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The Caenorhabditis elegans unc-32 Gene Encodes Alternative Forms of a Vacular ATPase a Subunit*

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Eukaryotes possess multiple isoforms of the a subunit of the V_o complex of vacular-type H^+-ATPases (V-ATPases). Mutations in the V-ATPase a^5 isoform have recently been shown to result in osteopetrosis, a fatal disease in humans, but no function has yet been ascribed to other isoforms. In Caenorhabditis elegans, the unc-32 mutant was originally isolated on the basis of its movement defect. We have isolated four new mutant alleles, the strongest of which is embryonic lethal. We show here that unc-32 corresponds to one of the four genes encoding a V-ATPase a subunit in the nematode, and we present their expression patterns and a molecular analysis of the gene family. unc-32 gives rise via alternative splicing to at least six transcripts. In the uncoordinated alleles, the transcript unc-32 B is affected, suggesting that it encodes an isoform that is targeted to synaptic vesicles of cholinergic neurons, where it would control neurotransmitter uptake or release. Other isoforms expressed widely during embryogenesis are mutated in the lethal alleles and would be involved in other acidic organelles. Our results indicate that V-ATPase a subunit genes are highly regulated and have tissue-specific function.

Vacuolar-type H^+-ATPases (V-ATPases) are ATP-dependent proton pumps involved in the acidification of intracellular compartments in eukaryotic cells and are important for many facets of cellular and organismal function (for review, see Refs. 1–3). As well as contributing to fundamental cellular processes

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†† The abbreviations used are: V-ATPase, vacuolar-type H^+-ATPase; bp, base pair; PCR, polymerase chain reaction; RT-PCR, reverse transcription-polymerase chain reaction; SL, spliced leader; GFP, green fluorescent protein; Unc, uncoordinated.

in the majority of eukaryotic cells, they also play specialized functions in certain cell types. In macrophages, the progressive acidification of phagosomes is an integral part of their maturation into lysosomes, whereas the acidification of early endocytic compartments is necessary for the recycling to the plasma membrane of transport molecules such as the transferrin receptor. In neurons, neurotransmitter trafficking is dependent on V-ATPases. The H^+-driven antiporters responsible for synaptic vesicle import of a specific neurotransmitter, such as the vesicular acetylcholine transporter, are known to be driven by proton gradients established by a V-ATPase localized to the same vesicle (4, 5).

The V-ATPases are composed of two complexes, V_1 and V_o, that serve different functions (for review, see Refs. 6, 7). V_1, a complex of 570 kDa composed of eight subunits (A–H), is located on the cytoplasmic side of the vesicle membrane and is responsible for ATP hydrolysis. V_o is a 260-kDa complex with five different subunits (a, d, c, c’, and c”) of 17–100 kDa. It is an integral membrane complex and is responsible for proton translocation from the cytoplasm to the lumen of the vesicle. In essence, the V-ATPases resemble, both in terms of sequence and structure, the ATP synthases of mitochondria. ATP synthases, however, couple the synthesis of ATP to the flow of protons from the mitochondrial matrix to the cytoplasm, and the overall topology of its two components is inverted, with the F_1 complex (analogous to the V_1 complex) being on the luminal side. Interestingly, there is no clear homologue for the V_o 100-kDa a subunit within the analogous F_0 complex, suggesting that it plays a specific role in the function of the V-ATPases. In yeast, there are two a subunit genes that encode proteins that have distinct subcellular localization (8). One, VPH1p, is associated with the central vacuole, whereas STV1p is thought to be a Golgi/endosome-specific form. Thus the two a subunits may be involved in targeting yeast V-ATPases to different intracellular membranes. Three murine a subunit genes have been characterized (a1, a2, and a3; Ref. 9). For a1 and a3, there are two and three alternative transcripts, respectively, giving the possibility of substantial variation in the structure of the murine V-ATPase. Intriguingly, the level of expression of the different transcripts of the murine genes varies among different tissues (9). This suggests that the V-ATPase complexes may have specialized compositions and structures, not only at the intracellular level but also in different tissues. The recent characterization of knockout mice for the a3 gene (ATP6i), which exhibit severe osteopetrosis due to loss of osteoclast-mediated extracellular acidification but no general defect in V-ATPase function, supports such a hypothesis (10). More recently, mutations in the human orthologue have been shown to

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be responsible for a subset of human osteopetrosis (11, 12).

Here, we describe the genetic and molecular identification of the Caenorhabditis elegans unc-32 gene, which encodes a set of V-ATPase 100-kDa a subunits. We show that the gene is one of a family of four V-ATPase a subunits encoding genes expressed in C. elegans. Furthermore, we show that a number of alternative unc-32 transcripts are produced by differential splicing of alternative exons and have characterized five mutants, two of which affect specifically one isoform. On the basis of our phenotypic and molecular analysis of the mutants, we postulate that specific isoforms may be involved in nervous system function, whereas others are required for viability.

EXPERIMENTAL PROCEDURES

Strains and Mutagenesis—All strains including the wild-type N2 Bristol and strains containing the unc-32 alleles and the transformants were grown and maintained at 20 °C as described (13). Four new ethyl methanesulfonate-induced alleles, unc-32(f120), unc-32(f121), unc-32(f123), and unc-32(f131), were isolated in two noncomplementation screens for new unc-32 alleles. The three former were induced on a dpy-17(e164) ncl-1(e1865) chromosome, the latter on an unc-36(e251) chromosome. Several thousand mutagenized genomes were tested against unc-32(e189) and scored for an uncoordinated phenotype. The new alleles were back-crossed at least four times to wild type, and genetic analysis was conducted using standard procedures. The two alleles, f121 and f123, were homozygous lethal and were kept heterozygous with the balancer qC1, dpy-19 glp-1, provided by J. Kimble. The integrated transformant els16(ZK637, unc-32+) was kindly provided by J. Sulston. The nonsense suppressor sup-7(st5) (14) as well as the smg-2(e808) suppressor involved in mRNA surveillance (15, 16) was combined with all alleles but f131.

Plasmid Constructions—All the plasmids used in this work were derived from the A1E5 plasmid (a gift from M. Cranton). A1E5 is a partial Sau3AI restriction fragment of the ZK637 cosmid (GenBank accession number Z11115), from 19434 to 32570, cloned in BlueScribe. E5-NS was obtained after Nhel and SpeI restriction and religation. E5-BK was constructed by a partial BamHI restriction to keep the BamHI site of the large exon of ZK637. E5-Pac corresponds to a deletion between the two PacI restriction sites of A1E5. E5.lin-9 was derived from E5-BK by using SauI and Sall restriction, T4 DNA polymerase treatment, and religation. E5-BK*, introducing a frameshift in exon 6, was derived from E5-BK by BamHI restriction, Klenow treatment, and religation. The green fluorescent protein (GFP) fusion in exon 4b, unc-32(e189)-GFP, was constructed by introduction of a 3600 bp SmaI-PstI restriction fragment from E5-NS in the HinDIII (blunted)-PstI-digested plasmid pPD95.81 (a gift from A. Fire).

Transgenic Lines—Plasmids for injection were prepared by a standard alkaline lysis procedure, followed by LiCl precipitation. Injections were performed as described (17). Subclones of ZK637 were injected at a concentration of 10 ng/ul with plasmid pRF4 (encoding rol6(su106), which confers a dominant rolling phenotype) at a concentration of 100 ng/ul. Some of the transformants also received either plasmid pPD93.97 (encoding a GFP reporter gene driven by the myo-3 promoter) or pPD20.97 (encoding a LacZ reporter gene driven by the myo-2 promoter) at a concentration of 20 ng/ul to check the presence of co-injected DNA directly by screening GFP in body wall muscle or β-galactosidase in the pharynx by 5-bromo-4-chloro-3-indolyl-b-D-galactopyranoside staining (both pPD plasmids were gifts from A. Fire). Clones were injected into e189 or f131 homozygotes and in unc-32(e189 or f131 or f121) dpy-17(e164)1(qC1).

Transcript Analysis—Partial sequences from 15 cDNAs were aligned on the genome, and three clones were fully sequenced (yk15a11, yk244e11, and yk5d7). There was evidence for alternative splicing, and the alternative transcripts from the three clones were designated as unc-32 A, B, and C, respectively. The differential splicing patterns were also analyzed by restriction digestion and/or sequencing of specific reverse transcription-polymerase chain reaction (RT-PCR) products. Total RNA from mixed populations of wild-type worms was prepared as previously described (18), reverse transcribed, and subjected to PCR with various sets of primers as well as the transspliced leaders SL1 and SL2 (19, 20). Amplification conditions and primers are available on request. The RT-PCR products were analyzed by restriction using enzymes cutting the transcripts only once, in exon 3 (XhoI), exon 4a (EcoRI), exon 4b (PstI), exon 7a (PvuI), and exon 7b (DraI). Some RT-PCR products were sequenced on an ABI machine using dye deoxyterminators (Prism; PerkinElmer Life Sciences). The traces were analyzed using the assembly program, developed by J. Thierry-Mieg and M. Potdevin.

When referring to the ZK637 cosmid, exon positions are: exon 1, 21653–21816 (coding sequence starting at 21664); exon 2, 21898–22074; exon 3, 22182–22310; alternative exon 4a, 22360–22714; alternative exon 4b, 23003–23109; alternative exon 4c, 23547–23468; exon 5, 23629–23583; exon 6, 23893–25147; alternative exon 7a, 25269–25591; alternative exon 7b, 26117–26257; exon 8, 26370–26589; exon 9, 26754–26925, and exon 10, 27145–27250. Transcripts were transspliced with SL1 at position 21653. Alternative polyadenylation occurs at similar frequencies at positions 27671 (atypical poly(A) signal atgaa 16–11 bp before the poly(A)) or 27784 (atypical poly(A) signal atagaa 17–12 bp before the poly(A)). Note that the existence, extent, and alternative status of exons 4a, 4b, 4c, 7a, and 7b were wrongly predicted for ZK637 in the data bases.

In Situ and GFP Expression Analyses—In situ hybridization analyses were performed as previously described (21). Transgenic worms carrying the unc-32e4::GFP construct were observed with a confocal microscope; the image in Fig. 6G is the projection of several Z serial sections.

Mutant Analysis—The genomic DNA of four mutant alleles (all but f123) was amplified with Takara TqQPCR products covering the segments from −629 to 832, 394 to 1761, 1687 and 2162 to 3804, 2768 to 4622, and 4605 to 5851 (where position 1 is the A of the initiator ATG) were fully sequenced on both strands. RNA was extracted from the three alleles mutated in a splice junction, e189, f120, and f131, and subjected to RT-PCR analysis (see above) and sequencing.

Sequence Comparisons—The nematode VHA-5 and VHA-6 proteins correspond respectively to the predicted F35H10.4 (GenBank accession number AAA18162) and VW02B12.1 (GenBank accession number CA909527) proteins, encoded by the expressed YK6569 and YK6534 genes. The nematode VHA-7 protein, corresponding to the predicted C26H9A.1 protein (GenBank accession number T19492), encoded by the YK778 expressed gene, was edited by removing the first 291 amino acids and the next to last exon, which does not occur in 5 of 5 independent cDNA clones. The 5′ region removed does not correspond to any available cDNA, nor does it resemble sequence in any of the well characterized 100 kDa a subunit proteins. Comparisons between the available human genomic sequence (GenBank accession number AC902557) and the published cDNA sequence for the human a1 gene (GenBank accession number NM_005177) allowed the deduction of the intron-exon boundaries. The intron-exon boundaries for the human a3 gene (OC116) were published previously (22). Amino acid sequences were aligned with the Multiple Alignment Program (developed by X. Huang). Phylogenetic trees were produced using the full-length protein sequences by neighbor joining using Clustalw alignment in Phylip format and the Treeview program.

RESULTS

Three unc-32 Alleles Are Viable but Uncoordinated—The unc-32(e189) allele was isolated by Brenner (13) as part of the first genetic screen for C. elegans mutants. Homozygous mutants are healthy and of normal size but exhibit a ventral coiler phenotype.
phenotype during backward movement (Fig. 1). This defect is seen from the L2 stage onward. The only additional defect observed in e189 is the brood size, smaller by one-third when compared with wild type, probably because of a reduction in number of fertile sperm. We carried out a genetic noncomplementation screen and isolated two more recessive viable alleles, f120 and f131. Although f120 shows a similar phenotype to e189 and was subsequently found to correspond to an identical molecular lesion (see below), f131 presents a much stronger uncoordinated (Unc) phenotype. The L1 larvae were small, thin, transparent, and slow-growing and had a brood size half that of the wild type. From the interactions with the nDf17 deletion, fully uncovering the region, that lead to a stronger phenotype, i.e. a high frequency of lethals (Table I), and the molecular evidence presented below, we believe that these three uncoordinated alleles, e189, f120, and f131, are partial loss-of-function alleles.

Two unc-32 Alleles Are Lethal, and the Locus Is Complex—The two further alleles, f121 and f123, isolated in the screen are zygotic lethal. f123 is equivalent to the deficiency in all interallelic combinations and therefore behaves as a null (Table I). In addition, f131 animals were small, thin, transparent, and slow-growing and had a brood size half that of the wild type. From the interactions with the nDf17 deletion, fully uncovering the region, that lead to a stronger phenotype, i.e. a high frequency of lethals (Table I), and the molecular evidence presented below, we believe that these three uncoordinated alleles, e189, f120, and f131, are partial loss-of-function alleles.

TABLE I

| unc-32 alleles | e189 or f120 | f131 | f121 | f123 |
|----------------|--------------|------|------|------|
| f121           | Almost WT    | L2, L3, or L4 Let | Extreme Unc, all stages |
| f123           | 10% extreme Unc, 90% L1 Let | L1 Let | L1 Let | L1 Let |
| nDf17         | 10% extreme Unc, 90% L1 Let | L1 Let | L1 Let | L1 Let |
| unc-32+ Transformants | WT | L1 Let | Emb Let | Emb Let |
| sup-7 interactions | None | ND | None | Emb Let |
| Nature of allele | Hypomorph | Hypomorph | Hypomorph | Hypomorph |
| Molecular lesion | g1393a | g4345a | g2874a | Gly(481/465/470) Glu |

* Transgenic worms with constructs containing the wild-type unc-32 locus. The Unc and Let phenotypes are rescued, but for the f121 and f123 alleles, as well as for f131 to a lesser extent, Mel and/or Ste (abnormal gametogenesis) phenotypes are revealed.

* sup-7 can suppress the embryonic lethality of f123, suggesting that this allele contains a premature stop codon.

* Positions of the molecular lesion, identified by sequencing the different alleles (position 1 is on the A of the initiator ATG); the position of the Gly amino acid mutated in the f121 allele is given for the different proteins containing, respectively, exons 4a, 4b, and 4c.

by each of the three Unc alleles, and conversely the Unc phenotype of these three alleles is partly rescued by f121. Such interallelic complementation defines unc-32 as a complex locus, a property often associated with alternative splicing.

unc-32 Encodes a 100-kDa a Subunit of V-ATPase—Genetic mapping had previously placed unc-32 between sma-2 and lin-12 in the center of LG III (13). Transformation rescue of unc-32(e189) by microinjection of cosmids from the candidate region of the physical map showed that the gene is contained within both ZK637 and T05B2 (Fig. 2A). We further refined the position of the gene by transformation rescue to a region of 9 kilobases (E5-BK, E5-NS, and E5-ln-9 constructs; see Fig. 2B), corresponding to the predicted gene ZK637.8. This gene encodes a protein similar in sequence to V-ATPase 100 kDa a subunits. Constructions in which part of ZK637 was deleted or in which a premature stop codon was introduced into the coding sequence of the gene failed to rescue the uncoordinated phenotype (E5-ΔPac and E5-BK* constructs; see Fig. 2B). Although a rescue of the zygotic lethality of f123 and f21 was obtained, the transformed worms were sterile (Table I and Fig. 2), possibly because of an absence of germ line expression of the transgenes as previously reported (23). Occasionally, the transforming plasmids rescued the various phenotypes, Unc, lethal, maternal effect lethal, and defect in gametogenesis, in a complex pattern. A particular transgenic array could rescue the Unc but not the lethal phenotype, whereas another, containing the same constructs, would do the opposite. This again presumably reflects the complexity of the unc-32 locus (see below).

Sequencing of the ZK637.8 genomic region from four mutant alleles revealed different molecular alterations: e189 and f120 contain a point mutation in the splice acceptor site of exon 4b, f131 in the splice donor site of exon 6 (see below), and f121 is a G to A transition in exon 6 (Table I). Together, these results confirm the identity of unc-32 as ZK637.8. The gene is the first in an operon containing two others, ZK637.9 and ZK637.10. The former encodes a protein similar in sequence to thiamine
Functions of the unc-32 V-ATPases in \textit{C. elegans}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure2.png}
\caption{Characterization of the \textit{unc-32} gene. \textbf{A}, Genetic and physical maps of the \textit{unc-32} region. The cosmids ZK637 and T05B2 were shown by J. Sulston to rescue the uncoordinated phenotype of \textit{unc-32(e189)} mutants. \textbf{B}, top, predicted and corrected genes in the indicated part of cosmids ZK637, with invariant exons indicated by open boxes; alternative exons in ZK637.8 are gray. The direction of transcription of ZK637.7, which corresponds to \textit{lin-9} (35), and of ZK637.11 is from right to left. For the other genes, it is from left to right. The positions of the point mutations of these \textit{unc-32} alleles are shown with asterisks: e189 is a G-A transition in the acceptor splice site in exon 4b; f131 is a G-A transition in the donor splice site in exon 6; and f121 is a G-A transition in exon 6. Bottom, insertions of the different clones tested for their ability to rescue the different \textit{unc-32} phenotypes. \textit{Let}, lethal; \textit{Mel}, maternal effect lethal. Transformation rescue experiments were performed in either e189, f131 or the lethal alleles, and only a summary of the rescue is presented. For example, the AE5 construct rescued the movement and fertility defects of f131 mutants but was associated with the appearance of an f123-like maternal effect lethality such that 85% of the eggs were dead. The clone E5\textsubscript{Δ}Pac contains a deletion corresponding to the major part of ZK637.8. The clone E5-BK\textsuperscript{*} harbors a point mutation resulting in a frame shift in the sixth exon and consequently a premature stop codon, the position of which is marked by an asterisk.}
\end{figure}

pyrophosphokinase (24). The latter encodes a putative mitochondrial thioredoxin reductase. By RT-PCR, we showed that these three genes are expressed, and that although the transcript of ZK637.8 is trans-spliced to SL1, both ZK637.9 and ZK637.10 can be trans-spliced to SL2 (data not shown; Ref. 25), as is normal for the case for genes in \textit{C. elegans} operons (26). Although in many operons the genes are functionally related, no evident link is apparent so far between the three genes of the \textit{unc-32} operon.

\textit{unc-32} Transcripts Are Alternatively Spliced—Sequencing of existing cDNAs and specific RT-PCR products revealed a number of alternative splice patterns for the \textit{unc-32} transcripts (Fig. 3, A and B). All \textit{unc-32} transcripts analyzed contain 10 exons. There are at least six different transcripts, using three variants for exon 4 (a–c) and two for exon 7 (a and b). Although the alternative exons 4a–c do not encode peptides that closely resemble sequences found in other V-ATPases, they occur at a position in the sequence where an alternative splicing pattern has also been observed for the murine \textit{a} gene (9), suggesting that they may have an important functional role. Significantly, the mutation in \textit{unc-32}(e189) is a G to A transition in the consensus splice acceptor site of exon 4b, meaning that in this allele, only the alternative transcripts with exon 4b should be affected. Consistent with this, anomalous \textit{unc-32} transcripts were found by RT-PCR amplification of the fourth exon region of total RNA extracted from \textit{unc-32(e189)} homozygotes (Fig. 3C). Cloning and sequencing of the forms that contain exon 4b in \textit{e189} homozygotes revealed the addition of 22 bp upstream of the exon 4b because of the use of a new acceptor splicing site. The resulting anomalous transcript contains a premature stop codon and would encode a nonfunctional protein. The other alternative forms containing exons 4a and 4c are still produced (Fig. 3C).

The alternative seventh exons, 7a and 7h, are similar in sequence and presumably arose from a recent duplication, because a single exon of similar sequence is present in other species (Fig. 3A). The \textit{f131} allele contains a G to A transition at the consensus splice donor site of the sixth exon, which affects the splicing of these alternative seventh exons (Fig. 3C). The large anomalous \textit{unc-32} transcript found by RT-PCR amplification of the seventh exon region of total RNA extracted from \textit{unc-32(f131)} homozygotes was sequenced. This revealed the presence of intron 6 between exon 6 and exon 7a. Because a stop codon is present in the sixth intron, the resulting mutant protein would be truncated in the eighth transmembrane domain. The other band with a size similar to that of wild type (Fig. 3C) corresponds to a mixture of alternate transcripts using cryptic donor sites. Because this allele is a hypomorph, not a null, we expect that some of these transcripts can be translated into a near full-length product that would differ from the wild type only in the relatively poorly conserved region around the exon 6-exon 7 junction.

\textit{V-ATPase} a Subunits Are a Large and Highly Conserved Family—\textit{UNC-32} belongs to a family of highly conserved \textit{V-ATPase} a subunit proteins. In other species there are multiple genes encoding \textit{V-ATPases} a subunits. There are three genes in mice (9) and other vertebrates, and searches in the available \textit{Drosophila} genomic sequence indicate that there are at least four members in the fly. In \textit{C. elegans}, in addition to \textit{UNC-32}, there are three other members of the family, named \textit{VHA-5}, \textit{VHA-6}, and \textit{VHA-7}. Each of the four genes has a distinctive intron-exon structure. We could not detect alternative splicing of these genes, as judged by the analysis of several cDNAs. The existence of shared intron-exon boundaries among the four nematode genes and the two human genes for which data are available supports their common ancestry (Fig. 4A). Each of these putative proteins contain nine potential membrane-spanning regions (Ref. 27 and Fig. 4B). The sequence identity (and

\begin{itemize}
\item GenBank™ accession numbers AAD34751, AAD34771, AAF53116, and AAF55550.
\item D. Thierry-Mieg and Y. Kohara, unpublished results.
\end{itemize}
FIG. 3. The unc-32 gene encodes several isoforms of V-ATPase α subunit. The different unc-32 transcripts were determined by sequencing different cDNA clones and RT-PCR products, and the respective proteins are named as follows: UNC-32 A–F contain, respectively, exons 4a/7b, 4b/7a, 4c/7b, 4a/7a, 4c/7a, and 4b/7b. Their GenBank accession numbers are, respectively, AF320899, AF320900, AF320901, AF320902, AF320903, and AF320904. Their predicted molecular masses, in kilodaltons, are, respectively, 103.4, 100.7, 102, 102.8, 101.3, and 101.1. Note that the different UNC-32 proteins differ from both predicted ZK637.8a and 8b proteins. A, UNC-32 A, F, and E protein sequences. Apart from the alternative exons (of variable size, hence marked with dashes), the sequences are identical and are not shown for the sake of clarity. Although the fourth exons are not similar, exons 7b and 7a in the variants F and E, respectively, are highly similar. Residues that are identical are black; those that are similar are gray. The vertical lines above the sequences mark the corresponding exon-intron boundaries. B, exon-intron structure of the six different alternative splice variants. The primers used for the RT-PCR presented in C are also shown. Alternative exons are gray. The relative abundances of different transcripts were evaluated after digestion of RT-PCR products made from mRNA from mixed stage wild-type worms and are shown as percentages. C, different patterns of mRNA splicing are revealed by RT-PCR in the unc-32 mutants, and their sequences are presented. Top panel, analysis of the e189 mutation. On the left, the amplicon containing the fourth exon was amplified with the primers 8.11 and 8.2 and digested with HaeIII. Transcripts containing exon 4a are cut in three fragments (466, 407, and 49 bp; the 49-bp band is not visible in this image). The e189 mutation removes a HaeIII site, which is normally formed by the splicing of exon 3 to 4b, and results in the appearance of a larger band (arrow) and the loss of the 49-bp band (not shown). The band at 2375 bp results from direct PCR amplification of genomic DNA, shown in lane gwt. On the right are presented the sequences of both wild-type and e189 transcripts. The e189 mutation removes the exon 4b acceptor splicing site; a new acceptor splicing site, located 22 bp upstream of the normal one, is used, introducing a stop codon. Bottom panel, analysis of the f131 mutation. On the left, the amplicon including the seventh exon was amplified with primers 8.5 and 8.6. Transcripts containing exon 7a (983 bp) or 7b (1001 bp) are not cut, whereas transcripts containing exon 4b are cut in three fragments (466, 407, and 49 bp; the 49-bp band is not visible in this image). The f131 mutation removes a HaeIII site, which is normally formed by the splicing of exon 6 to 7a, and results in the appearance of a larger band (arrow). The band at 2465 bp results from direct PCR amplification of genomic DNA, shown in lane gwt. On the right are presented the sequences of both wild-type and f131 transcripts. The f131 mutation removes the exon 6 donor splicing site; therefore, intron 6 is not spliced out and fuses to exon 7a, resulting in the introduction of stop codons. The bottom band is a mixture of various transcripts using cryptic splice sites.
similarity) between UNC-32A and other nematode and murine a subunit proteins are as follows: VHA-5, 50% (67%); VHA-6, 50% (65%); VHA-7, 46% (62%); a1, 55% (69%); a2, 43% (60%); and a3, 37% (55%). Phylogenetic analysis of the different members of this family indicates, in agreement with previous suggestions based on a more limited analysis (9), that the multiple genes appeared independently in vertebrates and invertebrates subsequent to their divergence from a common ancestor (Fig. 5). Interestingly, although the UNC-32 protein is closer in sequence to the a1 protein than to the other nematode proteins, the vha-7 gene shares more intron-exon boundaries with the human a1 and a3 genes than other nematode genes.

V-ATPase a Subunits Have Specific Expression Domains—The expression pattern of each of the four V-ATPase a genes in C. elegans was analyzed by in situ hybridization. Two of the genes, vha-6 and vha-7, have very limited and nonoverlapping domains of expression, whereas the two others, unc-32 and vha-5, are expressed more broadly, yet each with a distinctive pattern (Fig. 6). Transcripts for unc-32 were detected at high levels in early embryos (Fig. 6A) from the one-cell stage to the end of gastrulation. In larvae and adults, transcripts were...
clearly present in the gonad (Fig. 6, C and D), in the intestine, in many neurons in the head, and in motoneurons in the ventral cord (Fig. 6D). This expression pattern is consistent with the expression of a LacZ reporter gene driven by the upstream region of the unc-32 gene (28). Similarly, vha-5 showed precocious and widespread embryonic expression (Fig. 6E), followed by a more restricted pattern in larvae and adults. mRNAs are present in many tissues where unc-32 is also expressed, but the detailed patterns were different, because the gene is also expressed in the excretory cell and canals (Fig. 6F) and probably some somatic gonad cells (Fig. 6, G and H). In contrast, expression of vha-6 started late, in the intestine of the 2-fold embryo (Fig. 6I). It was strong but limited to the intestinal cells at all stages (Fig. 6, J–L). The vha-7 gene was silent during embryogenesis and the first larval stages (Fig. 6M). Its expression was first observed at the L3 stage and was limited to areas of the developing and mature gonad corresponding to the spermatheca (Fig. 6N) and the region where meiotic oocytes are in the pachytene and early diakinesis stages. It vanished abruptly in mature oocytes (Fig. 6, O and P). Although the patterns of expression of these other genes are suggestive, their functions remain to be investigated.

unc-32 Transcripts Containing Exon 4b Are Specifically Expressed in the Nervous System—In situ hybridization with the unc-32 gene revealed a large domain of expression, especially in the embryo where the expression is ubiquitous. To test the hypothesis that unc-32 alternative transcripts have tissue-specific expression patterns, we made a fusion construct, unc-32e4::GFP, in which GFP was inserted into the alternative exon 4b. GFP expression in wild-type strains carrying this construct was restricted to the nervous system (Fig. 6Q), including all the motoneurons in the ventral cord, the dorsal, the lateral, and all the sublateral cords, PLML/R, PVM, ALML/R, AVM, ALNL/R, PLNL/R, PVDL/R, PDEL/R, SDQL/R, and almost all the neurons in the head.

**DISCUSSION**

*unc-32(e189)* Is Affected in the Motor Circuit—The analysis of the uncoordinated phenotype of unc-32(e189) suggested a possible defect in specific classes of motoneurons (29). Serial reconstruction of the anterior part of the ventral nerve cord behind the retroversicular ganglion in the mutant, however, failed to show any wiring defect in the motoneurons. They revealed instead that synaptic vesicles had an abnormal and distinctive morphology compared with wild type. This strongly suggests that the function rather than the structure of the motor circuit is compromised in unc-32(e189) mutants. Consistent with this, mutant animals are resistant, at a low but consistent level, to inhibitors of acetylcholine esterase, as are mutants in other genes involved in acetylcholine synthesis, packaging, release, or signal reception (30). We have unambiguously identified unc-32 as being a nematode V-ATPase a subunit and show that unc-32 is expressed in some motoneurons and is thus likely to be directly responsible for the uncoordinated phenotype. We propose that in the unc-32(e189) mutant, V-ATPases are not functional in motoneurons, leading to a lack of import of the acetylcholine neurotransmitter into synaptic vesicles, a function that as been previously suggested for V-ATPase (4, 5).

**Alternative Splice Forms Are Expressed in Motoneurons**—There are at least six alternative transcripts for the unc-32 gene. The unc-32(e189) mutation affects the generation of two of the six transcripts, namely those containing exon 4b. In other terms, in the e189 allele, there is no transcript with exon 4b, or if a transcript contains exon 4b, it is nonfunctional. One hypothesis was that the isoforms with exon 4b are required for synaptic vesicle function in motoneurons but are dispensable in other tissues. Alternatively, motoneurons could force the splicing apparatus to always include exon 4b, thus generating a null allele in this cell type. The fact that a GFP fusion in exon 4b is specifically expressed in the nervous system favors the latter hypothesis.

*Other Alternative Splice Forms Are Required during Development*—The unc-32(f121) and the null unc-32(f123) alleles are lethals, so certain UNC-32 isoforms must be absolutely required for viability. Because unc-32(f121) corresponds to a point mutation in an exon common to all isoforms and will affect all alternative unc-32 transcripts, it remains to be determined which UNC-32 isoforms have an essential function. The lethality of those alleles is, however, consistent with the fact that unc-32 is widely expressed in early embryogenesis. Rescue of the e189 uncoordinated phenotype by the f121 lethal allele suggests that at least one of the f121 mutant forms is partially functional in the synaptic vesicles of the motor circuit. The f121 mutation results in a Glu for Gly substitution in the third transmembrane domain at a position that is not fully conserved between species. In motoneurons, this substitution in transcripts containing exon 4b might not perturb the function of the V-ATPase.

Our results have revealed that the function and regulation of unc-32 are complex. First, there are at least six alternative transcripts for this gene. Second, the three uncoordinated alleles are affected in the generation of these alternative transcripts. Third, the different mutant alleles display distinct phenotypes and unexpected interallelic complementation. These could be due to perturbations of an equilibrium between...
alternatively spliced forms of unc-32, each of which would encode a protein with a specific function. The smg-2 suppressor is involved in mRNA surveillance (15) and has been shown to modulate the level of alternatively spliced mRNAs (16). Significantly, the lethality of J21 was rescued by smg-2, but unc-32(f121); smg-2(e2008) adults did not lay eggs (data not shown).

Four V-ATPase a Subunits in the Nematode—The unc-32 gene corresponds to one of four nematode V-ATPase a subunits. Just like in vertebrates, where different a subunit isoforms are targeted to different tissues (9, 31), each of the C. elegans V-ATPase a genes has a specific, largely nonoverlapping, domain of expression. Tissue-specific expression of alternatively spliced forms of the murine a1 gene has also been reported (9). Furthermore, in yeast, the two a subunit isoforms have been implicated in the differential intracellular localization of the V-ATPase complex (8). This suggests that different isoforms of the a subunit, arising from alternative splicing or the expression of different genes, or both, have specific patterns of expression, both in terms of tissue distribution and targeting to individual intracellular membrane compartments. In the nematode, further variation of the structure of the V-ATPase is possible, because three c subunit genes (vha-1, -2, and -3) have been described (32–34). Given their conservation through evolution, future studies using C. elegans may allow a deeper understanding of the enormous potential for variation in the structure, function, and regulation of V-ATPases and their association with different pathologies in vertebrates (11).

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