Structure and Activation Mechanism of the Drosophila Initiator Caspase Dronc

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Activation of an initiator caspase is essential to the execution of apoptosis. The molecular mechanisms by which initiator caspases are activated remain poorly understood. Here we demonstrate that the autocatalytic cleavage of Dronc, an important initiator caspase in Drosophila, results in a drastic enhancement of its catalytic activity in vitro. The autocatalytic cleavage of Dronc forms a homodimer, whereas the uncleaved Dronc zymogen exists exclusively as a monomer. Thus, the autocatalytic cleavage in Dronc induces its stable dimerization, which presumably allows the two adjacent monomers to mutually stabilize their active sites, leading to activation. Crystal structure of a prodomain-deleted Dronc zymogen, determined at 2.5 Å resolution, reveals an unproductive conformation at the active site, which is consistent with the observation that the zymogen remains catalytically inactive. This study revealed insights into mechanism of Dronc activation, and in conjunction with other observations, suggests diverse mechanisms for the activation of initiator caspases.

Caspases, the executors of cell death, comprise two families, the initiator caspases and the effector caspases (1, 2). Caspase is synthesized as a single-chain zymogen and must undergo an activation process to acquire full catalytic activity. An effector caspase is activated through an intrachain cleavage mediated by a specific initiator caspase. For example, the effector caspase-3 and caspase-7 are processed by the initiator caspase-9 in the intrinsic apoptosis pathway or by the initiator caspase-8 in the extrinsic pathway (2, 3). An initiator caspase, on the other hand, is autoactivated in response to upstream death stimuli. The autoactivation of an initiator caspase is mediated by a specific adaptor protein complex in cells.

The activation mechanism for the effector caspases, as exemplified by caspase-7 (4, 5), is well characterized. Both the zymogen and the active form of caspase-7 are constitutively homodimeric. The active site of one caspase-7 monomer is formed by five surface loops: four from within, named L1, L2, L3, and L4, and the fifth one from the adjacent monomer, named L2'. This structural feature necessitates dimerization for activation of the effector caspases. In the caspase-7 zymogen, the L2' loop is covalently linked to its N-terminal sequences and is unable to adopt the conformation as required in the activated caspase-7. Consequently, the active site of the caspase-7 zymogen adopts an inactive conformation that does not support catalysis. The intrachain cleavage of caspase-7 zymogen allows the L2' loop to adopt the correct conformation.

In contrast to effector caspases, the mechanisms of activation for the initiator caspases are poorly understood (6). One of the most intensely studied initiator caspases is caspase-9, the activation of which requires the assembly of the apoptosome, a 1-MDa protein complex composed of Apaf-1, cytochrome c, and ATP/dATP (7). Interestingly, the autocleaved caspase-9 remains associated with the apoptosome as an active holoenzyme, and the isolated, cleaved caspase-9 is marginally active (8). The prodomain of caspase-9, which is not cleaved off the caspase unit in the activated enzyme, is primarily responsible for binding to Apaf-1. Both the zymogen and the mature caspase-9 exist as a monomer (9), and the autocatalytic cleavage does not play a critical role in the activation of caspase-9 (10). Despite recent progress (11, 12), the underlying mechanism by which the apoptosome activates caspase-9 remains enigmatic. In contrast to caspase-9, autocatalytic cleavage appears to play an essential role for the activation of caspase-8 (13, 14) and caspase-2 (15). In both cases, the autocatalytic cleavage facilitates the formation of a homodimer for the initiator caspases in vitro.

Dronc is the Drosophila ortholog of the mammalian initiator caspase-9 and is required for programmed cell death during the normal development of fruit flies (16–18). One important downstream target of Dronc is the effector caspase Drice, an ortholog of mammalian caspase-3. The activation of Dronc in Drosophila cells requires Dark/Hac-1/Dapaf-1 (19–21), the ortholog of mammalian Apaf-1. Analogous to caspase-9, the prodomain of Dronc is responsible for interaction with Dark/Hac-1/Dapaf-1 (2, 15). However, in contrast to caspase-9, the prodomain of Dronc is cleaved off the caspase unit in Drosophila (22), suggesting a different mode of activation. At present, the molecular mechanism by which Dronc is activated remains unknown.

In this study, we report the molecular mechanism for the activation of Dronc cleavage-induced stable dimerization. The Dronc zymogen exists exclusively as a monomer in solution. The intrachain cleavage triggers the preferential formation of a homodimer for the cleaved Dronc, which exhibits a drastically elevated level of catalytic activity when compared with the monomeric zymogen. The crystal structure of a catalytic mutant Dronc in its zymogen form reveals the basis for catalytic dormancy, an unproductive conformation in the active site. Together, these results define the precise molecular mechanisms by which Dronc is activated.

EXPERIMENTAL PROCEDURES

Protein Preparation—All constructs were generated using a standard PCR-based cloning strategy, and the identities of individual clones were verified through double-stranded plasmid sequencing. All proteins

* This work was supported by grants from the National Institutes of Health. The costs of publication of this article were defrayed in part by the payment of page charges. This article must therefore be hereby marked “advertisement” in accordance with 18 U.S.C. Section 1734 solely to indicate this fact.

† The atomic coordinates and structure factors (code 2FP3) have been deposited in the Protein Data Bank, Research Collaboratory for Structural Bioinformatics, Rutgers University, New Brunswick, NJ (http://www.rcsb.org/).

‡ The on-line version of this article (available at http://www.jbc.org) contains supplemental text on analytical ultracentrifugation and four supplemental figures.

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Mechanism of Dronc Activation

were overexpressed in Escherichia coli strain BL21(DE3) either as non-tagged protein using the vector pBB75 or as C-terminal His$_{6}$-tagged proteins using pET21b (Novagen). Proteins were purified to homogeneity as described (23).

Crystalization and Data Collection—Crystals of pro-domain-deleted Dronc (residues 136–450, C318A) were obtained by the hanging-drop vapor diffusion method by mixing the complex (10 mg ml$^{-1}$) with an equal volume of reservoir solution containing 100 mM imidazole, pH 9.0, and 1.1 mM ammonium mono-hydrogen phosphate. Cubic-shaped crystals appeared after 5 days and grew to a typical dimension of $0.2 \times 0.2 \times 0.2$ mm$^3$. They were in the space group I4132 and contained one molecule in each asymmetric unit. The unit cell dimensions are $a = b = c = 165.7$ Å. Selenomethionine-derived proteins were crystallized under a similar condition with the same space group and similar unit cell parameters. Crystals were equilibrated in a cryoprotectant buffer containing well buffer plus 25% glycerol and were flash-frozen in a $-170$ °C nitrogen stream. The native data were collected at the Cornell High Energy Synchrotron Source (CHESS) beamline A1, and seleno-multiple anomalous dispersion data were collected at the CHESS beamline F2 and processed using the software Denzo and Scalepack (24).

Structure Determination—The structure was determined by multiple anomalous dispersion using SOLVE (25). Seven selenium atoms per asymmetric unit were located. These selenium positions were further refined using MLFPHARE (26). A model was built using O (27) and refined at 2.5 Å resolution using CNS (28).

Gel Filtration Assay—Individual recombinant proteins were purified to more than 95% homogeneity and incubated in assay buffer (25 mM Tris, pH 8.0, 150 mM NaCl, and 2 mM dithiothreitol) for 10 min at 4 °C. 0.5 ml of the protein mixture was subjected to gel filtration analysis (Superdex 200, Amersham Biosciences) for each run. Samples taken from relevant fractions were applied to SDS-polyacrylamide gel electrophoresis (SDS-PAGE) and visualized by Coomassie Blue staining.

Dronc Activity Assay—Reactions were carried out at 22 °C in an assay buffer containing 25 mM HEPES, pH 7.5, 150 mM NaCl, and 2 mM dithiothreitol. Catalytically inactive DrICE zymogen (C196A) was used as the substrate at a concentration of ~100 μM. Mature Dronc and catalytically active but non-cleavable Dronc zymogen (E352A) were purified to homogeneity and used in each reaction at 100 nM. Reaction samples were taken out at the indicated time points and applied to SDS-PAGE followed by Coomassie Blue staining. Similar results were obtained under pH 6.8 and pH 8.0 as well as a variety of other ionic strength conditions.

Drosophila Genetics and Immunocytochemistry—GMR$^{3,}$-Dronc-WT, GMR-Dronc-C318S, and GMR-Dronc-E352A flies were generated by constructing wild-type (WT) Dronc, Dronc-C318S, and Dronc-E352A into the glass multiple reporter (GMR) P element vector and then introducing these constructs into the Drosophila germline using standard techniques. GMR-Hid, GMR-Reaper, and GMR-Grim were generated as described (29). Multiple lines were generated. Consistently, GMR-Dronc-E352A and GMR-Dronc-C318S showed alleviated ablated eye phenotype when compared with those of GMR-Dronc-WT flies. Anti-Dronc antibody staining was carried out on third instar eye discs from flies of various genotypes essentially as described (29).

Analytical Ultracentrifugation—Protein samples were prepared in 25 mM phosphate-buffered saline with pH 8.0 and 150 mM NaCl. All sedimentation equilibrium experiments were performed with a Beckman Optima XL-I at the Center for Analytical Ultracentrifugation of Macromolecular Assemblies (the University of Texas Health Science Center at San Antonio, Department of Biochemistry). See supplemental data for details.

RESULTS

Dronc Undergoes Autocatalytic Cleavage—To understand how Dronc is activated, we reconstituted an in vitro activation assay for Dronc. The full-length WT Dronc was expressed in E. coli and purified to homogeneity (Fig. 1A, lane 1). The purification procedures for Dronc were carried out at 4 °C so as to maintain the WT Dronc protein in its zymogen form. When the single-chain WT Dronc zymogen was incubated at 22 °C, autocatalytic cleavage at a slow yet detectable rate was observed as evidenced by the initial appearance of two bands below the zymogen (Fig. 1A, lanes 2–4). Further autocatalytic cleavage resulted in the generation of two additional products (Fig. 1A, lanes 5–7).

Analysis of the initial cleavage products by N-terminal peptide sequencing identified the primary autocatalytic cleavage site to be at residue Glu$^{132}$, which is the last amino acid of a catalytic Drice subunit sequence Thr$^{124}$-Glu$^{130}$-Thr$^{131}$-Glu$^{132}$ (124TQTE127). Although Dronc contains two putative caspase cleavage sites$^{110}$DESD113 and$^{122}$DIVD135 between its prodomain and the caspase unit, subsequent analysis by N-terminal peptide sequencing revealed that the additional cleavage products were due to a secondary cleavage site after residue Glu$^{134}$ following an$^{140}$EASE$^{143}$ sequence. This observation is consistent with the finding that Dronc, in contrast to all other known caspases, prefers substrate specificity of Glu at the P1 position (30). The accelerated cleavage of Dronc was not due to other contaminating protease(s) because no cleavage was observed for the catalytically inactive Dronc (C318A) even after prolonged incubation (Fig. 1A, lanes 8–9).

Our studies indicated that the maturation of Dronc has two sequential cleavage events. The first and primary cleavage after Glu$^{132}$ results in the separation of the caspase small subunit from the caspase large subunit and the prodomain, whereas the second cleavage after Glu$^{134}$ removes the prodomain from the catalytic core domain. The first cleavage after Glu$^{132}$ is required for the second cleavage after Glu$^{134}$ because the mutant Dronc E352A, which eliminated the first cleavage site but contained WT catalytic residues, remained as a single polypeptide even after prolonged incubation (Fig. 1A, lanes 10–11).

To further characterize the autocatalytic cleavage, we incubated the catalytically inactive Dronc C318A zymogen, which contains Glu$^{132}$, with Dronc E352A, which contains WT catalytic residues but was unable to cleavage itself due to the mutation E352A. The Dronc C318A zymogen was cleaved after Glu$^{132}$, resulting in two fragments (Fig. 1B). This result indicates that the autocatalytic cleavage of Dronc can occur in trans and further demonstrates that the mutant Dronc E352A is catalytically active, albeit the level of activity remains at the basal level (Fig. 1B).

Mature Dronc Exhibits Drastically Elevated Catalytic Activity—The autocatalytic cleavage of Dronc likely induces its activation. To confirm this conjecture, we reconstituted an in vitro Dronc activity assay to compare the catalytic activity of Dronc zymogen with that of the mature Dronc (Fig. 1C). The catalytically inactive Drice zymogen (residues 1–339, C196A) was used as a physiologically relevant substrate for Dronc. The fact that Dronc zymogen undergoes a process of autocleavage (Fig. 1C, left panel), the mature Dronc cleaved 90% of the

The abbreviations used are: GMR, glass multiple reporter; WT, wild-type.
substrate within the first hour of incubation (Fig. 1C, right panel). These experiments were performed under a condition that resembles the physiological pH and ionic strength (see “Experimental Procedures” for details). Similar results were obtained under other conditions (data not shown). Thus the autocleaved Dronc exhibits a catalytic activity that is drastically elevated over the zymogen.

**Autocatalytic Cleavage of Dronc Shifts a Monomer into a Dimer**—Although Dronc is the functional ortholog of mammalian caspase-9, our biochemical characterization demonstrates that they exhibit quite different patterns of activation. The prodomain of caspase-9 is required for the formation of an apoptosome holoenzyme, in which the activity of caspase-9 is allosterically regulated by Apaf-1 (8). Necessitated by this function, there is no caspase cleavage site between the prodomain and the caspase unit of caspase-9. Consequently, the prodomain and the large subunit of caspase-9 are a single polypeptide chain. In contrast, Dronc contains multiple caspase cleavage sites between its prodomain and the caspase unit, and the prodomain of Dronc is removed after a second autocatalytic cleavage event (Fig. 1A). These observations strongly suggest that, in contrast to activated caspase-9 as a holoenzyme, the mature Dronc protein may exist as a free enzyme. These observations further imply that Dronc may utilize a different mechanism for its activation.

To investigate the activation mechanism for Dronc, we first asked why the Dronc zymogen exhibited an extremely low level of catalytic activity through examination of the basal state of Dronc zymogen using gel filtration. Interestingly, the elution volume of the Dronc zymogen (residues 1–450, C318A) corresponded to an apparent molecular mass of $\sim 50$ kDa, consistent with that of a Dronc monomer (Fig. 2A, solid line). This observation suggests an explanation as to why the Dronc zymogen is marginally active since an unassisted caspase monomer is thought to be catalytically inactive due to the unproductive conformation of the active site (2).

Next, we examined the basal state of the autocleaved Dronc. Surprisingly, the elution of the cleaved Dronc (residues 1–352 and 353–450, second autocatalytic cleavage event (Fig. 1A). These observations strongly suggest that, in contrast to activated caspase-9 as a holoenzyme, the mature Dronc protein may exist as a free enzyme. These observations further imply that Dronc may utilize a different mechanism for its activation.

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Next, we examined the basal state of the autocleaved Dronc. Surprisingly, the elution of the cleaved Dronc (residues 1–352 and 353–450,
C318A) was shifted to earlier fractions (Fig. 2A, dotted line), with the elution volume corresponding to an apparent molecular mass of 100 kDa. This observation strongly suggests that the primary autocatalytic cleavage induces the formation of a stable Dronc homodimer. To definitively confirm this finding, we performed analytical ultracentrifugation experiments (Fig. 2B and Supplemental data). As anticipated, the single-chain Dronc zymogen (residues 136–450, C318A) existed exclusively as a monomer in solution, with a fitted molecular mass of 34.8 ± 0.3 kDa. In contrast, the two-chain Dronc (residues 136–352 and 353–450, C318A) existed primarily as a stable homodimer in solution, with a dissociation constant ($K_d$) of 0.27 μM. Thus Dronc activation appears to employ an interesting mechanism; the cleavage after Glu352 triggers the transition from an inactive monomer for the Dronc zymogen to a catalytically active dimer for the cleaved Dronc.

Structure of Dronc Zymogen Reveals an Unproductive Conformation in the Active Site

To elucidate the molecular mechanism by which Dronc zymogen remains catalytically inactive, we expressed, purified, and crystallized a prodomain-deleted Dronc zymogen (residues 136–450, C318A). The structure was determined by multiwavelength anomalous dispersion and refined to 2.5 Å resolution (Table 1 and Fig. 3A). Each asymmetric unit contains one monomer of the Dronc zymogen.

Similar to other known caspases, the Dronc zymogen contains a central six-stranded $\beta$ sheet, flanked on each side by three $\alpha$ helices (Fig. 3A, B and C). In contrast to other caspases, the Dronc zymogen exhibits a number of unique structural features. For example, a 39-residue fragment (residues 158–196) at the N terminus forms an extended and rigid surface loop that zigzags across one side of the zymogen (Fig. 3A and B). The fact that autocleavage after Glu352 leads to formation of a Dronc dimer suggests two possibilities: either the L2 loop interferes with dimer formation in the zymogen, or the cleaved L2 loop significantly strengthens dimer formation in the processed Dronc. In the crystals, the bulk of the L2 loop is disordered and does not allow a direct examination of the first possibility.

Four surface loops, L1, L2, L3, and L4, emanate from the structural core, forming a potential active site (Fig. 3B). The L2 loop appears to be highly flexible as the electron density after residue Ala318 becomes discontinuous. Loops L3 and L4 also exhibit some flexibility, as judged by their high temperature factors in the crystals. Previous studies on caspase-7 showed that the active site loops only become ordered upon activation cleavage and binding to substrate or inhibitors (4).

Structural analysis reveals that the active site of the Dronc zymogen exists in an unproductive conformation (Fig. 3D). In contrast to the active site conformation of the activated, inhibitor-free caspase-7 (Fig.

**FIGURE 2. Autocleavage of Dronc shifts a constitutive monomer into a stable dimer.** A, the Dronc zymogen appeared to be a monomer, but the autocleaved Dronc was primarily a dimer. Shown here is a superposition of the gel filtration chromatograms for the Dronc zymogen (solid line) and the autocleaved Dronc (dashed line). Both Dronc protein samples contained a mutation C318A on the catalytic residue Cys318 so as to eliminate a potential problem of protein degradation. The autocleaved Dronc was obtained by co-expression of the large (residues 1–352) and small (residues 353–450) subunits. Peak fractions were visualized by SDS-PAGE followed by Coomassie Blue staining (right panels). B, the Dronc zymogen exists exclusively as a monomer, but the autocleaved Dronc is primarily a dimer with a dissociation constant of 0.27 μM. These results were obtained from the analytical ultracentrifugation experiments (see Supplemental data).
Mechanism of Dronc Activation

Uncleavable Dronc Zymogen Is Dominant Negative in Vivo—Our biochemical characterization demonstrates that the Dronc zymogen is constitutively monomeric in solution and catalytically inactive due to an unproductive conformation at the active site. The autocleavage after Glu352 favors the formation of a Dronc dimer, which presumably results in the alteration of the unproductive active site conformation. In vivo, the autoactivation process of Dronc is likely facilitated by Dark/Hac-1/Dapaf-1 as Dronc activation in Drosophila cells requires Dark/Hac-1/Dapaf-1 (19–21). These analyses predict that an uncleavable Dronc zymogen E352A will likely act in a dominant negative fashion in vivo as this protein may compete with WT Dronc zymogen for binding to Dark/Hac-1/Dapaf-1.

To corroborate our biochemical and structural findings, we generated transgenic flies overexpressing the WT Dronc, the catalytic mutant Dronc (C318S), or the uncleavable Dronc (E352A) under the eye-specific GMR promoter. Indeed, although overexpression of the WT Dronc caused an ablated eye phenotype (Fig. 4A), overexpression of the catalytic mutant Dronc (Fig. 4B) or the uncleavable Dronc (Fig. 4C) had little impact on the eyes, suggesting that these Dronc variants were unable to become activated in vivo. The pro-apoptotic proteins Reaper, Hid, and Grim (RHG) function by antagonizing the activity of DIAP1 (31). Overexpression of the pro-apoptotic genes Reaper, Grim, and Hid led to induction of cell death and ablated eye phenotypes (Fig. 4, D–F). Co-expression of the Dronc catalytic mutant (C318S) (Fig. 4, G–I) or the uncleavable Dronc (E352A) (Fig. 4, J–L) partially suppressed the ablated eye phenotype. In Drosophila, continuous activity of Dark/Hac-1/Dapaf-1 is required for normal apoptosis (32). Presumably, the catalytically inactive Dronc zymogen or the uncleavable Dronc was able to bind to and saturate all available Dark/Hac-1/Dapaf-1 protein and thus had a negative impact on the autoactivation of WT Dronc protein mediated by Dark/Hac-1/Dapaf-1.

DISCUSSION

Previous studies on the activation of mammalian initiator caspases have revealed significant insights into the mechanisms. Zymogen of both caspase-8 and caspase-2 was found to exist mainly as a monomer by gel filtration (13–15). In contrast, the autocleaved wild-type caspase-8 exists in an equilibrium between monomers and dimers, with a dissociation constant of ~50 μM (14). The processed wild-type caspase-2 also preferentially formed homodimers (15). These findings strongly argue that dimerization is a crucial factor for the activation of caspase-8 and caspase-2 and that the autocalytic cleavage in these mammalian initiator caspases plays an important role for their activation.

In this study, we investigate the activation mechanism for the Drosophila initiator caspase Dronc. We demonstrate that the Dronc zymogen exists exclusively as a monomer in solution and forms a stable dimer following the primary autocalytic cleavage after Glu352. A cleavage mutant Dronc (E352A) does not undergo autocalytic cleavage and remains a monomer in solution. The structure of the Dronc zymogen, representing the first structure of an uncleaved initiator caspase zymogen, reveals an unproductive active site conformation. At present, we do not yet have a structure of the activated, dimeric Dronc, which is expected to reveal the precise conformation of the active site. Nonetheless, previous studies on dimeric caspases suggest that the active site of one Dronc monomer is likely stabilized by the adjacent monomer through the provision of a critical surface loop (L2') (33). Our results demonstrate that the autocalytic cleavage after Glu352 of Dronc greatly facilitates its dimerization, which is essential for its catalytic activity.

If Dronc zymogen exists exclusively as a monomer in solution, how can autocalytic cleavage occur? There are two possibilities. One is that
FIGURE 3. Structure of the Dronc zymogen reveals a molecular basis for the lack of catalytic activity. A, a representative portion of the electron density $2F_o - F_c$ map, contoured at 1.1 $\sigma$ and colored cyan, was shown in stereo around amino acid residues 158–230. B, overall structure of the Dronc zymogen in two perpendicular views. The structural core of Dronc zymogen is shown in brown, with the active site loops highlighted in magenta. The extended N-terminal fragment (residues 158–196) is colored cyan. The four loops defining the potential catalytic site are labeled. C, sequence alignment among Dronc, caspase-9 (Casp-9), and caspase-7 (Casp-7). Because the N-terminal fragment of Dronc exhibits no apparent sequence similarity with caspase-9 or caspase-7, only residues 185–450 of Dronc are shown here. The secondary structural elements of the Dronc zymogen are indicated above the sequences. Conserved residues are colored yellow. D, the appropriate substrate-binding groove is absent in the Dronc zymogen. When compared with the activated, inhibitor-free caspase-7 (middle panel), the Dronc zymogen does not have an appropriate substrate-binding groove (left panel). The active site of the caspase-7 zymogen is shown here as a control (right panel). E, comparison of the conformations of the active site loops among Dronc zymogen, the activated inhibitor-free caspase-7, and the caspase-7 zymogen. The green dotted line represents the L2' loop in the activated inhibitor-bound caspase-7. F, the active site conformation of the Dronc zymogen is quite different from that of the inhibitor-bound caspase-9 (inhibitor omitted for clarity). The active site conformation of caspase-9 is critically supported by the L2' loop, which is provided by the adjacent monomer. However, in Dronc zymogen, the monomeric and uncleaved nature prevents the provision of the L2' loop.
the monomeric zymogen can associate transiently into dimers, as suggested for the initiator caspase-2 (15). Then the transiently dimerized Dronc zymogens process each other in trans. The other possibility is that the monomeric Dronc zymogen can exhibit a very low level of catalytic activity for its own intramolecular cleavage. Although our current study does not allow differentiation of these two scenarios, the experimental evidence indicated that the autocatalytic cleavage of Dronc can occur in trans, suggesting that the cleavage might be a result of transient dimerization. However, our data are merely suggestive, and we do not know whether the autocatalytic cleavage can also occur intramolecularly. Nonetheless, our results unambiguously show that the autocatalytic cleavage greatly facilitates the formation of a homodimer for Dronc.

Although Dronc is the functional homolog of the mammalian caspase-9, their activation mechanisms appear to be quite different (Fig. 5). For caspase-9, the intrachain cleavage is not necessary for its activation as the fully processed caspase-9 in isolation is only marginally active, much the same way as the unprocessed caspase-9 zymogen (8, 10, 34). Rather, association with the apoptosome leads to a dramatic increase (up to 2000-fold) for the catalytic activity of the processed as well as the unprocessed caspase-9 (8, 10). Necessitated by association with apoptosome, the prodomain of caspase-9 remains covalently linked to the large subunit of the caspase unit. In contrast to caspase-9, the free, processed Dronc exhibits a catalytic activity that is several orders of magnitude higher than the Dronc zymogen (Fig. 1C). The extent of activity improvement is reminiscent of catalytic activation for other caspases (35). Thus the processed, free Dronc likely represents the fully activated state. This notion is supported by the observation that the prodomain of Dronc, which is responsible for binding to Dark/Hac-1/ Dapaf-1, is removed from the caspase unit in the mature enzyme. In addition, the presence of recombinant Dark/Hac-1/Dapaf-1 protein does not enhance the catalytic activity of the mature Dronc.2

**FIGURE 4. Uncleavable Dronc zymogen is dominant negative in vivo.** Genotypes are as follows: GMR-Dronc/+ (A), GMR-Dronc-C318S/+ (B), GMR-Dronc-E352A/+ (C), GMR-Rpr/+ (D), GMR-Grim/+ (E), GMR-Hid/+ (F), GMR-Rpr/GMR-Dronc-C318S (G), GMR-Grim/GMR-Dronc-C318S (H), GMR-Hid/GMR-Dronc-C318S (I), GMR-Rpr/GMR-Dronc-E352A (J), GMR-Grim/GMR-Dronc-C318S (K), and GMR-Hid/GMR-Dronc-E352A (L).
Isolated caspase-9 exists constitutively as a monomer in solution, both in its zymogen state and in the autocleaved form (9) (Fig. 5). Caspase-3 or caspase-7, on the other hand, exists constitutively as a homodimer in solution, both in its zymogen form and in the cleaved, activated state (Fig. 5). In contrast to both caspase-9 and caspase-3, Dronc exists exclusively as a monomer in the zymogen state and forms a stable dimer with a dissociation constant of 0.27 μM following the primary autocatalytic cleavage. The mammalian initiator caspsases, caspase-8 and caspase-2, exhibit an increased preference for the formation of homodimer following the autocatalytic cleavage. In this regard, the activation of the initiator caspsases, Dronc in Drosophila and caspase-8/1-2 in mammals, incorporates a feature reminiscent of effector caspsases (Fig. 5). In a more general sense, our study reveals another level of complexity in the activation of caspsases and suggests diverse mechanisms for the activation of initiator caspsases.

It should be pointed out that the model presented (Fig. 5) only highlights certain biochemical aspects of caspase activation. There are a number of other important considerations to caspase activation in vivo. For example, the activation of Dronc in Drosophila cells requires the presence of Dark/Hac-1/Dapaf-1 (19–21,32). A recent study showed that the Dark/Hac-1/Dapaf-1 protein forms a double-donut-shaped apoptosome, with each donut ring comprising eight molecules of Dark/Hac-1/Dapaf-1 (36). The Dark/Hac-1/Dapaf-1 apoptosome may greatly facilitate the recruitment and the activation of the Dronc zymogen by enhancing its dimerization. However, the mechanism by which the Dark/Hac-1/Dapaf-1 apoptosome activates the Dronc zymogen is likely to be different from that by which the Apaf-1 apoptosome activates caspase-9. The detailed activation mechanisms for initiator caspsases remain to be investigated by biochemistry and structural biology.

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FIGURE 5. A comparison of Dronc activation with activation of other caspsases. This comparison reveals that Dronc exhibits features of both initiator and effector caspsases. The inactive zymogens and the activated caspsases are colored green and orange, respectively. The thick and thin arrows denote the primary and secondary cleavage events, respectively.