

Research Article

Estimation of synteny conservation and genome compaction between pufferfish (*Fugu*) and human

Aoife McLysaght, Anton J. Enright[†], Lucy Skrabanek and Kenneth H. Wolfe*

Department of Genetics, University of Dublin, Trinity College, Dublin 2, Ireland

*Correspondence to:

K. H. Wolfe, Department of Genetics, University of Dublin, Trinity College, Dublin 2, Ireland.
E-mail: khwolfe@tcd.ie

[†]Current address:

Computational Genomics Group Research Programme, The European Bioinformatics Institute, EMBL Cambridge Outstation, Cambridge CB10 1SD, UK.

Abstract

Background: Knowledge of the amount of gene order and synteny conservation between two species gives insights to the extent and mechanisms of divergence. The vertebrate *Fugu rubripes* (pufferfish) has a small genome with little repetitive sequence which makes it attractive as a model genome. Genome compaction and synteny conservation between human and *Fugu* were studied using data from public databases.

Methods: Intron length and map positions of human and *Fugu* orthologues were compared to analyse relative genome compaction and synteny conservation respectively. The divergence of these two genomes by genome rearrangement was simulated and the results were compared to the real data.

Results: Analysis of 199 introns in 22 orthologous genes showed an eight-fold average size reduction in *Fugu*, consistent with the ratio of total genome sizes. There was no consistent pattern relating the size reduction in individual introns or genes to gene base composition in either species. For genes that are neighbours in *Fugu* (genes from the same cosmid or GenBank entry), 40–50% have conserved synteny with a human chromosome. This figure may be underestimated by as much as two-fold, due to problems caused by incomplete human genome sequence data and the existence of dispersed gene families. Some genes that are neighbours in *Fugu* have human orthologues that are several megabases and tens of genes apart. This is probably caused by small inversions or other intrachromosomal rearrangements.

Conclusions: Comparison of observed data to computer simulations suggests that 4000–16 000 chromosomal rearrangements have occurred since *Fugu* and human shared a common ancestor, implying a faster rate of rearrangement than seen in human/mouse comparisons. Copyright © 2000 John Wiley & Sons, Ltd.

Keywords: *Fugu rubripes* (pufferfish); comparative genomics; genome compaction; genome rearrangement; synteny conservation

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Introduction

Comparative genomics has great potential for maximizing the value of genome sequencing projects. Sydney Brenner and colleagues (Brenner *et al.*, 1993; Elgar *et al.*, 1996) proposed the pufferfish *Fugu rubripes* as a model genome for use in dissecting the human genome. As a vertebrate, *Fugu* is expected to have a similar gene repertoire to human. However, its genome, at ~400 Mb, is

approximately 7.5 times smaller than that of human. The reduced amount of repetitive sequence and high gene density make this small genome attractive to molecular biologists.

There are two main factors that will determine whether *Fugu* will be genuinely useful as a model vertebrate. *Fugu* genes must show sufficient similarity to their human orthologues to enable the isolation of a *Fugu* gene with a human (or other mammalian) DNA probe, and vice versa. Further-

more, knowledge of the extent of linkage conservation between the two genomes will advise as to the feasibility of positional cloning using map information extrapolated from one species to the other (Elgar *et al.*, 1996). Several regions of conserved synteny (but not necessarily conserved gene order) have already been reported between these two genomes (e.g. Baxendale *et al.*, 1995; Trower *et al.*, 1996; Elgar *et al.*, 1999; and references in Table 2).

Exploring the relationship between the human and pufferfish genomes in terms of the extent of synteny conservation and patterns of genome compaction could give insights into the evolution of vertebrate genomes, and could also provide more information on the usefulness of *Fugu* as a model genome. However, at present it is not known how large the syntenic regions are, or how well the gene order is conserved between *Fugu* and human. Recent research on the zebrafish (*Danio rerio*) indicated that for some groups of genes, synteny is conserved in the human but the order of the genes along the syntenic chromosome is different in the two species (Postlethwait *et al.*, 1998). Moreover, many mammalian genes have two zebrafish orthologues, and this is probably due to whole genome or chromosomal duplications that occurred in bony fish (including zebrafish and *Fugu*) after their divergence from the tetrapod lineage (Amores *et al.*, 1998; Gates *et al.*, 1999). It is also not known whether the compaction of the *Fugu* genome relative to the human is uniform throughout the genome, particularly in view of the uneven distribution of genes in the human genome (Ikemura and Wada, 1991; Duret *et al.*, 1995; Deloukas *et al.*, 1998).

Here we have made a comparative genomics study of *Fugu* and the human to investigate the phenomenon of genome compaction and to estimate the level of synteny conservation. There is no genetic map for *Fugu* (it is not possible to breed this fish in the laboratory), so gene linkage is only discernible at the level of genes that were sequenced on the same cosmid or other clone contig. We used two sources of *Fugu* sequence data: large contiguous genomic sequences determined by a variety of laboratories and obtained from GenBank; and 'cosmid skimming' data from the *Fugu* Landmark Mapping Project at the UK MRC HGMP-RC (Elgar, 1996; Elgar *et al.*, 1999). The human map data was obtained from two sources: the Online Mendelian Inheritance in Man database (OMIM

1999); and the physical map of about 30 000 genes (GeneMap '98) constructed from radiation hybrid data by Deloukas *et al.* (1998).

Materials and methods

Analysis of homologous introns from *Fugu* and human

The 22 genes included in this analysis were: RPS3, RPS24, DLST, STK9, PAX6, RPS7, APP (low GC3 group); SURF3, SMC1, RPL41, ARF3, CFOS, XLRS1, PCOLCE (medium GC3 group); CSFR1, GH, TSC2, HMOX1, WNT1/INT1, PKD1, G6PD, IT (high GC3 group). All sequences were obtained from GenBank.

Fugu sequence data

SwissProt version 37 (27 July 1999) contains 5406 human proteins. These were compared to the database of *Fugu* 'skimmed' cosmids using TBLASTN (Altschul *et al.*, 1990) using the BLOSUM62 scoring matrix and the SEG filter (Wootton and Federhen, 1996). To remove obvious paralogous hits, only the top hit for each query was retained (provided that it had $P \leq 10^{-15}$) as well as weaker hits that were within a factor of 10^5 of the top hit. The results of this BLAST search including human map information are available at <http://biotech.bio.tcd.ie/~amclysag/skimmed.html>

A 'skimmed' cosmid was adjudged to contain two genes if two non-overlapping subclones hit different mapped human proteins that are <40% identical in sequence and had $P \leq 10^{-15}$ in a BLASTP search. Overlapping *Fugu* cosmids were identified manually and reduced to one entry in Table 1.

Fugu proteins from completely sequenced cosmids were compared to the database of human sequences from GeneMap '98 by the TBLASTN programme applying the SEG filter. Only hits with a significance of $\leq 10^{-15}$, and that were no more than 10^5 less likely than the top hit, were accepted. Only the best hit per chromosome was included in further analysis.

Some of the limitations on the analysis of the skimmed cosmids become apparent when the results are compared with the fully sequenced cosmids. Cosmid 168J21 has been fully sequenced under Accession No. AJ010348 (Cottage *et al.*, 1999). The full sequence has three annotated proteins, all of

which had human homologues on chromosome 3. In the analysis of the skimmed cosmid sequence only one gene was found. As all three human orthologues are in the SwissProt database, it must be the case that the cosmid subclones do not include the coding sequences of the other two genes.

Human GeneMap '98 sequences

Deloukas *et al.* (1998) compiled a map (GeneMap '98) of human gene-based markers by radiation hybrid mapping. This includes approximately 30 000 genes. By electronic PCR (Schuler, 1997) they found the corresponding genomic sequence, mRNA and/or EST from the public databases. These results are updated weekly and were downloaded from the NCBI FTP site on 21 December 1998.

A BLAST database of human sequences represented on this radiation hybrid map was created. In order to have comparable map units, only the data from the GeneBridge4 panel (Gyapay *et al.*, 1996) were included. Some parts of the genome are represented more than once in the ePCR output because they have been sequenced more than once as genomic sequence, mRNA and/or EST. Redundancies of this kind were removed, preferentially keeping genomic sequences over mRNA over unfinished sequences over ESTs. The final database had 28 133 entries, totalling 226 506 753 nucleotides.

Some markers in GeneMap '98 are listed with several allocated map positions. In these cases the same position found from several independent experiments or the position with the highest confidence value, as determined by Deloukas *et al.* (1998), was used. Distances within the genome were estimated by counting the number of intervening genes in GeneMap '98. We then adjusted these values for missing data by multiplying this number by 80 000/30 000 (assuming the human genome contains 80 000 genes and the map contains 30 000 genes).

Computer simulation of genomic rearrangement

In order to make this simulation as realistic as possible, paralogues were assigned at the frequencies observed in the real data. Of the 91 *Fugu* proteins analysed, 78 had hits in the database of mapped human sequences. The distribution of hits is as follows: 47 hit one human sequence, 14 hit two, eight hit three, two hit four, and families of

seven, 11, 12, 15, 39, 42, and 59 human proteins were observed once each. More extensive human protein family size data from an intragenome comparison (Imanishi *et al.*, 1997) was used to confirm these results in an independent simulation.

Results

Compaction of *Fugu* introns

The *Fugu* genome is much smaller than the human genome, but by virtue of being vertebrate is presumed to have a similar gene repertoire (Brenner *et al.*, 1993). The difference in size must therefore be primarily due to differences in non-coding DNA, including both intergenic and intronic DNA. In vertebrate genomes there is a correlation between gene length and G+C content, with long genes being rare in G+C-rich isochores (Duret *et al.*, 1995). This suggested that there might be a correlation between base composition and the size difference between a human gene and its *Fugu* homologue.

Orthologous *Fugu* and human introns were identified by finding orthologous genomic sequences in GenBank, aligning the protein sequences using the Gap programme (with default settings) of the GCG package, and mapping intron locations onto the protein alignment. Introns were designated orthologous if they were in the same phase and occurred at precisely the same position in the protein alignment produced by Gap. No allowance was made for possible intron sliding during evolution. Using this method, 199 pairs of orthologous introns from 22 genes were found. There were only six cases where we could say with confidence that an intron had been gained or lost after the divergence of these two species. These were all cases where there was an unambiguous alignment of the two protein sequences, and where an intron was present in one sequence but there was no equivalent intron nearby or out of phase in the other organism. Non-coincident introns and introns in ambiguous alignments were excluded from further analysis. Recent research by Hurst *et al.* (1999) tentatively suggests that there may be a dichotomy in the relationship of synonymous GC content and intron size, with warm-blooded vertebrates showing a negative correlation, as previously observed, and cold-blooded vertebrates (including *Fugu*) showing a positive correlation. However, this is not borne

out here. In our dataset there is no correlation between intron size and GC3 content of the genes that house them.

Genes were assigned into three equal-sized groups according to their G+C content at codon third positions (GC3) in human, and the lengths of equivalent introns were compared (Figure 1A). The sum of the lengths of all 199 introns in *Fugu* was 59 392 bp, just over eight times smaller than the sum of the lengths of all the human introns (488 726 bp). The large introns of GC3-poor genes are seen to be severely compacted. The compaction averages are 2.9, 6.0 and 14.6, respectively, for the high-, medium-, and low-GC3 groups of genes (Figure 1A), which is broadly consistent with expectations. One-fifth of the *Fugu* introns (41 of the 199) are actually larger than their human counterparts (many only marginally so), and most of these are high-GC3 genes in the human (Figure 1B). However, for the majority of introns (Figure 1B) there does not appear to be any consistent relationship between intron lengths in the two species, or between these and GC3 in their host genes.

The compaction of individual genes, instead of individual introns, was also calculated (Figure 1C, D). Compaction was calculated by dividing the sum of the lengths of introns in a human gene by the sum of the lengths of their *Fugu* orthologues (excluding any non-coincident introns). The compaction values range from 46 (in the APP gene; Villard *et al.*, 1998) down to values of less than 1 in two genes (growth hormone and *int1/wnt1*), where the *Fugu* gene is larger than the human one. If the GC3 content of a gene and the compaction of its introns are related, then one would expect the greatest compaction to be between human genes with low GC3 and *Fugu* genes with high GC3. Rather surprisingly, there does not appear to be any relationship between the degree of compaction and the base composition in either species (Figure 1C), or the amount of interspecies difference in base composition (Figure 1D). The two most severely compacted genes have similar GC3 content in *Fugu* and human (Figure 1D).

Synteny conservation between *Fugu* and human

Synteny conservation between two species can be measured in two directions. We can ask, 'What

proportion of genes that are syntenic in species A are also syntenic in species B?', or conversely, 'What proportion of genes that are syntenic in B are also syntenic in A?'. These are two distinct quantities, as becomes obvious if one considers a hypothetical case where one of the species has only a single chromosome. The only syntenic genes that are known in *Fugu* are those that have been sequenced on the same clone; there are no large-scale maps of chromosomes. Therefore, we measured *Fugu*/human synteny conservation in terms of the proportion of neighbouring genes (from the same clone or GenBank entry) in *Fugu* that are syntenic in human. We also applied various limits to the physical distance permitted between the syntenic genes in human. Two separate datasets were analysed, as described below.

Synteny conservation—'cosmid skimming' data

The HGMP-RC *Fugu* landmark mapping project (Elgar, 1996; Elgar *et al.*, 1996; Elgar *et al.*, 1999) surveyed the *Fugu* genome by limited sequencing ('skimming') of a large number of genomic cosmid clones. Sets of shotgun sequence reads for 850 randomly chosen cosmids are publically available from their website (<http://fugu.hgmp.mrc.ac.uk/>). The data consist of 40 303 sequence reads, with an average of 47 reads per cosmid and 486 bp per read. Each read is assumed to contain no more than one gene.

Because these sequences are short and largely unannotated, we compared them to human data from SwissProt, rather than GeneMap '98 (which contains a large number of EST sequences). Cytogenetic map positions for 3963 of the 5406 human proteins in SwissProt were obtained by following links to OMIM. All 5406 proteins were searched against the *Fugu* cosmid database, using TBLASTN (Altschul *et al.*, 1990). Putative orthologous relationships were identified as described in Materials and methods.

A *Fugu* cosmid was considered 'informative' (i.e. it appeared to contain more than one gene, and so contained linkage information) if two different sequence reads hit two different mapped human sequences which did not themselves show significant sequence identity to one another. We identified 48 informative cosmids, containing 58 links between nearby *Fugu* genes (Table 1). For 26 of these links

Table 1. *Fugu* 'skimmed' cosmids containing homologues of at least two mapped human SwissProt sequences

Cosmid	Syntenic links ^a		Subclone	SwissProt name ^b	Description	OMIM location
	+	−				
002I16	0	1	bB8	CGB1	G ₂ /mitotic-specific cyclin B1	5q12
			bC1	UBCG	Ubiquitin-conjugating enzyme E2 G1	1q42
003A22	0	1	aD2	LCFD	Long-chain fatty-acid CoA ligase 4	Xq22.3
			aE9	API9	Clathrin coat assembly protein	Chr.7
018N05	0	1	cB3	COMT	Catechol O-methyltransferase	22q11.2
			cB7	RYK	Tyr-protein kinase RYK	3q22
020M06	1	1	bF2	FI6P	Fructose-1-6-bisphosphate	9q22.2-q22.3
			bG9	GAS1	Growth arrest specific protein	9q21.3-q22.1
			aE1	LMG2	Laminin γ -2 chain	1q25-q31
030J22	2	0	aF4	TRFE	Serotransferrin	3q21
			aF7	IF4G	Translation initiation factor 4 G	3q27
			aG1	CLC2	Chloride channel protein 2	3q26-qter
032I12	0	1	aD1	PA2Y	Cytosolic phospholipase A2	1q25
			aE3	TSP1	Thrombospondin 1	15q15
035P08	1	0	aC2	KPT1	Ser/thr protein kinase PCTAIRE-1	Xp11.3-p11.23
			aD5	HFC1	Host cell factor C1	Xq28
042H13	1	2	aE6	PIGF	Phosphatidylinositol-glycan synthase F	2p21-p16
			bA4	GCH1	GTP cyclohydrolase I	14q22.1-q22.2
			bD10	MSH2	DNA mismatch repair protein	2p22-p21
			bF8	CIKA	Voltage gated K channel KV21	20q13.2
050M16	1	0	bC5	CYCH	Cyclin H	5q13.3-q14
			bG2	GTPA	GTPase-activating protein (GAP)	5q13.3
055I13	0	1	bD9	A2MG	α -2-Macroglobulin	12p13.3-p12.3
			bE2	ECH1	δ 3,5- δ 2,4-Dienoyl-CoA isomerase	19q13
057B20	0	1	aC11	SCI4	sec-14-Like	17q25.1-q25.2
			aH1	GNT5	Glucoseaminyltransferase V	2q21
059A13	0	1	aD6	VLCS	Very long-chain acyl-CoA synthetase	15q21.2
			aE6	AMBP	AMBP protein	9q32-q33
060I09	0	1	ITA1	ITA1	Integrin α -1	Chr.5
			aG3	ROK	Het. nuclear ribonucleoprotein K	9q21.32-q21.33
063J19	1	1	aA5	AGAL	α -Galactosidase A	Xq22
			aD12	RL44	60S rpl44	Chr.14
			aH4	DDP	Dystonia protein	Xq22
068B10	1	0	aA9	MET	Hepatocyte growth factor receptor	7q31
			aC8	MGR8	Metabotropic glutamate receptor 8	7q31.3-q32.1
077E20	1	1	bB7	COGT	Matrix metalloproteinase-14	14q11-q12
			cC4	PKD2	Polycystin 2	4q21-q23
			cC5	AF4	AF-4 protein	4q21
081G09	1	0	aD12	CIK4	Voltage gated K channel protein	11q13.4-q14.1
			aF6	EAT2	Excitatory amino acid transporter 2	11p13-p12
082H05	0	1	aG5	KMLS	Myosin light-chain kinase	3cen-q21
			aH4	NED4	NEDD-4 protein	15q
082L03	1	0	aD12	MPCP	Mitochondrial PO ₄ carrier	12q23
			aF10	THPA	Thymopoietin- α	12q22
086H03	1	0	bC4	DOC2	Differentially expressed protein 2	5p13
			cE8	CO9	Complement component C9	5p13
096F11	0	1	aA7	WN11	WNT-11	11q13.5
			bC7	ACHD	Acetylcholine receptor δ chain	2q33-q34
103N12	1	0	aB9	RO52	Ro protein, 52 kDa	11p15.5
			bA6	COGM	Macrophage metalloelastase	11q22.2-q22.3
104N10	1	0	aD3	FER	FER Tyr protein kinase	5q21-q22
			bA12	MAN2	α -Mannosidase II	5q21-q22/20q11.2
107H09	0	1	aF11	RS12	40S rps12	6q
			aG6	EYA1	Eyes absent homologue 1	8q13.3
107N05	0	2	aG10	BCAM	Branched-chain aminotransferase	19q13
			aH4	GRN	Granulins	Chr.17

Table 1. Continued

Cosmid	Syntenic links ^a		Subclone	SwissProt name ^b	Description	OMIM location
	+	—				
110112	0	1	aF6	EAT2	Excitatory amino acid receptor 2	11p13-p12
			dA4	PAK1	Ser/thr protein kinase PAK- α	11q13-q14
			dD3	PET1	Oligopeptide transporter	13q33-q34
114M17	1	0	bB8	IHBA	Inhibin β a chain	7p15-p13
			bC3	EGFR	Epidermal growth factor receptor	7p12.3-p12.1
116E05	0	1	aB3	GNT2	Acetylglucosaminyltransferase	14q21
			aE6	HS9A	Heat shock protein 90- α	1q21.2-q22
118A15	0	1	cC8	PERT	Thyroid peroxidase	2p25
			cG3	VMD2	Bestrophin	11q13
122O20	0	1	cA4	CASR	Extracellular Ca-sensing receptor	3q13.3-q21
			cD1	CTR2	Low affinity cationic amino acid transporter	8p22
123I02	0	1	aC11	BTG1	B-cell translocation 1	12q22
			aE5	TEF	Thyrotroph embryonic factor	22q13
128G19	1	0	aC11	BTG1	B-cell translocation 1r	12q22
			aE5	TEF	Thyrotroph embryonic facto	22q13
137O18	0	1	aE10	CYA1	Adenylate cyclase, type I	7p13-p12
			bA4	BNA1	Amiloride-sensing brain Na ⁺ channel	17q11.2-q12
141H19	1	0	aH10	LDHH	L-Lactate dehydrogenase H chain	12p12.2-p12.1
			aH9	UGS2	Glycogen synthetase	12p12.2
143P11	1	0	aB6	ANK1	Ankyrin R	8p11.2
			aD6	NFM	Neurofilament triplet M protein	8p21
145K17	0	1	bF3	RHMI	Rhombotin-1	11p15
			cB1	AHR	AH receptor	7p15
147P16	2	0	aD1	DDP	Deafness dystonia protein	Xq22
			aF9	BTK	Tyr-protein kinase BTK	Xq21.3-q22
			aG7	GRA2	Gly receptor α -2 chain	Xp22.1-p21.2
155N11	1	1	bE7	SYB2	Synaptobrevin 2	17pter-p12
			bH3	MPP2	Maguk P55 subfamily member 2	17q12-q21
			aD7	UTY	Ubiquitously transcribed TPR on Y	Yq11
156P04	1	0	aH2	RO52	Ro protein, 52 kDa	11p15.5
			hC8	Z195	Zinc finger protein 195	11p15.5
157C15	0	1	aA3	RIR2	Ribonucleoside reductase M2	2p25-p24
			aD10	RL30	60S rPL30	Chr.8
159J19	1	0	aB1	MPK4	MAP kinase kinase 4	17p11.2
			aD11	MYSP	Myosin H perinatal skeletal muscle	17p13.1
164B03	0	1	aD11	ISL1	Insulin gene enhancer protein ISL-1	5q
			aD4	ETFA	Electron transfer flavoprotein- α	15q23-q25
165O08	0	1	aH7	UBA1	Ubiquitin-activating enzyme E1	Xp11.23
			bD10	DPOE	DNA polymerase ϵ , subunit A	12q24.3
171K15	0	1	bB10	DMK	Myotonin protein kinase	19q13.2-q13.3
			bB6	BMAL	Brain and muscle ARNT-like 1	11p15
174C18	1	1	aD11	G6PD	G6PD	Xq28
			bA1	CCB3	Ca ²⁺ channel β -3	12q13
			bB11	CYA6	Adenylate cyclase type VI	12q12-q13
176J15	1	0	aA8	DESM	Desmin	2q35
			aC5	PTPN	Protein-tyr phosphatase N	2q35-q36.1
192G14	1	0	aA2	ADG	γ -Adaptin	16q23
			aA7	RFP	Zinc finger protein RFP	Chr.6
222J11	1	0	bE3	NTTA	Taurine transporter	3p25-q24
			bC4	ACTQ	Ca ²⁺ -transporting ATPase	3p26-p25
Totals:	26	32				

^aThe '+' column refers to conserved linkages between *Fugu* and human, and the '-' column refers to non-conserved linkages.

^bAll SwissProt IDs are truncated, omitting '_HUMAN' from each one.

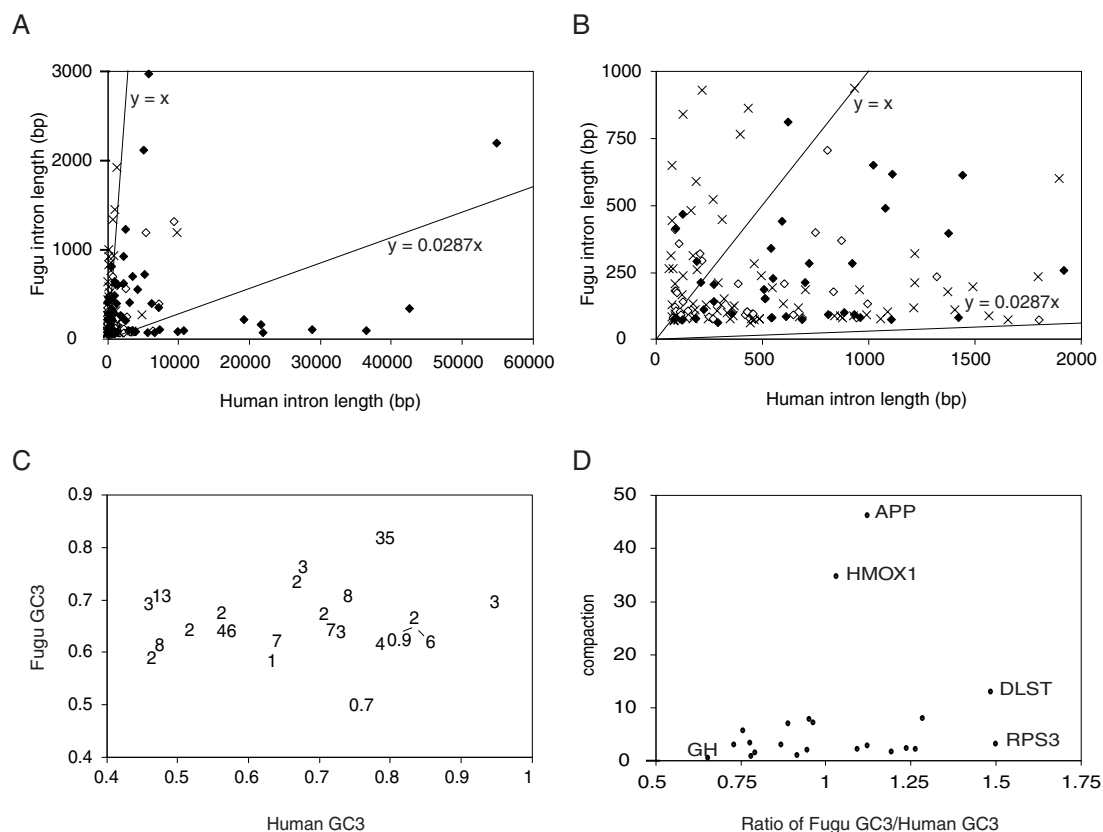


Figure 1. (A) Lengths of 199 orthologous introns from *Fugu* and human. The regression line for all data is shown, as is a line of slope 1. The symbols for the points represent different GC3 content categories in the human gene where the black diamond denotes low GC3 (<63.5%), the white diamond denotes medium GC3 (63.5–76%), and the cross denotes high GC3 content (>76%). The categories were designed in such a way as to have equal numbers of genes in each group. The 22 genes from which the introns are derived are named in Materials and methods. (B) Inset of (A) showing only the smaller introns. (C) GC3 content of the 22 orthologous genes whose introns were analysed. The points are replaced by values indicating relative gene compaction. Compaction was calculated by dividing the sum of the lengths of introns of a human gene by the sum of the lengths of their *Fugu* orthologues, ignoring non-conserved introns. (D) Compaction of 22 genes vs. the ratio of GC3 in *Fugu* to that in human. Outlying genes are labelled: APP, amyloid precursor protein; GH, growth hormone; RPS3, ribosomal protein S3; HMOX1, heme oxygenase; DLST, dihydrolipoamide succinyltransferase

(45%), the human homologues are on the same chromosome (i.e. synteny was conserved).

The same *Fugu* Landmark Mapping Project data were recently analysed by Elgar *et al.* (1999). They reported that ‘three-quarters’ of informative cosmid showed synteny to human. However, it is difficult to account for the differences between our results and theirs as they do not specify what stringency they imposed on the definition of orthology, neither do they indicate which cosmids displayed an orthologous relationship with which human sequences. Perhaps the greatest discrepancy between these analyses is in the number of

informative cosmids found (349 by Elgar *et al.* compared to 48 in this study). We expect that this difference is due to a greater stringency employed by us in the designation of orthologues (as described in Materials and methods).

Synten conservation—complete *Fugu* genomic sequences

We examined the GenBank annotation of all *Fugu* sequences greater than 5 kb long to look for sequences that coded for two or more proteins. The 21 GenBank entries that fit this criterion

Table 2. Details of the completely sequenced *Fugu* cosmids used in this analysis

Accession No.	Base pairs	Genes included	Reference
af056116	148 640	ACVR1B, ALR, fhf, Ikars-like, wnt1, wnt10b, ARF3, erbB3, PAS1, rpl41, LRP1	Gellner and Brenner, 1999
af094327	69 056	SCML2, STK9, XLR1, PPEF-1, KELCH2, KELCH1, PHKA2, API9, U2AF1-RS2	Brunner <i>et al.</i> , 1999
u90880	61 901	RNA-H, CAB3B, Adenyl Cyclase-VI, G6PD, LG3P, Na ⁺ channel 2	Riboldi Tunnicliffe GR <i>et al.</i> , unpublished
af016494	66 729	GABRB, P55, VAMP-1, PCOLCE, GRMP	Riboldi Tunnicliffe GR <i>et al.</i> , unpublished
af026198	63 155	LI-CAM, SMC1, CCA1	Riboldi Tunnicliffe GR <i>et al.</i> , unpublished
af083221	43 373	Neurotransmitter receptors, YDR140w homologue, glycineamide ribonucleotide transformylase	Reboul <i>et al.</i> , 1999
aj010317	39 410	GRM-7, TRIP, Sand, PRGFR3	Cottage <i>et al.</i> , 1999
y15170	10 753	EST00098 homologue, SURF2, SURF4, ASS	Armes <i>et al.</i> , 1997
aj010348	39 850	UBE1-like, PRGFR2, calmodulin binding protein kinase	Cottage <i>et al.</i> , 1999
af021880	37 170	IGFII, TH, NAP2	Chen E. <i>et al.</i> , unpublished
af021531	45 565	WT, Reticulocalbin, PAX6	Miles <i>et al.</i> , 1998
z93780	34 807	CPS3, MLC, MAP2	Schofield <i>et al.</i> , 1997
u92572	20 919	HOXC-9, HOXC-8, HOXC-6	Aparicio <i>et al.</i> , 1997
y15171	8 902	rpl7a (SURF3), SURF1, SURF6	Armes <i>et al.</i> , 1997
af013614	55 892	TSC2, PKD1	Sandford <i>et al.</i> , 1997
af022814	37 400	Zinc finger transcription factor, HMOX1	Gottgens <i>et al.</i> , 1998
af030881	5 645	gag, pol	Poulter and Butler, 1998
aj010316	10 959	Cav-2, Cav-1	Cottage <i>et al.</i> , 1999
u63926	23 196	PDGFR- β , CSF1R	How <i>et al.</i> , 1996
u92573	13 583	HOXA-10, HOXA-9	Aparicio <i>et al.</i> , 1997

Cosmids are listed in order of decreasing number of annotated proteins. The list of annotated proteins for each cosmid does not include putative proteins with no known human homologues at the time of submission to the sequence database.

(Table 2) total just under 0.9 Mb and encode 91 annotated proteins (some putative). Genes from the same GenBank entry have a known linkage relationship in the *Fugu* genome because they were sequenced contiguously.

The proteins encoded by these *Fugu* sequences were compared using TBLASTN to the database of human nucleotide sequences whose map positions are known in GeneMap '98 (Deloukas *et al.*, 1998). For some of the *Fugu* sequences, our results confirm previously published analyses (Sandford *et al.*, 1996; Aparicio *et al.*, 1997; Armes *et al.*, 1997; Schofield *et al.*, 1997; Miles *et al.*, 1998; Brunner *et al.*, 1999; Gellner and Brenner, 1999; Reboul *et al.*, 1999).

The results were examined to look for candidate conserved syntenous regions between human and *Fugu*. This was facilitated by a new method for displaying the relative positions of the homologues in the two species. In many cases, such as in the example shown in Figure 2, there was more than one candidate human chromosomal region for conserved synteny. In Figure 2 the *Fugu* sequence

(AF056116) appears to have conserved synteny with human chromosome 12 by virtue of having several top scoring BLAST hits to human genes that map close together on that chromosome, largely as described by Gellner and Brenner (1999). What is interesting is that regions on chromosomes 7, 17 and 2 also show synteny with this *Fugu* sequence (including matches to *Fugu* proteins not having homologues on chromosome 12—genes 3, 4, 6, and 14; Figure 2). These are the human chromosomes that contain the HOX clusters and this indicates that the similarity of these human chromosomes to each other extends beyond those clusters, as has been suggested by others (Ruddle *et al.*, 1994).

To examine synteny conservation in a quantitative way, instead of simply the presence or absence of genes on the same chromosome, we calculated the proportion of *Fugu* close neighbours (genes from the same GenBank entry) whose homologues were within a specified distance, x , of each other in human. We use the term 'proximity conservation' to denote this property of genes remaining within a specified distance of each other (regardless of gene

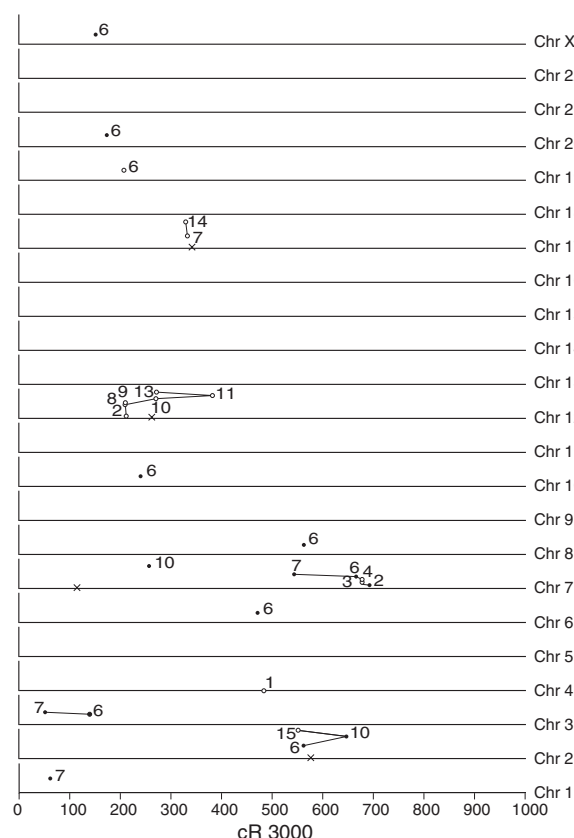


Figure 2. Graphical representation of the results of the TBLASTN search of the proteins from *Fugu* sequence AF056116 (Gellner and Brenner, 1999) against a database of mapped human sequences (GeneMap '98). The relative positions of the best hits of each of the 15 annotated *Fugu* proteins from this cosmid are shown for each chromosome in turn. The horizontal axis represents position (measured in centiRads) on the human chromosome in question, and each vertical axis represents the relative order (1–15) of the *Fugu* genes on the *Fugu* cosmid. White dots designate the top-scoring TBLASTN hit for each *Fugu* protein; black dots indicate weaker hits (that are within 10^5 of the strongest hit). The genes are in the following order in *Fugu*: 1, ACVR1B; 2, ALR; 3, fhf; 4, R05D3.2-like protein; 5, I38E3.2-like protein; 6 Ikaros-like; 7, wnt1; 8, wnt10b; 9, ARF3; 10, erbB3; 11, PAS1; 12, rpl41; 13, I78O23.1-like protein; 14, diaphanous-like protein; 15, LRP1. In addition to the matches shown here (based on data in GeneMap '98), genes 1, 7, 12 and 15 also have homologues on chromosome 12q13 (Kenmochi et al., 1998; Gellner and Brenner, 1999). The positions of Hox clusters ABCD are represented by crosses on chromosomes 7, 17, 12 and 2, respectively

order). To allow for the uneven distribution of genes in the human genome, the distance x was expressed in terms of the estimated number of

intervening genes, instead of in the physical map units (cR) that were used in GeneMap '98 (Deloukas et al., 1998). The number of intervening genes was estimated from GeneMap '98 by counting the number of intervening genes appearing on the map between the genes of interest and scaling by a factor of 80 000/30 000 to allow for unsequenced genes. This allows for gene density variation within and between chromosomes. Where more than one human sequence had been assigned to the same map position by Deloukas et al. (1998), these sequences were arbitrarily assigned an order.

The results are summarised in Table 3. Only 18% of *Fugu* neighbours have sequenced human homologues that are within 10 genes of one another. This increases to 39% within a limit of 200 intervening genes, and to a maximum of 47% within a limit of 4000 intervening genes (this is effectively no limit, because it is approximately the size of a chromosome). The last value is similar to the synteny estimate from Table 1 (which has no limit on the intervening distance).

Computer simulation of genomic rearrangement

We used computer simulations to try to relate the observed level of proximity conservation to the number of genomic rearrangements that have occurred since the divergence of *Fugu* and human. The simulation started with a linear array of 80 000 genes, representing the current gene order in *Fugu*. Varying numbers of rearrangements were made in a copy of this genome (representing human) by randomly choosing two endpoints in the genome and inverting the segment in between. To reflect the missing data in the human map, randomly chosen genes were marked 'unmapped' until only 30 000 remained (the number of genes in Deloukas et al., 1998). Pairs of genes that are neighbours in *Fugu* were then examined to see if they are neighbours in human, similar to the method of analysis in Tables 1 and 3.

To make the simulation more realistic, we modelled the presence of gene families. Because more than half of all human genes are still not included in the human gene map, there is a real possibility that if the human orthologue of a *Fugu* gene is not mapped, the *Fugu* gene would mistakenly be paired with a mapped human paralogue instead. This could reduce the estimated level of

Table 3. Observed levels of synteny conservation between completely sequenced *Fugu* cosmids and human

Fugu Accession No.	Annotated proteins	Proteins with human BLAST hits $P < 10^{-15}$	Maximum possible links	Human chromosome ^a	Number of links also present on human chromosome, at different values of x intervening genes ^b						
					5	10	20	50	200	1000	4000
AF056116	15	13	12	2	0	1	1	1	1	1	2
				3	0	0	0	0	0	1	1
				7	0	0	0	1	2	4	5
				12*	1	1	2	2	3	4	5
				17	0	0	0	0	1	1	1
AF094327	9	9	8	5	0	0	0	0	1	1	1
				X*	1	2	4	4	4	4	4
				2	0	0	0	0	0	0	1
				20*	0	0	0	0	1	2	2
				12	0	0	0	0	2	2	0
U90880	9	6	5	12	0	0	0	0	2	2	0
				3	0	0	0	0	0	0	0
				2	0	0	0	0	0	0	0
				3	0	0	0	0	0	0	0
				3	0	0	0	0	2	2	2
U72484	6	4	3	3	0	1	1	1	1	1	1
				9	0	0	0	0	0	0	1
				9	0	0	0	0	0	0	1
				11*	0	0	0	0	1	1	1
				12	0	0	0	0	0	0	1
AF016494	3	3	2	11	0	0	0	1	1	1	1
				2	0	0	0	1	1	1	1
				2	0	0	1	1	1	1	1
				2	0	2	2	2	2	2	2
				9	0	0	0	0	0	0	0
AF026198	3	2	1	16	0	0	0	0	0	0	0
				0	0	0	0	0	0	0	0
				0	0	0	0	0	0	0	0
				7	1	1	1	1	1	1	1
				4	1	1	1	1	1	1	1
AF083221	4	3	2	7	0	1	1	1	1	1	1
				4	1	1	1	1	1	1	1
				2	1	1	1	1	1	1	1
				2	2	2	2	2	2	2	2
				2	2	2	2	2	2	2	2
AJ010317	4	3	2	4	0	1	1	1	1	1	1
				2	1	1	1	1	1	1	1
				3	1	1	1	1	1	1	1
				3	2	2	2	2	2	2	2
				3	2	2	2	2	2	2	2
Y15170	4	2	1	12	0	0	0	0	0	0	0
				3	0	0	0	0	0	0	0
				3	0	0	0	0	0	0	0
				3	0	0	0	0	0	0	0
				3	0	0	0	0	0	0	0
AJ010348	3	3	2	12	0	0	0	0	0	0	0
				11	0	0	0	0	0	0	0
				2	0	0	0	0	0	0	0
				2	0	0	0	0	0	0	0
				2	0	0	0	0	0	0	0
AL021880	3	3	2	12	0	0	0	0	0	0	0
				11	0	0	0	0	0	0	0
				2	0	0	0	0	0	0	0
				2	0	0	0	0	0	0	0
				2	0	0	0	0	0	0	0
AL021531	3	3	2	11	0	0	0	1	1	1	1
				2	0	0	0	1	1	1	1
				2	0	0	1	1	1	1	1
				2	0	0	1	1	1	1	1
				2	0	0	1	1	1	1	1
Z93780	3	3	2	2	0	0	0	1	1	1	1
				2	0	0	1	1	1	1	1
				2	0	0	1	1	1	1	1
				2	0	0	1	1	1	1	1
				2	0	0	1	1	1	1	1
U92572	3	3	2	9	1	1	2	2	2	2	2
				16	0	0	0	0	0	0	0
				0	0	0	0	0	0	0	0
				7	1	1	1	1	1	1	1
				4	1	1	1	1	1	1	1
Y15171	3	2	1	7	0	1	1	1	1	1	1
				4	1	1	1	1	1	1	1
				2	1	1	1	1	1	1	1
				2	2	2	2	2	2	2	2
				2	2	2	2	2	2	2	2
AF013614	2	1	0	4	0	1	1	1	1	1	1
				7	1	1	1	1	1	1	1
				1	1	1	1	1	1	1	1
				2	1	1	1	1	1	1	1
				2	2	2	2	2	2	2	2
AF030881	2	2	1	7	0	1	1	1	1	1	1
				4	1	1	1	1	1	1	1
				2	1	1	1	1	1	1	1
				2	2	2	2	2	2	2	2
				2	2	2	2	2	2	2	2
AF022814	2	2	1	4	0	1	1	1	1	1	1
				7	1	1	1	1	1	1	1
				1	1	1	1	1	1	1	1
				2	1	1	1	1	1	1	1
				2	2	2	2	2	2	2	2
AJ010316	2	2	1	7	0	1	1	1	1	1	1
				4	1	1	1	1	1	1	1
				2	1	1	1	1	1	1	1
				2	2	2	2	2	2	2	2
				2	2	2	2	2	2	2	2
U63926	2	2	1	4	0	1	1	1	1	1	1
				7	1	1	1	1	1	1	1
				1	1	1	1	1	1	1	1
				2	1	1	1	1	1	1	1
				2	2	2	2	2	2	2	2
U92573	2	2	1	7	1	1	1	1	1	1	1
				4	1	1	1	1	1	1	1
				2	1	1	1	1	1	1	1
				2	2	2	2	2	2	2	2
				2	2	2	2	2	2	2	2
Totals					6	9	13	15	22	25	27
Proximity conservation (%)					11	16	23	26	39	44	47

^aIn cases where there is more than one candidate human chromosome, * marks the human chromosome with the highest numbers of top scoring BLAST hits, which was used in the calculation of the totals at the bottom. Some of these relationships to human chromosomes have previously been described by the original authors (Sandford et al., 1996; Aparicio et al., 1997; Arnes et al., 1997; Schofield et al., 1997; Miles et al., 1998; Bruner et al., 1999; Gellner and Brenner, 1999; Reboul et al., 1999).

^bThe quantity x is the largest allowed distance (in genes) between one of the human homologues and its nearest neighbour in the syntenous group. For the parts of the genome studied here, the intervals of x = 5, 10, 20, 50, 200, 1000 and 4000 genes correspond to average physical distances of 0.50, 1.27, 2.81, 7.05, 29.14, 144.01 and 494.04 cR, respectively.

synteny conservation. Simulating this problem requires knowledge of the distribution of gene family sizes, which we addressed in two ways. First, we used the distribution of the numbers of human BLAST hits to the *Fugu* proteins considered in Table 2 (plus annotated putative proteins, totaling 91) as an approximation of the distribution of family sizes. Second, we used the distribution calculated by Imanishi *et al.* (1997) from an all-against-all FASTA comparison of human proteins translated from mapped entries in DDBJ/EMBL/GenBank. In both cases the family sizes were scaled by a factor of 8/3 to account for unsequenced and unmapped genes. The latter (within-genome) method has the advantage that all the hits to a protein represent paralogues, whereas with between-genome comparisons the orthologues must be identified and removed before gene families can be examined. The results from the two methods

were similar and only those using the *Fugu* data are presented here.

Paralogous gene families were randomly assigned among the 80 000 genes in the simulated genome, according to the distributions described above. This process resulted in each simulated *Fugu* gene having one human orthologue, and possibly also a list of human paralogues, analogous to a list of BLAST hits. Some of the orthologues and paralogues could be 'unmapped'. Linkage conservation was measured by looking for the human homologues of 1000 pairs of adjacent *Fugu* genes, chosen at random. If the human orthologue of one (or both) of the *Fugu* genes in the pair was 'unmapped', a mapped paralogue from the list was used instead where possible. The extent of linkage conservation in human was then calculated, allowing various intervals between the human homologues. The simulation was run 30 times, looking at 1000 pairs of

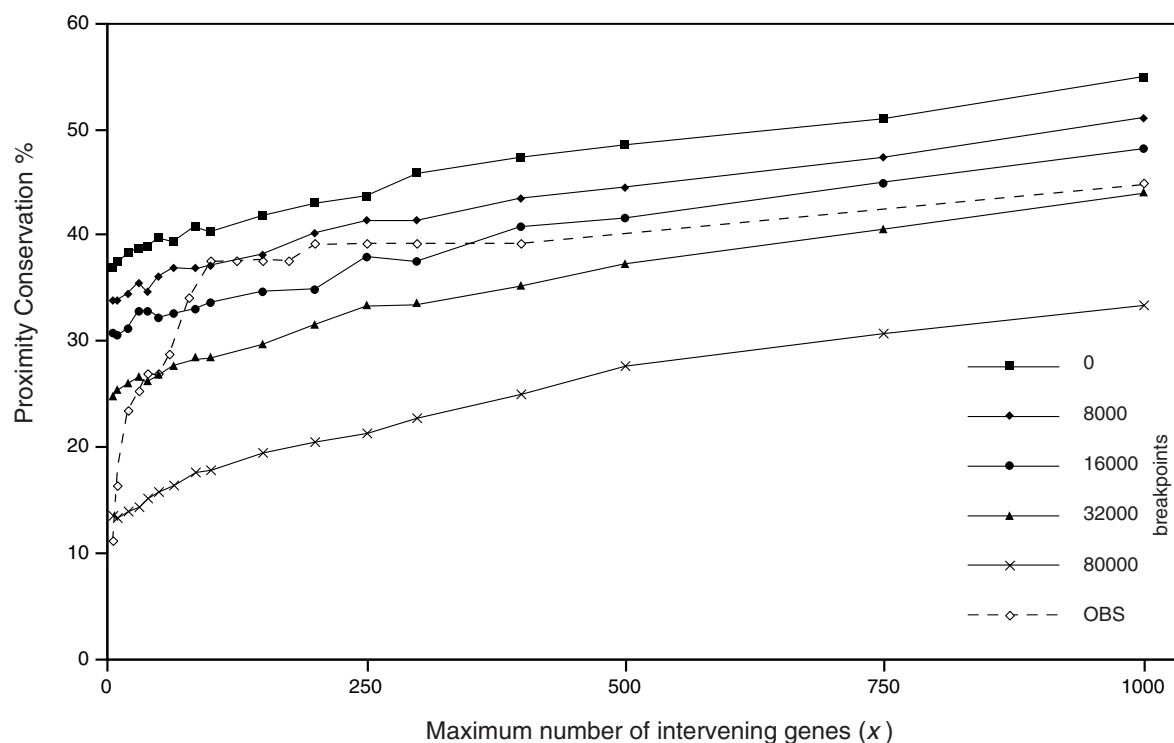


Figure 3. Extent of proximity conservation in real and simulated datasets. Proximity was measured allowing different gene distances between the human homologues of pairs of linked *Fugu* genes. The line marked 'OBS' graphs the observed data (Table 3). Average results for 30 computer simulations with 0, 8000, 16 000, 32 000 and 80 000 rearrangement breakpoints are shown (8000 breakpoints = 4000 rearrangements). The x axis is the limit used for the distance permitted between two human genes that are homologues of two *Fugu* neighbours, expressed in terms of the estimated number of intervening genes on the chromosome

genes each time, with the average results shown in Figure 3.

The most striking feature in the simulation results is that the presence of paralogues in an incompletely-sequenced genome has a substantial effect on the measured extent of linkage conservation. If there have been no genomic rearrangements (top curve in Figure 3), gene order conservation (and thus proximity conservation) should be 100%. However, the measured level is only 37%, because for many gene pairs, one or both of the human orthologues is 'unmapped' and a mapped paralogue at some other location in the genome has been used instead. This makes many linkages appear broken artefactually. Our measures of proximity conservation in the real data may also be underestimated to a similar degree (see Discussion). When the observed data from the fully-sequenced *Fugu* cosmids (Table 3) is plotted on the same axes, its initial slope is much greater than for the simulations (Figure 3). Possible reasons for this are discussed below. At large window sizes the line is approximately the same as the simulations with 8000–32 000 breakpoints.

Discussion

Although we confirmed the compaction of *Fugu* genes with respect to their human orthologues, we did not observe any strong relationship between gene compaction and the synonymous G+C content of the gene in either species. This may be an artefact of the sample analysed, or it may indicate true randomness in the compaction of the *Fugu* genome. There is an inverse relationship between the average compaction of the genes in each GC3 content category and their average GC3 content, which is consistent with expectations based on the lengths of genes in G+C rich isochores in vertebrate genomes (Duret *et al.*, 1995). However, this relationship is not strong enough to be predictive for individual genes.

The incomplete nature of the human genome data, and the uncertainty regarding whether homologues found in BLAST searches are orthologues or paralogues, reduces our power to examine synteny conservation between *Fugu* and human. The measured proximity conservation depends not only on whether the genes remain close or not, but also on whether they are mapped and sequenced, and if

there are paralogues of these genes in the mapped data. The simulations (Figure 3) suggest that the combination of incomplete sequence sets and the presence of gene families may cause the level of proximity conservation to be underestimated substantially, perhaps two-fold.

There is an obvious discrepancy between the slope of the graph of proximity conservation in real data from fully sequenced cosmids, as compared to the results from computer simulations (Figure 3). The observed proximity conservation rises steeply to 37% at a window size of 100 intervening genes, and then plateaus to a shape more like the simulated data. This suggests that the assumptions underlying the simulation are incorrect in some way.

The steep rise may be attributable to three primary factors. One possibility is that the real data is not a random sample of genes from the two organisms. A bias may result from *Fugu*'s role as a model vertebrate genome, inevitably influencing the selection of cosmids for complete sequencing. Cosmids with hypothesised synteny conservation with mammalian genomes may have been chosen preferentially. At least five of the *Fugu* complete sequences used had known synteny conservation with human chromosomes prior to sequencing (Aparicio *et al.*, 1997; Armes *et al.*, 1997; Sandford *et al.*, 1997).

Second, lack of resolution and incomplete data in GeneMap '98 data may affect the results. The arbitrary ordering of human genes that lie in the same radiation hybrid map interval could inflate apparent distances in human, although this effect is unlikely to be significant because the average number of genes per interval in the GeneMap '98 data used here is only 1.98. At least one distance in Table 3 has been overestimated due to missing data in GeneMap '98. This occurs with the genes TSC2 and PKD1 (*Fugu* Accession No. AF013614), which are neighbours in both species (Sandford *et al.*, 1996). However, PKD1 is not present in the map and instead our method identified a PKD1-like sequence elsewhere on chromosome 16 (Loftus *et al.*, 1999).

A third factor may be that our model of rearrangements is too simple. Our model assumed a random distribution of breakpoints throughout the genome, but comparative analysis of the human and mouse maps has shown that, although inter-chromosomal rearrangements seem to have random

endpoints, the number of intrachromosomal rearrangements is more than expected at random (Ehrlich *et al.*, 1997; Nadeau and Sankoff, 1998). The steep incline at the beginning of the graph may indicate a high frequency of small inversions or other small intrachromosomal rearrangements. Inversions of small segments of chromosome would disrupt gene adjacencies while preserving gene vicinities. This has been proposed by Gilley and Fried (1999), who noticed that some genes that are adjacent in *Fugu* are 2–4 Mb apart in human. Further examples from our study include *wnt10b*, *ARF3* and *erbB3*. These genes are adjacent in *Fugu* (Gellner and Brenner, 1999). In human, *wnt10b* and *ARF3* are adjacent but *erbB3* is separated from them by an estimated distance of 603 genes (226 mapped GenBank sequences scaled by 8/3 to allow for missing data) or 7.5 Mb [estimated from the map distance of 31 cR; chromosome 12 has an average of 234 kb/cR (Gyapay *et al.*, 1996)].

It is likely that the initial portions of the simulations in Figure 3 are not directly comparable with the observed data. However, as the window size gets larger the graph lines are approximately parallel to the plot of the observed data. From these an estimate of the extent of rearrangement since the divergence of these two lineages 400 million years ago is 8000–32 000 breakpoints (i.e. 4000–16 000 reciprocal translocations or inversions). This is higher than expected from comparisons of the human and mouse genomes, which diverged 100 million years ago and have only had an estimated 180 rearrangements (Nadeau and Taylor, 1984; Nadeau and Sankoff, 1998). Adjusting our simulations to incorporate a bias towards small rearrangements would only increase the estimated number of rearrangements since the *Fugu*–human divergence, making the discrepancy in rates even greater. Our estimate of the number of breakpoints depends somewhat on the estimated number of genes in the genome. Simulations based on a gene number of 61 000 instead of 80 000 (see Dunham *et al.*, 1999) led to an estimate of roughly 6000–12 000 breakpoints. Simulations using a gene number of 143 000 (recently suggested by Incyte Pharmaceuticals; see Dickson, 1999) produced the unexpected result that that no rearrangements were called for: the number of proximities observed in *Fugu* exceeded what would be expected due to the now-sparse sampling of genes, so either Incyte's estimate is unrealistic or

some of the orthologues listed in Table 3 are actually paralogues.

Another possible shortcoming of our analysis is the presence of short ESTs (which are not necessarily coding sequence) in the human DNA database used here, resulting in an overestimate of the frequency with which we can expect to find orthologues in this dataset from an amino acid level search. However, this is unlikely to have a great effect on the results, because we found that 78% of a random sample of over 500 human proteins submitted to TrEMBL after we downloaded GeneMap '98 were represented in the database. The gene family data is also likely to be oversimplified, as it is based on results from only 91 *Fugu* proteins. The Imanishi *et al.* (1997) data is from a larger set of proteins but is not as easy to relate to the human dataset used in this analysis.

Because we have approached the question of synteny conservation from the perspective of known gene adjacencies in *Fugu*, the proposed genome duplication in the bony fish lineage (Amores *et al.*, 1998; Meyer and Schartl, 1999), followed by differential gene loss, should not influence the results. If genes in the ancestral genome were ordered ABCD and this was duplicated in the fish lineage, differential gene loss could result in paralogous chromosomes, one bearing AC and another bearing BD. If synteny of these genes had not been disturbed, then the human genome would still contain the four genes arranged ABCD. If we were counting conservation of human linkages in *Fugu*, then we might plausibly have selected genes A and B for analysis and found that they are not syntenous in *Fugu*, an artefact of gene loss rather than genome rearrangement. However, as we are starting from the complementary viewpoint (given known relationships in *Fugu*), the only possible questions are 'Are A and C syntenous in the human genome?', and 'Are B and D syntenous in the human genome?', which is true in both cases. It is, however, possible that differential gene loss (after the genome duplication) in the *Fugu* lineage has contributed to the reduction of some intergenic distances as compared to human (e.g. the distance from A to C in the hypothetical example). This may also contribute to the steep initial slope seen in Figure 3. One example of apparent differential gene loss may already have been discovered in the case of the genes *IGF2* and *TH* (insulin-like growth factor and tyrosine hydroxylase), which are adjacent in

Fugu but separated by one intervening gene (insulin) in human (E. Chen *et al.*, unpublished; GenBank Accession No. AL021880; Lucassen *et al.*, 1993). Patterns of gene loss and gene order evolution should become clearer when more long homologous sequences from these species become available.

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