

1 **The molecular mechanism and evolutionary divergence of**
2 **caspase 3/7-regulated gasdermin E activation**

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19

20 **Abstract**

21 Caspase (CASP) is a family of proteases involved in cleavage and activation of
22 gasdermin, the executor of pyroptosis. In human, CASP3 and CASP7 recognize the
23 same consensus motif DxxD, which is present in gasdermin E (GSDME). However,
24 human GSDME is cleaved by CASP3 but not by CASP7. The underlying mechanism
25 of this observation is unclear. In this study, we identified a pyroptotic pufferfish
26 GSDME that was cleaved by both pufferfish CASP3/7 and human CASP3/7. Domain
27 swapping between pufferfish and human CASP and GSDME showed that the GSDME
28 C-terminus and the CASP7 p10 subunit determined the cleavability of GSDME by
29 CASP7. p10 contains a key residue that governs CASP7 substrate discrimination. This
30 key residue is highly conserved in vertebrate CASP3 and in most vertebrate (except
31 mammalian) CASP7. In mammals, the key residue is conserved in non-primate (e.g.,
32 mouse) but not in primate. However, mouse CASP7 cleaved human GSDME but not
33 mouse GSDME. These findings revealed the molecular mechanism of CASP7 substrate
34 discrimination and the divergence of CASP3/7-mediated GSDME activation in
35 vertebrate. These results also suggested that mutation-mediated functional alteration of
36 CASP probably enabled the divergence and specialization of different CASP members
37 in the regulation of complex cellular activities in mammals.

38

39 **Key words**

40 Immunity; Pyroptosis; Gasdermin E; Caspase 3; Caspase 7; Evolution

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42 **Introduction**

43 Pyroptosis represents a form of programmed cell death that provokes robust
44 inflammatory immune response (Bergsbaken, Fink, & Cookson, 2009; Tsuchiya et al.,
45 2019). Gasdermin (GSDM) serves as the direct executioner of pyroptosis. GSDM forms
46 transmembrane pores to permeabilize the cytoplasmic membrane, which leads to the
47 release of pro-inflammatory cytokines and, if these transmembrane pores persist,
48 pyroptosis (Kovacs & Miao, 2017; J. Shi, Gao, & Shao, 2017; Xia et al., 2021). Human
49 has six GSDM (HsGSDM) family members, HsGSDMA-E, and HsPJVK (Tamura et
50 al., 2007). All HsGSDMs (except for HsPJVK) adopt a two-domain architecture, the
51 N-terminal (NT) pore-forming domain and the C-terminal (CT) autoinhibitory domain
52 (De Schutter et al., 2021; Shao, 2021). Proteolytic cleavage of HsGSDM to remove the
53 autoinhibitory CT domain enables the binding of the lipophilic NT fragment to the cell
54 membrane, where the NT fragments oligomerize and form transmembrane pores to
55 induce osmotic cell lysis (Kuang et al., 2017; X. Liu et al., 2016).

56 Among the signaling pathways that activate GSDM-mediated pyroptosis, caspase
57 (CASP) 1/4/5/11-mediated GSDMD activation is well documented. In human,
58 HsGSDMD is specifically cleaved by the inflammatory HsCASP1 through multiple
59 inflammasome signaling pathways. HsCASP4/5 also cleave HsGSDMD in response to
60 lipopolysaccharide (LPS) stimulation. After cleavage by HsCASP1/4/5, the pore-
61 forming HsGSDMD-NT fragment becomes unconstrained and triggers pyroptosis (He
62 et al., 2015; Kayagaki et al., 2015; J. Shi et al., 2015). Structural and biochemical
63 analysis reveal that HsCASP1/4 β III/ β III' sheet forms an exosite that interacts with the
64 hydrophobic pocket of HsGSDMD-CT domain, rendering HsGSDMD cleavage
65 independent of CASP recognition of the tetrapeptide motif FLTD (K. Wang et al.,
66 2020). When the tetrapeptide FLTD is mutated to AAAD, HsCASP1/4 cleavage of
67 HsGSDMD is not affected. The residues that form the hydrophobic pocket of
68 HsGSDMD-CT domain are not conserved in HsGSDME, suggesting that HsGSDME
69 cannot be cleaved by HsCASP1/4 via interaction with an exosite (Z. Liu et al., 2020).

70 Actually, human HsGSDME is specifically cleaved by HsCASP3 at the consensus
71 motif DMPD, which is required for HsCASP3-mediated HsGSDME proteolysis.
72 Mutation of either the Asp residue of the DMPD tetrapeptide motif leads to cleavage
73 resistance of HsGSDME (Rogers et al., 2017; Wang et al., 2017), indicating that
74 HsCASP3-mediated HsGSDME activation paradigm is distinct from that of
75 HsGSDMD. It is intriguing that although HsCASP7 shares the same recognition motif
76 (DxxD) with HsCASP3, HsCASP7 cannot cleave HsGSDME (Agniswamy, Fang, &
77 Weber, 2009; Slee, Adrain, & Martin, 2001). The molecular mechanism underlying the
78 HsCASP3/7 cleavage discrimination of HsGSDME is unknown.

79 Historically, HsCASP3 and HsCASP7, which are 54% identical, are known to
80 possess similar structure and share overlapping protein substrate repertoire, and are
81 considered to have functionally redundant roles in regulating cell death (Crawford &
82 Wells, 2011; Kumar, 2007; Y. Shi, 2002). Both HsCASP3 and HsCASP7 consist of
83 two polypeptide subunits named p20 and p10 (1:1 ratio), which assemble to form active
84 heterotetramer (Timmer & Salvesen, 2007). HsCASP3/7 contain highly conserved
85 QACRG and SHG motifs in p20 subunit, and SWR and GSWF motifs in p10 subunit,
86 which participate in the catalytic reaction and substrate binding, respectively (Boyce,
87 Degterev, & Yuan, 2004; Cohen, 1997). In the apoptosis signaling pathway, death
88 receptor-mediated extrinsic pathway and mitochondria-mediated intrinsic pathway
89 ultimately converge to the activation of HsCASP3/7 (Budd, Tanneti, Lishnak, &
90 Lipton, 2000; Wajant, 2002). Active HsCASP3/7 cleave a series of protein substrates,
91 including poly(ADP-ribose) polymerase and DNA fragmentation factor 45, which
92 causes chromatin fragmentation and leads to apoptosis (Erener et al., 2012; Wolf,
93 Schuler, Echeverri, & Green, 1999; Zheng et al., 1998). Although HsCASP3 and
94 HsCASP7 exert almost indistinguishable proteolytic specificity to certain polypeptides,
95 there exist functional differences between these two HsCASPs in cleaving some
96 substrates (Brentnall, Rodriguez-Menocal, De Guevara, Cepero, & Boise, 2013;
97 Demon et al., 2009). For instance, HsCASP3 cleaves Bid much more efficiently than
98 HsCASP7, while HsCASP7 cleaves cochaperone p23 more efficiently than HsCASP3

99 (Slee et al., 2001; Walsh et al., 2008). These findings support the notion that HsCASP3
100 and HsCASP7 are enzymatically similar but functionally non-redundant.

101 Different from HsGSDME that is specifically cleaved by HsCASP3, teleost GSDME
102 is cleaved via various modes (Angosto-Bazarra et al., 2022; Yuan, Jiang, Qin, & Sun,
103 2022). In this study, we identified a functional GSDME (named TrGSDME) from
104 pufferfish *Takifugu rubripes*. We found that TrGSDME was specifically cleaved by
105 both HsCASP3/7 and TrCASP3/7, whereas HsGSDME was cleaved by TrCASP3/7
106 and HsCASP3, but not by HsCASP7. We examined the underlying mechanism of this
107 observation. We found that the GSDME-CT and the CASP7 p10 were critical for
108 CASP7 cleavage of GSDME. By a series of site-directed mutagenesis, we discovered
109 a previously unrecognized key residue (S234 in HsCASP7) in p10 that was responsible
110 for the discriminative cleavage of HsGSDME by HsCASP7.

111

112 **Results**

113 **Pufferfish GSDME is specifically cleaved by both human and fish CASP3/7**

114 It has long been observed that although human caspase (HsCASP) 3 and 7 share the
115 same consensus recognition motif DxxD, HsCASP3, but not HsCASP7, cleaves human
116 GSDME (HsGSDME) (at the site of DMPD) (Poreba, Strozyk, Salvesen, & Drag, 2013;
117 Wang et al., 2017). The underlying molecular mechanism is unknown. In this study, we
118 found that a pufferfish *Takifugu rubripes* GSDME (designated TrGSDME), which
119 belongs to the GSDMEa lineage of teleost GSDME ([Figure-figure supplement 1](#)), was
120 specifically cleaved by both HsCASP3 and 7 ([Figure 1A and B](#)). This intriguing
121 observation promoted us to explore the mechanism of HsCASP7 substrate
122 discrimination. We first examined whether TrGSDME could be cleaved by pufferfish
123 CASP (TrCASP) 3/7. The active forms of TrCASP3/7 were prepared ([Figure 1C](#)), both
124 exhibited high proteolytic specificity and activity towards the tetrapeptide DEVD
125 ([Figure 1D and E](#)), which is the conserved recognition motif of HsCASP3/7. When

126 incubated with TrGSDME, both TrCASP3 and TrCASP7 cleaved TrGSDME into the
127 NT and CT fragments, similar to that cleaved by HsCASP3/7, in a dose dependent
128 manner (Figure 1F and G). Accordingly, TrCASP3/7-mediated TrGSDME cleavage
129 was inhibited by the CASP3 inhibitor (Z-DEVD-FMK) and the pan-CASP inhibitor (Z-
130 VAD-FMK) (Figure 1H). Based on the molecular mass of the cleaved NT and CT
131 products, we inferred that the tetrapeptide motif ²⁵⁵DAVD²⁵⁸ in the vicinity of the linker
132 region of TrGSDME may be the recognition site of CASP3/7. Indeed, the D255R and
133 D258A mutants of TrGSDME were resistant to TrCASP3/7 and HsCASP3/7 cleavage
134 (Figure 1I and J, Figure 1-figure supplement 2). Taken together, these results
135 demonstrated that TrGSDME was cleaved by fish and human CASP3/7 in a manner
136 that depended on the specific sequence of DAVD (Figure 1K).

137

138 **CASP3/7-cleaved TrGSDME is functionally activated and induces pyroptosis.**

139 We next examined whether TrGSDME, like HsGSDME, is a functional pyroptosis
140 inducer. For this purpose, HEK293T cells were transfected with mCherry-tagged full
141 length (FL) or NT/CT domain of TrGSDME. The results revealed that TrGSDME-FL
142 and -CT were abundantly expressed in the cells, whereas TrGSDME-NT expression
143 was barely detectable (Figure 2A, Figure 2-figure supplement 1). No significant
144 morphological change or LDH release was observed in cells expressing TrGSDME-FL
145 or -CT (Figure 2B and C). By contrast, cells expressing TrGSDME-NT showed necrotic
146 cell death with massive LDH release and positive Sytox Green staining (Figure 2B and
147 C, Figure 2-figure supplement 2). TrGSDME-NT-induced cell death exhibited osmotic
148 cell membrane swelling, a typical feature of pyroptosis (Figure 2D). Time-lapse
149 imaging showed that TrGSDME-NT triggered rapid cell swelling and membrane
150 rupture, which led to release of cytoplasmic contents and eventually cell death (Figure
151 2E, Figure 1-video 1). To examine the pyroptotic activity of CASP3/7-cleaved
152 TrGSDME, TrCASP3/7 and TrGSDME were overexpressed in HEK293T cells (Figure
153 2F and G). The cells expressing TrCASP3, TrCASP7 or TrGSDME alone had no
154 apparent morphological change or LDH release, whereas the cells co-expressing

155 TrGSDME and TrCASP3 or TrGSDME and TrCASP7 underwent pyroptosis,
156 accompanying with massive LDH release, membrane rupture, and TrGSDME cleavage
157 (Figure 2G-I, Figure 2-figure supplement 3A). Consistently, the presence of the CASP3
158 inhibitor and pan-CASP inhibitor hampered TrGSDME-mediated pyroptosis (Figure 2J
159 and K). Mutation of the cleavage site (D255R and D258A) inhibited pyroptosis and
160 significantly decreased LDH release and TrGSDME cleavage (Figure 2L and M, Figure
161 2-figure supplement 3B and C). These results indicated that TrCASP3/7 cleavage was
162 required to activate TrGSDME-mediated pyroptosis.

163

164 **GSDME-CT domain mediates the recognition by CASP7.**

165 Since it is known that human GSDME is cleaved by CASP3 but not by CASP7, the
166 observation that TrGSDME was cleaved by both human and fish CASP3/7 intrigued us
167 to explore the underlying mechanism. We found that unlike TrCASP1, both HsCASP3
168 and 7 exhibited proteolytic activity towards the consensus CASP3/7 recognition motif,
169 DMPD, in HsGSDME, but HsCASP7 failed to cleave HsGSDME (Figure 3A and B,
170 Figure 3-figure supplement 1 and 2). Structure analysis revealed that compared to
171 TrGSDME, HsGSDME possesses two additional regions with the possibility to form a
172 loop (261-266 aa) and an α -helix (281-296 aa), respectively (Figure 3C). We tested
173 whether deletion of these two regions could confer HsCASP7 cleavage on HsGSDME.
174 The results showed that similar to the wild type HsGSDME, the loop deletion mutant
175 (Δ 261-266) and the α -helix deletion mutant (Δ 281-296) were cleaved by HsCASP3 but
176 not by HsCASP7 (Figure 3D). Since the NT and CT domains play different roles in
177 GSDM structure maintenance, we constructed GSDME chimeras consisting of
178 HsGSDME-NT plus TrGSDME-CT (named HsNT-TrCT), or TrGSDME-NT plus
179 HsGSDME-CT (named TrNT-HsCT) (Figure 3E). Compared with wild type
180 HsGSDME and TrGSDME, chimeric HsNT-TrCT was cleaved by HsCASP3/7,
181 whereas TrNT-HsCT was cleaved only by HsCASP3 (Figure 3F-I), suggesting that the
182 CT domain determined the cleavability of GSDME by HsCASP7.

183

184 **The p10 subunit determines the substrate specificity of CASP7.**

185 Since as shown above, unlike HsCASP7, TrCASP7 was able to cleave HsGSDME
186 (Figure 3H), we compared the sequences of TrCASP7 and HsCASP7. The two CASPs
187 share 66.24% sequence identity. In these CASPs, the catalytic motifs SHG and QACRG
188 in the p20 subunit and the substrate binding motifs SWR and GSWF in the p10 subunit
189 are conserved (Figure 4A). To explore their substrate discrimination mechanism, we
190 constructed CASP7 chimeras consisting of HsCASP7 p20 plus TrCASP7 p10 (named
191 Hsp20-Trp10) or TrCASP7 p20 plus HsCASP7 p10 (named Trp20-Hsp10) (Figure 4B,
192 Figure 4-figure supplement 1A). Compared to the wild types (HsCASP7 and TrCASP7),
193 the chimeras exhibited comparable enzymatic activities towards the tetrapeptide
194 substrates DAVD and DMPD (Figure 4C). Like the wild types, both chimeras cleaved
195 TrGSDME and HsNT-TrCT (Figure 4D and E). By contrast, the Hsp20-Trp10 chimera
196 cleaved HsGSDME, whereas the Trp20-Hsp10 chimera did not cleave HsGSDME
197 (Figure 4F). Similar cleavage pattern was observed towards TrNT-HsCT (Figure 4G).
198 These observations indicated that the p10 subunit dictated the cleavage specificity of
199 CASP7 toward GSDME.

200

201 **The S234 of p10 is the key to HsCASP7 discrimination on HsGSDME.**

202 Comparative analysis of the p10 sequences of TrCASP7 and HsCASP7 identified 13
203 non-conserved residues. To examine their potential involvement in GSDME cleavage,
204 a series of site-directed mutagenesis was performed to swap the non-conserved residues
205 between HsCASP7 p10 and TrCASP7 p10 (Figure 5A, Figure 4-figure supplement 1B
206 and C). Of the 13 swaps created, S234N conferred on HsCASP7 the ability to cleave
207 HsGSDME (Figure 5B), whereas its corresponding swap in TrCASP7 (N245S)
208 markedly reduced the ability of TrCASP7 to cleave HsGSDME (Figure 5C). None of
209 the 13 swaps affected the ability of HsCASP7 or TrCASP7 to cleave TrGSDME (Figure
210 5D, E). HsCASP7-S234N cleaved HsGSDME in a dose-dependent manner (Figure 5F),
211 and the cleavage was not enhanced by the additional mutation of S247N&I248V
212 (Figure 5G). Additionally, HsCASP7 and HsCASP7-S234N exhibited similar cleavage

213 capacities against other CASP7 substrates, such as PARP1 and gelsolin (Figure 5-figure
214 supplement 1) (Erener et al., 2012; Walsh et al., 2008). Previous studies showed that
215 human CASP1/4 cleaved GSDMD through exosite interaction (Z. Liu et al., 2020; K.
216 Wang et al., 2020). However, this exosite is not conserved in HsCASP7, in which the
217 corresponding region forms a coil, not a β sheet (Figure 5-figure supplement 1).
218 Structural modeling showed that the residues Q276, D278, and H283 are close to the
219 region corresponding to the HsCASP1/4 exosite (Figure 5-figure supplement 1). Since
220 these three residues are also not conserved between HsCASP7 p10 subunit and
221 TrCASP7 p10 subunit, they were selected for mutagenesis to examine whether there
222 existed an exosite interaction between HsCASP7 and HsGSDME. The mutation results
223 showed that, similar to the wild type, the mutant Q276W&D278E&H283S was unable
224 to cleave HsGSDME (Figure 5G), suggesting that, unlike human GSDMD, HsGSDME
225 cleavage by CASPs probably did not involve exosite interaction. In contrast to
226 HsGSDME, TrGSDME cleavage by HsCASP7 was not affected by the mutation of
227 S234N, S234N plus S247N&I248V, or Q276W&D278E&H283S (Figure 5H).

228 All the above observed GSDME cleavages by CASP7 were verified in a cellular
229 system. When co-expressed in HEK293T cells, HsGSDME was cleaved by TrCASP7
230 but not by HsCASP7, while TrGSDME was cleaved by both TrCASP7 and HsCASP7
231 (Figure 5-figure supplement 3). Additionally, both HsGSDME and TrGSDME were
232 cleaved by HsCASP7-S234N as well as TrCASP7-N245S (Figure 5I and J). An
233 apparent dose effect was observed in the cleavage of HsGSDME by HsCASP7-S234N
234 (Figure 5K).

235

236 **Divergent and GSDME-independent evolution of CASP7 leads to the loss of** 237 **GSDME-activation function in mammalian CASP7.**

238 Given the importance of N234 in p10 for HsCASP7 cleavage of HsGSDME, we
239 analyzed the sequence conservation of this locus in CASP3/7. For CASP3, a highly
240 conserved Asn residue (corresponding to HsCASP7 S234) was found immediately after
241 the SWR motif in vertebrates (Figure 6A and B). In HsCASP3, this conserved Asn is

242 at position 208. We examined whether this residue was functionally essential to
243 HsCASP3. Compared to the wild type, the N208S mutant exhibited much weaker
244 cleavage of HsGSDME (Figure 6C, Figure 4-figure supplement 1D). Similar weakened
245 cleavage was also observed in HEK293T cells co-expressing HsCASP3-N208S and
246 HsGSDME (Figure 6D). These results indicated that the Asn residue was also critical
247 for HsCASP3 cleavage of HsGSDME.

248 For CASP7, an Asn at the position corresponding to HsCASP7 S234 is highly
249 conserved in teleost, amphibians, reptiles and birds, but not in mammals (Figure 6A).
250 In mammals, the corresponding Asn is present in most non-primate species, such as
251 mouse *Mus musculus* and bovine *Bos taurus*, but is replaced by Ser in primate, such as
252 human *H. sapiens*, gorillas *Gorilla gorilla*, and chimpanzee *Pan troglodytes* (Figure
253 6E, Figure 6-figure supplement 1). We examined whether mouse CASP7 (MmCASP7)
254 could cleave GSDME. The results showed that MmCASP7 cleaved HsGSDME in a
255 dose-dependent manner, and the cleaving ability was abrogated by N234S mutation
256 (Figure 6G and H, Figure 4-figure supplement 1E). By contrast, neither MmCASP7 nor
257 MmCASP7-N234S was able to cleave mouse GSDME (MmGSDME) (Figure 6I).
258 Mouse CASP3 (MmCASP3), however, cleaved MmGSDME efficiently (Figure 6-
259 figure supplement 2). These results indicated the existence of an intra-species barrier
260 for GSDME cleavage by CASP7 in some mammals, suggesting independent evolutions
261 of CASP7 and GSDME.

262

263 Discussion

264 GSDME is an ancient member of the GSDM family existing ubiquitously in vertebrate
265 from teleost to mammals (Broz, Pelegrin, & Shao, 2020; De Schutter et al., 2021). In
266 human, GSDME is specifically cleaved by CASP3 at the consensus motif DMPD, but
267 it is not cleaved by CASP7, which recognizes the same DxxD motif. CASP3 cleavage
268 releases the pyroptosis-inducing NT fragment from the association of the inhibitory CT
269 fragment and switches cell death from apoptosis to pyroptosis (Rogers et al., 2017;
270 Wang et al., 2017). In this study, we found that different from human GSDME, a

271 pufferfish GSDME belonging to the teleost GSDMEa lineage was specifically cleaved
272 by both CASP3 and CASP7 at the site of DAVD to liberate the pyroptosis-inducing NT
273 fragment. In teleost, there generally exist two GSDME orthologs, named GSDMEa and
274 GSDMEb. Comparative study on the functions of GSDMEa and GSDMEb is scarce,
275 and it remains to be explored whether clear physiological roles are played by these two
276 orthologs in fish. As executors of pyroptosis, teleost GSDMEs are activated via
277 different mechanisms. In tongue sole *Cynoglossus semilaevis*, GSDMEb is preferably
278 cleaved by CASP1 to trigger pyroptosis (Jiang, Gu, Zhao, & Sun, 2019); in zebrafish
279 *Danio rerio*, GSDMEb is activated by caspy2 (CASP5 homolog) through the NLR
280 family pyrin domain containing 3 (NLRP3) inflammasome signaling pathway (Li et al.,
281 2020; Z. Wang et al., 2020). These observations indicate that similar to mammalian
282 GSDMD, teleost GSDMEb is activated by inflammatory CASPs. In contrast, teleost
283 GSDMEa is cleaved mainly by apoptotic CASPs. In zebrafish, GSDMEa is cleaved by
284 CASP3 to generate the pyroptotic NT fragment (Wang et al., 2017). In turbot
285 *Scophthalmus maximus*, GSDMEa is bi-directionally regulated by CASP3/7, which
286 activate GSDMEa, and CASP6, which inactivates GSDMEa (Xu, Jiang, Yu, Yuan, &
287 Sun, 2022). In common carp *Cyprinus carpio haematopterus*, GSDMEa is cleaved by
288 CASP3 and induces pyroptosis (Zhao et al., 2022). The complicated scenario of
289 GSDME-mediated pyroptosis signaling in fish is likely due to the reason that, unlike
290 mammals that have multiple GSDM members to induce pyroptosis under different
291 conditions, fish have only GSDME to induce pyroptotic cell death. Regulation by
292 different CASPs may represent a mechanism that enables fish GSDME to execute the
293 orders of different pyroptotic signals.

294 It is intriguing that, although HsCASP3 and 7 were indistinguishable in proteolysis
295 towards the consensus tetrapeptide, they differed remarkably in the cleavage of
296 HsGSDME and TrGSDME. HsCASP3, but not HsCASP7, cleaved HsGSDME,
297 whereas both HsCASP3 and 7 cleaved TrGSDME at the same site. NT/CT domain
298 swapping between HsGSDME and TrGSDME showed that chimeric GSDMEs
299 containing the TrGSDME-CT domain were cleaved by human as well as pufferfish

300 CASP7 regardless of the source of the NT, whereas chimeric GSDMEs containing the
301 HsGSDME-CT domain were resistant to HsCASP7. These results indicate that the CT
302 domain, which is well accepted to exert an inhibitory effect on the pore-forming NT
303 domain (Z. Liu et al., 2019; J. Shi et al., 2015), is the target of HsCASP7 discrimination,
304 and hence determines the cleavability of GSDME by HsCASP7. Recently, Wang and
305 Liu studied the molecular mechanism of human GSDMD recognition by CASP1/4, and
306 showed that GSDMD-CT interacted with CASP1/4 exosites through binding to a
307 hydrophobic pocket, which enhanced the recognition by CASP1/4 and contributed to
308 tetrapeptide sequence-independent cleavage of GSDMD (Z. Liu et al., 2020; K. Wang
309 et al., 2020). Different from GSDMD, we found that for TrGSDME, mutation of either
310 the P4 (D255) or P1 (D258) residue of the consensus motif ²⁵⁵DAVD²⁵⁸ made it
311 resistant to CASP3/7, implying a lack of DAVD-independent cleavage mechanism.
312 Similarly, a previous study observed that human and mouse GSDME harboring
313 cleavage site (P4 or P1) mutation resisted CASP3 cleavage (Wang et al., 2017). These
314 results suggest that, compared to GSDMD, GSDME has a distinct enzyme-substrate
315 engagement mode that, as demonstrated in the present study, involves GSDME-CT.

316 Unlike HsCASP7, which was unable to cleave HsGSDME, TrCASP7 effectively
317 cleaved HsGSDME. The swapping of p20 between HsCASP7 and TrCASP7 revealed
318 that the two catalytic motifs in p20, i.e., SHG and QACRG, were not involved in
319 HsCASP7 discrimination on GSDME. SWR and GSWF are known to be responsible
320 for CASP substrate binding (Chai et al., 2001; Riedl et al., 2001). However, in our
321 study, we found that these two motifs are conserved in human and pufferfish CASP7,
322 and they had no apparent effect on the substrate discrimination of HsCASP7. By
323 contrast, the chimeric HsCASP7 containing the p10 subunit of TrCASP7 acquired the
324 ability to cleave HsGSDME, indicating an important role of p10. By a series of residue
325 swapping and mutation analyses, we discovered that the residue in the position of 245
326 (in TrCASP7) or 234 (in HsCASP7) of the p10 subunit was the key that determined
327 CASP7 cleavage of HsGSDME. Sequence analysis revealed that N245 is highly
328 conserved in the CASP7 of teleost, amphibians, reptiles and birds. In contrast, this Asn

329 residue was changed to Ser in the CASP7 of mammals, especially primates, resulting
330 in divergence of CASP7 from CASP3 in substrate recognition and cleavage. Since,
331 unlike non-mammalian vertebrates, mammals possess multiple pyroptotic GSDM
332 members (GSDMA, B, C, D, and E), the functional divergence of CASP7 from CASP3
333 may have occurred as a result of specific and non-redundant regulation of programmed
334 cell death mediated by different executor molecules. One of the physiological
335 consequences is that CASP7 frees itself from the GSDME-mediated pathway and thus
336 is able to be engaged in other cellular processes. Another physiological consequence is
337 that GSDME activation is limited to CASP3 cleavage, thus restricting GSDME activity
338 to situations more specific, such as that inducing CASP3 activation. More physiological
339 consequences of CASP3/7 divergence in GSDME activation need to be explored in
340 future studies. With the exception of primates, the conserved Asn is present in a large
341 number of mammals, including mice. Mouse CASP7 was able to cleave HsGSDME,
342 further supporting the importance of the Asn in CASP7 recognition and cleavage. The
343 inability of mouse CASP7 to cleave mouse GSDME is consistent with a previous report
344 that CASP7 deletion had little effect on mouse GSDME-mediated pyroptosis (Sarhan
345 et al., 2018). These results suggest mutually independent evolution of GSDME and
346 CASP7 in mice, which likely has distanced GSDME and CASP7 from each other.
347 Similarly, human CASP7 and GSDME may also have undergone independent
348 evolution, which leads to the disengagement of GSDME from the substrate relationship
349 with CASP7.

350 In conclusion, we identified a teleost GSDME that can be cleaved by fish and human
351 CASP3/7 to trigger pyroptosis. The GSDME-CT domain and the CASP7 p10 subunit
352 are critical in the determination of CASP7 cleavage of GSDME. Within the p10 subunit,
353 a single residue plays a key role in CASP7 substrate recognition and cleavage. Our
354 results reveal the molecular basis of the functional divergence of CASP7 and CASP3,
355 and suggest separate evolutions of CASP7 and GSDME in mammals. These findings
356 add new insights into CASP-regulated GSDME activation in lower and higher
357 vertebrates.

358 **Materials and Methods**

359 **Animal, ethics, and cell line**

360 Clinically healthy pufferfish (*Takifugu rubripes*) were obtained from a local fish farm.
361 In the laboratory, the fish were maintained at 19-20°C in aerated seawater as reported
362 previously (Xu et al., 2022). For euthanization, the fish were immersed in excess
363 tricaine methane sulfonate (Sigma, St. Louis, MO). The animal experiments were
364 approved by the Ethics Committee of institute of Oceanology, Chinese Academy of
365 Sciences. HEK293T cells (ATCC, Rockville, MD, USA) were cultured in DMED
366 medium (Corning, NY, USA) supplemented with 10% fetal bovine serum (ExCell Bio,
367 Shanghai, China) at 37°C in a 5% CO₂ incubator.

368

369 **Sequence analysis**

370 The sequences of 742 CASP3 and 758 CASP7 used in this study were downloaded
371 from NCBI Orthologs. TrCASP7 and HsCASP7 sequences were aligned with Clustal
372 W program (www.ebi.ac.uk/clustalw/) and visualized with ESPript 3.0
373 (<http://espript.ibcp.fr/ESPript/cgi-bin/ESPript.cgi/>) (Robert & Gouet, 2014).
374 Conservation of S/N234 in CASP3/7 was analyzed via Weblogo3
375 (<https://weblogo.threeplusone.com/>) (Crooks, Hon, Chandonia, & Brenner, 2004), and
376 the sequences of CASP3/7 used are shown in [Table 1](#) and [Table 2](#). The phylogenetic
377 analysis of GSDME was performed as reported previously (Xu et al., 2022). The
378 phylogenetic relationship of major mammalian clades was estimated based on the
379 Mammals birth-death node-dated completed trees in VertLife (Upham, Esselstyn, &
380 Jetz, 2019). The phylogenetic tree was subsequently viewed and edited in iTOL
381 (Letunic & Bork, 2021). The icons representing phylogeny clades were retrieved from
382 the PhyloPic (<http://www.phylopic.org/>), with the detailed credentials provided in
383 [Table 3](#).

384

385 **Antibodies and immunoblotting**

386 Monoclonal anti-mouse (ab215191) and anti-human (ab221843) GSDME antibodies
387 were purchased from Abcam (Cambridge, MA). Antibodies against β -actin (AC026),
388 Flag (AE005), Myc (AE010), His (AE003), and mCherry (AE002) were purchased
389 from Abclonal (Wuhang, China). Mouse polyclonal antibody against TrGSDME was
390 prepared as reported previously (Xu et al., 2022). Anti-serum (1:1000 dilution) was
391 used for immunoblotting. Immunoblotting was performed as reported previously (Zhao
392 & Sun, 2022). Briefly, the samples were fractionated in 12% SDS-polyacrylamide gels
393 (GenScript, Piscataway, NJ, USA). The proteins were transferred to NC membranes,
394 immunoblotted with appropriate antibodies and visualized using an ECL kit (Sparkjade
395 Biotechnology Co. Ltd., Shandong, China).

396

397 **Gene cloning and mutagenesis**

398 Total RNA was extracted from pufferfish kidney using the FastPure Cell/Tissue Total
399 RNA Isolation Kit V2 (Vazyme Biotech Co. Ltd., Nanjing, China). cDNA synthesis
400 was performed with Revert Aid First Strand cDNA Synthesis Kit (Thermo Fisher
401 Scientific, Waltham, MA, United States). The coding sequences (CDSs) of pufferfish
402 GSDME (Accession number XP_029701077.1) and CASP3/7 were amplified by PCR.
403 The CDSs of mouse GSDME and CASP3/7 were synthesized by Sangon Biotech
404 (Shanghai, China). Site-directed mutagenesis was performed using Hieff Mut Site-
405 Directed Mutagenesis Kit (Yeasen, Shanghai, China). Truncates of TrGSDME were
406 created as reported previously (Xu et al., 2022) and subcloned into pmCherry-N1
407 (Clontech, Mountain View, CA, USA). For the construction of GSDME and CASP7
408 chimera, the CDSs of GSDME NT/CT and CASP7 p10/p20 were obtained by PCR and
409 ligated. The constructs of the truncate and chimeric proteins are shown in [Table 4](#). All
410 sequences/constructs were verified by sequencing analysis. The primers used are listed
411 in [Table 5](#).

412

413 **Plasmids and transient expression**

414 For expression in mammalian cell expression, the CDSs of GSDME (pufferfish, human
415 and mouse) were subcloned into pCS2-3×Flag (Wang et al., 2017) or pmCherry-N1
416 and CDSs encoding CASP (pufferfish, human and mouse) were subcloned into p-
417 CMV-Myc (Clontech, Mountain View, CA, USA) or pCS2-Myc (Wang et al., 2017).
418 All plasmids were prepared with endotoxin-free plasmid kit (Sparkjade Biotechnology
419 Co. Ltd., Shandong, China). For transient expression, HEK293T cells were cultured in
420 96- or 24-well plates (Corning, NY, USA) overnight, and then transfected with 100
421 ng/well (96-well plates) or 500 ng/well (24-well plates) indicated plasmids using
422 Liopfectamine 3000 (Invitrogen, USA) for 24 h or as specified. For GSDME
423 processing, the plasmids of GSDME and CASP were co-transfected in HEK293T cells
424 as described above and cultured continuously for 24 h or as specified. Cells were
425 harvested and lysed with RIPA Lysis Buffer (Beyotime, Shanghai, China) for
426 immunoblotting. The primer used are shown in [Table 5](#).

427

428 **Protein purification**

429 Recombinant GSDMEs and CASPs are all soluble and were purified as described
430 previously (Jiang, Zhou, Sun, Zhang, & Sun, 2020; Xu et al., 2022). Briefly, the CDSs
431 of TrGSDME and CASP variants were each cloned into pET30a (+), and the CDSs of
432 HsGSDME, MmGSDME, and chimeric GSDME were each cloned into pET28a-
433 SUMO. The recombinant plasmids were introduced into *Escherichia coli* Transetta
434 (DE3) (TransGen, Beijing, China) by transformation. The transformants were cultured
435 in Luria broth (LB) at 37°C until logarithmic growth phase. Isopropyl-β-D-
436 thiogalactopyranoside (0.3 mM) was added to the medium, followed by incubation at
437 16°C for 20 h. Bacteria were harvested and lysed, and the supernatant was collected for
438 protein purification with Ni-NTA columns (GE Healthcare, Uppsala, Sweden). The
439 proteins were dialyzed with PBS at 4 °C and concentrated.

440

441 **CASP activity**

442 To measure their proteolytic activity, recombinant CASPs were incubated separately
443 with various colorimetric substrates at 37°C for 2 h as described previously (Jiang et
444 al., 2020; Xu et al., 2022), and then monitored for released pNA at OD405. To compare
445 their substrate preference, TrCASP3/7 and HsCASP3/7 were incubated with Ac-
446 DAVD-pNA or Ac-DMPD-pNA (Science Peptide Biological Technology Co., Ltd,
447 shanghai, China) at 37°C for 30 min, and the released pNA was measured at OD405.
448 The CASP activity in cells transfected TrCASP3/7 was determined as described
449 previously (Xu et al., 2022).

450

451 **GSDME cleavage by CASPs**

452 GSDME cleavage assay was performed as described previously (Jiang et al., 2020; Xu
453 et al., 2022). Briefly, recombinant GSDME was incubated with 1 U of recombinant
454 HsCASP1 ,2, 3, 6, 7, 8 and 9 (Enzo Life Sciences, Villeurbanne, France) at 37°C for 2
455 h in the reaction buffer (50 mM Hepes (pH 7.5), 3 mM EDTA, 150 mM NaCl, 0.005%
456 (v/v) Tween 20, and 10 mM DTT). TrGSDME, HsGSDME or their chimeras were
457 treated with 1 U of TrCASP3/7, HsCASP3/7, or their chimeras for 2 h as above in
458 reaction buffer. After incubation, the samples were boiled in 5 x Loading Buffer
459 (GenScript, Piscataway, NJ, USA) and subjected to SDS-PAGE or immunoblotting
460 with indicated antibody.

461

462 **Cell death examination**

463 Cell death examination by microscopy was performed as reported previously (Jiang et
464 al., 2020; Xu et al., 2022). Briefly, HEK293T cells were plated into 35 mm glass-
465 bottom culture dishes (Nest Biotechnology, Wuxi, China) at about 60% confluency and
466 subjected to the indicated treatment for 24 h. To examine cell death morphology,
467 propidium iodide (PI) (Invitrogen, Carlsbad, CA, USA) and DiO (Solarbio, Beijing,
468 China) were added to the culture medium, and the cells were observed with a Carl Zeiss
469 LSM 710 confocal microscope (Carl Zeiss, Jena, Germany). To video the cell death
470 process, the cells were transfected with 1 µg pmCherry-N1 vector expressing

471 TrGSDME-NT for 16 h, and cell death was recorded using the above microscope. Cell
472 death measured by lactate dehydrogenase (LDH) assay was performed as reported
473 previously (Xu et al., 2022).

474

475 **Statistical analysis**

476 The Student's t test and one-way analysis of variance (ANOVA) were used for
477 comparisons between groups. Statistical analysis was performed with GraphPad Prism
478 7 software. Statistical significance was defined as $P < 0.05$.

479

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487

488 **Author Contributions**

489 **Hang Xu:** Methodology; Investigation; Visualization; Writing—original draft. ;**Zihao**
490 **Yuan:** Investigation. **Kunpeng Qin:** Investigation; **Shuai Jiang:** Conceptualization;
491 Methodology; Supervision; Writing—review & editing. **Li Sun:** Conceptualization;
492 Supervision; Writing—review & editing.

493

494 **Competing Interest Statement**

495 The authors declare that they have no competing interests.

496

497

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- 658
- 659

660 **Figure legends**

661 **Figure 1. Cleavage of TrGSDME by caspases.** (A, B) TrGSDME was treated with 1
662 U of different HsCASP3/7 for 2 h and then subjected to SDS-PAGE (A) and
663 immunoblotting (B). (C) SDS-PAGE analysis of purified TrCASP3/7. The p10 and p20
664 subunits are indicated. (D) TrCASP3/7 cleavage of different colorimetric CASP
665 substrates was monitored by measuring released pNA. The values are the means \pm SD
666 of three replicates. (E) TrCASP3/7 (0.25-8U) were incubated with Ac-DEVD-pNA,
667 and time-dependent release of pNA was measured. (F, G) SDS-PAGE (F) and
668 immunoblotting (G) analysis of TrGSDME cleavage by TrCASP3/7 (0.25-4U). (H)
669 TrGSDME cleavage by TrCASP3/7 was determined in the presence or absence of 20
670 μ M Z-DEVD-FMK or Z-VAD-FMK. (I, J) TrGSDME wild type (WT) and mutants
671 (D255R or D258A) were incubated with TrCASP3/7 for 2 h, and the cleavage was
672 determined by SDS-PAGE (I) and immunoblotting (J). (K) A schematic of TrGSDME
673 cleavage by TrCASP3/7. The arrow indicates cleavage site. For all panels, FL, full
674 length; NT, N-terminal fragment; CT, C-terminal fragment.

675 **Figure 2. The pyroptotic activity of TrGSDME.** (A-C) HEK293T cells were
676 transfected with C-terminally mCherry-tagged TrGSDME full length (FL) or truncate
677 (NT or CT) for 24 h, and then analyzed for TrGSDME expression (A), morphological
678 change (B), and LDH release (C). Scale bars, 50 μ m (A) and 20 μ m (B). (D) HEK293T
679 cells were transfected with the backbone vector or vector expressing Myc-tagged
680 TrGSDME-NT for 24 h. The cell nuclei and membrane were stained with PI and DiO,
681 respectively. Scale bar, 10 μ m. (E) HEK293T cells were transfected with mCherry-
682 tagged TrGSDME-NT for 16 h, and the progression of cell death was shown by time-
683 lapse microscopic imaging. Scale bar, 10 μ m. (F) HEK293T cells were transfected with
684 TrCASP3/7 for 24 h, and the proteolytic activity of TrCASP3/7 was assessed by
685 treatment with fluorogenic Ac-DEVD-AFC. (G) Phase-contrast images of HEK293T
686 cells transfected with the indicated vectors for 24 h. Scale bar, 20 μ m. (H, I) LDH
687 release (H) and TrGSDME cleavage and TrCASP3/7 expression (I) in the above

688 transfected cells were determined. For panel (I), FL, full-length; CT, C-terminal
689 fragment. **(J, K)** HEK293T cells co-expressing TrGSDME and TrCASP3/7 for 24 h in
690 the presence or absence (control) of 10 μ M Z-VAD-FMK or Z-DEVD-FMK were
691 subjected to microscopy (J) and LDH measurement (K). Scale bar, 20 μ m. **(L, M)**
692 HEK293T cells expressing TrCASP3/7 plus TrGSDME or TrCASP3/7 plus the
693 D255R/D258A mutant were subjected to microscopy (L) and LDH measurement (M).
694 Scale bar, 20 μ m. For panels (C, F, H, K and M), values are the means of three
695 experimental replicates and shown as means \pm SD. $**P < 0.01$, $***P < 0.001$.

696 **Figure 3. GSDME-CT domain determines the recognition by CASP7.** **(A)**
697 HsGSDME was treated with 1 U of various HsCASPs for 2 h, and the products were
698 analyzed by immunoblotting with anti-HsGSDME-CT antibody. **(B)** HsCASP3/7 and
699 TrCASP3/7 were incubated with Ac-DAVD- ρ NA or Ac-DMPD- ρ NA at 37°C for 30
700 min, and the proteolytic activity was determined. Data are expressed as the means \pm SD
701 of three replicates. **(C)** Structure analysis of the linker region from HsGSDME and
702 TrGSDME with mouse GSDMA3 (PDB:5b5r) as the template. Two regions that may
703 form α -helix and loop are indicated. **(D)** HsGSDME wild type (WT) and mutants
704 (Δ 261-266 and Δ 281-296) were treated with or without 1 U of HsCASP3/7 for 2 h, and
705 the cleaved fragments were subjected to immunoblotting with anti-HsGSDME-CT
706 antibody. **(E)** Schematics of the chimeric GSDME constructs. **(F-I)** TrGSDME (F),
707 HsNT-TrCT (G), HsGSDME (H) and TrNT-HsCT (I) were incubated with 1 U of
708 TrCASP3/7 or HsCASP3/7 for 2 h, and the cleavage was determined by
709 immunoblotting with anti-TrGSDME or anti-HsGSDME-CT antibodies. For panels (A,
710 D, and F-I): FL, full length; NT, N-terminal fragment; CT, C-terminal fragment.

711 **Figure 4. The p10 subunit of CASP7 is essential for discrimination on GSDME.**
712 **(A)** Sequence alignment of HsCASP7 and TrCASP7 with HsCASP7 (PDB:1K86) as
713 the template. α -helices and β -strands are indicated. The motifs involved in catalytic
714 reaction (SHG and QACRG) and substrate binding (SWR and GSWF) are boxed in red.
715 **(B)** Schematics of CASP7 chimeras. **(C)** The proteolytic activities of TrCASP7,
716 HsCASP7 and their chimeras were determined by cleaving Ac-DAVD- ρ NA and Ac-

717 DMPD-pNA. Data are expressed as the means \pm SD of three replicates. **(D-G)**
718 TrGSDME (D), HsNT-TrCT (E), HsGSDME (F) or TrNT-HsCT (G) were incubated
719 with TrCASP7, HsCASP7 and their chimeras for 2 h. The products were assessed by
720 immunoblotting with anti-TrGSDME or anti-HsGSDME-CT antibodies. For panels (D-
721 G): FL, full length; NT, N-terminal fragment; CT, C-terminal fragment.

722 **Figure 5. Functional importance of the non-conserved residues in the p10 of**
723 **HsCASP7 and TrCASP7.** **(A)** Sequence alignment of the p10 region in HsCASP7 and
724 TrCASP7. The arrows indicate non-conserved residues. **(B, C)** HsGSDME was treated
725 with wild type (WT) or mutant HsCASP7 (B) or TrCASP7 (C) for 2 h. The cleavage
726 was determined by immunoblotting with anti-HsGSDME-CT antibody. 205TME,
727 HsCASP7 with TME insertion at position 205; 213 Δ TME, TrCASP7 with TME deleted
728 at position 213. **(D, E)** TrGSDME was incubated with wild type (WT) or mutant
729 HsCASP7 (D) or TrCASP7 (E) for 2 h. The cleavage was determined by
730 immunoblotting with anti-TrGSDME antibody. **(F)** HsGSDME was incubated with
731 HsCASP7 or the S234N mutant (0.25-4U) for 2 h. HsGSDME cleavage was analyzed
732 as above. **(G, H)** HsGSDME (G) and TrGSDME (H) were treated with wild type or
733 mutant HsCASP7 for 2 h. The cleavage was determined as above. **(I, J)** TrGSDME (I)
734 or HsGSDME (J) was co-expressed with wild type or mutant TrCASP7/HsCASP7 for
735 24 h. GSDME cleavage, CASP7 and β -actin were determined as above. **(K)** HsGSDME
736 was co-expressed with HsCASP7 or HsCASP7-S234N for 48 h. Cleavage of GSDME
737 was detected as above. For panels (B-K): FL, full length; NT, N-terminal fragment; CT,
738 C-terminal fragment.

739 **Figure 6. Conservation and functional importance of S/N234 in CASP3/7.** **(A)**
740 Weblogo analysis of the conservation of S/N234 in CASP3/7 in Mammalia, Aves,
741 Reptilia, Amphibia and Osteichthyes. **(B)** Sequence alignment of HsCASP3/7. The S/N
742 residues are indicated by red arrow. **(C)** HsGSDME was treated with HsCASP3 or
743 HsCASP3-N208S for 2 h, and then subjected to immunoblotting with anti-HsGSDME-
744 CT antibody. **(D)** HsGSDME was co-expressed in HEK293T cells with different doses
745 of HsCASP3 or HsCASP3-N208S for 24 h. The cell lysates were immunoblotted to

746 detect HsGSDME, CASP3, and β -actin. **(E)** Sequence alignment of the S/N234 region
747 in the CASP7 of primate (shaded cyan) and non-primate mammals. The S/N residues
748 are indicated by red arrow. Asterisks indicate identical residues. **(F)** Alignment of
749 human and mouse CASP7 sequences. The S/N234 residues are indicated by arrow. **(G)**
750 HsGSDME was treated with different units of HsCASP7 or MmCASP7 for 2 h, and the
751 cleavage was assessed by immunoblotting with anti-HsGSDME-CT antibody. **(H)**
752 HsGSDME was incubated with different units of MmCASP7 or its mutant (N234S) for
753 2 h, and HsGSDME cleavage was analyzed as above. **(I)** MmCASP7 was treated with
754 HsCASP7, MmCASP7 or MmCASP7-N234S for 2 h. The cleavage was determined by
755 immunoblotting with anti-MmGSDME-NT antibody. For panels (C, D, and G-I), FL,
756 full length; NT, N-terminal fragment; CT, C-terminal fragment.
757

758 **Figure 1-figure supplement 1. Phylogenetic analysis of the TrGSDME used in this**
759 **study.** The phylogenetic tree was constructed using 15 fish GSDMEa (red) and 51 fish
760 GSDMEb (blue) collected from NCBI databank. The TrGSDME used in this study is
761 indicated by a red star.

762 **Figure 1-figure supplement 2. Cleavage of TrGSDME mutants by HsCASP3/7.**
763 TrGSDME wild type (WT) and mutants (D255R and D258A) were treated with or
764 without (-) HsCASP3/7 for 2 h, and the cleavage was determined by immunoblotting
765 with anti-His antibody. FL, full length; CT, cleaved C-terminal region.

766 **Figure 2-figure supplement 1. Ectopic expression of TrGSDME-FL and truncates.**
767 HEK293T cells were transfected with the backbone vector or the vector expressing
768 mCherry-tagged TrGSDME full length (FL) or truncates (NT or CT) for 24 h. The cell
769 lysate was then immunoblotted with antibody against mCherry or β -actin.

770 **Figure 2-figure supplement 2. The cell death-inducing ability of TrGSDME-FL**
771 **and truncates.** HEK293T cells were transfected with the backbone vector or the vector
772 expressing mCherry-tagged TrGSDME full length (FL) or truncate (NT or CT) for 24
773 h. The cells were then stained with Sytox Green and examined with a microscope. Scale
774 bar, 30 μ m.

775 **Figure 2-figure supplement 3. Activation of TrGSDME-mediated pyroptosis by**
776 **TrCASP3/7. (A)** HEK293T cells were transfected with the backbone vector or the
777 vector expressing the indicated protein for 24 h and then stained with Sytox Green.
778 Scale bars, 30 μ m. **(B, C)** HEK293T cells expressing wild type (WT) or mutant
779 TrGSDME together with or without (control) TrCASP3 or TrCASP7 were subjected to
780 Sytox Green staining **(B)** and TrGSDME cleavage analysis by immunoblot with
781 antibodies against mCherry (upper), Myc (middle), and actin (lower) **(C)**. Scale bar, 30
782 μ m.

783

784 **Figure 3-figure supplement 1. Cleavage of HsGSDME by caspases.** Recombinant
785 HsGSDME was incubated with or without (-) HsCASP1, 2, 3, 6, 7, 8, and 9 for 2 h, and
786 the cleavage was determined by SDS-PAGE. M, molecular markers. FL, full length;
787 NT, N-terminal fragment; CT, C-terminal fragment.

788 **Figure 3-figure supplement 2. Proteolytic activity of TrCASP1.** TrCASP1 was
789 incubated with colorimetric Ac-YVAD- ρ NA, Ac-DAVD- ρ NA, and Ac-DMPD- ρ NA
790 for 1 h, and the cleavage was determined by measuring the release of ρ NA. Values are
791 the means of triplicate experiments and shown as means \pm SD..

792 **Figure 4-figure supplement 1. Preparation of recombinant proteins.** (A) SDS-
793 PAGE analysis of purified Hsp20-Trp10 and Trp20-Hsp10. (B, C) Purified TrCASP7
794 (B) and HsCASP7 (C) were subjected to SDS-PAGE (upper panel) and immunoblotting
795 with anti-His tag antibody (lower panel). (D) HsCASP3 and its mutant (N208S) were
796 analyzed by SDS-PAGE. (E) SDS-PAGE analysis of purified MmCASP7 and its
797 mutant (N234S). In all panels, the p10 and p20 subunits are indicated.

798 **Figure 5-figure supplement 1. Cleavage of poly (ADP-ribose) polymerase 1**
799 **(PARP1) and gelsolin by HsCASP7.** HEK293T cells were transfected with C-terminal
800 Flag-tagged PARP1 (left panel) or gelsolin (right panel) for 24 h. The cell lysates were
801 incubated with or without (-) different units of HsCASP7 and HsCASP7-S234N for 2
802 h and then subjected to immunoblotting with anti-Flag antibody.

803 **Figure 5-figure supplement 2. Structural analysis of HsCASP1/4/7.** (A) The
804 structures of HsCASP1 (PDB: 6kn0, gray), HsCASP4 (PDB: 6kmz, green), and
805 HsCASP7 (PDB: 6kn0, purple) are superimposed. The β III/ β III' sheets of HsCASP1/4
806 are indicated by arrows; the residues Q276, D278, and H283 of HsCASP7 are shown
807 in cyan. (B) The sequence alignment of HsCASP1/4 β III/ β III' sheet with the
808 corresponding region of HsCASP7.

809 **Figure 5-figure supplement 3. GSDME cleavage by CASP7 in cellular system.**
810 TrGSDME (A) or HsGSDME (B) was co-expressed with different doses of
811 TrCASP7/HsCASP7 for 48 h in HEK293T cells. The expression/cleavage of

812 TrGSDME/HsGSDME and the expression of TrCASP7/HsCASP7 were determined by
813 immunoblotting. β -actin was used as a loading control.

814 **Figure 6-figure supplement 1. Conservation of S/N234 in mammalian CASP7.** The
815 four conserved amino acids are shown in LOGOs, with height representing the relative
816 proportion of the corresponding amino acid at that position. The S/N234 is presented
817 as the fourth LOGO and indicated with a red star.

818 **Figure 6-figure supplement 2. The cleavage of MmGSDME by MmCASP3.** (A)
819 SDS-PAGE analysis of purified MmCASP3. (B) MmGSDME was treated with 1U of
820 MmCASP3 for 2 h and then subjected to SDS-PAGE.

821

822

823 **Legends for Figure 1-video 1:**

824 **Figure 1-video 1 (separate file). TrGSDME-NT induces pyroptosis of HEK293T**
825 **cells.** HEK293T cells were transfected with the vector expressing mCherry-tagged
826 TrGSDME-NT, and time-lapse images of the cells were recorded with a confocal
827 microscope.

828

829 **Tables**

830 Table 1. The CASP3 sequences used for WebLogo analysis in this study.

831 Table 2. The CASP7 sequences used for WebLogo analysis in this study.

832 Table 3. The credits for the pictures used in this study.

833 Table 4. The constructs of the truncate and chimeric proteins.

834 Table 5. Primers used in this study.

835

836 **Table 1. The CASP3 sequences used for WebLogo analysis in this study.**

Class	Species	Accession Number
Mammalia	<i>Homo sapiens</i>	P42574
	<i>Mus musculus</i>	P70677
	<i>Sus scrofa</i>	Q95ND5
	<i>Canis lupus familiaris</i>	Q8MKI5
	<i>Saimiri boliviensis</i>	NP_001266895.1
	<i>Equus caballus</i>	NP_001157433.1
	<i>Mesocricetus auratus</i>	Q60431
	<i>Pan troglodytes</i>	XP_009446811.1
	<i>Ovis aries</i>	XP_014960045.1
	<i>Capra hircus</i>	NP_001273018.1
	<i>Pan paniscus</i>	XP_003823509.1
	<i>Ailuropoda melanoleuca</i>	XP_002914062.1
	<i>Oryctolagus cuniculus</i>	Q8MJC3
	<i>Loxodonta africana</i>	XP_023410784.1
	<i>Bos taurus</i>	Q08DY9
	<i>Macaca fascicularis</i>	Q2PFV2
	<i>Delphinapterus leucas</i>	XP_022426661.1
<i>Aotus nancymaae</i>	XP_012327995.1	
<i>Vulpes vulpes</i>	XP_025874057.1	

<i>Heterocephalus glaber</i>	EHB15384.1
<i>Bos mutus grunniens</i>	XP_010834058.1
<i>Balaenoptera acutorostrata</i>	XP_007177921.2
<i>Odobenus rosmarus divergens</i>	XP_012422221.1
<i>Tursiops truncatus</i>	XP_033704658.1
<i>Saimiri boliviensis</i>	AAV74267.1
<i>Theropithecus gelada</i>	XP_025242051.1
<i>Rhinopithecus roxellana</i>	XP_010376231.2
<i>Chlorocebus sabaeus</i>	XP_007998582.1
<i>Arvicanthis niloticus</i>	XP_034376775.1
<i>Castor canadensis</i>	JAV41953.1
<i>Urocyon parryi</i>	XP_026256383.1
<i>Neogale vison</i>	XP_044081570.1
<i>Lynx rufus</i>	XP_046947013.1
<i>Canis lupus dingo</i>	XP_025327613.1
<i>Balaenoptera acutorostrata scammoni</i>	XP_007177921.1
<i>Balaenoptera musculus</i>	XP_036694636.1
<i>Vicugna pacos</i>	XP_006198038.1
<i>Camelus bactrianus</i>	XP_010951604.1
<i>Eptesicus fuscus</i>	XP_008150114.1
<i>Pteropus alecto</i>	XP_006916432.1
<i>Ochotona princeps</i>	XP_004579072.1
<i>Equus quagga</i>	XP_046504640.1
<i>Talpa occidentalis</i>	XP_037382298.1
<i>Manis pentadactyla</i>	XP_036750218.1
<i>Gracilinanus agilis</i>	XP_044536929.1
Aves	
<i>Corvus hawaiiensis</i>	XP_048160132.1
<i>Taeniopygia guttata</i>	XP_030127125.3

<i>Pyrgilauda ruficollis</i>	XP_041335801.1
<i>Hirundo rustica</i>	XP_039921773.1
<i>Onychostruthus taczanowskii</i>	XP_041283907.1
<i>Passer montanus</i>	XP_039555833.1
<i>Molothrus ater</i>	XP_036238378.1
<i>Oxyura jamaicensis</i>	XP_035179722.1
<i>Melopsittacus undulatus</i>	XP_030904860.1
<i>Strigops habroptila</i>	XP_030349011.1
<i>Aythya fuligula</i>	XP_032043173.1
<i>Lonchura striata domestica</i>	OWK64686.1
<i>Geospiza fortis</i>	XP_005415590.1
<i>Camarhynchus parvulus</i>	XP_030803986.1
<i>Serinus canaria</i>	XP_030094646.1
<i>Manacus vitellinus</i>	XP_008923018.1
<i>Zonotrichia albicollis</i>	XP_005483896.1
<i>Apteryx rowi</i>	XP_025939844.1
<i>Numida meleagris</i>	XP_021249041.1
<i>Gallus gallus</i>	NP_990056.1
<i>Tyto alba</i>	XP_042663613.1
<i>Lagopus leucura</i>	XP_042726563.1
<i>Pygoscelis adeliae</i>	XP_009328950.1
<i>Calypte anna</i>	XP_030305365.1
<i>Aquila chrysaetos chrysaetos</i>	XP_029861401.1
<i>Haliaeetus leucocephalus</i>	XP_010568301.1
<i>Tyto alba</i>	XP_032854868.1
<i>Hirundo rustica</i>	XP_039921766.1
<i>Cygnus atratus</i>	XP_035413755.1
<i>Lepidothrix coronata</i>	XP_017664322.1

	<i>Amazona aestiva</i>	KQK77628.1
	<i>Chaetura pelagica</i>	XP_010000752.1
	<i>Antrostomus carolinensis</i>	XP_010171751.1
	<i>Nipponia nippon</i>	XP_009473093.1
	<i>Pelecanus crispus</i>	XP_009479867.1
	<i>Leptosomus discolor</i>	XP_009958677.1
	<i>Dryobates pubescens</i>	XP_009909179.1
	<i>Manacus vitellinus</i>	XP_008923018.1
	<i>Parus major</i>	XP_015479949.1
	<i>Falco rusticolus</i>	XP_037254769.1
	<i>Numida meleagris</i>	XP_021249038.1
Reptilia	<i>Pelodiscus sinensis</i>	XP_006128559.1
	<i>Gopherus evgoodei</i>	XP_030419000.1
	<i>Chelonia mydas</i>	XP_043402007.1
	<i>Dermochelys coriacea</i>	XP_043369528.1
	<i>Mauremys reevesii</i>	XP_039396469.1
	<i>Terrapene carolina triunguis</i>	XP_029766549.1
	<i>Trachemys scripta elegans</i>	XP_034627282.1
	<i>Python bivittatus</i>	XP_025018785.1
	<i>Pogona vitticeps</i>	XP_020658385.1
	<i>Zootoca vivipara</i>	XP_034967628.1
	<i>Podarcis muralis</i>	XP_028600040.1
	<i>Sceloporus undulatus</i>	XP_042323800.1
	<i>Lacerta agilis</i>	XP_033017209.1
	<i>Varanus komodoensis</i>	XP_044305785.1
	<i>Notechis scutatus</i>	XP_026522951.1
	<i>Pantherophis guttatus</i>	XP_034257179.1
	<i>Protobothrops mucrosquamatus</i>	XP_015687286.1

	<i>Eremias argus</i>	QCQ80689.1
	<i>Sceloporus undulatus</i>	XP_042323800.1
	<i>Gekko japonicus</i>	XP_015272894.1
	<i>Lacerta agilis</i>	XP_033017209.1
	<i>Notechis scutatus</i>	XP_026522951.1
	<i>Pseudonaja textilis</i>	XP_026553662.1
	<i>Sphaerodactylus townsendi</i>	XP_048364960.1
	<i>Thamnophis sirtalis</i>	XP_013910221.1
	<i>Crotalus tigris</i>	XP_039186723.1
	<i>Thamnophis elegans</i>	XP_032080611.1
	<i>Crocodylus porosus</i>	XP_019390228.1
	<i>Chelonoidis abingdonii</i>	XP_032656285.1
	<i>Chrysemys picta bellii</i>	XP_005282030.1
	<i>Mauremys mutica</i>	XP_044873973.1
Amphibia	<i>Xenopus laevis</i>	NP_001081225.1
	<i>Bufo gargarizans</i>	XP_044145804.1
	<i>Rana temporaria</i>	XP_040189928.1
	<i>Bufo bufo</i>	XP_040274658.1
	<i>Geotrypetes seraphini</i>	XP_033799250.1
	<i>Microcaecilia unicolor</i>	XP_030047368.1
	<i>Nanorana parkeri</i>	XP_018421176.1
	<i>Cynops orientalis</i>	AFN55260.1
	<i>Xenopus tropicalis</i>	NP_001120900.1
	<i>Rhinatrema bivittatum</i>	XP_029441076.1
	<i>Geotrypetes seraphini</i>	XP_033799250.1
Osteichthyes	<i>Paralichthys olivaceus</i>	XP_019958716.1
	<i>Danio rerio</i>	AWP39888.1
	<i>Dicentrarchus labrax</i>	ABC70997.1

<i>Larimichthys crocea</i>	NP_001290322.1
<i>Scophthalmus maximus</i>	AVW89178.1
<i>Cynoglossus semilaevis</i>	XP_016894801.1
<i>Cyprinus carpio</i>	XP_018965718.1
<i>Cyprinodon variegatus</i>	XP_015245088.1
<i>Cyprinodon tularosa</i>	XP_038127805.1
<i>Takifugu rubripes</i>	NP_001027871.1
<i>Salmo salar</i>	NP_001133393.1
<i>Oncorhynchus mykiss</i>	BAU69680.1
<i>Monopterus albus</i>	XP_020451082.1
<i>Nothobranchius furzeri</i>	KAF7225424.1
<i>Miichthys miiuy</i>	AHG06618.1
<i>Anguilla japonica</i>	AYC61977.1
<i>Scleropages Formosus</i>	XP_018591577.1
<i>Poecilia reticulata</i>	XP_008417983.1
<i>Poecilia formosa</i>	XP_007549682.1
<i>Oplegnathus fasciatus</i>	AFM09714.1
<i>Collichthys lucidus</i>	TKS72762.1
<i>Notothenia coriiceps</i>	XP_010794822.1
<i>Poeciliopsis prolifica</i>	JAO80483.1
<i>Haplochromis burtoni</i>	XP_005944370.2
<i>Cyprinus carpio</i>	AGU12796.1
<i>Cyprinus carpio</i>	AGU12795.1
<i>Electrophorus electricus</i>	XP_026884624.2
<i>Amphiprion melanopus</i>	AEA08874.1
<i>Archocentrus centrarchus</i>	XP_030591190.1
<i>Bagarius yarrelli</i>	TSN95700.1
<i>Oryzias melastigma</i>	XP_024138660.1

<i>Oryzias latipes</i>	NP_001098140.1
<i>Siniperca chuatsi</i>	ADK47519.1
<i>Kryptolebias marmoratus</i>	XP_017278585.1
<i>Astyanax mexicanus</i>	XP_022539261.1
<i>Miichthys miiuy</i>	AHG06618.1
<i>Xiphophorus couchianus</i>	XP_027872889.1
<i>Oreochromis niloticus</i>	ADJ57601.1
<i>Lutjanus peru</i>	QBY35776.1
<i>Mugil incilis</i>	QDK54780.1

837

838

839 **Table 2. The CASP7 sequences used for the WebLogo analysis in this study.**

Class	Species	Accession Number
Mammalia	<i>Homo sapiens</i>	P55210
	<i>Mus musculus</i>	P97864
	<i>Mesocricetus auratus</i>	P55214
	<i>Bos taurus</i>	XP_002698555.1
	<i>Pan troglodytes</i>	XP_003825650.1
	<i>Macaca fascicularis</i>	XP_011822901.1
	<i>Oryctolagus cuniculus</i>	XP_002718761.1
	<i>Ovis aries</i>	XP_042094975.1
	<i>Capra hircus</i>	XP_005698554.1
	<i>Ailuropoda melanoleuca</i>	XP_034519188.1
	<i>Saimiri boliviensis boliviensis</i>	XP_034519188.1
	<i>Equus caballus</i>	XP_014588811.1
	<i>Delphinapterus leucas</i>	XP_022424428.1
	<i>Sus scrofa</i>	XP_020928977.1
	<i>Pan paniscus</i>	XP_003825650.1
	<i>Loxodonta africana</i>	XP_010587277.1
	<i>Canis lupus familiaris</i>	XP_005637795.1
	<i>Oryctolagus cuniculus</i>	XP_002718761.1
	<i>Vulpes vulpes</i>	XP_025849552.1
	<i>Heterocephalus glaber</i>	XP_004838949.1
	<i>Uroditellus parryi</i>	XP_026244955.1
	<i>Equus asinus</i>	XP_014723324.1
	<i>Tachyglossus aculeatus</i>	XP_038614071.1
	<i>Camelus bactrianus</i>	XP_010968130.1
	<i>Balaenoptera acutorostrata scammoni</i>	XP_007173176.1
	<i>Myodes glareolus</i>	XP_048279394.1
<i>Dipodomys spectabilis</i>	XP_042545561.1	

	<i>Delphinapterus leucas</i>	XP_022424428.1
	<i>Eptesicus fuscus</i>	XP_028005288.1
	<i>Bison bison bison</i>	XP_010857610.1
	<i>Pteropus alecto</i>	XP_006920912.1
	<i>Rhinopithecus bieti</i>	XP_017747992.1
	<i>Balaenoptera musculus</i>	XP_036685506.1
	<i>Saimiri boliviensis boliviensis</i>	XP_010331177.1
	<i>Myotis brandtii</i>	XP_005874576.1
	<i>Ursus americanus</i>	XP_045659135.1
	<i>Neogale vison</i>	XP_044096500.1
	<i>Vulpes lagopus</i>	XP_041601933.1
	<i>Equus quagga</i>	XP_046510739.1
	<i>Oryctolagus cuniculus</i>	XP_002718761.1
	<i>Galemys pyrenaicus</i>	KAG8517409.1
	<i>Manis pentadactyla</i>	XP_036754817.1
	<i>Gracilinanus agilis</i>	XP_044521530.1
Aves	<i>Corvus hawaiiensis</i>	XP_048166914.1
	<i>Taeniopygia guttata</i>	XP_030132303.3
	<i>Pyrgilauda ruficollis</i>	XP_041317496.1
	<i>Hirundo rustica</i>	XP_039927077.1
	<i>Onychostruthus taczanowskii</i>	XP_041271849.1
	<i>Passer montanus</i>	XP_039565350.1
	<i>Molothrus ater</i>	XP_036241577.1
	<i>Oxyura jamaicensis</i>	XP_035186324.1
	<i>Turdus rufiventris</i>	KAF4791228.1
	<i>Melopsittacus undulatus</i>	XP_005154576.1
	<i>Strigops habroptila</i>	XP_030344254.1
	<i>Aythya fuligula</i>	XP_032047208.1

<i>Lonchura striata domestica</i>	XP_021381698.1
<i>Geospiza fortis</i>	XP_005416425.1
<i>Camarhynchus parvulus</i>	XP_030807062.1
<i>Serinus canaria</i>	XP_009085424.1
<i>Manacus vitellinus</i>	XP_029814357.1
<i>Zonotrichia albicollis</i>	XP_005481147.1
<i>Apteryx rowi</i>	XP_025925619.1
<i>Numida meleagris</i>	XP_021255526.1
<i>Gallus gallus</i>	XP_040530979.1
<i>Tyto alba</i>	XP_032844696.2
<i>Lagopus leucura</i>	XP_042719653.1
<i>Pygoscelis adeliae</i>	XP_009324216.1
<i>Corvus brachyrhynchos</i>	XP_008632648.1
<i>Calypte anna</i>	XP_030309482.1
<i>Apteryx mantelli mantelli</i>	XP_013797196.1
<i>Apaloderma vittatum</i>	XP_009876249.1
<i>Haliaeetus leucocephalus</i>	XP_010563034.1
<i>Tyto alba</i>	XP_032844693.2
<i>Lonchura striata domestica</i>	OWK56987.1
<i>Cygnus atratus</i>	XP_035403725.1
<i>Cyanistes caeruleus</i>	XP_023785639.1
<i>Amazona aestiva</i>	KQK80332.1
<i>Mesitornis unicolor</i>	XP_010180183.1
<i>Melopsittacus undulatus</i>	XP_005154576.1
<i>Athene cunicularia</i>	XP_026707509.1
<i>Camarhynchus parvulus</i>	XP_030807062.1
<i>Gallus gallus</i>	XP_421764.3
<i>Chaetura pelagica</i>	XP_010006222.1

	<i>Ficedula albicollis</i>	XP_005048826.1
	<i>Serinus canaria</i>	XP_009085424.1
	<i>Cuculus canorus</i>	XP_009568871.1
	<i>Corvus cornix cornix</i>	XP_039410202.1
	<i>Nipponia nippon</i>	XP_009474328.1
	<i>Pelecanus crispus</i>	XP_009483464.1
	<i>Balearica regulorum gibbericeps</i>	XP_010295946.1
	<i>Manacus vitellinus</i>	XP_017924402.1
	<i>Phalacrocorax carbo</i>	XP_009508983.1
	<i>Parus major</i>	XP_015489222.1
	<i>Falco rusticolus</i>	XP_037256573.1
	<i>Corvus hawaiiensis</i>	XP_048166914.1
Reptilia	<i>Pelodiscus sinensis</i>	XP_006135034.1
	<i>Gopherus evgoodei</i>	XP_030424754.1
	<i>Chelonia mydas</i>	XP_037760318.1
	<i>Dermochelys coriacea</i>	XP_038265833.1
	<i>Mauremys reevesii</i>	XP_039402417.1
	<i>Terrapene carolina triunguis</i>	XP_026506928.1
	<i>Trachemys scripta elegans</i>	XP_034632909.1
	<i>Python bivittatus</i>	XP_007441954.1
	<i>Pogona vitticeps</i>	XP_020635566.1
	<i>Zootoca vivipara</i>	XP_034988803.1
	<i>Podarcis muralis</i>	XP_028586060.1
	<i>Sceloporus undulatus</i>	XP_042313006.1
	<i>Anolis carolinensis</i>	XP_008112942.1
	<i>Varanus komodoensis</i>	KAF7253514.1
	<i>Lacerta agilis</i>	XP_033005524.1
	<i>Notechis scutatus</i>	XP_026526250.1

	<i>Notechis scutatus</i>	XP_026526259.1
	<i>Pantherophis guttatus</i>	XP_034296056.1
	<i>Python bivittatus</i>	XP_007441954.1
	<i>Protobothrops mucrosquamatus</i>	XP_015676047.1
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	<i>Anolis carolinensis</i>	XP_003223489.1
	<i>Varanus komodoensis</i>	KAF7253514.1
	<i>Varanus komodoensis</i>	XP_044295172.1
	<i>Lacerta agilis</i>	XP_033005524.1
	<i>Notechis scutatus</i>	XP_026526250.1
	<i>Pantherophis guttatus</i>	XP_034296056.1
	<i>Protobothrops mucrosquamatus</i>	XP_015676047.1
	<i>Pseudonaja textilis</i>	XP_026560589.1
	<i>Sphaerodactylus townsendi</i>	XP_048360798.1
	<i>Crotalus tigris</i>	XP_039208641.1
	<i>Thamnophis elegans</i>	XP_032081709.1
	<i>Chelonoidis abingdonii</i>	XP_032631043.1
	<i>Mauremys mutica</i>	XP_044881213.1
	<i>Chrysemys picta bellii</i>	XP_042711209.1
	<i>Trachemys scripta elegans</i>	XP_034632910.1
	<i>Crocodylus porosus</i>	XP_019411759.1
Amphibia	<i>Xenopus laevis</i>	BAA94748.1
	<i>Lithobates catesbeianus</i>	ACO51875.1
	<i>Bufo gargarizans</i>	XP_044152860.1
	<i>Rana temporaria</i>	XP_040217514.1
	<i>Bufo bufo</i>	XP_040291577.1

	<i>Geotrypetes seraphini</i>	XP_033799427.1
	<i>Microcaecilia unicolor</i>	XP_030058570.1
	<i>Nanorana parkeri</i>	XP_018419981.1
	<i>Cynops orientalis</i>	AFN55259.1
	<i>Xenopus tropicalis</i>	NP_001016299.1
	<i>Xenopus tropicalis</i>	CAJ82745.1
	<i>Rhinatrema bivittatum</i>	XP_029466303.1
Osteichthyes	<i>Paralichthys olivaceus</i>	XP_019965790.1
	<i>Danio rerio</i>	AWP39893.1
	<i>Dicentrarchus labrax</i>	CBN81450.1
	<i>Larimichthys crocea</i>	XP_010740374.1
	<i>Scophthalmus maximus</i>	XP_047183439.1
	<i>Cynoglossus semilaevis</i>	XP_024912944.1
	<i>Oncorhynchus kisutch</i>	XP_020329675.1
	<i>Cyprinus carpio</i>	XP_042591656.1
	<i>Cyprinodon variegatus</i>	XP_015245976.1
	<i>Fundulus heteroclitus</i>	XP_012710613.2
	<i>Cyprinodon tularosa</i>	XP_029690759.1
	<i>Takifugu rubripes</i>	XP_029690759.1
	<i>Salmo salar</i>	XP_014012537.2
	<i>Oncorhynchus mykiss</i>	QWC93456.1
	<i>Ictalurus punctatus</i>	XP_017338665.1
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	<i>Alosa sapidissima</i>	XP_041939757.1
	<i>Anabarrilius grahami</i>	ROI81868.1
	<i>Salvelinus alpinus</i>	XP_023842909.1

<i>Salvelinus alpinus</i>	XP_023861943.1
<i>Gadus morhua</i>	XP_030195895.1
<i>Labrus bergylta</i>	XP_020506858.1
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<i>Collichthys lucidus</i>	TKS89629.1
<i>Notothenia coriiceps</i>	XP_010765636.1
<i>Melanotaenia boesemani</i>	XP_041830322.1
<i>Clarias magur</i>	KAF5896512.1
<i>Anabas testudineus</i>	XP_026207209.1
<i>Chelmon rostratus</i>	XP_041817955.1
<i>Anguilla anguilla</i>	XP_035260198.1
<i>Trematomus bernacchii</i>	XP_033975792.1
<i>Pimphales promelas</i>	KAG1954935.1
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<i>Xiphophorus hellerii</i>	XP_032429481.1
<i>Sebastes umbrosus</i>	XP_037611938.1
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<i>Oryzias melastigma</i>	KAF6724972.1
<i>Megalops cyprinoides</i>	XP_036387401.1
<i>Salarias fasciatus</i>	XP_029954337.1
<i>Salvelinus namaycush</i>	XP_038866282.1

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842 **Table 3. The credits for the pictures used in this study.**

Icon	Figure Origin
Monotremata	Becky-Barnes
Didelphimorphia	Daniel-Stadtmauer
Diprotodontia	Gavin-Prideaux
Afrosoricida	Mo-Hassan
Macroscelidea	uncredited
Pilosa	Xavier-A-Jenkins
Primates	T-Michael-Keeseey
Rodentia	Jiro-Wada
Lagomorpha	Margot-Michaud
Eulipotyphla	Becky-Barnes
Perissodactyla	Mercedes-Yrayzoz-vectorized-by-T-Michael-Keeseey
Artiodactyla	DFoidl-modified-by-T-Michael-Keeseey
Carnivora	Chlo-Schmidt
Pholidota	Steven-Traver
Chiroptera	Margot-Michaud
<i>Homo sapiens</i>	NASA
Avian	Ferran Sayol
Reptilia	Gabriela Palomo-Munoz
Amphibia	Yusan Yang
Actinopterygii	Milton Tan

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844 **Table 4. The constructs of the truncate and chimeric proteins.**

Truncate/chimeric protein	Amino acid sequence
TrGSDME-NT	1-258 aa of TrGSDME
TrGSDME-CT	259-474 aa of TrGSDME
HsNT-TrCT	1-270 aa of HsGSDME + 259-274 aa of TrGSMDE
TrNT-HsCT	1-258 aa of TrGSDME + 271-496 aa of HsGSDME
Hsp20-Trp10	1-198 aa of HsCASP7 + 207-313 aa of TrCASP7
Trp20-Hsp10	1-206 aa of TrCASP7 + 199-303 aa of HsCASP7

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848 **Table 5. Primers used in this study.**

Primer	Sequence (5'-3')
Clone primers	
TrCASP3 forward	ATGTCGGCCAACGGACC
TrCASP3 reverse	GGAAAATAACATCTCTTTGGTCAGC
TrCASP7 forward	ATGCAGATGGCTGGAGAACC
TrCASP7 reverse	GTTAAAGTACAGTTCTTTTGTCAGCATCG
TrGSDME forward	ATGTTTTCCAAGGCCACGG
TrGSDME reverse	ATCGATAAAATCCGTTTCAGACTTTG
Recombination primers	
TrCASP3 forward	TAAGAAGGAGATATACATATGATGT CGGCCAACGGACC
TrCASP3 reverse	GTGGTGGTGGTGGTGGTCTCGAGGGA AAAATAACATCTCTTTGGTCAGC
TrCASP7 forward	TAAGAAGGAGATATACATATGATGCA GATGGCTGGAGAACC
TrCASP7 reverse	GTGGTGGTGGTGGTGGTCTCGAGGTTAA AGTACAGTTCTTTTGTCAGCATCG

MmGSDME-N234S forward	GTTATTACTCATGGAGGAGCCCAGG GAAAG
MmGSDME-N234S reverse	CTTTCCTGGGCTCCTCCATGAGTA ATAAC
HsGSDME-Δ261-266 forward	CCAGGATGGACATCCATTTGCGGAG CTGC
HsGSDME-Δ261-266 reverse	CAAATGGATGTCCATCCTGGGAAGAT ATCCCAT
HsGSDME-Δ281-296 forward	CCTGGTCTTT GACATGCCAGATGCTG CGCA
HsGSDME-Δ281-296 reverse	CTGGCATGTCAAAGACCAGGGGGTCC AGGTAGA
Overexpression primers	
TrGSDME-FL forward	TCAGATCTCGAGCTCAAGCTTATGTTT TCCAAGGCCACGG
TrGSDME-FL reverse	CATGGTGGCGACCGGTGGATCATCGA TAAAATCCGTTTCAGACTTTG
TrGSDME-NT forward	TCAGATCTCGAGCTCAAGCTTATGTTT TCCAAGGCCACGG
TrGSDME-NT reverse	CATGGTGGCGACCGGTGGATCGTCTAC AGCATCAGGAGACTCC
TrGSDME-CT forward	TCAGATCTCGAGCTCAAGCTTATGGGCC GGTGCCTGGA
TrGSDME-CT reverse	CATGGTGGCGACCGGTGGATCATCGA TAAAATCCGTTTCAGACTTTG
TrCASP7 forward	TTCTGAAGAGGACTTGAATTCAATGG CTGGAGAACCCACTGAG
TrCASP7 reverse	ACGACTCACTATAGTTCTAGATCAGT

	TAAAGTACAGTTCTTTTGTGTCAGC
TrCASP3 forward	TTCTGAAGAGGACTTGAATTCAAATG TCGGCCAACGGACC
TrCASP3 reverse	ACGACTCACTATAGTTCTAGAGGAA AAATACATCTCTTTGGTCAGC
HsGSDME forward	CAAGCTTGCGGCCGCGAATTCAATGT TTGCCAAAGCAACCAGG
HsGSDME reverse	ACGACTCACTATAGTTCTAGATCATG AATGTTCTCTGCCTAAAGC
HsCASP7 forward	TTCTGAAGAGGACTTGAATTCAATGG CAGATGATCAGGGCTG
HsCASP7 reverse	ACGACTCACTATAGTTCTAGACTATTG ACTGAAGTAGAGTTCCTTGGTG
HsCASP3 forward	TTCTGAAGAGGACTTGAATTCAATGG AGAACACTGAAAACCTCAGTGG
HsCASP3 reverse	ACGACTCACTATAGTTCTAGATTAGTG ATAAAAATAGAGTTCTTTTGTGAGC
MmCASP7 forward	TTCTGAAGAGGACTTGAATTCAATGA CCGATGATCAGGACTGTGC
MmCASP7 reverse	ACGACTCACTATAGTTCTAGATCAAC GGCTGAAGTACAGCTCTTTGG

Figure 1

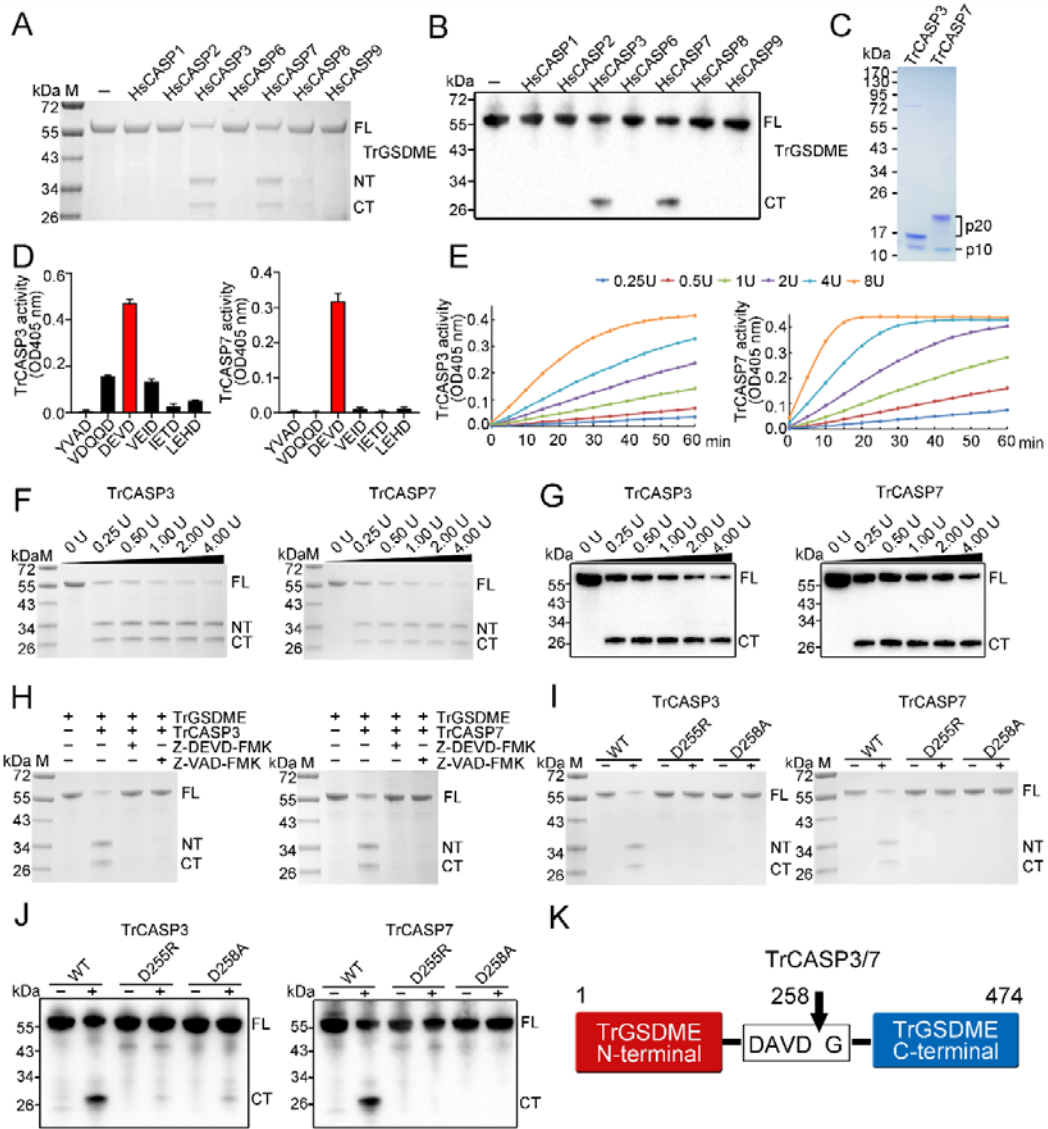


Figure 2

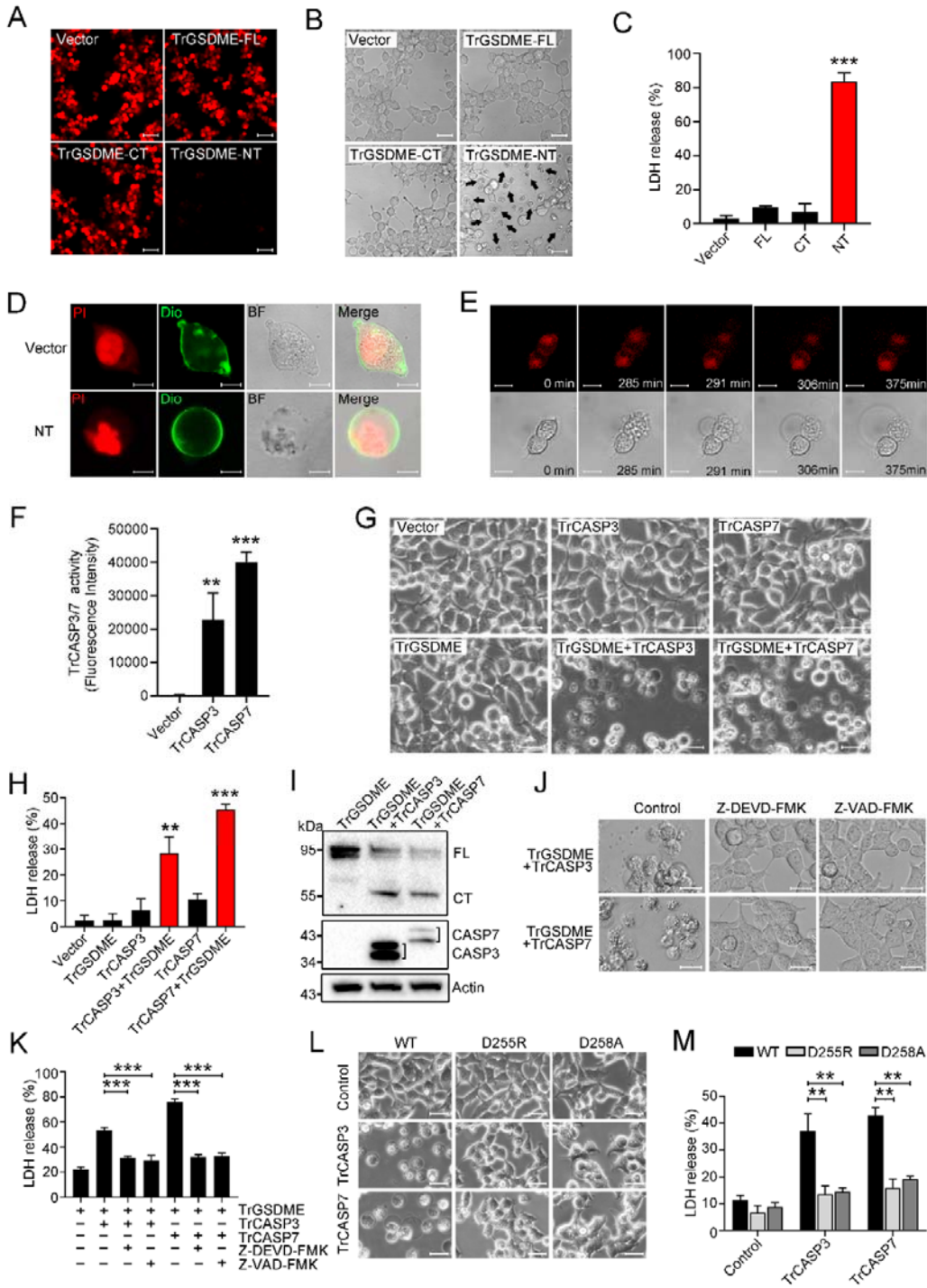


Figure 3

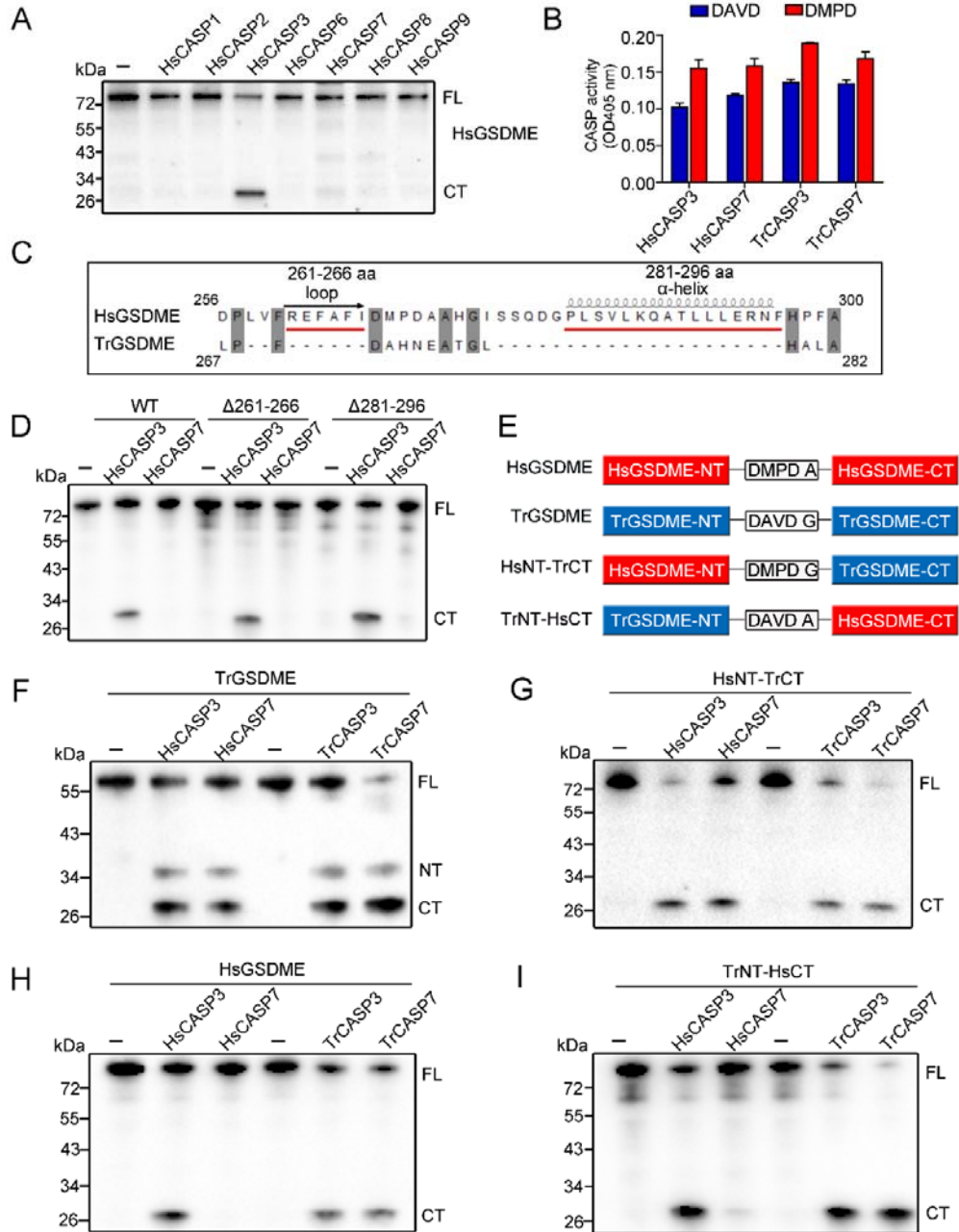


Figure 4

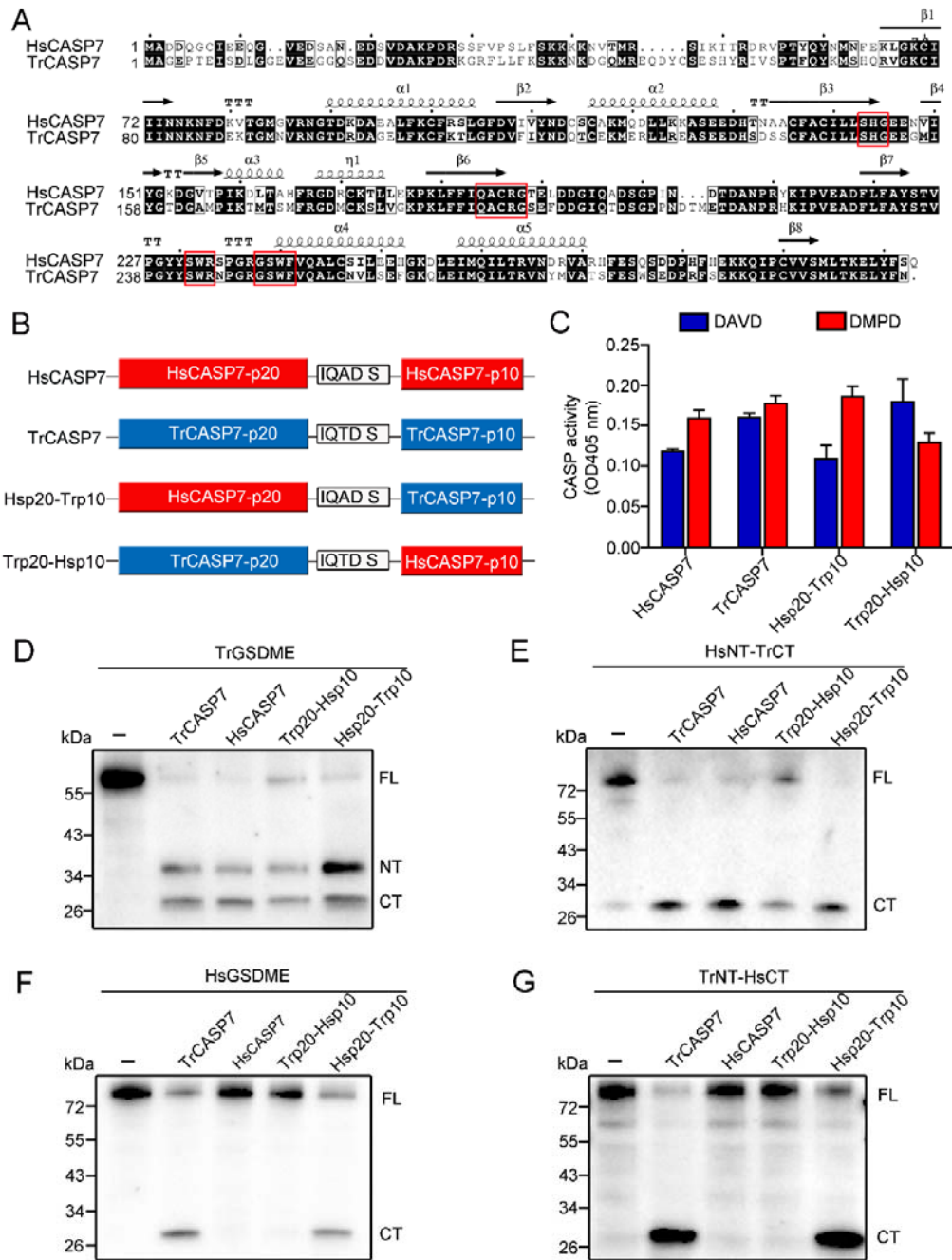


Figure 5

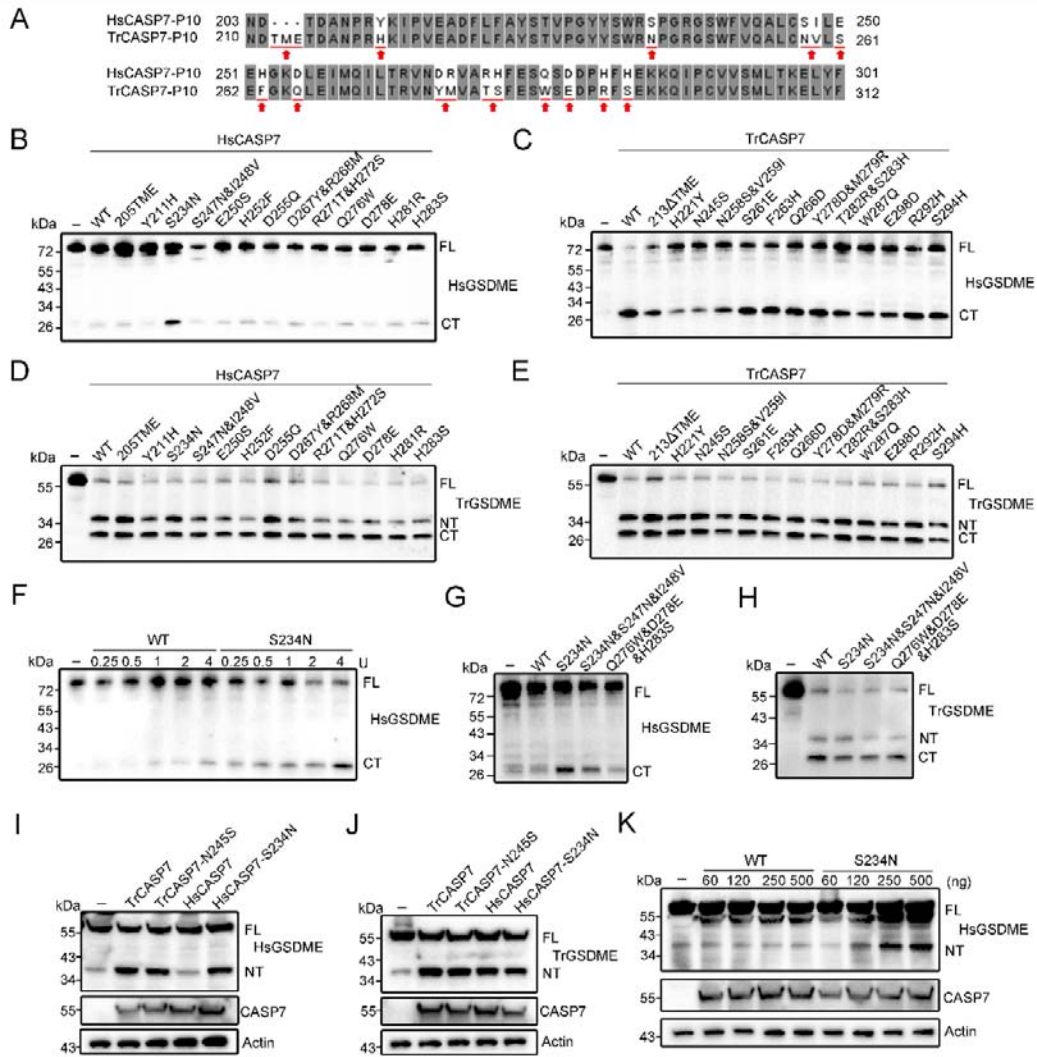


Figure 6

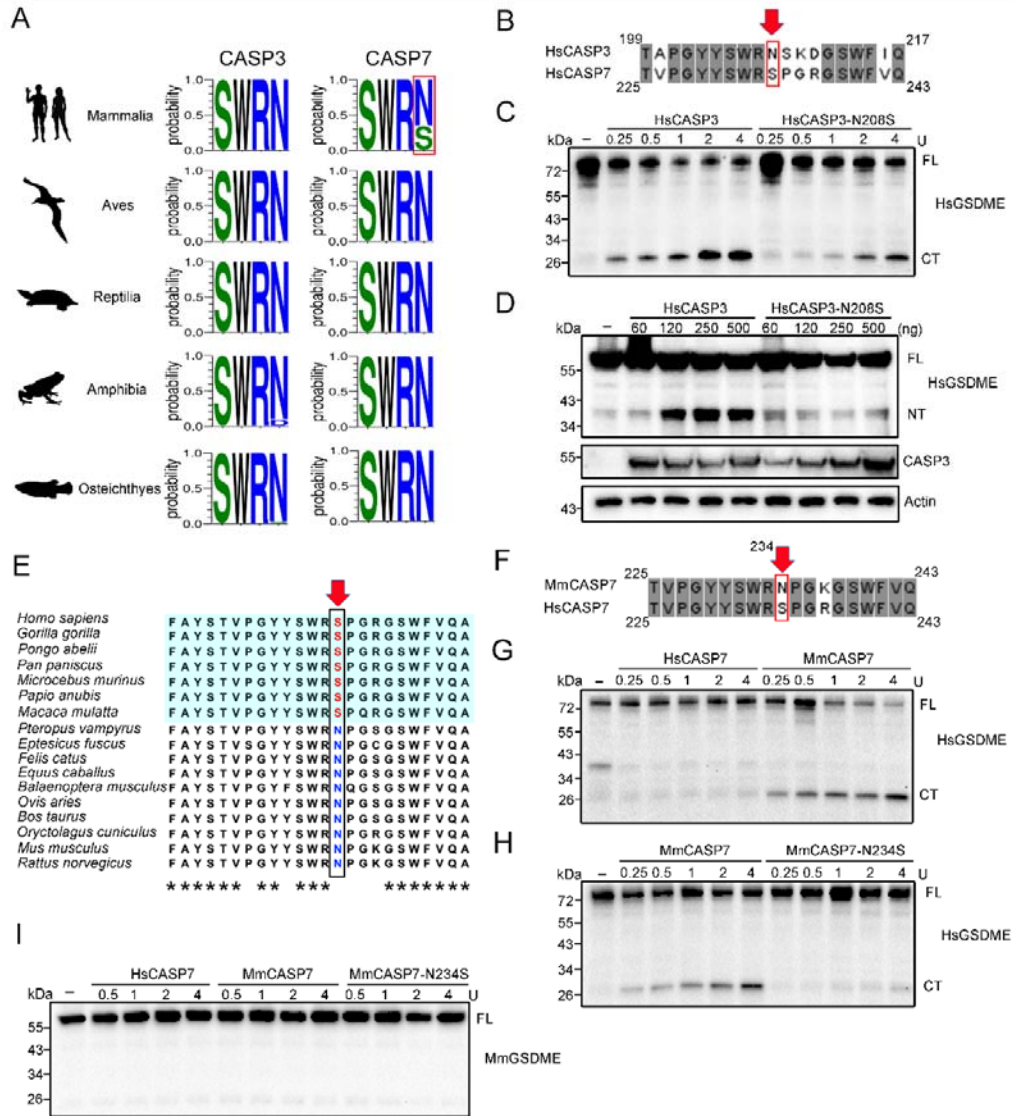


Figure 2-figure supplement 2

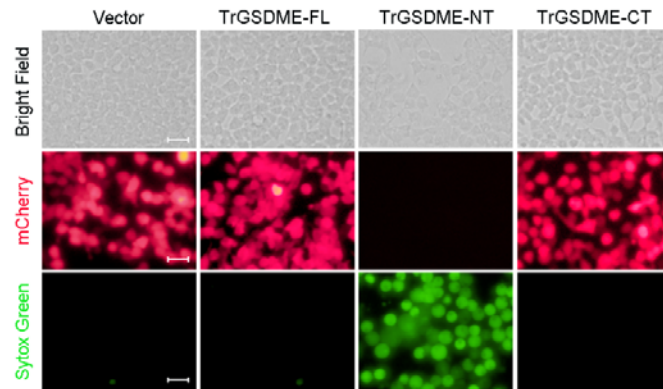


Figure 2-figure supplement 3

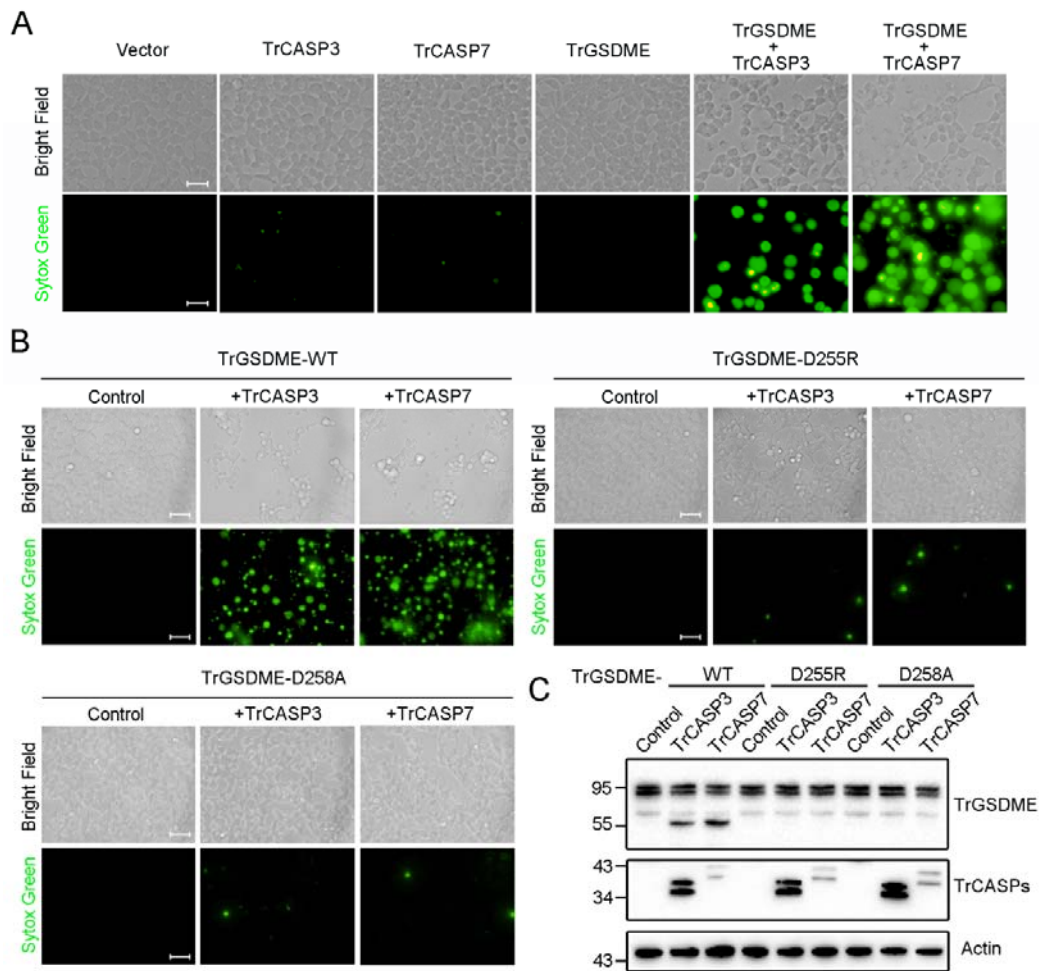


Figure 3-figure supplement 1

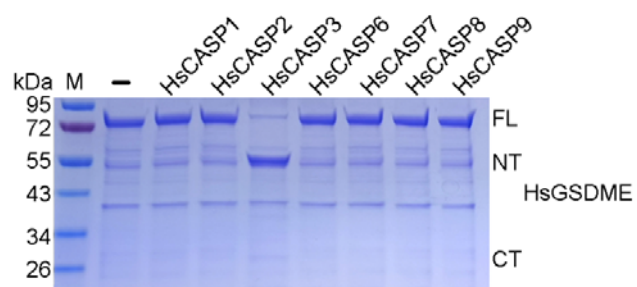


Figure 3-figure supplement 2

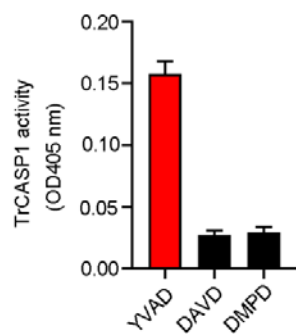


Figure 4-figure supplement 1

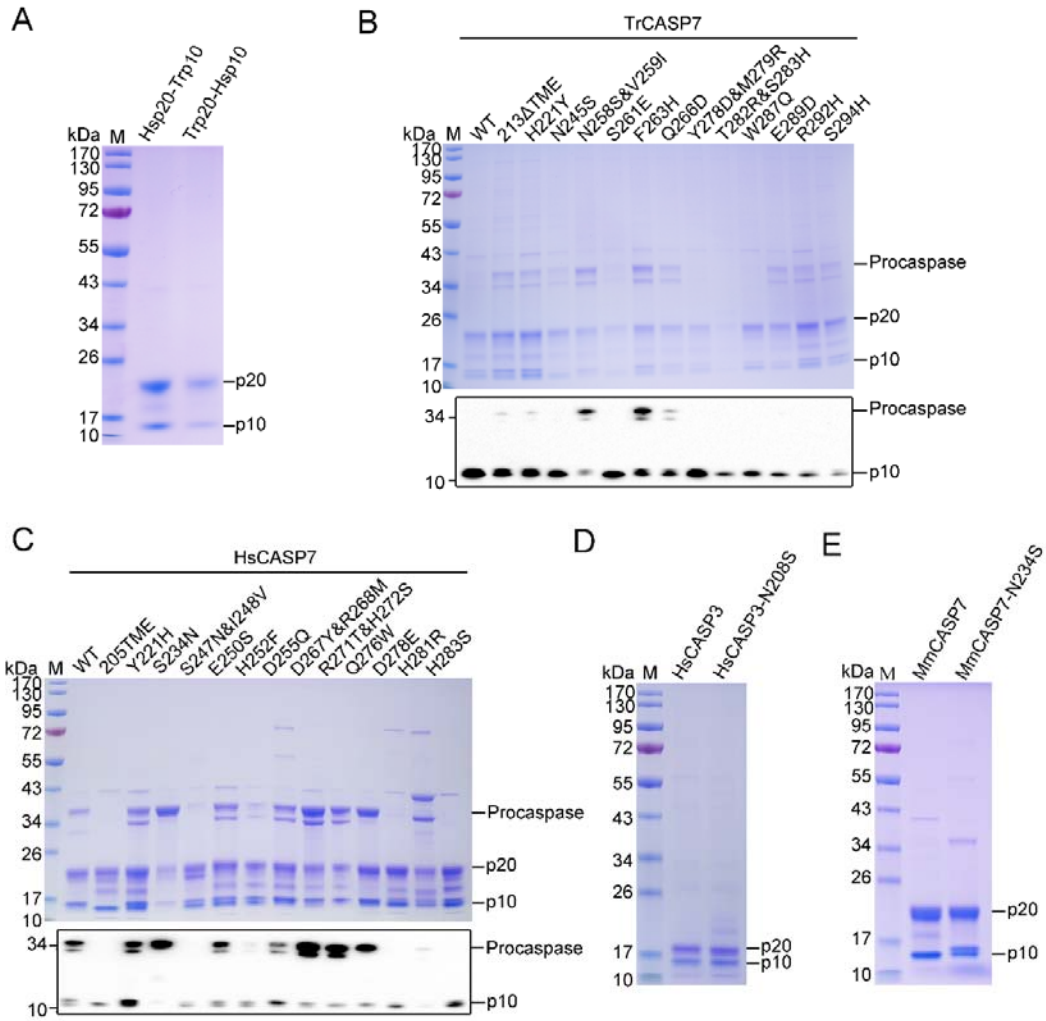


Figure 5-figure supplement 1

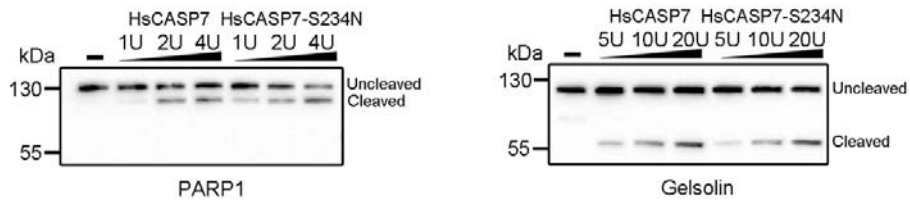


Figure 6-figure supplement 1

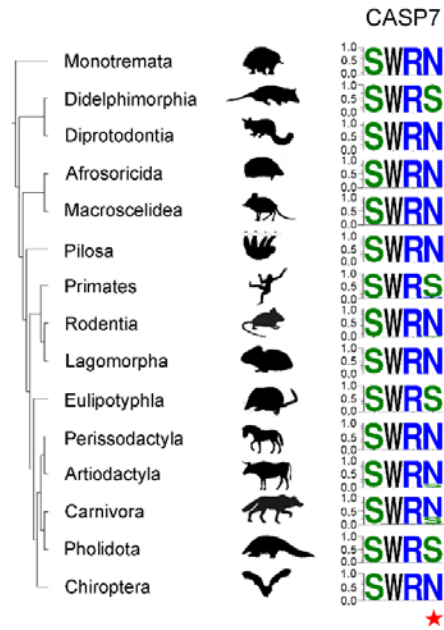


Figure 6-figure supplement 2

