

RESEARCH ARTICLE

Expansin Engineering Database: A navigation and classification tool for expansins and homologues

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Funding information

Federal Ministry of Education and Research, Germany, Grant/Award Numbers: 01DG17027, 031B0571A; National Research Foundation, South Africa, Grant/Award Number: 105873

Abstract

Expansins have the remarkable ability to loosen plant cell walls and cellulose material without showing catalytic activity and therefore have potential applications in biomass degradation. To support the study of sequence-structure-function relationships and the search for novel expansins, the Expansin Engineering Database (ExED, <https://exed.biocatnet.de>) collected sequence and structure data on expansins from *Bacteria*, *Fungi*, and *Viridiplantae*, and expansin-like homologues such as carbohydrate binding modules, glycoside hydrolases, loosenins, swollenins, cerato-platanins, and EXPNs. Based on global sequence alignment and protein sequence network analysis, the sequences are highly diverse. However, many similarities were found between the expansin domains. Newly created profile hidden Markov models of the two expansin domains enable standard numbering schemes, comprehensive conservation analyses, and genome annotation. Conserved key amino acids in the expansin domains were identified, a refined classification of expansins and carbohydrate binding modules was proposed, and new sequence motifs facilitate the search of novel candidate genes and the engineering of expansins.

KEYWORDS

CBM63, GH45, protein domains, sequence analysis, sequence motifs, standard positions

1 | INTRODUCTION

Expansins are plant cell wall loosening proteins without apparent catalytic activity, which have been identified in a broad range of organisms.¹⁻⁴ The loosening mechanism is still elusive, but it has been suggested that the non-covalent interactions between cellulose microfibrils are weakened and moved against each other, thus the tight cellulosic structure is loosened.¹ Plant expansins interact with the primary cell wall, which consists of cellulose, hemicellulose, pectins, and xyloglucans.³ Key sites of wall loosening may be limited contact points between cellulose microfibrils.⁵ How these interactions result in cell wall loosening still requires further investigation.⁶ Expansins

were first discovered in plants and were described as proteins mediating pH-dependent extension and stress relaxation of cell walls.⁷ Based on phylogenetic analysis, it has been proposed that expansins in *Bacteria* and *Fungi* resulted from multiple horizontal gene transfers from plants to microbes,⁸ but there is also the possibility that the microbial expansin subfamily evolved first in ancient marine microorganisms, and then diversified into distinct terrestrial plant subfamilies.⁹

Expansins consist of two tightly packed protein domains, connected by a short linker and preceded by a signal peptide¹⁰ (Figure 1). Both expansin domains need to be connected for effective wall extension activity and weakening filter paper.^{11,12} The C-terminal domain of EXLX1 (expansin-like X) from *Bacillus subtilis* dominates the binding

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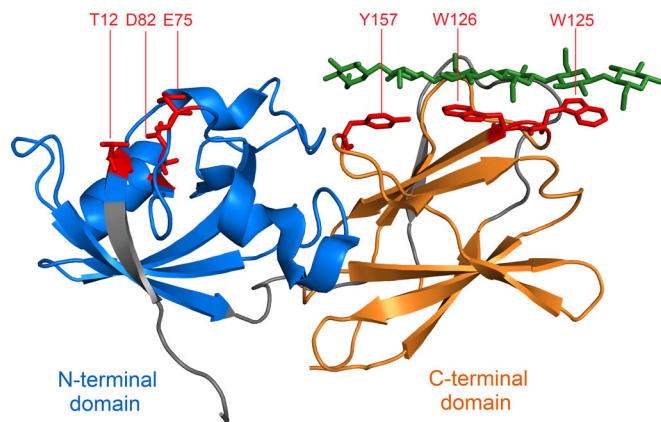


FIGURE 1 Functionally relevant positions in the expansin domains from the representative protein structure of *Bacillus subtilis* expansin BsEXLX1 (PDB entry 4fer, chain B) are labeled with standard position numbers (numbering according to Reference 14) and shown as sticks. The substrate cellohexaose is depicted above the C-terminal expansin domain in green [Color figure can be viewed at wileyonlinelibrary.com]

to cellulose and to matrix polysaccharides of cell walls through electrostatic or polar interaction.¹¹ The *Zea mays* β -expansin (ZmEXPB1) primarily binds glucuronoarabinoxylan, the major matrix polysaccharide in grass cell walls, and loosens it.¹³

Key amino acids in the N-terminal domain of *B. subtilis* expansin-like protein 1 (BsEXLX1) are two threonines at positions 12 and 14, a serine at position 16, two aspartates at positions 71 and 82, a tyrosine at position 73, and a glutamic acid at position 75,¹¹ numbered according to Reference 14. The threonine at standard position 12 is strongly conserved, but not essential for activity.¹¹ The aspartate at position 82 is crucial for activity; the threonine at position 14, the aspartate at position 71, and the tyrosine at position 73 are important for activity; and the serine at position 16 and the glutamic acid at position 75 play moderate roles in wall creep activity.¹¹ Three disulfide bridges can be found in the N-terminal domain of ZmEXPB1,¹⁵ and the six participating cysteines are highly conserved in the plant expansin groups, EXPA (expansin A), and EXPB (expansin B).¹⁵ An additional highly conserved cysteine pair is considered as a fourth disulfide bridge in plant α -expansins.¹⁶ In the expansin protein ScExl1 from the Basidiomycete fungus *Schizophyllum commune*, three disulfide bonds are predicted,¹⁷ whereas there is a lack of disulfide bridges in BsEXLX1¹⁴ and many other bacterial expansins.

The N-terminal expansin domain is formed by a six-stranded double- Ψ β -barrel¹⁴ that is shared by several protein superfamilies,¹⁸ for example, glycoside hydrolase family 45 (GH45).^{19,20} The expansin-like proteins found in Fungi such as loosenins, EXPNs, or cerato-platanins are single-domain proteins that resemble the N-terminal domain of expansins.^{21–23}

The C-terminal expansin domain is responsible for the binding to cellulosic material and is formed by two stacked β -sheets with an immunoglobulin-like fold.¹ The cellulose binding site on the protein surface consists of a linear arrangement of aromatic residues (tyrosines, phenylalanines, and tryptophans),¹⁴ which for BsEXLX1 includes

two tryptophans at positions 125 and 126, and a tyrosine at position 157.¹¹ A further key amino acid residue required for wall extension activity is a lysine at position 119.¹¹ The C-terminal domain of BsEXLX1 belongs to family 63 of carbohydrate binding modules (CBM63),¹¹ which mediate binding to polysaccharides.^{24,25}

In this article, we analyzed the similarity between “expansin-like proteins” (such as GH45s, loosenins, swollenins, cerato-platanins, EXPNs, and expansin-like proteins found in nematodes) and expansin domains on sequence level by establishing the Expansin Engineering Database (ExED), which collects characterized and putative expansin homologues. The protein sequences in the ExED were divided into different superfamilies (“Bacterial expansins,” “Fungal expansins,” and “Plant expansins”) according to sequence identity, and not by phylogenetic relationships of expansins, which were analyzed in Reference 9. By annotating the two expansin domains and using a continuous standard numbering scheme, conserved sequence motifs of the expansin protein family were identified that could be applied in the screening of genomic data for the identification of novel expansins.

2 | METHODS

2.1 | Sequence collection for the ExED

The ExED (<https://exed.biocatnet.de>) was built within the BioCatNet database system starting from 25 protein seed sequences (Table S1).²⁶ These seed sequences were used as queries for the basic local alignment search tool (BLAST)²⁷ using an e-value cutoff of 10^{-10} against the nonredundant protein database²⁸ of the National Center for Biotechnology Information (NCBI)²⁹ and the Protein Data Bank (PDB).³⁰ Two subsequent updates were performed to further enrich the ExED. For the first update, the sequences found by the initial search were clustered by UCLUST from the USEARCH package (version 11.0.667)³¹ by a threshold of 80% sequence identity, and the centroids (representative sequences) served as seed sequences for a BLAST search in the NCBI nonredundant protein database and the PDB. The seed sequences for the database updates of the ExED are available under <https://doi.org/10.18419/darus-622>. For the second update, profile hidden Markov models (HMMs) were generated for the N- and C-terminal expansin domains as described below. Further sequences were collected by searching with the *hmmsearch* command from the HMMER software package (version 3.1b2, <http://www.hmm.org>, Howard Hughes Medical Institute, Chevy Chase, Maryland).³² The hits were filtered by a minimal domain-based score of 35 (chosen after comparison with HMMER's domain-based “independent” e-values), a minimal hit length of 60 amino acids, and a maximal ratio of bias over domain-based score of 10%.

2.2 | Sequence hierarchy in the ExED

The initial 25 seed sequences comprise six bacterial, one fungal, and 17 plant expansins, as well as one expansin-like swollenin sequence.

The BLAST hits for each of these seed sequences were assigned to a corresponding superfamily named “Bacterial expansins,” “Fungal expansins,” “Plant expansins,” and “N-terminal domains.” Hence, the division of the identified protein sequences into the different superfamilies was based on sequence identity, and not on phylogenetic relationships. Herein, the term family refers to a group of sequences sharing a certain degree of similarity, that is, rather a cluster of similar sequences than a clade in a phylogenetic tree. Homologous families were created by a cutoff of 60% pairwise sequence identity as determined by the Needleman-Wunsch algorithm implemented in the EMBOSS software suite (version 6.6.0), with gap opening and extension penalties of 10 and 0.5, respectively.^{33,34} All sequence entries which shared at least 98% global sequence identity were assigned to a single protein entry. For each sequence entry, the respective superfamily, homologous family, and protein entry were annotated together with the identifiers of the original source database.

2.3 | Profile HMMs

A profile hidden Markov model (HMM)³² was derived for each expansin domain from a multiple sequence alignment built from 28 representative protein sequences, including 22 of the 25 seed sequences mentioned previously, two fungal sequences, and four sequences for which their structure was known (Table S2). To determine the region of the two domains in a multiple sequence alignment, four crystal structures of expansins were superimposed (PDB entries 1n10, chain A; 2hcz, chain X; 4fer, chain B; and 4jjo, chain A). The structure-based multiple sequence alignment (Figure S1) was generated by the Clustal Omega package³⁵ (version 1.2.1-1) and STAMP³⁶ (version 4.4), and visualized by PyMOL³⁷ (version 4.60, Schrödinger, New York, New York). Based on the structural alignment and on annotations of secondary structures in Pfam³⁸ (entries PF03330.17 for the N-terminal domain and PF01357.20 for the C-terminal domain), the respective domains were manually retrieved. The individual profile HMMs for the N- and C-terminal expansin domains were built by HMMER from the multiple sequence alignments. The input multiple sequence alignments were aligned against the derived output profile HMMs with the *hmmalign* command from HMMER in order to determine whether there are shifts between the input and output alignments. Shifted alignment columns were refined manually with respect to the positions of known secondary structure elements. The refined profile HMMs of the N- and C-terminal expansin domain comprise 95 and 75 positions, respectively (Figures S2 and S3), and are available together with their underlying alignments at <https://doi.org/10.18419/darus-623>.

2.4 | Standard numbering schemes

For the N- and C-terminal expansin domains, standard numbering schemes were introduced to annotate equivalent positions.³⁹ The *B. subtilis* expansin, BsEXLX1 (PDB entry 4fer), was used as the reference

sequence for the assignment of standard position numbers to the sequence entries in the ExED upon alignment against the respective profile HMM and subsequent transfer of position numbers: For both expansin domains, the standard positions range from 11 to 105 and 114 to 186. Insertions with respect to the reference sequence, such as loops, were specified by subsequent decimals. Thus, all position numbers mentioned herein are based on the reference BsEXLX1, unless otherwise stated. Due to insertions in the reference sequence of BsEXLX1, some regions in the underlying multiple sequence alignments of the standard numbering schemes appeared inaccurate, that is, these regions could not be aligned properly: In the N-terminal expansin domain, inaccurate positions are from 14.1 to 17, 39.1 to 47, and 104.1 to 105; in the C-terminal expansin domain, inaccurate positions are from 162.1 to 164 and 185.1 to 186.

2.5 | Conservation analyses

The two standard numbering schemes were used to analyze the amino acid frequencies for the two expansin domains. The domains were annotated by using *hmmscan* against all sequence entries of the ExED and deploying the match criteria mentioned previously. Each annotated domain position was analyzed for conserved amino acids. Groups of amino acids with similar biochemical properties, such as charge or polarity, were also taken into account.^{40,41} Conservation analyses were performed separately for each superfamily of the ExED, and additionally for EXPA, EXPB, EXLA (expansin-like A), and EXLB (expansin-like B) (Tables S3 and S4). An amino acid position was defined as conserved if it occurred in at least 70% of all annotated sequence entries. Conserved positions were compared with the positions in the structures of two bacterial expansins (PDB entries 4fer, chain B and 4jjo, chain A) and two plant expansins (PDB entries 1n10, chain A and 2hcz, chain X) to predict their functional relevance.

2.6 | Co-evolution of expansin domains

For comparison of the co-occurrence of the two expansin domains, all sequence entries from the ExED were aligned against the two profile HMMs for the expansin domains. Profile-to-sequence alignments were performed with the *hmmsearch* command from the HMMER software suite with the *max* option to collect all domain-based scores for each possible alignment. The lists of domain-based scores were sorted by sequence identifiers to ensure comparability, and in case of multiple hits, only the maximal bit score was kept. The bivariate histogram was visualized as heat map for bit scores greater than zero in MATLAB (version R2019a, The MathWorks, Natick, Massachusetts).

2.7 | Sequence length distributions

For comparing the lengths of the sequence entries in the ExED, histograms and boxplots were created with MATLAB to visualize

frequency distributions and to identify possibly fragmented or artificial sequences (version R2019a, The Mathworks, Natick, Massachusetts; version 2019a, Statistics and Machine Learning Toolbox version 11.5). The whisker length in a boxplot was chosen as 1.5 times the interquartile range.

2.8 | Protein sequence networks

Protein sequence networks visualize sequences as nodes in an undirected graph with edge weights to derive relationships between different clusters or communities. Protein sequence networks are useful for the identification of clusters of homologous sequences, that is, the visual representation of homologous families and their relationships to each other. The protein sequences in the ExED were sorted by decreasing sequence length and were subsequently clustered using the USEARCH algorithm (UCLUST) with a threshold of 90% sequence identity (without terminal gaps) to determine a reduced set of centroid sequences (representative sequences).³¹ For each centroid sequence, the N- and the C-terminal expansin domains were annotated by the two profile HMMs with the filter criteria mentioned above. Pairwise sequence identities between two sequences were derived from global Needleman-Wunsch alignments as described above and used as edge weights. Protein sequence networks were generated with edge weights of pairwise sequence identity, filtered by a pre-defined threshold. Metadata of the nodes (eg, the sequence ID) and of the edges (ie, the edge weights) were summarized in GraphML files by applying the NetworkX library in Python (version 1.9) for an automated assignment of node and edge attributes.⁴² The GraphML files are available at <https://doi.org/10.18419/darus-624>. Protein sequence networks were visualized with Cytoscape version 3.7.2⁴³ using a prefuse, force-directed layout with respect to the edge weights.

For the networks showing the relationships between CBM63s and expansin homologues, and between GH45s and the N-terminal expansin domain homologues, CD-HIT (version 4.7) was used with a clustering threshold of 90% and a word size of 5 (instead of UCLUST).^{44,45} The GH45 sequences were downloaded from the protein family database (Pfam, version 32.0, accession PF02015),⁴⁶ whereas the CBM63 sequences were downloaded from the carbohydrate-active enzymes (CAZy) database on June 3, 2019.⁴⁷ In the CAZy database, 633 individual CBM63 sequences were deposited, but only 582 NCBI accessions were available at the time of writing, as some of the records were moved or entries were merged. Members of CBM63 were annotated by the profile HMMs for the two expansin domains (<https://doi.org/10.18419/darus-625>).

2.9 | Homologous expansin-like domains in other proteins

In order to find similarities between the expansin family and the GH45 endoglucanase family, a structure-based multiple sequence

alignment was performed of the N-terminal expansin domains of five expansins (PDB entries 2bh0, 4jjo, 4fer, 1n10, and 2hcz) and the complete sequences of seven GH45s (PDB entries 1eng, 1hd5, 1oa9, 1wc2, 5h4u, 5kjo, and 5xbu). Five representative sequences of expansin-like proteins were analyzed: swollenin from *Trichoderma reesei* (NCBI accession AJ245918.1),⁴⁸ loosenin from *Bjerkandera adusta* (NCBI accession ADI72050.2),⁴⁹ EXPN from *Endogone* sp. FLAS-F59071 (NCBI accession RUS20349.1),^{50,51} an expansin-like protein found in nematode *Heterodera glycines* (NCBI accession ADL29728.1),⁵² and cerato-platanin from *Ceratocystis platani* (NCBI accession CAC84090.2).⁵³ The two profile HMMs of the N- and C-terminal expansin domains were used to search within these five expansin-like protein sequences for expansin domains with the filter criteria mentioned previously.

2.10 | Identification of expansin domains in actinobacterial genomes

Five actinobacterial genomes were selected to show the application of the ExED for the identification of expansin domains. An Illumina MiSeq sequencer was used to sequence the genomes (NGS facility, University of the Western Cape, South Africa). Due to the high G + C content of actinobacterial DNA, a 10% PhiX spike was included in the run.⁵⁴ The genomes were assembled using the A5-miseq pipeline.⁵⁵

The two newly created profile HMMs mentioned above were applied to search the five actinobacterial genomes for the occurrence of expansin domains. Nucleic acid sequences were translated using the default codon usage table available in the *transeq* tool from the EMBOSS software suite (version 6.6.0³⁴). Translated amino acid sequences with less than 60 subsequent amino acid symbols were discarded to reduce computation time.

The *hmmscan* tool from the HMMER software suite (version 3.1b2, <http://www.hmm.org>, Howard Hughes Medical Institute, Chevy Chase, Maryland) was used to scan the translated amino acid sequences with profile HMMs. The hits from *hmmscan* were filtered by a minimal domain-based score of 35 and a minimal coverage of 75% (defined as the ratio of hit length without insertions divided by the length of the profile HMM).

The matches for the profile HMMs of expansin domains were extended to find the adjacent start methionine and stop codon along the contig sequence of each match. The first or last available amino acid position in a contig was used to extend the hits, in case of a missing start or stop codon, respectively. The extended hit sequences are available for download under <https://doi.org/10.18419/darus-699>.

3 | RESULTS

3.1 | The ExED

The current version of the ExED contains 15 089 sequence entries, 12 400 protein entries, and 21 protein structures (Tables 1 and S5),

TABLE 1 Numbers of homologous families (Hfams), protein entries, sequence entries, and crystal structures in the different superfamilies (Sfam) of the ExED

Sfam	Sfam name	Hfams	Proteins	Sequences	Structures
1	Bacterial expansins	6	795	1172	10
2	Fungal expansins	1	421	543	0
3	Plant expansins	17	6636	8269	6
4	N-terminal domains	8	2102	2420	1
Total		32	9954	12404	17

Note: Superfamilies 1 to 3 were named after the kingdom of their most abundant source organisms. Superfamily 4, named “N-terminal domains,” comprises only proteins containing the N-terminal expansin domain.

which, based on global sequence similarity, were assigned to four superfamilies (comprising 12 404 sequence entries, 9954 protein entries and 17 structures). Three superfamilies include expansin homologues with two domains and were named according to their dominant source organisms: superfamily 1 “Bacterial expansins” (1172 sequences, 10 structures), superfamily 2 “Fungal expansins” (543 sequences, no structure), and superfamily 3 “Plant expansins” (8269 sequences, six structures). The members of superfamily 4 “N-terminal domains” consist of the N-terminal expansin domain only (2420 sequences, one structure). This superfamily comprises eukaryotic and bacterial sequences, for example, from *Magnoliophyta* (A, B, and C), *Actinobacteria*, *Oomycetes*, and *Basidiomycota*. The remaining number of 2685 sequences (corresponding to 2446 protein entries) and four structures could not be assigned to the four superfamilies and was thus collected in an unclassified fifth superfamily, which was omitted for further investigations.

The sequence lengths in the superfamilies “Bacterial expansins,” “Fungal expansins,” and “Plant expansins” vary between 40 and 1400 amino acids with a sharp peak between 250 and 270 amino acids and two minor peaks at 150 and at 600 amino acids (Figure S4). The sequence length distributions differ for each of the four superfamilies (Figure S5). For further analysis of whole expansin sequences and comparison with expansin-like proteins, only sequences with a length between 210 and 300 amino acids were considered (7706 sequences from the superfamilies “Bacterial expansins,” “Fungal expansins,” and “Plant expansins”) (Figure S4).

In the protein sequence network, which was built from global sequence alignments for the superfamilies “Bacterial expansins,” “Fungal expansins,” and “Plant expansins,” the latter are the most frequent group forming four large separate clusters, which consist of one or more homologous families (Hfams): cluster A has been classified as EXPA (Hfams 9-20), cluster B as EXPB (Hfams 21, 22), cluster C as EXLB (Hfams 24, 25), and cluster D as EXLA (Hfam 23) (Figure 2). These four clusters are followed by two clusters of the superfamily “Fungal expansins” (Hfam 7) and three clusters of the superfamily “Bacterial expansins” (Hfams 3, 4; 1, 2, 4, 6; and 3). Noteworthy, the “Plant expansins” clusters A and D also contain bacterial sequences. The protein sequence networks are a visual representation for the separation of expansins into different groups, and thus illustrate the assignment of the sequences in the ExED to families.

In our study, expansins were found in *Bacteria*, *Archaea*, and *Eukaryota*. When looking in detail at the major taxa in the tree of life

(after Figure 1 in Reference 9), expansins occur in *Gammaproteobacteria*, *Betaproteobacteria*, *Deltaproteobacteria*, *Acidobacteria*, *Bacteroidetes*, *Fibrobacteres*, *Ignavibacteria*, *Actinobacteria*, *Chloroflexi*, *Firmicutes*, *Cyanobacteria* (all *Bacteria*), *Euryarchaeota* (*Archaea*), *Metazoa*, *Fungi*, *Evosea*, *Discosea*, *Discoba*, *Embryophyta*, *Chloroplastida*, *Rhodophyta*, and *Stramenopiles* (all *Eukaryota*) (Table S6).

3.2 | Sequence space of expansin domains

Two profile HMMs for the N-terminal and the C-terminal expansin domains were derived and used for annotation of the two domains in all 12 404 classified sequences of the ExED (superfamilies 1, 2, 3, and 4), independent of their sequence lengths. For the superfamilies “Bacterial expansins,” “Fungal expansins,” and “Plant expansins,” the N- and the C-terminal expansin domains could be annotated in 9470 of 9984 sequences and in 8896 of 9984 sequences, respectively (Table S5). In 2182 of the 2420 sequences from the superfamily “N-terminal domains,” only the N-terminal expansin domain was annotated.

Based on the annotated domains in the classified superfamilies, two protein sequence networks were generated. The sequence network of N-terminal expansin domains is dominated by three large clusters (Figure S6): Homologues of cluster A classified as EXPA (Hfam 9-20), homologues of cluster B as EXPB (Hfam 21, 22), and homologues of cluster C as EXLB as well as fungal sequences (Hfam 3, 4, 8). These clusters are supplemented by clusters D (Hfam 24, 25; EXLB), E (Hfam 26; *Magnoliophyta* A), F (Hfam 23; EXLA), G (Hfam 7; Fungi), H (Hfam 27; *Magnoliophyta* B), and cluster I comprising N-terminal domains from different sources (Hfam 8, 11, 31, 32; Fungi, EXPA, *Basidiomycota*, Loosenin). The N-terminal domains of *Magnoliophyta* B, *Actinobacteria*, and *Oomycetes* form separate clusters. The sequences of CBM63 are within clusters of homologous families 3 and 4 from the superfamily “Bacterial sequences.”

The sequence network of the C-terminal expansin domain is dominated by six large clusters from “Plant expansins,” previously annotated as EXPA, EXPB, EXLB, and EXLA (clusters A-C and E-G), one cluster from “Fungal expansins” (D, Hfam 7), and three clusters from “Bacterial expansins” (H-J, Hfams 1, 3, 4, 6) (Figure S7). In each of the two domain-based networks, one bacterial sequence was found in a cluster from “Plant expansins,” *Streptomyces acidiscabies* (NCBI accession GAQ55178.1) in EXPA (Figure S6), and *Soehngenia saccharolytica* (NCBI accession TJX44964.1) in EXLA (Figure S7).

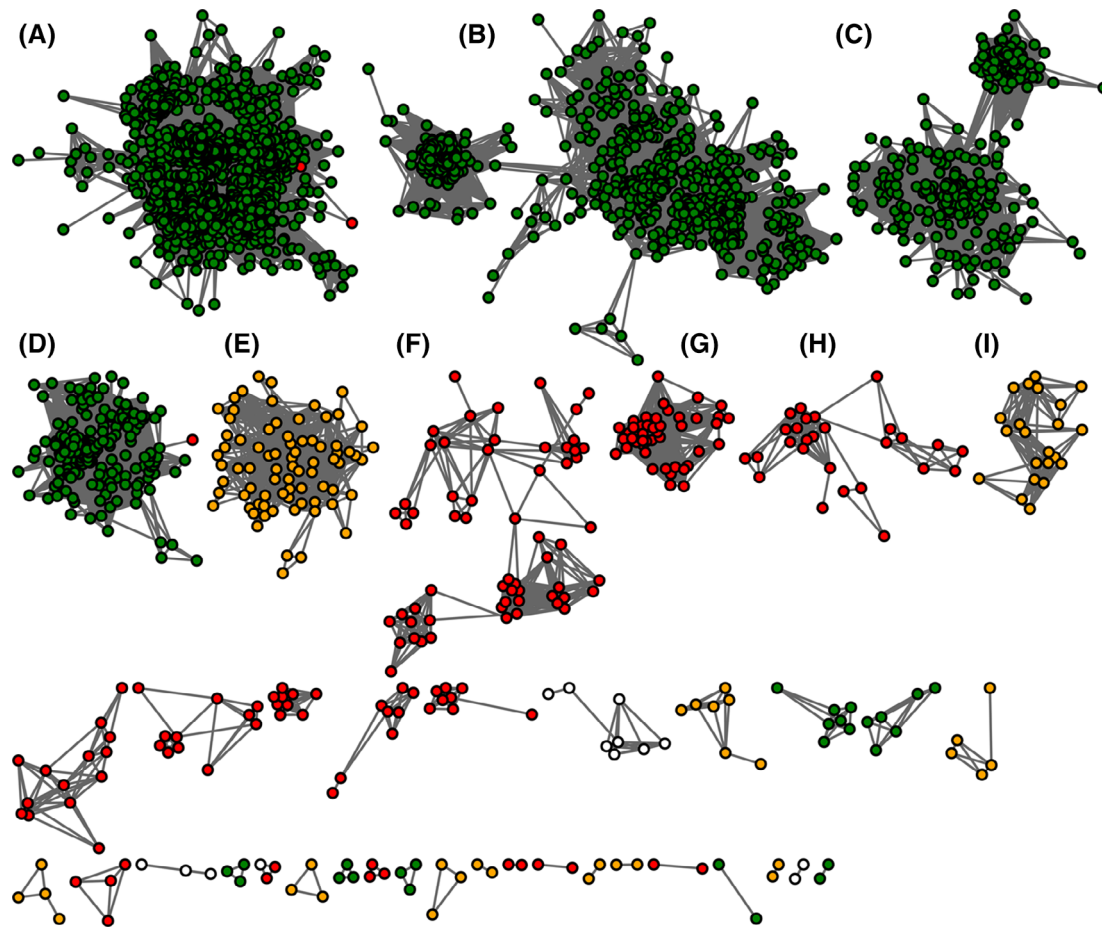


FIGURE 2 Protein sequence networks showing the sequence space of the expansin sequences in the ExED belonging to the superfamilies “Bacterial expansins,” “Fungal expansins,” and “Plant expansins”. These networks confirm the previous classification⁵⁷ that plant expansins can be divided into EXPA, EXPB, EXLA, and EXLB. All protein sequences presented in this network have a sequence length between 210 and 300 amino acids (Figure S4). The threshold for the nodes is 90% sequence identity (clustered with USEARCH) and the threshold for the edges is 50% pairwise sequence identity (determined by Needleman-Wunsch alignments). This network consists of 3504 nodes and 1 036 745 edges. With respect to the taxonomic lineages, the nodes from *Bacteria*, Fungi, *Viridiplantae*, and other origin are colored in red, orange, green, and white, respectively. The protein sequences of the nine biggest clusters belong to the following homologous families (Hfams) and expansin classifications: A (Hfams 9-20; expansin classification EXPA), B (21-22; EXPB), C (24-25; EXLB), D (23; EXLA), E (7, Fungi), F (3, 4; EXLX), G (1, 2, 4, 6; EXLX), H (3; EXLX), and I (7, Fungi). The bacterial sequences (red) in clusters A and D belong to *Streptomyces acidiscabies* (NCBI accession WP 050370046.1), *Kutzneria* sp. 744 (NCBI accession EWM10128.1, both Hfam 5, cluster A), and *Soehngenia saccharolyta* (NCBI accession TJX44964.1, cluster D)

The N- and C-terminal expansin domains have not evolved independently, but have co-evolved, as indicated by the correlation of sequence similarities of the two domains to the respective profile HMM (Figure 3). The shift in respect to the diagonal indicates a higher conservation for the N-terminal expansin domain than for the C-terminal expansin domain.

3.3 | Conserved positions in the two expansin domains

Standard numbering schemes for the N- and the C-terminal expansin domains (from positions 11 to 105 and 114 to 186, respectively) were applied to identify conserved positions ($\geq 70\%$ occurrence) in the

superfamilies “Bacterial expansins,” “Fungal expansins,” and “Plant expansins” (Table 2). In both expansin domains, glycine was the most frequently conserved amino acid (Tables S3 and S4). In the N-terminal expansin domain, nine positions were conserved in all three superfamilies (“Bacterial expansins,” “Fungal expansins,” and “Plant expansins”): T12, G21, A36, G53, P74, D82, L83, F88, and G97 (Table S3). In the superfamily “Bacterial expansins,” further seven positions were highly conserved ($\geq 90\%$ occurrence): V58, G60, G63, D71, S84, A87, and I91; in the superfamily “Fungal expansins,” C23, F25, W44, C52, C55, M68, and L81; and in the superfamily “Plant expansins,” A22, C23, G24, G49, C52, C55, C60, C61.8, T70, and F81.

Due to our conservation analysis, five of the previously proposed six cysteines¹⁵ were highly conserved in the superfamily “Plant expansins,” three conserved cysteines were found in the superfamily

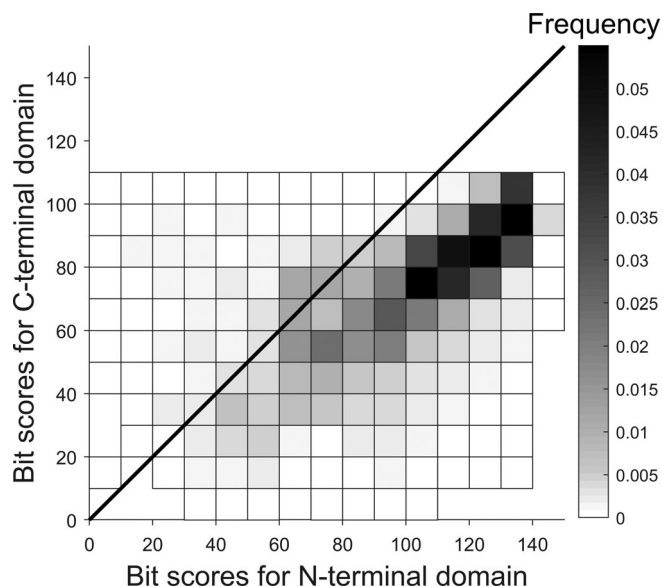


FIGURE 3 Bivariate histogram of co-occurring HMMER bit scores of the N- and C-terminal expansin domains. The greyscale bar represents the relative frequency of the bit scores. The black diagonal line is the bisecting line. The N-terminal expansin domain is more conserved than the C-terminal expansin domain, as the frequency distribution is shifted toward the x-axis. Furthermore, the HMMER bit scores for both expansin domains seem to correlate, which indicates co-evolution of both domains. The profile HMM of the N-terminal expansin domain is 20 positions longer than the one for the C-terminal expansin domain. Thus, larger bit scores can be observed for the N-terminal domain than for the C-terminal domain

“Fungal expansins,” and none in the superfamily “Bacterial expansins” (Table 2 and <https://doi.org/10.18419/darus-735>). The conserved cysteines at standard positions C23 and C52, C55, and a cysteine upstream of the N-terminal expansin domain standard numbering, and C60 and 61.8 were proposed to form disulfide bonds.¹⁵

In the C-terminal expansin domain, only two positions are conserved in the three superfamilies “Bacterial expansins,” “Fungal expansins,” and “Plant expansins”: W149 and G179 (Table S4). In addition, in the superfamily “Bacterial expansins,” highly conserved positions are K119, G121, W126, P137, Y157, N158, G166, T175, and D176; in the superfamily “Fungal expansins,” G121, S123, W126, F127, Q130, V131, N133, V143, S143.1, D146, R154, Y157, N158, F160, G164, V172, and T175; and in the superfamily “Plant expansins,” G136, G157, and W160. From the conserved positions, superfamily-specific motifs were derived. In the N-terminal domain of “Bacterial expansins,” these motifs are VpGP (standard positions 58–61, “p” is the abbreviation for polar amino acids selected from Reference 41) and HLDL (80–83) (Table 3); in the superfamily “Fungal expansins” the motifs T(F/W)YG (12–14 and 14.1), GTAnS (34–38, “n” is the abbreviation for nonpolar amino acids selected from Reference 41), VpGn (58–61), and HLDL (80–83) were identified; and in the superfamily “Plant expansins,” the motifs T(F/W)YG (12–14 and 14.1) in EXPA and EXPB, GGACGYG (20–26) in minor modifications in all four plant expansin groups, and HFDL (80–83) in EXPA and EXPB

were identified (Tables 3, S3, and <https://doi.org/10.18419/darus-735>). In the C-terminal expansin domain of the superfamily “Bacterial expansins,” the motif QVRNH (130–134) was conserved; in the superfamily “Fungal expansins” the motifs QVnN (130–133), LEVSTDGD (141–146, including 143.1 and 143.2 as insertions relative to the reference sequence), GGG (164–166), and VDVRVT (170–175) were identified. At the standard positions 170 to 175, the motif LSFpVT is included in sequences of EXPA. Sequences from the superfamilies “Bacterial expansins” and “Fungal expansins” were found to share the motif, KpG(S/T)S (119–123); and pGS exists also in EXPB. EXLA and EXLB were found to share the motif, YLA (126–128). The motif WGA exists in minor modifications in all four groups of “Plant expansins” (Tables 3, S4, and <https://doi.org/10.18419/darus-735>).

3.4 | Homologous expansin-like domains in other proteins

There is a considerable global sequence difference between all families of expansins and GH45 (Figure 4). Nevertheless, the GH45 protein sequences show conserved positions, which are also highly conserved in the superfamilies “Plant expansins” and “Fungal expansins”: the EXPA/EXPB motif HFDL (80–83), glycine 21, cysteine 23 and glycine 24 of the plant motif GGACGYG (21–26), threonine 12 and tyrosine 14 of the plant and fungal motif T(F/W)YG (12–14 and 14.1), and alanine 36 of the fungal motif GTAnS (34–38) (Figure S8). Thus, on a local sequence level, GH45 endoglucanases are more similar to the N-terminal expansin domain than expected from their different global protein sequences (Figure 4).

For further comparison, 582 protein sequences of the carbohydrate-binding module family 63 (CBM63) with a sequence length between 57 and 746 amino acids were downloaded from the CAZy database.⁴⁷ Interestingly, 510 of these sequences contained both expansin domains and were therefore already annotated in the ExED in the superfamilies “Bacterial expansins” and “Fungal expansins” (Table S10). Four CBM63 sequences contained only the C-terminal expansin domain, whereas 58 CBM63 sequences contained only the N-terminal expansin domain and shared a sequence identity of over 60% with N-terminal expansin domains of the superfamily “Bacterial expansins” (Figure S6). When using a lower threshold (minimal domain-based of 20 instead of 35) for filtering the expansin domains, 567 CBM63 sequences were found to contain both expansin domains. Nine sequences contained only the C-terminal expansin domain and one sequence (ATL67533.1) only the N-terminal domain (Table S10, <https://doi.org/10.18419/darus-625>). A protein sequence network including the whole CBM63 sequences and expansin sequences from the superfamilies “Bacterial expansins,” “Fungal expansins,” and “Plant expansins” revealed the similarity of CBM63 sequences to “Bacterial expansins” and also to “Fungal expansins” from homologous family 7, but also showed the distant relatedness to sequences of plant expansins (Figure 5).

The members of the superfamily “N-terminal domains” consist of the N-terminal expansin domain only. Similarly, loosenin (NCBI

TABLE 2 The conserved amino acids or groups of amino acids according to the standard numbering scheme for the N- or C-terminal expansin domain, with the sequence of *Bacillus subtilis* (PDB accession 4fer) as reference sequence

Standard position	Conserved amino acids in expansins from			Function and motif
	Bacteria	Fungi	Viridiplantae	
N-terminal expansin domain				
12	T (73%)	T (71%), S (3%)	T (82%), A (4%)	T(F/W)YG-motif
14	Polar (76%)	Y (86%), T (1%)	Y (85%), F (2%)	T(F/W)YG-motif
21	G (91%)	G (98%)	G (96%)	GGACGYG-motif
23	Non-polar (92%)	C (99%)	C (97%)	Disulfide bridge, ¹⁵ GGACGYG-motif
36	A (94%), G (4%)	A (79%), C (19%)	A (83%), G (9%)	Disulfide bridge ¹⁵
52	Non-polar (100%)	C (100%)	C (99%)	Disulfide bridge ¹⁵
53	G (100%)	G (100%)	G (99%)	Disulfide bridge ¹⁵
55	Y (70%), C (13%)	C (100%)	C (98%)	Disulfide bridge ¹⁵
60	G (92%), N (3%)	G (70%), Y (21%)	C (99%)	Important for wall extension activity ¹¹
61.8			C (96%), Y (1%)	Important for wall extension activity ¹¹
71	D (97%), N (3%)	D (78%), N (21%)	Polar (99%)	Moderate role for wall extension activity ¹¹
73		C (79%), T (19%)	C (79%), N (9%)	H(F/L)DL-motif, crucial for wall extension ¹¹
74	P (93%), G (2%)	P (73%), G (19%)	P (76%), Y (5%)	H(F/L)DL-motif
75	E (73%), G (11%)		G (78%), N (6%)	
82	D (99%)	D (99%), N (3%)	D (80%), V (8%)	
83	L (96%), M (3%)	L (81%), M (9%)	L (70%), M (26%)	
88	F (98%), Y (1%)	F (82%), W (16%)	F (82%), W (9%)	
97	G (96%)	G (95%)	G (93%), S (1%)	
C-terminal expansin domain				
119	K (97%), Q (1%)	K (89%), H (6%)	Polar (94%)	Important for wall extension activity ¹¹
125	W (63%), Y (30%)	Y (58%), N (15%)	Y (61%), N (14%)	Binding to cellulose material ¹¹
126	W (98%), Y (1%)	W (95%), F (3%)	F (57%), W (13%)	Binding to cellulose material ¹¹
149	W (88%), F (8%)	W (99%), Y (1%)	W (93%), C (3%)	Binding to cellulose material ¹¹
157	Y (90%), W (3%)	Y (93%), P (4%)	G (99%), S (1%)	
179	G (91%), H (5%)	G (95%), K (3%)	G (76%), R (12%)	

Note: All positions are listed separately for superfamilies 1 "Bacterial expansins," 2 "Fungal expansins," and 3 "Plant expansins" that are at least conserved to 70%. Positions marked in the standard numbering scheme as inaccurate are excluded (described in the Methods section). The last column names the function and the motif known from literature.^{10,56} If a single amino acid is at least conserved to 70%, the conservation of the respective amino acid group is not mentioned. Amino acid groups: nonpolar (A, C, F, G, I, L, M, P, V, W)⁴¹; polar (D, E, H, K, N, Q, R, S, T, Y).^{1,4}

TABLE 3 Known motifs from expansins in literature^{10,56} and the newly suggested motifs for the N- and C-terminal expansin domains based on the conservation analysis performed in this study (Tables S4 and S5, and <https://doi.org/10.18419/darus-735>)

Standard positions	Bacterial expansins	Fungal expansins	EXPA	EXPB	EXLA	EXLB
Motifs in the N-terminal expansin domain						
12 to 14 and 14.1		T(F/W)YG*	T(F/W)YG*	T(F/W)YG*		
20 to 26			GGACG*YG	GGACG	GACG*YG	GACG(Y/F)G
34 to 38		GTA _n S				
58 to 61	VpGP	VpG _n				
80 to 83	HLD*L	HLD*L	HFD* (L/I/V)	HFD*L		
Motifs in the C-terminal expansin domain						
119 to 123	KpG(S/T)S	KpG(S/T)S		pGS		
126 to 128					YLA	YLA
130 to 133 (and 134)	QVRNH	QV _n N				
141 to 146 (including 143.1 and 143.2)		LEVSTDGD				
156 to 158			WGp	WGA	pGA	(Y/F)GA
164 to 166		GGG				
170 to 175		VDVRVT	LSFpVT			

Note: Sequence motifs known from literature are marked with a star (*). Polar residues are abbreviated with "p" and nonpolar residues with "n." The expansins from *Viridiplantae* are separated into EXPA, EXPB, EXLA, and EXLB.

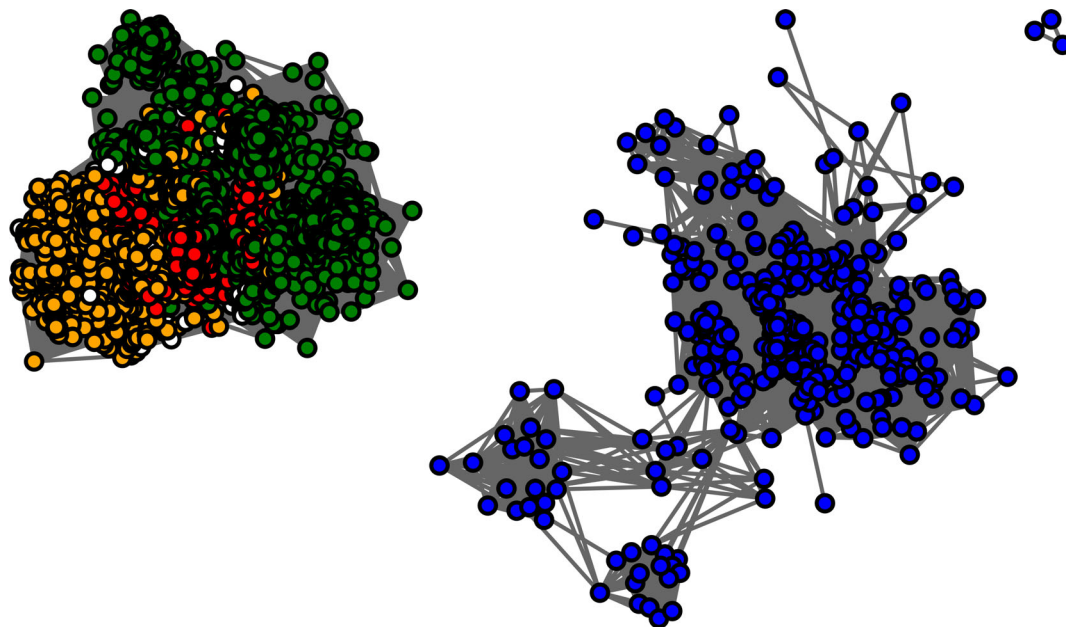


FIGURE 4 Protein sequence network showing the protein sequence space of GH45 sequences from Pfam⁴⁵ (accession PF02015) and the sequence regions annotated as N-terminal expansin domains from the superfamilies “Bacterial expansins,” “Fungal expansins,” “Plant expansins,” and “N-terminal domains.” The colors representing the origin of the expansin sequences correspond to the scheme in Figure 2 with GH45 sequences colored in blue. The threshold for the nodes is 90% sequence identity (clustered with USEARCH) and the threshold for the edges is 30% pairwise sequence identity (determined by Needleman-Wunsch alignments). This network consists of 4031 nodes and 2182,810 edges [Color figure can be viewed at wileyonlinelibrary.com]

ADI72050.2), EXPN from *Endogone* sp. FLAS-F59071 (NCBI accession RUS20349.1), the expansin-like protein found in nematode *Heterodera glycines* (NCBI ADL29728.1), and cerato-platanin from *Ceratocystis platani* (NCBI accession CAC84090.2) consist only of the N-terminal expansin domain (Table S7). At a threshold of 60% sequence identity, the N-terminal domains of loosenin and Basidiomycota cluster with fungal sequences from Hfam 7 and plant sequences from Hfam 11 (-Figure S6). In contrast, swollenin was found to possess only a distantly related C-terminal expansin domain (Table S7).

3.5 | Annotation of expansin domains in actinobacterial genomes

As a case study for the application of ExED in genome sequence annotation, actinobacterial genomes from various South African habitats were analyzed for the presence of expansin domains and conserved amino acid positions, using the profile HMMs of the expansin domains (Tables S8 and S9). In general, the sequence regions identified for the N-terminal expansin domains emerged with higher HMMER scores, whereas the C-terminal domains seemed less conserved (compare with Figure 3). Despite the lower scores for the C-terminal expansin domain, the coverage for the underlying profile HMM was still high (90%). One genome hit was identified in sediment samples collected at Gamka River in the Swartberg Mountain Range, which was identical to an expansin homologue from *Streptomyces swartbergensis* (NCBI accession WP_086602418), which matched well

the profile HMM of the N-terminal expansin domain (score: 60, 98% coverage) and moderately the profile HMM of the C-terminal expansin domain (score: 19, 89% coverage). The sequence from *S. swartbergensis* contains amino acids that are conserved in the superfamily “Bacterial expansins” (T12, G21, A36, G53, Y55, P74, D82, L83, F88, and G97 in the N-terminal expansin domain; K119, W126, W149, Y157, and G179 in the C-terminal expansin domain) and also amino acids that are conserved in the superfamilies “Fungal expansins” or “Plant expansins” (Y14, C23, C52, and C73).

4 | DISCUSSION

Expansins typically consist of about 225 amino acids (about 26 kDa) and an N-terminal signal peptide,² in total 250 to 275 amino acids,⁵⁶ which is in agreement with the average sequence length of 262 amino acids identified in this study. Thus, sequences shorter than 210 amino acids or longer than 300 amino acids were excluded from global sequence analyses (Figure S4). However, sequences with a length of about 600 amino acids contained replications of expansin domains as fusion proteins or due to sequencing errors, leading to expansin sequences that contained each domain two or three times. Since the two expansin domains have a length between 80 and 90 amino acids, shorter protein sequences can be considered as fragments or incomplete expansin domains.

The occurrence of expansins in major taxa in the tree of life (after Figure 1 in Reference 9 where a comprehensive phylogenetic analysis

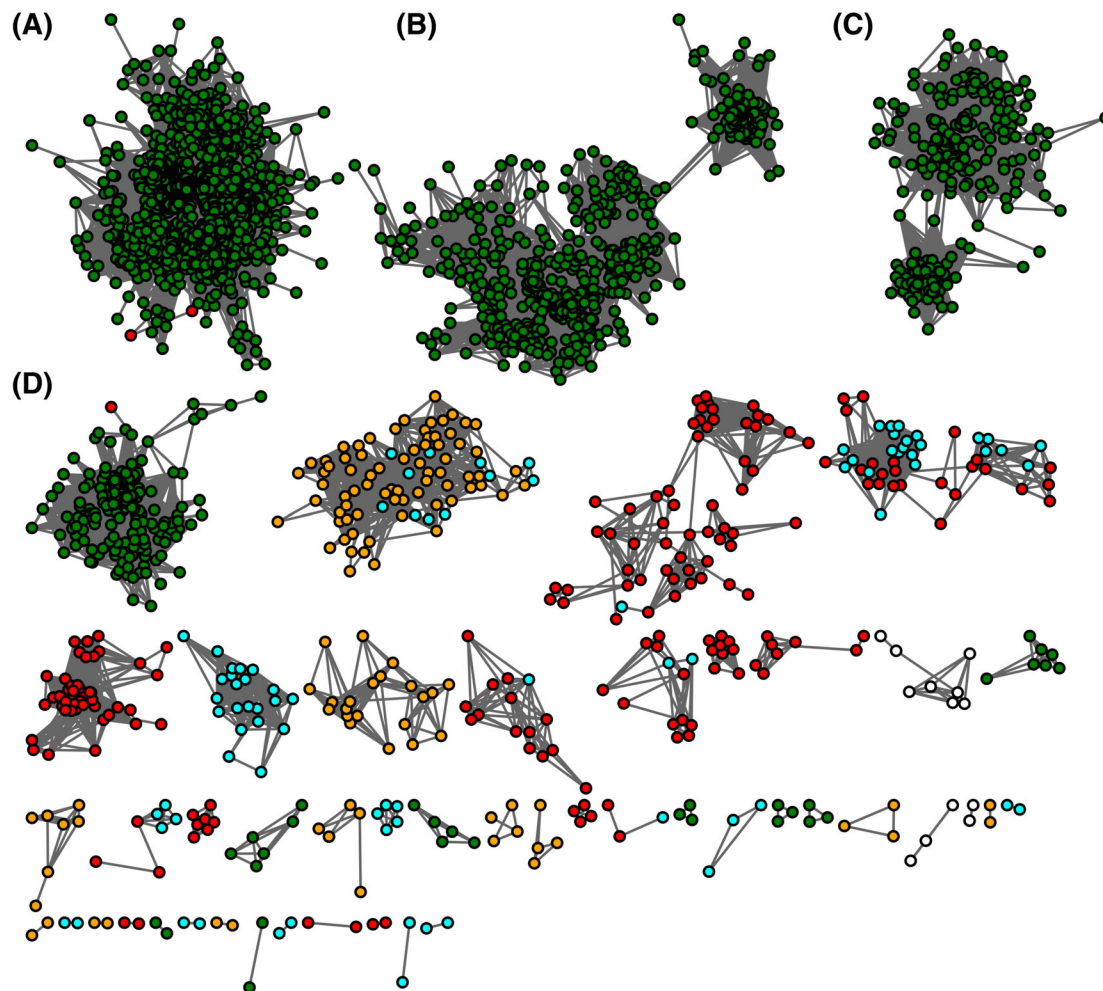


FIGURE 5 Protein sequence network showing the protein sequence space of CBM63 sequences from CAZy and the protein sequences of the superfamilies “Bacterial expansins,” “Fungal expansins,” and “Plant expansins” with a sequence length between 210 and 300 amino acids (Figure S4). In contrast to the four big clusters from “Plant expansins” (EXPA, A, EXPB, B, EXLA, C, and EXLB, D), where no CBM63 sequences can be found, the clusters from the superfamilies “Bacterial expansins” and “Fungal expansins” show many connections to sequences of CBM63. Therefore, sequences of CBM63 are similar to expansins from Fungi and *Bacteria* but have a distant relatedness to sequences of plant expansins. The colors representing the origin of the expansin sequences correspond to the scheme in Figure 2 with CBM63 sequences colored in cyan. The threshold for the nodes is 90% sequence identity (clustered with USEARCH) and the threshold for the edges is 50% pairwise sequence identity (determined by Needleman-Wunsch alignments). This network consists of 3344 nodes and 844 280 edges [Color figure can be viewed at wileyonlinelibrary.com]

of expansin genes across all kingdoms of life is shown) is comparable to the results obtained in this study (<https://doi.org/10.18419/darus-693>). For 12 of 90 groups that were compared, the results are different, for example, the archaeon *Halomicroarcula* sp. LR21 can be found in the ExED and contains one expansin homologue for which both expansin domains are annotated, whereas previous studies in Reference 9 did not find a putative expansin in *Archaea*. Other, apparently hitherto unknown, occurrences of putative expansins in the ExED include thirty-six sequences of *Fibrobacteres*, one sequence of *Ignavibacteria* in which both expansin domains can be found, seven sequences of *Discosea*, one sequence of *Discoba*, and one sequence of *Acidobacteria*, but without domain annotations. Further expansin sequences that were not included in this study but mentioned in Reference 9 are from the taxa *Verrucomicrobia*, *Chlorobi*, *Tubulinea*, *Glucophyta*, *Haptophyta*, *Dinoflagellata*, and *Phaeophyta*.

The protein sequence networks confirmed the nomenclature and classification of expansins into three kingdoms of *Bacteria*, *Fungi*, and *Viridiplantae* and the subclassification of plant expansins into EXPA, EXPB, EXLA, and EXLB⁵⁷ (Figure 2). Despite the differences on global sequence level, the protein sequence networks of expansins from *Bacteria* and *Viridiplantae* share similarities on a domain-based sequence level (Figures S1, S6, and S7). The N-terminal expansin domain is more conserved than the C-terminal expansin domain (Figure 3 and <https://doi.org/10.18419/darus-735>). When expansin homologues from more diverse backgrounds are discovered in the future, updated profile HMMs will show more insights into the possible co-evolution of both expansin domains.

A conservation analysis revealed and confirmed positions with an essential functional or structural role in expansin homologues. Glycine is structurally relevant, as it mediates the formation of short loops⁵⁸

and is frequently observed at the N- and C-caps of α -helices to increase helix stability.⁵⁹ As observed previously for other protein families,^{60,61} glycine is the most conserved amino acid in both expansin domains. In expansins, all four conserved glycines are located in loop regions (Table 2, compare with Figure 1). The conservation of threonine 12 and aspartate 82 in *Bacteria*, *Fungi*, EXPA, and EXPB confirms their functional role.¹¹ Interestingly, at standard position 75, which plays a moderate role in cell wall extension activity of *BsEXLX1*,¹¹ a glutamate is conserved in the superfamily "Bacterial expansins," and a glycine in EXPA and EXLB. In contrast, standard position 75 is not conserved in the superfamily "Fungal expansins," in EXPB, and in EXLA (Table 2 and <https://doi.org/10.18419/darus-735>). Aspartate 71, which has been proposed as important but not essential for wall extension activity of *BsEXLX1*,¹¹ is conserved in the superfamilies "Bacterial expansins" and "Fungal expansins," and in EXPB, EXLA, and EXLB (Table 2 and <https://doi.org/10.18419/darus-735>). However, three other proposed key amino acids for cell wall extension activity (threonine 14, serine 16, and tyrosine 73¹¹) are neither conserved in expansins from *Bacteria*, *Fungi*, nor *Viridiplantae* (Table S3 and <https://doi.org/10.18419/darus-735>), indicating the importance of an increased sample size for conservation analysis. The large number of expansin sequences investigated here also provided a deeper insight into the structural or functional relevance of disulfide bridges in the different superfamilies. Previously, three disulfide bridges were proposed to stabilize the tertiary structure of the N-terminal expansin domain of EXPA and EXPB.^{15,16} Five of the proposed six cysteines could be confirmed as highly conserved in the superfamily "Plant expansins" (Table 2 and <https://doi.org/10.18419/darus-735>). The sixth cysteine is located directly before the linker to the C-terminal expansin domain and therefore not included in our profile HMM for the N-terminal expansin domain. Against expectations, the additional highly conserved forth cysteine pair in plant α -expansins from¹⁶ was not found in our analysis (<https://doi.org/10.18419/darus-735>). Only three conserved cysteines were found in the superfamily "Fungal expansins," thus not all fungal expansin homologues possess three disulfide bridges, as concluded from the expansin *ScExlX1*.¹⁷ None of the six cysteines was conserved in the superfamily "Bacterial expansins" (Table 2), which is in accordance with previous observations of bacterial expansins lacking disulfide bridges.¹⁴ Due to the methodology of our conservation analysis, we have not investigated single bacterial expansins. From Reference 62 it is known, that a few bacterial expansins belonging to clades I and II in this respective paper have kept some of the conserved cysteines and conserved disulfides.

In the C-terminal expansin domain, the three aromatic residues at standard positions 125, 126, and 157, which mediate binding to cellulose,¹¹ are conserved in the superfamilies "Bacterial expansins" and "Fungal expansins," but are less conserved in the superfamily "Plant expansins" (Table S4 and <https://doi.org/10.18419/darus-735>). Lysine 119, which is important for cell wall-loosening activity,¹¹ is conserved in the superfamilies "Bacterial expansins" and "Fungal expansins," but not conserved in the superfamily "Plant expansins" (Table 2).

Through the use of conservation analysis, previously published family-specific motifs were confirmed: in the N-terminal expansin

domain, the T(F/W)YG motif was present in the two superfamilies "Bacterial expansins" and "Fungal expansins" (standard positions 12-14 and 14.1), and the motifs GGACG (20-24) and HFD (80-82) in the superfamily "Plant expansins"^{10,56} (Table 3). We suggest to extend the GGACG motif to a GGACGYG motif and the HFD motif to a HFDL motif in plant expansins. In bacterial and fungal expansins, these two plant motifs are slightly different: in the superfamily "Fungal expansins," the GGACGYG motif is shorter (GGxC), and in fungal and bacterial expansins the HFDL motif is replaced by HLDL. The HLD motif as well as the GGACS motif were already described for the fungal expansin *ScEXLX1*.¹⁷ Newly proposed motifs in the N-terminal expansin domain are VpGP (58-61) in the superfamily "Bacterial expansins" and GTAnS (34-38) in the superfamily "Fungal expansins" (Tables 3 and S3), where p and n denote polar and nonpolar amino acids, respectively. In expansins from *Fungi*, the proline of the VpGP-motif is replaced by a nonpolar amino acid. The previously described CDRC-motif at the amino terminus of EXLA⁵⁶ is located beyond the boundaries of our profile HMM for the N-terminal expansin domain.

No sequence motifs have been proposed yet for the C-terminal expansin domain, whereas we found eight novel motifs: KpG(S/T)S (119-123) and QVRNH (130-134) in the superfamilies "Bacterial expansins" and "Fungal expansins," where QVRNH shows slight modifications; LEVSTDGD (141-146, including 143.1 and 143.2), GGG (164-166), and VDVRVT (170-175) in "Fungal expansins"; YLA (126-128) in EXLA and EXLB; WGA (156-158) in EXPB and with slight modifications in EXPA, EXLA, and EXLB; and LSFpVT (170-175) in EXPA (Table 3). When annotating expansin sequences in future studies, these sequence motifs will help to assign unknown protein sequences (eg, metagenomic sequences) to the kingdoms *Viridiplantae*, *Bacteria*, or *Fungi*, and to distinguish the plant expansins EXPA, EXPB, EXLA, and EXLB (Tables 3, S3, and S4). Exemplary annotations were shown herein for actinobacterial genome samples from South Africa, including a putative expansin homologue from *S swartbergensis* (Tables S8 and S9).

The large number of expansin sequences used for analysis not only improved the identification of motifs, but also shed light on evolutionary relationships. Interestingly, when searching with the newly established profile HMMs for expansin domains within the CBM63 protein sequences from CAZy, 510 of the 582 CBM63 protein sequences were found to contain both expansin domains (Table S10). Only four sequences had a similarity to the C-terminal expansin domain, while missing the N-terminal expansin domain, as suggested previously,¹ and 58 CBM63 sequences contained only the N-terminal expansin domain. The discrepancies for the number of CBM63 homologues in comparison to the CAZy database result from the strict definitions of domain regions in the ExED (a minimal domain-based score of 35, a minimal hit length of 60 amino acids, and a maximal ratio of bias over domain-based score of 10%). This matter is further compounded when considering that the CAZy database does not only list CBM63 domains, but also includes proteins that have an associated CBM63 domain. When using a minimal domain-based score of 20 instead of 35, 567 of the 582 CBM63 protein sequences from CAZy were found to contain both expansin domains (Table S10, <https://doi.org/10.18419/darus-625>), nine sequences contained only

the C-terminal expansin domain, and one sequence only the N-terminal domain. Thus, the number of CBM63 sequences that contain only the N-terminal expansin domain decreases with less strict filter criteria. A possible way to resolve this would be to agree on a definition of the terms "CBM63" and "expansin," and to distinguish between domains and whole proteins. Such issues often occur with the description and classification of new proteins and will gradually be resolved over time as we learn more about these proteins from both functional studies as well as from analysis of the ever-increasing sequence space.

The observation of four bacterial sequences being found in clusters of plant expansins supports the hypothesis that microbial expansins were derived via horizontal gene transfer from plants to microbes⁸ (Figures 2, S6, and S7). The two bacterial sequences in clusters of the superfamily "Plant expansins" (Figure 5) are from the plant pathogens *Kutzneria* sp. 744 (NCBI accession EWM10128.1) and *Streptomyces acidiscabies* (NCBI accession WP 050370046.1), which are both actinobacteria, as described previously.²

With the chosen filter criteria, the sequence of the fungal swollenin does not contain any expansin domain. As the score for the C-terminal expansin domain is far below the chosen criteria, the swollenin sequence resembles a distantly related C-terminal expansin domain (Table S7), but we found no N-terminal expansin domain within the protein sequence of swollenin. This is due to the short N-terminal expansin domain in the swollenin from *Trichoderma reesei* and confirms the rather low sequence similarity between swollenin and expansins.⁴⁸

On a global sequence level, GH45s and N-terminal expansin domains share less than 30% pairwise sequence identity (Figure 4), and neither the profile HMM search of the N- and C-terminal expansin domains in the 542 GH45 sequences nor the profile HMM search of the GH45 profile HMM from Pfam (<https://pfam.xfam.org/family/PF02015/hmm>) in the 15 089 sequences of the ExED resulted in a match. In comparison to N-terminal expansin domains, GH45 sequences are longer due to several inserts and longer loop regions (179-208 amino acids as compared with 90-115 amino acids of the N-terminal expansin domains). Despite these differences, the evolutionary relationship between the two protein families is underlined by conserved amino acids. Both the conserved threonine and aspartate at standard positions 12 and 82, and the HFDL-motif (standard positions 80-83) were found in the GH45 protein sequences.

This study confirms the observation that microbial expansins comprise two protein domains and are widely distributed across diverse lineages of *Archaea*, *Bacteria*, *Fungi*, other eukaryotic microbes,⁹ and *Viridiplantae*. Therefore, the ExED can serve as a basis for a more detailed phylogenetic analysis in order to elucidate the origin of expansins and ancient evolutionary dynamics. Furthermore, the ExED can be used to search for expansin genes in virulent fungal and bacterial plant pathogens.

ACKNOWLEDGMENTS

We acknowledge funding by the Federal Ministry of Education and Research, Germany (BMBF, grants 01DG17027 and 031B0571A) and

the National Research Foundation, South Africa (NRF; UID: 105873). We thank Catharina Zeil for assisting with the database updates. Open access funding enabled and organized by Projekt DEAL.

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How to cite this article: Lohoff C, Buchholz PCF, Le Roes-Hill M, Pleiss J. Expansin Engineering Database: A navigation and classification tool for expansins and homologues. *Proteins*. 2020;1-14. <https://doi.org/10.1002/prot.26001>