Comparative Sequence Analysis of the X-Inactivation Center Region in Mouse, Human, and Bovine

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We have sequenced to high levels of accuracy 714-kb and 233-kb regions of the mouse and bovine X-inactivation centers (Xic), respectively, centered on the *Xist* gene. This has provided the basis for a fully annotated comparative analysis of the mouse Xic with the 2.3-Mb orthologous region in human and has allowed a three-way species comparison of the core central region, including the *Xist* gene. These comparisons have revealed conserved genes, both coding and noncoding, conserved CpG islands and, more surprisingly, conserved pseudogenes. The distribution of repeated elements, especially LINE repeats, in the mouse Xic region when compared to the rest of the genome does not support the hypothesis of a role for these repeat elements in the spreading of X inactivation. Interestingly, an asymmetric distribution of LINE elements on the two DNA strands was observed in the three species, not only within introns but also in intergenic regions. This feature is suggestive of important transcriptional activity within these intergenic regions. In silico prediction followed by experimental analysis has allowed four new genes, *Cnbp2*, *Ftx*, *Jpx*, and *Ppnx*, to be identified and novel, widespread, complex, and apparently noncoding transcriptional activity to be characterized in a region 5' of *Xist* that was recently shown to attract histone modification early after the onset of X inactivation.

[The sequence data described in this paper have been submitted to the EMBL data library under accession nos. AJ421478, AJ421479, AJ421480, and AJ421481. Online supplemental data are available at http://pbil.univ-lyonl.fr/datasets/Xic2002/data.html and www.genome.org.]

In mammals, dosage compensation of X-linked genes is achieved by the transcriptional silencing of one of the two X chromosomes in the female cell during early development, a process known as X inactivation. Initiation of X inactivation involves recognition of the number of X chromosomes present in the cell, ensuring that the single X chromosome remains active in the diploid male cell and that only a single X chromosome is inactivated in the female diploid cell. This process, which is known as counting, is thought to involve an evaluation of Xic number against ploidy. Initiation also includes a recognition process linked to the choice of the X chromosome to be inactivated. Initiation of X inactivation and other early events are regulated by a master control region, the Xic (X-inactivation center). The XIC/Xic is a unique region of the X chromosome situated in Xq13 in man and in the syntenic mouse region that is necessary for the counting, the choice, and the subsequent nucleation of silent chromatin on the presumptive inactive X. Silencing spreads bidirectionally from the Xic into linked sequences, which need not

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be of X-chromosome origin (Lee and Jaenisch 1997). The study of chromosomal rearrangements in human culminated in the identification of a 680-kb to 1.2-Mb candidate region that shows full XIC function (Rastan 1983; Rastan and Brown 1990; Brown et al. 1991a). Subsequent efforts to delimit the XIC/Xic have concentrated on the use of transgenesis in the mouse to determine the candidate region that is sufficient for Xic function. Using the stringent criteria that single-copy transgenes must show full Xic function, these experiments have not as yet defined the minimum size of the Xic necessary for ectopic function. Although a single copy of a 35-kb cosmid transgene recapitulates some aspects of Xic function (Herzing et al. 1997), other studies of transgene copy-number dependence have suggested that even a 450-kb region may not contain all the elements necessary for autonomous, ectopic Xic activity (Heard et al. 1999).

As defined cytologically, Xic has been shown to contain at least five genes (Heard et al. 1997; Avner and Heard 2001). One of these, the *Xist* (X-inactive specific transcript) gene, which is the only gene known to be specifically transcribed from the inactive X chromosome in female somatic cells, codes for a 17-kb spliced, polyadenylated noncoding RNA (Borsani et al. 1991; Brockdorff et al. 1991; Brown et al. 1991b). *Xist* is necessary and sufficient for the initiation and

spread of X inactivation but not for counting (Penny et al. 1996; Marahrens et al. 1997; Wutz and Jaenisch 2000).

At least some of the sequences required for counting must lie immediately 3' to Xist as a 65-kb deletion of a region extending 3' downstream of Xist exon 6 disrupts the counting process (Clerc and Avner 1998). This region contains a 17-mer minisatellite (Simmler et al. 1996), the Tsx gene (Cunningham et al. 1998), the DXPas34 locus, a CpG-rich minisatellite showing a highly characteristic pattern of hypermethylation on the active X chromosome (Courtier et al. 1995; Prissette et al 2001) and an associated CpG island that is the presumptive major initiation site for the Xist antisense transcript Tsix (Lee et al. 1999; Mise et al. 1999). Tsix may have a repressive role on Xist, with which it overlaps (Debrand et al. 1999; Lee and Lu 1999; Morey et al. 2001; Stavropoulos et al. 2001), and has also been implicated in imprinted X inactivation in murine extraembryonic tissues (Lee 2000; Sado et al. 2001). A spliced form of Tsix, initiating upstream of DXPas34, has been described (Sado et al. 2001) and it is likely that antisense transcriptional activity is widespread throughout the region (Debrand et al. 1999). The region between Xist and DXPas34, which includes the major Tsix initiation site, regulates Xist transcript accumulation and its retention at the site of transcription (Morey et al. 2001), probably through the activity of the Tsix antisense (Stavropoulos et al. 2001).

Another regulatory element within Xic is the X-controlling element, or *Xce* (Cattanach and Williams 1972). *Xce*, which lies 3' to *Xist* and the counting region (Simmler et al. 1993; M. Prissette, unpubl. data), influences the choice of which X chromosome is to be inactivated (Heard et al. 1997). *Xce* is, however, only one element regulating choice, and other regions lying both within the *Xist* gene and 3' to the *Xist* gene also influence chromosome choice (Avner and Heard 2001).

Several years ago, we obtained high-quality sequence of a 94-kb region encompassing the mouse *Xist* gene (Simmler et al. 1996) and the region lying 3' to *Xist*, which was subsequently shown to be involved in the counting process and in choice (Clerc and Avner 1998; Morey et al. 2001). More recently, the *Xist* gene and a small region immediately upand downstream have been sequenced in the vole (Nesterova et al. 2001) and compared with the homologous human and mouse sequences. The overall picture obtained from functional analysis of the mouse Xic is of a complex integrated locus in which functionally important elements may be located over several hundred kilobases both upstream and downstream of *Xist* itself. In this context, we decided to undertake a fully annotated comparative analysis of the human and mouse XIC/Xic based on the sequencing to high stan-

dards of a 714-kb region including the murine *Xist* gene, which was compared to the 2.3 Mb of available finished sequence for the orthologous region in human. Sequencing of a 233-kb core region of the bovine Xic around *Xist* to similarly high standards allowed a three-way species comparison of the core central region, including the *Xist* gene.

RESULTS

Global Description of the Region

We sequenced a region of 714 kb of the mouse X chromosome, centered on the Xist gene, and a region of 233 kb from the bovine Xist locus. The human sequence orthologous to the mouse locus was identified by a BLAST search and extracted from the Human Genome Project Working Draft Assembly (October 7, 2000, freeze; http://genome.ucsc.edu/; Lander et al. 2001). We searched for genes in the three species by using a combination of different approaches: ab initio exon prediction (GENSCAN, FGENESH), homology-based gene prediction (BLASTX, GENEWISE), ESTs, or cDNAs (BLASTN, SIM4; see Methods). GENSCAN or FGENESH exon predictions were considered as belonging to true genes only if they were confirmed by other evidence (cDNA sequencing or positive RT-PCR between adjacent exons or finding of ESTs covering adjacent exons). Mouse, human, and bovine sequences were compared with each other using SIM to identify conserved blocks corresponding to potential functional elements (coding or noncoding).

We identified 11 genes in the mouse Xic region. Seven of these genes were previously known: Xpct, Xist, Tsx, Tsix, Chic1 (formerly, Brx), Cdx4, and NapIl2 (formerly, Bpx). We characterized 4 new genes: Cnbp2, Ftx, Jpx, and Ppnx. Four of the 11 genes, Xist, Tsix, Ftx, and Jpx, are untranslated RNA genes (i.e., do not code for proteins). All the genes identified in mouse are conserved in human, except *Ppnx* and *Tsix* (see Fig. 1). In human, however, Tsx has become a pseudogene. The human XIC locus is considerably expanded compared to its mouse ortholog. It covers ~2300 kb (the exact size is not known because there is a gap in the human assembly between the CHIC1 and CDX4 genes), that is, about three times larger than in the mouse (Figs. 1,2,3). Despite this major change in locus size, the order and orientation of genes is perfectly conserved in human and mouse, except for Xpct, which is at the same location but in the inverse orientation (Fig. 1).

The XIC/Xic region is relatively GC-poor in all three species (42% in mouse, 40% in human, and 39% in bovine) and repeat elements (essentially LINEs and SINEs) make up 38%, 57% and 58% of the mouse, human, and bovine loci, respectively (Table 1). In mouse, we identified 22 pseudogenes (i.e.,

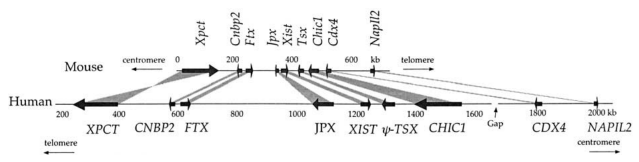


Figure 1 Comparative map of Xic region in mouse and human.

sequences with similarity to known functional genes but which contain stop codons or frameshifts in the coding region). The density of pseudogenes in the mouse Xic region (31 pseudogenes/Mb) is relatively high compared to estimates

from the human genome overall (6 per Mb; Goncalves et al. 2000). Protein-coding regions of confirmed genes, make up 0.84% of the Xic region in the mouse, 0.24% in the human XIC (compared with ~1.8% for the whole human genome; Lander et al. 2001). Thus, globally, the region is relatively G+C-poor, repeat-rich, and poor in protein-coding genes.

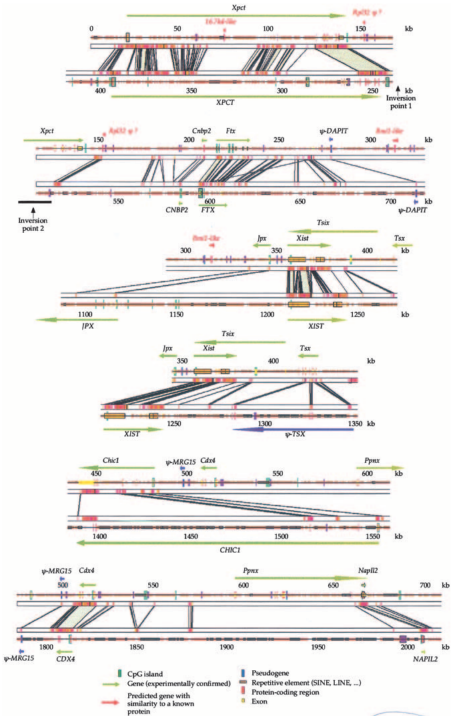


Figure 2 Comparison of mouse and human genomic sequences in Xic region. Genomic sequences were first analyzed with RepeatMasker to identify and mask repeated elements and then aligned with SIM. Conserved blocks with a similarity score >30 are displayed (see Methods). An electronic version of this figure is available at http://pbil.univ-lyon1.fr/datasets/Xic2002/data.html. Because the alignment is very long and because of an inversion of the *Xpct* gene, it was not possible to display the whole human—mouse comparison in a single continuous line. The alignment, therefore, is displayed in six overlapping fragments. The overlaps are designed to allow the continuity of the fragments to be better appreciated.

Gene Content

New Genes

Cnbp2: A Novel Gene Coding for a Zinc Finger Protein

This new gene, located ~150 kb 5' of the Xist gene, was predicted in the mouse by both GENSCAN and FGENESH and was confirmed by the identification in GenBank of a fulllength cDNA (accession number AK015789). The predicted open reading frame (ORF) encodes a protein of 170 amino acids with strong similarity (75% identity) to cellular nucleic acid binding protein (CNBP). CNBP is a zinc-finger DNAbinding protein of unknown function, highly conserved in vertebrates, having two possible forms due to alternative splicing (De Dominicis et al. 2000). In human, CNBP maps to chromosome 3 (3q13.3-q24) and consists of five exons. The new predicted protein aligns perfectly with the shortest of the two forms of CNBP (170 amino acids long) and hence is likely to have similar biochemical activities. We therefore named our gene Cnbp2. The Cnbp2 gene contains a single exon and is conserved in human (Fig. 2), in which we found several matching ESTs. The fact that Cnbp2 is conserved and expressed both in human and mouse suggests that it is a real gene and not a pseudogene despite its intronless nature suggestive of a retroelement. This is also supported by the comparison of the human and mouse orthologs: The ratio of synonymous over nonsynonymous substitution rate of 3.6, (i.e., much greater than one, indicating that this gene is under selective pressure). We found no evidence of other closely related CNBP ho-

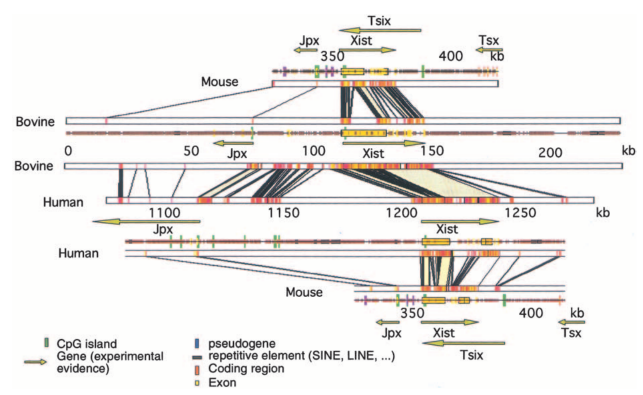


Figure 3 Three-way comparison of mouse, human, and bovine genomic sequences around *Xist*. See Fig. 2 legend. An electronic version of this figure is available at http://pbil.univ-lyon1.fr/datasets/Xic2002/data.html.

mologs in the human genome. An RT-PCR product was detected only in adult mouse testis RNA, suggesting that *Cnbp2* is not widely transcribed.

Ftx: A Novel Conserved Noncoding Gene

We found in GenBank a mouse cDNA (AK020989) that spans five exons of a new gene, located \sim 140 kb in 5′ of *Xist* and that we have named *Ftx*. We found several overlapping mouse ESTs (BB619145, BB628053, BB660326, BG920590, and BB305979) that identifed two additional exons and revealed

at least four different transcripts resulting from alternative splicing. The seven exons identified all have perfect consensus splice sites. We identified a human cDNA (AK057701) spanning four exons in the corresponding region and sharing the same orientation. The 5'-part of the gene is well conserved and contains a CpG island in both species (Fig. 2). Both the mouse and the human cDNAs start in this conserved CpG island, which suggests that it might correspond to the promoter region. The intron–exon structure of the gene is poorly conserved: Only four of the seven mouse exons are located in

Table 1. Frequency of the major classes of interspersed repeated elements (in term of percent of the sequence length) in the autosomes, the X chromosome, in the Xic region taken as a whole and in introns of genes located in the Xic region

		Length (kb analyzed)	LINE (%)	SINE (%)	LTR (%)	DNA (%)	Total (%)
Mouse	Autosomes ^a	223,057	10.5	9.4	6.9	0.5	27.4
	X chromosome ^a	10,000	16.2	9.1	6.5	0.7	32.6
	Xic region	714	14.5	14.5	8.5	0.3	37.9
	Xist introns	9	0.0	10.2	0.0	0.0	10.2
	Other introns ^c	190	17.2	13.2	3.1	0.4	33.9
Human	Autosomes ^b	348,845	17.0	14.3	6.8	2.4	40.6
	X chromosome ^b	56,667	30.2	9.6	9.0	2.5	51.2
	XIC region	2300	38.9	8.4	8.6	1.3	57.2
	XIST introns	18	8.2	8.9	0.5	1.0	18.6
	Other introns ^c	352	47.5	7.8	9.0	2.0	66.3
Bovine	XIC region	233	46.5	9.3	1.3	0.8	57.9
	XIST introns	17	17.8	5.8	0.6	0.0	24.2

^aSequences extracted from GenBank release 124 (June 2001).

^bData from Bailey et al. (2000).

^{&#}x27;Introns of protein-coding genes located in Xic region.





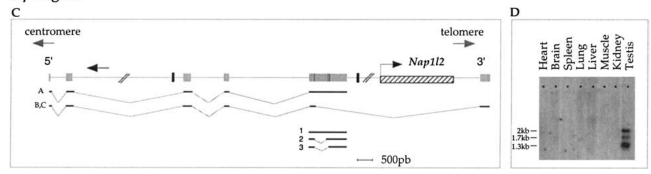


Figure 4 Cloning and characterization of *Ppnx* and *Jpx* genes. (*A) Jpx* gene. Schematic diagram showing the three exons (light grey boxes) of the murine *Jpx* gene, at positions 343745–343379 (exon 1), 340888–340796 (exon 2), 334465–334364 (exon 3), as defined by the three ESTs represented as dark grey boxes above the map. ESTs are identified by their GenBank accession number. The presence of an alternative splice at position 343626 nt in exon 1 generates either a 315- or 562-nt transcript. The presence of a poly(A) tail in EST BE994494 identifies the 3' end of the gene. The first exon may correspond to the 5' end of the gene, as a CpG island (diagonally hatched box) lies in the vicinity. The two last exons are LTR/MaLR-repeat elements (white boxes). (B) Northern analysis using a PCR probe corresponding to exon 1 of the Jpx gene after 5 d exposure. (C) Ppnx gene. Schematic diagram illustrating the six exons of the murine Ppnx gene. All exons and introns are represented to scale with the exception of introns 2 and 5 that are very large (21707 bp and >45389 bp, respectively). Each exon (grey) is identified by its coordinates: exon 1, 591535-591619; exon 2, 592146-592314; exon 3, 614021-614268; exon 4, 615349-615488; exon 5, 618116->619619 (with three alternative splice donor sites at position 618262, 618719, and 618758), and exon 6, 665008–665306. The black boxes represent two pseudogenes predicted by GENSCAN; the small black arrow shows an antisense transcription detected in intron 2. The single-exon Napll2 gene is shown diagonally hatched. Three independent clones A, B, and C obtained after screening a testis cDNA library are shown below the genomic structure. Clone A does not contain either a poly(A) consensus signal or a poly(A) track and has an in-frame stop codon present in its most 3' part. The end of the 3' UTR track is likely 300 bp downstream of the stop codon (unpubl.). Clones B and C possess a stop codon in exon 6 followed by a 3' UTR of 177 bp, containing both a poly(A) signal and a poly(A) tail. Three of the RT-PCR products suggesting the existence of alternative splice donor sites in exon 5 are also represented (1, 2, and 3). (D) Northern analysis of Ppnx expression. A mouse multiple tissue Northern blot was hybridized with a probe corresponding to the third exon of Ppnx. The Ppnx hybridization signals were obtained after 8 d exposure.

regions that are conserved in human, and only two of them overlap with the exons identified in human. We found SINE elements both in human and mouse transcripts, and we detected no similarity with any known protein (after masking repeated elements). The longest mouse ORF contains only 87 codons and is not conserved in human, which suggests that *Ftx* encodes an untranslated RNA. RT-PCR studies, using primers designed from within exon 7, suggest that the *Ftx* gene is ubiquitously expressed (undifferentiated ES cells, liver, brain, kidney, and testis have been tested; data not shown).

Jpx: A Novel Conserved Noncoding Gene Subject to X Inactivation

A small region proximal to the *Xist* gene, which is well conserved between mouse, human, and bovine, was identified (Fig. 3). As this region also contains a conserved CpG island, the presence of a transcript was suspected. Several ESTs were

found to match perfectly to this region in all three species, identifying a novel gene we have named *Jpx*. The murine *Jpx* gene, located in the mouse 10 kb upstream of Xist, is composed of three exons identified by three ESTs (accession numbers BE994494, AI391037, and BB564398), with alternative splice donor sites in the first exon generating 315-nucleotideand 562-nucleotide-long transcripts (Fig. 4). A polyadenylation consensus motif (AATAAA) is present in the third exon, followed in several of the ESTs by a poly(A) tail 16 nt downstream, indicating that the third exon likely represents the true 3' end of *Ipx*. The CpG island lies within the first exon, suggesting that exon 1 may represent the 5' end of the gene. This first exon is highly conserved in all three species. In both human and bovine, transcription of this exon with correct splice signals is confirmed by several matching ESTs. Human and bovine ESTs (AV714079, AW484353, and BE485548) indicate the presence of three exons as observed in the mouse, sharing the same orientation. All putative exon-intron junctions have a consensus splice-site sequence. The second human exon is a homolog of mouse exon 1. However, the last two mouse exons are not conserved. Indeed, in the three species, the last two exons (except human exon 2) correspond to repeat elements: The mouse second and third exons show perfect homology with LTR/MaLR elements, whereas the bovine second and third exons correspond respectively to LINE and SINE repeats and the human third exon to a SINE repeat. It is highly unusual to find repeat sequences functioning as exons as in Jpx. In the mouse, neither of the two alternative transcripts contains an ORF large enough to encode a protein: the longest ORF is only 129 bp long (43 amino acids) and is not conserved in human or bovine. This suggests that Jpx, such as Ftx, Xist, and Tsix, encodes an untranslated RNA. The fact that this gene is expressed and clearly conserved in all three species suggests a functional role for Jpx.

RT-PCR studies suggest that the *Jpx* gene is ubiquitously expressed, with Northern analysis revealing several low-level transcripts of different size (Fig. 4). Although we have not definitively identified the site of initiation of these transcripts, the presence of a conserved CpG island lying within the first exon of *Jpx* and the lack of conservation of genetic elements lying proximal to the *Jpx* region among all three species, suggests that they originate in exon 1 rather than more proximally. We cannot, however, formally rule out that these *Jpx* transcripts are not the 3' UTR sequences of an as-yet unidentified, ubiquitously expressed longer mRNA(s). It is unlikely however that these *Jpx* transcripts represent the 3' end of a variant *Tsix* transcript because the expression profiles are clearly distinct.

The mouse *Jpx* gene was shown to be subject to X inactivation (see Methods) and transcribed only from the active X chromosome. These findings allow the 5' end of the domain exclusively expressed from the inactive X, which includes *Xist* to be clearly delineated for the first time.

Ppnx: A Mouse-Specific Gene Expressed in Testis and ES Cells

Five of the six exons on the forward strand and lying in the vicinity of *Nap1l2* that characterize the *Ppnx* gene were predicted by both GENSCAN and FGENESH. Each of these exons was shown by random RT-PCR to be strongly expressed in both undifferentiated ES cells and the adult testis. More detailed analysis revealed the presence of these transcripts in the testis from 14 days postcoitum onward, suggesting expression in germ cells having entered meiosis. Strand-specific RT-PCR showed that the exons belong to a single gene, with a centromere 5′–3′ telomere orientation. Subsequent Northern blot analysis of adult and embryonic tissues confirmed the presence of *Ppnx* expression in the adult testis (Fig. 4).

The *Ppnx* gene contains two particularly large introns (22 and >45 kb, respectively, for introns 1 and 5) and several alternative exons, giving rise to alternative splice forms (see Fig. 4). Two different ORFs of 1815 and 873 nt were identified from the three cDNA clones we obtained, depending on the start and stop codons and the poly(A) consensus signal or poly(A) track used (for more details, see Fig. 4). Virtual translation of these clones gave protein products of 604 and 290 amino acids, respectively. Both putative proteins contained a common secretion-motif associated with the first 20 amino acids. In silico searches for protein similarities using BLASTP showed that both proteins share 20%–30% identity with trypsin due to the presence of a serine protease-like domain.

Ppnx is not conserved between mouse and human, at least within the Xic/XIC regions (Fig. 1). The only human homolog of *Ppnx* that could be detected when we compared the different *Ppnx* splice forms against protein databases (using BLASTP) and EST or genomic databases (using TBLASTN) located to the human Y chromosome. This human-coding sequence is interrupted by two stop codons and does not match to any human EST, suggesting that it probably corresponds to a pseudogene.

Potential Genes

We identified three putative genes, both predicted by GEN-SCAN and FGENESH, that showed significant similarity to other known proteins: a BMI1-like gene, a 16.7 kd-like gene, and a RPL32-like gene (Fig. 2). None of these putative genes is conserved in human. BMI1-like and 16.7 kd-like match several ESTs with ~90%–95% identity. However, the comparison of these ESTs with the assembly of the mouse genome (Ensembl database, v. 4.1.1; http://www.ensembl.org) revealed that they correspond to transcripts from closely related paralogous genes located in other regions of the X chromosome. In the absence of evidence of transcription activity, we are unable to conclude whether the BMI1-like and the 16.7 kd-like elements correspond to real genes. The RPL32-like element, on the other hand, is a retroelement that is 100% identical to the mouse ribosomal protein L32 (RPL32) mRNA but does not contain any introns and most likely corresponds to a recently inserted retropseudogene. This RPL32-like element is potentially functional, due to the fact that no frameshift or stop codon, which might stop it encoding a protein identical to RPL32, was noted.

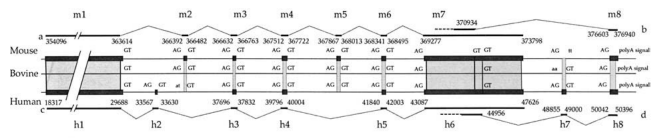
Previously Known Genes

Xpct: Alternative Splicing in First Intron

The *Xpct* gene (X-linked PEST-containing transporter; Debrand et al. 1998) encodes a protein of 613 amino acids that belongs to a family of monocarboxylate transporters (Lafreniere et al. 1994; Price et al. 1998). The gene contains 6 exons that are perfectly conserved in both species. *Xpct* spans 117 kb in mouse and 147 kb in human. The first intron is very long (104 and 135 kb in mouse and human, respectively), and contains remarkably large conserved regions (Fig. 2). In mouse, we found one EST (accession number AW106665) that matches exons 2 and 3 and an alternative exon (with consensus splice signals) located in a conserved region of the first intron. This suggests that *Xpct* might undergo alternative splicing or alternative promoter use in both species.

Xist: Evolution of Its Gene Structure in Mammals

Although the *Xist* gene that is conserved in mouse, human, and bovine (Fig. 3) has been studied by many different groups, its precise exon–intron structure is still not fully established, notably at the 3' end. We identified *Xist* exons in human and mouse by comparison with ESTs and with *Xist* cDNA sequences published in GenBank. Debate on the human *XIST* gene has concerned the seventh and eighth exons, these being found only in rare splice variants (Brown et al. 1992). Although Hong and colleagues (2000) failed to identify human exon 7, we found three ESTs from human hepatocellular carcinoma cells that clearly confirm the existence of *XIST* exons 7 and 8 (Fig. 5). In agreement with Hong et al. (1999), we found that the mouse cDNA published by Brockdorff et al. (1992) corresponds to seven exons rather than the



■ Exon (experimentally confirmed)

[] Conserved region corresponding to a known exon in at least one species

Figure 5 Comparison of Xist gene intron-exon structure in mouse, bovine, and human. Introns and the first exon are not drawn to scale. Conserved regions corresponding to known exons (in at least one species) are indicated. Consensus splice signals that align with boundaries of known exons are in capitals. In mouse and human, 8 exons have been identified (Brown et al. 1992; Sheardown et al. 1997). Note, however, that some exons in mouse have no counterpart in the human and vice versa. To avoid ambiguities in exon numbering, we used the prefix "h" to identify human exons, and "m" to identify mouse exons. Mouse exons m1, m3, m4, m6, m7, and m8 are conserved and correspond to human exons h1, h3, h4, h5, h6, and h8, respectively. In both species, we identified an alternative donor splice site in exon m7-h6, although it is not located exactly at the same position in human and mouse. With the exception of this splice site, splice signals (donor and acceptor) of these six exons, as well as the polyadenylation signal in the last exon, are perfectly conserved in mouse, man, and bovine (which suggests that they also correspond to true exons in the latter species). Human exon h2 is located in a region that is not conserved in mouse or bovine. Mouse exon m2 is conserved in the three species, with correct splice signals, except in human where the splice acceptor is mutated (AT instead of AG), which suggests that this exon is no longer functional in human. Mouse exon m5 is conserved in the three species with correct splice signals and might correspond to an unidentified alternative exon in humans. Human exon h7 is located in a region that is conserved in the three species, but consensus splice signals are not found in mouse and bovine. a, mouse Xist cDNA (GenBank accession number L04961; Brockdorff et al. 1992); b, mouse ESTs (BE626785, BE632200, and R74734); c, mouse exon m7a reported by Hong et al. (1999); d, human XIST cDNA (M97168; Brown et al. 1992); e, human exon h6a reported by Hong et al. (2000); f, human ESTs (AV699347, AV700119, and AV700677). Exon positions in mouse (bp): m1, 354096. . . 363614; m2, 366392...366482; m3, 366632...366763; m4, 367512...367722; m5, 367867...368013; m6, 368341...368495; m7a, 369277...376940; m7b, 369277...371600; and m8, 376603...376940. In human, exon positions are indicated relative to the X/ST genomic sequence (U80460): h1, 18317...29688; h2, 33567...33630; h3, 37696...37832; h4, 39796...40004; h5, 41840...42003; h6a, 43087...50396; h6b, 43087...44956; h7, 48855...49000; and h8, 50042...50396.

six exons originally suggested. Based on sequence conservation and RT-PCR analysis, it has also been proposed that mouse Xist contained an additional 3' exon, homologous to human exon 8 (Simmler et al. 1996; Sheardown et al. 1997). This eighth exon has been described in voles (rodent; Nesterova et al. 2001). We have confirmed the existence of mouse exon 8 by comparison with two ESTs from mammary gland (Fig. 5). As Hong and colleagues (1999, 2000) failed to identify this eighth exon, it may only be used, however, in rare splice variants. We conclude therefore on the presence of eight exons and of clear evidence of alternative splicing and poly(A) tail use at the 3' end of the Xist gene in both species. Although in mouse and human the longest Xist mRNA forms span 17.9 and 19.3 kb, respectively, the splice variants of the alternative 3'-end exons (m8, h7, and h8; for nomenclature, see Fig. 5 legend) result in RNAs that are at least 5 kb shorter.

We aligned the *XIST/Xist* genes in mouse, human, and bovine and compared the location of exons identified in mouse and human (Fig. 5). Seven out of the eight mouse exons are perfectly conserved in the three species. Six of these seven have been confirmed experimentally in human; one (exon m5) may correspond to an unidentified splice variant. The intron–exon structure, however, is not totally conserved. The human *XIST* gene includes two exons (h2 and h7) that are not present in mouse and bovine, and the exon m2 found in the mouse and bovine genes is probably not functional in human.

At the sequence level, *Xist* exons show on average 66% and 62% identity, respectively, between mouse and human, and mouse and bovine. This figure is close to the average conservation in 5'- and 3'-untranslated regions of human and mouse orthologous protein-coding genes (Makalowski and Boguski 1998). With the exception of several conserved

blocks, *Xist* introns are weakly conserved and cannot be aligned between the mouse and the two other mammals. By comparison with the rest of the Xic locus, the *Xist* gene is relatively devoid of LINE elements, not only in exons (LINE elements make up 0%, 0.4%, and 1.9% of *Xist* exons in mouse, human, and bovine, respectivley), but also in introns (see Table 1). Note that, unlike *Xist*, introns of protein-coding genes are generally rich in LINE repeats (Table 1). These observations suggest that the insertion of LINE elements within the *Xist* gene has been strongly counterselected (see below).

Tsix: No Evidence of Conservation in Other Mammals

The Tsix gene, identified in mouse, encodes an untranslated RNA antisense to Xist. Initially, Tsix was described as a 40-kb RNA encoded by a single exon, initiating close to a CpG island located 15 kb 3' of the previously described end of Xist (i.e., 12 kb 3' of the Xist m8 exon; Fig. 3; Lee et al. 1999). However, Sado and colleagues (2001) found that *Tsix* is, at least in part, subject to processing. They identified four exons, along with a major promoter upstream of Tsix exon 2 and a minor initiation site located upstream of Tsix exon 1 (i.e., ~28 kb 3' of the Xist m8 exon). Lee and colleagues (1999) have identified three conserved blocks between human and mouse, two of which are located close to the start of the Tsix transcription unit; this led them to suggest that the Tsix gene is conserved in human. The first conserved block corresponds to Xist exons m8 and h7 and h8. The two other blocks are located within Tsix intron 3, downstream of the major promoter. It is noteworthy that neither of these two latter blocks are conserved in bovine (Fig. 3). Tsix exon 4 overlaps with the 5' end of Xist exon 1. Although the 5' part of Tsix exon 4 that is common to Xist exon 1 is conserved in the three species, the 3' part is not conserved. Therefore, in contrast to Xist, which is well conserved in the three species, neither *Tsix* exons nor promoter regions are conserved in mammals. Thus, if *Tsix* does exist in human and bovine, it is clear that its primary sequence is not subject to a strong selective pressure.

Tsx: A Human Pseudogene

Tsx is a testis-specific gene of unknown function (Simmler et al. 1996). The mouse *Tsx* gene, which spans 10 kb, contains 7 exons and encodes a protein of 144 amino acids without homology to any known gene. The orthologous region in human spans 44 kb. Conserved blocks that correspond to exons 1, 3, 4, 5, and 6 are clearly identified (Fig. 2). Exon 2 is not conserved (although we found conserved blocks within introns 1 and 2) close to the exon-2 boundaries. Human exons 4 and 6 contain stop codons, exons 1 and 3 contain frameshifts, and intron 5 contains mutated splice-donor signals, all of which indicates that Tsx has become a pseudogene in human. We found no ESTs matching these conserved blocks in human, and no other regions with significant similarity to Tsx in the complete human genome sequence, which suggests that the inactivation of Tsx gene in the primate lineage has not been compensated for by another homologous gene located elsewhere in the genome. Note that the comparison of mouse and rat Tsx cDNAs has shown that this gene has evolved very fast in rodents, which suggests that it is under weak selective constraints (Simmler 1996).

Unexpectedly, a region with similarity to mouse *Tsx* exon 7 was found in human, translocated on to the opposite DNA strand. Interestingly, the region corresponds to *XIST* exon 8. The similarity is weak but spans the acceptor splice signal, the polyadenylation site, and a block within the last intron. This observation suggests a possible evolutionary relationship between the genes.

Chicl: A Revised Structure

Chic1 which belongs to the family of CHIC genes (cysteinerich hydrophobic proteins) is specifically expressed in the brain (Simmler et al. 1997). The 653 nt originally identified Chic1 coding sequence was suspected to be incomplete as no in-frame stop codon was detected upstream of the first ATG of the cDNA and because an 8-kb transcript using a specific Chic1 probe was detected by Northern blot. However, a comparison of the mouse and human genomic sequences 5' of this ATG failed to reveal conserved sequence with significant coding potential. Moreover, a comparison of Chic1 with others members of the family, all of which code for proteins of approximately the same size and which start with the same conserved motif, suggests that the 653-nt sequence almost certainly represents the complete coding region of Chic1. The 6-kb region distal to exon 6 of the Chic1 gene is highly conserved between human and mouse (Fig. 2) and sequence from this region match several ESTs, suggesting that the region is transcribed. However, the absence of an extended ORF suggests that the region is not translated. To test the hypothesis that this 6-kb conserved region corresponds to the 3' UTR of the Chic1 gene, strand-specific RT-PCR was carried out on female brain RNA, using primers positioned at several points within the 6-kb region. Our results show the presence of a continuous transcription unit with the same orientation as Chic1, extending from the last exon of Chic1 defined by Simmler and colleagues (data not shown), compatible with it being a 3' UTR. The complete Chic1 gene, including the six exons and the very long 3' UTR are perfectly conserved in human and mouse, even though the gene is highly expanded

in human compared to mouse. The *CHIC1* gene spans 40 kb in mouse and 163 kb in human (Fig. 2).

Cdx4

Cdx4 encodes a protein of 282 amino acids and belongs to the caudal family of homeobox genes (Gamer and Wright 1993). The gene spans ~8 kb in mouse and human and contains three exons that are perfectly conserved (Fig. 2).

Napll2: An Unusual Location in the Intron of a Gene Sharing the Same Orientation

Nap1l2 (nucleosome assembly protein 1-like 2) is a singleexon gene, encoding a protein of 460 amino acids conserved in human and mouse (Fig. 2), which is specifically expressed in neurons (Rougeulle and Avner 1996; Rogner et al. 2000). Surprisingly, we found the Nap112 gene to be entirely included in the last intron of *Ppnx*, a newly identified mouse gene (see above). Several other examples of genes located within introns have already been reported, but, to our knowledge, they are all oppositely orientated on the antisense strand, whereas Nap112 and Ppnx share the same orientation. Ppnx and Nap112, however, have different expression patterns. Although *Ppnx* is found only in the adult testis, *Nap1l2* is exclusively expressed in nervous tissue (Rougeulle and Avner 1996). We suppose that the absence of transcription interferences between these genes is due to their different expression pattern and the fact that Nap112 is composed of only one exon and thus contains no splicing site that could interfere with the splicing of *Ppnx*.

Patterns of Conservation

Conserved Genes

Overall, conserved blocks between the mouse and human Xic/XIC represent 76 kb (10.6% of the mouse locus; 3.3% of the human locus). Of these 76 kb, 62 kb are located within known genes (introns and exons of Xpct, Cnbp2, Ftx, Jpx, Xist, Tsx, Chic1, Cdx4, and Nap112). Thus, 35 out of the 48 identified exons in the Xic mouse sequence are located within conserved blocks. The 13 nonconserved exons correspond to the 6 exons of the *Ppnx* gene (absent in human), 2 of the 7 *Tsx* exons (a pseudogene in human), 3 out of the 7 Ftx exons, and 2 out of the 3 Jpx exons. Conserved regions span 28% of the total gene length within the mouse Xic, consistent with previous observations (Jareborg et al. 1999), whereas only 3.4% of intergenic regions are conserved. This is considerably lower than a recently published estimate (15.8%), which was based on the analysis of intergenic regions of 100 pairs of human and mouse orthologs (Shabalina et al. 2001). This discrepancy is probably due to the fact that the data set analysed by Shabalina and colleagues (2001) was biased toward gene-rich regions. In gene-rich regions, many of the conserved blocks in intergenic regions correspond to regulatory elements, often located in close vicinity to genes. Given the low density of conserved blocks within intergenic regions of the XIC/Xic locus, we believe it is unlikely that many other conserved genes have escaped detection.

Conserved Retropseudogenes

We identified in human and mouse a conserved pseudogene related to the MRG15 gene (morf-related gene 15 protein; Fig. 2; Bertram et al. 1999). In both species, this pseudogene (called MRG15-psi) contains stop codons and frameshifts and lacks introns (unlike its functional counterparts), which indi-

cate that it corresponds to a retropseudogene. MRG15-psi is located at the same position (downstream of *Cdx4*) and has the same orientation in both human and mouse. In both species, the retropseudogene corresponds to the 3' end of the MRG15 mRNA (i.e., it is truncated in the 5' part of the protein-coding region, probably as a consequence of incomplete retrotranscription). It is highly unlikely that two MRG15 retroelements have inserted independently at the same location during the evolution of the primate and rodent lineages as the MRG15-psi gene is not a member of a very large family of retropseudogenes. In all we identified only nine other MRG15-related pseudogenes in the complete human genome. This suggests that MRG15-psi has been a pseudogene since its insertion at this locus prior to the divergence of primates and rodents.

We identified another retroelement conserved in mouse and human, located at the same position and in the same orientation in both species, between the Xpct- and the Bmi1like genes (Fig. 2), which is related to the DAPIT gene (accession number AJ271158; H.H. Paivarinne, unpubl.). In both species, this element is intronless, which suggests that it has been derived from the intron-containing DAPIT gene by retrotranscription. Again, this element does not belong to a large family. We found only three other DAPIT-related sequences in the complete human genome. Thus, this DAPIT-related retroelement was most probably inserted in this locus before the divergence between rodents and primates. The mouse retroelement contains two stop codons, which indicates that it has become a pseudogene. The human retroelement contains a complete ORF. However, we found no matching ESTs and our phylogenetic analysis suggests a total absence of selective pressure on the coding sequence: The ratio nonsynonymous over synonymous substitution rates in the primate DAPITrelated retroelement is 1.5, close to the ratio of 1 expected for a pseudogene.

LINE Elements

Nonrandom Distribution of LINE Elements on the Two DNA Strands in Introns and Intergenic Regions

Smit (1999) noticed that L1 elements inserted within human introns were twice as frequent on the antisense strand compared to the sense strand (relative to the orientation of the gene). He proposed that this bias was due to the presence of a transcriptional termination site within L1 elements that interferes with transcription when inserted into an intron on the sense strand. Insertion of L1 elements is counterselected because it induces premature termination of transcription. Consistent with this model, SINEs and DNA transposons that have no or only weak transcriptional termination sites do not show strand bias (Smit 1999). The analysis of introncontaining genes from the XIC/Xic region confirms this trend not only in human, but also in mouse and bovine (Table 2). Overall, in human (for which we identified seven introncontaining genes), we observed a 2.1-fold excess of LINEs on the antisense strand of introns (197 vs. 93 L1 copies), whereas in mouse (9 genes) the excess is 5.3-fold (117 vs. 22) and in bovine (2 genes) 7.8-fold (31 vs. 4). A prediction of the model proposed by Smit (1999) is that in a locus that is transcribed from both strands, all LINE elements insertions should be counterselected. The very low density of LINE elements in the 3' part of the mouse *Tsix* gene that overlaps with *Xist* is in agreement with this model (see above and Tables 1, 2). Note

that both in human and in bovine, the density of LINE elements is also relatively low in this region (Table 2).

Interestingly, all the intergenic regions, including the *Xist–Tsx* intergenic region show an excess of L1 elements on one strand compared to the other. The orientation of the bias is conserved in human and mouse, although it is not always statistically significant in both species (Table 2). If the model proposed by Smit is correct, this would suggest that most of these intergenic regions are transcribed and with the same orientation of transcription in both species. Because the pattern of conservation between the human and mouse intergenic regions does not suggest the presence of many additional unidentified genes, these intergenic regions may contain nonconventional genes like *Tsix*, which are not well conserved in sequence and do not code for a protein, or other transcribed sequences that may play a regulatory role through their transcriptional activity.

It is also possible that the nonrandom distribution of LINE elements on the two DNA strands in intergenic regions is linked to factors other than transcription (e.g., replication). We are not, however, aware of any mechanism that could bias the orientation of LINEs during their integration, although this orientation could reflect some constraint on chromatin structure.

LINE Elements Distribution in the Mouse: No Support for Lyon's Model?

Lyon (1998, 2000) has proposed that LINE elements may be responsible for spreading the inactivation signal along the X chromosome. Consistent with such a model, it has been observed in human that the density of L1 elements on the X chromosome is about twice as high as that of the autosomes (26.5% vs. 13.4%) (Smit 1999; Bailey et al. 2000). It is known that overall LINE density is negatively correlated with the G+C content of isochores in which they are located (Duret et al. 1995) and that the G+C content of the human X chromosome is lower than the average G+C content of autosomes. However, the excess of L1 elements on the X is not due to its low G+C content: The X chromosome is 1.5- to 2-fold enriched for L1 elements over autosomal regions even when isochores of comparable G+C content are compared (Smit 1999). Furthermore, Bailey and colleagues (2000) have shown that the chromosomal bands encompassing the human XIC locus (Xq13, Xq21) are even more enriched for L1 elements (45% and 39%), essentially due to an excess of young L1 elements. Our analysis of the sequence of the human XIC region has confirmed the high density of LINE elements (39%; Table 1). The bovine Xic region is also particularly rich in LINE elements (46%). However, this feature is not observed in mouse. The L1 density of the mouse Xic region (14.5%) is about three times lower than in human or bovine and is close to the overall density measured for the mouse genome (Table 1). Moreover, the L1 density of the Xic region is relatively low compared to the average L1 density of mouse genomic sequences of comparable G+C content (25% of L1 for sequences of 40%–42% G+C content; Smit 1999). Thus, in contrast with human or bovine, there is no excess of LINE elements associated with the mouse Xic region.

We have also analyzed the mouse genomic sequences available in GenBank to compare the density of transposable elements throughout the X chromosome with that of the autosomes. Although LINE elements are slightly more frequent on the X chromosome than on autosomes, the excess is limited compared to that observed in human (Table 1). Indeed,

n .		Length	Gene			3	_
Region	Species	(kb)	orientation	LINE+	LINE –	χ²	P
Xpct gene	human	147	_	47	20	10.88	<1%
	mouse	117	+	11	51	25.81	<1%
intergene	human	186	intergene	31	63	10.89	<1%
Xpct-Cnbp2	mouse	69	<u> </u>	6	11	1.47	ns
ntergene	human	26	intergene	1	4	1.80	ns
Cnbp2-Ftx	mouse	7	<u> </u>	2	0	2.00	ns
Ftx gene	human	14	+	2	6	2.00	ns
3	mouse	16	+	1	3	1.00	ns
ntergene	bovine	62	intergene	63	20	22.28	<19
Ftx-Jpx	human	425	3	70	192	56.81	<19
71	mouse	114		9	13	0.73	ns
<i>lpx</i> gene	bovine	16	_	22	4	12.46	<19
r	human	68	_	49	30	4.57	<5%
	mouse	9	_	2	0	2.00	ns
ntergene	bovine	38	intergene	5	13	3.56	ns
px-Xist	human	95	cer gene	32	58	7.51	<19
pri riise	mouse	10		0	2	2.00	ns
Xist gene	bovine	35	+	0	7	7.00	<1%
ust gene	human	32	+	0	8	8.00	<19
	mouse	23	+	ő	0	nd	\17
intergene	bovine	82	intergene	49	31	4.05	<5%
Xist-Tsx	human	38	intergene	7	3	1.60	ns
(Tsix gene)	mouse	35	_	14	3	7.12	<19
Tsx gene	human	44	(pseudo)	26	4	16.13	<19
13A gene	mouse	10	(pscudo) _	2	i	0.33	ns
ntergene	human	70	intergene	31	7	15.16	<19
Tsx-Chicl	mouse	25	intergene	1	1	0.00	ns
Chicl gene	human	157	_	61	36	6.44	<59
criici gene	mouse	32		8	5	0.69	ns
ntergene	human		intergene	nd	nd	nd	113
Chicl-Cdx4	mouse	gap 27	intergene	1	18	15.21	<19
		7		0	1	1.00	
Cdx4-gene	human	8	_	1	0	1.00	ns
intergene Cdv4 Nanlla	mouse	8 193	intercone		84	1.00	ns <1%
ntergene Cdx4-NapII2	human		intergene	47			
ntergene <i>Cdx4-Ppnx</i>	mouse	77	intergene	10	42	19.69	<1%
<i>Ppnx</i> gene	mouse	74	+	1	36	33.11	<1%

 $LINE+, number of LINEs on the direct strand; \\ LINE-, number of LINEs on the complementary strand. \\ P; probability, ns; not significant, nd; not done. \\$

none of the major classes of interspersed repeats show a strong difference in density on the mouse X chromosome (and notably the Xic region) compared to the autosomes (Table 1). Our analysis therefore provides no support for Lyon's recent model of inactivation spreading.

DISCUSSION

Our sequencing of 714 kb of the mouse and 233 kb of the bovine Xic centered on the Xist gene has provided the basis for a fully annotated comparative analysis of the complete mouse Xic with the 2.3-Mb orthologous region in human and has allowed a three-way species comparison of the core central region, including the Xist gene. Comparative sequence analysis is an efficient way to identify functional elements within genomic sequences as selectively constrained regions generally evolve slower and hence are more evolutionary conserved than functionless sequences that rapidly diverge because of genetic drift. However, this approach requires that the compared sequences are sufficiently distantly related to allow constrained from nonconstrained sequences to be distinguished. Rodents and primates diverged ~100 million years ago (Myrs) and many studies have shown the efficiency of human-mouse comparison to identify functional elements (Ansari-Lari et al. 1998; Jang et al. 1999; Jareborg et al. 1999; Mallon et al. 2000; Pennacchio and Rubin 2001). Notably, it has been established that 95% of protein-coding exons are conserved between human and mouse (Batzoglou et al. 2000). It is important to emphasize that the comparative approach we have used is also efficient in identifying untranslated genes such as Xist, Ftx, or Jpx, which are not identified by programs designed to allow ab initio identification of proteincoding gene. The power of such comparative approaches for similarly identifying functionally conserved noncoding regions, whether they be small (<10 nt), such as the binding sites for protein factors, or larger regions such as the locus control region, has been clearly established by analysis of the mammalian β-globin clusters, as well as other regions (Duret et al. 1993; Gumucio et al. 1996; Duret and Bucher 1997; Kondrashov 1999; Bouck et al. 2000; Mohrs et al. 2001). The assertion that all conserved regions are functional (Shabalina et al. 2001), however, is likely to be exaggerated. Our finding of conserved pseudogenes in the XIC/Xic locus in human and mouse indeed suggests that functionless sequences may have retained significant similarities since the divergence of primates and rodents, although it is very difficult to exclude that such pseudogenes are not acting in one way or another as controlling elements for neighboring genes (see below). In this context, functional analysis of the MRG15-psi and DAPIT-related retroelements is of obvious interest. The use of a three-way species comparison such as we have used for the core Xic region, involving species separated by ~240-300 Myrs (the evolutionary distance between rodents and artiodactyls is as great as that between rodents and primates), increases the analytical power of such a comparative approach and contributes to avoiding the identification of similarities occurring purely by chance. In this context, it is important to note that mutation rates vary along the genome (Matassi et al. 1999; Perry and Ashworth 1999), and hence some regions may show greater conservation simply because they are less subject to mutations. We chose bovine as the third species because both rodents and primates are relatively distantly related to artiodactyls (~80–100 Myrs), because the role of Xist in X inactivation in cows has been studied (De La Fuente et al. 1999), and because of the ready availability of bovine BAC genomic libraries for sequencing.

To further increase the specificity of such comparative approaches, it is necessary to consider even more distantly related species, such as birds or fish. We were constrained, however, by the functional specificity of our system since the XIC/Xic has no functional homolog in nonmammalian vertebrates. Indeed, the very mechanism of dosage compensation by *Xist*-induced X inactivation is specific to mammals, even if other species have alternative systems of dosage compensation which may present similarities at certain levels (Avner and Heard 2001; Park and Kuroda 2001)

The usefulness of adding a third species to the comparative sequence analysis is well illustrated by the short region conserved between human and mouse corresponding to the first exon of Jpx (lying 5' of Xist). In the context of the repeat elements composing the second and third exons, Jpx would probably have escaped observation if we had not also detected conservation in the bovine (Fig. 3). The conservation of exon 1 taken together with the expression of the *Ipx* gene strongly suggests a functional role for Jpx. Taken together with the nonconservation of the repeats themselves between species, this may suggest that evolutionary constraints are acting on the promoter region itself rather than on the whole gene. However, the close-on 4% frequency of transposable element (TE) integration within human genes (Nekrutenko and Li 2001) makes it unlikely that the conserved involvement of repeats, albeit of differing classes, as in exons 2 and 3 of this gene, is totally fortuitous. Indeed, it could suggest a major role for the repeat elements in *Ipx* function. Such hypotheses are reinforced both by the key role of repeats in some other forms of genes silencing and the intriguing position of Jpx immediately upstream of Xist itself.

A second example of the utility of adding in a third species comes from the analysis of the conservation pattern of *Tsix*. Lee and colleagues (1999) identified two conserved blocks between human and mouse, close to the promoter region of *Tsix*, which led them to conclude that *Tsix* was conserved in human. However, these blocks are short and are conserved neither between mouse and bovine, nor between human and bovine. Thus, despite the mutational evidence from the mouse clearly establishing a role for *Tsix* antisense transcript and the associated *DXPas34* locus in *Xist* regulation (Maxfield Boumil and Lee 2001) and in imprinted X-inactivation and -chromosome choice, it appears that the underlying sequence is not selectively constrained and is conserved in human and mouse simply by chance. Surprisingly, it turns out that the blocks of sequence conserved between hu-

man and mouse do not correspond to Tsix exons (Sado et al. 2001). Although the recent publication of Migeon and colleagues (2001) is suggestive of transcriptional activity antisense to XIST within the human XIC this activity presents few of the characteristics originally assigned to the murine Tsix antisense transcript. Our results raise specific questions as to the role of *Tsix* in species other than mouse and more generally as to the degree to which specific aspects and parameters of X inactivation may vary between mammalian species, even when the overall function such as that of the Xic or X inactivation is globally conserved (Looijenga et al. 1999). Differences between rodent and human X inactivation on one hand and marsupial X inactivation on the other include both the preferential inactivation of the paternal X chromosome and the tissue and species-specific nature of inactivation. Indeed, Xist has yet to be identified in marsupials. Differences between primates are likely less radical but several publications attest to differences in X inactivation occurring in human and murine extra-embryonic tissues and to potential differences in the regulation of the human and mouse XIST/Xist genes. It is interesting in this context to note that differences in the expression status of imprinted genes in mouse and human have been identified, and in the case of IGF2R/Igf2r this may be related to the nonexpression of the AIR antisense transcript in human (Oudejans et al. 2001).

Our work, although clearly showing the value of adding a third mammalian species (distantly related to the first two) in increasing the specificity of the comparative approach, does not clarify whether the choice of the species itself was optimal. Indeed the choice of bovine as a third species led to considerable difficulties in both the selection of bacterial artificial chromosomes (BACs) and sequencing due to the extremely high density of repetitive elements in the bovine Xic region. It would be useful to determine whether the genome of other mammalian model organisms, such as rabbit or dog, are less rich in repetitive elements, and thus better candidates for such sequencing.

Conserved elements identified include CpG islands and, more surprisingly, conserved pseudogenes. We identified 19 CpG islands (CGIs) in the mouse sequence, and 40 in the human one, that is, 2.7 and 1.7 CGIs per 100 kb, respectively, which is close to the density observed for the whole human genome (2.2 CGIs/100 kb; data from Lander et al. 2001). Five of these CGIs correspond to promoter regions of known gene (Xpct, Xist, Chic1, Cdx4, and Nap1l2), and three other to the putative promoter regions of *Jpx* and *Ftx* (NB: in mouse *Ftx* 5' region there are two CpG islands very close to each other, that correspond to a single large CpG island in human). Thus, respectively, 42% (8/19) and 18% (7/40) of mouse and human CGIs correspond to known or putative promoter regions. This latter figure is consistent with a recent statistical analysis showing that 20% (373/1846) of human CGIs overlap with transcription start sites (Ponger et al. 2001). There are only nine CGIs that are conserved between human and mouse (Fig. 2). Interestingly, the eight CGIs mentioned previously are all conserved. Thus 89% (8/9) of conserved CGIs correspond to known promoter regions. The last conserved CGI, located downstream of Xpct (close to its 3' end), might well correspond, moreover, to an unidentified promoter region. Although our gene sample is too small to be statistically reliable, these observations suggest that conserved CGIs frequently correspond to promoter regions. Comparative analysis of human and mouse CGIs, therefore, seems to be an efficient approach for identifying promoter regions.

The observation of conserved pseudogene orthologs raises the question of how such sequences can remain conserved since the time of divergence of rodents and primates and indeed what this might imply for their eventual functionality. Generally, retrotranscribed mRNAs are inactive from the moment of their insertion because they lack the promoter elements necessary for their expression. However, it can happen that a retroelement is expressed by chance after insertion downstream of an active promoter. This is unlikely to be the case for either the MRG15-psi or DAPIT-related retroelements we have studied in mouse or human as we found no matching EST. However, we obviously cannot exclude that they are expressed at a low level or in limited specific tissues or developmental stages that are not represented in the EST datasets. Under this scenario one might imagine that the DAPIT-related retroelement remained functional for a long period during primate and rodent evolution (which would explain their conservation) and only recently became a pseudogene in the rodent (and maybe also in primates, as suggested by the high ratio of nonsynonymous over synonymous substitution rates). However, in the case of MRG15-psi, it is clear that the coding region must have been truncated at the moment of its insertion. Thus a putative function (if any) cannot be related to protein-coding capacity. An RNA function or indeed a function in epigenetic control as, for example, a center of methylation, would not, however, necessarily require either conservation of protein-coding capacity nor indeed conservation of the entire genetic element. Functional tests will be necessary to further elucidate this.

Repeat elements and LINE (L1) elements, in particular, have had a major place in models seeking to explain various facets of the X-inactivation process and more recently imprinting (Greally 2002). Our computational analysis of the distribution of repeat elements in general and of LINE repeats, in particular, within the mouse Xic region, when compared to the rest of the genome did not however give support to the hypothesis of a role of these repeat elements in the spreading of X inactivation (Lyon 1998, 2000). Interestingly, though, our studies on L1 element distribution revealed that most intergenic regions within the mouse Xic, including the Xist-Tsx intergenic region, show an excess of L1 element on one strand compared to the other. If the model proposed by Smit (1999) is correct, this would suggest that most of these intergenic regions are transcribed. The orientation of strand bias is moreover conserved in human and mouse, reinforcing the notion of some underlying fundamental biological constraint. These intergenic regions could contain nonconventional genes like Tsix, Ftx, or Jpx, which are not well conserved in sequence and do not code for a protein but may play a regulatory role through their transcriptional activity. The intergenic region between Cnbp2 and Jpx contain many ESTs that might correspond to alternative transcripts of the Ftx and Jpx noncoding genes. ESTs corresponding to the other intergenic regions have yet to be found. The absence of ESTs, however, does not allow conclusions to be drawn systematically as to the absence of transcriptional activity associated with a genomic sequence, as ESTs are almost always derived from poly(A)+ cDNA libraries and generally correspond to the 3' end of transcripts. The proven transcription activity at the Tsix locus does not, for instance, match that of any known EST.

Whether or not the increasing number of noncoding RNAs encoded within the Xic, other than Xist, play regulatory roles remains to be established. Such transcripts may turn out, even in mammals, to be part of a more widespread and widely

used mechanism of epigenetic control. A region lying 50 kb upstream of the 5' end of the murine *Xist* gene is one intergenic region where atypical transcriptional activity has recently attracted our attention during studies originally undertaken to validate GENSCAN predictions unsubstantiated by FGENESH. The region turned out to be characterized by multiple (as judged by their distinct expression profiles) apparently noncoding transcripts originating alternatively from both DNA strands. Strandspecific RT-PCR has indicated that these transcripts, unlike the *Tsix* and *Xist* transcripts, are not encoded over many kilobases of contiguous genomic DNA. Intriguingly this region was recently shown to correspond to part of a constitutive hot spot of Lys 9 methyl histone H3 activity that is associated with the onset of the X-inactivation process (Heard et al. 2001).

It is interesting to note that 3 of the 11 genes characterized in the murine Xic, including the newly characterized *Ppnx* and *Cnbp2* genes, are exclusively or almost exclusively expressed in the testis. Although we have yet to confirm expression of the latter in male germ cells, the high proportion of Xic genes expressed in the testis is compatible with the X chromosome having a prominent role in male germ-cell development (Wang et al. 2001).

Our annotated sequence provides an excellent informative base for further genetic and molecular probing of the Xic. Comparison of our results for the Xic with chromosomal regions involved in other epigenetic phenomena, such as imprinting (Paulsen et al. 2001), may in turn allow other structural features to be identified and be subjected to functional analysis. Conversely, functional analysis of the Xic and further experimental validation of some of our computational observations is likely to improve both the understanding of the importance and role of some of the currently more poorly understood structural features we have identified and in the long term aid in identifying other regulatory motifs.

METHODS

Contig Construction and Sequencing

To sequence the murine Xic region, we established a contig of BAC clones from the 129/Sv mouse strain BAC library (Research Genetics, Inc.) by PCR screening using known primers. Overlapping BAC-end sequences were identified and the order verified by PCR. The entire mouse contig extending through from BAC 474E4 to 211B4 covers ~1 Mb. A minimal tiling path set of five clones collectively spanning the region were selected for genomic sequencing. Although the telomeric and centromeric ends of clones 399K20 and 334L11 do not overlap, their ends are only 1178 pb apart and lie within the previously sequenced 94-kb *Xist* region.

Bovine BACs were identified by PCR screening of a bovine library maintained at the UR339, Unité de Génétique Biochimique et Cytogénétique, Institut National de la Recherche Agronomique. Primers corresponding to the 5' region of the bovine *Xist* gene were designed to amplify a 650-bp product. End sequences of positive clones were used to rescreen the library, and three clones overlapping the 5' end of the initial clone identified. Two BACs, covering a 233-kb interval from the bovine *Xist* region were retained for sequencing. The primers used are shown in supplemental data available at www.genome.org.

Sequencing of the clones was performed by ligating mechanically sheared 3-kb fragments of the BAC DNA into the pCDNA vector (Invitrogen), followed by random shotgun sequencing to 10-fold coverage. To increase sequence contiguity and facilitate assembly additional 10-kb fragment-size subclone libraries were prepared for mouse BACs 155J2 and bo-

vine BAC 834C6. All plasmid clones were sequenced from both ends, using Licor sequencing technology. After assembly, sequence gaps and ambiguities were resolved using standard finishing techniques. Difficulties linked to the nature of the sequence meant that 436, 288, 384, 125, 198, 134, and 45 pairs of primers were used for finishing BACs 334L11, 399K20, 155J2, 561P13, 474E04, 356E2, and 834C6, respectively.

Sequence Analysis

The sequence of the human XIC region was extracted from the Human Genome Project Working Draft Assembly (October 7, 2000, freeze; http://genome.ucsc.edu/; Lander 2001). The human, mouse, and bovine sequence was analyzed for repeat sequences (LINEs, SINEs, etc.) using RepeatMasker (A.F. Smit, unpubl.; http://ftp.genome.washington.edu/cgibin/RepeatMasker) and species-specific collection of repeat sequences from Repbase-Update (Jurka 2000). Exons of previously known genes were located with SIM4 (Florea et al. 1998). After masking of repeat sequences and already known exons, new genes were searched for using three complementary approaches. First, GENSCAN (Burge and Karlin 1997) and FGENESH (http://genomic.sanger.ac.uk/gf/gf.shtml) were used to identify potential exons. Exons predicted by GENSCAN were tested experimentally by RT-PCR. Secondly, genomic sequences were compared to protein databases with BLASTX (Altschul et al. 1997). Homologous proteins detected with BLASTX were then aligned to the genomic sequences with GENEWISE (Birney et al. 1996) to identify potential genes (http://www.sanger.ac.uk/Software/Wise2/). GENEWISE gene predictions were also performed by comparison of the genomic sequences to the PFAM database of protein domains. Finally, genomic sequences were compared with BLASTN (Altschul et al. 1997) to mRNA (EST or cDNA) sequences from GenBank (release 123). Detected mRNA sequences were then aligned to the genomic sequences with SIM4 to locate intronexon junctions. Sequences with similarity to a functional genes but containing stop codons or frameshifts in the coding region were considered to be pseudogenes. Potential tRNA genes were searched for with tRNAscan-SE (Lowe and Eddy 1997).

Following Ponger et al. (2001), CpG islands are defined here as regions >500 bp with a G+C content >50% and a ratio CpGo/e (CpGo/e, number of observed CpG over number of expected CpG) of >0.6.

Pairwise local alignments of genomic sequences were performed with SIM (Huang and Miller 1991). SIM is a space-efficient implementation of the Smith and Waterman algorithm. It is much slower but also much more sensitive than heuristic methods such as BLAST or FASTA. We used SIM with the following scoring scheme: match = 1, mismatch = -1, gap opening penalty = 6, and gap extension penalty = 0.2. Only local alignments with a score >30 were retained (e.g., a score of 30 corresponds to a gap-free alignment with 100% identity >30 bp, or 80% identity >50 bp, or 65% identity >100 bp). Pairwise local alignments were visualized with LALNVIEW (Duret et al. 1996).

RT-PCR Analysis

Reverse transcription (RT) was performed on total RNA isolated with RNABle (Eurobio) and treated with RNAse-free DNAse I (Pharmacia; 10U/µg of RNA) for 30 min at 37°C to destroy genomic DNA. Genomic DNA contamination artifacts were controlled for in all RT reactions by including an RNA sample without reverse transcriptase. Random primed RT was performed on 10 µg of RNA by using SuperScriptII reverse transcriptase as recommended by the manufacturer (GIBCO-BRL) with random hexamers (Pharmacia) to prime first-strand cDNA synthesis in a 50-µL reaction volume for 1 h at 42°C. To determine the orientation of transcription, strand-specific RT was performed. For strand-specific RT, 20 µg of RNA was divided into five aliquots and reactions were per-

formed in a 50-µL volume for 1 h at 50°C with specific primer (26 pM) with or without reverse transcriptase (RTase) as follows: (1) forward primer (plus strand), RTase present, to detect transcript with a telomere-centromere orientation; (2) forward primer, RTase absent; (3) reverse primer (minus strand), RTase present, to detect transcript with an opposite orientation; (4) reverse primer, RTase absent; and (5) no-primer control, RTase present. PCR amplification was performed on 2 µL of the RT reaction products and involved 40 cycles with forward and reverse primers under standard PCR conditions. To amplify DNA fragments >2 kb, the Expand Long Template PCR system (Boehringer) was used. Fifteen microliters of the 50-µL reaction mixture was loaded onto an ethidium bromide-stained agarose gel. Initial testing of gene predictions involved testing against RNAs from undifferentiated and differentiated male and female ES cells, adult somatic cells, and testis.

Inactivation Status

Female T16/Mai F1 mice undergo nonrandom X inactivation: Genes subject to X inactivation are expressed only from the T16H allele (*Mus musculus domesticus*) and not from the Mai allele (*Mus musculus musculus*). To determine the inactivation status of the *Jpx* gene reverse-transcribed mouse brain RNAs from the T16H, Mai, and T16H/Mai mouse strains (Rougeulle and Avner 1996) were PCR amplified using the Jpx 1Up and Jpx 1Lo primers (see below for sequences). RT-PCR products were purified using the Qiaquick PCR Purification kit (Qiagen) and digested with MspA1l (Promega) to reveal a restriction site known to be polymorphic between the Mai and T16H strains. Digestion products were analyzed by loading onto a 4% ethidium bromide-stained agarose gel (NuSieve Agarose, Tebu).

Northern Analysis

Poly(A)*mRNA Northern blots from Clontech were hybridized using the Express-Hyb solution (Clontech), as recommended by the manufacturer. Hybridizations exploited ³²P random-primed double-stranded DNA probes (Megaprime DNA labeling kit, Amersham) obtained from PCR products amplified using the Jpx1Up/Jpx1Lo and Ppnx3Up/Ppnx3Lo primer pairs (Jpx1Up, 5'CGGCGTCCACATGTATACGTCC3'; Jpx1Lo, 5'TAGGAATGAGCCTCCCCAGCCT3'; Ppnx3Up, 5'AACCGTTATACCTGGACATTTC3'; Ppnx3Lo, 5'CATAACAGCTCTTGTATTGGCA 3').

Isolation and Sequencing of Ppnx cDNA Clones

Three positive clones were isolated from an adult 129/Sv mouse testis cDNA library kindly provided by Colin Bishop and 2.5 \times 10⁵ lambda gt10 clones were plated out and colony lifts made with Hybond N $^{+}$ membranes (Amersham). Screening was performed with the radioactive labeled 232-bp PCR fragment described in Northern analysis (see above) and hybridization carried out in 0.45 M sodium phosphate (pH 7.2), 1 mM EDTA, and 7% SDS at 65°C as modified from Church and Gilbert (1984), followed by washing in 2 \times SSC, 0.1% SDS at 65°C. Single-pass sequencing of the three positive cDNA clones was carried out commercially.

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