Identification of Four Highly Conserved Genes between Breakpoint Hotspots BP1 and BP2 of the Prader-Willi/Angelman Syndromes Deletion Region That Have Undergone Evolutionary Transposition Mediated by Flanking Duplicons

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Prader-Willi and Angelman syndromes (PWS and AS) typically result from an ∼**4-Mb deletion of human chromosome 15q11-q13, with clustered breakpoints (BP) at either of two proximal sites (BP1 and BP2) and one distal site (BP3).** *HERC2* **and other duplicons map to these BP regions, with the 2-Mb PWS/AS imprinted domain just distal of BP2. Previously, the presence of genes and their imprinted status have not been examined between BP1 and BP2. Here, we identify two known (***CYFIP1* **and** *GCP5***) and two novel (***NIPA1* **and** *NIPA2***) genes in this region in human and their orthologs in mouse chromosome 7C. These genes are expressed from a broad range of tissues and are nonimprinted, as they are expressed in cells derived from normal individuals, patients with PWS or AS, and the corresponding mouse models. However, replication-timing studies in the mouse reveal that they are located in a genomic domain showing asynchronous replication, a feature typically ascribed to monoallelically expressed loci. The novel genes** *NIPA1* **and** *NIPA2* **each encode putative polypeptides with nine transmembrane domains, suggesting function as receptors or as transporters. Phylogenetic analyses show that NIPA1 and NIPA2 are highly conserved in vertebrate species, with ancestral members in invertebrates and plants. Intriguingly, evolutionary studies show conservation of the four-gene cassette between BP1 and BP2 in human, including** *NIPA1/2, CYFIP1,* **and** *GCP5***, and proximity to the** *Herc2* **gene in both mouse and** *Fugu***. These observations support a model in which duplications of the** *HERC2* **gene at BP3 in primates first flanked the four-gene cassette, with subsequent transposition of these four unique genes by a** *HERC2* **duplicon-mediated process to form the BP1–BP2 region. Duplicons therefore appear to mediate genomic fluidity in both disease and evolutionary processes.**

Introduction

Prader-Willi syndrome (PWS [MIM 176270]) and Angelman syndrome (AS [MIM 105830]) are imprinted genetic disorders that arise most commonly from an ∼4-Mb deletion of human chromosome 15q11-q13 (Nicholls and Knepper 2001). Molecularly, there are two common classes of deletions in patients with PWS/AS, one from breakpoint (BP) 1 to BP3 and the other from BP2 to BP3 (Knoll et al. 1990; Amos-Landgraf et al. 1999; Christian et al. 1999) (fig. 1*a*). PWS arises from the

functional loss of a set of paternally expressed genes, whereas AS is associated with the loss of maternally expressed genes. Within a 2-Mb imprinted domain distal to BP2, five paternally expressed genes encoding polypeptides have been identified; the genes include *MKRN3* (Jong et al. 1999), *NDN* (Jay et al. 1997; MacDonald and Wevrick 1997), *MAGEL2* (Boccaccio et al. 1999; Lee et al. 2000), and the polycistronic *SNURF-SNRPN* (Gray et al. 1999). This last locus also encodes five classes of snoRNAs (Cavaillé et al. 2000; de los Santos et al. 2000; Runte et al. 2001). Adjacent to the telomeric end of the paternally expressed domain are two maternally expressed genes, with loss of function of *UBE3A* underlying the neurological phenotype of AS (Jiang et al. 1998; Nicholls and Knepper 2001) and loss of the imprinted *ATP10C* locus (Herzing et al. 2001; Meguro et al. 2001) putatively associated with obesity in AS (Dhar et al. 2000; Lossie et al. 2001; Nicholls and Knepper 2001). Genes in 15q11-q13 are homologous to mouse chromosome 7C (fig. 1*a*), with the exception

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of the paternally expressed *Frat3* gene, a recent evolutionary acquisition in the mouse (Chai et al. 2001). These imprinted loci in human and mouse are controlled in *cis* by a centrally located imprinting center (IC), a complex genetic element that acts bidirectionally over distances of at least 0.5–1.0 Mb to establish and/or maintain the maternal and paternal imprint (reviewed by Nicholls and Knepper 2001).

In addition to imprinted genes, several nonimprinted genes have been identified in the human and mouse PWS/AS region (fig. 1*a*). These genes may be responsible for other genetic disorders, and some of them may modify the PWS or AS phenotype. An example is the pinkeyed dilution (*p*) gene on mouse chromosome 7C and the corresponding oculocutaneous albinism type II (*OCA2*) locus in human. Although *OCA2* is recessive in humans and mice, hypopigmentation is a frequent finding in PWS and AS patients with the common deletion (Spritz et al. 1997; Nicholls and Knepper 2001). Other recessive disease loci identified within the mouse region homologous to PWS/AS are neonatally lethal cleft palate (*cp1*) due to loss of *Gabrb3* (Culiat et al. 1995; Homanics et al. 1997; Hagiwara et al. 2003) and juvenile development and fertility (*jdf2*) associated with *Herc2* gene mutations (Lehman et al. 1998; Ji et al. 1999; Walkowicz et al. 1999).

Nevertheless, the complete extent of the imprinted domain is not known in human 15q11-q13 or mouse 7C. The telomeric boundary of the 15q11-q13 imprinted domain—and corresponding region in the mouse—may be between *ATP10C* and *GABRB3,* since genetic evidence in both species indicates that the latter is not imprinted (Nicholls et al. 1993; Culiat et al. 1995; Homanics et al. 1997; Nicholls 1999; Nicholls and Knepper 2001). At the other end of the imprinted domain in humans, it has not been clear whether functional genes occur centromeric of *MKRN3*, but Ritchie et al. (1998) proposed that the imprinted domain might extend significantly farther by identifying asynchronous DNA replication at a partial duplication of the *GA-BRA5* locus, which maps proximal to deletion breakpoints BP1 and BP2 in the pericentromeric region of 15q11.1. Replication asynchrony is a feature of imprinted domains (Kitsberg et al. 1993; Knoll et al. 1994; Simon et al. 1999) and other domains with monoallelic gene expression (Chess et al. 1994; Hollander et al. 1998; Mostoslavsky et al. 2001). To define the extent of this imprinted domain, we searched for novel genes centromeric of *MKRN3* and describe here four unique genes in 15q11.2, the imprinting status of which was tested in human and mouse. These genes include the novel *NIPA1* (nonimprinted in Prader-Willi/Angelman syndrome 1) and *NIPA2* genes (symbols approved by the gene nomenclature committee), which encode highly conserved polypeptides with possible function as recep-

tors or transporters. Both genes—as well as the flanking *CYFIP1* (Kobayashi et al. 1998; Schenck et al. 2001) and *GCP5* (Murphy et al. 2001) genes—are candidate modulators of the PWS/AS phenotypes and/or may play a role in other disorders mapping to this region in humans and mice. Furthermore, our phylogenetic studies have identified an evolutionary transposition of these four genes from the BP3 region of ancestral vertebrates to form the human BP1–BP2 region. We propose that this transposition was mediated by the flanking *HERC2* and/or other duplicated sequences.

Material and Methods

Isolation of BAC Clones and STS Mapping of BAC and YAC Clones

The *CYFIP1* (*SRA1*) gene was identified by an *in silico* search of National Center for Biotechnology Information (NCBI) GeneMap '98. Of all EST clusters found in proximal 15q, the cluster containing the 4.4-kb cDNA KIAA0068 was the most robust, with numerous ESTs in different tissues, and was subsequently identified as *SRA1*/*CYFIP1* (Kobayashi et al. 1998; Schenck et al. 2001). GeneMap '98 presented compiled mapping data from two radiation hybrid panels, GB4 and G3; however, because the relative order of markers was not consistent between these panels, genes were further mapped, by PCR, to BAC and YAC contigs (see below). CITD BAC 3242E18 was identified by a BLAST search of the GenBank Genome Survey Sequence (GSS) database using *CYFIP1* cDNA sequence (GenBank accession number D38549). Exons 7 and 8 are present in the 3242E18 end sequence (AQ203319) in a $5' \rightarrow 3'$ orientation compared with the end sequence. Similarly, partially sequenced BAC clones RP11-26F2 (GenBank accession number AC011767) and RP11-289D12 (GenBank accession number AC090764) were identified by BLAST with *CYFIP1* cDNA sequence and STS markers (Christian et al. 1999) *D15S18,* located in *CYFIP1* intron 1 close to exon 2, and microsatellite markers *D15S1035, D15S541,* and *D15S542* that are clustered within *CY-FIP1* intron 1 ∼15 kb from exon 1.

For murine *Cyfip1,* which was originally identified as *Shyc* (Köster et al. 1998), primers RN1102 and RN1103 were used to screen a Genome Systems mouse BAC library (129/Sv) by PCR, using 35 cycles at 94 \degree C for 2 min, 55°C for 30 s, and 72°C for 1 min, with a 10-min final extension at 72°C. Mouse RPCI-22 BAC 252P22, which was identified by the PCR screen, was end sequenced, allowing a PCR product (amplified by RN1235 and RN1236) from one end of this BAC to be used to further screen the RPCI-22 BAC library by hybridization. This approach identified BACs 508F17, 402D17, 373J21, and 526E9, and BACs 466F3 and 304H15 were

Figure 1 Genetic and physical maps. *a*, Schematic maps of human chromosome 15q11-q13 and the homologous mouse chromosome 7C region. PWS/AS deletion BP hotspots 1–3 associated with *HERC2* duplicons (dup) are shown, and the regions studied in the present work are boxed and shaded. Two BAC clones spanning the four novel BP1–BP2 genes are indicated. Black and dark gray circles represent paternally and maternally expressed imprinted genes, respectively, and white circles are biallelic, nonimprinted genes (the mouse *Atp10c* gene is likely, but not proven, to be maternally expressed [*light gray circle*]). Genes or regions associated with PWS, AS, oculocutaneous albinism type II (*OCA2*), and a transgene insertion-deletion mouse model of PWS/AS [TgPWS/AS(del)] are indicated. *b,* PCR mapping of *NIPA1* and *GCP5* STS markers within YACs spanning the BP1–BP2 region. *c,* PCR detection of a *CYFIP1* STS marker in BP1–BP2 YACs. *d,* Orientation of BP1–BP2 genes, with detailed map of YACs and STS markers spanning BP1–BP2. Markers from this study are shown by large black circles, and data for the *HERC2*-dup CpG island probe (*black squares, Hin*dIII fragment sizes in kb also shown) and 254RL2 probe (*gray squares*) are from Amos-Landgraf et al. (1999). The left (L) and right (R) YAC ends (Ye), and deleted fragment (?) are also shown. *e,* Schematic of class I and II PWS and AS deletions, with detection by FISH using unique BAC clones spanning *CYFIP1*. *f,* Fine mapping and BAC contig of six nonimprinted genes in mouse 7C. Each gene is shown proportionally by the length of black arrows, with direction of transcription from 5' to 3'. The map was determined by STS analysis derived from exons (e) and BAC ends in a series of BACs and mouse 10-kb clones (*thin black bars under genes*). Black and white squares represent STSs confirmed by PCR assay or identified by database DNA sequence analysis without further PCR assay, respectively.

identified with a PCR probe from the 195C6 T7 BAC end (table 1). BLAST searches of GSS, with the use of cDNA sequences of *Cyfip1, Nipa1, Gcp5, Herc2,* and *p*, were conducted, identifying BACs RPCI-23 (C57BL/ 6) 88I18, 452K16, 363F10, 471F18, 335K21, 49E2,

195C6, 396N13, 256L9, and 181P23 and RPCI-24 247O4, 150F2, 354P8, and 265E12. DNA from all BACs was isolated according to the minipreparation protocol recommended by Research Genetics. To establish BAC contigs and to determine gene order for the

a

mouse loci, several pairs of primers were designed from gene exons or BAC ends (table 1). PCR was conducted under conditions of 94°C for 2 min, 31 cycles of 94°C for 30 s, 60°C for 30 s, and 72°C for 30 s and a 10min final extension at 72°C.

DNA from YAC clones spanning the BP1–BP2 region was isolated as described elsewhere (Amos-Landgraf et al. 1999). STS primers from single exons of the *NIPA1, GCP5,* and *CYFIP1* genes (table 1) were used for PCR typing of these YACs, using 10 ng DNA for each PCR. PCR used the following cycling conditions: 94°C for 2

min, then 35 cycles of 94°C for 30 s, 55°C for 30 s, 72°C for 30 s, and a final extension for 10 min at 72°C.

FISH

For human studies, metaphase chromosome preparations were made from lymphoblastoid cell lines derived from class I and class II AS deletion patients (I: WJK29, WJK43; II: WJK18, WJK35, WJK48; Knoll et al. 1990), as well as normal individuals, according to standard protocols. BAC 3242E18 was used as the probe

PCR Primers (5^{'-3'})

for the human chromosome FISH studies. BAC DNA was labeled with biotin or digoxigenin, by nick translation. Chromosomal hybridizations were detected as described elsewhere (Knoll and Lichter 1994; Amos-Landgraf et al. 1999). Ten to twenty metaphase cells were analyzed per cell line. To identify the chromosomal location of murine *Cyfip1,* BAC 252P22 was digoxigenin labeled and hybridized to murine metaphase chromosomes derived from Tg^{PWS(del)} splenocytes, along with a mouse chromosome 7 centromere probe (kindly provided by A. G. Matera), which also hybridizes to the telomere of chromosome 5, labeled with biotin, as described elsewhere (Gabriel et al. 1999). Probes were detected by fluoroscein-labeled antidigoxigenin and Texas red avidin, with DAPI (4',6-diamino-2-phenylindole) counterstaining.

DNA-replication–timing studies were performed as described elsewhere (Greally et al. 1998). A mouse splenocyte culture was pulsed with bromodeoxyuridine (BrdU, 100 μ M) 90 min prior to harvesting. The cells were then harvested, fixed with methanol–acetic acid and spread onto slides (Henegariu et al. 2001). The BAC probes (452K16 and 252P22) were labeled by nick translation using digoxigenin, Texas red, and biotin.

FISH was performed by denaturing the slides, using 0.05N NaOH at room temperature for 4 min, followed by suppressive hybridization using probes preannealed with mouse Cot-1 DNA. The haptens were detected using antidigoxigenin-rhodamine and avidin-CY5, while simultaneously detecting S-phase cells with anti-BrdU-FITC (Pharmingen), followed by counterstaining with DAPI. Multiple image planes were acquired for each cell, to ensure that all signals were identified. Replication patterns in S-phase cells were assigned as prereplicative if a single signal was seen at a locus and were assigned as postreplicative if more than one signal was evident. The replication patterns for 50 nuclei were analyzed for each probe. For the parent of origin-specific patterns at the *Nipa1*-*Nipa2*-*Cyfip1* locus, simultaneous probing with RPCI-22 BAC 434N7 was performed with a different fluorophore on *c32DSD* mice with a *Tyr* deletion on their paternal chromosome (Rinchik et al. 1993). The 434N7 signal from the maternal chromosome colocalized with one of the *Cyfip1* homologues in S-phase cells, allowing the confident assignment of parental origin in $>90\%$ of cells. The few cells in which the *Tyr*signal was not clearly associated with one chromosome were not included in our analyses.

Identification of NIPA1 *and* NIPA2 *cDNAs*

BLAST search of the GenBank GSS database, using *Cyfip1* cDNA sequence (GenBank accession number AF072697), matched sequence at one end of a mouse 10 kb plasmid 1M0204M02F (GenBank accession number AZ424789). The other end of this plasmid had a segment 99% identical to the *Mus musculus* hypothetical protein MNCb-2146 (renamed *Nipa2*) mRNA (GenBank accession number NM_019997; 1,867-nt sequence), 85% to several human cDNA clones (GenBank accession numbers HSU90904, BC011775, and BC000957), and 83% to a *Gallus gallus* cDNA (GenBank accession number AL588904) in the nonredundant GenBank database. A full-length human *NIPA2* cDNA (2,421 bp) spanning exons 1–7 was then assembled *in silico* on the basis of these cDNAs and partial genomic sequences from BACs 26F2 and 289D12 (see above). The cDNA sequences included one clone with exons 1 and 3–7 (2247-bp, BC011775), and one 5' clone with exons 1, 2, 2b, and 3 (GenBank accession number BF203654). Exon 2b is a rare alternatively spliced exon in the 5' UTR of human *NIPA2*, but it is not conserved and is likely nonfunctional. Subsequent BLAST searches confirmed numerous mouse and human ESTs with the predominant structure including sequence spanning seven exons. *NIPA2* is oriented tail to tail with *CYFIP1*, with 1,747 nt between the two genes. A *Gallus gallus Nipa2* cDNA clone (GenBank AL588904) from a chicken brain library was obtained from the Roslin Institute, United Kingdom, and was sequenced completely through the DNA Sequence Facility at the University of Pennsylvania School of Medicine. The 5' end was from a chicken EST (GenBank accession number AJ452290). BLAST searches using human and mouse Nipa2 coding sequence in the *Fugu* Project Database identified five exons spanning the conserved ORF sequence of *Fugu Nipa2* (FT:T000059 Scaffold 59; these data were generously provided by the *Fugu* Genome Consortium for use in this publication only).

Human *NIPA1* was identified by *in silico* methods, beginning with identification, within partial BAC 26F2 genome sequence, of a second strong CpG island not associated with the 5' end of *NIPA2*. This unique sequence matched with three human ESTs (GenBank accession numbers AW962805, AA355571, and D81972), which identify an exon-intron structure by comparison with BAC 26F2 sequence, suggesting the presence of a unique gene that further analyses identified as *NIPA1.* The AW962805 EST sequence covers part of exons 1– 4 of *NIPA1,* as well as an exon termed 3b. Exon 3b is alternatively spliced (see below), but it has an in-frame stop codon, is not present in other EST clones, occurs at lower levels in RT-PCR products than does product with exons 2–4 (see below), and is not conserved in the mouse. These facts led us to conclude that exon 3b is

nonfunctional. BLAST search of the GenBank EST database also identified an initial fourth human EST (Gen-Bank accession number BE003132), which spans part of exons 4 and 5 and the 3- UTR of *NIPA1*, as well as three putatively orthologous mouse ESTs (GenBank accession numbers BF118592, BE654018, and BF539166). Genome sequence within BAC 26F2 that lay 3' of the NIPA1 ORF was then used, combined with BLAST searches of the EST database, to identify numerous cDNA clones containing the two 3' ends of *NIPA1* predicted on the basis of the mRNA size from northern blots (2.2 and 7.5 kb) and the presence of a polyadenylation signal and polyA tract at one end of the EST sequences.

The mouse BF539166 cDNA clone was purchased from Research Genetics and was sequenced to completion; it represents a full-length 1.9-kb mouse *Nipa1* cDNA. Several mouse *Nipa1* gene exon-intron boundary sequences were initially predicted on the basis of sequences identified by BLAST search of the mmtrace database and were subsequently confirmed by direct sequencing (data not shown) and by *in silico* analyses of BAC and shotgun sequence from the public genome sequence, as well as Celera databases. BLAST search of the *Fugu* Project Database with mouse NIPA1 amino acid sequence identified all five exons for the ORF sequence of the orthologous *Fugu Nipa1* (as above for *Nipa2*).

Analyses of NIPA1 *and* NIPA2 *Polypeptide Sequences*

In addition to the *NIPA1* and *NIPA2* genes identified as above for human, mouse, *Fugu*, and chicken (*NIPA2* only), using translated BLAST searches of cDNA and genome project databases, we identified an *NIPA2* ortholog in *Xenopus* (encoding a putative polypeptide of 362 amino acids), as well as a single ancestral gene in *Drosophila* (385 amino acids), *Anopheles gambiae* (351 amino acids), and *Caenorhabditis elegans* (358 amino acids), and a representative paralog in *Arabidopsis* (343 amino acids). CLUSTAL W (Thompson et al. 1994) was used to align NIPA1, NIPA2, and ancestral gene amino acid sequences, with comparison of NIPA1/2 orthologs and paralogs by an identity/similarity matrix program, while phylogenetic analyses used drawtree to plot an unrooted tree diagram. Putative transmembrane helices were predicted based on hydrophobicity plots using the TMHMM (v. 2.0) program. It may be noted that a WW protein-protein interaction domain was predicted to occur in mouse and human NIPA1 proteins; however, we assume this was a chance similarity, since it spans a strong transmembrane (TM) helix (H) domain and is not conserved.

Determination of Gene Expression by RT-PCR and Northern Blot Analyses

Total RNA samples were extracted using TRIzol (Invitrogen), according to the manufacturer's protocol, from lymphoblast cell lines derived from normal individuals, unaffected and affected patients with an imprinting defect from the PWS-U (Buiting et al. 1995) and AS-J families (Saitoh et al. 1996), and rodent-human somatic cell hybrids with a single maternal (20L-28) or paternal (A59-3az) human chromosome 15 (Gabriel et al. 1998). For the hybrids analysis, PCR primers were human specific. In addition, RNA was extracted from homogenized mouse brain from three genotypes of mice: wild-type, transgenic PWS, and AS (TgPWS(del) and Tg^{AS(del)}), the last two of which have an ~4-Mb deletion of mouse chromosome 7C (Gabriel et al. 1999). From each sample, $1-10 \mu$ g of total RNA was pretreated with DNase I (GIBCO-BRL), with half of this used to synthesize first-strand cDNA, using SuperscriptII reverse transcriptase (GIBCO BRL) and an oligo dT primer plus random hexamers $(RT+)$, and the other half was reverse transcriptase free (RT-). One twenty-fifth of each reaction was used subsequently for PCR amplification, with 10 mM Tris-HCl (pH 8.4), 50 mM KCl, 2 mM MgCl2, 0.2 mM dNTPs, 2 U *Taq* DNA polymerase (GIBCO BRL), and 200 nM each of primers, in a $25-\mu$ l reaction volume. To assess imprinting, we utilized 35 cycles of 94°C for 2 min, 55°C–60°C for 30 s, and 72°C for 1 min, with a 10-min final extension at 72°C. Primers used for each locus (table 1) were as follows: human and mouse *NIPA1* exons 2–4; human and mouse *NIPA2* exons 3–6; human and mouse *GCP5* exons 10 and 11; human *CYFIP1 3'* UTR, as well as exons 17–25; mouse *Cyfip1* exons 4–7; human *SNURF-SNRPN* exons 2 and 3; mouse *Snurf-Snrpn* exons 3–9; and mouse *Mkrn1*.

RT-PCR products obtained from human and mouse *NIPA1* (167 bp), *NIPA2* (339 bp), and *GCP5* (399 bp), human *CYFIP1* 3' UTR (505 bp), and mouse *Cyfip1* 3' UTR (258 bp, RN1102, and RN1103), as described above, were subcloned into a TA vector and used for preparation of probes. The probe for *SNURF-SNRPN* was similarly prepared from TA-cloned PCR product of exons 2 and 3 by amplification with RN420 and RN423. Probes were labeled with α -³²P-dCTP and were hybridized to human multiple tissue and brain multiple tissue northern blots (Clontech), human 12-lane MTN blot (BD Biosciences), or mouse multiple tissue northern blot-2 (Sigma), according to the manufacturer's protocol. b*-ACTIN* or *SNURF-SNRPN* and *Gapdh* were used as controls for human and mouse northern blots, respectively. Hybridizations were performed using ExpressHyb (Clontech) at 65°C and were washed at high stringency in $0.1 \times$ SSC + 0.1% SDS at 60°C-65°C. Membranes were exposed to autoradiographic film at -80° C.

Results

Identification and Fine Mapping of Novel Genes in the BP1–BP2 Region of Human 15q11.2

To identify novel genes in the BP1–BP2 region, we first used BLAST searches with the chromosome 15q11.2 STS marker D15S18 and microsatellite markers D15S1035, D15S541, and D15S542 (Christian et al. 1999) to identify partially sequenced (subsequently completed) BACs RP11-26F2 and RP11-289D12, which span this region (fig. 1*a*). From GeneMap '98, we identified the KIAA0068 EST, subsequently recognized as *SRA1*/ *CYFIP1* (Kobayashi et al. 1998; Schenck et al. 2001), with exons in each of the two BP1–BP2 region BACs, a location confirmed by PCR (see below). Subsequently, we identified a second gene, *GCP5* (Murphy et al. 2001), adjacent to *CYFIP1,* by BLAST searches with genomic sequence from BAC 289D12. Gene structures for these two genes have not been previously determined. The human *CYFIP1* gene spans 111.4 kb and has 31 exons (table 2) overlapping BACs 26F2 (exons 23–31) and 289D12 (exons 1–24) (fig. 1*a*), with short 5' and 3' UTRs of 52 nt and 562 nt, respectively. Exon 1 is noncoding, with the AUG start codon present at the $5'$ end of exon 2. Mouse *Cyfip1* has a similar 31-exon gene structure. Human *GCP5* (also known as *TUBGCP5*) spans 46.8 kb and 23 exons with a 54-nt 5' UTR and a 643-nt 3' UTR (table 3), with a similar structure in the mouse (data not shown). Although 45% of the human *GCP5* 3' UTR and 17% of the mouse 583-nt 3' UTR are an *Alu* and a truncated B2 repetitive element, respectively, these short interspersed nuclear elements (SINEs) have different evolutionary origins and are inserted at different positions and orientations, which suggests coincidental insertion and a lack of selection on the 3' UTR in each species. The *CYFIP1* and *GCP5* genes are arranged in a head-to-tail orientation in human and mouse (fig. $1a$), with the $5'$ ends of both genes associated with strong CpG islands. In addition, two novel CpG island sequences not associated with either known gene were identified from genomic sequence of BAC RP11-26F2, and these are characterized in detail below as being located at the 5' ends of the *NIPA1* and *NIPA2* genes.

Because of the repeated nature of *HERC2*- and other duplicons in BP1 and BP2 and because of poor BAC and DNA sequence coverage in these regions, such analyses could not determine the orientation of BACs 26F2 and 289D12 or of the four BP1–BP2 region genes with respect to the centromere. Using primers derived from the 5' end of *NIPA1* (fig. 1*b*), both 5' and 3' ends of *GCP5* (fig. 1*b*) and the 3' UTR (exon 31) of the *CYFIP1* gene (fig. 1*c*), we performed PCR on a panel of YACs known to span the BP1–BP2 region (fig. 1*d*). This YAC panel was characterized previously by STS content and

Table 2

Exon-Intron Structure of the Human *CYFIP1* **Gene**

^a Lowercase letters refer to intron sequences, and uppercase letters refer to exon sequences. Underlined letters "ag" and "gt" represent consensus splice acceptor and donor sequences, respectively. The underlined "AGT" represents the sequence coding for the translation initiation codon in the corresponding mRNA.

 b NA = not applicable.

HERC2-duplicon mapping (Amos-Landgraf et al. 1999). Both *NIPA1* exon 1 and the 3' UTR of *CYFIP1* were detected on YACs 931C4 and A124A3 (fig. 1*b, c,* and *d*), confirming that these genes map proximal to BP2 of *MKRN3* and were therefore proximal to the known PWS imprinted domain. Furthermore, both ends of *GCP5* are within YAC 931C4 but not A124A3 or any BP2 region YACs (fig. 1*b* and 1*d*), unambiguously defining the order as cen–*NIPA1*–*CYFIP1*–3- -*GCP5*-5- – BP2–tel.

FISH Mapping of BP1- and BP2-Associated Rearrangements

BAC clone 3242E18, containing exons 7–31 of *CY-FIP1* from the BP1–BP2 region (fig. 1*e*), was utilized for FISH studies on metaphase chromosomes from three normal human lymphoblastoid cell lines, with hybridization to a single region of proximal chromosome 15 in band 15q11.2 (fig. 2*a*). No other chromosomal hybridizations were detected, including in interphase cells, confirming that BAC 3242E18 does not contain duplicon sequences. Therefore, we investigated the pattern of hybridization of BAC 3242E18 against a panel of cell lines from patients with AS deletions. Deletions in patients with AS and PWS (fig. 1*e*) are either class I, which extends from BP1 $(15q11.2)$ to BP3 $(15q13)$, or class II, which extends from BP2 (15q11.2) to BP3 (Knoll et al. 1990; Amos-Landgraf et al. 1999). As expected, the 3242E18 FISH assay distinguishes class I and class II deletions in patients with AS, because the BAC is present on both chromosomes 15 in cells of three class II patients and is present only on the normal chromosome 15 in cells from two class I patients (fig. 1*e;* data not shown).

Physical Map Spanning Six Genes at the Centromeric End of Mouse Chromosome 7C

The human *CYFIP1* gene is orthologous to the mouse *Shyc* (*Cyfip1*) gene (Köster et al. 1998; Schenck et al. **Table 3**

Exon	cDNA Position (bp)	Splice Acceptor Intron Exon ^a	Exon Size (bp)	Splice Donor Exon Intron ^a	Intron Size (bp)	
1	$1 - 200$	5'UTR (1-54)	200	CCAACTTCAGgtgtggggcacg	2,244	
2	$201 - 254$	tgtctttaatagATTTCATCGT	54	CAATCGAAGGgtatgccttcaa	92	
3	$255 - 363$	cttatattttagAATTTATGAA	109	GGAAATAAAGgtatgccttcaa	4,075	
4	364-460	tttgttttaaagACAGATGCAC	97	AAAGAAGTGGgtatgttactag	696	
5	$461 - 540$	gttacattttagAAAAGAAAGA	80	GGACACACCAgtaagtggtagt	928	
6	541-675	tttttaaatcagAATTGGTCTG	135	AGTTGGAAAGgtattatgtgtt	3,582	
7	667-791	cagatgagccagATGAGCCAGA	125	CTGCTGTCTGgtaagaagctaa	985	
8	792-881	gctgtctcatagGGACCAACAC	90	AAACCCTATGgtaagaaatgtt	1,284	
9	882-975	atctcatttcagGTTACTTTCA	94	TTTAACACATgtaagttatttg	542	
10	$976 - 1,222$	cttgtgttgtagAGCTGTTTAC	247	ATCAATAATGgtattaaatgtt	1,784	
11	1,223-1,425	tgatatttgcagATACTACAAT	203	TGAGCAAACCgtaagtgagact	2,623	
12	1,426-1,539	tctctgcctcagGTCTCCCTCC	114	TCATCCAGAGgtgagctgcagc	7,177	
13	1,540-1,809	gaatctctttagAAACAAAAAT	270	GGAGCCAGAGgtacttgttccg	6,824	
14	1,810-2,009	tttatgtttcagATGCAGAAAG	200	ACTTTGCAAGgtgagacacacc	998	
15	$2,010 - 2,198$	ttttatcttgagGATGTATTTG	189	AAGATTACAGgtaaaagagtaa	1,063	
16	2,199-2,381	ctctccattcagGTTGGTAGAA	183	ATAGTTCACAgtaaaatatgga	2,348	
17	2,382-2,466	ttctttctttagTCTATCTATA	85	CAGCTACAAGgtatatgctgat	94	
18	$2,467 - 2,587$	tttgtttcatagGTCCCATGGC	121	CTTTTTGGTGgtaagatttgtt	440	
19	2,588-2,766	aatgttttatagAACTGGTTAG	179	CATGACCAGGgtttgtgcctct	1,203	
20	2,767-2,892	ttctcttccaagATTCTACACA	126	GAGAGAAAAGgtacatgagatg	947	
21	2,893-2,981	tttggtctgcagGTCAGCTTTG	89	GCACTTGGCGgtaagtatgtga	2,394	
22	2,982-3,082	ttttattcctagAATGGAATCT	101	TTTCCCCATTgtgagtatatca	701	
23	3,083-3,771	ttcttcccacagTGGAATCTCT	689	3' UTR (3127-3771)	NA^b	

Exon-Intron Structure of the Human *GCP5* **Gene**

^a Lowercase letters refer to intron sequences, and uppercase letters refer to exon sequences. Underlined letters "ag" and "gt" represent consensus splice acceptor and donor sequences, respectively.

 b NA = not applicable.

2001). We isolated the *Cyfip1* 129/Sv RPCI-22 252P22 BAC clone (fig. 1*f*) by PCR screening with a 3' UTR STS. FISH was then performed on metaphase chromosomes from a mouse model of PWS with a chromosome 7C deletion (Gabriel et al. 1999). BAC 252P22 (*Cyfip1*) localizes to the normal chromosome 7C region and within the mouse transgene-induced deletion (fig. 2*b*), which expands the known syntenic relationship between human 15q11-q13 and mouse 7C (fig. 1*a*).

To generate a physical map, seven unsequenced 129/ Sv and 14 partially sequenced C57BL/6 BACs were identified (see "Material and Methods" section) and typed by PCR with STSs derived from the *p, Herc2, Nipa1, Nipa2, Cyfip1,* and *Gcp5* genes and with BAC end sequences created in this study, allowing generation of an overlapping BAC contig map (fig. 1*f*). These data indicate that the genes in this region are arranged in the order cen–*Gcp5*–*Cyfip1*–*Nipa2*–*Nipa1*–*Herc2*-*p*–tel (fig. 1*f*). The map is anchored at one end by an STS marker, D7Mit70, which maps close to the *l71Rl* locus involved in peri-implantation survival (Wu et al. 2000), that is ∼22 kb centromeric to *Gcp5*.

Replication Analyses of the Nipa1*-*Nipa2*-*Cyfip1 *Locus*

Replication-timing studies of imprinted regions have demonstrated a parent-of-origin–dependent asynchrony,

using FISH and quantitative PCR techniques (Kitsberg et al. 1993; Knoll et al. 1994; Simon et al. 1999). For two BACs of different sizes that collectively span the *Nipa1, Nipa2,* and *Cyfip1* genes, we see a pattern of asynchrony that is similar to that at the imprinted *Mkrn3* and *Snurf-Snrpn* loci (fig. 2*c*). To test whether the pattern at *Nipa1*-*Nipa2*-*Cyfip1* was determined by parent of origin, we used mice heterozygous for a deletion at the mouse *Tyr* (*c*) locus (c^{32DSD}) (Rinchik et al. 1993) to examine which chromosome was replicating earlier in each cell. The relatively close linkage of *Tyr* to the imprinted 7C region (∼14 cM) makes this a useful marker for in situ studies of this region while being sufficiently distant that the likelihood that *cis* effects of the deletion cause disruption of imprinting is very small. We found a pattern of replication in these cells that was asynchronous but was not due to parent-of-origin influences. In other words, the proportion of cells that showed the paternal chromosome replicating early was equal to the proportion that showed the maternal chromosome replicating early (fig. 2*c*).

Characterization of the NIPA1 *Gene in Vertebrates*

Human *NIPA1* maps within BAC 26F2 and was identified, on the basis of a strong CpG island, by visual inspection of genomic DNA sequence, followed by

 C

Figure 2 FISH mapping of *CYFIP1* in human and mouse, as well as DNA replication-timing asynchrony at the mouse *Nipa1-Nipa2- Cyfip1* locus. *a,* BAC 3242E18 spanning *CYFIP1* (fig. 1*e*) is unique and maps to human chromosome 15q11.2 (*arrows*). *b,* BAC 252P22 (fig. 1*f*) maps to mouse chromosome 7C (*green signal, yellow arrow*) in metaphase chromosomes from Tg^{PWS(del)} mice. A control probe identifies the chromosome 7 centromere (*red signal, white arrow*) but also hybridizes to the telomere of chromosome 5 (*white arrowhead*). *c,* Probes from imprinted (BACs RP-23 371M8 and 266F22 for *Snurf-Snrpn* and *Mkrn3*, respectively) and nonimprinted (BACs 452K16 and 252P22 for *Nipa1*- *Nipa2-Cyfip1*; see fig. 1*f* and fig. 5) loci exhibit asynchronous replication (a high singlet/doublet [1/2] proportion of hybridization foci in Sphase cells, comparable to prior studies [Kitsberg et al. 1993; Greally et al. 1998; Simon et al. 1999]).

assembly of a cDNA sequence contig, using BLAST searches extending 3' from the CpG island (see "Material and Methods" section for details). This led to an *NIPA1* cDNA sequence of 2,247-bp that spanned five exons and 38.8 kb in genomic DNA (fig. 3*a*) and encoded a putative polypeptide of 329 amino acids (fig. 4*a;* see below). The five *NIPA1* exon-intron boundaries conform to consensus splice acceptor and donor sequences (table 4), with the translational AUG start codon present in exon 1, the stop (TGA) codon in exon 5, and in the 3' UTR an ATTAAA variant polyadenylation signal at cDNA position 2221–2226 (fig. 3*a;* table 4). Although single base changes of the hexanucleotide (AATAAA) poly(A) signal sequence greatly reduce polyadenylation, the $A\rightarrow T$ substitution at position $+2$ of the noncanonical ATTAAA hexamer is the mildest mutation and corresponds to the most common variant (Sheets et al. 1990). Nevertheless, multiple *NIPA1* ESTs have polyA tails beginning 13–17 nt 3' of this polyA signal, suggesting this as a bona fide 3' end of low-abundance *NIPA1* transcripts, which was supported by northern analysis of gene expression, although these studies also identify a larger neuronal transcript (see below). Therefore, we examined genomic sequence further 3' both visually and by BLAST against the EST database, which allowed identification of the major human *NIPA1 3'* UTR polyadenylation signal (AATAAA) at cDNA positions 6540–6545. The latter is embedded in a region that is highly conserved in five eutherian mammals and that is diagnostic for the *NIPA1* 3' end (fig. 3*b*).

Within the GC-rich (75%) exon 1 of *NIPA1,* we identified a (GCG) _n repeat, encoding polyalanine (fig. 4*a*), which we examined for polymorphism by one or both of following methods: direct sequencing of exon 1 PCR products or detection of PCR products by use of a ^{32}P dCTP labeled primer. This analysis showed polymorphism in 135 chromosomes of European and Asian origin with the number of repeats ranging from 6 to 10 (data not shown). The two most frequent alleles, $(GCG)_8$ and $(GCG)_{7}$, have allele frequencies of 0.78 and 0.2, respectively. Additional studies are needed to determine whether expansions occur in the *NIPA1* (GCG)_n repeat and are disease associated.

The mouse *Nipa1* cDNA was identified by *in silico* methods, using the human coding sequence as well as sequencing of a 1.9-kb cDNA clone. By comparison with mouse genome sequence, the *Nipa1* locus was found to span 41.1 kb of genomic DNA and to include five exons that are highly conserved with the human *NIPA1* gene (fig. 3*a;* table 4). Mouse *Nipa1* encodes a putative polypeptide of 323 amino acids (fig. 4*a*), with an N-terminal polyalanine sequence encoded by a (GCG) ₃ $GCT(GCG)$ ₃ sequence in exon 1. In a manner similar to that in humans, two polyadenylation signals occur in the mouse *Nipa1* 3' UTR at cDNA positions 1866 nt and 6840 nt

of the cDNA (fig. 3*a*), both of which are AATAAA signals and correspond to the two *Nipa1* transcripts (see below). We also identified *in silico* the complete orthologous *Nipa1* gene from the *Fugu* Project Database by translated BLAST analyses, using human and mouse amino acid sequences. This allowed us to predict a virtual cDNA sequence of 932 bp, with five exons of similar sizes and conserved consensus splice acceptor and donor sequences, as for the human and mouse orthologous genes (fig. 3*a;* table 4), and encoding a putative polypeptide of 304 amino acids (fig. 4*a*). As expected, the genomic locus of *Fugu Nipa1* is much smaller than that of mammalian *NIPA1* loci and spans only ∼1.4 kb, not including the UTRs.

Characterization of the NIPA2 *Gene*

As with *NIPA1,* the human *NIPA2* gene was identified by *in silico* methods using BLAST of BAC 26F2 genomic sequence, as well as identification of a second CpG island in this BAC that represents the 5' end of *NIPA2*. We compiled a synthetic *NIPA2* cDNA sequence of 2,450 bp, with an AATAAA poly(A) signal at positions 2444– 2449 utilized for polyadenylation (fig. 3*c*). *NIPA2* spans 29 kb in the human genome and has seven exons (fig. 3*c*), each with consensus exon-intron boundaries (table 4), and for which exons 3–7 encode a NIPA2 putative polypeptide of 360 amino acids (fig. 4*b*). The mouse *Nipa2* cDNA (see the "Material and Methods" section) is very similar to human *NIPA2,* having seven conserved exons (fig. 3*c;* table 4), encoding a putative polypeptide of 359 amino acids in exons 3–7 (fig. 4*b*), and with this locus spanning 29.8 kb in the mouse genome. Two forms of the mouse *Nipa2* cDNA (1,867 bp and 3,200 bp; fig. 3*c*) differ in the positions of alternative polyadenylation sites in the 3' UTR, created through the use of $poly(A)$ signals at cDNA positions 1846–1851 (ATTAAA) and 3131–3136 (AATAAA). Both transcripts are observed in different tissues (see below).

We also sequenced and assembled a 1,805-bp *Gallus gallus Nipa2* cDNA, encoding a 361–amino acid putative polypeptide. Adjacent to the *Fugu Nipa1* gene, we identified the *Fugu Nipa2* ortholog, encoding a 366– amino acid putative polypeptide. There are five coding exons that correspond to exons 3–7 of the orthologous human and mouse genes (fig. 3*c*) with conserved exonintron boundaries (table 4), although putative noncoding exons 1 and 2 have yet to be identified in *Fugu*. The *Fugu Nipa1* and *Nipa2* genes were found in conserved positions with respect to the mouse locus (fig. 1*f*), and *Fugu* orthologs for the *p, Herc2, Cyfip1,* and *Gcp5* genes were also identified in conserved positions (data not shown), providing strong evidence that each represents the ortholog of the mammalian genes. Finally, the position of the splice donor and acceptor sequences for the fourth intron of *NIPA2* is conserved in all homologs analyzed (fig. 3*c*), including the single intron present in the insect "ancestral" gene, intron 2 for the "ancestral" *C. elegans* gene (total of six exons), and intron 2 of an *Arabidopsis* paralog with nine total exons.

Comparison of human *NIPA2* and mouse *Nipa2* cDNAs demonstrates two peaks of homology, one that begins at the NIPA2 start codon in exon 3 and the other that is a short region (∼120 bp) in exon 1 (fig. 3*d*). The latter encodes a potential 36-to-40–amino acid upstream ORF (uORF) that is conserved in mammals (figs. 3*d* and 3*e*), and several additional uORFs occur in exons 1–3. Distantly related uORF sequences also occur in the *NIPA2* 5' region in chicken (where it is the second uORF) and *Xenopus* (fig. 3*e*). A conserved uORF, such as that observed here in *NIPA2* loci, is likely to regulate NIPA2 translation (Morris and Geballe 2000).

Polypeptide and Phylogenetic Analyses of the NIPA1 and NIPA2 Family

Analysis of the human, mouse, chicken, *Xenopus,* and *Fugu* NIPA1 and NIPA2 polypeptide sequences (figs. 4*a* and 4*b*), as well as the ancestral invertebrate and related plant members (fig. 4*b*), identified nine hydrophobic segments in each that likely encode putative transmembrane helices (TMH) (fig. 4*c*). Although the TMHMM program assigned a lower probability for TMH6 in human and mouse NIPA1, it is nevertheless clear that there is a hydrophobic domain of sufficient length to span the lipid bilayer and that TMH6 is clearly predicted in *Fugu* NIPA1, as well as in all related proteins (fig. 4*c*). We conclude that the *NIPA1* and *NIPA2* gene family encode polypeptides that have nine transmembrane-spanning helices.

Alignment of amino acid sequences of the NIPA1 polypeptide from human, mouse, and *Fugu* reveals these to be highly conserved (fig. 4*a;* table 5). Indeed, the human and mouse sequences display 98% identity, with a single amino acid change other than the length of the polyalanine tract, whereas NIPA1 in *Fugu* is ∼55% identical to that in mammals. Alignment of amino acid sequences of the NIPA2 polypeptide from human, mouse, chicken, *Xenopus,* and *Fugu* shows identities ranging between 78% and 96%, with the C-terminus being the most divergent (fig. 4*b;* table 5). Remarkably, NIPA1 and NIPA2 polypeptide sequences are related, with 32%–36% identity (and 47%–50% similarity) between any paralogous pair within these five species. Similarly, the ancestral *Drosophila, C. elegans,* and *Anopheles* polypeptides and a representative *Arabidopsis* member are highly related, with 40%–49% identity to vertebrate NIPA2 and lower identity (29%–34%) to vertebrate NIPA1 (table 5). The phylogenetic relationships are best seen in a tree diagram, which clearly illustrates the

grouping of NIPA2 and NIPA1 homologs within vertebrates, and the distant ancestral members in invertebrates and *Arabidopsis*. The phylogenetic distance between NIPA1 and NIPA2 paralogs is as great as that between them and ancestral members, strongly suggesting that NIPA1 and NIPA2 have evolved related but different functions.

Analyses of Imprinting for BP1–BP2 Genes and Mouse Orthologs

The most proximal known imprinted gene in 15q is *MKRN3,* which is located just distal to BP2, although the presence of replication asynchrony in the pericentromeric region (see the "Introduction") and at mouse *Nipa1*-*Nipa2*-*Cyfip1* (fig. 2*c*) raises the possibility that the imprinted domain could extend further. Consequently, we tested whether the four BP1–BP2 region genes might be imprinted in human, using RNA from cells from patients with imprinting mutations who have two maternally (PWS) or two paternally (AS) imprinted chromosomes 15 (Buiting et al. 1995; Saitoh et al. 1996), as well as a somatic cell hybrid imprinting assay in which the rodent cell lines retain either a maternal or a paternal human chromosome 15 (Gabriel et al. 1998). RT-PCR shows that human *NIPA1* (both the major mRNA detected by a 167-bp band and a larger form with the alternatively spliced exon 3b), *NIPA2, GCP5,* and *CY-FIP1* are each expressed in lymphocyte cells from a normal individual and PWS and AS imprinting mutation patients (figs. 5*a* and 5*b*), whereas *SNURF-SNRPN,* a paternally expressed imprinted gene in the PWS region as a control is only expressed in the normal and AS individuals (fig. 5*a*). In addition, *CYFIP1* is expressed from both the maternal and paternal chromosome 15 in somatic cell hybrids (fig. 5*b*). Therefore, the four BP1– BP2 genes are nonimprinted in peripheral blood lymphocytes and in fibroblasts.

To test imprinting in the mouse, we used a transgenic PWS and AS mouse model that has a paternally or maternally derived chromosome 7C deletion, respectively, spanning at least from *Frat3* through *Herc2* (Gabriel et al. 1999; Chai et al. 2001). On the basis of the deletion of BAC 252P22 (fig. 2*b*), the 3- UTR of *Cyfip1* and all of *Nipa1* and *Nipa2* are included in the deletion (see fig. 1*f*), as are 5' Cyfip1 and Gcp5 (J.M.G., J-H.C., and R.D.N., unpublished data), allowing this model to be used to test imprinting of these genes. All four genes (*Nipa1, Nipa2, Cyfip1,* and *Gcp5*) are expressed in wildtype, TgPWS(del), and TgAS(del) mice, resembling the control *Mkrn1* profile (fig. 5*c*). In contrast, the imprinted *Snurf-Snrpn* gene is not expressed in the PWS mouse, as expected (fig. 5*c*). Therefore, mouse *Nipa1, Nipa2, Cyfip1,* and *Gcp5* genes are expressed from both the paternal and maternal alleles and are nonimprinted in the brain.

b

Figure 3 Vertebrate *Nipa1* and *Nipa2* gene structures. *a,* Schematic genomic structure of *NIPA1* in human, mouse and *Fugu*. Coding regions (*shaded rectangles*) and untranslated (5' UTR and 3' UTR) regions (o*pen rectangles*) are shown. The horizontal black bars are CpG islands, the arrows below the exons (e) are primers used for RT-PCR, and the vertical dotted lines are functional polyA signals (predicted in *Fugu*). Alternative polyadenylation generates two different 3' ends for human and mouse *Nipa1* (with distance between the polyA signals shown). b, Highly conserved 3' ends of the mammalian *NIPA1* 7.5-kb mRNA sequences from five eutherian species, aligned by CLUSTAL W. Black nucleotides with gray background agree with the consensus, and polyA signals are shown as white letters on black background. GenBank accession numbers for the *NIPA1 3'* ESTs are as follows: human, BF439642; pig, BI339387; cow, BE685351; mouse, BE946294; and rat, AW533027. *c,* Genomic structure of orthologous human, mouse, and *Fugu NIPA2* genes, as well as conserved intron placement in ancestral genes from *Drosophila* and *Anopheles*. Symbols as for panel *a*. The shaded box in exon 1 represents uORF1, but putative 5' noncoding exons in *Fugu (dotted line)* have not been identified. *d, Conserved ORFs in the 5'* UTR (exons 1–3) of the vertebrate *NIPA2* transcripts. White letters on black background represent sequences conserved in all species shown, black letters with gray background have one mismatch, and the initiation codons of *NIPA2* and uORF1 have bold white letters on black background. Also shown are exon (ex) positions for the mouse (mu) gene, and the stop (TGA or TAA) codons for the uORF1. GenBank accession numbers for the *NIPA2 5'* ESTs are as for fig. 4*b*,, and the GenBank accession number for dog is BM538411. Alignments were generated with CLUSTAL W. *e,* Amino acid sequence of the putative exon 1 uORF1 from human, mouse, cow, dog, chicken (uORF2), and *Xenopus NIPA2* mRNAs. White letters on black background represent sequences conserved by comparison with the mammalian consensus, and black letters with gray background represent sequences conserved in fewer species, and italics represent chemically similar amino acids. GenBank accession numbers are as for fig. 4*b,*and the GenBank accession number for chicken is AJ452290.

a

d

human2 CTAGGTCCCGGCAGCGGTGGTGACGGCGGTGCCGGAGGTTGTCCTTGGCAG 60 **TT** ----------GGTGCTGGCCGCGGTGCGGCCGAAGGCTGTCCGA<mark>GGT</mark>AGGC<mark>T</mark>CCT $mouse2$ 48 dog₂ -- GCACGAGGCC $AC($ 22 start uORF1 human2 CATGGAGGAGGC г٣ 120 ïС GATGC $mouse2$ CCATGGAGGAGGCG GAACTT C **GC** ĦТ 103 **ATGGAGGAGO** dog₂ ГT 82 human2 171 mouse2 $-GC$ 154 dog₂ GGCGGCGGCCC 142 stop uORF1 human₂ F<mark>TGAGCCCCCGTG</mark> 180 TCGCC $-CCG$ C
G mouse₂ AGC CGGGC GT GGCZ GAGCCCACTC 209 FTC-30 GC GGACCTGACGAGCCCCAGCGACTGGA dog2 AAGC 202 נה AGCCC start ex2 (mu) 3GGCTGGAGC<mark>TACT</mark>CGGCCAGCGTTTAAGA
AGGGTGGAAG<mark>TACT</mark>CTGCCGTGGTTAAATA human2 ATGA 232 YТ ACGGCCCTAAGGGTGGAACTA mouse2 TTAAATGG 269 CT GAAACCTACCATTGGAAGTATTTAAGAAGCGAATGCCTCTGGATTTTG chicken2 46 dog₂ GCCCGGG 215 GAGTGAATAAG<mark>T</mark>CA human2 TCTCTAACCTCTCCCA 352 ATGAGAATGAATAAATCATCTCTG-AAA mouse2 **AGCGIVINI** CCAA **EGA** $--- 319$ chicken2 TGCTC -- TGGCTGTTCTCAA- $- - -$ 95 dog2 $\sqrt{\frac{3}{2}}$ start ex3 (mu) human2 GCCTTTTTTTCATA CCATATTAAACTT **ACATETTY** 412 TTTTTTAAAAGA $mouse2$ CCATATTAAACTTACATGTTT $CTTC$ 374 chicken2 -GCTCTGTGA 129 $-CTAACTT$ TG CAATGTTI dog₂ 237 human2 472 ATCAATGTGATTCACAGGAACTCGTTAAC 472
ATCAATGTCACTCACAGAACT - - TAAGC 431
ATCAATGTG - - - ACAGTCCCC - - - - AC 182
C<mark>TCAATGTGACTCACA</mark>AGAACTTCTTTAC 297 "AATG" mouse2 chicken2 נידי **CTGTT** eathe $dog2$ **TIGTTTTTGTC** Nipa2 start codon human₂ **ATGA CCAGGGC** 509 CGTGG AATATGA mouse2 ACGA **ATGA** 468 CGTGG **G**ATATGA chicken2 ACGA **PATGZ** CGTGG 228 $doq2$ **ATATGA** 334 $\mathbf e$ R<mark>A</mark>AAAPNALASFCTATKALAARA<mark>FE</mark>AAARAACVGV Dog IEEAVRTAAA<mark>L</mark>DA<mark>H</mark>TSHWT<mark>E</mark>TKAVA<mark>A</mark>RAQEWVAGAACVGV Cow MEEAVRTA<mark>S</mark>APNARISHWTATKAVAVRAQ*D*W **AACVGV** Human MEEAALTA-VSDARTSRGTSPETVAVCAQER AACVGV* Mouse AEHRHSDEVSDPHLKSVD----ESEEKGINVGPTQA* Xenopus

MELNRNLPLEVFKKRMPLDFGTNCEFMWKRCCGCSQAL*

Expression Analyses of BP1–BP2 Genes and Mouse Orthologs

ChickuORF2

The tissue patterns of expression for the four BP1– BP2 region genes and their mouse orthologs were studied by northern blot analysis. As predicted from cDNA and genomic analyses (see above), a single 2.4-kb transcript is seen for human *NIPA2* (fig. 6*a*), whereas 1.9 kb and 3.2 kb transcripts occur for mouse *Nipa2* (fig. 6*b*), with constitutive expression in both species including pronounced expression in the placenta (figs. 6*a* and 6*b*). In

mouse, a 1.9-kb *Nipa1* transcript is also constitutive but at relatively low levels, whereas a larger (7.5-kb) mRNA is seen in several tissues, with both mRNA isoforms enriched in brain (fig. 6*b*). Similarly, the human *NIPA1* locus expresses a short (2.2-kb) and a long (7.5-kb) mRNA at low levels, with enrichment of the larger isoform in neuronal tissues (Rainier et al. 2003 [in this issue]). Similar studies with *CYFIP1* demonstrate a 4.4 kb constitutively expressed mRNA in human (fig. 6*c*) and mouse (fig. 6*d*), with highest levels in placenta (figs.

Figure 4 NIPA1 and NIPA2 polypeptides. *a,* Amino acid alignment of NIPA1 orthologs in human, mouse, and *Fugu*. White letters with black background represent identical residues, black letters with light gray shading indicate conserved substitutions. Putative transmembrane (Tm) domains are designated by brackets, and the positions of exon (e)-intron boundaries are shown. GenBank accession numbers are: human NIPA1, BK001020; mouse Nipa1, AY098645; and *Fugu* Nipa1 was predicted from genomic sequence (see the "Material and Methods" section). *c,* Amino acid alignment of NIPA2 orthologs from human (GenBank accession number BK001120), mouse (GenBank accession number BK001121), chicken (GenBank accession number AY099502), *Xenopus* (GenBank accession number BK001125), *Fugu* (predicted as for panel *a*), ancestral genes from *Drosophila* (GenBank accession number AE003637), *Anopheles* (predicted from GenBank accession number AAAB01008980), *C. elegans* (GenBank accession number AC006804), and a representative *Arabidopsis* homolog (GenBank accession number AY046035). Symbols as for panel *a*. *c,* Representative transmembrane helices based on hydrophobicity plots. *d,* Phylogenetic comparison of vertebrate NIPA1 and NIPA2 orthologs, ancestral invertebrate homologs and a representative *Arabidopsis* homolog. The tree was constructed using CLUSTAL W, and branch lengths are proportional to sequence divergence. Each polypeptide is designated as the species, and the number 1 or 2 designates NIPA1 or NIPA2, respectively, except for ancestral members.

 ${\sf d}$

Exon-Intron Structures of *NIPA1* **and** *NIPA2* **in Human, Mouse, and** *Fugu*

^a Lowercase letters refer to intron sequences, and uppercase letters refer to exon sequences. Underlined letters "ag" and "gt" represent consensus splice acceptor and donor sequences, respectively.

b Exon encoding AUG start codon.

 ϵ Low level alternatively spliced exon; NA = not applicable.

^d First polyA signal (human, mouse) or stop codon (*Fugu*).

^e Noncoding 5' exon (may encode uORF).

Table 5

Phylogenetic Comparison of NIPA2 and NIPA1 Orthologs/Paralogs

Phylogenetic Comparison of NIPA2 and NIPA1 Orthologs/Paralogs

Figure 5 Imprinting assay of BP1–BP2 genes by RT-PCR in human and mouse. *a, NIPA1*, *NIPA2,* and *GCP5* mRNA expression was examined in lymphoblast cell lines derived from a normal individual and from patients who have PWS (PWS-U) and AS (AS-J) with imprinting defects. An imprinted control gene, *SNURF-SNRPN*, shows paternal-only expression, as detected in the AS but not the PWS cell line. + = RT present; - = RT minus control. *b, CYFIP1* imprinting analysis in families PWS-U and AS-J with imprinting defects and in somatic cell hybrids with a single maternal (Mat) or paternal (Pat) human chromosome 15. *c, Nipa1, Nipa2, Cyfip1, Gcp5,* imprinted control *Snurf-Snrpn,* and control *Mkrn1* mRNA expression in mouse. Brain mRNA from wild-type (WT), or transgenic-deletion mouse models of PWS (Tg^{PWS}) and AS (Tg^{AS}) were used for RT-PCR. *Mkrn1* is a nonimprinted control gene from mouse chromosome 6A (Gray et al. 2000). Paired lanes show reactions with $(+)$ or without $(-)$ reverse transcriptase.

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Figure 6 Expression of BP1–BP2 genes in human and mouse tissues. *a*, Expression of human *NIPA2* in various tissues by northern analysis, with b*-ACTIN* as a control. *b,* Constitutive expression of *Nipa2* and brain-enriched expression of *Nipa1* in mouse tissues by northern blot analysis, with *Gapdh* as a control. *c,* Expression of *CYFIP1* in different human tissues and regions of brain, with *SNURF-SNRPN* or b*-ACTIN* as controls. *d,* Mouse *Cyfip1* is widely expressed and enriched in placenta. *Gapdh* expression serves as a control. *e,* Human *GCP5* is expressed at high levels in muscle and at lower levels in other tissues. *f,* Mouse *Gcp5* is constitutively expressed in different tissues. In each case, radioactivity from each blot was stripped prior to successive hybridizations. $B = brain$; $C = colon$; $Cl = cerebral$ cortex; Cm = cerebellum; $F =$ frontal lobe; H = heart; K = kidney; Leu = peripheral blood leukocyte; Li = liver; Lu = lung; M = medulla; O = occipital lobe; Pa $p =$ pancreas; Pl = placenta; Pu = putamen; S = spinal cord; Si = small intestine; Sm = skeletal muscle; Sp; spleen; T = testis; Th = thymus; and $Tl =$ temporal lobe.

6*c* and 6*d*) and equal levels throughout the brain (fig. 6*c*). *GCP5* has a 3.7-kb transcript in human (fig. 6*e*) and mouse (fig. 6*f*), with the former enriched in muscle tissues, whereas a smaller (2.8-kb) *GCP5* mRNA is also seen in some tissues. Mouse *Gcp5* is also enriched in placenta (fig. 6*f*), as is true for *NIPA2* and *CYFIP1* in human and mouse.

Discussion

Chromosome rearrangements involving BP1 or BP2 in 15q11.2 include proximal deletion end points for PWS/ AS deletions (Amos-Landgraf et al. 1999; Christian et

al. 1999), duplications, and triplications (Roberts et al. 2002) and for inverted duplicated chromosomes (inv dup [15]) (Cheng et al. 1994; Leana-Cox et al. 1994; Huang et al. 1997). Numerous duplicons mapping to the BP1, BP2, and BP3 segments appear to be transcribed pseudogenes, including *HERC2* duplicons, a truncated copy of poly(A)-specific ribonuclease from 16p13, LCR15 duplicons that include the golgin-like protein gene and *SH3P18* also mapped at 15q24 and 15q26, and sequences encoding an ATP-binding cassette protein, a BEM-1/BUDS suppressor-like protein, and *MYLE* (Buiting et al. 1999**;** Amos-Landgraf et al. 1999; Christian et al. 1999; Ji et al. 1999; Pujana et al. 2001).

Similarly, only large duplicated and rearranged blocks of sequences derived from other chromosome locations have been mapped centromeric of BP1 (Ritchie et al. 1998; Fantes et al. 2002). Here, we have unequivocally demonstrated that four highly conserved genes (*NIPA1, NIPA2, CYFIP1,* and *GCP5*) are located in a genomic domain between BP1 and BP2. Our observations have significant implications for the delineation of the PWS imprinted domain; for understanding the basis of nonimprinted genetic diseases that map to this chromosomal region in human and the homologous chromosome region in mouse, including modification of the PWS or AS phenotype; and for an understanding of the fluidity of the genome during evolution and the way this mirrors events that occur mechanistically as de novo events in human disease.

Implications for Genomic Imprinting Mechanisms

All four genes between BP1 and BP2 in 15q11.2 were shown here to be nonimprinted in lymphoblastoid cell lines and human-rodent somatic cell hybrids, with the mouse orthologs nonimprinted in brain. The latter observation is important because most, if not all, the imprinted loci in the PWS/AS imprinted domain are predominantly expressed and imprinted in the brain (Nicholls and Knepper 2001). We conclude that the mammalian *NIPA1, NIPA2, CYFIP1,* and *GCP5* genes are nonimprinted. Although the four mouse genes map ∼2 Mb from the imprinted domain, the human orthologs are separated from the PWS imprinted domain solely by BP2 sequences; however, the size of BP2 is poorly defined because of the complex and incompletely known composition of duplicons. Since the *HERC2* duplicons within BP2 include the CpG-rich promoter and are derived from a nonimprinted locus (Amos-Landgraf et al. 1999; Ji et al. 1999), we suggest that one model to explain the proximal imprint boundary might simply be the evolutionary integration of "nonimprintable" promoter sequences adjacent to the imprinted domain. In contrast, when genes such as mammalian *MKRN3* and rodent *Frat3* were duplicated by retroposition, they had promoters that were subject to the imprint process when these genes integrated into or adjacent to the imprinted domain (Chai et al. 2001).

Several studies have shown replication asynchrony in domains with monoallelic gene expression, including those imprinted (Kitsberg et al. 1993; Chess et al. 1994; Knoll et al. 1994; Hollander et al. 1998; Simon et al. 1999; Mostoslavsky et al. 2001) and with a transition from imprinted to nonimprinted sequences at mouse *H19,* defined by an asynchronous to synchronous replication change (Greally et al. 1998). The use of marked mouse chromosomes 7 gave us an extremely useful tool for analysis of replication patterns at the *Nipa1*-*Nipa2*-*Cy-* *fip1* domain, where we found asynchrony that was random with respect to parental origin. A similar pattern has been described elsewhere for the *P* (OCA2) locus in human 15q13 (Knoll et al. 1994), whereas imprinted loci replicate the chromosome derived from one parent (usually the paternal allele) earlier than the other (Kitsberg et al. 1993; Knoll et al. 1994; Simon et al. 1999). The OCA2 locus is ∼1.5–2 Mb from the imprinted domain, whereas the mouse region studied here is ∼2–2.5 Mb from the mouse 7C imprinted domain. It is possible that the transition from asynchrony determined by parent of origin to the pattern of random asynchrony that we have demonstrated here is detectable only with the use of a marked chromosome such as that used in this study. Our finding that the four genes in the BP1–BP2 region are nonimprinted suggests that the asynchronous DNA replication observed proximal to deletion breakpoint BP1 (Ritchie et al. 1998) may be another example of the same phenomenon and does not indicate a more extensive imprinted domain. Rather, replication asynchrony may identify those chromosomal regions with the capacity for monoallelic gene expression: however, we propose that other factors, including DNA methylation and/or elements of the histone code (Jenuwein and Allis 2001; Turner 2002), are mechanistically needed to complete the silencing or activation of individual alleles.

Functional Considerations of a New Gene Family

Two of the four genes identified in the BP1–BP2 region are the related *NIPA1* and *NIPA2* genes. Phylogenetic studies indicate that both paralogous gene members are ancient and likely arose 450–600 million years ago, around the time of the origin of vertebrates. Despite the adjacent chromosome linkage of *NIPA1* and *NIPA2* and retention of this syntenic relationship in all examined vertebrates, there is no evidence that one served as the precursor to the other. Indeed, four additional NIPA1/ 2-related paralogous family members are dispersed in vertebrate genomes, and each appears to have arisen early in vertebrate evolution (J-H.C. and R.D.N., unpublished data). Alternatively, *NIPA1* or other family members may be evolutionarily younger, and, after their origin, they may have gone through a period of rapid mutation, followed by fixation and a dramatic slowdown in mutation rate. Further study of this gene family in extant vertebrate species will shed light on the evolutionary history of this novel gene family.

NIPA1 and *NIPA2* differ in gene structure and expression patterns, reflecting their ancient origins; however, in both cases, there are five coding exons, and these show homology in intron placement. In particular, the only intron in the insect *NIPA1/2* gene is conserved in position in all vertebrate *NIPA2, NIPA1,* invertebrate, and plant family members, despite variation in exon number. During evolution of *NIPA2*, two 5' noncoding exons arose along with the coding potential for a small uORF. The presence of one or more uORFs in *NIPA2* suggests that this gene undergoes translational regulation (Morris and Geballe 2000) and that translation of *NIPA2* may involve an internal ribosome entry site mechanism (Fernandez et al. 2002), although further work will be necessary to examine these possibilities. In contrast to the constitutive expression of *NIPA2* and the smaller *NIPA1* transcript in all tissues of human and mouse, *NIPA1* shows high levels of brain-enriched expression of a 7.5-kb transcript (as shown here and in the article by Rainier et al. 2003 [in this issue]). The mechanism for this may reflect stabilization of the mRNA in brain for the long form of *NIPA1,* transcriptional activation plus coordinate polyA site selection in a neuron-specific manner, or silencing of the latter transcription-polyadenylation pattern in nonneural tissues, as occurs with the neuron-restrictive silencer factor (Zhao et al. 1999; Lunyak et al. 2002; Proudfoot et al. 2002; Worthington et al. 2002).

Our prediction of a nine-TMH helix domain structure for the NIPA1 and NIPA2 polypeptides suggests that they function as either receptors or transporters. NIPA1/ 2 show low homology, spanning three TM domains (∼29% over ∼93 amino acids), with some G-protein coupled receptors, such as extracellular calcium receptors and olfactory receptors (data not shown); however, this may reflect the hydrophobic nature of TM domains, and any functional or evolutionary relationship remains conjecture. Indeed, polypeptides with nine TM spanning passes of the lipid bilayer are rare. Exceptions are glucose-6-phosphatase (Pan et al. 1998), a cardiac Na^{+/-}- Ca^{2+} exchanger (Qiu et al. 2001), and the TM9 domain superfamily (TM9SF) (Chluba-de Tapia et al. 1997); TM9SF members are localized to endosomes (Schimmöller et al. 1998), bind the synthetic ligands cyanopindolol and the β -adrenergic agonist SM-11044, and have roles in colon relaxation and eosinophil chemotaxis (Sugasawa et al. 2001), or are involved in adhesion and phagocytosis in *Dictyostelium* (Cornillon et al. 2000). However, the TM9SF family is unrelated in primary sequence to the NIPA1/2-family. Furthermore, whereas TM9SF members have a signal peptide (type I topology), members of the NIPA1/2-family do not, and the latter must have internal membrane targeting sequences. Consequently, the function of the NIPA1/2 family remains unknown. Nevertheless, the presence of distinct family members in vertebrates, invertebrates, plants, and bacteria (the present article; J-H.C. and R.D.N., unpublished data) indicates that these conserved polypeptides are likely to be critical for signaling within or between cells. Studies with antibodies and with animal models deficient in or overexpressing these genes, as well as iden-

tification of ligands for these polypeptides, are now necessary to determine their functions.

Implications of BP1–BP2 Region Genes for Hereditary Disease

Since blocks of duplicated sequences that are implicated in homologous recombination flank the BP1–BP2 region genes, chromosome rearrangements involving the BP1–BP2 region may lead to dosage imbalance or rearrangements of the *NIPA1, NIPA2, CYFIP1,* and *GCP5* genes. This includes loss of one allele in the larger class I chromosome deletions in PWS and AS, or gain of one or two alleles, in 15q11-q13 duplications and triplications as well as for inv dup (15) marker chromosomes (Knoll et al. 1990; Cheng et al. 1994; Leana-Cox et al. 1994; Huang et al. 1997; Amos-Landgraf et al. 1999; Christian et al. 1999; Roberts et al. 2002). On the basis of the finding of small inv dup (15) marker chromosomes with the BP1–BP2 region but not the PWS/ AS region in a series of patients with generally normal phenotypes, Huang et al. (1997) concluded that no genes in these small markers have clinically relevant dosage effects. In contrast, recent studies by Butler et al. (2003) found that PWS subjects with class I deletions (BP1– BP3) have a more severe phenotype than those with class II deletions (BP2–BP3), including greater self injurious behavior, deficits in adaptive behavior (including motor skills), obsessive-compulsive behavior, and difficulties with reading, mathematics skills, and visual-motor integration. These data provide evidence that deletions of the four genes in the BP1–BP2 region may be associated with dosage-sensitive behavioral and psychological phenotypes. We propose that some individuals may have a condition with just these types of clinical phenotypes and a deletion or duplication limited to the BP1–BP2 region due to recombination between duplicated sequences within BP1 and BP2. Further study of BP1–BP2 region genes and chromosome rearrangements will determine whether the presence of four highly conserved genes flanked by unstable DNA sequences can have a significant phenotypic impact.

A chromosome 15 rearrangement limited to the unique 250-kb BP1–BP2 region was recently found in a subject with PWS due to a 15q11-q13 deletion that arose on the paternally inherited chromosome containing a familial duplication limited to BP1–BP2 (Butler et al. 2002). Consequently, duplications, deletions, or inversions limited to the BP1–BP2 region may predispose to subsequent larger rearrangements of proximal 15q. Indeed, a polymorphic 1.5-Mb inversion within a similar class of duplicated sequences in chromosome 7q11.23 is associated with susceptibility to deletion of the 1.5- Mb region in Williams-Beuren syndrome (Osborne et al. 2001). Similarly, inversions of BP2–BP3 were identi-

Figure 7 Model for evolutionary transposition of BP1-BP2 genes by flanking duplicons. Genes are shown as horizontal arrows or arrowheads indicating the direction of transcription. Black arrows and white arrowheads represent nonimprinted and imprinted genes, respectively. *HERC2* and duplicons (dup) derived from *HERC2* are shown as gray arrows or bars when transcriptional direction is unknown. The rectangle indicates a putative transposition of a four- gene cassette with flanking *HERC2*-duplicons, in an ancestral primate. See text for further details. Not shown in this schematic is a cluster of three GABA_A receptor genes and *ATP10C*, located between the *UBE3A* and *P* loci in human and mouse, nor additional duplicons that are interspersed with *HERC2* duplicons but that are poorly characterized to date (see Nicholls and Knepper 2001).

fied in four of six mothers of patients with AS deletion (Gimelli et al. 2003), which supports the previous finding of a familial 15q11-q13 inversion in the mother and father of deletion patients with AS and PWS, respectively (Clayton-Smith et al. 1993), consistent with the large inversion being a predisposition allele. Indeed, BP1–BP2 deletions or other rearrangements may be more common than the PWS/AS deletions, which occur at an overall frequency of ∼1/10,000 newborns, since the distance between the putative recombining segments is only a fraction of that of the 4–4.5-Mb PWS/AS deletions.

All four BP1–BP2 region genes are candidates for dominantly inherited spastic paraplegia, locus 6 (SPG6 [MIM 600363]). SPG6 is characterized by insidiously progressive lower-extremity spasticity in which axonal degeneration affects primarily the longest axons in the CNS and is caused by a mutation within a 7.3-cM seg-

ment spanning the PWS/AS region (Fink at al. 1995). The finding of obligate recombinants for markers within the cluster of $GABA_A$ receptor genes in distal $15q11-q13$ ruled these out (Fink et al. 1996). As a dominant disorder, SPG6 is not expected to be associated with imprinted genes, ruling out most other genes in the PWS/ AS deletion region. Since no functional genes are known proximal to BP1, the BP1–BP2 region genes, particularly *CYFIP1* and *GCP5*, became prime candidates. CYFIP1 associates with Rac1 and F-actin (Kobayashi et al. 1998), with Rac1 implicated in regulation of the actin cytoskeleton, including axon growth, guidance, and branching (Ng et al. 2002). CYFIP1 also associates with FMRP (Schenck et al. 2001), the fragile X mental retardation protein, which is implicated in neurite extension, guidance, and branching (Morales et al. 2002). In addition, *GCP5* encodes a protein that is part of the human γ -tubulin complex required for microtubule nucleation at the centrosome (Murphy et al. 2001). Microtubules are critical in axonal transport, and defects in this process are known in spastic paraplegias (Crosby and Proukakis 2002). However, although *NIPA1* and *NIPA2* are of unknown function, the CNS-specific expression of *NIPA1* made it an attractive and correct candidate (Rainier et al. 2003 [in this issue]).

Mutation studies of the mouse orthologs of the four BP1–BP2 region genes will shed light on the potential phenotypic role of recessive loss-of-function mutations in these genes. Indeed, a locus required for embryo implantation, *l71Rl,* was identified just proximal to the pink-eyed dilution (*p*) gene on the basis of recessively inherited deletions of the region (Wu et al. 2000). Genetic complementation and molecular mapping define the critical interval for *l71Rl* as a small region between *D7Mit70* and 5' *Herc2* (Wu et al. 2000), to which we have mapped the mouse *Nipa1, Nipa2, Cyfip1,* and *Gcp5* genes, making one or more of these four genes likely candidates for *l71Rl*. Although the homologous human chromosome BP1–BP2 region is flanked by unstable DNA sequences, as discussed above, the mouse *l71Rl* data indicate that it is unlikely that individuals will exist who are homozygously deleted for the BP1– BP2 region, since such homozygous deletions would be predicted to be associated with a failure of the embryo to implant. Should BP1–BP2 deletions occur in association with a normal or mild neurobehavioral phenotype (see above), their inheritance from each parent might be associated with early pregnancy loss and thus may represent an important genetic counseling issue.

Evolutionary Transposition of BP1–BP2 Region Genes Mirrors Recombination Events in Genomic Disease

We have shown that a block of at least six genes (*OCA2, HERC2, NIPA1, NIPA2, CYFIP1,* and *GCP5*) are syntenic in human, mouse, and *Fugu;* however, in human, the latter four genes have transposed to a site ∼4 Mb away from the OCA2 and *HERC2* genes. These and previously published data allow us to propose a model for the evolution of human $15q11-q13$ (fig. 7). Mouse and *Fugu* are representative of the ancestral vertebrate arrangement of these genes (fig. 7). The evolutionarily young imprinted domain was subsequently added in mammals, adjacent to this ancestral domain (fig. 7) (Chai et al. 2001; Nicholls and Knepper 2001). During evolution of an ancestral primate, 25–60 million years ago (Amos-Landgraf et al. 1999; Christian et al. 1999; Ji et al. 1999; Locke et al. 2001), the *HERC2* gene began its evolutionary odyssey, and we propose that duplications of this gene first formed duplicons flanking a *NIPA1*-*NIPA2*-*CYFIP1*-*GCP5* cassette at a position equivalent to BP3 (fig. 7). Subsequently, the ∼250-kb

four-gene unique cassette (with all required *cis* regulatory elements) and flanking *HERC2* duplicons were transposed by a duplicon-mediated process in an ancestral primate to a site ∼4 Mb away, on the other side of the imprinted gene domain. Here, the flanking *HERC2* duplicons would now have formed the BP1 and BP2 domains, as found in human (fig. 7). This transposition may have involved *HERC2* duplicons directly, as supported by the location of a *HERC2* duplicon in BAC 26F2 immediately adjacent to *NIPA1* at BP1 (data not shown). Alternatively, other duplicons have been added and interspersed with *HERC2* duplicons at unknown times during primate evolution (Buiting et al. 1999; Christian et al. 1999; Pujana et al. 2001) and may have contributed to the genomic transposition. Genomic studies of primates will help determine the timing of these events and mechanisms, although secondary rearrangements within and between duplicated sequences in different extant primate species might complicate interpretation of the exact evolutionary events. Nevertheless, it is clear that an evolutionary genetic instability has been conferred on this domain by the origin and expansion of large blocks of duplicated DNA sequences, now mirrored by the events that mediate chromosome rearrangements in genomic diseases involving 15q11-q13.

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Electronic-Database Information

Accession numbers and URLs for data presented herein are as follows:

- Biological Software, Institut Pasteur, http://bioweb.pasteur.fr/ intro-uk.html (for the CLUSTAL W identity/similarity matrix program and the drawtree program to plot unrooted tree diagrams)
- *Fugu* BLAST Server, http://fugu.hgmp.mrc.ac.uk/blast/blast.cgi (for BLAST searches to identify the *Fugu p*, *Herc2*, *Nipa1*, *Nipa2*, *Cyfip1*, and *Gcp5* gene orthologs)
- GenBank, http://www.ncbi.nlm.nih.gov/Genbank/ (for mouse *Nipa1* cDNA [accession number AY098645], for chicken *Nipa2* cDNA [accession number AY099502], for human

NIPA1 cDNA [accession number BK001020], for human *NIPA2* cDNA [accession number BK001120], for mouse *Nipa2* cDNA [accession number BK001121], and for *Xenopus Nipa2* cDNA [accession number BK001125])

GeneMap '98, http://www.ncbi.nlm.nih.gov/genemap98/

- NCBI BLAST, http://www.ncbi.nlm.nih.gov/BLAST/ (forBLAST searches)
- Online Mendelian Inheritance in Man (OMIM),http://www .ncbi.nlm.nih.gov/Omim/ (for PWS, AS, and SPG6)
- TMHMM Server v. 2.0, http://www.cbs.dtu.dk/services /TMHMM-2.0/ (for TMH prediction)
- Trace Archive Database, http://www.ncbi.nlm.nih.gov /blast/mmtrace.html (for mouse genome sequences)

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