Structural and functional studies of *Arabidopsis thaliana* legumain beta reveal isoform specific mechanisms of activation and substrate recognition

Authors:

Elfriede Dall $^{\dagger*\#}$, Florian B. Zauner †* , Wai Tuck Soh † , Fatih Demir § , Sven O. Dahms † , Chiara Cabrele † , Pitter F. Huesgen $^{\S|_1}$, Hans Brandstetter $^{\dagger\#}$.

Affiliation:

[†]Department of Biosciences, University of Salzburg, 5020 Salzburg, Austria.

§Central Institute for Engineering, Electronics and Analytics, ZEA-3, Forschungszentrum Jülich, 52428 Jülich, Germany.

CECAD, Medical Faculty and University Hospital, University of Cologne, 50931 Cologne, Germany

¹Institute for Biochemistry, Faculty of Mathematics and Natural Sciences, University of Cologne, 50674 Cologne, Germany

*These authors contributed equally to this work.

*Corresponding authors:

Elfriede Dall: elfriede.dall@sbg.ac.at

Hans Brandstetter: hans.brandstetter@sbg.ac.at

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Abstract

The vacuolar cysteine protease legumain plays important functions in seed maturation and plant programmed cell death. Because of their dual protease and ligase activity, plant legumains have become of particular biotechnological interest e.g. for the synthesis of cyclic peptides for drug design or for protein engineering. However, the molecular mechanisms behind their dual protease and ligase activities are still poorly understood, limiting their applications. Here we present the crystal structure of Arabidopsis thaliana legumain isoform B (AtLEGβ) in its zymogen state. Combining structural and biochemical experiments, we show for the first time that plant legumains encode distinct, isoform-specific activation mechanisms. While the autocatalytic activation of isoform y (AtLEGy) is controlled by the latency-conferring dimer state, the activation of the monomeric **AtLEGB** is concentration independent. Additionally, in AtLEGB the plant-characteristic two-chain intermediate state is stabilized by hydrophobic rather than ionic interactions as in AtLEGy, resulting in significantly different pHstability profiles. The crystal structure of AtLEGB reveiled unrestricted non-prime substrate binding pockets, consistent with the broad substrate specificity as determined by degradomic assays. Further to its protease activity, we show that AtLEGB exhibits a true peptide ligase activity. While cleavage-dependent transpeptidase activity has been reported for other plant legumains, AtLEGβ is the first example of a plant legumain capable of linking free termini. The discovery of these isoform specific differences will allow to identify and rationally design efficient ligases with application biotechnology and in drug development.

Introduction

The plant cysteine proteases of the legumain family (C13 family, EC 3.4.22.34) have an important role in processing and maturation of seed storage proteins within the vacuole and are therefore also referred to as vacuolar processing enzymes (VPEs) (1). Plant legumains are structurally related to the mammalian caspases and exhibit a strong substrate sequence preference for cleavage after asparagine and, to a lesser extent, aspartate residues (2,3).

Therefore, they are also synonymously referred to as the asparaginyl endopeptidases (AEP). In contrast to mammals, where only one functional legumain isoform is expressed, Arabidopsis thaliana contains four genes coding for legumains $(\alpha, \beta, \gamma, \delta$ -VPE) and other plants even up to eight functional variants (4). Plant legumains are expressed primarily in seeds and vegetative organs, consistent with their phylogenetic grouping into two angiosperm clades, the seed type (β-VPE) and non-seed or vegetative type VPEs (α -, γ - and δ -VPE) (5-8). Vegetative legumains are found in lytic vacuoles and have been suggested to play critical roles in plant programmed cell death (PCD) and may functionally substitute the caspases, which are absent in plants (9). Seed type legumains like thaliana legumain Arabidopsis isoform (AtLEGβ) play important functions in the processing and maturation of seed storage proteins within storage vacuoles (10,11). The importance of legumains is especially illustrated in Arabidopsis mutant strains missing all four legumain genes (a, β, γ, δ), which were shown to accumulate aberrantly processed seed storage proteins (12). Importantly, AtLEG β can compensate missing vegetative α and γ proteins, further confirming that AtLEGβ is the main player in precursor protein processing in seeds (10). Known substrates of AtLEGB include the pro12S globulin and pro2S albumin proteins (5,10,12,13).

On top of that, several plant legumains possess peptide ligase and cyclase activity (14-20). Recently we could show that the vegetative type AtLEGy harbors ligase activity (21). However, it is still unknown whether this is also true for the other three A. thaliana legumain isoforms, especially the phylogenetically more distant seed type AtLEGB. Cyclic peptides are important for plants' defense against pathogens (16,17,22,23).Well characterized examples include the kalata B1 peptide found in Oldenlandia affinis which has proven antimicrobial and insecticidal activities and the Sunflower trypsin inhibitor 1 (SFTI) (22,24). Cyclic peptides are very resistant to extremes in pH and temperature, making them ideal scaffolds for biotechnological applications and drug design (25-27). Peptide cyclisation in plants is typically catalyzed by legumains. Consequently, there is a high interest in understanding the ligation mechanism, specificity and efficacy of different plant legumain isoforms. Recent studies led to the discovery of a marker of ligase activity (MLA) and

a gatekeeper residue (Cys247, *Oldenlandia affinis* numbering) that allow to predict ligase activity based on sequence information (20,28). However, in order to validate these marker regions, experimental data on ligase activity of different legumain isoforms is indispensable.

Structural analysis of plant legumains showed that they are synthesized as inactive zymogens composed of a caspase-like catalytic domain with the AEP activity (AEP domain) and a C-terminal death domain like prodomain (LSAM domain, Legumain Stabilization and Activity Modulation domain) that are connected by an activation peptide (AP) harboring the α 6-helix (20,21,29,30). Although this tripartite domain architecture (AEP-AP-LSAM) is conserved in mammalian and plant legumains, the activation process of vegetative-type proAtLEGy (Arabidopsis thaliana prolegumain isoform γ) significantly differs from that of human legumain (31,32). Importantly, proAtLEGy is present in an enzymatically latent dimer state that is mediated by AP-LSAM – AP'-LSAM' interactions and is dependent on pH and protein concentration (21). Furthermore, we have previously shown that conversion to the active, monomeric AEP form, i.e. release of the prodomain, proceeds via a previously unknown two-chain intermediate state. Two-chain AtLEGy results from cleavage at the N-terminal side of the α 6-helix within the AP and is suppressed by high protein concentration where AtLEGy dimerization is favored. Even after an initial cleavage within the AP, an enzymatically latent, dimeric two-chain AtLEGy intermediate form remains stable at neutral pH-environment. Only at acidic pH the dimer dissociates to monomeric twochain legumain, which may further release the LSAM domain and thereby convert to the mature AEP form. The identification of the dimer and twochain states allowed developing a pH-dependent four step activation-model mechanism of plant legumains, i.e. single chain – two chain conversion; α6-helix destabilization; dimer – monomer dissociation; and AEP – LSAM release. However, given the subtle regulation of these conversions, isoform specific differences in activation are to be expected, with experimental data still lacking.

Here we present the crystal structure of zymogenic pro $AtLEG\beta$ which led to the discovery of a distinct activation mechanism, contrasting that of $AtLEG\gamma$. Combining structural and biochemical information, we show for the first time that plant legumains follow isoform specific autocatalytic activation

mechanisms and differential strategies of activity regulation and stability. Furthermore, we provide evidence that seed type AtLEG β is an active ligase capable of peptide cyclisation. AtLEG β ligase activity is not strictly linked to peptide bond cleavage but enables also the efficient joining of free N- and C-termini. To our knowledge, AtLEG β is the first example of a plant legumain for which we could demonstrate the ligation of free peptide termini.

This study broadens our understanding of isoformspecific differences in plant legumains and their relevance in plant physiology. Furthermore, the study discloses new avenues to rationally design peptide ligases with applications in biotechnology and drug development.

Results

Crystal structure of proAtLEGB

To understand isoform specific differences between different AtLEGs we determined the crystal structure of seed-type proAtLEGβ to a resolution of 2.0 Å (Table S1). The asymmetric unit of the tetragonal space group contained 12 independent molecules. Like isoform γ, proAtLEGβ comprises an N-terminal caspase-like catalytic domain and a C-terminal Legumain Stabilization and Activity Modulation (LSAM) domain with death domainlike topology (Fig. 1 and S1). The AEP and LSAM domain are connected by an activation peptide that harbors the α6-helix. Overall, the structure of proAtLEGB closely resembles the structure of the homologous two-chain AtLEGγ indicated by a Cαrmsd of 0.49 Å. However, inspecting the individual sub-domains unraveled specific differences. While the catalytic AEP domains of AtLEGB and y superimpose very well with an overall Cα-rmsd of 0.39 Å, we observed bigger differences in the LSAM domains with a Cα-rmsd of 0.78 Å (determined with Pymol). This observation is also in agreement with a higher sequence identity of the β and γ catalytic domains (67% identity) compared to the LSAM domains (56% identity). Furthermore, we observed an isoform specific glycosylation at Asn309, located at the bottom of the enzyme, which is also conserved in human legumain (Fig. 1A and 2A).

proAtLEG β forms atypical dimers in the crystal and is monomeric in solution

An important feature of proAtLEGy is that it exists in a latent dimer state in solution, which is mediated by AP-LSAM - AP'-LSAM' interactions. This dimer controls both the activation and activity of AtLEGy (21). Similarly, in the crystal structure of proAtLEGB we found all twelve independent protomers in the crystallographic asymmetric unit to engage in symmetric dimer contacts which were mediated by LSAM - LSAM' interactions (Fig. 2A). However, these interactions were mediated by different amino acids and led to an approximately 90° tilted orientation of the monomers within the β and y-dimer, respectively (Fig. 2B, C). Indeed, detailed analyses of the β and γ dimer interfaces revealed significant, isoform specific differences. The proAtLEGy dimer is mediated primarily by three symmetric anchoring sites, $\alpha 6$ and $\alpha 7$ helices and a conserved cyclic protein recognition motif (cPRM) on the c341-loop. The α 6 and α 7 helices form a 4-helix bundle that is stabilized around a symmetric hydrophobic core formed by $W363^{\gamma}$ and Val383^γ, L384^γ, respectively (AtLEGγ numbering; Fig. 2C, E). This hydrophobic core is further stabilized by a network of salt bridges on the N- $(R355^{\gamma} - E371^{\gamma})$ and $D356^{\gamma} - K376^{\gamma}$ and Cterminal (K376 $^{\gamma}$ – D356 $^{\gamma}$ ' and E371 $^{\gamma}$ – R355 $^{\gamma}$ ') ends of the α6-helices. By contrast, the proAtLEGβ dimers in the crystal structure were predominantly mediated by the α7 helix. This interaction was formed around the symmetric $H384^{\beta}$ ($H392^{\gamma}$) and further stabilized by one symmetric salt bridge $(E390^{\beta} - K383^{\beta})$ as well as by a hydrophobic contact of the α 7 C-terminal LFG motif (396 β -398 β) with W355 β ' centered in the α 6 helix (Fig. 2D, E). The hydrophobic core of the $\alpha 6-\alpha 7$, $\alpha 6'-\alpha 7'$ four helix bundle was missing as was any stabilization by the conserved cPRM, despite key residues important for proAtLEGy-like dimer formation being conserved in proAtLEGβ (Fig. S1). However, modeling a proAtLEGy-like dimer uncovered repulsive charge densities of α 7- α 7' helix contact residues in AtLEGB (R380 – R380', K373 – K383', D369 – D386') that will prohibit this γ -mode of dimerisation (Fig. 2F and S2). Together, these findings suggest that the observed β -dimer is weak and probably only transient in solution. To test this conclusion, we performed size exclusion chromatography (SEC) experiments. As expected, at pH 7.0 proAtLEGβ migrated at the expected size of a monomer, similar to human legumain (Fig. 2F). Accordingly, proAtLEGB was a monomer in solution.

Conserved Gln346 keeps proenzyme in latent state

Comparing the crystal structure of proAtLEGB with YVAD-cmk inhibited AtLEGy we found that the AP binds to the non-prime substrate binding sites in a substrate-like orientation, similar as we previously observed in mammalian prolegumain (Fig. 1B and Fig. 3A,B). Thereby the AP is blocking substrate access, keeping the proenzyme in a latent, inactive state. Additionally, we observed a conserved Gln346 (AtLEGß numbering) on the N-terminal end of the α6 helix. Gln346 is binding into the S1 pocket in an unproductive orientation and thereby preventing cleavage of the AP and further blocking substrate access to the active site (Figs. 1B and 3B). This interaction was similarly observed in the crystal structure of A.thaliana legumain isoform y; additionally, Gln346 is conserved throughout the plant VPE sequences, strongly suggesting that the Gln346-S1 binding forms a conserved mechanism in plant legumain activity regulation. Additionally, this interaction is further strengthened by the neighboring Arg347, which forms ionic interactions to Glu212, directly next to the catalytic Cys211 (Fig 3C).

Similar to two-chain AtLEG γ and mammalian prolegumains, the LSAM domain is further stabilized by two conserved disulfide bonds (Figs. 1B, and 3B). On the C-terminal end of the LSAM domain, AtLEG β harbours a potential VSS (vacuolar sorting signal), which is however not structured and therefore not visible in the electron density (Fig. 1A).

Activation proceeds via two-chain intermediate state

In an effort to unravel the basic principles of proAtLEGβ activation, we analyzed the interdomain interfaces of AEP and LSAM domains. Interestingly we found that the interface has a hydrophobic character with only two salt bridges identified by PDBe Pisa, R347-E212 and K422-D187, which are also conserved in proAtLEGγ (R355γ-E220γ and K432γ-D195γ; Fig. 4A). This is in stark contrast to proAtLEGγ where the interdomain interface has a mixed charged-hydrophobic character, which is reflected by eight interdomain salt bridges and a hydrophobic cluster localized to the prime substrate binding sites (Fig. 4B). Interestingly, the conserved D358γ-R74γ, (D348β -R66β) and D358γ-H177γ (D348β -H169β)

form salt bridges in proAtLEG γ , but not in proAtLEG β due to a local reorientation of the $\alpha 6$ helix. The residues involved in other AtLEG γ -specific interdomain salt bridges are not conserved in AtLEG β , i.e. $K365^{\gamma}$ -E109 $^{\gamma}$ (M357 $^{\beta}$ -L101 $^{\beta}$), R375 $^{\gamma}$ -E109 $^{\gamma}$ (K367 $^{\beta}$ -L101 $^{\beta}$), R375 $^{\gamma}$ -E264 $^{\gamma}$ (K367 $^{\beta}$ -I256 $^{\beta}$) and R490 $^{\gamma}$ -D136 $^{\gamma}$ (L482 $^{\beta}$ -S129 $^{\beta}$). Combined with the differences in oligomerization state, these findings led to the hypothesis that there will be pronounced differences in the activation and pH-stability profiles of the two *A.thaliana* legumain isoforms.

Since the interaction between the catalytic domain and the LSAM domain in proAtLEGβ is primarily hydrophobic in nature, we expected that its activation would be rather independent of pH. Surprisingly, an SDS-PAGE based, pH-dependent activation assay uncovered that the activation profile of AtLEGB closely resembles that of mammalian legumain, with complete activation only occurring at very acidic pH (4.0) (Fig. 5A). Consequently, we hypothesized that autocatalytic activation requires conditions that will destabilize the LSAM domain in order to gain accessibility to the active site. Indeed, we found complete degradation of the LSAM domain at pH \leq 4.0, indirectly indicating that the LSAM domain is destabilized at acidic pH conditions (Fig. 5A). Interestingly, upon incubation at pH proAtLEGB was split into catalytic (AEP) and LSAM domain. However, the LSAM domain was not degraded but remained stable on SDS-PAGE. This suggested to us that AtLEGB might form a two-chain state, where cleavage between LSAM and catalytic domain occurred but both domains remained bound to each other. To test this, we performed SEC experiments using proAtLEGB activated at pH 5.0. Indeed, we found a mixture of two-chain state and isolated AEP domain (Fig. S3). Importantly, there was no dimeric two-chain intermediate state of AtLEGB observed in SEC after activation.

Proteolytic activation is initiated by cleavages in the AP

Using mass spectrometry, we could identify two main autocatalytic cleavage sites, Asn333 and Asn345 on the AP (Fig. 1). These sites were similarly observed in proAtLEG γ and seem to be equally accessible to cleavage. Upon incubation at pH < 5.0, we observed additional cleavage sites on the LSAM domain, including Asp363, Asp416 and

Asp417 (Fig. 1B). Due to the architecture of the S1pocket, cleavage after Asp is restricted to low pH conditions (< 5.0), in line with the observed cleavage pattern. Interestingly, Asp363 is localized between the α 6- and α 7-helices and could, in combination with processing at Asn333/345, therefore allow the selective release of the α 6-helix (fragment Gln346-Asp363) as observed in mammalian legumain (31). Asp416 and Asp417 are localized within the V⁴¹⁵DDW⁴¹⁸ motif, right before the α 9-helix (Fig. 1B and S1). This motif is conserved within plant legumains and cleavage within this sequence was previously shown to be critical for the autocatalytic activation of castor bean legumain (33). Taken together, activation of At LEG β at pH < 5.0 goes along with cleavage at the aforementioned Asn and Asp sites, which finally results in the complete removal of the AP (including the α 6-helix) and the LSAM domain. Thereby rendering the active site accessible for substrates.

In addition to cleavage on the AP and LSAM domain, we observed another processing at the Nterminal end of the protein. Here it is important to note that our proAtLEG expression constructs typically carry an N-terminal His6-tag followed by a TEV-recognition site (ENLYFQG; TEV: tobacco etch virus protease). We found that AtLEGB was capable of cleaving after the Asn residue within the TEV-recognition site and thereby removing the His6-tag, as evidenced by a Western-blot using an Anti-His-Antibody (Fig. 5B,C). Based on SDS-PAGE experiments we propose that the primary cleavage at the Asn333/345 cleavage site can be catalyzed by the two-chain form. However, since N-terminal processing within the TEV-recognition motive was only observed at very acidic pH conditions we suggest that the latter cleavage is performed by the fully activated AtLEGβ. Importantly, N-terminal cleavage is not a physiological event as the relevant sequence is not present in native proAtLEGβ (Fig. S1).

(pro)AtLEGβ is stable at intermediate pH

Based on the remarkable variances we observed at the AEP – LSAM interfaces of proAtLEG β and γ we hypothesized that they would translate into differences of their pH-stability profiles. Indeed, when we measured the thermal stability of proAtLEG β using differential scanning fluorimetry we found a stability optimum at pH 5.0 (Fig. 4C). This is very different to proAtLEG γ and

mammalian legumain where the stability optimum of the proenzyme is at neutral pH (21,32). Even more interestingly, we found that AtLEGB activated at pH 4.0 and thereby lacking the LSAM domain, similarly showed a maximum in pHstability at pH 5.0 (Fig. 4D). This is in stark contrast to AtLEGy and also mammalian legumain, where the AEP domain is most stable at pH ~4. However, this difference becomes clear considering the hydrophobic interaction between AEP and LSAM domain in proAtLEGB. Mammalian legumain and AtLEGy harbor a highly charged electrostatic stability switch (ESS) on the AEP surface, located at the area surrounding the active site (32). At neutral pH conditions, the ESS causes electrostatic destabilization of the isolated AEP domain because of the high negative charge density which is not compensated by the LSAM domain. In human legumain and AtLEGy the isolated AEP can be stabilized by protonation of the excess acidic residues, hence the maximum stability at pH 4. The AEP in AtLEGβ lacks the pronounced ESS, explaining why a strong acidic pH is not necessary for charge neutralization, in agreement with the pH optimum at 5.0. The interaction of the AEP with LSAM generally stabilizes the protein. In proAtLEGB, the AEP-LSAM interaction and stabilization do not depend on neutral pH, whereas the tight electrostatic clamping of these domains in proAtLEGy and human prolegumain depend on neutral pH. Consequently, proAtLEGB is most stable at the pH which is also favorable for the isolated AEPβ.

Overall topology of AEP domain is highly conserved

Previous studies showed that the AEP domain in prolegumain is present already in an active conformation (21,34). Zymogenicity resulted solely from the steric blockage of the active site by the AP and LSAM domain. Therefore, we can use the crystal structure of proAtLEGβ to also analyze the active AtLEGß state. When we superimposed the AEP domains of AtLEGB and y we found that their fold is highly conserved (Fig. 6A). AtLEGB exhibits a caspase-like topology, i.e. a 6-stranded central β-sheet that is surrounded by 5 major αhelices (Figs. S1, S4) (35). Furthermore, AtLEGβ harbours the c341- and c381-loops, which form the non-prime substrate binding sites. The c341-loop encodes a plant VPE specific disulfide bond that is stabilizing the proline-rich insertion that is

extending the c341-loop compared to mammalian legumain (Fig. 6B). Mutation of Cys244 or Cys258 resulted in a complete loss of protein expression, confirming that the disulfide is also critical for folding. Furthermore, we observed 2 cis-imide peptide bonds (Thr180-Pro181 and Asn248-Pro249) with relevance for stabile bend and turn formation (Fig. 3B) (36). Interestingly, both turns are located to the substrate binding sites. The Asn248-Pro249 cis-peptide bond is on the c341-loop (non-prime side) and presents the Asn248 carbonyl oxygen as main chain recognition site for the P4 amide. Thr180-Pro181 is part of the eastern rim of the S2' pocket.

AtLEGβ has a wide S3-S4 pocket

When looking into the active site, we found that also the active site residues Cys211, His168 and Asn64 superimpose very well with the related AtLEGy (Fig. 6B). Furthermore, the residues forming the S1-specificity pocket, Arg66, His67, E209 and D261 adopt identical conformations as observed in AtLEGy. The highly conserved architecture of the active site suggested similar substrate specificity and catalytic activity of AtLEGβ and γ. However, when we compared the catalytic activity towards the fluorogenic Ala-Ala-Asn-AMC substrate, we observed a surprisingly low catalytic activity for AtLEGβ as compared to γ (Fig. 6C). Since the positioning of the active site residues were basically identical in β and γ , we did not expect this difference in activity to originate from a kcat effect, but rather from differences in substrate affinity (K_M). Beyond the highly similar S1-pocket, we indeed identified major differences on the c341- and c381-loops on the non-prime side (Fig. 6B and S5). Variations in sequence and conformation resulted in a narrow S3 – S4 pocket in AtLEGy but a rather wide pocket in AtLEGB (Fig. 4A,B, Fig. 6 and S5). To test whether these differences were a result of induced fit of the YVAD-cmk inhibitor, we superposed the crystal structures of proAtLEGβ, two-chain (pro)AtLEGγ and active YVAD-AtLEGy and compared their active sites. Interestingly, we found that the conformations of the substrate specificity loops c341 and c381 of proAtLEGB most closely resembled the active state of AtLEGy. Thereby we could exclude that induced fit was a main regulator of substrate affinity (Fig. S5). However, the situation might be different in AtLEGy, where we observed pronounced conformational changes of

the c381-loop between the proenzyme and the YVAD-cmk inhibited form. Modeling a peptidic substrate, based on the YVAD-cmk-AtLEGy crystal structure, we found tight interactions in AtLEGγ but less interactions to AtLEGβ. While AtLEGB offered an open, broad surface to accommodate the YVAD substrate, AtLEGy was tightly embracing the peptidic substrate as visible in Fig. 4A,B and 6B. We could assign $Tyr240^{\beta}/Trp248^{\gamma}$ on the c341-loop and Gly 300^{β} /Tyr 307^{γ} on the c381-loop as the main determinants for this difference. Together, this suggested to us, that small peptidic substrates would bind with lower affinity to AtLEGB as compared to gamma, because of missing enzymesubstrate interactions. Indeed, when we determined K_M values for AtLEG β and γ towards the AAN-AMC substrate we found high affinity binding (K_M = 57 \pm 3 μ M) to AtLEGy but low affinity for AtLEGB ($K_M = 337 \pm 3 \mu M$) (Fig. 6D). Importantly, we found similar k_{cat} (AtLEGβ: 4.5 x 10⁻³ min⁻¹ and AtLEGy: $6.3 \times 10^{-3} \text{ min}^{-1}$) and V_{max} values (AtLEGβ: 0.9 x 10⁻³ μmol/min and AtLEGγ: 1.1 x 10⁻³ µmol/min) for both enzymes. These findings confirmed that the difference in catalytic activity between AtLEGβ and γ was explained by differences in substrate affinity. Interestingly, when we used a VAN-AMC substrate instead of AAN-AMC we observed a reduction in enzymatic activity both for AtLEG β and γ (Fig. 6C). Accordingly, the smaller alanine is preferred over the branched valine at the P3 position in both AtLEG isoforms. Furthermore, we found an activity optimum for AAN-AMC turnover at pH 5.5 which is also in agreement with the pH-stability requirements of the AEP-domain (Fig. S6).

c381-loop is variable in length and sequence

Together, these observations made us hypothesize that the c341- and c381-loop might serve as $K_{\rm M}$ -switch. To analyze this further, we superposed all plant legumain structures available in the PDB. While the main structural elements superimposed very well in all available structures, we observed big differences on the c381-loops. It is variable in length, sequence and may even contain a glycosylation site (Fig. 6E and S4). Together these findings suggested that the c381-loop is a main determinant of the proteolytic activity of legumains, similar to caspases. The relevance of the c381-loop for legumain activity is further supported

by a previous analysis suggesting it as a marker of ligase activity (MLA) (28).

AtLEGβ substrate specificity is pH-dependent

To further analyse the substrate specificity of AtLEGβ, we carried out PICS experiments, which use proteome-derived peptides as substrate libraries (37,38). Here we used a peptide library that was generated from an E.coli proteome by digestion with trypsin for AtLEG specificity profiling at three different pH conditions. As expected, we observed a strong preference for Asn in P1 position at all investigated pH values (Fig. 7A). Interestingly, we also observed an increasing frequency of cleavage at Asp residues upon prolonged incubation times (18h). This time-dependence illustrates that substrates with Asn in P1 are kinetically favored over Asp. The substrate preference was also pHdependent, i.e. the turnover rate of P1-Asp substrates increased with lower pH values, which nicely agrees with the bi-polar architecture of the S1-specificity pocket and with previously published data for human legumain (Fig. 6B) (32).

AtLEGβ has a strong preference for hydrophobic residues in P2'

Furthermore, we observed a slight preference for small, polar residues in P1' position which was especially visible at the shorter incubation times (Fig. 7A), suggesting that P1'-Gly is kinetically preferred. Additionally, we found a pronounced preference for Leu in P2' position. Leucine has previously been proven to be beneficial at P2' position in legumain ligase substrates (16). Together, these results are in nice agreement with the architecture of the S1' and S2' binding sites: While the S1' binding site is flat and not allowing much interaction with the enzyme, the S2'-binding site forms a pronounced pocket (Fig. 4A) (29). Small residues in P1' position will facilitate the simultaneous binding of the P1 and P2' residues into the respective S1 and S2' binding pockets while still maintaining enough flexibility to allow efficient cleavage of the scissile peptide bond. The bottom of the S2' pocket is formed by Gly176 and the eastern wall by His182 (Fig. 7B). Gly176 is conserved in all plant legumains that have been structurally characterized so far (Fig. 7C). The eastern wall is mostly histidine and tyrosine, with some exceptions. Interestingly, mammalian legumain harbors a valine at position 176, making the S2'-pocket shallower and thereby less specific at this position (39). Furthermore, AtLEG δ has the glycine replaced by alanine (Fig. 7C), suggesting that it will also have a less pronounced specificity at P2' position. To test the relevance of His182 for prime side substrate specificity, we repeated the PICS experiments using AtLEG γ , which has a tyrosine at the equivalent position (Fig. S7). Interestingly, we found highly similar preferences on the non-prime and prime substrate binding sites, further confirming that Gly176 is the main determinant at the S2' site.

AtLEGβ has a strong preference for small residues in P1' position in protein substrates

In a next step we analyzed the substrate specificity of AtLEGB towards protein substrates, using proteome extracts isolated under non-denaturing conditions from leaves of the A. thaliana vpe0 mutant lacking expression of all four VPE isoforms as a substrate library. After incubation with recombinant AtLEGβ, recombinant AtLEGγ or buffer control, free N-terminal α-amines where labeled with three different formaldehyde isotopologues, and cleavage sites determined using the HUNTER N-termini enrichment and mass spectrometry (40). Based on three biological replicates we identified 381 N-terminal peptides significantly accumulating after incubation with AtLEGB at pH 6.0 (Figure 8A,B Supplementary Table 1), matching to 363 unique cleavage sites (Fig. 8C) in 289 proteins (Fig. 8D). As expected, we found a pronounced preference for Asn at P1 position (Fig. 8B). Furthermore, we observed a stronger preference for small and polar residues in P1' position, suggesting that the accessibility of the scissile peptide bond is enhanced when it is flanked by a small residue. Additionally, we also noticed a slightly increased preference for the more bulky and charged Asp and Glu amino acids. As in the peptide-based PICS experiment, we again observed a preference for hydrophobic amino acids in P2' position. For AtLEGy, identified significantly we 412 accumulating N-terminal peptides (Fig. Supplementary Table 1). These matched 390 unique cleavage sites (Fig. 8C) in 304 proteins (Fig. 8D) that exhibited a very similar cleavage profile in line with our observations using peptide substrates (Fig. 8F). Notably, a vast majority of 313 of the cleavage sites in 257 proteins were cut by both enzymes, while only 50 cleavages in 32 proteins were strongly preferred substrates of AtLEGβ and 77 cleavages sites of 47 proteins were selectively cut by AtLEG γ (Fig. 8C,D and G).

AtLEGβ is a broad spectrum transpeptidase

To characterize the cyclase activity of AtLEGβ we co-incubated it with different SFTI-derived linear peptides and measured the formation of the cyclic product using mass spectrometry. Indeed, we found that AtLEGB could cleave the SFTI-GL precursor peptide to the linear L-SFTI (lacking GL) version and further cyclize it to cyclic SFTI (c-SFTI) (Fig. 9A). Cyclisation worked most efficient at pH 6.0, which is in agreement with the previously reported pH-requirements of legumain ligase activity (29,41). Using the SFTI-GL precursor peptide, which harbors an Asp at P1 position, we observed a product formation rate of about 60%. This is less than compared to AtLEGy, which resulted in approx. 80% product formation (29). Interestingly, when the P1 residue was replaced by Asn, as is the case in SFTI(N14)-GL, AtLEGB was still able to catalyze peptide cyclisation, contrasting the situation of AtLEGy. When we replaced the P1'-P2' Gly-Leu by His-Val residues, which is the preferred sequence found for butelase-1 (C. ternatea legumain), we observed a similar cyclisation efficiency (Fig. S8) (16). Showing us that albeit optimized for butelase-1 the HVdipeptide is not facilitating peptide ligation in AtLEGβ.

AtLEGB is a broad spectrum ligase

Along that line we also tested whether AtLEGB would be able to cyclize linear L-SFTI and L-SFTI(N14) peptides, which are lacking amino acids on P1' and P2' positions of the protease substrate. Surprisingly, AtLEGB was indeed able to join the free termini and form the cyclic product, suggesting that AtLEGβ is not only a transpeptidase but a real ligase (Fig. 9A and B). Using the SFTI peptides carrying Asn at P1 position (N14), cyclisation worked equally well with or without preceding cleavage of prime side residues. In case of Asp at P1 position, transpeptidation (cleavage-linked ligation) was preferred as compared to joining free ends. Again, product formation was pH dependent, working best at near neutral pH conditions. So far, there was not a single report of a (plant) legumain capable of efficiently linking free peptide termini.

Discussion

Dimerization is a critical regulatory event for caspase-like proteins. In case of the apoptotic caspases, dimerization is mediated primarily by the β6 strand on the catalytic domain and is associated with structural rearrangements that render the caspase active (Fig. S1). Similarly, dimerization was also observed in plant legumains. The crystal structures of OaAEP1 (pdb entry 5hoi) and AtLEGy (5nij) both showed a dimer state that was mediated by the $\alpha 6$ and $\alpha 7$ helices on the LSAM domain. However, in these cases dimerization was not associated with activation but rather with inactivation. Under conditions where dimerization is maintained, such as high protein concentration, the proenzyme will not auto-process to the active AEP form. Additionally, there is a two-chain intermediate state, which is active to some extent. In this study we show for the first time that there are isoform specific differences in the activation and activity regulation of A.thaliana legumains. Firstly, we observed that proAtLEGB is monomeric in solution. In this respect, autocatalytic activation of proAtLEGβ resembles more the mechanism known from mammalian legumain, which also lacks a stable, latency-conferring dimer state (Fig. 10). We should point out, however, that in the crystal we found six equivalent proAtLEGB dimers per asymmetric unit. Nonetheless, this atypical dimer interaction is transient and short-lived, hence could not be observed in solution experiments. Secondly, we found that the AEP - LSAM interface is rather hydrophobic and not charged in nature. Consequently, the stability profile of AtLEGB differs from AtLEGy and mammalian legumains (Fig. 10). Thirdly, AtLEGB encodes autocatalytic cleavage sites on both ends of the α6-helix (Asn345 and Asp363), which in principle allows the selective removal of the AP, like in mammalian legumain (32). While N-terminal cleavage was observed at pH < 6.0, cleavage on the C-terminal end of the α 6-helix is restricted to pH < 5.0, which is in agreement with the charge requirements of the S1 pocket (Fig. 5A and 6B). Additionally, at acidic pH the ionic clamp that is linking the N-terminal end of the α 6-helix (Arg347) to the active site (Glu212), will loosen (Fig. 3C), which will further facilitate the release of the AP (α 6-helix). Therefore, an AEP-LSAM complex might represent a critical intermediate state, which initiates the complete removal of the LSAM

domain by proteolytic degradation and/or conformational destabilization. However, as we did not observe a stable AEP–LSAM complex in our experiments, it will only be short lived (Fig. 5 and 10).

These unique characteristics provide another new regulatory mechanism distinct from that of AtLEG γ . Different oligomerisation states will cause AtLEG β to favor activation at high local concentrations but will favor the latent two-chain state in AtLEG γ . On the other hand, the transient dimers observed in the AtLEG β crystal might possibly play a role in cooperative substrate processing. Together these findings suggest that AtLEG β and γ represent examples of two distinct classes of plant legumains, not only concerning their physiological function but also with regard to completely different mechanisms of zymogenicity, activation and stability.

All plant legumains are specific for cleaving after P1-Asn. However, we could show that subtle differences in the non-prime substrate binding sites translate into pronounced kinetic differences. Consequently, different legumain isoforms will feature kinetically-driven substrate preferences, which may be modulated by the amount and time of substrate availability (Fig. 7A). We provide evidence that the c381-loop can encode such kinetic differences. The corresponding sequences and significantly in plant conformations differ legumains, making it the single most variable region within the plant legumain catalytic domain. Differences in substrate affinity (K_M) can be kinetically assayed using specific substrates. PICS assays with proteome-derived peptide libraries are typically insensitive to such differences due to the mixed and unknown concentration of individual peptide substrates. However, if the substrate affinity is extremely different, such preferences can become apparent. Indeed, using time-series experiments we show that P1-Asp is a low affinity legumain substrate at increasing pH values. Presenting a P1-Asp may consequently serve as a strategy to kinetically regulate substrate turnover, i.e. to release a certain cleavage product in a slow, and pH-controlled manner. An example includes the autocatalytic activation of proAtLEGB which critically depends on cleavage after Asp residues on the LSAM domain and which is thereby restricted to low pH. Together this indicates that the differences in the c381-loop among the plant legumains will have an impact on cleavage kinetics rather than on sequence specificity. In line with these observations, we found mostly overlapping AtLEG β and γ cleavage sites in protein substrates in vitro.

Previously the c381-loop was described as marker of ligase activity (28). More precisely, a deletion in that region was associated with an increase in ligase activity. However, both AtLEGB and y encode relatively long c381-loops, yet both are active ligases. Furthermore, we could show that not only the sequence but also the conformation of this loop can be quite different, although it might be similar in length (Fig. S4). Therefore, we suggest that the c381-loop is primarily a determinant of protease activity. Since protease and ligase activities are inversely coupled, the c381-loop may be an indirect marker of ligase activity: If the affinity of the nonprime (protease) substrate is low, the affinity of the prime-side ligase substrate might be relatively high comparison. Such a situation favors transpeptidation over substrate hydrolysis. Furthermore, low affinity of non-prime substrates may also result in less re-cleavage of cyclic products, and thereby again indirectly favor ligation. This hypothesis also fits to our observation that AtLEGB, which has a non-prime binding site optimized for low affinity binding, is a ligase with broad substrate specificity.

In general, we found that SFTI-derived peptides harboring Asp at P1 position are better ligase substrates resulting in most efficient formation of cyclic product. This observation fits with the notion that poor (high K_M) non-prime substrates are more likely to find a prime ligase substrate at the active site, which in turn excludes the catalytic water molecule from the active site. In concert, the poor non-prime affinity should substrate favor aminolysis of prime substrate over hydrolysis – by the catalytic water that is excluded from the active site. For P1-Asp substrates this is particularly true at near neutral pH, where ligation is favored (Fig. 10). Additionally, the residence time of the ligation product is very short, making re-cleavage of the cyclic product unfavorable and consequently indirectly stabilizing the cyclic product. However, L-SFTI, which lacks P1' and P2' residues, resulted in less formation of cyclic product, indicating that P1-Asp will only be tolerated as a substrate at near neutral pH if coupled to prime side amino acids. P1-Asn as a free C-terminal end worked better, probably because Asn is in general a better K_M substrate at pH 6.0. Likely, the K_M will also be

influenced by prime side residues. As a result, a substrate with P1-Asp linked to prime side amino acids will have a critically superior (lower) K_M as compared to C-terminally free Asp, giving the P1-Asp substrate the possibility for binding and transpeptidation to happen. Looking at the prime substrate binding sites, we found that AtLEG β and γ encode nearly identical substrate binding sites. Taken together, differences in ligation efficiency between AtLEG β and γ might therefore be explained by their different non-prime substrate binding sites optimized for low and high affinity binding respectively.

In addition to the marker of ligase activity, Cys247 (O. affinis numbering) was identified as a gate keeper residue for ligase activity (20). Mutation to Ala247 resulted in an enzyme with superior ligase activity. Since all AtLEG isoforms harbor a glycine at the equivalent position (Gly241, AtLEGB numbering), this residue cannot explain the observed isoform specific differences in ligase activity. Similarly, the sequence motif Gly171-Pro172 (AtLEGβ numbering; Fig. S1) which is located close to the S1' pocket and was recently found to be beneficial for ligase activity is conserved in both A.thaliana legumains (30). However, directly next to Gly241 is Tyr240, which is a critical part of the non-prime binding site (S2 – S3) and which is different in AtLEGy (Trp248). Based on this observation and the differences in the close-by c381-loop, we suggest that it is rather the overall architecture of the non-prime substrate binding sites that affect substrate affinity and might positively affect ligase activity.

The ability of AtLEG β to join free ends is also interesting from a biotechnological point of view, as it will allow to link targets without the necessity of introducing artificial cleavage sites. However, still with the prerequisite of a P1-Asn or Asp. Furthermore, it also highlights that joining free termini is a general feature encoded in selected plant legumain isoforms. Given that all plants express a variety of different legumain isoforms, it is very likely that there is an AtLEG β -like enzyme present in every plant.

Previously we could show that the two-chain state observed in AtLEG γ is especially interesting with regard to ligase activity, as it is stable at neutral pH-environments where ligase activity is favored. Since two-chain AtLEG β has the same pH-stability profile as active AEP with a pH stability optimum at 5.0, two-chain AtLEG β will most likely not be a

superior ligase. However, it may implement differences in substrate specificity and catalytic efficacy. Indeed, we could previously demonstrate that human two-chain legumain with the C-terminal LSAM domain still present, exhibits carboxypeptidase activity rather than endopeptidase activity (32). The carboxypeptidase activity is structurally encoded by LSAM-derived arginine residues, which anchor the carboxy-terminus at the primed recognition site. Interestingly, we observed a slight preference for aspartate and glutamate residues in the P1' position of protein substrates (Fig. 8B) together with a relative depletion of basic residues (K, R), which could similarly indicate carboxypeptidase activity of two-chain AtLEGβ. However, this observation has to be taken with some caution, as the relative increase in specificity for Asp and Glu at the P1'-position was low. Finding out whether or not two-chain AtLEGB indeed harbours carboxy-peptidase activity will require further experiments and may be the subject of future studies.

Experimental procedures

Protein preparation

The Arabidopsis thaliana vacuolar processing enzyme (VPE, legumain) isoform β (AtLEGβ) fulllength clone U12200 (locus AT1G62710) was obtained from the Arabidopsis Biological Resource Center (ABRC). Using this as a template we subcloned an N-truncated variant missing the Nterminal signal sequences into the pLEXSY-sat2 (Jena Bioscience, Germany) vector using PCR amplification and XbaI and NotI restriction enzymes. The final expression construct carried an N-terminal signal sequence for secretory expression in the LEXSY supernatant and an Nterminal His6-tag followed by a TEV recognition site. Furthermore, we prepared a C211A dead mutant using the round-the-horn site directed mutagenesis technique, which is based on the inverse PCR method (42). Primers were designed which allowed the amplification of the cyclic plasmid template, harbouring the proAtLEGB wildtype insert, to a linear full-length PCR product carrying the desired mutation on one end of the PCR product. Following gel extraction of the PCR product and blunt end ligation, an intact plasmid carrying the desired mutation was generated and transformed into *E.coli* XI2(blue) cells. The C211A-mutant was used for crystallization experiments. Correctness of all constructs was confirmed by DNA sequencing. The so generated expression constructs were stably transfected into the LEXSY P10 host strain and stable cell lines were grown as described previously (21). Protein expression and purification was performed as described elsewhere (21,31). The final proAtLEG β protein was stored in a buffer composed of 20 mM Hepes pH 7.0 and 50 mM NaCl. ProAtLEG γ was prepared following the same protocol.

Crystallization, data collection and refinement

Initial screening was performed using the sittingdrop vapour-diffusion method utilizing a Hydra II Plus one liquid-handling system. Crystals of proAtLEGβ were obtained in a condition composed of 0.5 M ammonium sulfate, 1 M Lithium sulfate and 0.1 M trisodium citrate. Crystals grew within 2 weeks at a protein concentration of 10 mg/ml. To prevent autocatalytic activation we used a C211A dead mutant. Following pre-incubation in a cryoprotectant solution containing 0.8 M ammonium sulfate, 1.5 M lithium sulfate, 0.1 M trisodium citrate and 10% sucrose. Crystals were flash frozen in liquid nitrogen and subjected to Xray measurements. A high-resolution data set was collected at the ESRF on beamline ID30B. The beamline was equipped with a Pilatus 6M detector. Data collection was performed at a wavelength of 0.94 Å, 0.037 s exposure time and 15.3% transmission. 1,000 images were collected at an oscillation range of 0.1° and 100 K. Diffraction images were processed using xds and scala from the CCP4 program suite (43.44). An initial model could be obtained by molecular replacement using PHASER (45), using the crystal structure of twochain AtLEGy combined with the sequence of proAtLEGB. Following iterative cycles of model building in coot (46) and refinement in phenix (47) a final model was obtained and coordinates and structure factors were deposited to the PDB under the accession code 6YSA.

Electrostatic surface potentials were created with APBS (48) after assigning charges at pH 7.0 using Pdb2pqr (49). Surface potentials were contoured at +/-5 kT/e.

Autoactivation

To test the pH-dependence of auto-activation of proAtLEG β we incubated it at 0.4 mg/ml concentration in buffer composed of 100 mM buffer substance (pH 3.5 – 6.0: citric acid; pH 6.5: MES; pH 7.0: Hepes), 100 mM NaCl and 2 mM DTT for 1 h at 25 °C. Reactions were stopped by the addition of 10 mM MMTS (S-methyl methane thiosulfonate; Sigma-Aldrich) before subjecting them to SDS-PAGE.

To generate active AtLEG β in large scale we incubated the proenzyme in a buffer composed of 100 mM citric acid pH 4.0, 100 mM NaCl and 2 mM DTT at 25 °C for 1 h. Completion of autoactivation was analyzed by SDS-PAGE. Active AtLEG β was buffer exchanged using a NAP column (GE Healthcare) pre-equilibrated in a buffer composed of 20 mM citric acid pH 4.0 and 50 mM NaCl. Active AtLEG γ was prepared following the protocol described in (21).

Enzymatic activity assays

The enzymatic activity of active AtLEGB was investigated using the peptidic Z-Ala-Ala-Asn-7amino-4-methylcoumarin (Z-AAN-AMC; Bachem) and Z-Val-Ala-Asn-AMC (VAN-AMC) substrates. Activity was measured in assay buffer composed of 100 mM citric acid pH 5.5, 100 mM NaCl, 2 mM DTT and 100 µM of the respective substrate at 25 °C after adding the enzyme at 60 nM concentration. Assays were carried out in an infinite M200 plate reader (Tecan). Increase fluorescence was measured at 460 nm upon excitation at 380 nm. K_M values were determined upon incubation of AtLEGβ or γ with serial dilutions of the AAN-AMC substrate in assay buffer. Kinetic data was processed using GraphPad and K_M values were calculated using implemented algorithms.

Characterization of oligomerization state

To test the oligomerization state of proAtLEG β , 200 µl of sample were loaded on a S200 10/300 GL column (GE Healthcare) equilibrated in a buffer composed of 20 mM Hepes pH 7.5 and 100 mM NaCl. To test the oligomerization state of pH 5.0 activated AtLEG β , we loaded it on a S200 column pre-equilibrated in buffer composed of 20 mM citric acid pH 5.0 and 100 mM NaCl. BSA served as a size standard.

Determination of melting temperatures

To access the thermal stability of proAtLEG β and pH 4.0 activated AtLEG β we used the Thermofluor method. Experiments were setup as described previously (50). The investigated assay buffers were composed of 100 mM buffer substance (pH 4.0, 5.0, 6.0: citric acid; pH 7.0: Hepes) and 100 mM NaCl. Fluorescence data was analyzed as described elsewhere (51).

Western blot

Protein samples to be analysed were separated on an SDS-PAGE gel. Subsequently, proteins were blotted onto a Amersham Protran 0.45 NC membrane (GE Healthcare) using a Trans-Blot SD semi-dry transfer cell (Bio-Rad). The membrane was blocked with 1 x TBST supplemented with 5% (w/v) nonfat dry milk. Subsequently, the membrane was incubated with 5% milk-TBST supplemented with 1:10,000 (v/v) Anti-His-HRP antibody (ROTH). Chemiluminescent detection of Histagged protein was performed by using the Amersham ECL prime Western blotting detection reagent (GE Healthcare) together with an Odyssey Fc imaging system (Li-Cor).

Substrate specificity profiling

To test the substrate specificity of AtLEG β and $-\gamma$ we carried out Proteomic Identification of protease Cleavage Sites (PICS) assays using peptide libraries generated from Escherichia coli Bl21 cells (37,38). The peptide library was prepared as described previously (52). The proteome (2.2 mg/ml) was digested with trypsin at a ratio of 1:100 in 100 mM Hepes pH 7.5 overnight at 37 °C. The peptide library (2 mg/ml) was incubated with AtLEG proteases (10 µg/ml) in assay buffer composed of 50 mM buffer substance (pH 4.0 and 5.5: citric acid; pH 6.5: MES) and 100 mM NaCl at 25 °C. Samples were taken after 1 h and 18 h of incubation. Protease treated samples were stable isotope-labeled with 20 mM heavy formaldehyde (13CD₂O) and 20 mM sodium cyanoborohydride and control reactions with 20 mM light formaldehyde (CH2O) and 20 mM sodium cyanoborohydride for 2 h and quenched with 100 mM Tris pH 8.0 for 1h. Protease-treated and control samples were mixed and purified using C18 StageTips.

Substrate specificity profiling of AtLEG β and γ using intact A. thaliana leaf proteome

A. thaliana VPE quadruple mutant (VPE0, (53)) was obtained from the Nottingham Arabidopsis Stock center (accession N67918). Leaves were harvested from 8 week old plants grown on soil under short day conditions (9 h/15 h photoperiod, 22°C/ 18°C, 120 µmol photons m⁻² s⁻¹). The harvested leaves were homogenized with a Polytron PT-2500 homogenizer (Kinematica, Luzern, Switzerland) in extraction buffer containing 0.05 M MES pH 6.0, 0.15 M NaCl, 10% (w/v) sucrose, 0.01 M DTT and HALT protease inhibitor cocktail (ThermoFisher, Dreieich, Germany) on ice. The lysate was then filtered through Miracloth (Merck, Darmstadt, Germany), followed by centrifugation at 4000 x g at 4 °C, for 5 min. The protein concentration was determined by the Bradford assay using BSA as a reference.

Recombinant AtLEGB, recombinant AtLEGy or buffer control were added to the isolated proteome at a protease to proteome (1 mg) ratio of 1:100 (w/w) in the extraction buffer and incubated in parallel at 25 °C for 3 hours. The reactions were terminated by addition of 50 µM caspase-1 inhibitor (YVAD-cmk, Bachem, Switzerland). The reaction mixtures were purified by chloroformmethanol precipitation (54) and resuspended in 6 M GuaHCl, 0.1 M HEPES pH 7.5. The protein concentrations were determined using the BCA assay (ThermoFisher, Dreieich, Germany). The digested proteomes were reduced with 5 mM DTT at 56 °C for 30 min followed by alkylation with 15 mM iodoacetamide for 30 min at 25°C and quenched by addition of 15 mM DTT for 15 min. The three samples were differentially dimethyl labeled with 20 mM light formaldehyde (12CH₂O) and 20 mM sodium cyanoborohydride (light label), 20 mM medium formaldehyde (12CD2O) and 20 mM sodium cyanoborohydride (medium label) or 20 mM heavy formaldehyde (13CD₂O) and 20 mM sodium cyanoborodeuteride. After 16 hours incubation at 37 °C, the same amounts of fresh reagents were added and incubated for another 2 hours. The reactions were quenched with 0.1 M Tris (final concentration) at pH 7.4 and 37 °C for 1 hour. Equal amounts of protein were pooled, purified by chloroform-methanol precipitation resuspended in 0.1 M HEPES pH 7.4. The sample was then digested with trypsin in a 1:100

protease:protein ratio (SERVA Electrophoresis, Heidelberg, Germany) at 37 °C for 16 hours. Enrichment of N-terminal peptides was performed according to the HUNTER method (40). In brief, trypsin-digested sample was tagged with undecanal at a ratio of 50:1 (w/w) in 40% ethanol supplemented with 20 mM sodium cyanoborohydride at 50 °C for 45 min. Additional 20 mM sodium cyanoborohydride was added for another 45 min under the same condition. The reaction was then acidified with a final concentration of 1% TFA and centrifuged at 21,000 x g for 5 min. Next, the supernatant was injected through a pre-activated HR-X (M) cartridge (Macherey-Nagel, Düren, Germany). The flowthrough containing N-terminal peptides was collected. Remaining N-terminal peptides on the HR-X (M) cartridge were eluted with 40% ethanol containing 0.1% TFA, pooled with the first eluate and subsequently evaporated in the SpeedVac to a small volume suitable for C18 StageTip purification prior to mass spectrometric analysis. The assays were performed in three biological triplicates.

Mass spectrometry data acquisition

Samples were analyzed on a two-column nano-HPLC setup (Ultimate 3000 nano-RSLC system with Acclaim PepMap 100 C18, ID 75 µm, particle size 3 µm columns: a trap column of 2 cm length and the analytical column of 50 cm length, ThermoFisher) with a binary gradient from 5-32.5% B for 80 min (A: H₂O + 0.1% FA, B: ACN + 0.1% FA) and a total runtime of 2 h per sample coupled to a high resolution Q-TOF mass spectrometer (Impact II, Bruker) as described (55). Data was acquired with the Bruker HyStar Software (v3.2, Bruker Daltonics,) in line-mode in a mass range from 200-1500 m/z at an acquisition rate of 4 Hz. The Top17 most intense ions were selected for fragmentation with dynamic exclusion previously selected precursors for the next 30 sec unless intensity increased three-fold compared to the previous precursor spectrum. Intensitydependent fragmentation spectra were acquired between 5 Hz for low intensity precursor ions (> 500 cts) and 20 Hz for high intensity (> 25k cts) spectra. Fragment spectra were averaged from tstepped parameters, with 50% of the acquisition time manner with split parameters: 61 µs transfer time, 7 eV collision energy and a collision RF of 1500 Vpp followed by 100 µs transfer time, 9 eV collision energy and a collision RF of 1800 Vpp

Mass spectrometry data analysis

Acquired mass spectra were matched to peptide sequences at a FDR of 0.01 using MaxQuant (56) v.1.6.0.16 using standard Bruker OToF instrument settings. For PICS experiments, the UniProt E.coli K12 proteome database (downloaded Nov 2015, 4313 entries) with appended common contaminants was used. Search parameters considered semispecific tryptic peptides, light (+28.031300) and heavy (+36.075670) dimethyl labeling at peptide N-terminal or Lys side chain amines, Cys carbamidomethylation as fixed and Met oxidation as variable modification. Identified peptides that showed at least a fourfold increase in intensity after protease treatment compared to the control treatment or were exclusively present in the protease-treated condition were considered as putative cleavage products. An in-house Perl script was used to remove putative library peptides (trypsin specificity on both sides of the identified peptide) and to reconstruct the full cleavage windows from the identified cleavage products as described (38) and visualized as IceLogos using software version 1.3.8 (57).

For HUNTER experiments, the *A. thaliana* UniProt proteome database (downloaded Dec 2018, 41592 entries) with appended list of common laboratory contaminants was used for searches that considered C-terminal cleavage by ArgC as digestion enzyme. Further search parameters included isotope labeling by light (+28.031300), medium (+32.056407) or heavy (+36.075670) dimethylation of peptide N

termini or Lys residues, Cys carbamidomethylation as fixed and Met oxidation, N-terminal acetylation (+42.010565) or N-terminal pyroGlu formation from Glu (-18.010565) or Gln (-17.026549) as variable modifications. Further statistical data analysis, filtering and annotation was performed with the Perl script MANTI.pl 3.9.7 (https://manti.sourceforge.io).

Peptide cyclisation assay

SFTI-derived peptides were synthesized and analysed as described previously (29). Subsequently, cyclization experiments were carried out using 500 μM of the respective linear peptide and 0.5 μM AtLEGß in a buffer composed of 100 mM NaCl and 50 mM Tris, Bis-Tris, citric acid pH 4.0 or pH 6.0. Reactions were incubated at 30 °C for 12 h. Subsequently the reactions were desalted using ZipTip C18 tips (Merck Millipore) and analyzed by MALDI-TOF-MS (Autoflex, Bruker Daltonics, matrix: α -cyano-4-hydroxycin-namic acid).

Data availability: The coordinates and structure factors presented in this paper have been deposited with the Protein Data Bank (PDB) with the accession code 6YSA. MS data have been deposited with the PRIDE (https://www.ebi.ac.uk/pride/archive/) (58) repository with the accession codes PXD019220 for the PICS dataset and PXD19276 for the HUNTER N-terminome dataset. All remaining data are contained within the article.

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Conflict of Interest

The authors declare that they have no conflict of interest with the contents of this article.

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FOOTNOTES

The abbreviations used are: proAtLEGβ, *Arabidopsis thaliana* prolegumain isoform beta; AtLEGβ, *Arabidopsis thaliana* legumain isoform beta; AEP, asparaginyl endopeptidase; LSAM, Legumain Stabilization and Activity Modulation domain; AP, activation peptide; VPE, vacuolar processing enzyme; SFTI, sunflower trypsin inhibitor; cPRM, cyclic protein recognition motif; rmsd, root mean square deviation

Table 1. Xray data collection and refinement statistics

	proAtLEGβ (6ysa)
Data collection	
Space group	$P4_1$
Cell dimensions	
a=b, c (Å)	170.4, 196.5
Resolution (Å) ^a	49.6–2.0 (2.04–2.01)
$R_{ m merge}$	0.12 (1.42)
R_{pim}	0.08 (0.99)
CC (1/2) (%)	0.99 (0.22)
I/oI	6.8 (0.7)
Completeness (%)	90.2 (86.3)
Redundancy	2.8 (2.6)
Refinement	
Resolution (Å)	49.6–2.0
No. unique reflections	336594
$R_{ m work}$ / $R_{ m free}$	20.8/21.8
No. atoms	
Protein	39124
Ligand/ion	763
Water	2254
Overall B-factor (Å ²)	36.0
R.m.s deviations	
Bond lengths (Å)	0.01
Bond angles (°)	1.15
Ramachandran plot	
No. outliers (%)	0.0
No. favored (%)	97.9

The structure was determined from a single crystal.

The resolution cutoff was set by applying the CC1/2 criterion (59).

[[]a] Highest resolution shell is shown in parentheses.

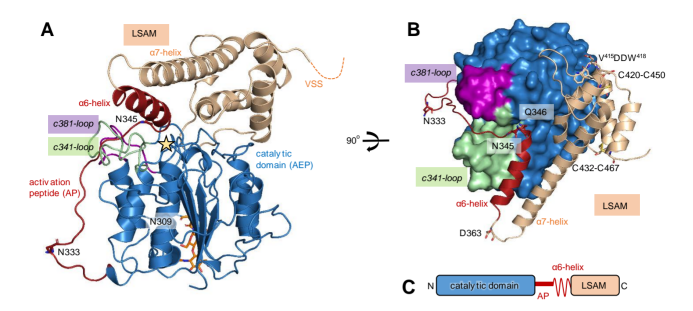


Figure 1. proAtLEGβ shares the typical prolegumain-like architecture. A. Cartoon representation of proAtLEGβ with the catalytic AEP domain shown in blue, the activation peptide harbouring the α 6-helix in red and the LSAM domain in beige. Asn333 and 345 autocatalytic cleavage sites and the Asn309 glycosylation site are indicated as sticks, an asterisk is labeling the active site, the C-terminal vacuolar sorting signal (VSS) is indicated by a dashed line. C341- and c381-specificity loops are colored green and purple respectively. **B.** Top-view on the active site in standard orientation (substrate binding from left to right). Gln346 (red sticks) on the AP binds to the S1-pocket. Disulfide bonds on the LSAM domain are shown as sticks. The autocatalytic processing sites Asp363 and Asp416 (within the V⁴¹⁵DDW⁴¹⁸ motif) are indicated. **C.** Schematic representation of proAtLEGβ domain architecture.

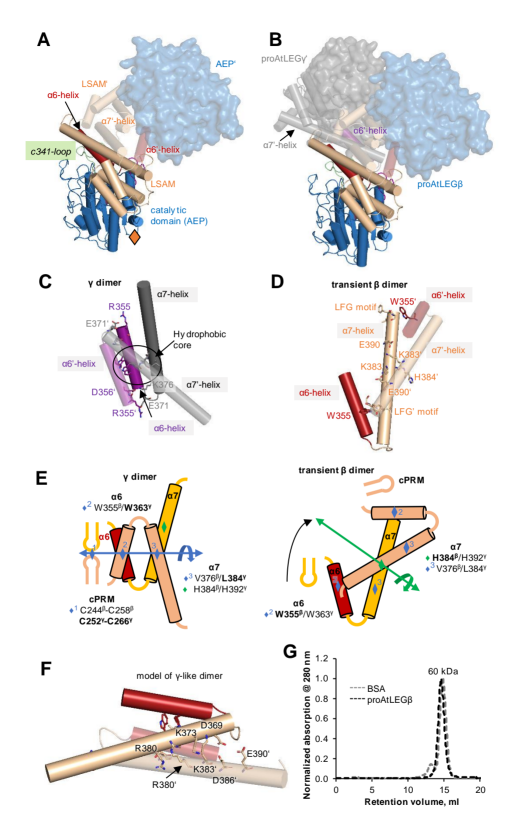


Figure 2. proAtLEGβ is monomeric in solution. A. Crystal packing induced proAtLEGβ dimerization. Monomer 1 is shown in cartoon representation, monomer 2 is labeled with a prime symbol (AEP' in surface representation). Interactions were mainly mediated by 2 symmetric salt bridges on the α 7-helices (LSAM domain). The location of the Asn309 glycosylation site is indicated with an orange diamond. **B.**

Superposition of A) (proAtLEG β dimer observed in the crystals) with dimeric two-chain AtLEG γ (pdb entry 5nij). Dimerization led to different spatial orientation of the AEP domains. C. Zoom-in view on the 4-helix bundle as observed in two-chain AtLEG γ . Interaction is mediated by a hydrophobic core that is surrounded by electrostatic interactions. D. Zoom-in view on the 4-helix bundle observed in proAtLEG β . Interaction is mediated by a symmetric E390 – K383 salt bridge localized on the α 7-helix and hydrophobic interactions between the LFG motif (Leu396 β -Gly398 β) on α 7 helix and W355' on the α 6' helix. Relative to B), the views in C) and D) are rotated by 90° along the y-axis. E. Schematic representation of the 4-helix bundle as observed in AtLEG γ and β . F. Model of an AtLEG γ -like dimerization mode in proAtLEG β . AtLEG γ -like dimerization is not favored because of electrostatic repulsion of R380 – R380', K373 – K383' and D369 – D386' – E390' pairs. G. Size exclusion runs confirming monomeric state of proAtLEG β . BSA served as a size standard.

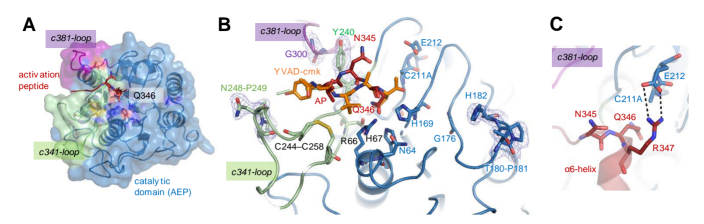


Figure 3. The activation peptide binds canonically to the active site. A. Top view on the active site of proAtLEG β . The activation peptide (AP) harbouring the autocatalytic Asn345 cleavage site and Gln346 that is occupying the S1-pocket are shown in red. B. Zoom-in view on the non-primed and primed substrate binding sites with a YVAD-cmk peptide modeled based on the crystal structure of the YVAD – AtLEG γ complex (pdb entry 5obt). cis-imide peptide bonds (Thr180-Pro181 and Asn248-Pro249) are shown as sticks. For selected residues a 2Fo-Fc composite omit map is displayed at a contour level of 1 σ. C. Zoom-in view on the active site of proAtLEG β . The ionic clamp (R347–E312) that is a linking the α6-helix to the active site is indicated.

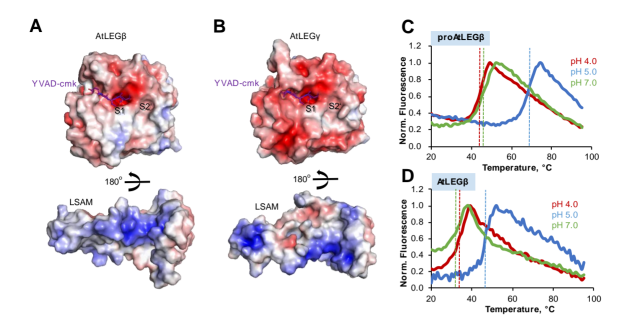


Figure 4. The AEP – LSAM interaction in proAtLEG β is mostly hydrophobic. A. Color-coded electrostatic surface potential of AtLEG β AEP and LSAM domains based on the crystal structure of proAtLEG β (blue: positive charge, red: negative charge) calculated at pH 7.0 and contoured at +/- 5 kT/e. The LSAM domain has been rotated by 180° relative to the AEP domain. The YVAD-cmk inhibitor has been modeled based on the crystals structure of the AtLEG γ inhibitor complex (pdb entry 50bt). B. same as A., but calculated for AtLEG γ in complex with YVAD-cmk inhibitor. C. Melting curves of proAtLEG β at indicated pH values show highest thermal stability at pH 5. Melting points are indicated by dashed lines. D. Melting curves of active AtLEG β showing highest stability at pH 5.0.

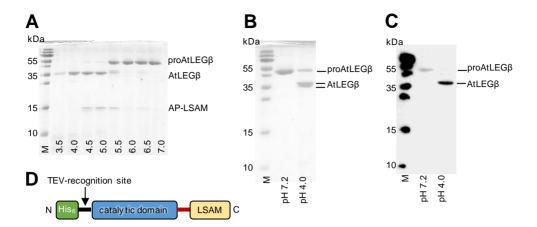


Figure 5. Auto-catalytic activation of AtLEG β is pH-dependent and results in a two-chain intermediate state (pH 5.0) and active AEP state (pH 4.0). A. ProAtLEG β after 1h incubation at indicated pH values. 'AtLEG β ' corresponds to the catalytic domain up to the autocatalytic cleavage site Asn333, and 'AP-LSAM' corresponds to the Gln346–Ala486 C-terminal fragment, that is generated by cleavage after Asn345). B. SDS-PAGE showing proAtLEG β (pH 7.2) and AtLEG β following activation at pH 4.0. Activation results in a double band at around 36 kDa. C. Western-blot using an Anti-His-HRP antibody, showing that only one AtLEG β activation product harbours the N-terminal His₆-tag. D. Scheme illustrating the domain architecture of the recombinant expression construct.

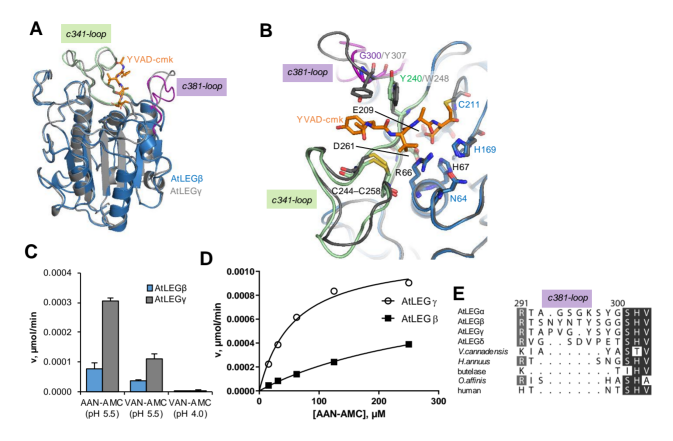


Figure 6. AtLEGs differ in their substrate specificity loops. A. Superposition of AtLEGβ (blue) and γ (grey) AEP domains. The YVAD-cmk inhibitor bound to AtLEGγ is shown in orange sticks, the c341-loop in green and the c381-loop in purple. **B.** Zoom in view on the active site. Catalytic residues are labeled in blue, residues forming the S1 specificity pocket in black. **C.** Catalytic activities of AtLEGβ and $-\gamma$ towards peptidic AAN-AMC and VAN-AMC substrates at indicated pH values. **D.** K_M-determination for AtLEGβ and $-\gamma$ towards the AAN-AMC substrate. **E.** Sequence alignment of the c381-loops of indicated (plant) species. Sequences were derived from structures deposited to the PDB, where applicable; AtLEGα (P49047), AtLEGβ (Q39044), AtLEGγ (5nij), AtLEGδ (Q9LJX8) *V.canadensis* (*Viola Canadensis*; 5zbi), *H.annuus* (*Helianthus annuus*; 6azt), butelase (*Clitoria ternatea*; 6dhi), *O.affinis* (*Oldenlandia affinis*; 5hoi).

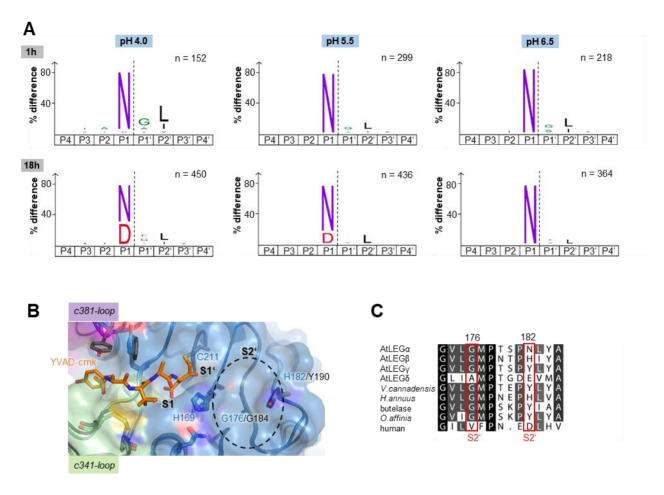


Figure 7. AtLEGβ has a pH-dependent substrate specificity. A. Cleavage site specificity determined by the PICS assay, using peptides generated by tryptic digest of an *E.coli* proteome as substrate library. iceLogos visualize the substrate preference surrounding the cleavage sites (p = 0.05) based on peptides cleaved by AtLEGβ after incubation at indicated pH values and times. Number of non-redundant cleavage sites used to generate the iceLogos are indicated. B. Top-view on the AtLEGβ substrate binding site. Binding of the YVAD-cmk inhibitor was modeled based on the crystal structure of the YVAD – AtLEGγ complex (pdb entry 50bt). C. Sequence alignment of the residues forming the prime-substrate binding site. Sequences used are the same as in Figure 5.

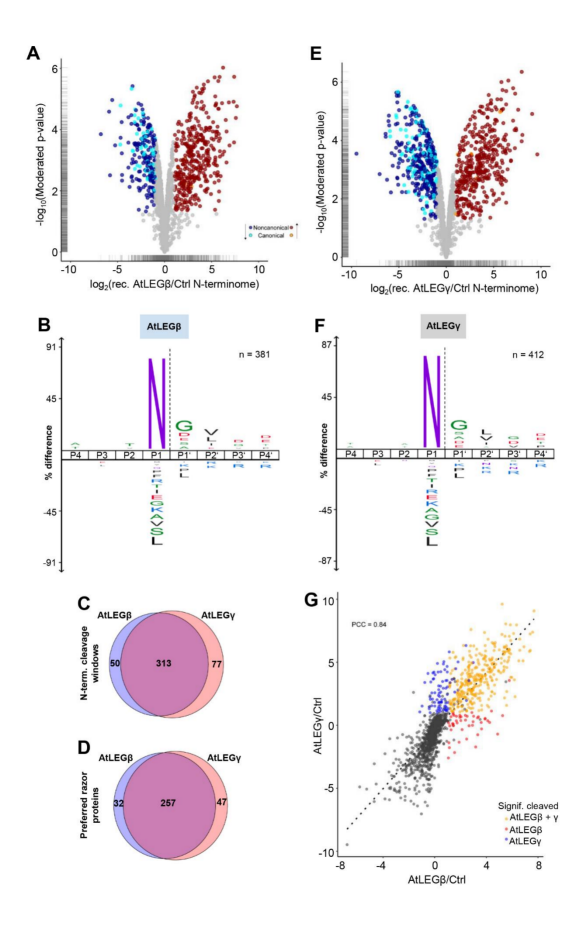


Figure 8. Substrate specificity of AtLEGβ and γ towards intact proteins extracted from A. thaliana leaves. Volcano plots identify protein N-terminal peptides significantly changing in abundance (greater 2-fold change in abundance supported by LIMMA-moderated t-test p-val <0.05) after in vitro incubation of A. thaliana vpe0 proteome with recombinant A) AtLEGβ or E) AtLEGγ. Log₂ fold-change is the mean of 3 biological replicates. Accumulating N-terminal peptides indicative of AtLEGβ/γ cleavage are highlighted red, depleted peptides cleaved within their sequence in blue. iceLogos visualize the substrate preference surrounding the cleavage sites for B) AtLEGβ and F) AtLEGγ (p = 0.05). Number of non-redundant cleavage sites used to generate the iceLogos are indicated. Venn diagrams showing the overlap of C) cleavage sites and D) proteins cleaved in the vpe0 proteome after incubation with AtLEGβ or γ. G) Correlation of N-terminal peptide abundance in both experiments (dimethylated N-terminal peptides quantified in at least 2 out of 3 replicates). Significantly accumulating dimethylated N-terminal peptides (log₂FC > 1, LIMMA-moderated t-test p-value < 0.05) indicate cleavage by AtLEGβ (red), AtLEGγ (blue), or both (orange). The linear fit confers a Pearson correlation coefficient (PCC) of 0.84 indicating a very high degree of overlap among the putative substrates.

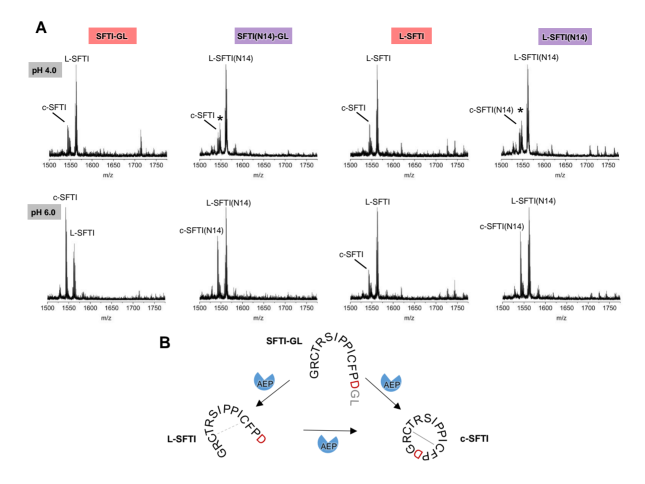


Figure 9. Cyclisation of SFTI-derived peptides by AtLEGβ is pH-dependent. A. Reactions were carried out at indicated pH values. An unidentified species is labeled with an asterisk. **B.** Reaction scheme of AtLEGβ catalyzed cyclisation of SFTI-GL peptides. The pre-cursor peptides SFTI-GL and SFTI(N14)-GL were synthesized in the reduced form, and were also observed mostly reduced in the assays. The linear L-SFTI and L-SFTI(N14) cleavage products were observed both in the reduced and oxidized forms, with the Cys3–Cys11 disulfide bond formed. c-SFTI and c-SFTI(N14) were mostly oxidized.

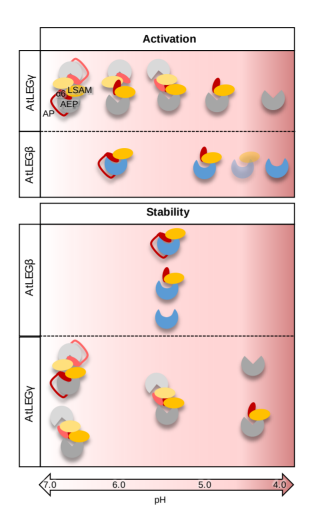


Figure 10. Activation and stability of AtLEGβ and AtLEGγ are pH dependent. In the zymogen forms of proAtLEGβ and γ the LSAM domain (orange) and activation peptide (AP, red line) that harbors the α 6-helix (red ellipsoid) sit on top of the active site and thereby block access to the substrate binding sites. ProAtLEGγ forms a dimer at neutral to intermediate pH conditions, and is mostly present in its two-chain state, which is generated upon cleavage at the N-terminal end of the α 6-helix. By lowering pH, the interaction of the α 6-helix with the AEP domain gets weaker, as it is mainly mediated by electrostatic interactions. At pH < 4.5, the two-chain state will disassemble and thereby allow degradation of the α 6-helix and the LSAM domain. By contrast, proAtLEGβ is a monomer in solution. Activation proceeds via cleavage after (1) Asn333/345 on the N-terminal end of the α 6-helix at intermediate pH, followed by (2) multiple cleavages after aspartic acid residues at pH < 4.5, which finally result in AP-LSAM degradation. Activation likely proceeds via a short-lived intermediate state, that has the α 6-helix selectively removed, but the LSAM domain still bound to the AEP domain (indicated by transparent coloring). While all AtLEGβ activation states show highest conformational stability at intermediate pH, two-chain (pro)AtLEGγ is stable at neutral to slightly acidic pH and monomeric two-chain AtLEGγ as well as the AEP domain are most stable at acidic pH.