fugu rubripes*, which we have called *SAND*, as its location is next to the plasminogen related growth factor receptor (PRGFR) thought to be the orthologue of SEA (EMBL:AJ010317) (Cottage *et al.*, 1999). BLAST (Altschul *et al.*, 1997)

searches of cDNA and EST sequences have revealed that the *SAND* gene product is expressed in diverse eukaryotes, in various developmental stages in plants, and in many tissue types in vertebrates. We have identified two *SAND* paralogues in mammals, which appear to be the result of a taxonomy class specific duplication. In an attempt to assign function to this new protein family, we have carried out a computational analysis including similarity searches, multiple sequence alignments, domain identification and characterisation, secondary structure predictions, solvent accessibility predictions and automatic protein fold recognition analyses.

## Materials and methods

### Characterisation of the *F. rubripes* cosmid 165K09

The *F. rubripes* cosmid 165K09 (EMBL: AJ010317) had previously been identified as encoding a PRGFR. The cosmid was sequenced and annotated as reported (Cottage *et al.*, 1999). Additionally, RT PCR was used to verify the predicted *SAND* gene product.

### Bioinformatic analysis of *SAND* nucleic acid sequences

BLASTn (version 2.0.12) and tBLASTx (version 2.0.12) (Altschul *et al.*, 1997) searches of EMBL release 66 (EMBL) (Stoesser *et al.*, 2001) (and unfinished human genome contigs (<http://www.ncbi.nlm.nih.gov/BLAST/>) and BLASTx (version 2.0.12) (Altschul *et al.*, 1997) searches of SWISS-PROT release 39 (SP) and SWISS-PROT TrEMBL release 16 (SPTR) (Bairoch and Apweiler, 2000) with *F. rubripes SAND* were used to identify homologous sequences. Where gene sequences had not been identified, these were determined from genomic sequences using NIX, a WWW tool to view the results of many DNA analysis programs (Williams, G., Woollard, P. and Hingamp P. unpublished data, <http://www.hgmp.mrc.ac.uk/Registered/Webapp/nix/>). Analysis of the promoter regions, in order to determine potential TATA boxes and conserved transcription factor binding sites, was facilitated by the use of Theatre. Theatre is a tool for obtaining the results for a set of DNA sequences, from various DNA sequence analysis programs (with emphasis of finding transcription factor binding

sites), and producing a clear graphical display of this information in a comparative context (Edwards and coworkers; <http://www.hgmp.mrc.ac.uk/Registered/Webapp/theatre/>). tBLASTn version 2.0.12 (Altschul *et al.*, 1997) searches of EST databases were used to generate an *in silico* expression profile of *SAND* homologues.

### Bioinformatic analysis of *SAND* protein sequences

When gene sequences had been determined, the derived protein sequences were aligned, together with the mRNA and EST translations using ClustalW version 1.74 (Thompson *et al.*, 1994). Percentage sequence identity and specific conserved residues were determined by the use of Belvu, an alignment viewer written by Erik Sonnhammer (<http://www.cgr.ki.se/cgr/groups/sonnhammer/Belvu.html>). Composite analysis of each protein was performed using PIX, a WWW tool to view the results of many protein analysis programs for a query sequence (<http://www.hgmp.mrc.ac.uk/Registered/Webapp/pix/>). Further database searches using PSI BLAST version 2 (Altschul *et al.*, 1997) were used to identify potential domains by accepting the default and running several iterations. *SAND* was also submitted for PSI BLAST analysis in overlapping contigs of 100 residues. A consensus secondary structure and associated solvent accessibility was generated using Jpred2 (Cuff *et al.*, 1998) from a ClustalW (Thompson *et al.*, 1994) alignment of seven full length *SAND* sequences in MSF format. The protein fold recognition program Threader (Jones *et al.*, 1992) was used to score protein sequence compatibility against known protein folds. Sequence threading against a structural databank of 1902 known protein folds was performed for the seven *SAND* sequences. Threadings were computed in terms of 1. pairwise interaction energies, 2. solvation potential energies and 3. their weighted sum, in order to evaluate the fit of each sand sequence to a particular fold conformation, and represented as Z-scores ( $Z = (\text{Energy} - \text{Mean}) / \text{Standard Deviation}$ ). Provided there is greater than 50% sequence and structure matching, the Z scores were sorted for input into a program called SumThreader (Edwards and Perkins, 1996) in order to summarise the outcome of the searches. For each of the three Z scores, the average rank of each fold was calculated from the seven values determined for the individual *SAND* sequence

threadings. The average ranked position for each fold for seven sequences were calculated.

## Results and discussion

### Identifying sequence homologues

The Japanese puffer fish, *F. rubripes*, was chosen as a suitable vertebrate model in which to identify genes of interest. Evidence suggests that the *F. rubripes* gene complement is about the same as that of mammals, and that gene order and structure may be preserved, but within a remarkably compact genome of only 400 Mbp (Brenner *et al.*, 1993). Several gene families have been studied in this vertebrate, including the *Hox* genes (Aparicio *et al.*, 1997), the pheromone receptors (Naito *et al.*, 1998) and the surfait genes (Armes *et al.*, 1997). Analysis of the genomic sequence from the *F. rubripes* cosmid 165K09 (EMBL:AJ010317) identified a novel sequence, *SAND* (Fr\_SAND) (SPTR:Q9YGN1), with sequence identity to hypothetical proteins predicted in *S. cerevisiae* (Sc\_SAND) (SP:P53129) and *Sz. pombe* (Sp\_SAND) (SP:Q10150), and to part of a hypothetical protein predicted in *C. elegans* (Ce\_SAND) (SPTR:Q20298). The *F. rubripes* sequence also has sequence identity to the *D. melanogaster* CG11926 gene product (Dm\_SAND) (SPTR:Q9VR38) and a mouse cDNA clone (Mm\_SAND1) (EMBL:AK013387). tBLASTx searches of *A. thaliana*'s chromosome 2 sequences identified a sequence with similarity to SAND (At\_SAND) (EMBL:AC006283). Two human SAND homologues were identified: a cDNA clone KIAA0872 (Hs\_SAND2) (SPTR:O94949), and a predicted open reading frame on chromosome 3p21 (Hs\_SAND1) (EMBL:AC068701). Whilst BLAST (Altschul *et al.*, 1997) searches reveal the expression of at least two SAND genes in a number of mammals (see Table 1A and 1B), including human, mouse, pig and cow, only one SAND gene can be detected in any non-mammalian organism (see Table 1C).

### Intron/exon structure

*SAND1* is predicted to comprise two exons in *Sz. pombe* (see Table 2A) and one in *S. cerevisiae*, however the hypothetical protein produced from the reported *S. cerevisiae* prediction introduces gaps into an alignment with other SAND proteins. The *S. cerevisiae* sequence is approximately 644 residues

by comparison to the 520 residues encoded by the other sequences. A better alignment with the other SAND sequences was achieved by removing 54 residues from the N terminus and introducing an intron which removes residues 537–577 (SP:P53129), experimental confirmation is needed of this prediction and this sequence was not used for secondary structure predictions.

Gene sequences were determined from the *C. elegans* genomic data by Genefinder (P. Green and L. Hillier unpublished software). These predictions were confirmed or adjusted to account for protein, cDNA and EST matches (The *C. elegans* sequencing consortium, 1998). The *C. elegans* SAND

Table 1. Expression profile of SAND generated by EST database searches

#### IA. Expression profile of SAND1 in mammals

Organism	EMBL Accession number	Expression
<i>Homo sapiens</i>	CNSL1EA9	Placenta
	CNSL1FVF BE793894	T cells/T cell leukemia Lung
<i>Mus musculus</i>	BF321829	Mammary infiltrating ductal carcinoma
	BF179283 BF584835	Mammary, gross tissue Colon
	AI853696 AU06752	Brain Brain
<i>Rattus norvegicus</i>	AI237869 BE111711	Normalized spleen Cardiac tissue
	<i>Sus scrofa</i>	BE234656 BF193484
<i>Bos taurus</i>	AW655266 AW653632 BF664794	Pooled Pooled Pooled

#### IB. Expression profile of SAND2 in mammals

Organism	EMBL Accession number	Expression	
<i>Homo sapiens</i>	BE785702 BF313087 BE885699 CNSLTOYVF	Lung Neuroblastoma Leiomyosarcoma T cell leukemia	
	<i>Mus musculus</i>	AK023374 AW626447 AW322902	Ovarian tumour Lung Lung
	<i>Sus scrofa</i>	AW785756 AW322902	Pooled Pooled
		<i>Bos taurus</i>	BF602206

### 1C. Expression profile of SAND in non mammalian eukaryotes

Organism	EMBL accession number	Expression
<i>Gallus gallus</i>	GGA395913	Bursa of fabricus
	GGA392697	Bursa of fabricus
<i>Danio rerio</i>	AW343114	Mixed tissue
	AW115566	Mixed tissue
<i>Drosophila melanogaster</i>	AI294091	Larvae
	AI516700	Embryo 0–24 hrs
	AI389145	Head
<i>Caenorhabditis elegans</i>	CEI2C12	Mixed
	AU111676	Whole
	AU112292	Whole
<i>Dictyostelium discoideum</i>	DDC4407	Gamete
	DDC4408	Gamete
<i>Arabidopsis thaliana</i>	BE522449	Developing seeds
	AV536086	Flower buds
	AV555362	Green siliques
<i>Glycine max</i>	AW201401	Cotyledons
<i>Lycopersicon esculentum</i>	AI777002	Callus
	AW222181	Pericarp
	AW222182	Fruit
	AI487936	Ovary
<i>Medicago truncatula</i>	CN506BLP	Arbuscular mycorrhiza
<i>Solanum Tuberosum</i>	BF459706	Tuber
<i>Pinus taeda</i>	BG318737	Xylem
<i>Secale cereale</i>	BE588169	Root tip
<i>Gossypium hirsutum</i>	AW187646	Cotton fibre
<i>Hordeum vulgare</i>	BG300416	Seedling shoot

protein (F41H10.4 SP:Q20298) is predicted by Genefinder to be encoded by 13 exons; however it would appear that exons 1–4 belong to a neighbouring gene and it is this sequence that shows similarity to a *Plasmodium yoelii* rhopty protein as specified in the annotation. Residues encoded by exon 8 introduced a gap in an alignment with other SAND proteins, so this was removed from the gene prediction. We therefore expected that *C. elegans* SAND is encoded by 6 exons, although the exact gene start could not be determined from the given sequence, possibly due to sequencing/assembly errors and so is not shown in Figure 1.

The predicted gene structure of *D. melanogaster* did not require further adjustment. The *D. melanogaster* genome annotation used Genscan (Burge and Karlin, 1997), and a version of Genie that uses expressed sequence tag data (Reese *et al.*, 2000), plus the results of cDNA and protein database searches, followed by review by human annotators (Adams *et al.*, 2000). *D. melanogaster* SAND is encoded by

three exons. In contrast, SAND is encoded by 12 exons in *A. thaliana*; the genes small intron and exon sizes, coupled with low G+C content (42%), may explain why this gene was missed by the gene prediction programs used in this genomic project, even though the annotation involved both DNA and protein database searches and gene prediction with the programs Genscan (Burge and Karlin, 1997), Genefinder (P. Green and Hillier unpublished) and Grail (Uberbacher *et al.*, 1991).

Human and mouse SANDs have the highest G+C content (>60%), followed by *F. rubripes* (53%) and *D. melanogaster*. *C. elegans*, both yeast and *A. thaliana* all have a lower G+C composite at

**Table 2.** Comparison of intron/exon sizes and exon phase between SAND sequences in *A. thaliana* (At), *C. elegans* (Ce), *F. rubripes* (Fr), Human (Hs), *D. melanogaster* (Dm), *S. cerevisiae* (Sc) and *S. pombe* (Sp). The question mark indicates unresolved or ambiguous boundaries in gene annotation

#### 2A. Size of exons in base pairs

Ex.	At	Ce	Fr	Hs1	Hs2	Dm	Sc	Sp
1	186	155?	124	127	148	593	1449?	1295
2	101	213	384	486	327	152	204	247
3	106?	155	766	766	820	842		
4	72	371	148	148	148			
5	166	407	143	141	201			
6	122	277						
7	233							
8	83							
9	263							
10	92							
11	65							
12	96							

#### 2B. Size of introns in base pairs

Ex.	At	Ce	Fr	Hs1	Hs2	Dm	Sc	Sp
A	57	269?	146	1208	1818	66	120	86
B	66	58	89	637	557	56		
C	122	298	129	718	380			
D	249	50	439	98	2425			
E	96	51						
F	366							
G	162							
H	251							
I	81							
J	90							
K	103							

## 2C. Phase of introns

At	Ce	Fr	HsI	Hs2	Dm	Sc	Sp
0						0	
	0	0	0	0			
0							
0							
0							

around 40%. Analysis of the promoter regions by the use of Theatre (Edwards and coworkers; unpublished material) determined that the promoters appear to be largely TATA-less; no significant sequence identity could be determined between them. Comparisons between the promoter regions proved difficult due to the variable length of the N terminus of the encoded proteins and the lack of sequence identity between them across this region.

Both intron/exon number and size, as well as intron phase, show no conservation between yeasts, fly, worm and plant (see Table 2). Analyses of *SAND* in these organisms may elucidate some interesting clues to the dynamics of intron formation/loss. Vertebrate evolution, however, appears to have taken a more concerted route: *F. rubripes* *SAND* and human *SAND1* and *SAND2* are all encoded in five exons, and the intron phases of the three genes are conserved (see Table 2C). Exon 4 is the same size in all three genes *Fr\_SAND* (148 bp), *Hs\_SAND1* (148 bp) and *Hs\_SAND2* (148 bp) (see Table 2A).

## Genomic loci

The orientation of genes neighbouring *SAND* and their accession numbers are given in Table 3. Human *SAND2* protein KIAA0872 has been mapped to chromosome 16 (Nagase *et al.*, 1998). BLAST (Altschul *et al.*, 1997) searches of the human *SAND2* locus (EMBL:AC009139) have shown its 5' neighbour on the direct strand to be a putative pheromone receptor, whilst its 3' neighbour on the reverse strand encodes a hypothetical protein. Hypothetical proteins are also predicted as neighbours to *SAND* in *D. melanogaster*, *C. elegans*, and *A. thaliana*, but these sequences do

not appear to be homologous. Of interest are the ribosomal proteins predicted 5' to *SAND* and on the opposite strand in both *D. melanogaster* and *S. cerevisiae*. The 5' neighbour (on the reverse strand) of *F. rubripes* *SAND* is the TRAF interacting protein TRIP. The 3' neighbour, on the direct strand, is the plasminogen related growth factor receptor (PRGFR3). This gene organisation is conserved on the genomic segment of chromosome 3p21 encoding human *SAND1*. Human TRIP (SPTR:O00467) is on the reverse strand 5' of *SAND1*, and the plasminogen related growth factor receptor (SP:Q04912) is approximately 10Kbp on the direct strand 3' to it. Considerable interest has been focused on 3p21 in man as a cancer hot spot; it is particularly implicated in all major types of lung cancer (Kok *et al.*, 1987).

## Expression profile

Expression of the human *SAND* protein KIAA0872 has been assayed by means of RT PCR from mRNA derived from heart, brain, lung, liver, smooth muscle, kidney, pancreas, spleen, testis and ovary. KIAA0872 was shown to be expressed predominately in brain, kidney and ovary (Nagase *et al.*, 1998). In this study, RT PCR analysis confirmed expression of *F. rubripes* *SAND* in brain, kidney and ovary, reflecting the expression profile seen in human. Although some expression was seen in other tissues assayed such as heart, lung, liver, smooth muscle, spleen and testis. An expression profile was generated for the *SAND* transcripts by BLAST (Altschul *et al.*, 1997) searches with all identified *SAND* sequences and is shown in Tables 1A, 1B and 1C. Human *SAND2* (KIA00872) had been identified in pooled tissues as well as lung. The transcript had also been identified in four neoplastic tissues, of particular interest is the lack of exon 3 from EMBL:AK023374 identified from an ovarian tumour. Expression of *Hs\_SAND1* appears more widespread and includes neurological, lymphatic and cardiac tissue, whilst once again neoplastic tissues are also represented. The expression of *SAND* in other eukaryotes appears to occur at various developmental stages, and even in specialised tissues such as the bursa of fabricus in chicken. In plants, expression of *SAND* is seen in both monocots and dicots, and is even to be found expressed in the xylem of pine. *SAND* is expressed in many different tissue types in plants, including: flowers, seeds, tubers, leaves, shoots and roots.

HS\_SAND1 MATDMQRKRSSSECLDGLTLPDQGSMEARASPTPGMAQGMPEAGQEGAMFVHARSYEDLSEDEGAAAGSDHKEGTRGPPPLPTDMRQI
MM\_SAND1 MAADMQRKRSSSECEPGLTLPSPNGQSVRAESPTPGLTQCTE PGAGQEGAMFVHARSYEDLSEDEGAAAGSDHKEGTRGPPPLPTDMRQI
FR\_SAND MDGEAQN--SVACEKATLAPVDRLRNRAESPTPGIVEGTEPGAVQKSAFFAQAQSFEDLTAEEAEKAEQEGVVEDSG--
HS\_SAND2 -----MEVGGDTAAPAGGAELEDTQFP--SEEAREGGVHAVPPDEDEGLEETGSKDKDQPPSPSP-----
DM\_SAND -----MEVQTSVRSNTNCEYLDAEGDPESPNLYQEADPDQAEQONHSI I SELRDGLTMRDNSALSPEPG-----
AT\_SAND -----MEAGEPPSPSTSSYDGERGSSGGATSTYKADGSEDEITREANVDGDTASQHEAAWLPGRKRFKHLL-----
SP\_SAND -----MEFTSEHSSIKEEVENDNVHRSHSECEGSLLLNPGNVLMAAPSVSEDDQEVSRSTPELRSHVENVEQ-----
Secondary
Access.

(L1)

HS\_SAND1 SQDFSELSTQLTGVARDLQEEMLPGSSEDDWLEPPGAVGR--PATEPPREGTTEGDEEDATEAWRLHOKHVFVLSSEAGKPVYSRYGSEEALS
MM\_SAND1 SQDFSELSTQLTGVARDLQEEMLPGSSEDDWLEPPGAAAGRAATEPSEQEGSTEGEEDATEAWRLHOKHVFVLSSEAGKPVYSRYGSEEALS
FR\_SAND --QTEDELQVLVG--ERTITEQHAQDVS-----PQSRKSE--EDMSSEAWRSRHKHVFVLSSEAGKPVYSRYGSEEALS
HS\_SAND2 --PQSEALSSTSRVWSPAAAPENSPTCS-----PESSSGGGGDPDEEWSRQKHKHVFVLSSEAGKPVYSRYGSEALS
DM\_SAND --QENKGLAASVESLALSTSTSAKTEDS-----IGGLEEEYDYQHDLSWQOQKHKHVFVLSSEAGKPVYSRHLGNEKCLA
AT\_SAND --VLSLLVFSFRHGLYGFSRNAVEN-----VEHIENMKTQDDASTSWRKRKHKHVFVLSSEAGKPVYSRYGDEHKLA
SP\_SAND --LLSDILHDNSSLPLMVSTSVSSSSNNT-----AVDEIIKLLSLISFDLAKQKRTYLFSSSGKPVYSNIVDDSIETP
Consensus
Secondary
Access.

(A1) L2 (B1) L3 (B2) L4

HS\_SAND1 STMGVMMALVSVLEADKN-----AIRSIHADGYKVVVFRSRPLVAVAV--RTQSAQELAQELLYIYQILSLTGAQLSHIFQOKQ
MM\_SAND1 STMGVMMALVSVLEADKN-----AIRSIHADGYKVVVFRSRPLVAVAV--RTQSAQELAQELLYIYQILSLTGAQLSHIFQOKQ
FR\_SAND STMGVMMALVSVLEADKN-----AIRSIHADGYKVVVFRSRPLVAVAV--RTQSAQELAQELLYIYQILSLTGAQLSHIFQOKQ
HS\_SAND2 ATMGVMMALVSVVQAGD-----AIRAIYAEDHKLVFLQGGPPLLVA--RTSQSAQELRQELLAHVHAQYVSTLTRASVARIFAHKQ
DM\_SAND TLFQVTVQALVSVVQAGD-----ATTSIHAGGKVFQFQKRSLLVAVAS--RSNMSVQQLQQLGQVYVQILSLITYSMTKIFERRK
AT\_SAND GFSATLQAILISFVENGKITALKKITIIILKMRFPQGIQVFLVQKGPVLYVLCIS--CTDETYEYLRGQLDLYGQMLLITLTKS--IDRCFEKNA
SP\_SAND STVGALQAILISFVSKS-----ELTSTFSTFNVVLSKPNLYLVGVSFSTLSAAYLSELENLYYQILSLITYSMTKIFERRK
Consensus
Secondary
Access.

(A2) L5 (B3) L6 (B4) L7 (B5) L8 (A3)

HS\_SAND1 NYDLRRLLSGSERTIDNLLQL----MARDP--SFLMGAARCLPLAAVAVRTVSASLQO--ARARSVLSVILLARNQVALVVRK----D
MM\_SAND1 NYDLRRLLSGSERTIDNLLQL----MARDP--SFLMGAARCLPLAAVAVRTVSASLQO--ARARSVLSVILLARNQVALVVRK----D
FR\_SAND NYDLRRLLSGSEYLTIDNLLHR----LDRDP--GLLSAVTCLPLNSVDRVSSVSSLOA--AKAKNLSVILLAGDLRVLTVRKK----D
HS\_SAND2 NYDLRRLLSGSERTLDRLDLS----MEQDPGALLLGAVRCVPLARPLRDLGALLR--CTAPGLALSVLAVGCRVLTAAQERNVLAE
DM\_SAND NYDLRRLLSGSERTLFYLLANDSSAKVSNNIFFLTNSIRVFPPTTIRSQITSAIQSNCSKIKNLVAVLITANNKLTALVRMK----K
AT\_SAND KFDNTPPLGGTDAVSSLVHS----FSWNP--ATFLHAYTCLPLPYALROATGTILQE--VCASGVLSLIMCRHKVKNPCRKG----K
SP\_SAND NYDLRRLLSGSEYLFKELCDQ----LNDYELVPTLNATISPLPLRSSFRDQLSOLLR--ETPKLLETFIATRGRLVCMVKAK----Q
Consensus
Secondary
Access.

L9 (A4) L10 (B6) L11 (A5) L12 (A6) L13

HS\_SAND1 QFLHPIDLHLLFNLISSSSSFR--GEAWTPVCLPKFNAAGFFFAHISYLEPDTDLCLLLVLS--TDREDFAVSDCRRRFQERLRKRGAHLA
MM\_SAND1 QFLHPIDLHLLFNLISSSSSFR--GEAWTPVCLPKFSAAGFFFAHISYLEPDTDLCLLLVLS--TDREDFAVSDCRRRFQERLRKRGTHLA
FR\_SAND QFLHPIDLHLLMNLVGSSSSFR--GEGWTPICLPKFNATAGFFFAHISYLESADLCLLLVLS--TDREDFNMSDCKQRFLERLTKRTAYQA
HS\_SAND2 CRLDPADLRLLLDWWG--APAFAA--GEAWAPVCLPRFNPDPGFYAYVARLD--AMPVCLLLG--TQREAFHAMAACRRLVEDGPHALGAMRA
DM\_SAND YSIHPADLRLIINLVCESESEFKS--SENNSPICLPKFDMMNGYLVHAHVSYLADDCQACLLVLS--VDRDAFFTAEAKAKITFKLRKSHCLEA
AT\_SAND SFSAWSKIIVNDRVYVQVSVL--AESFSPICLPRYNAQAFHAYVHFVFDVSQLAVLLKSNILSVVQRSAEAGGMVDEVPIDRRRRSS
SP\_SAND LLLHANDLYLLELSLFTQSFNDMSMEHWVVPVCFPTLNPDAYIYIYSYFLC--KDTV--LTMGS--SESGVFFEMQSVKCVQAQEQDHDGWLKK
Consensus
Secondary
Access.

(A7) L14 (A8) L15 (B7) L16 (A9) L17 (B8) L18 (A10) L19

HS\_SAND1 LREAL-----RTPYYSVAQVGIPIDLRHFLYK-----SKSSGLFTSPEIEAPYTSSEEQERLLGLYQVLSHRAHNA--SRPLKTIYY
MM\_SAND1 LREAL-----RTPYYSVAQVGIPIDLRHFLYK-----SKSSGLFTSPEIEAPYSSEEQERLLGLYQVLSHRAHNA--SRPLKTIYY
FR\_SAND LKEAL-----KCPYSVEQVGIPIELRHFLYK-----SKSSGLYTSPEFPELYQSDDEEQERLMGLYQDLHSHLHP--VRPLRFYR
HS\_SAND2 LGEAASFSNASSAPAYSQAVGAPCLRHFLYKPLDIPDHRQLPQFTSPELEAPYSREERQERLSDLYHRLHARLHST--SRPLRLIYH
DM\_SAND INEELQOP-----FNAKLYQVQVGIPIELRHFLYK-----PKSTAQLCLPMLRHPYKSLTELELEAITYCDLHRLHNS--SRPLKLIYH
AT\_SAND TTNQEQDS-----PGPDISVGTGGPFGLWHFMRYR-----SIYLDQYISSEFSPPVTSRQOKRLRYRAYQKLYASMHVKGLGPHKTQYR
SP\_SAND LIYCEEMD-----RTTPRNPGSPICLSHYLFY-----SKKYSQFYTPGYSFSTPNF--NTRTLVYAYASLHDQAFHK--KNSFNSIMT
Consensus
Secondary
Access.

(A11) L20 (B9) L21 (A12) L22 (B10)

HS\_SAND1 TGNPNENLLAWVTGAFELYMYSPLGTKASAVSAIHKLMRWIRKEEDRLFILTPITY-----
MM\_SAND1 TGNPNENLLAWVTGAFELYMYSPLGTKASAVSAIHKLMRWIRKEEDRLFILTPITY-----
FR\_SAND CTNKENLLAWVTNGFQYLYCPSPLGTKAMAVSAVNIKLLRWIRKEEDRLFILTPITY-----
HS\_SAND2 VAEKETLLAWVTSKFELYTCLSPLVTKAGAILVVTIKLLRWVKEEDRLFIRYPPKYSTPPATSTDQAAHNGLFTGL
DM\_SAND MKEREVLLAWATGTIYELYAIFEPVVDKATVIKYVDKLIKRWIEKEYDYVYFTRNHATF
AT\_SAND RDENYTLVAVTDFELYAAAFDPLADKAMAKI CNQVQORVQVENEVFLQAGSPFSW-----
SP\_SAND VHESLLFTWSTASDFDHCIANATTSSQLLIANVNIKLLRWIRREENRLFITQNLFS
Consensus
Secondary
Access.

L23 (B11) L24 (B12) L25 (A13) L26 (B13) L27

Table 3. Characterisation of SAND loci

5'			3'	
	Accession number	Identification	Accession number	Identification
Fr_SAND	SPTR:Q9YGN2	TRIP (-)	SPTR:Q9YGN0	PRGFR3 (+)
Dm_SAND	SPTR:Q9VR37	Ribosomal protein S2 (-)	SPTR:Q9VR39	Hypothetical protein (+)
Ce_SAND	SPTR:Q20297	Hypothetical protein (+)	SPTR:Q20299	Similarity to GRM5 (+)
Sc_SAND	SP:P25443	Ribosomal protein S4 (-)	SP:P53128	MetI (-)
Sp_SAND	SPTR:Q10147	Probable T complex protein (+)	EMBL:Z69239.1	p65 (+)
At_SAND	SPTR:AAD20686	Hypothetical protein (+)	SPTR:AAD20658	Hypothetical protein (+)
Hs_SAND2	EM:AC009139	Probable pheromone receptor (+)	EM:AC009139	Hypothetical protein(-)
Hs_SAND1	SPTR:O00467	TRIP (-)	SP:Q04912	MSP Receptor RON (+)

The orientation of the gene neighbouring SAND is given in parenthesis, (+) for the leading strand and (-) for the reverse strand.

### Protein structure and function

Inferring function from sequence data is the new dogma in DNA research, and the main focus of molecular biologists and bioinformaticians following in the wake of the large genomic projects. Despite current innovations in bioinformatics, this endeavour is non trivial and requires validation.

### Search for a structural homologue using basic search methods

In order to determine if SAND has sequence similarity to proteins with known 3D structures, BLASTP (version 2.0.12) (Altschul *et al.*, 1997) was used to search NRL-3D (<http://pir.georgetown.edu/pirwww/dbinfo/nrl3d.html>), a databank of protein sequences whose structures have been experimentally determined (Garavelli *et al.*, 2001). No structural homologues were identified by searches with the seven full SAND sequences reported in this communication.

PSI-BLAST (Altschul *et al.*, 1997) – Position Specific Iterated BLAST uses an iterative search in which sequences identified in the first round of searching are used to build a score model for the next round. PSI BLAST has been used to successfully identify homologous protein sequences that have known 3D structures, even when the query

and subject sequences have less than 20% sequence identity overall (Bork *et al.*, 1999). Successive PSI BLAST 2 iterations with the seven full length SAND protein sequences resulted in convergence after the 4<sup>th</sup> iteration.

### Search against the databank of Pfam profiles

Profile Hidden Markov Models (HMMs) built from Pfam alignments can be useful for automatically recognising that a new protein contains an existing protein domain, even if the sequence similarity is weak. Pfam HMMs (Sonnhammer *et al.*, 1998) were searched with the nine SAND sequences using the search package HMMER; no significant matches were reported. A seed alignment of SAND is available in PfamB (pfam b8448) but only contains 5 partial sequences from *H. sapiens* (SAND2), *F. rubripes*, *C. elegans*, *Sz. pombe* and *S. cerevisiae*. No homologous structural domains are reported for these proteins.

### PIX analysis of the nine SAND protein sequences

Composite analysis of the SAND proteins for protein motifs and structural elements was facilitated by using PIX (<http://www.hgmp.mrc.ac.uk/Registered/Webapp/pix/>). Matches to various

**Figure 1.** A ClustalW alignment of seven full length SAND amino acid sequences. The following abbreviations are used Hs (*H. sapiens*), Fr (*F. rubripes*), Dm (*D. melanogaster*), At (*A. thaliana*) Mm (*Mus musculus*) and Sp (*S. pombe*). Shown below the alignment is the Jpred2 consensus secondary structure prediction (H – helix, E – strand, – loop) and solvent accessibility (b – buried, e – exposed). Shown in bold underlined type are two cysteines located on helix A10 and strand B12, they are present only in vertebrates and yeast, they may form a disulphide bridge in the mature peptide. SAND has 13 conserved sequence motifs. Motif 3 contains 4 and 6 hydrophobic residues which coincide with beta strands B4 and B5. Motif 5 contains the conserved residues NYDLRRL encoded on loop L9 which are also present in Ipii. Motif 13 contains charged C terminal residues predicted on loop L26

features, for example transmembrane domains, coiled coils, signal peptides and peptide cleavage sites were reported in individual sequences, but the threshold at which these features were determined was not significant. Furthermore, these features were not present in the majority of the sequences. With the exception of *A. thaliana*, PSORT KNN predictions only marginally predict SAND as a nuclear localised protein ahead of the prediction as a cytoplasmic protein. The *A. thaliana* SAND is predicted as a plasma membrane located protein with a score of 42%. There were no significant matches to domain databases, e.g. SBASE (Murvai *et al.*, 2001), ProDom (Corpet *et al.*, 1998), BLOCKS (Henikoff *et al.*, 1999), PRINTS (Attwood *et al.*, 1997) and PROSITE (Hofmann *et al.*, 1999). Although comprehensive protein database searches have not revealed a possible function for the SAND proteins, they firmly establish SAND as a new protein family.

#### Alignment and prediction of secondary structure and solvent accessibility

The *C. elegans* and *S. cerevisiae* SAND sequences were not included in the alignment (Figure 1) due to problems with their annotations as outlined in this manuscript. The alignment comprises seven sequences that have low sequence identity overall, this typically being less than 30% between pairs of sequences. There is no shared sequence similarity at their N terminus (sequence lengths vary between 104–151 residues). However, following the N terminus, several regions of highly conserved residues are apparent (see Figure 1). These provide evidence for at least two domains within the SAND proteins.

Jpred2 was used to predict the secondary structure and solvent accessibility of the SAND proteins. Three types of input can be supplied to Jpred; a single protein sequence, an unaligned set of protein sequences, or a multiple protein sequence aligned in MSF format. The ClustalW alignment shown in Figure 1 was used as input to Jpred. The Jpred2 consensus secondary structure and solvent accessibility predictions are also shown in Figure 1. The N terminal region described previously is predicted not to contain either alpha helices or beta strands. The SAND proteins are predicted to contain thirteen alpha helices and thirteen beta strands, plus a total of 27 loops. All beta strands are predicted to be largely solvent inaccessible, as well as four helices A2, A7,

A8 and A9. Eight of the thirteen helices display an amphipathic pattern (A3, A4, A5, A6, A10, A11, A12 and A13); these helices are likely to be located on the outer surface of the protein with one side of the helix facing the solvent and the other the hydrophobic interior. The lengths of secondary structure elements predicted are typical of those observed in known protein structures and A3 is the only helix predicted to be over 20 residues. Given that the proteins share a low level of sequence identity, the accuracy of the prediction will be affected. The Jpred secondary structure prediction is expected to be about 70% accurate at the amino-acid residue level (Cuff *et al.*, 1998).

#### Automatic protein fold recognition

The identification of a protein fold was attempted using Threader (Jones *et al.*, 1992). SumThreader (Edwards and Perkins, 1996) was used to summarise the outputs. The sequences in Figure 1 minus the N terminal residues with the absence of regular secondary structure predicted were used as input for threading analysis.

The SAND sequences were matched 'favourably' with protein structures (protein structure codes and domain annotations are given and the average Z score are given in parentheses): 1pii00(2.65), 1pdz02(2.65), 1eedP0(2.62), 2exo00 (2.56), 1mpp00 (2.55) and 1hpm00(2.52). The Z scores of these matches approach the threshold score of 2.7. In the Threader user guide, this score is specified as "borderline significant, possibly correct". 1pii consists of two alpha beta TIM barrel domains. 1pii is the experimentally determined structure of the bifunctional *E. coli* enzyme phosphoribosyl-anthranilate isomerase: indoleglycerolphosphate synthase. The 1pdz lyase and the 2exo hydrolase structures both adopt an alpha beta TIM barrel fold. 1hpm is a heat shock protein ATPase fragment adopting an open alpha beta alpha sandwich structure. 1eedP0 and 1mpp00 are aspartate proteinases, adopting beta barrel structures. This beta barrel structure is not compatible with the SAND secondary structure prediction results, that comprise equal numbers of alpha helices and beta strands. The predicted secondary structure of SAND is only a fair match with the largely alternating pattern of helices and strands of 1pii. However, the secondary structure and solvent accessibility predictions for SAND indicate mostly amphipathic helices and predominately solvent

inaccessible strands, these characteristics are typical properties observed in known alpha beta protein fold types like the alpha beta sandwich structures or the closed alpha beta TIM barrels.

## Conclusions

Lack of conformity and standards in annotation has resulted in variability between genomes in how accurately genes are predicted. In this example, SAND has been partially missed from the *A. thaliana* genome annotation, mis-predicted in *C. elegans* and *S. cerevisiae*, whilst Hs\_SAND1 has been not annotated in the human genome. Whilst there are at least two mammalian SAND genes, and at least one SAND gene is present in all eukaryotes, the function of these genes' products is still completely unknown. It is possible that at least one of the mammalian paralogues may play a role in neoplastic disease following loss of exon 3. Using bioinformatic analyses, this study has illustrated the complexities of determining what the structure and function of these proteins might be. When faced with a totally new gene coding for a protein product of unknown function, this task is non trivial and in this example after considerable database searching and analysis we have made only small advances in this endeavour. SAND is only one gene product in approximately 60% of the 19, 000 in *C. elegans* that have an unknown function. A complete eukaryotic genome sequence is today's achievement, but a functional understanding of it is still a distant goal.

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