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Published in final edited form as:

Title: Caspase-3 and RasGAP: a stress-sensing survival/demise switch. Authors: Khalil H, Bertrand MJ, Vandenabeele P, Widmann C Journal: Trends in cell biology Year: 2014 Feb Volume: 24 Issue: 2 Pages: 83-9 **DOI:** 10.1016/j.tcb.2013.08.002

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Faculté de biologie et de médecine

1	Caspase-3 and RasGAP: a stress-sensing survival/demise switch			
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13	Keywords: Caspase-3, RasGAP, Cellular stress, Sensors			
14				

16 Abstract

17 The final decision on cell fate, survival versus cell death, relies on complex and tightly regulated checkpoint mechanisms. The caspase-3 protease is a 18 19 predominant player in the induction of apoptosis. However, this protease 20 is more than a blind killer. It can gauge the degree of cellular stress and 21 damage by differential processing of p120 RasGAP to generate the 22 appropriate cellular response (survival or cell demise). Partial cleavage of 23 RasGAP by caspase-3 initiates the anti-apoptotic Akt pathway. In contrast, 24 full cleavage of RasGAP abrogates this survival response, hence favoring cell death. Therefore, rather than relying on separate sensors, cells can 25 26 modulate a given set of proteins to generate, depending on the intensity of 27 the input signals, opposite outcomes (survival vs. death).

29 Introduction

30 Proper functioning of cells requires that they maintain their biochemical parameters within rather narrow limits. This process, called homeostasis, is vital 31 32 for cells and they spend a considerable energy to maintain it. For example, the 33 appropriate gradient of sodium and potassium ion concentrations between the 34 inside and the outside of brain cells via the action of the sodium/potassium 35 ATPase pump consumes close to half of the ATP used at the basal state [1]. If the 36 extracellular conditions are changing too much or if the cellular biochemistry is 37 exceedingly altered, such as following viral infection [2,3] or oncogene 38 expression [4], than homeostasis maintenance is compromised. This 39 perturbation is sensed by cells as cellular stress. Situations and conditions that 40 can generate cellular stress are extremely diverse. They include environmental 41 pollutants, metabolic stress induced by toxic lipids in obese individuals, UV-42 mediated stress generated by prolonged exposure of the epidermis to sun 43 exposure, endoplasmic reticulum stress caused by sustained chronic increase in 44 insulin production by pancreatic beta cells in type 2 diabetes patients, chemical 45 or ionizing stress that cancer patients experience when treated with chemo- or 46 radiotherapy, or physicochemical stress provoked by heat or cold exposure.

When encountering a stress condition, cells have two options, either to cope with it, which requires an adaptive response, or to give up and die, which involves the initiation of an active and programmed cell death process. Cells are in fact well equipped to sense the extent of stress and this allows them to decide whether they should try to restore homeostasis or whether they should commit suicide because stress is evaluated as too extensive and potentially deleterious. Various switch mechanisms can be used by cells to evaluate stress and insults to either

promote cell survival or apoptosis. We will focus here on a newly described
physiologic stress-sensing cellular system based on the differential cleavage of a
specific caspase-3 substrate.

57

58 **Caspase-3, from reaper to friendly protease**

59 At the end of the nineties, a consensus emerged that apoptosis (Box1) was 60 carried on by the executioner members of the caspase family (Box1) (caspase-3, 61 -6, and -7) [5-8]. Caspase-3 has been considered as the central executioner 62 protease due to its high catalytic turnover and greediness to cleave substrates 63 [9]. Gene targeting in mice has revealed a certain level of redundancy in the 64 function of executioner caspases in apoptosis induction [10], and more 65 importantly that these executioner caspases fulfill other non-apoptotic functions 66 as well [11].

67 Historically however, non-apoptotic functions of pro-apoptotic caspases were first discovered for caspase-8, an initiator caspase. It was shown in 1998 that 68 69 mice lacking caspase-8 were embryonic lethal [12]. This lethality was later 70 shown to result from degeneration of the vasculature of the yolk sac [13]. This 71 suggested that caspase-8 is required for the proper development and/or the 72 maintenance of vascular structures in the early embryo, which is most certainly 73 distinct from its ability to mediate apoptosis via the extrinsic cell death pathway. 74 Also in 1998, it was shown that cell expressing CrmA, a caspase-8 inhibitor, were 75 paradoxically sensitized to necrotic cell death [14], suggesting that caspase-8 76 could also promote cell survival [15]. Non-apoptotic functions were then 77 progressively reported for other caspases and a series of reviews have been 78 describing these in the past decade [11,16-21]. Caspase-3 in particular appears

to regulate the differentiation of skeletal muscle cells, osteoblasts, lens epithelial cells, neural stem cells, and several hematopoietic lineages (T and B lymphocytes, erythrocytes, macrophages, platelets). Caspase-3 has also been reported to participate in stem cell maintenance, inhibition of B-cell proliferation, dendritic cell maturation, and proliferation of brain cells and keratinocytes [22-25].

85 The levels of activated caspase-3 that stimulate the non-apoptotic biological 86 responses should obviously be lower than during apoptosis otherwise this cell 87 death process would not be avoided. Numerous substrates of caspase-3 have 88 been determined [9,26-28]. However their role in apoptosis has only been 89 explored in a few cases. Additionally, the impact of caspase-mediated cleavage 90 for the vast majority of substrates has not been explored by studying the 91 phenotype of cells or mice in which the wild-type substrate is replaced with a 92 non-cleavable knock-in form.

93 Different executioner caspases can target the same substrates while a given 94 caspase can cleave some substrates better than others [9,29]. In particular, 95 caspase-3 appears to cleave most substrates more efficiently than other caspases 96 [29]. It is conceivable that the high K_m value of caspase-3 for some of its 97 substrates is related to its non-apoptotic functions while apoptosis may imply 98 the less effective substrates. In other words, low activation levels of caspase-3, 99 leading to restricted cleavage of high affinity substrates, would allow survival 100 while higher levels of caspase-3 activation would result in the cleavage of a much 101 wider spectrum of substrates, including low affinity ones, ultimately leading to 102 the induction of apoptosis.

103 In addition to the possible differential affinity of caspase-3 toward the anti- and 104 pro apoptotic substrates, there are other mechanisms that can keep caspase-3 105 activation levels in check during stressful situations. One possibility is that cells 106 have a built-in system that prevents amplification of caspase-3 activation. The 107 best-known family of proteins that can negatively control executioner caspases 108 is the inhibitor of apoptosis (IAPs) protein family. IAPs are characterized by one 109 to three <u>baculovirus</u> <u>IAP</u> repeat (BIR) zinc-binding domain, which, at least for 110 some IAP family members, is required for their ability to bind caspases [30]. 111 However, contrary to initially speculated, X-linked IAP (XIAP) is the only IAP that can function as a direct inhibitor of caspase-3, -7 and -9 [31,32]. Cellular IAP1 112 113 and 2 (cIAP1/2) instead promote cell survival by functioning as E3 ubiquitin 114 ligases that promote the degradation of caspase-3 and -7 [33] and positively 115 regulate activation of the canonical NF-kB pathway [34-36]. In addition, cIAP1/2 116 function as regulators of survival signaling in cancer cells by preventing RIPK1 117 from becoming a pro-death stress sensing molecule [37]. Hence, IAP family 118 members allow cells to cope with stress to a certain limit if they are expressed at 119 sufficiently high levels either by activating NF-kB, preventing RIPK1 from 120 becoming a pro-death molecule or by keeping caspase-3 activity in check.

Another possibility that can explain why activation of caspase-3 does not lead to apoptosis in all cases is that caspase-3 itself, when activated at low levels, turns on an anti-apoptotic response that acts as a negative feedback loop preventing further caspase-3 activation (i.e. an amplification of caspase-3 activity). A recent study [38] provides genetic evidence for this second possibility. In the C57BL/6 background, mice lacking caspase-3 are viable and fertile [39]. The vast majority of development programs occur normally in these mice. They do however 128 display abnormal inner ear development [40]. It is not known whether this 129 defect results from a defect in apoptosis or from alterations of other functions 130 controlled by caspase-3. Apoptosis in these mice can proceed normally or be 131 inhibited, depending on which tissues and organs are investigated. For example, 132 female germ cell apoptosis is not affected by the lack of caspase-3 while 133 granulosa cell apoptosis is not taking place during follicular atresia when this 134 caspase is missing [41]. Caspase-3 knock-out mice are unable or strongly 135 impaired in activating the anti-apoptotic Akt kinase (box 2) in response to a 136 variety of stresses such as UV-B exposure of the skin, doxorubicin-mediated 137 cardiac damage, and experimentally-induced colitis, an inflammation of the large 138 intestine. Akt activation is also compromised in wild-type mice injected with the 139 O-VD-OPh caspase inhibitor suggesting that the activity of caspase-3, and not a 140 potential adaptor function of the protein, is required for its capacity to activate Akt. As detailed below, the defect in Akt activation in mice lacking caspase-3 is 141 142 accompanied by increased cell death responses. Caspase-3 therefore appears to 143 stimulate survival responses in some situations.

144 As caspase-3 is a main executioner caspase, its absence generally compromises 145 or abrogates apoptosis induction by various stress inducing conditions. For 146 example, pancreatic beta cells lacking caspase-3 are fully resistant against 147 streptozotocin-induced death [42]. Even though absence of caspase-3 generally 148 reduces the apoptotic response, this does not mean necessarily that the death 149 response is lowered. For example, UV-B-induced apoptosis is decreased in the 150 epidermis of caspase-3 KO mice but the overall death response is not, as in the 151 absence of apoptosis induction cell death proceeds in caspase-independent cell 152 death with necrotic morphology [38].

153 In some cases, caspase-7 can compensate, at least partially by enhanced 154 expression levels in C57Bl6 strains for the lack of caspase-3 and allow apoptosis 155 to proceed [10]. Doxorubicin is an efficient anti-cancer drug that is currently 156 used against various tumors. However, this drug also induces serious cardiac 157 side effects. Cardiomvocytes are indeed rather sensitive to doxorubicin and 158 readily undergo apoptosis when exposed to the drug. Unexpectedly, caspase-3 159 knock-out mice experience even more cardiomyocyte apoptosis than wild-type 160 mice when treated with doxorubicin, leading to increased mortality [38]. This 161 suggests that the apoptotic role of caspase-3 is redundant while its prosurvival 162 role in stressed tissues and organs is not.

163

164 **Cellular sensor of caspase-3 activity**

165 There is only a handful proteins including p120 RasGAP[43], p27^{kip1} [44], Lyn [45], Synphilin-1 [46], nucleophosmin [47], Rb [48], and the Drosophila IMD 166 167 protein [49] that may activate survival pathways once cleaved by caspases. The 168 first of these, RasGAP, is of particular interest in the present context because the 169 N-terminal moiety generated by cleavage of RasGAP at position 455 by caspase-3 170 activates Akt and protects cells *in vitro* [50,51]. In cells, RasGAP does not seem to 171 be cleaved by caspases other than caspase-3 [51]. RasGAP bears another 172 caspase-3 cleavage site at position 157 that is made accessible only following the 173 first cleavage event or that is much less efficiently recognized by caspases [43]. 174 Consequently, the second site is not used when caspase-3 is mildly activated [52] 175 (Figure 1, upper left panel). However, after an apoptotic insult, concentrations of 176 activated caspase-3 further increases, reaching a threshold allowing the N-177 terminal fragment of RasGAP to be cleaved at position 157, abrogating Akt

178 activation. The second RasGAP cleavage thus favors death [52] (Figure 1, upper 179 right panel). RasGAP is the only caspase-3 substrate that shows this differential 180 cleavage-mediated control of two opposite outcomes (survival vs apoptosis). 181 Interestingly, RasGAP has no obvious anti-apoptotic activities as a full-length 182 protein [53]. It only acquires cell protective functions when cleaved at position 183 455. The physiological role of caspase-3-mediated cleavage of RasGAP has 184 recently been assessed using a knock-in mouse strain homozygous for a 185 mutation in RasGAP at position 455 that prevents its cleavage by caspase-3. The 186 RasGAP knock-in mice phenocopied the caspase-3 knock-out stress-induced Akt 187 activation defect in response to various patho-physiological stresses (Figure 1, 188 lower panels). The knock-in mice also experienced increased tissue damage and 189 organ dysfunction in response to these insults in comparison to wild-type mice 190 [38]. These results provide genetic evidence for the importance of caspase-3-191 mediated p120 RasGAP cleavage as a defense mechanism to protect organisms 192 against damage induced by diverse pathological conditions.

193 It can therefore be proposed that caspase-3 itself activates the anti-apoptotic Akt 194 kinase following certain types of stresses (e.g. starvation, DNA damage)[51] and 195 this seemingly operates via the efficient cleavage of only one of its substrates, the 196 ubiquitous p120 RasGAP protein. The model drawn for these studies is that 197 RasGAP bears two cleavage sites with differential sensitivities for caspase-3 198 proteolytic activity. RasGAP acts as a "sensor like" protein that reflects the levels 199 of caspase-3 activation, which correlates with the extent of cellular stress. This 200 differential cleavage aids determining the fate of stressed cells: activation of a 201 protective Akt-dependent pathway following the first cleavage of RasGAP or

inactivation of this protective pathway following the second cleavage of RasGAP(Figure 1, upper panels).

204

205 **Perspectives: one set of signaling proteins, different outcomes**

206 There are other comparable decision-making systems assessing the magnitude 207 of cellular stress, where within a given signaling pathway, differential regulatory 208 events dictate distinct cell signaling events. These "sensor-like" responses 209 include the unfolded protein response (UPR) in the endoplasmic reticulum and 210 the activation of p53 transcription factor pathway by DNA damage. What 211 determines the cellular outcome once a given switch is triggered relies on the 212 duration and the extent of the stress that allows or not the accumulation of pro-213 apoptotic genes, either as a consequence of differential stability of the anti- and 214 pro-apoptotic proteins or as a result of post-translational modifications of the 215 switch components (Figure 2). For instance, the goal of the UPR initially is to try 216 restoring the folding capacity of the ER while diminishing protein load in order 217 to reestablish ER homeostasis [54-56]. Only a strong and sustained ER stress 218 allows for the accumulation of pro-apoptotic labile proteins such as the 219 transcription factor CHOP [56,57]. Similarly, distinct post-translational 220 modifications (e.g. acetylation on lysine 150 or phosphorylation of serine 46) 221 generated by various degrees of DNA damage, determines whether p53 222 preferentially stimulates a repair pathway versus a cell death response [58-60]. 223 From these examples one could propose a general paradigm were cellular stress 224 is sensed by molecules and pathways that stimulate either cell survival or 225 apoptosis based on the extent of their stimulation (as in the case of the UPR) or 226 as a result of stress-mediated differential post-translational modifications (e.g.

cleavage of RasGAP, acetylation/phosphorylation of p53). Hence, cells may use
the same set of sensor proteins to fine-tune an appropriate cellular response
either resulting in cell survival or cellular demise, rather than relying on
separate sensors for either response.

231 Figure legends

232

Figure 1. RasGAP cleavage, a sensor of caspase-3-activity controlling the survival and the death of cells.

235 RasGAP bears two cleavage sites with different sensitivities towards caspase-3 236 activity. Site 1 at position 455 is used at low levels of caspase-3 activity, while 237 site 2, at position 157, is only recognized at high levels of caspase-3 activity. 238 **Upper left panel**. In situations of mild stress, caspase-3 is activated to low levels 239 and this leads to the partial cleavage of RasGAP into fragment C and fragment N. 240 The latter activates an Akt-mediated anti-apoptotic response that inhibits 241 further amplification of caspase-3 activity. **Upper right panel**. When cells are 242 facing higher stress, caspase-3 is more strongly activated resulting in further 243 cleavage of fragment N and abrogation of Akt activation. This favors cell death.

Lower panels. When cleavage of RasGAP cannot occur as a result of a mutation at its first caspase-3 cleavage site, the prosurvival kinase Akt is not stimulated. In this condition, the feedback loop that prevents an initially mild caspase-3 activity is not activated and this results in an amplification of caspase-3 activity. Consequently, mild stresses generate elevated caspase-3 activation comparable to when cells experience stronger stresses. This eventually leads to cell death.

250

Figure 2. Examples of mechanisms allowing single stress sensors to activate survival or death responses

A given protein or set of proteins can determine the fate of a cell (survival vs death). This can result as a consequence of a differential cleavage of a protease substrate (e.g. cleavage of RasGAP by caspase-3) (**upper panel**), the

- accumulation above a certain threshold of a labile transcription factor (e.g CHOP
 translation following ER stress) (middle panel), or differential posttranslational modifications (e.g. as occurring on p53 in response to varying
 degrees of DNA damage) (lower panel).

261 **Box 1. Apoptosis and caspases**

262 Apoptosis (apo=for, ptosis=falling) is a type of cell death that was formally 263 described by Kerr, Wyllie and Currie in the early 1970s [61]. Apoptosis 264 participates in the control of tissue size and shape during development. It is 265 involved in the elimination of cells that may represent a threat to the organism 266 such as pre-malignant cells and activated immune T cells. Apoptosis is 267 characterized by a series of organized and finely regulated biochemical and 268 cellular modifications that process the dying cells for efficient elimination by 269 phagocytosis by neighboring cells or macrophages. This insures that no 270 cytoplasmic compounds leak from the cells that would otherwise generate an 271 inflammatory response [62]. The hallmarks of apoptosis include cell shrinkage, 272 membrane blebbing, nuclear and cytoplasmic condensation, DNA fragmentation, 273 cytoskeleton proteolysis and breakup of organelles. Cellular components are 274 sorted into a number of vesicles known as apoptotic bodies. Only a few of these events are diagnostic for apoptosis (DNA cleavage, formation of apoptotic 275 276 bodies). Intact apoptotic cells display "eat-me" flags on their surface such as 277 phosphatidylserine that tag them as targets for phagocytosis. Phosphatidylserine 278 is usually confined to the inner side of the plasma membrane but becomes 279 exposed on the surface of apoptotic cells [63,64].

The cellular and biochemical features of apoptosis are triggered by members of the caspase family of proteases. Caspases are a family of cysteine aspartic acid proteases present in healthy cells as inactive precursor enzymes (zymogens). All caspases have a similar domain structure (pro-polypeptide, large and small subunits) [65]. Domains within the pro-polypeptide, such as caspase recruitment domain (CARD) and death effector domain (DED), by homophylic interactions,

286 can recruit caspases to activation platforms. Not all caspases are implicated in 287 apoptosis but those that are can be divided into initiator (caspase-2, -8, and -9) 288 or executor caspases (caspase-3, -6, and -7). The former auto-activate when 289 brought together by receptors belonging to the death receptor family (e.g. Fas). 290 The latter are activated by the former by proteolysis and execute the proteolysis 291 events that are seen during the demolition phase of apoptosis [62,66]. 292 Executioner caspases cleave hundreds of substrates during the process of 293 apoptosis but only for a minority of these is the physiological function of their 294 cleavage understood. These include ICAD, the inhibitor of CAD, the DNAse that 295 cleaves the DNA between nucleosomes. Caspase-mediated cleavage of ICAD leads 296 to CAD activation, generating ~180 base pair multimeric DNA fragments, the 297 diagnostic laddering in apoptotic cells.

298

299 Box 2. The generally pro-survival Akt pathway

300 Akt is a serine/threonine kinase involved in the regulation of cell survival, 301 proliferation, and metabolism and is activated by phosphorylation. Three 302 isoforms exist (1, 2, and 3) that all contain an N-terminal pleckstrin homology 303 (PH) domain, a central kinase domain containing a phosphorylation site within 304 the activation-loop (threonine 308 in human Akt1), and a conserved regulatory 305 serine phosphorylation site in a hydrophobic motif near the C terminus (position 306 473 in human Akt1). The interaction of the Akt PH domain with 3'-307 phosphoinositides causes Akt translocation to the plasma membrane, inducing 308 conformational changes that allow Akt to expose its phosphorylation sites [67]. 309 Phosphoinositide-dependent kinase-1 (PDK-1) is also recruited to the plasma 310 membrane after PIP3 generation. PDK-1 is the kinase that phosphorylates Akt on

threeonine 308, which stabilizes the activation loop in an active conformation.
Serine 473 is then phosphorylated by the PDK2 kinase activity which is
predominantly carried out by mammalian target of rapamycin (mTOR) complex
2 (mTORC2) or DNA-PK. Phosphorylation of threeonine 308 is a prerequisite for
kinase activation. Phosphorylation of serine 473 appears to further increase Akt
kinase activity. Recent evidence suggests that this phosphorylation event also
controls the target specificity of the kinase [67].

318 Among the Akt substrates that have been identified in mammalian cells, many 319 are regulators of apoptosis or cell growth [67]. The Akt substrates are 320 phosphorylated within the same basic motif RXRXXS/T. The anti-apoptotic 321 response induced by Akt involves NF-kB- or CREB-mediated up-regulation of 322 anti-apoptotic proteins such as c-IAP1/2, Mcl-1 and Bcl-2, or the direct 323 phosphorylation and inhibition of pro-apoptotic proteins such as Bad and 324 caspase-9. It is worth noting however that the Akt phosphorylation site in caspase-9 is not conserved in mammals [68]. Akt also inhibits members of the 325 326 Forkhead family of transcription factors (FOXO transcription factors). This 327 prevents the expression of pro-apoptotic genes such as Fas ligand. Additionally, 328 murine double minute 2 (MDM2), an E3 ubiquitin ligase targeting p53 for 329 degradation, is stabilized by Akt-mediated phosphorylation. This will therefore 330 diminish p53-induced pro-apoptotic signaling in cells. Therefore, Akt promotes 331 survival in most cells either through direct phosphorylation of targets, or 332 through the induction of anti-apoptotic genes [67]. There are some cell types 333 however in which Akt activation can lead to death. In pancreatic beta cells for 334 example, Akt activation leads to NF-κB-dependent apoptosis [69].

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Cells or animals expressing wild-type RasGAP

Cells or animals expressing a caspase-3 cleavage-resistant RasGAP mutant



Switches	Survival	Death
Differential cleavage of a protease substrate	anti-apoptotic signal	abrogation of the anti-apoptotic signal
Levels of unstable pro-apoptotic proteins	degraded under the threshold of apoptosis induction	
Differential modulation of the specificity of a transcription factor	post-translational modification X	post-translational modification Y pro-apoptotic genes

Figure 2