Review

Caspase-dependent non-apoptotic processes in development

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Caspases are at the core of executing apoptosis by orchestrating cellular destruction with proteolytic cascades. Caspase-mediated proteolysis also controls diverse nonlethal cellular activities such as proliferation, differentiation, cell fate decision, and cytoskeletal reorganization. During the last decade or so, genetic studies of Drosophila have contributed to our understanding of the in vivo mechanism of the non-apoptotic cellular responses in developmental contexts. Furthermore, recent studies using C. elegans suggest that apoptotic signaling may play unexpected roles, which influence ageing and normal development at the organism level. In this review, we describe how the caspase activity is elaborately controlled during vital cellular processes at the level of subcellular localization, the duration and timing to avoid full apoptotic consequences, and also discuss the novel roles of non-apoptotic caspase signaling in adult homeostasis and physiology.

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Facts

- Caspase activation is locally regulated in subcellular compartments so that cellular remodeling during terminal differentiation is accomplished without causing cell death.
- Transient caspase activity is utilized for cell fate determination in mammalian stem cells and Drosophila neural progenitor cells in which no organelle degeneration occurs.
- Temporal control of the caspase activity mediated by IAP turnover regulates the cytoskeletal dynamics during cellular morphogenesis and cell migration in Drosophila.
- Caspase-mediated signaling at the organism level affects animal-wide phenotypes, such as ageing and normal development in C. elegans.

Open Questions

- What triggers caspase activation in nonlethal cellular processes in diverse developmental contexts?
- How is caspase activation spatiotemporally controlled and how are specific substrates recognized by active caspases during non-apoptotic processes?
- How does caspase signaling integrate cellular responses and affect organism-level phenotypes?
- What does the past-caspase activation in normal healthy cells tell us? Are there novel non-apoptotic functions of caspases or does developmental anastasis occur as a physiological event?

Caspases, a family of cysteine proteases, are highly conserved throughout the metazoa and are mostly known as executioners of apoptosis. Apoptosis is a genetically encoded suicide program that manifests distinct morphological features including cell shrinkage and nuclear fragmentation followed by phagocytic clearance. In vivo, apoptosis is widely observed and it plays a pivotal role in sculpting or removing tissues by eliminating unwanted cells. Apoptosis also assists morphogenesis and organogenesis by providing mechanical forces. In contrast, the inhibition of apoptosis causes developmental defects, such as aberrant heart formation and exencephaly. Furthermore, dysregulation of cell death signaling leads to human diseases including cancer and inflammatory disorders. Thus, caspase-mediated apoptosis is a fundamental cellular response in development, homeostasis, and pathophysiology.

Although caspases have been recognized as killer enzymes, studies over the last 15 years have indicated that the same enzymes also have vital functions. Such non-apoptotic functions of caspases are diverse, ranging from immune response, cell proliferation, cellular remodeling, cell fate determination to cytoskeletal reorganization. Studies using genetically tractable animals, especially the fruit fly Drosophila melanogaster, have provided mechanistic insights into non-apoptotic caspase signaling, which is under local, transient or temporal control in different developmental scenarios. In addition, recent studies using the nematode C. elegans have revealed novel functions of caspase signaling at the organism level. In this review, we exclusively focus on caspase-dependent non-apoptotic processes in development, aiming to introduce the...
progress in understanding the mechanism that controls the nonlethal function of caspases and to discuss future directions of this field of research. As has been previously proposed, we herein consider development as the entire life span of the organism, including the so-called ‘egg to organism’ and the following adulthood stage.

Conserved Caspase-Mediated Pathway in Development

The core caspase signaling pathway has been studied using model animals, *C. elegans*, *Drosophila*, and the mouse. The first cell death gene was identified by the genetic study of *C. elegans*, in which *ced-3* mutant exhibited the survival of 131 cells that are destined to die. In *C. elegans*, four genes (**ced-3**, **ced-4**, **ced-9**, and **egl-1**) control apoptotic cell death as a linear pathway; **CED-3** is a caspase and **CED-4** is its activator, whereas **CED-9** and **EGL-1** belong to the BCL-2-like anti-apoptotic proteins and BH3-only pro-apoptotic proteins, respectively. When cells undergo apoptosis, **EGL-1**, which is transcriptionally upregulated, displaces **CED-9** from its complex with **CED-4** at mitochondria, and the released **CED-4** interacts with **CED-3** to activate the proteolytic activity, resulting in cell death (Figure 1a).

Caspases are synthesized aszymogens that are processed at selective aspartate residues for their activation. In flies and mammals, caspase signaling is negatively controlled by the apoptotic inhibitors X-linked IAP (XIAP), DIAP1, and DIAP2. XIAP is the mammalian ortholog of DIAP1. 

In flies and mammals, caspases are subdivided into two categories: initiator caspases (ex. **Dronc** in fly; **caspase-9** in mammals) and effector caspases (ex. **Drice** and **Dcp-1** in fly; **caspase-3** and **caspase-7** in mammals). Initiator caspases have long amino-terminal prodomains that provide the molecular platform for caspase activation. In mammals, upon the release of cytochrome c (**Cyt-c** from mitochondria, the adapter protein apoptosis activating factor 1 (**Apaf-1**) forms the apoptosome complex with **CED-4** at mitochondria, and the released **CED-4** interacts with **CED-3** to activate the proteolytic activity, resulting in cell death (Figure 1a).

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Caspase signaling is also negatively controlled by the inhibitor of apoptosis proteins (**IAPs**). Several **IAPs** carry E3-ubiquitin ligase activity that promotes the degradation of cell death regulators by the ubiquitin-proteasome system. Upon apoptosis signaling stimuli, **IAPs** are inhibited by IAP antagonists, which leads to caspase activation. In *Drosophila*, **Reaper**, **Hid**, and **Grim**, called RHG proteins, which localize to mitochondria to inhibit *Drosophila* IAP1 (**DIAP1**), promote DIAP1 self-degradation, leading to the release of caspases. The mammalian ortholog of DIAP1 is X-linked IAP (**XIAP**), which has E3 ligase activity, and IAP antagonists such as **Smac**, **HtrA2**, and ARTS trigger caspase activation by binding to **IAPs** or promoting their degradation (Figures 1b and c).

In summary, caspase activity is under tight control by both the activating and the inhibiting mechanisms, and these pathways are further regulated by different layers of upstream signaling networks. As illustrated in the following, non-apoptotic events appear to utilize the same set of core caspase signaling, although caspase activation is regulated through its localization, duration, and timing to avoid deleterious consequences. Caspases thus have acquired the elaborate mechanisms to activate their irreversible proteolytic ability for multiple purposes.

Localized Caspase Activation in Cellular Remodeling

During apoptosis, caspases orchestrate the cellular destruction procedure by cleaving their respective substrates, allowing the demolition of cytoarchitecture as well as the degradation of cytoplasmic organelles. Such caspase-mediated proteolysis can be effective for removing specific organelles if the caspase activity is spatially controlled inside the cells. Terminal differentiation of some vertebrate cell types, such as lens epithelial cells, keratinocytes, and erythrocytes, accompanies cellular remodeling that includes the elimination of nuclei and other organelles, in which the caspase activity is required for organelle loss. During lens fiber differentiation, caspase-3-like activity, which is detected in the equatorial epithelium, is necessary for the initiation of the differentiation and enucleation process. Therefore, a loss of organelles may be attributed to partial cellular destruction mediated by local caspase activation. Similar cellular-remodeling events occur in sperm differentiation in both mammals and flies.
Especially, spermatid terminal differentiation in male *Drosophila* gives compelling evidence of a nonlethal level of localized caspase activation with cellular and molecular details.

**Sperm individualization in Drosophila.** *Drosophila* spermatogenesis occurs within individual units, called cysts, where 64 haploid spermatids are initially interconnected by cytoplasmic bridges. Later in sperm development, an actin-based cytoskeletal-membrane complex, 'individualization complex (IC)' is formed, which caudally slides along the spermatid from the head to the tail, disconnecting the cytoplasmic bridges and separating the syncytium into individual sperms (Figure 2a). This final step of spermatogenesis, termed individualization, includes disassembling of organelles and dumping most cytoplasmic components into cystic bulges, which eventually accumulates in waste bags (Figure 2a). Because morphological changes of spermatids accompany organelle destruction and cytoplasmic expelling, this series of events is reminiscent of apoptosis.

The pioneering work by Arama et al. showed that effector caspase activity is required for proper sperm individualization processes. They first identified apoptotic-like corpses in cystic bulges and waste bags during spermatid individualization, whereas caspase-3-like activity is detected once IC is formed and it is highest in cystic bulges, but undetectable in the post-individualized region. Furthermore, both in cultured testes and *in vivo* spermatids, the inhibition of caspases blocked IC movement and the removal of the bulk cytoplasm. Consistent with the involvement of effector caspases, their activator, apoptosome components (Dronec, Ark and Cyt-c) are expressed in and required for individualizing spermatids.

A recent report further demonstrated that Tango7, a previously identified cell death effector, physically interacts with Dronec and Ark, and regulates the apoptosome-dependent caspase activity during sperm differentiation. Of note, spermatids individualization defects are often correlated with male sterility, implying that caspase activity during sperm differentiation is critical for fertility. Collectively, these studies suggest that multiple components of the caspase signaling pathway tightly regulate non-apoptotic caspase activation in individualizing spermatids.

How is the caspase activity in spermatids spatially controlled and maintained at a certain level to avoid cell death while allowing cellular remodeling? In addition to pro-apoptotic genes, the giant IAP-like protein dBruce was identified to protect spermatids from excessive caspase activity that causes nuclear degeneration. From genetic screening, a testis-specific isoform of Cullin-3 was identified to be a component of an E3-ubiquitin ligase complex. Cullin-3 interacts with the small RING protein Roc1b and BTB-containing protein Khi10, forming a Cullin-3-based RING ubiquitin ligase (CRL3) that is required for effector caspase activation. Subsequent work from the same group identified an inhibitor of the CRL3 complex, Soti, which can compete with dBruce to bind Khi10 protein. Interestingly, Soti is expressed in a gradient form in the early elongating spermatids descending from the tail to the head, which is followed by dBruce protein distribution at the time of individualization (Figure 2b). The CRL3 complex and Soti are necessary for the spatial gradient of dBruce distribution, which in turn leads to an inverse gradient of caspase activity in spermatids from proximal to distal, allowing for the gradual accomplishment of individualization and protecting the distal region from excessive caspase activity (Figure 2b). The most recent study further identified a Krebs cycle component, β subunit of the ATP-specific form of the succinyl-CoA synthetase (A-Sβ) that specifically binds to the CRL3 complex. The long, testis-specific isoform of A-Sβ is localized to mitochondria and competes with Soti, contributing to the CRL3 activation at the surface of the mitochondria. The CRL3 complex consists of Cullin-3, Khi10 and Roc1b that recruits an ubiquitin (Ub)-conjugating enzyme (E2)
Altogether, spatiotemporal restriction of the CRL3 complex ensures low levels of caspase activation that avoids the rapid activation of caspases, but allows the proper removal of the cytoplasmic contents (Figure 2b). In murine spermatids, apoptosis-like events in subcellular compartments and caspase activity are also observed during late stage of spermatogenesis.\textsuperscript{45,46} Intriguingly, sperms from mutant mouse for Sept4 gene that encodes the IAP antagonist ARTS, exhibit the retention of cytoplasm, which is similar to the defect in \textit{Drosophila} sperm individualization.\textsuperscript{45} Moreover, the mammalian Klhl10, which is required for mouse spermatogenesis, interacts with Cullin-3 that is expressed in spermatids.\textsuperscript{37} Given that the orthologous proteins function as the CRL3 complex in flies, conserved mechanisms likely regulate localized caspase activation during cellular remodeling of differentiating sperms in mammals.

**Transient Caspase Activation in Cell Fate Determination**

Caspase-mediated proteolysis is an irreversible process that causes apoptosis as well as cellular remodeling. The same irreversible signals can be useful for determining cell fate without causing organelle degeneration if caspases cleave only specific substrates. Caspases potentially have hundreds of substrates so that the level and the duration of caspase activity must be controlled to allow for the cleavage of a few substrates. In some mammalian stem cells, the caspase-3 activity mediates the decision of self-renewal or differentiation.\textsuperscript{48–51} In embryonic stem cells, an increased caspase activity plays a critical role in differentiation by cleaving Nanog, a core transcription factor that controls their pluripotency.\textsuperscript{50} Transient caspase activity is also required for the efficient induction of induced pluripotent stem cells from human fibroblasts, which is a process of cell fate reprogramming.\textsuperscript{52} In \textit{Drosophila} neural cell lineage, the caspase activity contributes to cell fate decision, which provides a foundation of understanding the mechanism of how to control transient caspase activity and its substrate specificity.

**SOP cell specification in \textit{Drosophila}.** \textit{Drosophila} macrochaetae, or large bristles, are external sensory organs located on the adult fly notum and are derived from asymmetric cell division of sensory organ precursor (SOP) cells. SOP cells of the macrochaetes originally appear in the wing imaginal disc during larval stage and are selected from cells. SOP cells of the macrochaetes originally appear in the asymmetric cell division of sensory organ precursor (SOP) located on the adult fly notum and are derived from chaetes, or large bristles, are external sensory organs in \textit{Drosophila} specific orthologs.\textsuperscript{54} One of the isoforms, Sgg46, can be cleaved in a caspase-dependent manner, and after the processing, it can function as an active kinase Sgg10. Sgg10 then negatively regulates Wg/\textit{Wnt} signaling through the phosphorylation and degradation of Armadillo/\textit{β}-catenin\textsuperscript{60} or directly phosphorylating Scute and its activator Pannier,\textsuperscript{61} contributing to SOP cell specification. A subsequent study by Kuranaga \textit{et al.}\textsuperscript{62} bolstered the idea that nonlethal level of caspase activity regulates precise SOP cell numbers. The \textit{Drosophila} IKK-related kinase (DmlIKK\textsubscript{ε}) phosphatrolyzes DIAP1 and causes its degradation, thereby regulating the caspase activity through the control of DIAP1 level. RNAI-mediated knockdown of DmlIKK\textsubscript{ε}, which stabilizes DIAP1 and suppresses the caspase activity, induces extra macrochaetae due to extra SOP cells. Of note, the depletion of DmlIKK\textsubscript{ε} affects the caspase activity without contributing to naturally occurring cell death. These results suggest that...
DmIKKe determines SOP cell numbers by controlling the non-apoptotic level of transient caspase activity.\textsuperscript{62}

How does transient caspase activation lead to the cleavage of a specific substrate during SOP cell fate decision? A recent study revealed that the unconventional myosin Crinkled (CK) physically interacts with Dronc, which is required for the Dronc-dependent non-apoptotic processes.\textsuperscript{53} CK, the Drosophila ortholog of mammalian non-muscle MYO7A, was identified from proteomic screening using the Dronc protein complex. Although CK controls different Dronc-dependent phenotypes, it does not affect naturally occurring apoptosis. Depletion of CK in the scutellum phenocopies the effect of RNAi for dronc, DmIKKe, and sgg, and the co-expression of dominant-negative form of CK (CK\textsuperscript{DN}) and DmIKKe-RNAi enhanced the extra macrochaete phenotype. These observations suggest that CK regulates the non-apoptotic function of caspases and contributes to the control of SOP cell numbers. Mechanistically, CK binds to both Dronc and inactive isoform Sgg46, but not to active Sgg10, suggesting that CK acts as an adapter that brings the substrate Sgg46 to the proximity of Dronc (Figure 3).\textsuperscript{63} Such a mechanism may confer substrate specificity of caspases, and allow the activation of downstream signaling depending on CK localization.

Caspase signaling may contribute to cell fate determination as a fine-tuning mechanism with additional factors. Indeed, the H3K9 histone methytransferase SETDB1 has been identified to cooperatively regulate the SOP cell numbers with caspases.\textsuperscript{64} Importantly, the mechanisms that control non-apoptotic caspase signaling discovered in flies are conserved in mammals. As DmIKKe phosphorylates DIAP1, the mammalian IKK-related kinase NAK can phosphorylate XIAP.\textsuperscript{62} The mammalian CK counterpart, MYO7A, binds to caspase-8 in the presence of RIPK1 and suppresses the formation of ripoptosome complex by cleaving RIPK1.\textsuperscript{65} Given that IKK-related kinases and RIPK1 are important for the mammalian immune system, such conserved non-apoptotic caspase signaling may perform diverse physiological functions including cell fate determination.

**Temporal Control of Caspase Activation in Cytoskeletal Dynamics**

Caspases are responsible for altering the morphological changes associated with apoptosis, such as cell rounding and membrane blebbing. Cytoskeletal remodeling during nonlethal events can be regulated by the caspase activity if it targets cytoskeletal regulators under strict temporal control. In mammals, during apoptosis, caspase-3 can cleave Rho-associated protein kinase 1 (ROCK1), which results in its activation and the following actin cytoskeletal rearrangement.\textsuperscript{65–67} Caspase-3-mediated cleavage of ROCK1 is also observed during macrophage polarity formation, a non-apoptotic process.\textsuperscript{68} Moreover, caspase-8 and caspase-11 are implicated in cell migration in non-apoptotic circumstances,\textsuperscript{69–72} suggesting that caspases are indeed the regulators for actin cytoskeleton dynamics. For controlling the caspase activity, IAPs are conserved determinants of both apoptotic and non-apoptotic functions.\textsuperscript{27,72} In particular, cellular shaping and border cell migration in Drosophila are the processes that provide mechanistic insights into the temporal control of caspase activation through IAP turnover, which influences actin cytoskeleton dynamics.

**Cellular shaping and border cell migration in Drosophila.** The Drosophila antenna arista has a feather-like structure that consists of a central core and a series of lateral branches (Figure 3).\textsuperscript{73} Mutant flies for DIAP1 and its antagonist Hid exhibit abnormal arista morphology, while lacking lateral branching and ectopic branching, respectively.\textsuperscript{74} Although diap1 overexpression was sufficient to induce extra branches, p35 overexpression that blocks effector caspases did not affect the arista morphology, raising the possibility that DIAP1-mediated non-apoptotic caspase signaling could influence arista morphogenesis. Oshima et al.\textsuperscript{75} showed that the DIAP1-degrading kinase DmIKKe negatively regulates F-actin assembly. Expression of the dominant-negative form of DmIKKe (DmIKKe\textsuperscript{DN}) in arista causes an abnormal excess-branching morphology, which is suppressed by diap1 knockdown and is enhanced by diap1 overexpression, suggesting that this phenotype is sensitive to the DIAP1 dosage. Because the DIAP1 level is critical for the control of Dronc, the extra branching of arista may be attributed to the suppression of Dronc activity. Indeed, dronc mutant adult escapers exhibit extra arista branches,\textsuperscript{76} and the lateral branching phenotype caused by DmIKKe\textsuperscript{DN} expression is enhanced by the knockdown of dronc or its activator ark.\textsuperscript{75} A recent study further showed that the mutant allele for the unconventional myosin CK that controls Dronc activity exhibits extra arista branches in the same manner as flies expressing DmIKKe\textsuperscript{DN}.\textsuperscript{63} Although how the Dronc activity affects F-actin dynamics is still unclear, these results suggest that DIAP1-mediated control of non-apoptotic caspase signaling is crucial for the proper cellular shaping of arista (Figure 3).

Cytoskeletal dynamics regulated by non-apoptotic caspase signaling is also featured in border cell migration during Drosophila oogenesis. In the developing egg chamber, a group of cells detach from the follicle epithelium and become migratory border cells (Figure 3).\textsuperscript{77} From genetic screening, Geisbrecht and Montell\textsuperscript{78} found that the overexpression of diap1 rescues the border cell migration defect caused by a dominant-negative form of Rac (Rac\textsuperscript{DN}) expression. Mutant flies for DIAP1 also exhibit migration defects without showing cell death, suggesting that the endogenous level of DIAP1 is important for normal border cell migration. The migration defect caused by Rac\textsuperscript{DN} expression was also rescued by the overexpression of Dronc\textsuperscript{DN} or hypomorphic alleles of ark.\textsuperscript{78} In contrast, the overexpression of DmIKKe that leads to the activation of Dronc, inhibits border cell migration.\textsuperscript{75} These observations suggest that the level of DIAP1 and the subsequent control of the Dronc activity are crucial for border cell migration. During SOP cell specification, the unconventional myosin CK acts as a substrate adapter that brings the GSK-3\textsuperscript{β} precursor Sgg46 to Dronc, facilitating its cleavage.\textsuperscript{63} The expression of CK\textsuperscript{DN} or a non-cleavable form of Sgg46 also rescues the migration defect caused by Rac\textsuperscript{DN} expression, suggesting that the activation of Sgg suppresses border cell migration downstream of Dronc activation.\textsuperscript{63} Although the underlying mechanism of how Sgg regulates the actual migration process remains elusive, it might affect the dynamics of the cytoskeletons or cell adhesions that are
required for proper migration. Taken together, these results suggest that CK-regulated Dronc activity under the control of DIAP1 negatively regulates the Rac-mediated cell migration of border cells (Figure 3). The above examples illustrate the role of the proper amount of DIAP1 protein or its metabolism, which is critical for the control of Dronc activity. How can DIAP1 affect both the apoptotic and non-apoptotic functions of caspases? Using a fluorescent probe for DIAP1 dynamics, which reflects its endogenous degradation, Koto et al. examined the protein turnover of DIAP1 in the external SOP lineage of Drosophila pupa. The SOP cells divide asymmetrically to give rise to the shaft, socket, sheath cells, and the neurons that all constitute the shaping of this sensory organ. Live imaging of the SOP lineage indicated that DIAP1 turnover is controlled depending on cell type and maturity. By genetically manipulating the DIAP1 levels, this study addressed the physiological role of DIAP1 dynamics. DmIKK-ε-knockdown flies exhibited a delayed DIAP1 degradation in the shaft cell, which caused the shorter and thicker bristle phenotype. In contrast, the knockdown of diap1 or overexpression of rpr resulted in a defect of DIAP1 stabilization associated with the shaft cell death and subsequent bristle loss phenotype. These results suggest that temporal regulation of DIAP1 turnover determines whether caspases function in apoptosis or in cellular shaping as a non-apoptotic process. Given that mammalian IAPs affect cytoskeletal remodeling besides the control of caspase activity, similar mechanisms might play a role in switching apoptotic and non-apoptotic roles in different organisms.

**Caspase Activation at the Organism Level in Homeostasis and Physiology**

During animal life from embryo to adult, organisms must cope with various environmental stresses and have developed stress–response programs at different levels. At the cellular level, cells that have deleterious damage undergo apoptosis to remove any potentially harmful cells. In contrast, at the organism level, a coordinated stress response can enhance the resistance to further stress. Interestingly, in *C. elegans*, mutations in ced-3 gene confer stress resistance to a variety of stresses, such as ER and osmotic stress, suggesting that animal-wide stress responses are coordinated by the same signaling pathway that causes apoptosis at the cellular level. In this section, we introduce recent studies about the unexpected contribution of the apoptotic caspases to organism-level phenotypes such as ageing and whole-body development.

**Increased longevity and robust development in *C. elegans***

As the first long-lived mutants were identified, genetic pathways that control ageing have been vigorously studied using *C. elegans*. Among the mutants that show increased longevity, missense mutations in isp-1 and nuo-6 exhibit elevated reactive oxygen species (ROS) generation. The isp-1 and nuo-6 genes encode subunits of the complex III and complex I of the mitochondrial respiratory chain, respectively. Such a long-lived phenotype is phenocopied by treatment with a low dose of superoxide generator paraquat for wild-type. These results suggest that mitochondrial ROS (mtROS) is associated with increased longevity. A recent study addressed the mechanism of how an increase in mtROS promotes longevity by focusing on intrinsic apoptotic signaling. The conserved intrinsic apoptosis pathway components, CED-9 and CED-4, are physically associated with mitochondria, and mtROS triggers the apoptotic pathway in vertebrates. Adding mutations in ced-9 (gain of function), ced-4 or ced-3 in isp-1 and nuo-6 mutants significantly suppressed their prolonged life span. Similar effects on the suppression of increased longevity were observed in those mutants following exposure to paraquat. Interestingly, mutations in egl-1 that encodes the BH3-only protein required for apoptosis does not affect the life span of mitochondrial mutants and wild-type animal after paraquat treatment, suggesting that suppression of the increased longevity is not the results of inhibition of apoptosis. Instead, another BH3-only protein CED-13 is required for the increased longevity and functions in the same pathway with CED-9, CED-4, and CED-3. These observations suggest that in the context of mitochondrial dysfunction, mtROS is sensed by the intrinsic apoptotic pathway, and this non-apoptotic function serves as a protective mechanism to maintain organism survival, leading to the increased life span (Figure 4).

Another interesting observation came from the study of microRNAs (miRNAs) on robust animal development. miRNA-mediated gene silencing is critical for diverse cellular processes by ensuring dynamic gene expression, which often works as a large complex, miRNA-induced-silencing-complex (miRISC). From an attempt to investigate genes that act with miRNA, Weaver et al. performed genome-wide RNAi screening and identified the caspase ced-3 as an interactor of ain-1, a homolog of the miRISC component. Double mutants for ced-3 and ain-1 have multiple defects in development including delays in larval growth, abnormal adult morphology, and embryonic lethality. Mutations in ain-1 do not...
affect cell death phenotypes, suggesting that the functions of ced-3 with ain-1 are non-apoptotic. Subsequent experiments revealed that ced-3 specifically functions with miRNAs in let-7 family members that targets mRNA of pluripotency factors LIN-14 and LIN-28. Of note, ced-3 and ain-1 double-mutant phenotypes were suppressed by downregulating lin-14 or lin-28. These results suggest that CED-3 normally represses these pluripotency factors, leading to the hypothesis that proteolytic cleavage by CED-3 regulates their expression. In vitro assays confirmed their CED-3-dependent cleavage, and in vivo experiments demonstrated the dynamic turnover of LIN-28, which is negatively regulated by CED-3. Finally, the expression of the non-cleavable form of LIN-28 alone is sufficient to induce developmental delay and temporal cell fate patterning associated with the abundance of LIN-28, suggesting that the CED-3-dependent cleavage of LIN-28 plays a role in ensuring correct development.95,96 Taken together, these results suggest that the non-apoptotic caspase activity functions as a backup mechanism that allows normal development in C. elegans. It would be interesting to clarify whether higher organisms may have similar mechanisms to adapt to both internal abnormalities and environmental challenges, and precisely elucidate how caspase signaling integrates cellular responses to the organism level.

Conclusions and Perspectives
Recent research progress has provided a clearer view of the mechanism how killer enzymes regulate vital cellular processes. The apoptotic signaling pathway is utilized in a way that regulates the nonlethal level of caspase activity at the level of initiator caspases or by conferring substrate specificity for active caspases. In order to avoid apoptosis, it is also important to restrict the caspase activity into subcellular compartments to allow local organelle degeneration. Although it still remains elusive, caspase signaling in non-apoptotic processes must be spatiotemporally controlled, as previously shown in the cases of developmental apoptosis.91–94 Moreover, little is still known regarding what developmental signal triggers non-apoptotic caspase activation in different contexts. Dissecting the precise mechanism of when, where, and how caspase activation initiates and propagates in both apoptotic and non-apoptotic processes will provide a comprehensive view on the control of caspase activation in vivo.

Using caspase reporters in vivo, Drosophila genetic studies illustrated that the effector caspase activity has an execution threshold: cells can tolerate and survive the caspase activity below this threshold, whereas they commit suicide if the caspase activity increases beyond that threshold.95,96 Consistent with this finding, studies using the new biosensors, which detect past-caspase-3-like activities as a history, showed that large numbers of healthy cells in adult flies have experienced effector caspase activation under normal physiological conditions.97,98 This observation implies the existence of many cases of non-apoptotic caspase activation with potential novel roles. It is also possible that some cells can recover from stimulus that could cause apoptosis. This phenomenon, named anastasis, has been reported in mammalian cell cultures and in the Drosophila egg chambers under stress conditions.97,99 It is currently unclear how apoptotic cells can survive and whether anastasis is a normal developmental event with physiological significance. Looking ahead, future studies will examine the mechanism and the role of unappreciated non-apoptotic caspase activity in homeostasis and in physiology. Deciphering the physiological role of anastasis or cell recovery from apoptotic events in vivo will be another important avenue for future research. A better understanding of caspase signaling in non-apoptotic processes in vivo will lead to the precise control of the caspase activity in space and time, and the medical applications of this phenomenon may lead to potentially new treatments for diseases caused by aberrant cell death signaling.

Conflict of Interest
The authors declare no conflict of interest.
Non-apoptotic roles of caspases in development
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