Fusion of the Human Gene for the Polyubiquitination Coeffector UEV1 with Kua, a Newly Identified Gene

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UEV proteins are enzymatically inactive variants of the E2 ubiquitin-conjugating enzymes that regulate noncanonical elongation of ubiquitin chains. In <em>Saccharomyces cerevisiae</em>, UEV is part of the RAD6-mediated error-free DNA repair pathway. In mammalian cells, UEV proteins can modulate c-FOS transcription and the G2-M transition of the cell cycle. Here we show that the UEV genes from phylogenetically distant organisms present a remarkable conservation in their exon–intron structure. We also show that the human UEV gene is fused with the previously unknown gene Kua. In <em>Caenorhabditis elegans</em> and <em>Drosophila melanogaster</em>, Kua and UEV are in separated loci, and are expressed as independent transcripts and proteins. In humans, Kua and UEV are adjacent genes, expressed either as separate transcripts encoding independent Kua and UEV proteins, or as a hybrid Kua–UEV transcript, encoding a two-domain protein. Kua proteins represent a novel class of conserved proteins with juxtamembrane histidine-rich motifs. Experiments with epitope-tagged proteins show that UEV1A is a nuclear protein, whereas both Kua and Kua–UEV localize to cytoplasmic structures, indicating that the Kua domain determines the cytoplasmic localization of Kua–UEV. Therefore, the addition of a Kua domain to UEV in the fused Kua–UEV protein confers new biological properties to this regulator of variant polyubiquitination.

Described recently as a class of proteins structurally related to the ubiquitin-conjugating enzymes (E2), a distinctive feature of UEV proteins is that they are inactive variants of E2 enzymes, lacking a recognizable catalytic center (Koonin and Abagyan 1997; Ponting et al. 1997; Sancho et al. 1998). These proteins are well conserved in sequence and structure in all eukaryotic organisms, and this results in the sharing of specific functions, such as protection of cells from DNA damaging agents (Broomfield et al. 1998; Thomson et al. 1998) and enhancement of transcription from the c-FOS promoter (Xiao et al. 1998), by UEV proteins from distant organisms. The biochemical mode of action of UEV proteins in the yeast <em>Saccharomycyes cerevisiae</em> has been established by Hoffman and Pickart (1999). The S. cerevisiae UEV protein, also known as Mms2, interacts with the E2 enzyme Ubc13p, and the resulting heterodimer is competent for the elongation of polyubiquitin chains. A novel feature of the polyubiquitin chains thus formed is that Lys 63, instead of the ca-
nonical Lys 48, is used for the Gly–Lys isopeptide bonds between ubiquitin moieties (Hoffman and Pickart 1999). Modification of proteins by this variant polyubiquitin chain may be reversible, and could modulate the function of target proteins, without directing them for degradation (Spence et al. 2000).

In *S. cerevisiae*, UEV genes are part of the error-free DNA repair pathway regulated by RAD6 (Broomfield et al. 1998; Hoffman and Pickart 1999). In human cells, UEV1 (Rothofsky and Lin 1997) and UEV2/Mms2 (Xiao et al. 1998) promote the transcriptional activity of c-FOS, possibly through interactions with as yet unidentified DNA-binding transcriptional regulators (Xiao et al. 1998). Overexpression of UEV1 in human colon cancer cells induces the accumulation of cells in G2-M and polyploidy, apoptosis, and inhibition of cell differentiation (Sancho et al. 1998). How the participation of UEV proteins in all these processes relate to the activity of UEV proteins as coeffectors in the polyubiquitination of target proteins has yet to be determined. In humans, there are two different UEV proteins encoded by separate genes, UEV1 or CROC1 (Rothofsky and Lin 1997; Sancho et al. 1998), and UEV2 or MMS2 (Sancho et al. 1998; Xiao et al. 1998). The UEV1 gene codes for two isoforms generated by alternative splicing, which share a common phylogenetically conserved UEV domain (Sancho et al. 1998). The isoform UEV1B contains a unique 82-residue amino-terminal extension, the B domain (Sancho et al. 1998).

The ubiquitin-conjugating enzymes are a large group of proteins, of which many variants exist in all eukaryotic organisms (for review, see Hershko and Ciechanover 1998). Although a common protein sequence and structural theme is shared between all E2 enzymes, attempts to assign primordial ancestors as the origin of one or more branches have not been met with success, mainly due to the great interspecies variability of functionally equivalent proteins. Being a new family of proteins with strong structural and functional links to the long-known ubiquitin-conjugating enzymes, UEV proteins and their genes could be useful to study the origins of the E2 proteins and their genes. Here we have analyzed the structure of the UEV genes in a number of organisms, and found that it is very conserved between phylogenetically distant organisms. As a relevant consequence of this analysis, we have found that the human UEV1 gene is part of a hybrid gene that results from the fusion of UEV1 with a second, previously unknown gene, which we have named *Kua*. We also show that, in humans, *Kua* and UEV1 can be expressed either as independent transcriptional units and proteins, or as a hybrid *Kua–UEV* transcript and protein. In contrast, in flies and worms the gene for *Kua* is unlinked to the gene for the corresponding UEV protein, and *Kua* and UEV are always expressed as separate proteins.

**RESULTS**

**Precise Conservation of the Positions of Introns in UEV Genes from Distant Organisms**

We have analyzed the architecture of UEV genes in those organisms for which the complete sequences of the loci are available. Exons were predicted by applying gene prediction algorithms (Guigó et al. 1992; Socolovsky et al. 1994; Burge and Karlin 1997) to DNA sequences from genomic cosmid clones, and validated by stringent alignments with expressed sequence tags. This analysis yielded the exon arrangements schematically depicted in Figure 1A, confirming the structure predicted for the human *UEV1* gene (Sancho et al. 1998), with three exons coding for the common domain of the protein (C domain), one for the domain specific of isoform UEV1A (A domain), and two for the domain specific of isoform UEV1B (B domain).

The common domain of the UEV protein is encoded by one exon in *S. cerevisiae*, four in *Schizosaccharomyces pombe*, two in *Plasmodium falciparum*, three in *Arabidopsis thaliana*, two in *Caenorhabditis elegans*, three in *Drosophila melanogaster* and three in *Homo sapiens* (Fig. 1A). These exons are preceded by an initial exon (exon 1; exon A in *H. sapiens*), not conserved in sequence between different organisms. The exon structure for the common domain of UEV is identical in *H. sapiens*, *A. thaliana*, and *D. melanogaster*. *S. pombe* is the organism in which the UEV gene has the most exons (five). *S. cerevisiae* is the only organism in which the common domain of UEV is not interrupted by introns. Introns 2 and 3 of *S. pombe* UEV have equivalents in all other organisms, except *S. cerevisiae*. The positions of these introns within the corresponding UEV genes are identical, regardless of the organism (Fig. 1). These introns lie between codons, and are thus in phase zero. Also, they correspond to boundaries between structural domains within the UEV protein (Fig. 1). Therefore, they have features predicted for “early” introns (Gilbert 1987; de Souza et al. 1998). Intron 4 of *S. pombe* UEV is unique to this organism, and has no correspondence in any other organism. This intron is in phase 2, and it interrupts a sequence coding for an α-helical domain within the protein (Fig. 1). Therefore, this intron would be more compatible with being a “late” intron (Gilbert 1987; de Souza 1998).

In contrast to the strong sequence conservation for the rest of the gene, the first exon of the UEV genes (exon A in *H. sapiens*) is not conserved to a significant degree between different organisms. The size of this initial exon varies from 4 bp (1 and 1/3 codons), in *C. elegans*, to 88 bp (29 and 1/3 codons) in *A. thaliana*, or 91 bp (30 and 1/3 codons) in *H. sapiens*. Intron 1, separating the first exon and the exons for the common domain of UEV, is placed in different phases in different organisms: phase 0 in *S. cerevisiae* and *P. falciparum,* and
phase 2 in *S. pombe*, and phase 1 in *C. elegans, A. thaliana, D. melanogaster*, and *H. sapiens* (Fig. 1). The 3' boundaries set by this intron impose different 5' ends for the common domain of UEV in different organisms, which results in one to three codon differences between UEV proteins from these organisms at the amino-end of this domain (Fig. 1). Similarly, there are no sequences resembling the exons coding for the B domain of human UEV1 in the vicinity of the UEV genes of any other organism. It has been suggested that
the B domain of human UEV1B could confer specific functions to this isoform (Sancho et al. 1998). We thus set our efforts to explore the evolutionary origin of the sequences coding for the B domain of UEV1B.

**A New Gene in *C. elegans* and *D. melanogaster* Coding for a Protein Containing a UEV1 B Domain-Like Sequence**

A search for B domain-like sequences in genomic DNA databases yielded two small fragments of significant similarity within *C. elegans* clone Y53C10 and *D. melanogaster* clone DS00863 (Fig. 2A). In the *C. elegans* genome, the segment included in Y53C10 is in chromosome 1, ~2.5 Mb away from the location of the UEV gene in clone F39B2, also in chromosome 1 (http://www.sanger.ac.uk). We have found no evidence for UEV-like sequences within *C. elegans* Y53C10 (86 Kbp) or *D. melanogaster* DS00863 (78 Kbp) genomic clones. Therefore, in contrast to the B domain-like sequences in *H. sapiens*, the B domain-like sequences in *C. elegans* and *D. melanogaster* do not appear to be part of the corresponding UEV genes.

We hypothesized that, in *C. elegans* and *D. melanogaster*, these sequences belonged to a second gene, unrelated to UEV. Mapping of known cDNAs and ESTs onto the genomic sequence indicated the presence of known genes upstream from position 17,000 and downstream from position 29,000 in Y53C10, and upstream from position 54,500 and downstream from position 62,400 in DS00863 (data not shown). Reciprocal TBLASTX searches (Altschul et al. 1990) and dot-plot analyses between the segments in Y53C10 and DS00863 that lie within these positions (denoted here Y53C10-B and DS00863-B) indicated that they shared other conserved segments in the vicinity of the B domain-like sequence, and delineated a tentative exonic structure for a gene in Y53C10-B and DS00863-B (Fig. 2B). The exonic structure of the putative gene in Y53C10-B was further refined by means of stringent alignments with ESTs from a nonredundant database, and computational gene identification programs (Fig. 2C). A recent addition to GenBank of a *C. elegans* EST (accession no. AV182903) provided strong support for this analysis, and a confirmation of the existence of a gene in this region. Thus, we predict in *C. elegans* Y53C10-B a new gene consisting of seven exons, with the potential to encode a 319-amino acid protein (Fig. 2E). This tentative structure was supported by a number of ESTs with distant, but significant similarity to sequence segments in DS00863-B (Fig. 2E). The exonic structure thus predicted was highly compatible with that delineated by the conserved sequence segments shared with Y53C10-B for the 3′ end of the putative gene. The final analysis resulted in a five-exon gene, with the potential to encode a 326-amino acid protein (Fig. 2F).

The *C. elegans* EST confirmed that the gene predicted is indeed expressed. However, evidence of this kind was lacking for the *D. melanogaster* gene. Therefore, we designed primers specific for the exons predicted for the *D. melanogaster* gene, for use in RT–PCR reactions. All sets of primers yielded specific amplification products from adult and larval RNA (Fig. 2G). Sequencing of the products showed that they correspond to processed RNA, formed by joining of the predicted exons, of which exons 4 and 5 are the B domain-like segments (Fig 2F). These experiments confirmed the exon structure for the new the gene, as predicted by computational methods. We have given the name *Kua* to this new gene (see Acknowledgments).

Clone DS00863, harboring the *Drosophila* *Kua* gene, is localized to segment 38B2–38C1, on chromosome 2 (Hartl et al. 1994; http://flybase.bio.indiana.edu/). There is no experimental data for the cytogenetic localization of the *Drosophila* UEV gene. Therefore, we performed in situ hybridization on polytene chromosomes, using a probe specific for *Drosophila* UEV. This permitted the cytogenetic assignment for the *D. melanogaster* UEV gene to 64D, on chromosome 3 (Fig. 2H). With posteriority to this analysis, the sequence of the *D. melanogaster* genome was released, confirming our assignment of *D. melanogaster* UEV to 64D (Adams et al. 2000).

In conclusion, in both *C. elegans* and *D. melanogaster* the “B domain”-like sequences contained in clones Y53C10 and DS00863, respectively, correspond to exons located within a new gene, *Kua*, expressed in worms and in flies. In both organisms, the genes *Kua* and UEV are in widely separated loci.

**H. sapiens: Fusion of UEV with Kua**

The entire human UEV1 gene, including exons coding for the B domain, is contained within the genomic PAC clone dJ1185N5 from chromosome 20. Conserved segments spanning a 32-Kbp region within this clone were found to correspond to most of *D. melanogaster* and *C. elegans* *Kua*. Identical or strongly related mouse and human EST matches fully covered this region. This, together with the application of gene prediction programs allowed us to predict a six-exon human *Kua* gene (Fig. 3A) with the potential to code for a 270-amino acid protein. Exon 1 is predicted to contain a 5′ untranslated region (UTR) of unknown size, and exon
Figure 2  A new gene in Caenorhabditis elegans and Drosophila melanogaster coding for B domain-like sequences. (A) Alignment of the human UEV1 B domain with TBLASTN-identified segments from C. elegans Y53C10 and D. melanogaster DS00683 genomic clones. The aligned segments are discontinuous in Y53C10, as indicated by the positions of nucleotides from the database entry. (B) Dot-plot analysis of reciprocal TBLASTX comparisons of genomic clones Y53C10-B (C. elegans) and DS00683-B (D. melanogaster). (Bottom) Scores of sequence identities for each diagonal of conserved segments, shown as horizontal bars. (C) Predicted structure of the new gene in C. elegans (middle) supported by gene prediction algorithms (top) and alignment with ESTs (bottom). (D) Predicted C. elegans protein containing a B domain-like segment (shaded). (E) Predicted structure of the new gene in D. melanogaster (middle), supported by gene prediction algorithms (top) and alignment with ESTs (bottom). (F) Predicted D. melanogaster protein containing a B domain-like segment (shaded). (G) Expression analysis by RT-PCR, with forward primers corresponding to exons 1, 2, 3, and 4 of the predicted gene, and a reverse primer corresponding to exon 5, using as templates embryo mRNAs. (H) Cytogenetic assignment of D. melanogaster UEV gene to chromosome 3 segment 64D. Digoxigenin-labeled cDNA probes were used for hybridization on wild-type Drosophila polytene chromosomes.
Figure 3  A hybrid Kua–UEV gene in Homo sapiens genomic PAC clone dJ1185N5. (A) Prediction of the exon structure of human Kua (middle), supported by gene prediction algorithms (top), and alignments with expressed sequences (bottom). (B) Schematic representation of the relative arrangement of exons in the human Kua–UEV locus, based on predictions for clone dJ1185N5. (C) RT–PCR analysis showing expression of human Kua (K1F + KR), a hybrid Kua–UEV transcript (K2F + UR), and UEV1A (A + UR) in six cell lines. RT–PCR was performed with primers for the putative 5′ UTR (K1F), the 3′ UTR (KR), or the 5′ UTR of Kua (KR), and the end of exon C3 of UEV1 (UR), using as templates RNAs from the human cell lines HT–29 (lane 1), SW480 (lane 2), HeLa (lane 3), SK–PC–3 (lane 4), T24 (lane 5), and HEL (lane 6). (Arrows) Specific amplification products; (asterisk) nonspecific amplification products. The faster-migrating band in the right panel corresponds to an alternative form of UEV1A (Sancho et al. 1998). (D) Expression analysis by RT–PCR of Kua–UEV hybrid transcripts and Kua-only transcripts on RNAs from cell lines Jurkat (lane 1), SK–NS–H (lane 2), N2A (lane 3), PZ–HPV7 (lane 4), CA–HPV10 (lane 5), PC–3 (lane 6), and SK–PC–1 (lane 7). Amplification with primers for transcripts corresponding to the housekeeping S14r ribosomal protein was used to normalize for input RNA.

6 is predicted to have a 3′ UTR extending 1268 bp from the stop codon up to the first potential polyadenylation signal (Fig. 3A,B). The exons predicted for the Kua gene lie 5′ to the exons for the common- and A-domains of UEV1 (Sancho et al. 1998; Fig. 3B).

Next we designed primers corresponding to the predicted exons of human Kua (Fig. 3B) for use in RT–PCR reactions (Fig. 3C). An amplification product was present in all of the cell lines analyzed, of the size expected (964 bp) for the joining of exons 1–6 of Kua (Fig. 3C). Sequencing of the amplification products confirmed that they contain all of the predicted exons, including those encoding the B domain of UEV1B (Fig. 4A,B). In the original reports (Rothofsky and Lin 1997; Sancho et al. 1998), the 5′ end of the transcript for UEV1B was described as a 5′ UTR. In our analysis, it is part of exon 4 of Kua, the B domain being encoded by exons 4 and 5. RT–PCR with primers specific for UEV1A yielded two products in all samples (Fig. 3C, right), corresponding to alternatively spliced isoforms described previously (Sancho et al. 1998).

The UEV1B isoform of UEV1 was isolated independently by two groups using either RACE or RT–PCR (Rothofsky and Lin 1997; Sancho et al. 1998). Our analysis shows that what was then described as a domain specific for the UEV1B isoform of UEV1 is encoded by two exons that are also part of a second gene, located upstream from UEV1, which we now call Kua. Thus, we hypothesized the existence of continuous transcriptional units joining the exons predicted for Kua and UEV1. Shown in Figure 3C are RT–PCR experiments performed with primers for the predicted exon 2 of Kua (forward primer) and exon C3 of UEV (reverse primer) yielding a specific product in the majority of the cell lines analyzed. Sequencing confirmed that these products correspond to the joining of exons 2, 3, 4, and 5 of Kua, and exons C1, C2, and C3 of UEV1 (Fig. 4). RT–PCR reactions with primers corresponding to regions in Kua that were not supported by our gene prediction analyses to be part of exons did not yield any specific amplification products (data not shown). Therefore, this analysis demonstrates the existence of a hybrid Kua–UEV transcript containing all exons from Kua and UEV1, except exon 6 of Kua and exon A of UEV1. This transcript has the potential to encode a protein with two distinct domains, Kua at its amino half, and UEV at its carboxyl half.
To determine the relative levels of Kua and Kua–UEV transcripts, RT–PCR was performed under nonsaturating conditions on RNAs from seven different cell lines. For normalization, amplification was performed with primers for transcripts for the ribosomal protein S14. In these analyses, the ratio of Kua–UEV to Kua amplification products ranged from 0.1 (samples 1 and 2) to 0.02 (samples 5 and 7), in those samples with visible Kua–UEV amplification products (Fig. 3D). Two samples did not yield measurable levels of Kua–UEV hybrid amplification products under these conditions (30 cycles), although products were observed with 35 amplification cycles or more (data not shown).

**Kua: A New Family of Conserved Proteins**

Several organisms express Kua transcripts, including *H. sapiens, Mus musculus, D. melanogaster, C. elegans, Trypanosoma cruzi*, each of which was predicted to express one Kua protein, and the plant *A. thaliana*, in which we predict three different Kua proteins corresponding to distinct genes (Fig. 5A). The three animal Kua proteins show identities of 94%, 67%, and 61% with respect to the human protein (comparison with the sequence between positions 83 and 238 of the human protein), whereas the three plant Kua proteins, strongly related to each other, show 29%–34% identities with respect to the human protein. Although yeasts have UEV proteins (Broomfield et al. 1998; Sancho et al. 1998), they do not have potential Kua proteins.

A pattern of histidine residues conserved in all Kua proteins is reminiscent of certain histidine-rich motifs in which specific residues are coordinated by transition metals (Shanklin et al. 1994). Membrane-bound fatty acid desaturases and hydroxylases contain several histidine-rich motifs, with the general pattern H-x(2,3)-(x-H)-H (Fox et al. 1993; Mitchell and Martin 1997). Kua proteins contain two of these motifs (Fig. 5A). Although all Kua proteins contain fatty acid hydroxylase-like His-rich motifs, only plant Kua proteins show a marginal degree of homology (E values in the order of $10^{-3}$) to fatty acid hydroxylases within the segment of the protein containing such motifs. Nevertheless, a PHI–BLAST search of a nonredundant database with a sequence profile generated based on the consensus sequence for Kua proteins yielded bonafide fatty acid hydroxylases (Fig. 5B).

In fatty acid hydroxylases and desaturases, the protein segments containing these motifs are located on the cytoplasmic face of the ER membrane, such that the relevant histidine residues become closely apposed for the coordination of a di-iron cluster (Mitchell and Martin 1997). A prediction of transmembrane domains for the Kua proteins indicates an overall topology similar to that of fatty acid hydroxylases (Mitchell and Martin 1997) (Fig. 5C, left panel), especially when comparing the transmembrane prediction for the plant Kua proteins (e.g., F19K23.12) with that for *A. thaliana* Fah1 (Fig. 5C, left panel). The positions of the histidine-rich motifs relative to the predicted transmembrane domains of Kua proteins and fatty acid hydroxylases are diagrammatically represented in Figure 5C (right panel).

**Subcellular Localization of UEV1A, Kua, and Kua–UEV**

A partial UEV1 protein has been reported to localize to the nucleus of mammalian cells (Rothofsky and Lin 1997).
**Figure 5**  Kua, a new family of conserved proteins. (A) Alignment of Kua proteins predicted from PCR-generated cDNAs (Homo sapiens and *Drosophila melanogaster*), ESTs (Mus musculus and *T. cruzi*), or genomic sequences (Caenorhabditis elegans and Arabidopsis thaliana, F19K23.12, F10M23.370, and T20K9.10). The prediction for the *A. thaliana* F10M23.370 genomic sequence is supported by EST AA712589. Color codes: blue background, His; yellow background, Gli and Pro; grey background, hydrophobic residues; blue, charged residues. Key for consensus sequences: a, aromatic (F, H, W, Y); c, charged (D, E, H, K, R); h, hydrophobic (F, L, M, V, W, Y, I); l, aliphatic (I, L, V); o, alcohol (S, T); p, polar (C, D, E, H, K, N, Q, R, S, T); s, small (A, C, D, G, N, P, S, T); u, tiny (A, G, S); x, helix breaking (G, P); , negative (D, E); +, positive (H, K, R). Potential transmembrane domains are boxed in the consensus sequence. (B) Alignment of the segments of proteins containing the two histidine-rich motifs detected by PHI-BLAST search of GenBank nr database with the profile H-x-(YWF)-x(8,25)-(RK)-x(2)-H-x(2)-H-H, generated after the consensus sequence for Kua proteins in this region. Shown are fatty acid hydroxylases from *D. melanogaster* (AC004280), *C. elegans* (C25A1.5), *A. thaliana* (AtFAH1), *Schizosaccharomyces pombe* (SpFAH1) and *Saccharomyces cerevisiae* (ScFAH1). (Bottom) Secondary structure predictions for yeast FAH (cylinders, alpha helix; arrow, beta sheet). (C) Transmembrane domain predictions and compared topological models for Kua proteins and fatty acid hydroxylases. Left panel, predictions of transmembrane domains for human (AF155120), worm (Y53C10A.5), and plant (F19K23.12), Kua and worm (C25A1.5), and plant (At Fah1) fatty acid hydroxylases. Right panel, diagrams representing the positions of the histidine-rich motifs in the same proteins, relative to the predicted transmembrane domains. Diagrams are not drawn to scale.
1997). The above analysis of the polypeptide sequence of Kua predicted its localization to endomembranes. To experimentally determine the subcellular localization of Kua and Kua–UEV, we generated constructs for the expression of Kua, Kua–UEV, and UEV1A in COS-7 cells, bearing in-frame a hemaglutinin tag at their carboxyl termini. The epitope-tagged full-length UEV1A isoform of human UEV1 showed a nuclear localization, with a uniform pattern and nucleolar exclusion (Fig. 6A). The full-length Kua protein localized mainly to cytoplasmic structures, with a pattern compatible with its association with the endoplasmic reticulum (Fig. 6B), thus providing experimental evidence supporting the predicted localization of Kua to endomembranes. Finally, the hybrid Kua–UEV protein was also associated with cytoplasmic structures, with a clear nuclear exclusion, in a pattern very similar to that displayed by Kua (Fig. 6C). In conclusion, two alternative forms of UEV1 are targeted to distinct subcellular localizations, nucleus, or cytoplasm. This differential targeting is determined by the sequence present at the amino terminus of UEV1, with the Kua domain in Kua–UEV directing its localization to cytoplasmic structures.

DISCUSSION
The analysis performed in this study allows us to postulate an evolutionary history of the genes for the polyubiquitination coeffecter UEV, and how they have become fused, in humans, to a second gene, which codes for a new class of proteins.

Evolution of UEV Gene Introns
The strict conservation in the positions of introns interrupting the coding sequences for the common domain of UEV in organisms as phylogenetically distant as yeasts and humans is a remarkable fact. The localization of introns in positions that are phylogenetically invariant has been observed in genes such as globin or alcohol dehydrogenase, with the development of additional insertions occurring as primordial genes have diverged (Naito et al. 1991; Sherman et al. 1992). For a given conserved protein, a certain degree of variation is frequently present in the location of introns relative to protein sequence, which is taken as supporting the “introns-late” theory (Cavalier-Smith 1991), as well as the occurrence of sliding and mutations in equivalent exon–intron junctions between genes from different organisms (Long et al. 1995; Gilbert et al. 1997). The position of introns 2 and 3 of S. pombe UEV is strictly invariant in distant organisms, P. falciparum (intron 3), A. thaliana (introns 2 and 3), C. elegans (intron 2), D. melanogaster (introns 2 and 3), and H. sapiens (introns 2 and 3). The maintenance of the positions of these introns through ∼5 × 10^8 years of evolution suggests an early insertion of these introns, an argument further supported by the observations that these introns are in phase zero, and placed between structural domains (Gilbert 1987; de Souza et al. 1998).

Conversely, intron 4 of S. pombe UEV interrupts an α-helical domain in the S. pombe UEV protein, which would be more consistent with a late insertion of this intron (de Souza et al. 1998). The absence of the equivalent of this intron in all other organisms suggests either the removal through splicing and reinsertion of processed and reverse-transcribed RNA (Fink 1987), or an insertion event specific to S. pombe, not transmitted evolutionarily. Absence of other S. pombe UEV introns in other organisms (all introns in S. cerevisiae, intron 2 in P. falciparum, intron 3 in C. elegans) could also be due to retrotransposition, or the result of independent origins and evolution of each intron. The conservation in S. cerevisiae of the 5′-most intron, in C. elegans of the two 5′-most introns, or the loss of the 3′-most intron in A. thaliana, D. melanogaster, and H. sapiens, would also be consistent with a scenario of retrotranscription followed by homologous recombination as a mechanism for the loss of introns in the evolution of the UEV genes (Fink 1987). In any case, it appears that both early and late scenarios (Trotman 1998) could apply to different UEV gene introns.

Figure 6 Subcellular localization of UEV1A (A), Kua (B), and Kua–UEV (C) proteins tagged with a hemaglutinin epitope. Liposome-mediated transient transfection was performed on COS-7 cells with plasmids engineered for the expression of the corresponding proteins bearing a hemaglutinin epitope at their carboxyl termini, and processed for indirect immunofluorescent confocal microscopy (× 400).
The analysis of intron 1 of the UEV genes suggests that it originated as a result of a process distinct from the ones discussed above for the introns that interrupt the common domain. Intron 1 is inserted in different phases in different organisms, and it generates different carboxyl- and amino-ends in the flanking exons. Also, exon 1 (exon A in humans), placed 5' to this intron, is different for all organisms. This suggests that exon 1 evolved separately from the rest of the UEV gene.

**Fusion of the Kua and UEV Genes**

We also show that, in humans, one of the two UEV genes in this organism, **UEV1**, is adjacent to an unrelated gene, which we have named **Kua**, and can be expressed as a hybrid **Kua–UEV** transcript and protein. In contrast, in *D. melanogaster* and *C. elegans*, **Kua** and **UEV** are independent genes, coding for separate transcripts and proteins. Therefore, the combination of computational and experimental approaches used here show how two genes, which are unrelated and located in separate loci in worms and insects, converge into a hybrid gene and protein in humans. The identification of **Kua** in *C. elegans* and *D. melanogaster* was possible without any prior knowledge of expressed sequences, applying ab initio biocomputational methods on genomic sequences. Of these methods, the use of dot-plot analyses of reciprocal TBLASTX alignments was sufficient to infer the structure of **Kua** in both organisms, later confirmed with alignments with ESTs and experimental data. Therefore, this could be a very useful tool for the identification of genes when expression data are not available.

New genes are thought to originate through events such as gene duplications (Ohno 1970; Ohta 1989), exon shuffling (Gilbert 1978), or the generation of processed genes (McCarrey and Thomas 1987). In metazoans, fusion of genes generally involves a number of intermediate processes, such as duplication and shuffling of exons (Bazan et al. 1989; Simmer et al. 1990; Chen et al. 1997; Coppock et al. 1998). Secondary events, such as retrotransposition with exon capture (Long and Langley 1993) or genetic hitchhiking associated with selective sweeps can generate chimeric genes (Nurminsky et al. 1998; Long et al. 1999). These events often involve extensive refashioning of coding and noncoding regions (Nurminsky et al. 1998). The **Kua–UEV** fusion does not appear to involve such promiscuous changes, and it rather suggests the occurrence of a direct fusion of loci. The likely scenario in the **Kua–UEV** fusion would be a two-step process, duplication of the **UEV** gene, followed by fusion of the duplicated gene to **Kua**. This model is supported by the fact that, in humans, there are two **UEV** genes, of which the **UEV1** gene is fused to **Kua** in chromosome 20, whereas the **UEV2** gene is on chromosome 8, without any evidence for this type of fusion (B. Cormand and T.M. Thomson, unpubl.). Therefore, **UEV2** would correspond to the gene in the original locus, and **UEV1** to the duplicated gene, which would undergo subsequent rearrangement with a head-to-tail fusion to **Kua**.

The generation of three different classes of transcripts from the **Kua–UEV** locus represents a unique strategy aimed at the modular expression of two genes, coding either for two separate polypeptides, or as a combination of both to yield a single two-domain polypeptide. The generation of a **Kua–UEV** hybrid transcript could be the result either of **cis**-splicing directed by canonical splice sites, or **trans**-splicing. **Trans**-splicing has been shown to occur in mammalian cells, either in artificial (Bruzik and Maniatis 1992) or natural (Caudevilla et al. 1998; Kingzette et al. 1998; Kopian et al. 1999; Li et al. 1999; Zaphiropoulos 1999) settings. Although **trans**-splicing in mammalian cells usually occurs between transcripts from genes in separate loci or chromosomes, it has been reported to occur also between transcripts from clustered genes (Zaphiropoulos 1999). It has also been shown that both **cis**- and **trans**-splicing can be concomitant mechanisms for the generation of hybrid transcripts from the same genes in mammalian cells (Eul et al. 1995). Our observations do not provide sufficient information to infer the splicing mechanism prevalent in the generation of hybrid **Kua–UEV** transcripts. However, there are indirect arguments against **trans**-splicing as the major mechanism for the generation of these transcripts. First, **trans**-splicing in mammalian transcripts appears to be regulated by sequences at the acceptor exon, with the consensus GAAGAAG(G/C) (Caudevilla et al. 1998). Sequences fully compatible with this consensus are present in exon C1 of **UEV1**, and also at equivalent positions in exon C1 of **UEV2**, but only **UEV1**, and not **UEV2**, is involved in hybrid **Kua–UEV** transcripts (T.M. Thomson, unpubl.). A more speculative argument would be based on the teleological nature of the fusion of **Kua** and **UEV** from the standpoint of the evolution of these genes. The driving force for the evolutionary rearrangement and fusion of **Kua** and **UEV** into a single locus would be stronger for **cis**-splicing being a major mechanism for the generation of hybrid transcripts than it would be for a **trans**-splicing mechanism.

The close proximity of **Kua** to **UEV1** could raise the question whether the mere juxtaposition of two genes with the same transcriptional direction is sufficient for the generation of detectable run-off transcription from the upstream gene. To test whether this is a common situation, we have performed a survey of all genes on human chromosome 22 with a distance between genes of ≤25 Kb. Of 546 genes annotated on this chromosome, 221 correspond to pairs that are within a distance of ≤25 Kb and with the same transcriptional direction. Five of these pairs correspond to overlapping
genes. Of the remaining 216 gene pairs, BLASTN searches of EST databases have identified two with transcripts matching both genes in the pair, that could correspond to transcripts bridging the two genes. For one of these two gene pairs, PNUTL1 and GP1BB, experimental evidence for the existence of hybrid transcripts, as well as single-gene transcripts, has been reported (Zieger et al. 1997; Yagi et al. 1998). The second gene pair with a potential hybrid transcript has not been characterized, and corresponds to genes predicted for a hypothetical protein (transcript dJ1194E15.3) and an EST cluster (dJ1104E15.5). The PNUTL1–GP1BB fusion transcript is predicted to contain two open reading frames (ORFs), and appears to result from defective truncation of the upstream PNUTL1 transcript due to an imperfect polyadenylation signal sequence (Zieger et al. 1997). Therefore, the approach used in this analysis can detect potential gene-bridging hybrid transcripts in 1% of the gene pairs analyzed. This is probably an underestimate of all instances of gene fusions expressing hybrid transcripts, because not all fused gene pairs will be represented by ESTs matching both genes in expression databases. Also, some of the annotated genes could have been misrepresented as single genes, especially if they have been predicted on the basis of matching ESTs. One conclusion of this type of analysis, relevant to the present study, is that transcript fusions between two adjacent genes, although not infrequent, are observed only in a subset of closely associated gene pairs.

In contrast to the Kua–UEV fusion, the PNUTL1–GP1BB gene fusion does not result in a fusion of proteins (Zieger et al. 1997). A second difference is that the mature PNUTL1–BP1BB fusion includes the terminal exon from the upstream gene, with its transcriptional truncation signal. Because this signal is apparently inefficient, the PNUTL1–BP1BB hybrid transcript could be the consequence of a genuine transcriptional run-off. In contrast, the mature processed transcripts of the Kua–UEV fusion have spliced out the terminal exon of the upstream gene, Kua, as well as the first exon of the downstream gene, UEV1A, which contains a 5′ untranslated sequence. RT–PCR experiments aimed at detecting fused transcripts that contain these two exons have failed to yield any amplification products. In the Kua–UEV fusion, therefore, a continuous primary transcript between both genes is subjected to specific splicing events that allow the expression of a two-domain protein. This also implies that a truncation of transcripts at the terminal exon of Kua must proceed at a rate that is sufficiently slow to allow the subsequent splicing events for the maturation of the Kua–UEV transcript. A carefully orchestrated balance between splicing and truncation of transcripts has been shown to occur in lower organisms (Ull et al. 1993).

A summary of the different classes of transcripts generated at the Kua–UEV locus is depicted in Figure 7. The locus for H. sapiens UEV1, on chromosome 20q13.2, contains two contiguous genes, Kua and, immediately downstream, UEV1. We predict that each gene has its own promoter, an initial exon with a 5′ UTR, and a terminal exon with a 3′ UTR. The genes Kua and UEV1 can be expressed as separate transcriptional units from their independent promoters, yielding, respectively, the proteins Kua and UEV1A. By means of alternative splicing, the transcript originating from the Kua promoter can generate a hybrid Kua–UEV transcript, with the potential to code for a two-domain Kua–UEV protein. Productive processing of the latter transcript to include the exons coding for the common domain of UEV requires removal of exon K6 by splicing, before truncation occurs, signalled by the polyadenylation signal present in that exon. For such a transcript to produce an uninterrupted reading frame yielding a two-domain Kua–UEV protein, exon A of UEV1 must be spliced out. In this scheme, the isoform UEV1B (or CROC1B), as originally described, corresponds to a partial, rather than a full-length, transcript and the B domain is a segment of Kua.

It has been shown that evolutionary fusion of proteins often indicates a functional interaction between them (Enright et al. 1999; Marcotte et al. 1999). In our hands, yeast two-hybrid experiments have not provided unequivocal evidence for direct interaction between Kua and UEV proteins from any of the organisms studied (N. Loukili and T.M. Thomson, unpubl.). Nevertheless, it remains possible that these two proteins perform functions in the same biochemical pathway, such that a requirement for functional interaction would provide a driving force for their evolutionary
fusión. One of the functional consequences of the fusion of Kua with UEV in humans is that the domain regulating polyubiquitination is not redirected to cytoplasmic structures, rather than the nucleus. As a consequence of their different subcellular localizations, nuclear (UEV1A) and cytoplasmic (Kua–UEV) forms of UEV1 could target different substrates for variant (K63) polyubiquitination. Thus, an endomembrane-associated form of UEV1 could preferentially direct the variant polyubiquitination of substrates closely associated with the cytoplasmic face of the ER, possibly, although not necessarily, in conjunction with membrane-bound ubiquitin-conjugating enzymes (Sommer and Jentsch 1993). The substrates could be either proteins specifically targeted for ubiquitination by these UEV–E2 complexes, or misfolded ER-associated proteins that are dislocated into the cytoplasm for subsequent ubiquitination (Kopito 1997).

METHODS

Biocomputational Analysis

Putative conserved coding regions between genomic sequences were identified with reciprocal TBLASTX (Altschul et al. 1990), the output processed with MSPCrunch (Sonhammer and Durbin 1994), and conserved segments visualized with apolot (http://www1.imim.es/~jabril/GFFTOOLS/APLOT.html). For prediction of protein coding genes, three different ab initio gene prediction programs were used, Genescan (Burge and Karlin 1997), Geneid (Gigué et al. 1992), and Fgenes (Solovyev et al. 1994). To support the resulting exonic structures, BLASTN searches were performed against the EST division of GenBank (Benson et al. 2000). A minimum subset of ESTs covering the matches was selected, all other EST matches being either identical or included therein. Each of these ESTs was aligned with the matching genomic clone sequence using the program est_genome (Mott 1997). Putative coding domains were recorded using the gff format (http://www.sanger.ac.uk/Software/formats/GFF/GFF_Spec.shtml), and plotted with gff2ps (available at http://www1.imim.es/~jabril/GFFTOOLS/GFF2PS.html). Protein sequences were aligned with ClustalW (Thompson et al. 1994), and secondary structures predicted with the PHD package (Rost 1996). Transmembrane domains were predicted with TMPred (http://www.ch.embnet.org/software/TMPRED_form.html). Searches for distant homologies were performed with PSI−BLAST (Altschul et al. 1997).

Expression Analysis by RT−PCR

RNAs from human cell lines and mouse tissues were isolated by the acid phenol procedure (Chomczynski and Sacchi 1987). RNAs from D. melanogaster embryos were isolated by guanidium isothiocyanate extraction and CsCl gradients, and enriched for mRNA on oligo dT-cellulose columns (Sambrook et al. 1989). RNAs were resuspended in diethyl pyrocarbonate-treated H2O and 20 µg were treated with 1 unit RNase-free RQI DNase I (Promega, Madison, WI), in a reaction containing 10 mM Tris·HCl at pH 8.0, 5 mM MgCl2, and 40 units RNasin (Promega). For Drosophila samples, RT−PCR was done by a single-tube procedure (Life Technologies, Barcelona), with forward primers 1 (5′-AATGACATCAACGAACGTC-3′), 2 (5′-CCTATGGTTGACCTTCCGGAT-3′), 3 (5′-CCTGTTGCAGCATATAACGGG-3′), 4 (5′-TGCCATAACATTTGCTGGCTA-3′), and reverse primer 5′-CGATGAGCTCGAATGGTTG-3′ (see Fig. 2G). For human samples, a two-step RT−PCR procedure was used, with reverse transcription of 1 µg RNA in a reaction containing 1× first-strand buffer, 200 µM dNTPs, 500 ng oligo-dT(12−18), and 200 units RNase H(−) RT (Life Technologies) at 42°C for 1 h. Aliquots were used as templates for hot-start PCR in 25-µl reactions containing 1× buffer, 130 µM dNTPs, 5 pmol of each primer and 0.2 units Taq polymerase (Ecogen, Barcelona). Amplification products were gel purified and aliquots used for nested or seminested PCR, and products sequenced from both strands by cycle sequencing and resolution in a ABI Prism 310 automatic sequence analyzer (Applied Biosystems). Forward primers used for RT−PCR and nested PCR were K1F (5′-TCTATGCGGCGTGTACCT-3′) or KF1n (5′-AGCTGGACAGGACGAG-3′) for exon 1 of human Kua, K2F (5′-CAGCGTCATCGGCCCCACCACCC-3′) for exon 2, and K4F (5′-ATGGCCCTCAAGTTCCGAC-3′), for exon 4. Reverse primers were KR (5′-GCCAGATGGCCCTCGGTTCGG-3′), for exon 6 of Kua, or UR (5′-CTAAGGGGAGAAGGCGAGA-3′), for exon C3 of UEV (see Fig. 3B). Amplification products were gel purified and sequenced as above. To determine relative levels of amplification of Kua−UEV and UEV transcripts, RT−PCR products were electrophoresed in ethidium bromide-containing agarose gels, and intensities (arbitrary units) determined for specific bands, normalized relative to the intensity of RT−PCR amplification products of the same RNAs with primers for the ribosomal protein gene S14r.

In Situ Hybridization on Drosophila Polytene Chromosomes

Drosophila polytene chromosome spreads were obtained from third instar wild-type larvae (Canton S strain) salivary glands. cDNA for D. melanogaster Kua was generated by RT−PCR with specific primers (primers 1 and reverse; see above and Fig. 2G), in single-tube RT−PCR reactions (Life Sciences) using as a template mRNA from D. melanogaster Kua embryos. cDNA for D. melanogaster UEV corresponded to the insert in IMAGE clone LD23138 (Research Genetics), cloned in pOT2 (plasmid vector). Different primer sets (primers 1 and reverse; see above and Fig. 2G), were gel purified and sequenced as above. To determine relative levels of amplification of Kua−UEV and UEV transcripts, RT−PCR reactions (Life Sciences) using as a template mRNA from Drosophila embryos. cDNA for D. melanogaster UEV corresponded to the insert in IMAGE clone LD23138 (Research Genetics), cloned in pOT2 (plasmid vector) and products sequenced from both strands by cycle sequencing and resolution in a ABI Prism 310 automatic sequence analyzer (Applied Biosystems). Forward primers used for RT−PCR and nested PCR were K1F (5′-TCTATGCGGCGTGTACCT-3′) or KF1n (5′-AGCTGGACAGGACGAG-3′) for exon 1 of human Kua, K2F (5′-CAGCGTCATCGGCCCCACCACCC-3′) for exon 2, and K4F (5′-ATGGCCCTCAAGTTCCGAC-3′), for exon 4. Reverse primers were KR (5′-GCCAGATGGCCCTCGGTTCGG-3′), for exon 6 of Kua, or UR (5′-CTAAGGGGAGAAGGCGAGA-3′), for exon C3 of UEV (see Fig. 3B). Amplification products were gel purified and sequenced as above. To determine relative levels of amplification of Kua−UEV and UEV transcripts, RT−PCR products were electrophoresed in ethidium bromide-containing agarose gels, and intensities (arbitrary units) determined for specific bands, normalized relative to the intensity of RT−PCR amplification products of the same RNAs with primers for the ribosomal protein gene S14r.

Expression Constructs and Transient Transfection Experiments

Full-length Kua, Kua−UEV, and UEV1A were amplified by RT−PCR using as a template total RNA from the cell lines HT-29 or Jurkat, and Expand High Fidelity polymerase (Boehringer-Mannheim), and the products subcloned in pGEM-T (Promega). The resulting inserts were amplified with primers for subcloning into pgEM112HA, bearing sequences coding for the hemaglutinating epitope, such that this sequence was placed in-frame at the carboxyl termini of the cDNAs. The resulting HA-tagged cDNAs were subcloned into pcDNA3.1 (InVitrogen). For transient transfection, 1 µg of plasmid DNA was transfected with Lipofectamine Plus (Life Sciences) into
COS-7 cells grown on glass coverslips. As a control, pcDNA3.1 vector DNA was used. Twenty-four hours after transfection, cells were washed, fixed in 4% paraformaldehyde/PBS, and permeabilized with 1% saponin/2% BSA/PBS. Cells were incubated with rat monoclonal anti-HA antibody (Boehringer-Mannheim) for 2 h, washed, and further incubated for 1 h with FITC-conjugated goat anti-rat Ig (Dako), washed, mounted in Immuno-Fluore (ICN), and observed under a Leica confocal microscope (Wetzlar, Germany). Transfection efficiencies ranged from 10%–15%.

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