Hyperactivation of the Mammalian Degenerin MDEG Promotes Caspase-8 Activation and Apoptosis**

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Intracellular calcium overload plays a critical role in numerous pathological syndromes such as heart failure, brain ischemia, and stroke. Hyperactivation of the acid-sensing ion channels including degenerin/epithelial amiloride-sensitive sodium (DEG/ENaC) channels has been shown to elevate intracellular calcium and cause subsequent neuronal cell death that is independent of the canonical Egl-1/Ced-9/Ced-4/Ced-3 apoptotic pathway in Caenorhabditis elegans. In mammalian cells, hyperactivation of the DEG/ENaC channels can also lead to cell death, although the underlying mechanism remains largely unknown. Here, we use a tetracycline-inducible system to express the hyperactivation mutant of a mammalian DEG/ENaC channel protein, MDEG G430F, in murine kidney epithelial cells deficient in the key mitochondrial apoptotic proteins Bax and Bak. Remarkably, expression of MDEG G430F induces increased intracellular calcium, reactive oxygen species (ROS) production, and cell death. The MDEG G430F-induced cell death is blocked by the intracellular calcium chelator 1,2-bis(o-aminophenoxy)ethane-N,N,N′,N′′-tetraacetic acid (acetoxymethyl ester), ROS scavengers, and the caspase inhibitor z-VAD-fmk (where z and fmk are benzyloxycarbonyl and fluoromethyl ketone). Mechanistically, the intracellular calcium overload and ROS increase lead to the inhibition of proteasomal and autophagic protein degradation, which promotes the accumulation of protein aggregates containing caspase-8 and subsequent caspase-8 activation. As protein aggregation upon the inhibition of proteasomal and autophagic degradation pathways is mediated by the ubiquitin-binding protein SQSTM1/p62 and the autophagy-related protein LC3, silencing of p62 and LC3 promotes cells from MDEG G430F-induced cell death. Our results uncover a new mechanism of caspase-8-mediated apoptosis induced by intracellular calcium overload that is dependent on the autophagy-related proteins LC3 and p62 upon hyperactivation of DEG/ENaC channels.

The degenerin/epithelial sodium channels (DEG/ENaC)2 are a family of amiloride-sensitive cell surface channels that allow apical entry of cations including Na+ and Li+ into cells. Some of these channels are nonselective and are permeable to larger ions such as K+ and Ca2+. A subset of these channels is activated by extracellular protons and is hence termed acid-sensing ion channels. Proper function of these channels is critical for maintaining cellular ion homeostasis and osmotic balance. Although previously thought to be mostly expressed in fluid absorptive epithelia and in neurons, these channels have since been found to be expressed in most tissues (1, 2). Dysregulation of DEG/ENaC channels has been implicated in numerous pathological conditions ranging from hypertension, anxiety disorder, and neurodegenerative disorder to cancer, many of which are associated with hyperactivation-induced cytotoxicity. Modulation of DEG/ENaC channel activity has become an emerging therapeutic opportunity (2–4).

The cytotoxicity of DEG/ENaC hyperactivation was initially recognized in Caenorhabditis elegans. The gain-of-function mutants of DEG-1, MEC-4, and MEC-10 led to necrosis-like cell death in C. elegans sensory neurons (5, 6). It has been suggested that the rise in intracellular Ca2+ induced by the gain-of-function mutants of the DEG/ENaC channels is the cause for cell death (7–9). This DEG/ENaC hyperactivation-induced cell death in C. elegans is independent of the canonical apoptosis regulators EGL-1, CED-9, and CED-3, but rather is mediated by the Ca2+-dependent activation of calpain and aspartyl proteases (10).

The mammalian degenerin homolog (MDEG) was originally cloned from human and rat brains (11). Similar to the gain-of-function DEG-1(d), MEC-4(d), and MEC-10(d) mutants in C. elegans, MDEG overexpression and hyperactivation leads to cell death (12). Our results and those of others (12) indicate that the cellular mechanisms of DEG/ENaC hyperactivation-induced cell death are conserved from nematodes to mammals.

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The abbreviations used are: DEG/ENaC, degenerin/epithelial sodium channels; MDEG, mammalian degenerin homolog; ROS, reactive oxygen species; BAPTA-AM, 1,2-bis(o-aminophenoxy)ethane-N,N′,N′′,N′′′-tetraacetic acid (acetoxymethyl ester); 3-MA, 3-methyladenine; LC3, light chain 3; TEMPO, 4-hydroxy-TEMPO; Triton, 4,5-dihydroxy-1,3-benzenedisulfonic acid disodium salt monohydrate; DOX, doxycycline hyyclate; PI, propidium iodide; z, benzyloxycarbonyl; CM-H2DCFDA, 5-(and-6) chloromethyl-2',7’ dichlorodihydrofluorescein diacetate acetyl ester; fmk, fluoromethyl ketone; PARP, poly(ADP-ribose) polymerase; Ub, ubiquitin; BMK, baby mouse kidney; WB, Western blot.

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C. elegans, the substitution of glycine 430 in MDEG with an amino acid bulkier than serine such as phenylalanine leads to its constitutive activation. Ectopic expression of the MDEG G430F mutant led to cell death in Xenopus oocytes, human kidney cell line HEK293, and C. elegans sensory neurons (11). However, the molecular mechanism underlying this MDEG G430F-induced cell death in mammalian cells remains unknown. In the present study, we provide evidence that MDEG G430F triggers protein aggregation-mediated caspase-8 activation and apoptosis.

**EXPERIMENTAL PROCEDURES**

**Cell Lines, Culture, and Transfection**—HEK293T and bax−/− bak−/− baby mouse kidney (BMK) cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% FBS, 100 units/ml penicillin, and 100 μg/ml streptomycin (Invitrogen). Lipofectamine 2000 (Invitrogen) was used for transfection. Rat MDEG and MDEG (G430F) expression constructs were a gift from Dr. Michel Lazdunski (11). MDEG and MDEG (G430F) sequences were amplified by PCR with primers (5′-AGTTGACATGACCTCAGAGGGACCCAGTTGA-G-3′ and 5′-AATCGATTTAGTAGTGATGTGTGTAGT-GTCCGACCAGATCTCTCAGGTTGC-3′) and were inserted into the pRev-TRE vector. cDNA sequences with MDEG and MDEG G430F fused with GFP at their C terminus were also inserted into the pRev-TRE vector. The retrovirus encoding rtTA2-M2 activator, TRSID silencer, and puromycin selection marker (pLIB-rtTAM2iresTRSIDiresPuro, a gift from Dr. Michele Lazdunski) (12) and the pRev-TRE retrovirus encoding MDEG, MDEG (G430F), MDEG-GFP, or MDEG (G430F)-GFP were used to co-infect cells. The infected cells were selected by puromycin (0.3 μg/ml) and the lentiviral expression plasmid (3 μg) using Lipofectamine 2000. The medium was replaced with fresh medium 4 h after transfection. 20 h later, the virus-containing cell culture supernatant was filtered through a 0.45-μm nylon filter and subsequently used to infect bax−/− bak−/− BMK cells. The infection procedure was repeated two more times every 12 h. 24 h after the last infection, cells were selected with 2 μg/ml puromycin. Retroviral infection was performed as described previously (13). Briefly, retrovirus was produced in HEK293T cells after transfection with LHC retroviral constructs (2.5 μg) and helper viral construct (1.5 μg). The same infection procedure as lentiviral infection was used, except that the viral supernatant was supplemented with 10 μg/ml Polybrene (Sigma).

**His-Ub Assay for Mammalian Cells**—3 × 10^5 cells were plated into a 10-cm culture dish. After overnight recovery, cells were transfected with 2 μg of pMT107 plasmid encoding poly-histidine-tagged Ub. 1 day after transfection, cells were collected, and the same number of cells was plated into multiple wells for treatment. After treatment, cells were collected, washed twice with phosphate-buffered saline (PBS), and resuspended in 1 ml of buffer A (6 mM guanidine-HCl, 0.1 M NaHPO4/NaOH, 10 mM imidazole, pH 8.0). After 10 s of sonication, 50 μl of nickel-nitrilotriacetic acid-agarose (Qiagen) equilibrated with buffer A was added to each sample and incubated for 3 h at room temperature with agitation. Agarose was precipitated by centrifugation and washed twice with buffer B (10 mM Tris-Cl, pH 8.0, 8 M urea, 0.1 M NaHPO4) and three times with 1:4 diluted buffer B. The precipitates were resuspended in 100 μl of 2X Laemmli buffer (4% SDS, 20% glycerol, 10% 2-mercaptoethanol, 0.004% bromphenol blue, 0.125 M Tris-HCl, pH 6.8) with 200 mM imidazole, heated at 95 °C for 10 min, and subjected to Western blotting.

**Measurement of Proteasome Activity**—2 × 10^4 cells were plated into a 24-well plate. The ubiquitin-luciferase bioluminescence imaging reporter (Ub-FL) (14) and *Renilla* luciferase control plasmids were transfected into cells, and medium was replaced with fresh medium 5 h later. 16 h after transfection, the transfected cells were left untreated or treated with DOX or other indicated combinations. 24 h after treatment, the luciferase activities were measured by the Dual-Luciferase reporter assay system.

**Detection of Intracellular ROS**—5 × 10^5 cells were plated into 6-cm dishes. After overnight recovery, cells were untreated or treated with DOX together with other agents. 24 h after treatment, cells were incubated with CM-H_2DCFDA (10 μM final concentration) for 30 min in the dark and collected. The collected cells were resuspended in DMEM without phenol red
MDEG Hyperactivation Induces Caspase-8 Activation

and analyzed with a flow cytometer using the CellQuest program.

Caspase-8 Activity Assay—Caspase-8 activity was determined using the Caspase-Glo 8 assay kit (Promega G8200), which uses the luminesogenic caspase substrate, following the manufacturer's instructions. Luciferase activities were read by a luminescence reader (SpectraMax M5, Molecular Devices).

Measurement of Cell Death—For cell viability, two methods were used. For PI exclusion assay, cells (including the detached ones) were collected and resuspended in culture medium with PI at 1 μg/ml. Cell viability was determined by flow cytometry using a FACSCalibur. For trypan blue staining, 0.5 ml of cells was counted under a phase contrast light microscope. For trypan blue staining, 0.5 ml of cells (around 1 × 10^6 cells/ml) was mixed with 0.1 ml of 0.4% trypan blue and incubated for 5 min at room temperature. Cells were counted under a phase contrast light microscope.

Calcium Imaging and Quantification—MDEG G430F was treated with DOX (0.5 μg/ml). To perform time-course experiments, cells expressing wild-type MDEG and MDEG G430F were cultured on 25-mm diameter glass coverslips in 6-well plates, and cells were loaded with the cytosolic Ca^{2+} indicator Fluo-4/AM (Invitrogen, 5 μM) at room temperature for 30 min in extracellular medium containing 121 mM NaCl, 5 mM NaHCO_3, 10 mM Na-HEPES, 4.7 mM KCl, 1.2 mM KH_2PO_4, 1.2 mM MgSO_4, 2 mM CaCl_2, 10 mM glucose, and 2% bovine serum albumin (BSA), pH 7.4, in the presence of 100 μM sulfipyrazone and 0.003% pluronic acid. After dye loading, cells were washed and resuspended in the experimental imaging solution (extracellular medium containing 0.25% BSA), and images were acquired using the temperature-controlled Carl Zeiss LSM510 META confocal imaging system with 40×/1.3 NA oil objective. Cytosolic Fluo-4 changes were analyzed and quantified using ImageJ software (National Institutes of Health) as described previously (15).

GFP-LC3 Puncta Observation and Quantification—Quantification of GFP-LC3 puncta was performed as described previously (16). Briefly, cells stably expressing GFP-LC3 were fixed in 4% paraformaldehyde in PBS and observed under an inverted deconvolution microscope (Axiovert 200M; Carl Zeiss, Inc.) using the 63× oil objective lens. The cells with more than 10 obvious puncta were selected for quantification. ~100 cells were randomly selected and counted.

Statistics—Data from cell death assays are presented as mean ± S.D.

Image Processing and Densitometry Measurement—Images taken from deconvolution and confocal microscopes were viewed and processed by AxioVision LE and Zeiss LSM image browser, respectively. Images were processed in Adobe Photoshop to enhance the brightness and contrast. Densitometry of immunoblot bands was determined by ImageJ software, unless indicated otherwise.

RESULTS

Expression of MDEG G430F Leads to an Increase in Intracellular Calcium, ROS Generation, and Apoptosis—Because MEC-4(d)-induced cell death is not inhibited by the Bcl-2 homolog Ced-9 (17), we reasoned that MDEG hyperactivation may also induce cell death independent of mitochondrial apoptosis in mammalian cells. Therefore, we used the BMK cells with genetic deletion of both Bax and Bak (18), the two key regulators of mitochondrial apoptosis (19–21).

GFP-tagged wild-type and the G430F mutant MDEG were constructed into a tetracycline-regulated expression system and stably transfected into bax^−/− bak^−/− BMK cells. The addition of doxycycline induced progressive expression of MDEG proteins (Fig. 1A). As expected, MDEG was predominantly localized to the plasma membrane (Fig. 1B). We were unable to obtain a clear subcellular localization pattern for MDEG G430F due to the rapid induction of cell death (Fig. 1, C and D). Although cells expressing wild-type MDEG appeared normal, expression of MDEG G430F led to massive cell death after 48 h as indicated by cell morphology, nuclear fragmentation, and loss of plasma membrane integrity (Fig. 1, C and D). Furthermore, MDEG G430F-induced cell death is blocked by amiloride, the chemical inhibitor of ENaC (Fig. 1E), indicating that this cell death is specifically mediated by hyperactivation of MDEG.

Because hyperactivated MEC-4(d) mutant was reported to increase intracellular Ca^{2+} (9), we assessed whether MDEG G430F could also induce intracellular calcium rise. Indeed, ectopic expression of MDEG G430F, but not wild-type MDEG, led to a significant increase in intracellular Ca^{2+} levels (Fig. 2, A and B). Furthermore, the cell-permeable calcium chelator BAPTA-AM blocked cell death (Fig. 2C), whereas the cell-impermeable chelators BAPTA and EGTA did not (Fig. 2D), suggesting that the intracellular Ca^{2+} rise is a result of endoplasmic reticulum Ca^{2+} emptying. We went on to determine how cytoplasmic Ca^{2+} overload promotes cell death. To this end, ROS production is considered a main molecular consequence of intracellular Ca^{2+} rise (22). Indeed, MDEG G430F induced ROS production, which was inhibited by amiloride and the ROS scavenger TEMPO (Fig. 2E). Importantly, BAPTA-AM also inhibited ROS production (Fig. 2F), indicating that the cytoplasmic Ca^{2+} overload was responsible for the increased level of ROS. Furthermore, MDEG G430F-induced cell death was significantly inhibited by ROS scavengers Tiron and TEMPO (Fig. 2F). These results indicate that hyperactivation of the amiloride-sensitive ENaC MDEG can lead to intracellular Ca^{2+} rise, ROS overproduction, and cell death.

MDEG G430F-mediated Apoptosis Is Mediated by Caspase-8—Although the cell death induced by MDEG G430F occurs in Bax/Bak doubly deficient cells, the morphological appearance and nuclear fragmentation indicate that this cell death is still apoptotic (Fig. 1C). In agreement with this notion, the pancaspase inhibitor z-VAD-fmk at least partially blocked cell death (Fig. 3A), indicating the involvement of caspase activation. Moreover, immunoblotting showed that the caspase cascade was activated by MDEG G430F expression, indicated by apoptotic cleavage of caspase-8, caspase-3, and PARP (Fig. 3B). The activation of caspase-8, an initiator caspase, accompanied by its cleavage, was determined using the Caspase-Glo 8 luminesogenic assay kit (Fig. 3C). Consistent with the inhibition of apoptosis by amiloride, BAPTA-AM, ROS scavengers, and z-VAD-fmk (Figs. 1 and 2), these agents inhibited caspase-8 activation (Fig. 3D). To determine whether the activation of caspase-8 contributes to cell death induced by MDEG G430F, caspase-8 was knocked down using shRNA (Fig. 3E), which led
to a significant lower activity of caspase-8 (Fig. 3F) and conferred protection against MDEG G430F expression indicated by both PI exclusion and reduced cleavage of caspase-3 and PARP (Fig. 3, G and H). These data indicate that cell death induced by MDEG G430F is mediated by activation of caspase-8.
**FIGURE 2. MDEG G430F induces cell death via intracellular calcium rise and elevated ROS.**

A and B, a time course of intracellular Ca^{2+} in cells expressing wild-type MDEG or the MDEG G430F mutant following treatment with DOX. A, representative Fluo-4 fluorescence images were acquired using confocal microscopy at the indicated time points. B, quantification of Fluo-4 fluorescence intensity in A is expressed as fluorescence arbitrary unit (f.a.u.). Cumulative data are the mean ± S.D. of three independent experiments, each consisting of triplicate analyses of 20–40 cells.

C, MDEG G430F-expressing cells were treated with DOX in the absence or presence of the cell-permeable calcium chelator BAPTA-AM. Cell death was measured by PI exclusion. The average of three independent assays ± S.D. is shown.

D, MDEG G430F cells were treated with DOX alone or together with the cell-impermeable calcium chelators BAPTA or EGTA. Cell death was measured by PI exclusion. The average of three independent assays ± S.D. is shown.

E, MDEG G430F cells were treated with DOX alone, or together with amiloride, BAPTA-AM, or TEMPOL. 24 h after treatment, cells were incubated with CM-H_2DCFDA for 30 min in the dark and then resuspended in DMEM without phenol red and analyzed by flow cytometry. Note increased ROS production upon DOX treatment, which is reduced in the presence of amiloride, BAPTA-AM, or TEMPOL.

F, MDEG G430F cells were treated with DOX alone or in the presence of the ROS scavengers Tiron or TEMPOL. Cell death was measured by PI exclusion. The average of three independent assays ± S.D. is shown.
MDEG G430F Leads to Proteasome Inhibition and Accumulation of Polyubiquitinated Caspase-8—We then went on to study how caspase-8 is activated upon hyperactivation of MDEG. We recently reported that caspase-8 can be activated, independently of death receptors, by the inhibition of intracellular protein degradation machineries (23, 24). Indeed, expression of MDEG G430F led to an accumulation of polyubiquitinated proteins (Fig. 4B), indicating an inhibition of intracellular protein degradation machineries. Intracellular calcium rise and ROS have been shown to inhibit proteasome activity (25–27). Because MDEG G430F could induce intracellular calcium rise (Fig. 2, A and B) and ROS generation (Fig. 2D), and amiloride, BAPTA-AM, and ROS scavengers could block caspase-8 activation (Fig. 3G), it is possible that MDEG G430F can lead to caspase-8 activation via proteasome inhibition. Indeed, increased levels of slow migrating caspase-8 species were observed in SDS-PAGE upon MDEG G430F expression (Fig. 3B). To determine whether MDEG G430F can lead to proteasome inhibition, we utilized a ubiquitin-luciferase reporter construct (14). This reporter construct is rapidly degraded by the 26 S proteasome, leading to low luciferase activity. Upon inhibition of proteasomal protein degradation, luciferase levels increase (14). Although wild-type MDEG did not alter proteasome activity, MDEG G430F expression led to strong proteasome inhibition (Fig. 4A). This proteasome inhibition was alleviated by amiloride, BAPTA-AM, TEMPOL, and Tiron (Fig. 4A), indicating that it was mediated by MDEG hyperactivation-induced intracellular calcium rise and ROS production. Consistent with proteasome inhibition shown by the luciferase reporter assay, MDEG G430F expression led to accumulation of polyubiquitinated proteins that was blocked by amiloride, BAPTA-AM, and TEMPOL, but not by z-VAD-fmk (Fig. 4B).
Consistent with the literature showing that profound proteasomal inhibition can lead to the inhibition of the autophagolysosomal degradation (28, 29), MDEG G430F-expressing cells showed decreased autophagy flux indicated by reduced LC3 accumulation upon the treatment with the lysosomal inhibitor bafilomycin A1 (Fig. 4C).

To demonstrate that this MDEG G430F-induced inhibition of protein degradation can lead to an accumulation of polyubiquitinated caspase-8, His-tagged ubiquitin was transfected into cells expressing either wild-type MDEG or MDEG G430F. His pulldown was performed and showed increased caspase-8 ubiquitination only upon MDEG G430F expression (Fig. 4D), which was reduced by amiloride, BAPTA-AM, and TEMPOL (Fig. 4E). The amount of ubiquitinated caspase-8 correlated with apoptotic cleavage of caspase-8, caspase-3, and PARP (Fig. 4F). Together, these results indicate that MDEG G430F induces apoptosis by promoting proteasome inhibition and accumulation of ubiquitinated caspase-8.

MDEG G430F Leads to Caspase-8 Aggregation and Association with LC3 and p62—Inhibition of protein degradation has recently been shown to promote caspase-8 oligomerization via its interaction with p62 and LC3 (23, 24, 30). Proteasome inhibition leads to accumulation of ubiquitinated caspase-8. The aggregates are recruited by the ubiquitin-binding protein SQSTM1/p62 and shuttled to autophagosomes for degradation via the interaction between p62 and the autophagy “receptor” protein LC3. As excessive proteasome inhibition also leads to the inhibition of autophagic degradation (31, 32), it eventually leads to the accumulation of protein aggregates that promote caspase-8 oligomerization, activation, and apoptosis (24, 30).

Because we observed proteasome inhibition and accumulation of ubiquitinated caspase-8 upon MDEG G430F expression (Fig. 4), we went on to determine whether MDEG G430F can lead to caspase-8 association with LC3 and p62. Indeed, immunofluorescence showed that MDEG G430F expression induced large-sized
punctate structures containing caspase-8 and GFP-LC3, which at least partially co-localized (Fig. 5A). Co-immunoprecipitation showed that MDEG G430F expression induced association of p62, LC3, and caspase-8 (Fig. 5B). These LC3-, p62-, and caspase-8-containing structures were reminiscent of the protein aggregates observed upon proteasome inhibition, which were distinct from starvation-induced autophagic puncta (24). The formation of the LC3-containing protein aggregates was blocked by amiloride, BAPTA-AM, and TEMPOL, but not by z-VAD-fmk (Fig. 5C). Together, these data indicate that the protein aggregation is mediated by proteasome inhibition downstream of MDEG hyperactivation-induced intracellular rise of calcium and ROS and is upstream of caspase activation.

**MDEG G430F-induced Apoptosis Is Dependent on LC3 and p62**—The above results indicate that MDEG G430F-induced cell death resembles the recently described caspase-8-mediated...
apoptosis induced by proteasome inhibition, which is dependent on the autophagy-related proteins LC3 and p62 (24, 30). Therefore, we went on to determine whether MDEG G430F-induced apoptosis is also mediated by LC3 and p62. Silencing of LC3 (A) and p62 (B) was verified by immunoblotting. C, MDEG G430F cells with stable shNTC, shLC3, and shp62 were treated with DOX. Cell lysates were examined by immunoblotting for the indicated proteins. D and E, cell death of MDEG G430F cells with the stable expression shLC3 (D) and shp62 (E) was measured by PI exclusion. The average of three independent assays ± S.D. is shown. F, MDEG G430F cells with stable expression of GFP-LC3 were left untreated or treated with DOX or DOX plus 3-MA. 24 h later, cells were observed under a deconvolution fluorescence microscope. The percentage of cells with more than 10 obvious puncta was obtained from five blind counts. Representative images are shown. G and H, MDEG G430F cells were treated with DOX alone or together with 3-MA. 24 h later, caspase-8 activity was measured (G), and cell death was measured by PI exclusion (H). The average of three independent assays ± S.D. is shown. I, MDEG G430F cells were cultured with DOX alone or together with chloroquine (CQ). Cell death was measured by PI exclusion. The average of three independent assays ± S.D. is shown.

**DISCUSSION**

In this study, we show that expression of the hyperactivation mutant of the ENaC channel, MDEG G430F, leads to intracel-
lular calcium rise, ROS generation, and proteasome and autophagy inhibition. This proteasome inhibition leads to an accumulation of polyubiquitinated proteins including caspase-8. Polyubiquitinated proteins are normally recognized by p62, and via its interaction with LC3, are delivered to the autophagosome for degradation. However, as autophagic degradation is blocked by MDEG G430F, the LC3- and p62-containing protein aggregates accumulate and facilitate caspase-8 oligomerization, activation, and apoptosis. According to this model, one would predict that silencing of p62 or LC3, or preventing of LC3 accumulation on autophagosome membranes, would lead to decreased protein aggregation and suppressed apoptosis. This is indeed the case as shRNA silencing of p62 or LC3, or inhibition of LC3 membrane localization by 3-MA, suppressed MDEG G430F-induced cell death (Fig. 6). The caspase-8 activation upon protein aggregation is consistent with the literature showing that expanded polyglutamine repeats induce a caspase-8-mediated cell death (33) and is reminiscent of recent studies including those of our own group showing that pharmacological or genetic inhibition of proteasomal and autophagolysosomal protein degradation pathways leads to caspase-8-mediated apoptosis (23, 24, 30). In addition, our results are consistent with a study in C. elegans showing that inactivation of unc-51, bec-1, and lgg-1, the worm counterparts of autophagy proteins Atg1, Atg6, and Atg8 (LC3), suppresses neural degeneration induced by hyperactivation mutants of MEC-4 and DEG-1 (34).

Our results show that MDEG G430F can cause intracellular Ca$^{2+}$ rise (Fig. 2, A and B), and that MDEG G430F-induced cell death can be blocked by BAPTA-AM (Fig. 2C). This is consistent with the hyperactivation mutant of the MEC-4 Na$^{+}$ channel MEC-4(d), which has been found to induce C. elegans neuronal cell death by promoting intracellular calcium rise (8). In C. elegans, MEC-4(d) induces Ca$^{2+}$ influx and promotes endoplasmic reticulum Ca$^{2+}$ release (8, 9). However, in mammalian cells, the DEG/ENaC members including MDEG G430F are Ca$^{2+}$-impermeable (7, 11, 35). Hence, the MDEG G430F-induced intracellular Ca$^{2+}$ increase is unlikely to be mediated by extracellular Ca$^{2+}$ entry; rather, it is likely to be mediated through endoplasmic reticulum Ca$^{2+}$ release provoked by excess Na$^{+}$ entry. This notion is supported by our observation that MDEG G430F-induced cell death was blocked by the cell-permeable Ca$^{2+}$ chelator BAPTA-AM (Fig. 2C), but not by the cell-impermeable chelators BAPTA and EGTA (Fig. 2D). The precise mechanism for MDEG G430F-induced intracellular Ca$^{2+}$ rise remains to be determined.

Our results also show that the increased redox stress plays a critical role in MDEG G430F-induced cell death. Increased ROS generation was observed upon MDEG G430F expression (Fig. 2E). This increased ROS is likely to be caused by the intracellular Ca$^{2+}$ rise as BAPTA-AM could inhibit the ROS increase in MDEG G430F-expressing cells (Fig. 2E). It is well documented that cytoplasmic Ca$^{2+}$ rise can lead to increased ROS production, most likely via mitochondrial Ca$^{2+}$ overload that stimulates mitochondrial metabolic activity (22). In agreement with the critical role of redox stress in MDEG G430F-induced cell death, ROS scavengers could block cell death (Fig. 2, E and F), as well as the upstream events including proteasome inhibition, protein aggregation, and caspase-8 activation upon MDEG G430F expression (Figs. 3, 4, and 5). To our knowledge, this is the first report of a link between redox stress and cell death induced by DEG/ENaC hyperactivation.

As both cytoplasmic Ca$^{2+}$ rise and ROS generation play critical roles in MDEG G430F-mediated cytotoxicity, our results further show that the cytotoxic effect is mediated by the inhibition of proteasomal and autophagolysosomal degradation pathways, which leads to the accumulation of caspase-8-containing protein aggregates and subsequent caspase-8 activation and apoptosis. Both cytosolic Ca$^{2+}$ rise and ROS elevation have been implicated in the regulation of protein quality control and degradation machineries. Cytosolic Ca$^{2+}$ levels are known to modulate proteasome activity (25, 26). Accumulation of oxidatively damaged proteins can lead to a reduction in proteasome activity (27). Although the role of Ca$^{2+}$ in regulating autophagy remains controversial (36–39), some recent studies suggest that Ca$^{2+}$ inhibits autophagy (40–43). The discrepancy could be due to the extent and duration of intracellular Ca$^{2+}$ increase as well as cell type specificity. Our current study shows that autophagy flux is inhibited by MDEG G430F expression (Fig. 4C). The inhibition of autophagy may be caused by intracellular Ca$^{2+}$ increase and/or through profound proteasomal inhibition that can lead to inhibition of autophagy possibly by “clogging” the autophagic degradation system (24, 31, 32). Nevertheless, our finding that inhibition of autophagy flux is involved in MDEG G430F-mediated cell death is consistent with the model that LC3 serves as a platform for the formation of protein aggregates that contain p62 and polyubiquitinated caspase-8 upon inhibition of protein degradation (23, 24, 30).

Our study provides a previously unknown mechanism for the cytotoxicity induced by hyperactivation of DEG/ENaC in mammalian cells, namely through a Ca$^{2+}$-mediated protein degradation inhibition and caspase-8 activation. This is different from the neural degeneration mediated by the activation of calpain proteases and aspartyl proteases upon the channel hyperactivation in C. elegans. Although we do not exclude the possibility that Ca$^{2+}$-dependent activation of lysosomal hydrolases may be involved in MDEG G430F-mediated cytotoxicity, our findings provide a novel mechanism that involves the activation of caspase-8. This will help with the understanding of ion channel hyperactivation-related pathological conditions and provide novel therapeutic strategies to target protein aggregation by silencing p62 and/or LC3 for DEG/ENaC-related diseases such as heart failure and stroke.

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MDEG Hyperactivation Induces Caspase-8 Activation

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