### **REVIEW**

# **Plant Proteases Involved in Regulated Cell Death**

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**Abstract**—Each plant genome encodes hundreds of proteolytic enzymes. These enzymes can be divided into five distinct classes: cysteine-, serine-, aspartic-, threonine-, and metalloproteinases. Despite the differences in their structural proper ties and activities, members of all of these classes in plants are involved in the processes of regulated cell death – a basic fea ture of eukaryotic organisms. Regulated cell death in plants is an indispensable mechanism supporting plant development, survival, stress responses, and defense against pathogens. This review summarizes recent advances in studies of plant pro teolytic enzymes functioning in the initiation and execution of distinct types of regulated cell death.

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The influence of external stimuli, both mechanical and physicochemical, often leads to uncontrolled death of living cells. This type of cell death is generally referred to as "accidental cell death". However, in many cases cell death is genetically programmed, although this program can be cancelled by means of specific genetic or pharma cological factors. Such "regulated cell death" (RCD) can occur as a part of physiological programs or can be acti vated at the level of the whole organism by intracellular events or some extracellular factors. Mechanisms of reg ulated cell death can also be activated in response to external stimuli. According to the latest recommenda tions formulated by the Nomenclature Committee on Cell Death (NCCD), cellular death occurring as a part of physiologic program in ontogenesis (i.e. a particular case of RCD) should be referred to as programmed cell death (PCD) [1].

The term "programmed cell death" was first pro posed by Richard Lockshin in the mid-1960s [2]. Intensive studies of PCD phenomenon during last 50 years has led to the discovery of different types of cell death and different mechanisms involved. Such a number of mechanisms necessitated the development of a special

classification of cell death types. A classification based on morphological features distinguishing main types of cell death was proposed ten years ago [3]. However, it soon became clear that morphological features alone were not sufficient for exhaustive classification of all types of cell death. Therefore, it was suggested to complement mor phological features with the unique molecular and bio chemical features specific for certain types of RCD [4]. One of the major molecular markers of cell death is the activity of proteolytic enzymes, which control the initia tion and play a role in execution of RCD [4, 5].

Studies of RCD phenomenon in plants have not been as intensive as studies of this process in human and animal cells. Since information about molecular mecha nisms of RCD in plants was lacking, the classification of various types of RCD in plants was based only on mor phological features [6]. Studies of RCD in plants revealed some similarities with these processes in animals, but also pointed out some differences [6]. One of the main differ ences was the absence of caspases in plants, while in ani mals these enzymes are key players in initiation and mediation of apoptosis [5]. At the same time, in plants there are many other proteolytic enzymes involved in var ious stages of RCD, including enzymes with caspase-like activities and caspase-like functions.

In plants, as in animals, RCD can be induced by both biotic and abiotic factors. Different pathogens (viruses, bacteria, or fungi) can trigger RCD, in a form of hypersensitive response, for example, which is a protec-

*Abbreviations*: ER, endoplasmic reticulum; NCCD, Nomen clature Committee on Cell Death; PCD, programmed cell death; PS-SCL, positional scanning substrate combinatorial library; RCD, regulated cell death; VPE, vacuolar processing enzyme.

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Plant proteolytic enzymes involved in RCD

Protease type	Family	Localization	RCD inducer	Activate RCD $(\uparrow)$ or inhibit RCD $(\downarrow)$	Reference
Cysteine proteases					
<b>VPE</b>	C13	vacuole	biotic factors abiotic factors developmental		$[22, 23, 30-33]$ $[13-15, 34-38]$ $[39-43]$
Metacaspases, type I	C14	cytoplasm, nucleus	biotic factors abiotic factors developmental	$\begin{array}{c}\uparrow,\downarrow\\ \hline \downarrow\\ \hline \downarrow \end{array}$	[59, 60] [60]
Metacaspases, type II	C14	cytoplasm, nucleus, apoplast, protoplasts of dead cells	biotic factors abiotic factors developmental	↑↑↑	$[61-64]$ [64, 65] [47, 57, 66, 67]
CEP1-like proteases	C1	ER, ricinosomes, apoplast (?)	biotic factors abiotic factors developmental	↑ $\frac{1}{\uparrow}$	$[99]$ $[79, 81, 101 - 106]$
RD21A- and XBCP3-like proteases	C1	ER derivatives, vacuole, cytoplasm	biotic factors abiotic factors developmental	↑	$[110]$ [111]
XCP2-like proteases	C1	vacuole, protoplasts of dead cells	biotic factors abiotic factors developmental	↑	$[17]$
CTB3(CathB)-like proteases	C1	vacuole, apoplast	biotic factors abiotic factors developmental	↑	[116, 117] [116]
RD19A-like proteases	C1	vacuole, nucleus	biotic factors abiotic factors developmental	$\bar{\uparrow}$	$[118]$ [119]
SAG12-like proteases	C1	peripheral vacuoles	biotic factors abiotic factors developmental	$\uparrow, \downarrow$	$[122]$ [120, 122]
Autophagins (ATG4)	C54	cytoplasm	biotic factors abiotic factors developmental	$\uparrow$ (?) $\hat{\Gamma}(?)$ $\uparrow$ (?)	
Serine proteases					
Subtilisin-like proteases	S8	cytoplasm, apoplast	biotic factors abiotic factors developmental	$\uparrow$ $\overline{\phantom{0}}$	[129, 143] $[142]$
Aspartic proteases					
Atypical proteases	A1	ER	biotic factors abiotic factors developmental	$\uparrow, \downarrow$	$[148-151]$
Threonine proteases					
Proteasome components	T <sub>1</sub>	cytoplasm, nucleus	biotic factors abiotic factors developmental	↑ ↑	[155] [154]
Metalloproteinases					
Matrix metalloproteinases	M10	plasma membrane (?), apoplast (?), ER(?)	biotic factors abiotic factors developmental	$\uparrow$	$[159]$

tive reaction of a plant. In this case, cell death can be induced either by elicitors – compounds specific to a pathogen, recognized by the immune system of a plant – or by pathogen-specific protein products, whose expres sion in a plant cell causes ER stress leading to the devel opment of RCD [7-10]. Among the abiotic factors trig gering the development of RCD in plants are metal ions, oxidative stress, salt stress, UV radiation, and heat shock [11-15]. At the same time, some cells undergo RCD to ensure normal development of a plant. For example, such phenomenon can be observed during the development of xylem [16, 17].

In this review, we summarize our knowledge about proteolytic enzymes involved in regulation and execution of regulated cell death, caused by biotic and abiotic fac tors, or necessary for normal plant development (table). Proteolytic enzymes involved in particular types of RCD associated with senescence of plant organs will be dis cussed less extensively.

#### CYSTEINE PROTEASES

**Vacuolar processing enzymes (VPE).** The detection of caspase-1-like proteolytic activity in dying cells during the development of the hypersensitive response of tobac co to the infection induced by tobacco mosaic virus [18] stimulated active search for a plant enzyme with such properties. Meanwhile, other examples of caspase-like activities in plants were revealed by studies of different types of RCD [19]. However, sequencing of the whole genomes of *Arabidopsis thaliana* L. and rice *Oryza sativa* L. [20, 21] showed in plants there are no caspase genes that can be identified by means of simple homology search. Therefore, some groups focused their attention on plant proteolytic enzymes that possess caspase-like activ ities without being caspase orthologs.

The first identified protease with caspase-1 like activity that is involved in RCD is a vacuolar processing enzyme (VPE) [22, 23]. A VPE is a legumain-like cys teine protease that belongs to the C13 family of the CD clan according to the classification of proteolytic enzymes represented in the MEROPS peptidase database [24]. Like many proteases, VPE is translated as an inactive zymogene containing *N*- and *C*-terminal propeptides that are autocatalytically cleaved during activation of the enzyme. The *N*-terminal fragment contains a signal pep tide that guides VPE into the vacuole, where the autocat alytic processing takes place [25, 26].

Homologs of VPEs are widespread throughout the plant kingdom; they are found both in mosses and ferns, as well as in higher plants. A VPE homolog known as asparagine endopeptidase (AEP) [27, 28] is also found in animals. There are four VPE genes in the genome of *A. thaliana*: α*VPE*, β*VPE*, γ*VPE*, and σ*VPE*. Expression of <sup>α</sup>*VPE* and γ*VPE* is detected in vegetative organs of the

plants, while β*VPE* is expressed in embryos, and σ*VPE* is expressed during the formation of the seed coat [27, 28].

Collapse of the vacuole induced by a VPE is consid ered one of the key factors in RCD in plants. A plant VPE was shown to be involved in different types of RCD, including RCD induced by biotic and abiotic factors. Moreover, VPEs play a role in RCD indispensable for the normal development of a plant [28]. A VPE is necessary for the development of a hypersensitive response induced by tobacco mosaic virus in tobacco plants bearing a resist ance gene *N*, as it stimulates degradation of a vacuole, DNA fragmentation, and the development of a necrotic reaction [22, 29]. A VPE is also involved in morphologi cally similar mechanisms of RCD induced by other virus es, fungi, bacteria, and their toxins [23, 28, 30-32]. It should be noted that VPEs also take part in the develop ment of a special type of RCD induced by ER stress, which is the result of interaction between cells of *A. thaliana* with the fungus *Piriformospora indica* [33]. VPEs also play a role in the development of RCD induced by a number of abiotic factors, such as thermal shock [15], salt stress [14, 34], oxidative stress [34, 35], UV radiation [13], and metals [36-38]. Moreover, they are players in the mechanisms of RCD supporting ontogenesis and aging, such as formation of seed coat [39], death of peri carp and nucellus cells in the ovule [40, 41], and senes cence of leaves [42] and petals [43].

In spite of the fact that VPEs are key players in the development of some types of cell death in plants, sub strates of these enzymes remain largely unknown. Therefore, the conclusion that VPEs in plants have the same functions as caspases in animal cells is premature.

**Metacaspases.** Only special bioinformatic tools allowed researchers to find in plant genomes the very dis tant relatives of caspases – metacaspases [44]. Publication of these data stimulated studies of plant metacaspases with the aim to show their functional simi larities with caspases. However, soon it became clear that substrate specificity of metacaspases differs dramatically from that of caspases: metacaspases appear to be arginyl/lysyl-specific endopeptidases, while caspases are aspartate-specific proteolytic enzymes [45-47].

Metacaspases are cysteine proteases belonging to peptidase family C14 of the CD clan according to the classification of proteolytic enzymes in the MEROPS database [24]. Metacaspases, similarly to caspases, con sist of a large (p20) and a small (p10) subunit. As in cas pases, catalytic His and Cys residues in metacaspases are located in the p20 subunit, while the p10 subunit takes part in the formation of a substrate-binding pocket [48, 49]. Plant metacaspases are subdivided into two main types. Type I metacaspases often contain an additional *N* terminal proline-rich prodomain with a zinc finger motif. Type II metacaspases never have such *N*-terminal prodomain and, unlike type I metacaspases, the p20 and p10 subunits of type II enzymes are separated by a linker

sequence [48, 50]. Metacaspases found in some phyto plankton species were united into type III. Unlike type I and type II metacaspases, the p10 domain of type III metacaspases is located closer to the *N*-terminus of the protein, while the p20 domain is closer to the *C*-terminus [51].

Plant genomes generally contain about ten genes coding metacaspases, although in some species the num ber of genes can reach twenty [50]. There are nine genes in the *A. thaliana* genome: three genes coding type I metacaspases (*AtMC1-AtMC3*) and six genes coding type II metacaspases (*AtMC4-AtMC9*). All nine genes are expressed in various tissues of the plant [44, 48].

Maturation of metacaspases, similarly to maturation of many others cysteine endopeptidases, is a result of an autocatalytic cleavage of the zymogen. However, auto proteolytic hydrolysis of metacaspases is a  $Ca^{2+}$ -dependent process, with rare exceptions [45, 47, 52]. Moreover, as shown for the metacaspase AtMC4, during  $Ca^{2+}$ induced autocatalytic proteolysis the cleavage sites can vary slightly depending on the conditions of the reaction. This probably reflects the existence of a special mecha nism that can increase or decrease protease activity at the early stage of its activation [53, 54]. Nitric oxide (NO) can also regulate the activity of metacaspases. It was shown that the catalytic Cys residue of metacaspase AtMC9 could be S-nitrosylated, which can be the result of excess NO and can lead to inhibition of the autopro cessing of the enzyme [55].

Intracellular localization of metacaspases can slight ly vary depending on the enzyme, but these proteases are generally localized in the cytoplasm and/or in the nucle us [56]. Moreover, the intracellular localization of an enzyme can change during the development of RCD, as shown, for example, for the Norway spruce mcII-Pa metacaspase: at initial stages of RCD, the enzyme was found in the cytoplasm, while at later stages – in the nucleus [47]. AtMC9 metacaspase of *A. thaliana* can be found not only in the nucleus and cytoplasm, but also in the apoplast [57, 58].

Plant metacaspases were shown to be key players and regulators of RCD processes induced by different biotic and abiotic factors and in RCD supporting normal plant development. It was shown that type I metacaspase AtMC1 of *A. thaliana* participates in the activation of hypersensitive response of the plant to infection by the oomycete *Hyaloperonospora arabidopsidis* or bacterium *Pseudomonas syringae*, while another type I metacaspase AtMC2 was acting as an antagonist of AtMC1 [59]. Although AtMC2 blocked the development of RCD, it should be noted that in this case this function of the enzyme was not related to its proteolytic activity [59, 60]. Later, it became clear that AtMC1 also can interfere with the development of RCD; a pro-survival role of AtMC1 metacaspase during aging was shown [60]. Expression of the metacaspase-9 gene (CaMC9) in *Capsicum annuum*

L. is increased in response to infection with *Xanthomonas campestris*. Silencing of the CaMC9 gene inhibits the development of pathogen-triggered RCD, while the over expression if this gene, in contrast, increases the proba bility of cell death [61]. Metacaspases TaMC4 in wheat and NbMCA1 in *Nicotiana benthamiana* L. also partici pate in the development of RCD as protective mecha nisms against infection of wheat with the fungus *Puccinia striiformis* or infection of tobacco plants with the fungus *Colletotrichum destructivum* and bacterium *P. syringae* [62, 63]. Plants with knockout of the type II metacaspase AtMC4 gene showed reduced sensitivity to RCD-induc ing mycotoxin fumonisin B1 and other oxidative stress inducers, while overexpression of the AtMC4 gene, in contrast, increased plant sensitivity towards these factors [64]. Overexpression of the metacaspase AtMC8 gene stimulated RCD induced by UV irradiation or  $H_2O_2$  in protoplasts, while silencing of the AtMC8 gene inhibited it [65]. Norway spruce metacaspase mcII-Pa induces autophagy, which triggers RCD mechanisms during the terminal differentiation of embryonic suspensor cells, and, on the other hand, participates in further develop ment of RCD [47, 66]. Being localized in extracellular space, AtMC9 metacaspase can be considered as an effector of RCD activation, as it cleaves a peptide of 11 amino acids from GRI protein, which, once bound to the membrane receptor PRK5, initiates RCD [67]. Moreover, it turned out that one of the functions of the secreted AtMC9 metacaspase is degradation of xylem cell content after the collapse of the central vacuole. Thus, we conclude that in this particular case the functional role of the enzyme is manifested after the fact of cell death [57]. Is should be mentioned that the genome of *A. thaliana* contains a gene coding for the protein inhibitor of AtMC9 metacaspase – *AtSerpin1*, which is at the same time a sub strate of AtMC9, since before the formation of a covalent bond with an enzyme, an irreversible inhibitor from the serpin family has to be cleaved by a protease [68]. However, the physiological role of such inhibition of AtMC9 in RCD is not yet clear.

Besides AtSerpin1, several dozens of other substrates of plant metacaspase AtMC9 were identified [58]. However, the functional role of proteolytic cleavage cat alyzed by AtMC9 *in vivo* was demonstrated only for one of them – phosphoenolpyruvate carboxykinase 1 (PEPCK1), which is one of the key enzymes involved in gluconeogenesis in plants. It was shown that processing of PEPCK1 catalyzed by AtMC9 led to increase in enzyme activity [58]. Unfortunately, analysis of identified sub strates of metacaspase AtMC9 does not allow making any conclusions about RCD mechanisms that involve AtMC9.

Studies of RCD during the terminal differentiation of embryonic suspensor cells in Norway spruce showed that physiological substrate of mcII-Pa metacaspase was the evolutionarily conserved multifunctional protein

Tudor-SN [69]. Genes coding for Tudor-SN protein were found not only in plants, but also in animals including humans. The protein itself, being a component of many ribonucleoprotein complexes, is involved in a number of functional processes related to transcription, splicing, RNA interference, RNA editing, and RNA degradation [70-74]. Interestingly, during apoptosis in human cells Tudor-SN was recognized as a substrate by caspase-3 [69]. Both in plants and in animals, proteolytic cleavage of Tudor-SN led to its inactivation [69]. It was shown recently that not only plant metacaspases have common substrates with caspases from animal cells, but also meta caspases from fungi. Thus, PaMCA1 metacaspase of *Podospora anserina* is able to cleave *in vivo* a classical cas pase substrate – PARP [75]. These data relaunched dis cussion of whether metacaspases can be considered as caspases. A number of researchers suppose that metacas pases can be considered as functional analogs of caspases, and RCD mechanisms are so conserved that common features can be found in RCD processes in different king doms of living organisms.

**Papain-like enzymes.** According to the mechanism of catalysis, papain-like endopeptidases are cysteine pro teases. Taking into account their phylogenetic features, papain-like C1A proteases (C1 family of CA clan) were placed with enzymes similar to L-, B-, H-, and F-cathep sins from animals [24, 76].

Papain-like proteases are relatively stable proteins. They are often found in rather harsh environment, such as the apoplast, the vacuole, and lysosomes [77]. These proteases are globular proteins, composed of two domains, both of which are involved in substrate binding pocket formation. This pocket is able to bind substrates, and the radicals of the catalytic triad composed of Cys, His, and Asn residues are located in this pocket [78].

Activity and specificity of plant C1A peptidases *in vitro* were thoroughly studied by means of various tests using proteins, fluorogenic synthetic peptide substrates, and peptide inhibitors. Moreover, new bioinformatics methods allowing modeling of molecular interactions between endopeptidases and their substrates are being developed. Data accumulated so far enable us to conclude that plant papain-like proteases have rather low specifici ty. Nevertheless, studies of substrate specificity of these peptidases showed that their preferential substrate should have a nonpolar (including Pro) or aromatic amino acid residue in P2 position, and in some cases, it can be an Arg residue [79-83].

To be targeted to a specific intercellular compart ment, protease precursors carry a signal peptide, whereas the presence of an autoinhibitory prodomain prevents premature activation of the enzyme [80, 84]. Upon trans lation, the polypeptide chain of the inactive proenzyme enters the ER lumen; the majority of C1A proteases is addressed through *trans*-Golgi into the vacuole, lyso somes, or is secreted into the apoplast [76]. However,

some C1A proteases carrying a *C*-terminal signal for retention in the endoplasmic reticulum K/HDEL can be addressed into other specific compartments, such as rici nosomes, which are derivatives of the endoplasmic retic ulum (ER) [79, 85, 86]. Further cleavage of the prodomain can occur *in cis* via intramolecular interac tions or *in trans* via intermolecular interactions. Moreover, some data show that the prodomain can be cleaved by other proteases, which suggests that papain like proteolytic enzymes can be a part of proteolytic cas cades [80, 87].

A peculiar feature of plant papain-like proteases when compared to other proteolytic enzymes involved in RCD is that some of these enzymes can be reversibly inhibited by natural peptides, for example, by plant cys tatins. Plant cystatins can interact with members of the papain-like C1A family of cysteine proteases; phytocys tatins were placed in a separate subfamily within the cys tatin family [24, 76]. Generally, plant cystatins are rather small proteins with molecular mass of 12-16 kDa. Nevertheless, high molecular weight phytocystatins with molecular mass of 85-87 kDa and containing several cys tatin domains were also found [88-91]. Moreover, *in vitro* experiments showed that for inhibition of legumain-like C13 peptidases, phytocystatins should carry additional *C* terminal extensions, which increase the molecular mass of the protein up to 23 kDa [92, 93]. As mentioned earli er, VPEs belong to the C13 family of cysteine peptidases, although there are no data so far that VPE activity can be regulated by protein inhibitors. Another peptide inhibitor of papain-like C1A cysteine proteases is a member of the serpin family, AtSerpin1, which is able to inhibit RD21A protease [94, 95].

Preproenzymes of plant papain-like proteases always consist of an *N*-terminal signal peptide, a prodomain, and a proteolytic domain containing the catalytic triad Cys- His-Asn. Especially for plants, an additional classifica tion of papain-like proteases was proposed [80] that takes into account structural particularities of plant enzymes, such as an additional *C*-terminal granulin domain, pro line-rich domain, *C*-terminal ER-retention signal (K/HDEL), vacuole localization signal NPIR in the beginning of the prodomain, and some others [79, 80, 96]. According to this classification, there are nine sub families of papain-like proteases in plants [80]. Data obtained so far have shown that members of at least seven subfamilies (CEP1-, RD19A-, RD21A-, XCP2-, XBCP3-, SAG12- and CTB3-like proteases) are impli cated in the development of various types of RCD. However, there is no evidence of any special role of papain-like proteases belonging to the AALP- and THI1 like subfamilies in the development of RCD.

*CEP1-like proteases.* Members of CEP1-like subfam ily of proteases are characterized by the presence of the *C* terminal ER-retention signal KDEL. Interestingly, genes coding for homologs of KDEL-containing plant proteases were not found in the yeast genome or in animals [80, 97]. Proteases carrying ER-retention signal are often addressed into special structures, ricinosomes, which are the derivatives of ER but separated from it. Ricinosomes do not merge with the central vacuole, but as RCD is ini tiated they collapse and release proteases and hydrolases they contain [81, 85, 98]. Under biotic stress, CEP1 pro tease in *A. thaliana* is accumulated in ER and is probably further released in the apoplast [99].

Many members of CEP1-like protease subfamily were shown to participate in the development of RCD in various plants [100, 101]. The best-studied members of this subfamily are the CEP1, CEP2, and CEP3 proteases of *A. thaliana*, which are expressed in roots, stems, flow ers, and green siliques [97]. Under biotic stress conditions, expression of CEP1 was also observed in leaves [99]. Participation of this protease in the development of regu lated death of epidermal cells, caused by the infection with the ascomycete *Erysiphe cruciferarum* was studied in detail [99], as well as the role of CEP1 protease in RCD mecha nisms during maturation of pollen in tapetum, a special layer that covers sporangia and anthers [101]. CEP2 pro tease was shown to participate in the development of RCD during the formation of a root cap [81]. Moreover, in other plant species KDEL-containing proteases participate in RCD processes involved in petal senescence in daylily (*Hemerocallis* sp.) [102], in orchid seed coat formation in *Phalaenopsis* sp. [103] and *Jatropha curcas* L. [104], in RCD in the endosperm of *Ricinus communis* L. [79], tomatoes [105], and *J. curcas* [104], as well as in the anther of tomatoes [106]. It should be emphasized that some KDEL-containing endopeptidases were able to cleave the hydroxyproline-rich glycoprotein extensin, which is a component of the plant cell wall [97].

Thus, a large experimental dataset has accumulated so far, showing the role of proteolytic CEP1-like enzymes in different types of RCD, including RCD mechanisms supporting normal plant development and providing defense against the pathogens.

*RD21A- and XBCP3-like proteases.* Proteolytic enzymes belonging to the RD21A- and XBCP3-like pro tease subfamilies (members of these two subfamilies differ mainly in primary structures of the prodomains) are char acterized by the possible presence of an additional *C*-ter minal fragment composed of a proline-rich domain fol lowed by a granulin-like domain [80, 107]. In animals, granulins play a role as extracellular growth factors. They are expressed as pro-granulin, which comprises several copies of cysteine-rich granulin modules [108]. In plants, granulins were found only within proenzymes of some papain-like proteases. The function of granulins in plants remains unknown [80].

Sorting of *A. thaliana* RD21 protease occurs through the Golgi apparatus, where the protein is fucosylated, then the enzyme is accumulated in compartments derived from ER, and after fusion of these compartments with the vacuole the enzyme stays in the vacuole [87, 107, 109, 110]. It is generally thought that cleavage of the prodomain of RD21 *in vivo* can be catalyzed by one or several other proteases, which point out the possible implication of RD21 in proteolytic cascades. However, granulin-like domain cleavage occurs by autocatalysis [87]. Interestingly, RD21 protease purified from plants can be activated with SDS. This might indicate that RD21 protease is accumulated *in vivo* as a complex with a reversible endogenous inhibitor [87]. Moreover, it was shown that activity of RD21 could be inhibited by the irreversible endogenous inhibitor AtSerpin1, which belongs to the serpin family  $-$  a family of peptide inhibitors of proteases [94]. Within cells, AtSerpin1 is localized in the cytoplasm, so probably the inhibition of RD21 by AtSerpin1 can occur during translocation of the enzyme to the cytoplasm [110]. Elicitors of RCD in plants, such as benzothiadiazole (a salicylic acid agonist) or oxalic acid (a toxin of pathogenic fungi, such as *Botrytis cinerea* and *Sclerotinia sclerotiorum*), stimulate changes in the permeability of vacuole membranes, which can lead to the translocation of RD21 into the cytoplasm with formation of inactive RD21-AtSerpin1 complexes [110]. If the level of RD21 expression is decreased, or the level of AtSerpin1 expression is increased, development of elicitor-triggered RCD slows considerably, which led to the conclusion that RD21 is a stimulator of RCD, while AtSerpin1 reduces the activity of this effector [110].

NtCP14 protease from tobacco is closer to XBCP3 protease than to RD21A from the point of view of phy logeny. Nevertheless, it was shown that NtCP14 protease along with its inhibitor cystatin NtCYS has similar func tion to that of RD21 protease and its inhibitor AtSerpin1 [110, 111]. It was shown that in the majority of cells of tobacco embryos, NtCP14 protease is not active because it forms a complex with NtCYS, while in embryonic sus pensor cells, where RCD starts to develop, the level of NtCYS decreases, leading to the appearance of NtCP14 protease molecules that are not bound to NtCYS and therefore active. The presence of active NtCP14 protease ensures further development of regulated death processes of suspensor cells [111].

Thus, we conclude that plant proteolytic enzymes bearing a granulin sequence within the proenzyme together with their inhibitors play a regulatory role in dif ferent types of RCD induced either to ensure ontogenesis or to assure the activation of defense mechanisms. Involvement of these enzymes in proteolytic cascades accompanying RCD cannot be excluded.

*XCP2-like proteases.* One of the main properties of plant C1A proteases belonging to the XCP2-like protease subfamily is the presence of a conserved putative glycosy lation site in the protease domain [80]. Particularities of secondary structure of these proteolytic enzymes can be discussed since the crystal structure of papain, which also belongs to the XCP2-like protease subfamily, was defined

some time ago [112]. The genome of *A. thaliana* contains genes coding for two members of this subfamily: XCP1 and XCP2. Genes of these two proteases along with a gene coding for the subtilisin-like serine protease XSP1 were initially identified in a xylem cDNA library [113]. It was shown later that during xylogenesis, XCP1 and XCP2 are transported into the central vacuole of the cell, where they participate in micro-autolysis preceding the macro autolysis induced by tonoplast breakage [17, 114]. It is considered that after tonoplast breakage during formation of xylem tracheary elements, both enzymes continue fur ther degradation of cellular content [17]. The hypothesis of implication of XCP1 and XCP2 enzymes in providing defense from pathogens that live and propagate in plant xylem was suggested recently [115].

*CTB3(CathB)-like proteases.* The primary structure of CTB3-like proteases is close to that of human cathep sin B. Their distinctive feature is the presence of four additional disulfide bonds and a conservative putative gly cosylation site [80]. The genome of *A. thaliana* contains genes coding for three members of this subfamily: CTB1 (AtCathB1), CTB2 (AtCathB2), and CTB3 (AtCathB3) [80, 116]. The level of expression of these genes increases with senescence and during contacts with pathogens. Moreover, in mutant *A. thaliana* with deletion of all three genes of the CTB3-like proteases, significant delay in the development of senescence is observed, as well as the reduction of hypersensitive response induced by *P. syringae* [116]. In *N. benthamiana* plants, silencing of a CTB3-like protease (NbCathB) also leads to reduction in hypersensitive response to bacterial infections caused by *Erwinia amylovora* and *P. syringae* [117]. Moreover, NbCathB was found to be secreted into the apoplast, where it can be activated even in absence of pathogens [117]. Thus, one can conclude that plant cathepsin B-like proteolytic enzymes are implicated in processes related to RCD indispensable for ontogenesis, or for RCD provid ing resistance to pathogens.

*RD19A-like proteases.* RD19A-like proteases are characterized by the presence in the prodomain of an ERFNAQ motif and four additional cysteine residues, which are supposed to form disulfide bonds for additional stabilization of enzyme structure, as well as by the pres ence of a conserved putative glycosylation site [80]. As shown for *A. thaliana*, RD19 participates in biochemical pathways that assure resistance of the plant to bacterial infection caused by *Ralstonia solanacearum*. RD19 pro tease is translocated from the mobile structures associated with the vacuole into the nucleus [118]. However, direct implication of RD19 in RCD processes in *A. thaliana* has not been reported. On the other hand, data about the involvement of a close homolog of RD19, SmCP protease, in RCD processes in eggplant (*Solanum melongena* L.) was published. SmCP was also proposed to be implicated in RCD processes indispensable for ontogenesis, particularly during formation of xylem tracheary elements [119].

*SAG12-like proteases.* The majority of SAG12-like proteases contain an additional Cys residue before the cat alytic Cys residue (motif CGCCWAFS) [80]. AtSAG12 protease of *A. thaliana* (from senescence-associated genes) was identified as one of the products encoded by genes associated with senescence [120]. It was shown that this proteolytic enzyme is localized in special vacuoles that are found in peripheral cytoplasm separately from the central vacuole in aging cells of the mesophyll. However, the phenotype of *A. thaliana* with deleted AtSAG12 gene did not differ from that of the wild type [121]. At the same time, expression of the gene coding for a homologous pro tease in rice (*OsSAG12-1*) was induced not only by senes cence, but also during RCD supporting the response to a biotic stress [122]. Silencing of the *OsSAG12-1* gene in rice led to accelerated senescence and increased RCD in case of infection with bacterial pathogen *Xanthomonas oryzae*, compared to the wild type plants, suggesting that OsSAG12-1 is an RCD suppressor [122].

**Autophagins (ATG4).** Describing the key role of cys teine proteases in regulated cell death in plants, it is not possible to omit autophagins or Atg4 – cysteine proteases that are key factors initiating autophagosome formation [123, 124]. It is important to note that autophagy in plants can be observed during different types of RCD, both before and after the point of no return [125, 126]. Thus, key factors of autophagy, including Atg4, can be considered as legitimate players and regulators of RCD processes in plants.

Atg4 proteases belong to the C54 family of cysteine proteases from the CA clan [24, 123]. The primary struc ture of these enzymes does not contain signal peptide, which defines their cytoplasmic localization. The main function of these enzymes is the processing of Atg8 pro tein, which gains thereafter the ability to bind phos phatidylethanolamine and to subsequently initiate autophagosome formation [124]. Unlike yeast, bearing one copy of Atg4 and one copy of Atg8 genes, mammals and plants have several homologous genes of Atg4 and Atg8 [127, 128]. Two genes coding for Atg4 (*AtAtg4a* and *AtAtg4b*) and nine genes coding for Atg8 (*AtAtg8a*-*AtAtg8i*) were found in the genome of *A. thaliana* [128]. One can not exclude the possibility that two Atg4 genes of *A. thaliana* encode protein products with different functions. Indeed, *in vitro* experiments have shown that AtAtg4a demonstrated higher proteolytic activity than AtAtg4b, although the substrate specificity of the two enzymes was quite similar [128]. Such differences might have function al importance for regulation of a RCD program.

#### SERINE PROTEASES

Attempts to reveal the sources of caspase-like activi ty in plant cells during regulated cell death led to charac terization of one more proteolytic RCD-related enzyme.

It is a member of a subtilisin-like family of proteases named phytaspase after its source (from  $\varphi \circ \varphi$  – plant in Greek) and manifested activity (aspartate specific protease) [129]. Phytaspase was shown to participate in the development of hypersensitive response in *N. tabacum*, bearing a resistance gene *N*, after infection of plants with tobacco mosaic virus [129]. Moreover, activity of this enzyme was also detected in response to mechanical damage in mono- and dicotyledonous plants [130].

Subtilisin-like proteases, also called subtilases, are serine proteases. These enzymes are characterized by the presence of a catalytic triad composed of Asp, His, and Ser residues [131]. According to the MEROPS classifica tion of proteolytic enzymes, subtilisin-like enzymes belong to the S8 family, clan SB [80]. Plant genomes con tain a large number of genes coding for various subtilases. For example, in the genome of *A. thaliana* there are 56 genes coding for these enzymes [132, 133]. Subtilases, like many other proteases, are translated as inactive zymogens composed of a signal peptide, prodomain, and peptidase domain, in which the protease-associated domain is localized [133-135]. Secondary structure of subtilases should resemble that of tomato SlSBT3 subti lase that was solved [136]. In plants, subtilases have vari ous functions; some of them are related to the response to pathogen infection [135]. Substrate specificity for many of these enzymes is not very high [132, 133].

Unlike other characterized subtilases, phytaspases from rice and *A. thaliana* showed rather high specificity, recognizing, for example, caspase-6 classical cleavage site VEID. It turned out that phytaspases are also able to cleave a number of fluorescent peptide substrates cleaved by cas pases, such as YVAD, VAD, IETD, LEHD, and some oth ers. However, it should be noted that phytaspases were not able to cleave one classical caspase-3 substrate DEVD [129, 137], but cleaved with high efficiency IWLD peptide, which differs considerably from common caspase sub strates [138]. Study of rice phytaspase using the positional scanning substrate combinatorial library (PS-SCL) [139] showed that unlike caspases, this enzyme prefers substrates containing hydrophobic amino acid residues in positions P4-P2 [138]. Moreover, it was shown that phytaspase cleaves more efficiently full-length proteins, which proba bly signifies that interactions between phytaspases and its substrate are not limited to the P4-P1 positions [138].

Overexpression of the phytaspase gene increased sensitivity of plants towards inducers of RCD, while silencing of this gene inhibited RCD, suggesting that phy taspase actively participates in processes of cell death [129]. However, the most remarkable feature of phytaspases is their localization. It was shown that constitutive ly expressed phytaspase zymogen is processed and the mature enzyme is secreted to the apoplast. When RCD is induced by biotic or abiotic factors, active phytaspase is translocated back to the cytoplasm, where it starts to hydrolyze proteins [129, 140]. Unfortunately, only one

natural substrate of phytaspase – VirD2 protein from the plant pathogenic bacterium *Agrobacterium tumefaciens* – has been characterized so far. VirD2 is responsible for the delivery of a fragment of bacterial DNA in the nucleus of an infected cell. After being processed by phytaspase, VirD2 loses its *C*-terminal signal peptide, responsible for nuclear localization, which might be a part of the mech anism actively protecting the plant cell from undesirable transformation [141]. However, this process is not direct ly related to RCD. Thus, questions about natural phytas pase substrates that are processed during RCD and about functional consequences of this hydrolysis remain open.

It was shown that subtilisin-like AtSBT1.1 protease from *A. thaliana* is able to cleave the prepropeptide of the phytosulfokine AtPSK4 [142]. Active hormone produced upon this cleavage is able to stimulate differentiation of xylem tracheary elements [115]. In the cDNA library from xylem tissue of *Arabidopsis*, the gene of another sub tilisin-like protease XSP1 was found [113]. However, functions of proteolytic enzyme XSP1 have not been investigated yet.

At least two more putative subtilisin-like functional analogs of animal caspases were found in oats. Both enzymes, named saspases (SAS-1 and SAS-2), showed high substrate specificity similar to that of caspases [135, 143]. These proteases were shown to be secreted into intercellular space during the development of RCD induced by *Cochliobolus victoriae* toxin victorin and accompanied by DNA fragmentation and mitochondrial dysfunction [143]. However, their functional role in RCD was not characterized. Their natural substrates also remain unknown.

#### ASPARTIC PROTEASES

The genome of *A. thaliana* contains more than fifty genes coding for aspartic proteases [144]. Despite of this large number of genes (the human genome codes for only eight aspartic proteases [145] and the genome of *Caenorhabditis elegans* – only twelve [146]), functions of corresponding protein products generally remain unknown. According to MEROPS classification of prote olytic enzymes, proteases from families A1, A3, A11, and A12 of AA clan and proteases from A22 family of AD clan were found in plants [80]. Generally, plant aspartic pro teases are divided into three groups: typical aspartic pro teases (group A), nucellin-like (group B), and atypical proteases (group C) [144, 147]. Typical aspartic proteases from plants usually contain an additional *C*-terminal domain that is cleaved during enzyme maturation. Nucellin-like aspartic proteases are homologs of nucellin – a protease found in cells of the nucellus in bar ley. Atypical aspartic proteases demonstrate various inter mediate characteristics specific for typical and nucellin like proteolytic enzymes [144, 147].

Data showing involvement of aspartic proteases in RCD of plant cells is rather limited. Nevertheless, it was shown that expression of genes coding for two atypical proteases (OsAP25 and OsAP37) in tapetal cells of rice was regulated by transcription factor EAT1. Proteolytic enzymes OsAP25 and OsAP37 are involved in mecha nisms of regulated death of these cells [148]. Taking into account that expression of nucellin in barley is limited to nucellar cells in which RCD occurs, we can also hypoth esize involvement of this representative of aspartic pro teases in processes of cell death [149].

It should also be noted that among aspartic proteas es, similarly to papain-like proteases, proteolytic enzymes capable of inhibiting the development of RCD were found. For example, in *A. thaliana* atypical aspartic protease PCS1 carries an *N*-terminal serine-rich frag ment and is localized in ER. This enzyme can inhibit the development of RCD in some types of cells that normal ly die. Moreover, lack of expression of active PCS1 leads to massive RCD in gametophytes of both types, as well as during embryogenesis [150]. Another example from *A. thaliana* is the atypical protease UNDEAD, which can inhibit the development of programmed death of tapetal cells [151].

Thus, it was shown that some aspartic proteases par ticipate in regulation of at least some types of RCD need ed for normal ontogenesis. Moreover, these enzymes can act both as positive and negative regulators of RCD.

#### THREONINE PROTEASES

In eukaryotes, 26S proteasome is the main proteolyt ic component of the ubiquitin-dependent protein degra dation system. The 26S proteasome (in plants as well) consists of two components: 20S proteasome and two 19S regulatory particles (RPs), which interact with ubiquiti nated proteins assuring substrate specificity of 26S-com plex. In *A. thaliana*, three β-subunits out of seven sub units of the 20S proteasome (PBA, PBB, and PBE) belong to the threonine peptidase family (T1 family, PB clan) and possess proteolytic activity [152]. In plant cells, there are both 26S proteasomes and free 20S proteasomes capable of ubiquitin-dependent and ubiquitin-independ ent cleavage of proteins, respectively [152, 153].

It was recently shown that 20S proteasomes are responsible for caspase-3-like activity detected during xylem development in *A. thaliana* and poplar [154]. Hatsugai with coauthors showed that proteolytically active β-subunit of 20S proteasome PBA1 is partially responsible for caspase-3-like activity that can be detect ed during the development of hypersensitive response of *A. thaliana* to bacterial infection caused by *P. syringae* [155]. Taking into account these data, proteasomes and/or their components were proposed to be involved in RCD mechanisms in plants.

#### METALLOPROTEINASES

Plant genomes also contain genes coding matrix metalloproteinases. For example, in the genome of *A. thaliana* there are five such enzymes [156, 157]. According to the MEROPS classification of proteolytic enzymes, these proteases belong to M10 family from MA clan [80]. As in other organisms, in plants matrix metal loproteinases are composed of a signal peptide, prodomain, and catalytic domain containing a zinc-bind ing motif. Activation of the enzyme occurs after physical separation of the prodomain from the catalytic site, which can be reached by means of proteolytic cleavage of the prodomain [158]. It is believed that plant matrix metallo proteinases are localized either in plasma membrane or in intercellular space [156, 157]. Moreover, At4-MMP met alloproteinase from *A. thaliana* bears non-cleavable *N* terminal signal peptide that can target this enzyme to the ER [156]. Implication of plant matrix metalloproteinases in RCD processes was not demonstrated, except the fact that the gene coding for matrix metalloproteinase Cs1- MMP of cucumber is expressed *de novo* at final steps of aging processes in cotyledon before initiation of RCD [159]. This fact suggests that plant matrix metallopro teinases might be involved in RCD processes, although functions of these enzymes still need to be studied.

Plant genomes bear genes coding for hundreds of proteases. In *A. thaliana*, their number reaches 800 [24]. Among proteins encoded by plant protease genes, repre sentatives of all five main classes of proteolytic enzymes are found: cysteine, serine, aspartic, threonine, and met alloproteinases. Now it is clear that representatives of all these classes of proteolytic enzymes are involved in devel opments of various types of regulated cell death (RCD) (table).

This review has summarized data about proteolytic enzymes of plants able to regulate initiation and further developments of regulated cell death caused by biotic and abiotic factors or by particularities of ontogenesis. However, this review does not discuss (with rare excep tions) participation of plant proteolytic enzymes in senes cence, since it is still difficult to distinguish processes pre ceding regulated cell death and RCD itself. Moreover, the role of plant proteolytic enzymes in senescence was reviewed earlier [160, 161].

In fact, accumulation of data about the functional role of each proteolytic enzyme implicated in RCD mechanisms in plants has just started. Only a few physio logical substrates of these enzymes have been identified so far. Obviously, we still need to characterize the majority of these substrates. There is some evidence that proteolytic cascades can be involved in initiation and regulation of RCD in plants, as well as in animals. Identification of particular endopeptidases involved in these cascades will undoubtedly be subject of further research. In plants, several types of RCD demanding participation of several proteolytic enzymes were already described; however, the question of regulation of their interactions remains with out answer so far.

One of the reasons for intensive studies of RCD in humans and other animals is the direct connection of RCD in humans with processes of carcinogenesis. It is now clear that RCD in plants is tightly linked with processes of development, as well as with resistance mechanisms against various stresses and pathogens. Thus, studies of molecular mechanisms of regulated cell death in plants are not only important for basic research; they might provide new tools for regulation of defense mecha nisms and ontogenesis in plants.

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