Molecular cloning and sequence analysis of expansins—a highly conserved, multigene family of proteins that mediate cell wall extension in plants

cellulose/cellulose binding proteins/plant cell enlargement/pollen allergens


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ABSTRACT Expansins are unusual proteins discovered by virtue of their ability to mediate cell wall extension in plants. We identified cDNA clones for two cucumber expansins on the basis of peptide sequences of proteins purified from cucumber hypocotyls. The expansin cDNAs encode related proteins with signal peptide sequences directing secretion to the cell wall. Northern blot analysis showed moderate transcript abundance in the growing region of the hypocotyl and no detectable transcripts in the non-growing region. Rice and Arabidopsis expansin cDNAs were identified from collections of anonymous cDNAs (expressed sequence tags). Sequence comparisons indicate at least four distinct expansin cDNAs in rice and at least six in Arabidopsis. Expansins are highly conserved in size and sequence (60–87% amino acid sequence identity and 75–95% similarity between any pairwise comparison), and phylogenetic trees indicate that this multigene family formed before the evolutionary divergence of monocotyledons and dicotyledons. Sequence and motif analyses show no similarities to known functional domains that might account for expansin action on wall extension. A series of highly conserved tryptophans may function in expansin binding to cellulose or other glycans. The high conservation of this multigene family indicates that the mechanism by which expansins promote wall extension tolerates little variation in protein structure.

The growing cell wall in vascular plants consists of a complex network of cellulose microfibrils glued together by noncovalent interactions with matrix polymers (1). As a result of high turgor pressure, the cell wall bears considerable tensile stress. Growing cells enlarge by "loosening" the wall to allow slippage of the cellulose microfibrils and the adhering matrix polysaccharides and proteins (2). This results in relaxation of wall stress and turgor pressure and, consequently, an uptake of water to enlarge the cell and expand the wall (3). Despite considerable attention, the biochemical nature of wall loosening has long eluded definition (1).

Recently, we identified a class of wall proteins, called expansins, which have the remarkable ability to promote the extension of isolated plant cell walls (4). When growing cells are disrupted, for example by freezing or direct protein secretion to the cell wall. Northern blot analysis showed moderate transcript abundance in the growing region of the hypocotyl and no detectable transcripts in the non-growing region. Rice and Arabidopsis expansin cDNAs were identified from collections of anonymous cDNAs (expressed sequence tags). Sequence comparisons indicate at least four distinct expansin cDNAs in rice and at least six in Arabidopsis. Expansins are highly conserved in size and sequence (60–87% amino acid sequence identity and 75–95% similarity between any pairwise comparison), and phylogenetic trees indicate that this multigene family formed before the evolutionary divergence of monocotyledons and dicotyledons. Sequence and motif analyses show no similarities to known functional domains that might account for expansin action on wall extension. A series of highly conserved tryptophans may function in expansin binding to cellulose or other glycans. The high conservation of this multigene family indicates that the mechanism by which expansins promote wall extension tolerates little variation in protein structure.

which derives its mechanical strength from hydrogen bonding between cellulose fibers. Because expansins have an acidic optimum, they are ideal candidates for the protein mediator of "acid growth" in plants. Acid growth refers to the propensity of plant cells to grow faster when the wall pH is reduced (7, 8). This property is also observed in isolated cell walls under tension and has been reported in a wide variety of plant species.

In this study, we used peptide sequences from purified expansins to identify cDNA clones encoding these proteins. Sequence analysis of expansins from cucumber, Arabidopsis, pea, and rice indicates that these are unique proteins comprising an ancient, highly conserved, multigene family.

MATERIALS AND METHODS

Protein Purification and Peptide Sequencing. We purified two expansin proteins from cucumber hypocotyls as described (4). In brief, walls from the growing hypocotyl of etiolated cucumber seedlings (Cucumis sativus L., cv. Burpee Pickler) were collected and extracted for ionically bound proteins, which were precipitated with ammonium sulfate and fractionated by HPLC on C8 and sulfopropyl columns. Two active fractions, S1 and S2, were separated by SDS/PAGE, electroblotted onto a poly(vinylidene difluoride) membrane, and sent to the Wistar Protein Microsequencing Laboratory, Philadelphia, PA, for total amino acid composition and automated sequencing of tryptic peptides and undigest protein (9).

RNA Extraction, PCR Amplification, and cDNA Library Screening. RNA was extracted from 3-cm apical hypocotyl sections from 4-day etiolated cucumber seedlings. The segments were frozen in liquid nitrogen and extracted according to published methods (10). Poly(A)+ RNA was selected by standard procedures (11) and used for cDNA synthesis with reverse transcriptase (RT) (Gibco/BRL; type M-MLV). A cDNA library was constructed with the λ ZAP II vector, using the Stratagene ZAP cDNA synthesis kit.

Degenerate primers based on the N-terminal amino acid sequence from the cucumber S1 expansin were used to amplify a 95-bp cDNA by PCR (sense primer, 5'-GAYTAYGGNGNT-GGCA-3'; antisense primer, 5'-TANARRTNCCRTANCC-3'). This product was used as template for nested amplification with the primer 5'-CAYGGCNACNTTYTAYGG-3' and the same antisense primer, resulting in a 71-bp product, which was cloned into the Invitrogen TA cloning vector. The 71-bp product was radiolabeled by PCR with [α-32P]dCTP, purified by centrifugation through a Bio-Gel P-6 column, and used to screen the

Abbreviations: RT, reverse transcriptase; EST, expressed sequence tag.

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*The sequences reported in this paper have been deposited in the GenBank data base (accession nos. U0382, U30460, U30476, U30477, U30478, U30479, U30480, and U30481).
cucumber cDNA library. Immobilization of bacteriophage and hybridization were carried out as described (11). Bluescript SK phagemids containing positive inserts were rescued from λ ZAP II using Exassist helper phage (Stratagene).

For S2 expansin, we designed S2-specific degenerate primers; the sense primer 5'-TYAANCNGWSNGGNTGC3'- was based on the S2 N-terminal peptide; the antisense primer 5'-l CNCCKDTRATDNC3'- was based on two peptide sequences (S2-22 and S2-55), which we predicted to be conserved on the basis of homology with the S1 sequence (see Results and Discussion). Amplification with these primers was followed with a nested PCR amplification with the same antisense primer and a different sense primer (5'-CNCCKNCCKRTADNC3'), which was based on the most conserved nucleotide sequence in the alignments of cucumber S1 expansin with several Arabidopsis and rice expansin cDNAs (see Results and Discussion). The amplification yielded a product of the predicted size (~300 bp), which was cloned and sequenced to confirm an S2-specific partial clone of 312 bp. This fragment was used to screen the cDNA library.

DNA Sequencing. Plasmid DNA containing cDNA inserts was prepared by the Terrific Broth method (12) and used for dyeoxyxynucleotide sequencing with Sequenase (United States Biochemical). We used subcloning and primer walking strategies to obtain complete sequences.

In Vitro Translation and Western Blotting. DNA template was prepared by PCR amplification of the cucumber S1 expansin cDNA. Sense primer included the T7 RNA polymerase promoter followed by ATG and the first seven codons of the mature S1 expansin protein (5'-TAATACGACTCACTATAGGACCCATGACTACGCGTGGTGGAGCACGAGCCGCTGGCGACTACGATGGGAGAGCGTGTTTCACAGCGCGAGTCAGC3'). The antisense primer corresponded to the sequence following the stop codon (5'-CAGACTCATGCTGCACGGCGAGTCTGCACGGCGAGTCTGCA3'). DNA was purified with Promega's Wizard minicolumns and translated in vitro using Promega's Tnt coupled wheat germ extract system. S1 protein synthesis was assayed by [35S]methionine incorporation (data not shown) and by Western blot analysis using rabbit polyclonal antibodies directed against the cucumber S1 expansin (13).

Northern Blot Analysis. RNA was extracted as described above from the apical (growing) and basal (nongrowing) hypocotyl regions of 5-day etiolated cucumber seedlings. Fifteen micrograms of RNA was separated by electrophoresis in a 1.2% agarose gel with formaldehyde and Mops buffer and blotted onto Hybond-N membrane. S1 expansin cDNA was radiolabeled with [γ-32P]dCTP by random priming and hybridized to the filter (Amersham hybridization buffer tablets) at 65°C for 18 h and washed at the same temperature with 2× SSC for 15 min, then with 2× SSC/0.1% SDS for 15 min, and twice for 30 min. As a control, a blot was probed with a radiolabeled ubiquitin cDNA from Arabidopsis thaliana.

RESULTS AND DISCUSSION

Peptide Sequences. The following tryptic peptide sequences were obtained from purified S1 and S2 expansins: TGWQSMSR

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Fig. 1. Alignment of amino acid sequences predicted from cucumber S1 and S2 expansin cDNA clones. Underlined regions in boldface indicate the peptide sequences obtained from the purified proteins. A break is inserted between the predicted hydrophobic signal peptide and the start of the mature protein. Breaks in the alignment of the mature protein are indicated by dashes. Dotted line over S2 marks deleted region in the cDNA clone.
originally estimated from SDS/PAGE analysis of the purified protein (4). The predicted protein has a pl of 8.1, which resembles our previous estimate of 8.5 based on elution from ion-exchange columns (4). The amino acid composition of the S1 expansin is not remarkable, except that it is somewhat high in glycine (14.5%) and low in charged residues. Also, in comparison with average proteins (15), the S1 expansin is somewhat high in tryptophan (3%), cysteine (3.5%), and asparagine (9.25%). These properties are consistent with amino acid composition analysis of the purified protein (data not shown).

Analysis of the S1 expansin sequence by PROSITE (16) and BLOCKS (17) programs did not reveal any functional motifs that might account for the protein’s biochemical activity—e.g., no consensus sequences for glucanohydrolase domains or cellulose binding domains. This is consistent with previous results indicating the protein acts by an unusual mechanism (4, 6, 18).

The expansin protein does not contain repetitive domains, as judged by dot plots, and secondary structure predictions do not indicate any extraordinary features (data not shown).

S2 Expansin cDNA. We obtained a 312-bp partial clone of cucumber S2 expansin cDNA by RT-PCR using degenerate oligonucleotide primers based on the S2 expansin peptides (see Materials and Methods). This fragment was used to screen the cDNA library, resulting in a single isolate. Sequencing revealed that the clone apparently had intact 5’ and 3’ ends but contained a 109-bp deletion within the coding region, a deletion that resulted in a frameshift error and that omitted peptide S2-64 and part of peptide S2-59 (see Fig. 1). The sequence for the deleted region was obtained from cucumber hypocotyl mRNA by RT-PCR, using nested primers based on the region flanking the deletion and specific to the S2 sequence. A 337-bp PCR fragment containing the 109-bp deletion was amplified, cloned, and sequenced.

The predicted S2 expansin protein sequence is shown in Fig. 1. The primary translation product is 258 aa long (28.1 kDa). The (−3, −1) rule (14) predicts a signal peptide of 24 aa, cleavage of which should result in a mature protein of 234 aa (25.1 kDa) with an N-terminal peptide exactly matching that obtained from the purified protein. The mature protein has an overall amino acid identity to cucumber S1 expansin of 69%. As was the case with S1 expansin, no functional domains were identified by PROSITE and BLOCKS searches.

Expansin Homologs in Arabidopsis, Rice, and PEA Reveal a Highly Conserved Multigene Family. Using the S1 expansin amino acid sequence to search the nonredundant GenBank data base with the TBLASTN program (19), we identified a number of anonymous, partial cDNA sequences from the Arabidopsis and rice expressed sequence tag (EST) collections that appear to encode homologs of the cucumber expansins.

These were identified by their high sequence similarity to the cucumber expansins (≈75% identity at the amino acid level). We identify the Arabidopsis expansins as AtEx1, AtEx2, etc., and the rice expansins as RlExA, RlExB, etc., in the order of their identification. The cucumber S1 and S2 expansins are referred to as CuExS1 and CuExS2. These names are not intended to imply specific homologies between individual expansin genes among these species.

In rice, there are at present 20 EST sequences with high similarity to cucumber expansin. They fall into five sequence classes, four of which encode distinct members of the expansin gene family: RlExA (GenBank accession no. D42004, with 3 redundant entries), RlExB (accession no. D24200, with 11 redundancies), RlExC (accession no. D47507), and RlExD (accession no. D23861). The fifth class (accession no. D40668 with 2 redundancies) appears to be a partial clone missing the N-terminal half of the coding region. Because the ESTs are only partially sequenced, it is not yet possible to determine whether the partial clone represents a fifth expansin in rice or overlaps with one of the other four expansins.

Upon sequencing RlExB and RlExD, we found the rice expansins to be remarkably similar to cucumber expansins (Fig. 3). For example, CuExS1 and RlExD have 79% amino acid identity, which is greater than that shared by the two cucumber expansins.

In a similar manner, we identified nine Arabidopsis ESTs that encode at least six distinct expansins, represented by the following names and GenBank accession nos.: AtEx1 (Z37585), AtEx2 (Z29931), AtEx5 (T45214), AtEx6 (T45828), AtEx7 (F14125), and AtEx9 (R29778). In addition to these six, there are currently three ESTs that appear to be partial cDNAs, missing the N-terminal region of the protein: AtEx3 (Z29178), AtEx4 (Z24541), and AtEx7 (T76481); their sequences reveal them to be distinct from each other, but it is not yet possible to determine whether they overlap the other ESTs. Four of the Arabidopsis ESTs were obtained and sequenced (Fig. 3); like the rice expansins, they closely resemble the cucumber sequences.

Thus, in rice there appear to be at least four distinct expansin genes and in Arabidopsis there are at least six. It should be noted that the rice ESTs come from a single cDNA library made from whole seedlings, whereas the Arabidopsis ESTs were obtained from multiple cDNA libraries made from several different plant stages and tissue cultures (but all of the Columbia ecotype). It seems likely that there are additional expansins still to be found in rice and Arabidopsis. The smaller number found in cucumber probably reflects the greater selectivity of the cDNA library (growing hypocotyls only).

Besides the anonymous cDNAs (ESTs) that appear to be expansins, the next closest group of sequences in the GenBank data base appear to be proteins identified as type 1 pollen allergens from several species (e.g., Lolium and Phleum, accession nos. M57447 and X78813). They show ≈25% amino acid sequence identity to expansins and are conserved in size and in several structural features of the expansin protein (see Fig. 3 and below). Statistical tests using the program RDF2 (20) confirm that the sequences for expansins and the type 1 pollen allergens are significantly related. Also in GenBank (accession no. X85187) is a sequence for a pea cDNA that is identified as a pollen allergen-like protein, but its sequence identifies it as an expansin homolog (Fig. 3).

The high conservation among the various expansins from rice (a monocotyledon) and cucumber, Arabidopsis, and pea (all dicotyledons) suggests several important points: (i) that the expansin multigene family arose by gene duplication before the evolutionary divergence of monocotyledons and dicotyledons some 150 million years ago; (ii) that the expansin protein has strict functional constraints that limit the structural modifications possible while maintaining function; and (iii) that the function of the protein is important to the normal development
and physiology of angiosperms. These points are further supported by the fact that the signal peptide is not highly conserved among expansins; likewise, the 3' and 5' untranslated regions of expansin cDNAs are not conserved. Furthermore, the rate of nonsynonymous substitutions within the cDNA coding region for the mature protein is less than the rate of synonymous substitutions, indicating that negative (purifying) selection has worked against changes in the amino acid sequence of the protein.

To explore these points further, we constructed a phylogenetic tree (Fig. 4) based on nonsynonymous nucleotide substitutions using the MEGA program (21). The tree was rooted using the pollen allergen as outgroup. We note that the rice and cucumber expansins fall on either side of the root. This tree supports the idea that the duplication and functional specialization of expansin genes predated the divergence of monocotyledons and dicotyledons.

Phylogenetically Conserved Sites Suggest Functional Regions. Although PROSITE and blocks analyses did not reveal how expansins might act, several phylogenetically conserved features of the expansin sequence merit comment. First, we note that the N-terminal half of the expansin protein contains a series of 8 conserved cysteines with spacing similar to that of conserved cysteines in the chitin-binding domain of wheat germ agglutinin (22). The cysteines in this lectin form a series of intramolecular disulfide bonds that stabilize the protein's structure. We suggest that the N-terminal half of the expansin protein may be folded and stabilized by disulfide bonds in an analogous manner.

Second, we note that the C-terminal one-third of the expansin protein contains four conserved tryptophans whose spacing resembles that of tryptophans in the cellulose-binding domains of cellulase (23, 24). Site-directed mutagenesis confirms the significance of tryptophan residues for cellulose binding (25). Moreover, tryptophan and the structurally related

FIG. 3. Amino acid sequence alignments for nine expansin cDNAs from cucumber, rice, Arabidopsis, and pea. Also shown are aligned conserved residues from the Phleum type I pollen allergen (nonconserved amino acids are not shown). Residues identical to the consensus sequence in boldface, and nonconservative substitutions in lowercase. Numbers refer to the aligned consensus sequence, not individual sequences. Also shown on the number line are the conserved cysteines, tryptophans, and basic and acidic residues. Dashes indicate alignment gaps, and x indicates uncertain amino acid. Mature proteins, after signal peptide cleavage, are shown. Cleavage sites were predicted by the (−3, −1) rule with cgene software. The cleavage site for AtEx1 was not identified because the 5' end of the cDNA is missing.

FIG. 4. Phylogenetic tree of the nine expansin sequences from Fig. 3. This tree was constructed by neighbor joining for nonsynonymous nucleotide differences using the mEGA program. Numbers indicate bootstrap P values. The trees were rooted with the PhP1 pollen allergen as outgroup. Similar results were obtained when trees were constructed using γ distances and amino acid distances with Poisson correction.
lated amino acids phenylalanine and tyrosine are important for sugar binding in other proteins, and we note that these residues are particularly conserved in the C-terminal half of expansin. We therefore speculate that this region may be responsible for expansin binding to cellulose and related wall glycans (18).

Third, although charged residues are relatively underrepresented in the expansin protein, they are not randomly distributed along the protein backbone. The N-terminal half is nearly devoid of charged amino acids, whereas a short middle region (residues 119–139) contains the majority of the basic lysine and arginine residues (Fig. 3). These residues may be important for stabilizing protein structure by salt bridges or perhaps they interact with the carboxyl groups of wall pectins. Aspartic and glutamic acid residues may be particularly important for expansin function since the protein's acidic optimum (4) probably depends on protonation of the carboxyl groups of these residues. We point out in this regard that four glutamic acid residues (at positions 15, 106, 161, and 205; Fig. 3) are highly conserved among expansins.

Finally, we note that the conserved cysteines, tryptophans, and glutamic acids also appear as conserved residues between expansins and the pollen allergens (Fig. 3). This similarity suggests to us that the type I pollen allergen may function within the pollen cell wall as an expansin. This idea is supported by the fact that the N-terminal 23 aa of the Phleum PhlF1 pollen allergen is recognized as a signal peptide by the (−3, −1) rule. Regarding the limited conservation between expansins and the type I pollen allergens, we note that the pollen spore coat and the pollen tube wall are composed of polymers different from those that make up the cell walls of the plant body and that the genes that function in the haploid and diploid phases of plant life may have diverged long ago.

Summary and Perspectives. Our previous studies of expansins and their mediation of acid growth in higher plants indicate that expansins promote sustained slippage between wall polymers without hydrolytic activity (4, 6, 18). We now have shown that expansins constitute an ancient multigene family that arose before the evolutionary divergence of monocotyledons and dicotyledons. If the mechanism of wall extension in land plants is monophyletic, as seems reasonable given the similarity in wall structure in these groups, we predict that expansins will be found in all groups of land plants. This prediction is strengthened by the observation of acid growth mechanisms in mosses (26), ferns (27), and gymnosperms (28) as well as in angiosperms.

At present, the function of the multiple forms of expansin (e.g., at least six in Arabidopsis) is unclear, but three possibilities occur to us. The various expansin genes may be expressed in different growing organs or in different cell types. The fact that we can find only two expansin proteins and corresponding cDNAs in the cucumber hypocotyl, yet there are at least six expansins in Arabidopsis, lends some support for this idea. We assume that cucumber has as many expansins as has Arabidopsis, but they are found in tissues other than the hypocotyl. An alternative idea (not mutually exclusive with the first) is that there are different forms of expansin that act on different components of the complex wall. This idea gains some support from the fact that the two cucumber expansins possess different biochemical properties (pH dependence, action on wall stress relaxation; see ref. 18). A variation on this idea is that two or more expansins may be expressed in the same cell but be regulated by different stimuli, such as hormones, light, and environmental stresses.

The novelty of expansin’s sequence and its unique ability to induce plant cell wall extension present an unusual system for further study of how proteins may interact with polysaccharide networks to alter their rheological properties. By protein engineering and by analysis of more divergent expansins, it may be possible to identify the protein regions responsible for polysaccharide binding, pH dependence, and disruption of polysaccharide associations.

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