The C. elegans Cell Death Gene ced-3 Encodes a Protein Similar to Mammalian Interleukin-1β-Converting Enzyme

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Summary

We have cloned the C. elegans cell death gene ced-3. A ced-3 transcript is most abundant during embryogenesis, the stage during which most programmed cell deaths occur. The predicted CED-3 protein shows similarity to human and murine interleukin-1β-converting enzyme and to the product of the mouse nedd-2 gene, which is expressed in the embryonic brain. The sequences of 12 ced-3 mutations as well as the sequences of ced-3 genes from two related nematode species identify sites of potential functional importance. We propose that the CED-3 protein acts as a cysteine protease in the initiation of programmed cell death in C. elegans and that cysteine proteases also function in programmed cell death in mammals.

Introduction

Cell death occurs as a normal aspect of animal development as well as in tissue homeostasis and aging (Glucksmann, 1950; Ellis et al., 1991a). Naturally occurring or programmed cell death can act to regulate cell number, to facilitate morphogenesis, to remove harmful or otherwise abnormal cells, and to eliminate cells that have already performed their functions. In many cases, gene expression within dying cells is thought to be required for these cells to die, since the cell death process can be blocked by inhibitors of RNA and protein synthesis (Stanisic et al., 1978; Cohen and Duke, 1984; Martin et al., 1988).

During the development of the nematode Caenorhabditis elegans, 131 cells undergo programmed cell death (Sulston and Horvitz, 1977; Sulston et al., 1983). Fourteen genes have been identified that function in different steps of the genetic pathway of programmed cell death in C. elegans (Hedgecock et al., 1983; Ellis and Horvitz, 1988, 1991; Ellis et al., 1991b; Hengartner et al., 1992; reviewed by Ellis et al., 1991a). Two of these genes, ced-3 and ced-4, play essential roles in either the initiation or execution of the cell death program, since recessive mutations in these genes prevent almost all of the cell deaths that normally occur during C. elegans development. Genetic mosaic analysis indicates that ced-3 and ced-4 most likely function within cells that die or within their close relatives to cause cell death (Yuan and Horvitz, 1990). The ced-4 gene encodes a novel protein that is expressed primarily during embryogenesis, the period during which most programmed cell deaths occur (Yuan and Horvitz, 1992).

To understand how the ced-3 gene acts to cause cell death, we have cloned this gene. As deduced from the sequence of a ced-3 cDNA clone, the CED-3 protein is 503 amino acids in length and contains a serine-rich middle region of about 100 amino acids. We compared the sequences of the CED-3 protein of C. elegans with the inferred CED-3 protein sequences from the related nematode species C. briggsae and C. vulgaria. This comparison revealed that the carboxy-terminal portions of these proteins are most conserved. The non-serine-rich portions of the CED-3 protein are similar to human interleukin-1β (IL-1β)-converting enzyme (ICE), a cysteine protease that can cleave the inactive 31 kd precursor of IL-1β to generate the active cytokine (Cerretti et al., 1992; Thornberry et al., 1992). In addition, the carboxy-terminal portions of both the CED-3 and ICE proteins are similar to the mouse Nedd-2 protein, which is encoded by a messenger RNA (mRNA) expressed during mouse embryonic brain development and is down-regulated in adult brain (Kumar et al., 1992). We suggest that CED-3 acts as a cysteine protease in controlling the onset of programmed cell death in C. elegans and that members of the ced-3/ICE/Nedd-2 gene family might function in programmed cell death in vertebrates.

Results
ced-3 Is Not Essential for Viability

All previously described ced-3 alleles were isolated in screens designed to detect viable mutants in which programmed cell death did not occur (Ellis and Horvitz, 1986). Such screens might systematically have missed classes of ced-3 mutations that result in inviability. Since animals of the genotype ced-3 deficiency are viable (Ellis and Horvitz, 1986), we designed a screen that would allow us to isolate recessive lethal alleles of ced-3, if such alleles could exist (see Experimental Procedures). We obtained four new ced-3 alleles (n1163, n1164, n1165, and n1286) in this way. All four of these mutants are viable as homozygotes. These alleles were isolated at a frequency of about 1 in 2500 mutagenized haploid genomes, approximately the frequency expected for the generation of loss-of-function mutations in an average C. elegans gene (Brenner, 1974; Brenner and Horvitz, 1980).
ment that contained Tc1 (Emmons et al., 1983), that was present in the C. elegans Bristol strain N2 but not in the C. elegans Bergerac strain EM1002, and that was closely linked to ced-3. We named this restriction fragment length polymorphism (RFLP) nP35. Using Tc1 to probe Southern blots of cosmids derived from N2 genomic DNA and known to contain Tc1 (G. Ruvkun, personal communication), we identified two cosmids (MMM-C1 and MMM-C9) that contained this 5.1 kb EcoRI fragment. These cosmids overlapped an existing cosmid contig that had been defined as part of the C. elegans genome project (Coulson et al., 1986, 1988; A. J. Coulson et al., personal communication). We used cosmids from this contig to identify four additional Bristol-bergerac RFLPs (nP33, nP34, nP36, and nP37). By mapping these RFLPs between the Bristol and Bergerac strains with respect to the genes unc-30, ced-3, and unc-26, we oriented the contig with respect to the genetic map. These experiments narrowed the region containing the ced-3 gene to an interval spanned by the three cosmids C48D1, W07H6, and C43C9 (Figure 1A).

These three cosmids were microinjected (Fire, 1986; J. Sulston, personal communication) into ced-3 mutant animals to test for rescue of the mutant phenotype. Specifically, a candidate cosmid and cosmid C14G10, which contains the wild-type unc-37 gene (R. Hoskins, personal communication), were co-injected into ced-1(e1735); unc-31(e928) ced-3(n717) hermaphrodites, and non-Unc progeny were isolated and observed to see whether the non-Unc phenotype was transmitted to the next generation, thus establishing a line of transgenic animals. Young first larval stage (L1) progeny of such transgenic lines were examined for the presence of cell deaths using Nomarski optics to see whether the Ced-3 phenotype was rescued. Cosmid C14G10 alone does not confer wild-type ced-3 activity when injected into a ced-3 mutant (data not shown). ced-1 was used to facilitate scoring of the Ced-3 phenotype (see Experimental Procedures), and unc-31 was used as a marker for cotransformation (R. Hoskins, personal communication; Kim and Horvitz, 1990).

As indicated in Figure 1A, of the three cosmids tested, only C48D1 rescued the CED-3 mutant phenotype. Two non-Unc transgenic lines were obtained, the insertion line nls1 and the extrachromosomal line nEx2. Both were rescued. Specifically, L1 ced-1 animals contained an average of 23 cell corpses in the head, and L1 ced-1; ced-3 animals contained an average of 0.3 cell corpses in the head (Ellis and Horvitz, 1986). By contrast, L1 ced-1; unc-31 ced-3; nts1 and L1 ced-1; unc-31 ced-3; nEx2 animals contained an average of 16.4 (n = 20) and 14.5 (n = 20) cell corpses in the head, respectively. From these results, we concluded that C48D1 contains the ced-3 gene.

To locate ced-3 more precisely within cosmids C48D1, we subcloned this cosmid and tested the subclones for their abilities to rescue the CED-3 mutant phenotype (Figure 1B). From these experiments, we localized ced-3 to a DNA fragment of 7.5 kb (pJ7.5).

ced-3 Is Transcribed Primarily during Embryogenesis and Independently of ced-4 Function

We used the 7.6 kb pJ107 subclone of C48D1 (Figure 1B)
to probe a Northern blot of poly(A)\(^+\) RNA derived from the wild-type C. elegans strain N2. This probe hybridized to a 2.8 kb transcript (data not shown; also see Figure 2). Although this transcript was present in all 11 different ethyl methanesulfonate (EMS)-induced ced-3 mutants strains (data not shown; the mutant n1764 was not tested), subsequent analysis revealed that all 11 mutants contain mutations in the genomic DNA that encodes this mRNA (see below), thus establishing this RNA as a ced-3 transcript. The ced-3 transcript was most abundant during embryogenesis, when most programmed cell deaths occur (Sulston and Horvitz, 1977; Sulston et al., 1983), and was also detected at later stages (Figure 2).

Since ced-3 and ced-4 are both required for programmed cell death in C. elegans and since both are highly expressed during embryonic development (Yuan and Horvitz, 1992; see above), it is possible that one of these genes regulates the mRNA levels of the other. Previous studies showed that ced-3 does not regulate ced-4 mRNA levels (Yuan and Horvitz, 1992). To determine whether ced-4 regulates ced-3 mRNA levels, we probed a Northern blot of RNA prepared from ced-4 mutant embryos with the ced-3 cDNA subclone pJ178. The abundance and size of ced-3 transcript were normal in the ced-4 mutants n1762 (glutamine to ochre nonsense mutation at codon 40), n1416 (AC4 insertion into exon 5), n1894 (tryptophan to opal nonsense mutation at codon 401), and n1920 (G to A mutation in the intron 3 splice donor site) (Figure 3A). None of the ced-4 mutants tested has detectable ced-4 RNA (Figure 3B; Yuan and Horvitz, 1992). Thus, ced-4 does not seem to affect the steady-state levels of ced-3 mRNA.

### ced-3 cDNA and Genomic Sequences

To isolate ced-3 cDNA clones, we used the ced-3 genomic DNA clone pJ40 (see Figure 1B), which rescued the Ced-3 mutant phenotype, when microinjected into ced-3 mutant animals, as a probe to screen a cDNA library prepared from the C. elegans wild-type strain N2 (Kim and Horvitz, 1990). The 2.5 kb cDNA clone pJ87 was isolated in this way. On Northern blots, pJ87 hybridized to a 2.8 kb transcript, and on Southern blots, it hybridized only to bands to which pJ40 also hybridized (data not shown). Thus, pJ87 was derived from an mRNA transcribed entirely from pJ40. To confirm that pJ87 corresponds to a ced-3 cDNA clone, we made a frameshift mutation in the unique SalI site of pJ40, which corresponds to the unique SalI site in the pJ87 cDNA clone and disrupts the putative ced-3 open reading frame (see Experimental Procedures; see also Figure 4A). Constructs containing this frameshift mutation failed to rescue the C3d-3 phenotype when microinjected into ced-3 mutant animals (six transgenic lines were examined; data not shown), suggesting that ced-3 activity was eliminated by mutating the putative ced-3 open reading frame.

We determined the DNA sequence of pJ87 (Figure 4). pJ87 contains an insert of 2482 bp that can encode a protein of 503 amino acids. pJ87 contains 953 bp of 3'
untranslated sequence, not all of which is essential for ced-3 function, since genomic constructs that lack the last 380 bp of the 3'-most region (pJ107) and its derivatives; see Figure 1B) rescued the Ced-3 mutant phenotype. The pJ87 cDNA clone ends with a poly(A) sequence (data not shown), suggesting that the 3' end of pJ87 corresponds to the extreme 3' end of the ced-3 transcript. The 5' end of pJ87 does not contain trans-spliced sequences (Bektash et al., 1988; Huang and Hirsh, 1989) and therefore might or might not include the 5' end of the ced-3 transcript.

To determine the 5' end of the ced-3 transcript, we performed primer extension experiments (Figure 5). Two primers containing sequences separated by 177 bp in the genomic DNA sequence (see below) and by 123 bp in the cDNA sequence were used for the primer extension reactions: Pex1, starting at position 2305 of the genomic sequence, and Pex2, starting at position 2482 of the genomic sequence. The Pex2 reaction yielded two major products of 283 nt and 409 nt, whereas the Pex1 reaction gave one product of 160 nt. The 160 nt product of the Pex1 reaction corresponds to the 283 nt product of the Pex2 reaction, since these products differ in size by 123 nt. Products of these lengths are consistent with the presence of a ced-3 transcript that is trans-spliced to a 22 bp C. elegans splice acceptor (Bektash et al., 1988) at a consensus splice acceptor site at position 2166 of the genomic sequence.
Table 1. Sequences of ced-3 Mutations

<table>
<thead>
<tr>
<th>Allele</th>
<th>Wild-Type Sequence</th>
<th>Mutant Sequence</th>
<th>Substitution or Splice Site Change</th>
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</thead>
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<tr>
<td>n717</td>
<td>tttgca/CAA</td>
<td>tttgca/CAA</td>
<td>Exon 7 acceptor</td>
</tr>
<tr>
<td>n718</td>
<td>GGA</td>
<td>AGA</td>
<td>G65R</td>
</tr>
<tr>
<td>n1040</td>
<td>GCA</td>
<td>GTA</td>
<td>A448V</td>
</tr>
<tr>
<td>n1129.</td>
<td>GCA</td>
<td>GCA</td>
<td></td>
</tr>
<tr>
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<td>TCC</td>
<td>TTC</td>
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</tr>
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<td>Q404amber</td>
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<tr>
<td>n2433</td>
<td>GCC</td>
<td>AGC</td>
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</tr>
</tbody>
</table>

Amino acid positions correspond to the numbering in Figure 4A.

was cloned into the PCR1000 vector (Invitrogen, San Diego, California), and its sequence was determined (data not shown). This sequence confirmed that at least some ced-3 transcripts are trans-spliced to the C. elegans spliced leader SL1 at a consensus splice acceptor at position 2166 of the genomic sequence. Similar experiments using a primer containing the SL2 spliced leader sequence (Huang and Hirsh, 1988) failed to identify an SL2 spliced mRNA. Since the primer extension experiments identified a major ced-3 mRNA that probably is trans-spliced, it seems likely that a large proportion of ced-3 mRNA is SL1 trans-spliced. Based upon these observations, we propose that the translational start codon of ced-3 is the first in-frame ATG downstream of the SL1 splice acceptor site at position 2232 of the genomic sequence and that the CED-3 protein is 503 amino acids in length (see Figure 4A).

To define the structure of the ced-3 gene, we determined the genomic sequence of the ced-3 gene from the plasmid pJ107. The insert in pJ107 is 7535 bp in length (see Figure 4A). Comparison of the ced-3 genomic and cDNA sequences revealed that the ced-3 gene has seven introns that range in size from 84 bp to 1195 bp. The four largest introns, as well as sequences 5' of the start codon (see below), contain repetitive elements, some of which have been previously identified in noncoding regions of other C. elegans genes, such as fem-1 (Spence et al., 1990), in-12 (Yochem and Greenwald, 1989; J. Yochem and I. Greenwald, personal communication), gip-1 (Yochem and Greenwald, 1989), and htk-1 (Krause et al., 1990), as well as in the cosmids ZK643 and B0303 (Sulston et al., 1992) (see Figure 4B). Genomic sequence analysis of a ced-3 homolog from the related nematode C. briggsae (J. Y. and S. L., unpublished data) revealed that these repeats are not present in this nematode species, suggesting that the repeats do not have a role in regulating ced-3 expression. It is possible that such repeats represent active or inactive transposable elements.

The predicted CED-3 protein is hydrophilic (256 of 503 residues are charged or polar) and does not contain any obvious potential transmembrane domains. One region of
the CED-3 protein is rich in serines: from amino acid 107 to amino acid 205, 32 of 99 amino acids are serines.

We determined the sequences of 12 EMS-induced ced-3 mutations (see Figure 4A; Table 1). Eight are missense mutations, three are nonsense mutations, and one alters a conserved G at the presumptive splice acceptor site of intron 6.

To identify functionally important regions of the CED-3 protein, we cloned and determined the genomic sequences of the ced-3 genes from the related nematode species C. briggsae and C. vulgaris. Sequence comparisons showed that amino acids corresponding to residues 1–205 of the C. elegans CED-3 protein are less conserved among the three nematodes (68% identical) than are amino acids corresponding to residues 206–503 of the C. elegans CED-3 protein (84% identical) (see Figure 7A). All eight EMS-induced missense mutations in ced-3 (see above) altered residues that are conserved among the three species (see Figure 7A). Interestingly, six of these eight mutations alter residues within the last 100 amino acids of the protein, and none affects the serine-rich region (see Figure 4A; Figure 6). These results suggest that the carboxyl region is important for ced-3 function and that the serine-rich region might be unimportant or that different residues within it might be functionally redundant.

CED-3 Protein Is Similar to the Mammalian ICE and Nedd-2 Proteins

A search of the GenBank, PIR, and SWISS-PROT data bases revealed that the non-serine-rich regions of the CED-3 protein are similar to the human and murine ICEs (Figure 7A). ICE is a cysteine protease that cleaves the inactive 31 kd precursor of IL-1β between Asp-116 and Ala-117, releasing a carboxy-terminal 153 amino acid polypeptide known as mature IL-1β (Kostura et al., 1989; Black et al., 1989). The CED-3 proteins from the three Caenorhabditis species and the human ICE protein share 29% amino acid identity. The most highly conserved region consists of amino acids 246–360 of the CED-3 protein and

Figure 7. CED-3 Protein
(A) Comparison of the CED-3 protein sequences from C. elegans, C. briggsae, and C. vulgaris with the human and mouse ICEs and with the mouse Nedd-2 protein. Amino acids are numbered to the right of each sequence. Dashes indicate gaps in the sequence to allow optimal alignment. Residues that are identical among more than half of the proteins and between nematode and mammalian sequences are boxed. Missense ced-3 mutations are indicated above the comparison blocks, showing the residue in the mutant CED-3 protein and the allele name. Asterisks indicate potential aspartate-specific cleavage sites in the CED-3 protein. Circles indicate known aspartate-specific cleavage sites in human ICE. Residues indicated in boldface correspond to the highly conserved pentapeptide containing the active cysteine in ICE.
(B) Comparison of structural features of the CED-3 protein and human ICE. The predicted proteins corresponding to the ICE proenzyme and CED-3 are represented. The active site in ICE and the predicted active site in CED-3 are indicated by the closed rectangles. The four known cleavage sites in ICE flanking the processed ICE subunits (p24), which was detected in low quantities when ICE was purified (Thornberry et al., 1992), p20, and p10) and two conserved presumptive cleavage sites in the CED-3 protein are indicated with solid lines and linked with dotted lines. Five other potential cleavage sites in the CED-3 protein are indicated with dashed lines. The positions of the aspartate (D) residues at potential cleavage sites are indicated below each diagram. The carboxyl terminus of p24 has not been determined and is indicated by a dotted arrow.
amino acids 166–287 of the human ICE protein: 49 of 115 residues are identical (43% identity). Cys-285 is thought to be an essential component of the active site of ICE (Thornberry et al., 1992). The amino acid pentapeptide QACRG containing this active cysteine is the longest peptide conserved among the murine and human ICE proteins and the CED-3 proteins of the three nematode species.

Active human ICE is composed of two subunits (p20 and p10) that appear to be proteolytically cleaved from a single proenzyme by the mature enzyme (Thornberry et al., 1992). Four cleavage sites in the proenzyme have been defined. Only p20 and p10 are necessary for the in vitro enzymatic activity of ICE, suggesting that the three additional fragments resulting from ICE cleavage are not required for ICE function. Two of these cleavage sites (aspartic acid-serine dipeptides at positions 103–104 and 297–298 of ICE) are conserved in CED-3 (positions 131–132 and 371–372, respectively) (Figure 7).

The carboxy-terminal portion of the CED-3 protein and the p10 subunit of ICE are similar to the protein product of the murine gene nced-2 (Figure 7A), which is highly expressed during embryonic brain development and is down-regulated in adult brain (Kumar et al., 1992). The C. elegans CED-3 protein and the Nced-2 protein are 27% identical, as are the ICE and Nced-2 proteins (Figure 7A). The Nced-2 protein apparently does not contain the QACRG peptide found at the active site of ICE (Figure 7A). Six of the eight known ced-3 missense mutations (n718, n1040, n1129, n1164, n2426, and n2433) alter amino acids that are identical among the three nematode CED-3 proteins and human ICE. For example, the mutation n2433 introduces a glycine to serine substitution at an absolutely conserved glycine near the putative active site cysteine (see Figure 4A; Figure 7A). Four mutations (n1129, n1163, n1164, and n2426) alter amino acids that are identical among the nematode CED-3 proteins and the Nced-2 protein (Figure 7A).

Discussion

The genes ced-3 and ced-4 are the only genes known to be required for programmed cell death to occur in C. elegans (Ellis and Horvitz, 1986). Our genetic and molecular studies of the ced-3 gene have revealed that this gene shares a number of features with ced-4: like ced-4 (see Yuan and Horvitz, 1992), ced-3 is not required for viability and is expressed mostly during embryogenesis, the stage during which 113 of the 131 programmed cell deaths occur (Sulston et al., 1983). Furthermore, just as ced-3 gene function is not required for ced-4 gene expression (Yuan and Horvitz, 1992), ced-4 gene function is not required for ced-3 gene expression. Thus, these two genes do not appear to control the onset of programmed cell death by acting sequentially in a transcriptional regulatory cascade.

The CED-4 protein is novel in sequence, and the only hint concerning its function is that two regions of the protein show some similarity to the EF-hand motif, which binds calcium (Yuan and Horvitz, 1992). For this reason we have suggested that the CED-4 protein and hence programmed cell death in C. elegans might be regulated by calcium. However, no direct evidence for this hypothesis has yet been obtained. The CED-3 protein similarly contains a region that offers a clue about possible function: a region of 99 amino acids contains 32 serines. Since serines are common phosphorylation sites (Edelman et al., 1987), it is possible that the CED-3 protein and hence programmed cell death in C. elegans are regulated by phosphorylation. Phosphorylation has previously been suggested to function in cell death (McConkey et al., 1990). Although the precise sequence of the serine-rich region varies among the three Caenorhabditis species studied, the relatively high number of serines is conserved (32, 31, and 33 in C. elegans, C. briggsae, and C. vulgari, respectively). None of the mutations in ced-3 affects the serine-rich region. These observations are consistent with the hypothesis that the presence of serines is more important than the precise amino acid sequence within this region.

Much more striking than the presence of the serine-rich region in the CED-3 protein is the similarity among the non-serine-rich regions of CED-3 and the human and murine ICEs. Human ICE was identified as a substrate-specific protease that cleaves the 31 kd pro-IL-1β between Asp-116 and Ala-117 to produce the mature 17.5 kd IL-1β. IL-1β is a cytokine involved in mediating a wide range of biological responses, including inflammation, septic shock, wound healing, hematopoiesis, and the growth of certain leukemias (Dinarello, 1991; Di Giovine and Duff, 1990). A specific inhibitor of ICE, the crmA gene product of cowpox virus prevents the proteolytic activation of IL-1β (Ray et al., 1992) and inhibits the host inflammatory response (Ray et al., 1992). Cowpox virus carrying a deleted crmA gene is unable to suppress the inflammatory response of chick embryos, resulting in a reduction in the number of virus-infected cells and less damage to the host (Palumbo et al., 1989). These observations indicate the importance of ICE in bringing about the inflammatory response.

A region of 115 amino acids (residues 246–360 of CED-3) shows the highest identity (43%) between the C. elegans CED-3 protein and the human ICE protein. This region contains a conserved pentapeptide QACRG (positions 356–360 of the CED-3 protein), which contains a cysteine known to be essential for ICE function. Specific modification of this cysteine in human ICE results in a complete loss of activity (Thornberry et al., 1992). The ced-3 mutation n2433 alters the conserved glycine in this pentapeptide and eliminates ced-3 function, suggesting that this glycine is important for ced-3 activity and might be an integral part of the active site of ICE. Six of the other seven identified ced-3 missense mutations also affect highly conserved residues that are likely to be important for the actions of both CED-3 and ICE. Interestingly, the mutations n718 (position 65 of CED-3) and n1040 (position 27 of CED-3) eliminate ced-3 function in vivo yet alter conserved residues that are not contained in either the mature p10 or p20 subunits of ICE (Thornberry et al., 1992). It is possible that these residues have a noncatalytic role in both CED-3 and ICE function, for example, in maintaining a proper conformation for proteolytic activation. The hu-
man ICE proenzyme (p45) can be proteolytically cleaved at four sites (Asp-103, Asp-119, Asp-297, and Asp-316 of ICE) to generate two peptides (p20 and p10) necessary for in vitro activity (Thornberry et al., 1992) and three other peptides with as yet undefined functions. At least two of these cleavage sites are conserved in CED-3, indicating that the CED-3 protein might be processed as well.

The similarity between the CED-3 and ICE proteins strongly suggests that CED-3 functions as a cysteine protease in controlling programmed cell death by proteolytically activating or inactivating a substrate protein or proteins. A potential substrate for CED-3 might be the product of the cod-4 gene. The CED-4 protein contains six aspartate residues that might be targets of the CED-3 protein. Four of these aspartates are followed by a serine (Asp-151, Asp-184, Asp-192, and Asp-541), and two are followed by an alanine (Asp-25 and Asp-459); of the four ICE cleavage sites in the ICE proenzyme, two are aspartic acid-serine and one is aspartic acid-alanine. Alternatively, the CED-3 protein might directly cause cell death by proteolytically cleaving proteins that are crucial for cell viability.

The similarity between CED-3 and ICE defines a novel protein family. Thornberry et al. (1992) suggested that the sequence GDSGP at position 287 of ICE resembles a GX(S/C)GXG motif found in serine and cysteine protease active sites. In the three nematode CED-3 proteins, however, only the first glycine of this sequence is conserved, and in mouse ICE the S/C is missing, suggesting that the CED-3/ICE family shares little sequence similarity with known protease families.

The similarity between CED-3 and ICE suggests not only that CED-3 might function as a cysteine protease but also that ICE might function in programmed cell death in vertebrates. Consistent with this hypothesis, after murine peritoneal macrophages were stimulated with lipopolysaccharide and induced to undergo programmed cell death by exposure to extracellular ATP, mature active IL-1β was released into the culture supernatant; by contrast, when cells were injured by scraping, IL-1β was released exclusively as the inactive proenzyme (Hoghoqiet et al., 1991). These results suggest that ICE might be activated upon induction of programmed cell death. A role for ICE in programmed cell death need not be mediated by IL-1β but rather could be mediated by another ICE substrate. ICE transcripts have been detected in cells that do not make IL-1β (Ceretti et al., 1992), suggesting that other ICE substrates might well exist. Alternatively, members of the CED-3/ICE family other than ICE might function in vertebrate programmed cell death.

The p10 subunit of ICE and the carboxy-terminal portions of the CED-3 protein are similar to the protein encoded by the murine nedd-2 gene, which is expressed during early embryonic brain development (Kumar et al., 1992). Since the Nedd-2 protein apparently lacks the OACRG active site region and is similar to ICE, primarily in the region of the p10 subunit of ICE, nedd-2 might function noncatalytically to regulate an ICE or ICE-like p20 subunit. Interestingly, three ced-3 missense mutations alter residues conserved among the Nedd-2 and CED-3 proteins. nedd-2 gene expression is high during embryonic brain development (Kumar et al., 1992), when much programmed cell death occurs (Oppenheimer, 1981). These observations suggest that Nedd-2 might function in programmed cell death.

The C. elegans gene ced-9 protects cells from undergoing programmed cell death by directly or indirectly antagonizing the activities of ced-3 and ced-4 (Hengartner et al., 1992). The vertebrate gene bcl-2 acts functionally similarly to ced-9: overexpression of bcl-2 protects or delays the onset of apoptotic cell death in a variety of vertebrate cell types as well as in C. elegans (Vaux et al., 1988; Nunez et al., 1990; Garcia et al., 1992; Senman et al., 1992; Strasser et al., 1991; Vaux et al., 1992; M. Hengartner and H. R. H., unpublished data). Thus, if ICE or another CED-3/ICE family member is involved in vertebrate programmed cell death, an intriguing possibility is that bcl-2 could act by modulating its activity. Furthermore, since bcl-2 is a dominant oncogene (overexpression of bcl-2 as a result of chromosomal translocation occurs in 85% of follicular and 20% of diffuse B cell lymphomas; Fukuura et al., 1979; Lovine et al., 1985; Yunis et al., 1987), ICE and other ced-3/ICE family members might be recessive oncogenes: the elimination of such cell death genes could prevent normal cell death and promote malignancy, just as overexpression of bcl-2 does.

Experimental Procedures

General Methods and Strains

The techniques used for culturing C. elegans were as described by Brenner (1974). All strains were grown at 20°C. The wild-type strains were C. elegans varity Bristol strain N2, Bergerac strain EM1002 (Emmons et al., 1983), C. briggsae, and C. vulgans (V. Ambros, personal communication). Genetic nomenclature follows the standard C. elegans system (Horvitz et al., 1979). The mutations used have been described by Brenner (1974) and by Hodgkin et al. (1988) or were isolated by us. These mutations are listed below.

LG I: ced-1(3735).

LG II: unc-3(928), unc-30(e191), ced-3(n717), n178, n1049, n129, n1163, n1164, n1165, n1286, n1949, n2426, n2430, n2432, unc-26(205).

LG V: egl-1(n487), n688.

LG X: dpy-2(n237).

Isolation of Additional Alleles of ced-3

A noncomplementation screen was designed to isolate new alleles of ced-3. Because animals carrying ced-3(n717) in trans to a deficiency are viable (Elia and Horvitz, 1986), we expected animals carrying a complete loss-of-function ced-3 allele generated by mutagenesis to be viable in trans to ced-3(n717), even if the new allele caused inviability in homozygotes. We used an egl-1 mutation in our screen. Dominant mutations in egl-1 cause the two hermaphrodite-specific neurons known as the HSNs to undergo programmed cell death (Trent et al., 1988). The HSNs are required for normal egg-laying, and egl-1 hermaphrodites, which lack HSNs, are egg-laying defective. The mutant phenotype of egl-1 is suppressed in a ced-3; egl-1 strain because mutations in ced-3 block programmed cell deaths. egl-1(n688) males were mutagenized with EMS (20 mM for 4 hr; Sigma, St. Louis, Missouri) and crossed with ced-3(n717) unc-26(e205); egl-1(n487); dpy-2(n237) hermaphrodites. Most cross progeny hermaphrodites were egg-laying defective because they were heterozygous for ced-3 and homozygous for egl-1. Rare egg-laying competent animals were picked as candidates for carrying new alleles of ced-3. Four such animals were isolated from about 10,000 hermaphrodite F1 cross progeny of EMS-mutagenized animals. These animals could have carried either a dominant suppressor of the egg-laying defect of egl-1 or a recessive mutation in ced-3. To distinguish between the two possibilities,
sites, cod-3 (new allele), egf-1 males were mated with egf-1 hermaphrodites, and the progeny (cod-3+/egf-1) were scored for suppression of the Egl phenotype. In each case, all of the progeny were egg-laying defective, suggesting that the mutation was recessive and thus likely to be a cod-3 allele.

Cosmid Libraries
Two cosmid libraries were used extensively in this work. A Sau3A1 partial digest genomic library of 7000 clones in the vector pHC79 was a gift from G. Benian (personal communication) and was used to isolate the cosmid MmM-C1 and MmM-C9. A Sau3A1 partial digest genomic library of 6000 clones in the vector pJBB (S. H. Hoekzema and Burke, 1989) was kindly provided by J. Coulston (Coulston et al., 1986) and was the source of cosmids Jc8, C4801, and C43C9. W07H16 is in the vector Lorid 6.

Identification and Mapping of RFLPs
To place cod-3 on the physical map, we sought to identify Tc1 elements flanking the region of interest in the leftmost part of the Bristol N2 or Bergerac EM1002 strains. We mated Bristol cod-3 unc-26+ males or Bristol unc-30 cod-3/+/males with Bergerac hermaphrodites. We isolated recombinant phenotypes Cod-3 non-Unc-26, Unc-26 Cod-3, Unc-30 non-Cod-3, and Cod-3 non-Unc-30 from the progeny of the Cod-3 unc-26 (Bristol)X Cod-3 (Bergerac) and unc-30 Cod-3 (Bristol)X Cod-3 (Bergerac) crosses, respectively. In this way, we established 13 ordered lines containing copies of chromosome IV derived from both the Bristol and Bergerac strains and recombinant in the region of cod-3.

By probing DNA from these strains with the Tc1 insert of plasmid pC2001 (Emmons et al., 1983), we identified a 5.1 kb EcoRI Tc1-containing restriction fragment specific to the Bristol strain and closely linked to cod-3. We named this RFLP nP35. nP35 was localized to cosmid MmM-C1 and MmM-C9, which were used to identify a cosmid contig in the cod-3 region (see text for details).

To identify additional RFLPs in the cod-3 region, DNAs from the recombinant inbred Bristol and Bergerac strains were digested with various restriction enzymes and probed with different cosmids to look for RFLPs between these strains. nP33 is a HindIII RFLP detected by the “right” end of cosmid Jc8, which is from the Coulston and Sulston library. The right end of Jc8 was made by digesting Jc8 with EcoRI and self-ligating. nP34 is a HindIII RFLP detected by the “left” end of Jc8. The left end of Jc8 was made by digesting Jc8 with SalI and self-ligating. nP36 and nP37 are HindIII RFLPs detected by the cosmids T10H5 and B0564, respectively.

We mapped the RFLPs nP33, nP34, nP35, nP36, and nP37 with respect to the genes unc-30, cod-3, and unc-26. The location of unc-30 on the physical map was determined by R. Hoskins (personal communication). Southern blots using DNA from the Cod-3 non-Unc-26 and Unc-26 non-Cod-3 recombinant strains described above were used to map nP33, nP34, and nP35 (data not shown). Three of these RFLPs, nP33, nP34, and nP36, mapped to the same region in the genome (data not shown). In this way, we established 13 ordered lines containing copies of chromosome IV derived from both the Bristol and Bergerac strains and recombinant in the region of cod-3.

Isolation of cod-3 cDNAs
To isolate cod-3 cDNA clones, the insert of pJ40 was used as a probe to screen a cDNA library from wild-type N2 hermaphrodites (R. K. Burdick, 1996). Seven cDNA clones were isolated, of which four were 3.5 kb in length (e.g., pJ85) and three were 2.5 kb in length (e.g., pJ87). One cDNA clone of each size class was subcloned and analyzed further. Two experiments showed that pJ85 contained DNA derived from a cod-3 transcript fused to DNA derived from an unrelated transcript: first, on a Northern blot containing N2 RNA, the pJ85 insert hybridized to two transcripts, one of which did not hybridize to the pJ40 insert; second, on a Southern blot containing N2 DNA, the pJ85 insert hybridized to one band in addition to those to which pJ40 hybridized (data not shown). Plasmid pJ87 contained a 2.5 kb cDNA clone and was determined to contain the complete coding region for cod-3 (see text).

Construction of a cod-3 Frameshift Mutation
The Sall site in pJ40 is located at position 5850 of the genomic sequence (Figures 1B and 4A). The construct FSA was obtained by cloning pJ40 with Sall and filling in the resulting gaps with the Klenow enzyme. By determining the sequence of FSA, we confirmed that it contained the sequence GTGCGTAAGC instead of GTGCGAC at the Sall site and so had a frameshift mutation that should result in the premature termination of protein synthesis at a UGA codon at position 6336.

Miniprep DNA was prepared from 1.5 ml of an overnight bacterial culture grown in superbroth (Sambrook et al., 1989). DNA was extracted by the alkaline lysis method, as described by Sambrook et al., 1989, and was treated with RNAase A (37°C for 30 min) and then with proteinase K (55°C for 30 min), phenol-extracted and then chloroform extracted, precipitated twice (first in 0.5 M sodium acetate [pH 5.2] and then in 0.1 M potassium acetate [pH 7.2]), and resuspended in 5 μl of injection buffer as described by Fire (1986). The DNA concentrations used in injections were between 100 μg/ml and 1 mg/ml.

Subclones of C4801 were generated as follows. C4801 was digested with BamHI and self-ligated to generate subclone C4801-2B. C4801-43 was generated by partial digestion of C4801-2B with BglII. pJ40 was generated by digestion of C4801-43 with Apal and BamHI. pJ107 was generated by partial digestion of pJ40 with BglII. pJ7.5 and pJ7.4 were generated by Exoll deletion of pJ107.

All transformation experiments used a cod-107735; unc-3(9298) cod-3(n1717) strain. The cod-3 mutation was present to facilitate scoring of the Cod-3 phenotype. Mutations in cod-1 block the engulfment of corpses generated by programmed cell death, causing the corpses of dead cells to persist much longer than in the wild type (Hedgecock et al., 1983). Thus, the presence of a corpse indicates a cell that has undergone programmed cell death. The Cod-3 phenotype was scored by counting the number of corpses present in the head of young L1 animals. Cosmids C4801 or plasmid subclones were mixed with C14G10 (which contains the unc-37(+) gene) at a weight ratio of 2:1 or 3:1 to increase the chances that an unc-37(+) transformant would contain the cosmid or plasmid being tested. Usually, 20-30 animals were injected in one experiment. Non-Unc F1 progeny of injected animals were isolated 3-4 days later. About one half to one third of the non-Unc progeny transmitted the non-Unc phenotype to the F2 generation and could be used to establish a line of transgenic animals. The young L1 progeny of such non-Unc transgenics were checked for the number of dead cells present in the head using Nomarski optics, as described by Ellis and Horvitz (1988). Animals of the transgenic line n1 transmitted the non-Unc phenotype to 100% of their progeny, implying that the cod-3 and unc-37 genes had integrated into the genome. Non-Unc animals of the transgenic line n62 transmitted the non-Unc phenotype to 50% of their progeny, implying that the cod-3 and unc-37 genes were present on an extrachromosomal array that is maintained as an unstable free duplication (Way and Chalfie, 1988).
Determination of the ced-3 Transcription Initiation Site

Two primers, Pex1 (5'-TCTACGACTTATGAGCATAGAAACTC-3') and Pex2 (5'-GTGGACACGTTCCACGTCCTCCGTTCT-3'), were used for primer extension experiments, which were performed as described by Triezenberg (1991) with minor modifications. The primers were used to amplify total RNA with the polymerase chain reaction were SL1 (5'-GTTAATTACCAAGTTTACG-3') and logS (5'-CCG-GTGACACTTGTCGACACT-3'). Amplification was performed using the GeneAmp kit (Perkin-Elmer Cetus, Norwalk, Connecticut). The products were reapplied using the primers SL1 and oligo10 (5'-ACT-ATTCACAGTCTG-3'). See text for additional details.

DNA Sequence Analysis

For DNA sequencing, serial deletions were made according to the procedure of Hanaford (1984). DNA sequences were determined using the Sequenase kit (U. S. Biochemical Corporation, Cleveland, Ohio) and protocols obtained from the manufacturer.

The CED-3 protein sequence was compared with protein sequences in the GenBank, PIR, and SWISS-PROT data bases at the National Center for Biotechnology Information using the BLAST network service.

Cloning of ced-3 Genes from Other Nematode Species

The C. briggsae and C. vulgare ced-3 genes were isolated from corresponding phage genomic libraries (C. Link, personal communication) using the ced-3 cDNA probe p1118 as a probe under low stringency hybridization conditions (5 × SSPE [Sambrook et al., 1989], 20% formamide, 0.2% Ficoll, 0.02% bovine serum albumin, 0.02% polyvinylpyrrolidone, 1% sodium dodecyl sulfate at 40°C overnight) and washed in 1 × SSPE and 0.5% sodium dodecyl sulfate twice at room temperature and twice at 42°C for 20 min for each wash.

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