A classic feature of apoptotic cells is the cell-surface exposure of phosphatidylserine (PtdSer) as an “eat me” signal for engulfment. We show that the Xk-family protein Xkr8 mediates PtdSer exposure in response to apoptotic stimuli. Mouse Xkr8−/− cells or human cancer cells in which Xkr8 expression was repressed by hypermethylation failed to expose PtdSer during apoptosis and were inefficiently engulfed by phagocytes. Xkr8 was activated directly by caspases and required a caspase-3 cleavage site for its function. CED-8, the only Caenorhabditis elegans Xk-family homolog, also promoted apoptotic PtdSer exposure and cell-corps engulfment. Thus, Xk-family proteins have evolutionarily conserved roles in promoting the phagocytosis of dying cells by altering the phospholipid distribution in the plasma membrane.

One Sentence Summary: Xkr-family transmembrane proteins mediate the externalization of phosphatidylserine, a signal that triggers the engulfment of apoptotic cells.
Phospholipids are distributed asymmetrically between the outer and inner leaflets of plasma membranes (1): PtdSer and phosphatidyethanolamine (PtdEtEtn) localize exclusively to the inner leaflet, whereas 60-70% of phosphatidylcholine (PtdCho) and sphingomyelin (SM) are found on the outer leaflet. This asymmetric distribution is disrupted during apoptosis, and exposed PtdSer on dying cells serves as an “eat me” signal to facilitate phagocytosis (2, 3). PtdSer exposure and the more general transfer of phospholipids between the inner and outer leaflets are likely mediated by phospholipid scramblases (1), the identities of which are disputed (4).

We previously generated a mouse Ba/F3 pro-B cell line (Ba/F3-PS19) with a high level of PtdSer exposure, constructed a cDNA library (of clones > 2.5 kb) and discovered TMEM16F, a transmembrane protein required for Ca\(^{2+}\)-dependent phospholipid scrambling but not apoptosis-dependent PtdSer exposure (5, 6). To identify molecules that mediate apoptotic PtdSer exposure, we introduced a Ba/F3-PS19 cDNA library (of clones 1.0-2.5 kb) into Ba/F3 cells, serially enriched for cells with high PtdSer exposure, and established a cell line (LD-PS5-2-2) with a high level of PtdSer exposure (Fig. 1A). LD-PS5-2-2 cells carried a cDNA encoding Xkr8, a member of the evolutionarily conserved XK protein family (7) (figs. S1 and S2). Analyses of the amino acid sequences of vertebrate Xkr8 orthologs using the programs Transmembrane Prediction (TMpred; www.ch.embnet.org) and Transmembrane Hidden Markov Model (TMHMM; www.cbs.dtu.dk) suggested that Xkr8 contains six transmembrane regions flanked by cytosolic N- and C-termini (fig. S3).

We transformed mouse T cell lymphoma WR19L cells with Fas (8) (WR-Fas). Fas ligand (FasL) efficiently induced apoptosis of the WR-Fas cells, accompanied by caspase-3 activation and PtdSer exposure (Fig. 1B, and fig. S4). The introduction of mouse Xkr8-GFP (mXkr8 fused to green fluorescent protein) but not mTMEM16F-GFP increased the fraction of PtdSer-exposing cells generated by FasL (Fig. 1B). The expression of mXkr8 short hairpin RNAs (shRNA) in WR-Fas cells decreased the amount of the endogenous mXkr8 mRNA by 76-82% (fig. S5) and the fraction of cells with FasL-induced PtdSer-exposure (Fig. 1C) but not levels of caspase-3 activation (fig. S5). The transformation of mXkr8 shRNA-expressing cells with human (h)Xkr8 cDNA, which is not recognized by the mXkr8 shRNAs, restored FasL-induced PtdSer exposure. hXkr8-GFP expressed in human 293T cells localized primarily to the plasma membrane (Fig. 1D), suggesting that Xkr8 functions at the cell surface to promote apoptotic PtdSer exposure.

Human PLB-985 leukemia and Raji lymphoma cells do not expose PtdSer during apoptosis (9, 10). Real-time RT-PCR indicated that the amount of Xkr8 mRNA in PLB-985 and Raji cells were 8% and 9%, respectively, of those in Namalwa cells (Fig. 2A). PLB-985 or Raji cell transformants expressing hXkr8 responded to apoptotic stimuli by exposing PtdSer (Fig. 2B). PtdSer exposure is necessary for the recognition of apoptotic cells by phagocytes (3, 10, 11). Accordingly, whereas apoptotic PLB-985 cells were rarely engulfed by macrophages, their Xkr8 transformants were frequently internalized (Fig. 2C). Caspase-3 activation, DNA fragmentation, cell death, and cell shrinkage occurred similarly in PLB-985 cells with or without Xkr8-expression, indicating that Xkr8 and PtdSer exposure had no obvious effects on other aspects of the apoptotic process (fig. S6). The program “CpG island searcher” (www.cpgislands.usc.edu) identified two CpG islands near the transcription start site of the hXkr8 gene (fig. S7). Bisulfite DNA sequencing (12) indicated that none of the 23
CpGs between -232 and +4 of hXkr8 gene was methylated in peripheral blood leukocyte (PBL), Jurkat or Namalwa cells (fig. S7). By contrast, these CpGs were methylated with more than 90% probability in PLB-985 and Raji cells. Treatment of PLB-985 cells with 5-aza-2'-deoxycytidine (DAC) increased Xkr8 mRNA levels (Fig. 2D). After 7 days of DAC treatment, all CpGs were demethylated (fig. S7), and Xkr8 mRNA levels were 91% of that in Namalwa cells. Accordingly, DAC-treated PLB-985 cells exposed PtdSer upon UV irradiation (Fig. 2E). We suggest that the methylation of CpG islands in the Xkr8 promoter in PLB-985 and Raji cells blocks Xkr8 gene expression and prevents apoptotic PtdSer exposure.

We assayed staurosporine-treated PLB-985 or Xkr8-expressing PLB-985 cells for PtdEtn exposure using RO09-0198, and for PtdCho and SM internalization using 1-oleoyl-2-[6-[(7-nitro-2-1,3-benzoxadiazol-4-yl)amino]hexanoyl]-sn-glycero-3-phosphocholine (NBD-PC) and NBD-sphingosine-1-phosphocholine (NBD-SM), respectively. Inhibitor of caspase-activated DNase (ICAD) was cleaved equally well in PLB-985 and its hXkr8 transformants following staurosporine treatment (fig. S8). By contrast, apoptotic hXkr8-expressing cells (but not parental cells) stained with RO09-0918 and internalized NBD-PC and NBD-SM (fig. S8), indicating that Xkr8, like TMEM16F (5), promotes the scrambling of multiple lipid species. Unlike TMEM16F, Xkr8 had no effect on the Ca^{2+}-induced exposure of PtdSer (fig. S9), suggesting distinct pathways control Ca^{2+}-induced phospholipid scrambling and apoptosis-induced scrambling. These findings are consistent with reports that B-cell lines from Scott patients, who carry a null mutation in TMEM16F (5), respond to apoptotic stimuli by exposing PtdSer (13), and that mouse Bak<sup>−/−</sup>Bax<sup>−/−</sup> platelets, which do not undergo apoptosis, expose PtdSer upon Ca^{2+} ionophore treatment (14).

An analysis of Xkr8 sequences from six vertebrates using the program CASVM (www.casbase.org) identified a conserved caspase 3-recognition site near the Xkr8 C-terminus (fig. S1). We generated a mutant version of hXkr8 (2DA) in which the putative caspase-recognition sequence at position 355 was changed from PDQVDG to PAQVAG (fig. S3). PLB-985 cells expressing wild-type hXkr8-GFP exposed PtdSer in response to staurosporine (Fig. 3A), accompanied by the loss of a 52-kD hXkr8-GFP band on polyacrylamide gels and the appearance of a 29-kD band detected with anti-GFP antibodies (Fig. 3B). Following staurosporine treatment, hXkr8(2DA)-GFP failed to promote PtdSer exposure and was not proteolytically processed; ICAD was cleaved in cells expressing either the wild-type or 2DA mutant of hXkr8, indicating similar caspase-3 activity in both cell lines. Processing of mXkr8-GFP at the caspase-recognition site during apoptosis was also observed in WR-Fas cells after treatment with FasL (Fig. 3C). The solubilized membrane fraction from cells expressing hXkr8-GFP was then incubated with human caspases. Western blot analysis with anti-GFP showed that caspases-3 and 7 cleaved the wild-type but not 2DA mutant hXkr8 (Fig. 3D). Thus, mammalian Xkr8 is activated to expose PtdSer via caspase-mediated cleavage of its cytosolic C-terminus.

mXkr8 mRNA was detectable in most mouse tissues (fig. S10), with notably high expression in the testes. We established mXkr8-conditional knock-out mice (fig. S11), from which we prepared mouse embryonic fibroblasts (MEF). After treatment with staurosporine, Xkr8<sup>+/−</sup> but not Xkr8<sup>−/−</sup> MEFs exposed PtdSer (Fig. 4A). Similarly, Xkr8<sup>flx/flx</sup> and TMEM16F<sup>−/−</sup> but not Xkr8<sup>−/−</sup> fetal thymocyte (IFET) cell lines exposed PtdSer in response to FasL (Fig. 4B), although caspase-3 was activated similarly in these cell lines (fig. S12). The transformation of Xkr8<sup>−/−</sup> IFETs with mXkr8 restored PtdSer exposure in response to FasL.
CED-8 is the only C. elegans homolog of Xk proteins and was previously shown to control the timing of programmed cell deaths (15) (fig. S2). To determine if CED-8 (like Xkr8) promotes phagocytosis, we examined ced-8 eggs for “floater” cells, which are generated in embryos defective in engulfment; floaters are a subset of apoptotic cells that if not engulfed (e.g., in ced-1, -2, -5, -6, -7, -10 or -12 mutants) detach from the embryo (Fig. 4C; fig. S13) (16, 17). ced-8 eggs contained floaters, and ced-8 mutations synergistically enhanced the number of floaters in engulfment mutants. This enhancement was dependent on the caspase gene ced-3 (fig. S14), which is required for apoptosis partially defective in engulfment. The PtdSer-binding protein MFG-e8::Venus (18) associated with 94% of apoptotic cell corpses in the ventral cords of wild-type animals but only with 21% of those in ced-8 mutants (Fig. 4D). Similarly, PtdSer was exposed on newly detached floaters from ced-1 but not ced-8 or ced-1; ced-8 embryos (fig. S15). Since 21% of ced-8 ventral cord cell corpses have normal PtdSer exposure, additional factors likely contribute to this process.

In short, the Xk-related proteins Xkr8 and CED-8 promote caspase-dependent PtdSer exposure during apoptosis. Based on the following observations, Xkr8 and CED-8 likely act at a late step in PtdSer exposure, possibly in phospholipid scrambling: i) Xkr8-deficient cells expose PtdSer in response to Ca²⁺, indicating Xkr8 is dispensable for steps prior to PtdSer exposure, including PtdSer biogenesis and localization; ii) Xkr8 is directly activated by caspase cleavage, suggesting Xkr8 does not function prior to the onset of apoptosis; and iii) Xkr8 and CED-8 are transmembrane proteins at the plasma membrane and therefore positioned to effect -- or interact with partners that effect -- the externalization of PtdSer during apoptosis.

Although intracellular concentrations of Ca²⁺ increase during apoptosis (19, 20), the involvement of Ca²⁺ in apoptotic PtdSer exposure is unclear (4) and our observations do not support a generalization. We found that FasL-induced PtdSer exposure was Ca²⁺-dependent in WR19L but not Ba/F3 cells and that when WR19L cells were transformed with Xkr8, they lost the Ca²⁺ requirement for the apoptotic exposure of PtdSer. These results, together with the constitutive activity of overexpressed Xkr8 in Ba/F3 but not other cells, suggest that Xkr8 might cooperate with Ca²⁺-regulated proteins in some cell-specific contexts.

The swift clearance of dead cells is essential for maintaining homeostasis, and the masking of PtdSer on apoptotic cells or the failure of the engulfment system can cause autoimmune disorders like systemic lupus erythematosus (3, 21). Our finding that in cancer cells Xkr8 is epigenetically repressed suggests a mechanistic link among inflammation, autoimmunity and cancer (22).

References and Notes:


19. D. L. Bratton et al., Appearance of phosphatidylserine on apoptotic cells requires calcium-mediated nonspecific flip-flop and is enhanced by loss of the
doi:10.1074/jbc.272.42.26159


doi:10.1038/nrrheum.2010.46


doi:10.1016/j.cub.2012.05.052

doi:10.1016/j.cub.2012.06.004

doi:10.1016/S0092-8674(85)80053-2


doi:10.1016/j.exphem.2003.07.005


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Fig. 1. Xkr8-mediated PtdSer exposure. (A) Staining of Ba/F3 and LD-PS5-2-2 cells with Cy5-Annexin V. (B) WR-Fas cells and transformants expressing mXkr8-GFP or mTMEM16F-GFP were treated with FasL and stained with Cy5-Annexin V. (C) WR-Fas cells transformed with scrambled shRNA, mXkr8 shRNA, or both mXkr8 shRNA and hXkr8 cDNA were treated with FasL and stained with Cy5-Annexin V. Representative FACS profiles for each group are shown. (D) 293T cells expressing hXkr8-GFP were observed by fluorescence microscopy. Scale bar, 10 μm.

Fig. 2. Epigenetic repression of Xkr8 in human cancer cell lines. (A) Abundance of hXkr8 mRNA relative to β-actin mRNA was determined by real-time RT-PCR. (B) The indicated cell lines and hXkr8 transformants were treated with apoptotic stimuli, and stained with Cy5-Annexin V. (C) PLB-985 cells and hXkr8-transformants were treated with UV, labeled with pHrodo, and incubated with peritoneal macrophages. FACS profiles for pHrodo-positive cells in CD11b+ cells are shown. Shown is the average % of pHrodo+ cells from three experiments. Right, macrophages (arrows) engulfing apoptotic cell were observed by fluorescence microscopy. (D) PLB-985 cells were treated with DAC, and Xkr8 mRNA was quantified relative to GADPH mRNA by real-time RT-PCR. (E) PLB-985 cells were treated with DAC for 5 days, exposed to UV, and stained with Cy5-Annexin V and PI.

Fig. 3. Activation of Xkr8 by caspase cleavage. (A and B) PLB-985 and transformants expressing hXkr8-GFP or hXkr8 2DA-GFP were treated with staurosporine (STS) and stained with Cy5-Annexin V (A). In (B), the cell lysates were analyzed by western blotting with anti-GFP and anti-ICAD antibodies. (C) WR-Fas and transformants expressing GFP, mXkr8-GFP or mXkr8 2DA-GFP were treated with FasL. Cell lysates were analyzed by western blotting with anti-GFP. (D) The membrane fraction of PLB-985 cells expressing hXkr8-GFP (W) or the 2DA mutant was incubated with human caspases (C1 to C10, caspase-1 to caspase-10) and analyzed by western blotting with anti-GFP antibody.

Fig. 4. Promotion of PtdSer exposure and cell-corpse engulfment by mouse Xkr8 and C. elegans CED-8. (A) MEFs from Xkr8+/− and Xkr8−/− embryos were treated with staurosporine (STS) or control buffer for 8 h and stained with Cy5-Annexin V. (B) IFETs of the indicated genotypes were treated with FasL and stained with Cy5-Annexin V and PI. (C) The number of floater cells per egg was counted for each genotype. Error bars, standard deviation; ***, p < 0.0001 in a Student’s t-test for each pair-wise comparison between ced-x and ced-8; ced-x double mutants. (D) PtdSer was detected using MFG-e8::Venus. Shown is the cell corpse of P12.aap (arrowhead), which undergoes apoptosis and is engulfed by P12.pa (arrow). The death of P12.aap is accompanied by PtdSer exposure in wild-type but not in ced-8 (n1891) animals. In addition, P12.pa, like other C. elegans engulfing cells (23, 24), exposes PtdSer on its outer plasma membrane during engulfment.
Suzuki et al. Figure 1
Suzuki et al. Figure 2
Suzuki et al. Figure 3
Suzuki et al. Figure 4
Supplementary Materials for

Xk-related protein 8 and CED-8 promote the exposure of phosphatidylserine on the surfaces of apoptotic cells


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This PDF file includes:

Materials and Methods
Figs. S1 to S15
Materials and Methods

Cell Lines, Recombinant Proteins, Antibodies, and Materials

Mouse interleukin-3 (IL-3)-dependent Ba/F3 cells (25) were maintained in RPMI-10% fetal calf serum (FCS, Gibco), 45 units/ml mouse IL-3, and 50 µM β-mercaptoethanol. Human PLB-985 (26), Jurkat, Namalwa, and Raji cells were grown in RPMI1640-10% FCS and 50 µM β-mercaptoethanol. Plat-E packaging cells (27) were grown in DMEM-10% FCS. Mouse IL-3 (28), and human FasL (29) were prepared as described. Rabbit anti-activated caspase-3 mAb was from Cell Signaling. Mouse anti-human ICAD mAb was from Medical & Biological Laboratories, and Alexa 488-labeled goat anti-rabbit IgG Abs were from Invitrogen. Staurosporine was provided by Kyowa Hakko Kirin.

Construction of the cDNA Library, and Identification of Xkr8

Using poly(A) RNA from Ba/F3-PS19 cells, cDNA was synthesized with random hexamers as primers, and a BstXI adaptor was attached as described (5). DNA fragments 1.0 to 2.5 kb in length were size-fractionated by electrophoresis through a 1% agarose gel and ligated into a BstXI-digested pMXs vector (30). Approximately 1.3 x 10^6 clones were produced with E. coli DH10B (Invitrogen) by electroporation. Using plasmid DNA from the cDNA library, retrovirus was produced in Plat-E cells, concentrated by centrifugation, and used to infect Ba/F3 cells as described (5). Cells treated with A23187 were stained on ice for 15 min with Cy5-Annexin V (Biovision) and for 2 min with 5 µg/ml Propidium Iodide, and sorted with a FACSAria (BD Biosciences). The cDNA integrated into the retroviral vector was identified by PCR (5) as the Xkr8 cDNA. Real-time PCR indicated that the amount of Xkr8 mRNA in Ba/F3-PS19 is similar to that in the parental Ba/F3 cells, suggesting that the strong PtdSer exposure in Ba/F3-PS19 cells is due to the mutation introduced into TMEM16F gene (5).

Expression Plasmids for Mouse and Human Xkr8, and Their Mutants

The coding sequences for mXkr8 (GenBank NM_201368) and hXkr8 (GenBank NM_018053) were prepared by RT-PCR from Ba/F3 and Namalwa cells, respectively. The pMXs puro c-GFP was constructed by inserting the GFP sequence into pMXs puro. The Xkr8 cDNAs were then inserted into pMXs puro c-FLAG (5) or pMXs puro c-GFP to express proteins tagged with FLAG or GFP at the C-terminus. To generate the D351A/D354A (2DA) mutant of mXkr8, and the D352A/D355A (2DA) mutant of hXkr8, the mouse and human Xkr8 cDNAs were mutated by recombinant PCR (31) using primers carrying the mutated nucleotides.

Primers used to prepare the mXkr8 and hXkr8 cDNA were as follows (in each primer, the BamHI or EcoRI recognition sequence is underlined): mXkr8, 5’- ATATGGATCCATCATGCCTCTGTCCGTGCACCA-3’ and 5’- ATATGAATTCGAGGACTCCATTCAGCTGCA-3’; hXkr8, 5’- ATATGGATCCGCACTCAGCTGCACTGCA-3’; hXkr8, 5’- CTAGGGTCCCAGCCACGAGGGTCCGCT-3’ and 5’- CTAGGGTCCCAGCCACGAGGGTCCGCT-3’. Primers used to generate the D351A/D354A (2DA) mutants of mXkr8, and the D352A/D355A (2DA) mutants of hXkr8 were: mXkr8 2DA, 5’- GGGACCTGCCCCTCCTGGTGACCTGGACCCCTAG-3’, and 5’- CTAGGGTCCCAGCCACGAGGGTCCGCT-3’, and hXkr8 2DA, 5’- AAGCCCGACCTCGCCAGGTAGCCCGGGGGCC-3’ and 5’- GGCCCGGGCTACCTGGCGGAGGGTCCGCTT-3’.
shRNA

Four shRNA expression plasmids for mXkr8 and scrambled shRNA in a pRS vector were purchased from OriGene. Among the four mXkr8 target sequences, the most effective was 5’-GAATCTGTGCCATCGCTTTTCAGCT-3’. WR19L cells were transfected by electroporation, and stable transformants were selected in medium with 1.0 µg/ml puromycin, and subjected to cloning by limited dilution. The Xkr8 mRNA was quantified by real-time RT-PCR.

Establishment of Xkr8 Knock-out Mice

Xkr8 conditionally targeted mice were generated by Unitech as a custom order. In brief, a neo-loxP cassette carrying the phosphoglycerate kinase promoter-driven neomycin-resistance gene flanked by FRT sequences was inserted into intron 3 of the Xkr8 gene. A 1.0-kb fragment containing exon 3 was replaced with a fragment carrying the corresponding sequence and a loxP sequence. The diphtheria toxin A-fragment driven by the thymidine kinase promoter was inserted at the 5’ end of the vector. Mouse Bruce-4 embryonic stem cells (32) were transfected with the targeting vector, and the G418-resistant clones were screened for homologous recombination by PCR. Positive clones were used to generate Xkr8+/NeoFRT mice, which were crossed with transgenic mice carrying cytomegalovirus enhancer-chicken b-actin hybrid promoter (CAG)-driven Cre recombinase gene (CAG-CRE) (33) or CAG-driven flippase variant (FLPe) gene (CAG-FLPe) (34). The resulting mice were backcrossed to the wild-type C57BL/6 to generate Xkr8+/− or Xkr8+/flox mice. Intercrosses of Xkr8+/− mice generated Xkr8−/− mice in a normal Mendelian ratio. All the mice were housed in a specific pathogen-free facility at Kyoto University, and all the animal experiments were carried out in accordance with protocols approved by Kyoto University.

Mouse Embryonic Fibroblasts and Fetal Thymocyte Cell Lines

Fibroblasts were prepared form E14.5 embryos of Xkr8+/− and Xkr8−/− mice, and cultured in DMEM containing 10% FCS. A fetal thymocyte cell line (IFET) was established by immortalizing fetal thymocytes with H-rasV12 and c-myc as described (35, 36). In brief, Xkr8+/lox mice were intercrossed, and thymocytes were obtained on embryonic day 14.5. Retrovirus carrying the genes for H-rasV12 and c-myc was produced in Plat-E cells with the pCX4 vector (37), and bound to RetroNectin-coated plates (Takara Bio) by centrifugation at 2,000 x g for 2-3 h. The thymocytes were attached to the retrovirus-coated plate by centrifugation at 400 x g for 5 min, and cultured in DMEM-10% FCS, 1 x non-essential amino acids, 10 mM Hepes-NaOH buffer (pH 7.4), 50 µM β-mercaptoethanol, 5 ng/ml mouse IL-7 (38) (PeproTech), and GlutaMax™ (Gibco). The resultant IFET cells were infected with Adeno-Cre (Adenovirus Cre/loxP, Takara Bio), and subjected to cloning by limited dilution. The TMEM16F−/− IFET cell line was described previously (6).

Transformation of Human and Mouse Cells

Retroviruses carrying a Xkr8 cDNA were produced by introducing the pMX-puro vector into Plat-E cells, concentrated by centrifugation, and used to infect Xkr8−/− IFET cells. Stable transformants were selected in medium containing 2.0 µg/ml puromycin, and the expression of recombinant protein was confirmed by Western blotting with an anti-Flag Ab (Clone M2, Sigma) or anti-GFP Ab (Clone JL8, Clontech). Mouse Fas cDNA (39) was introduced into IFET cells by retrovirus-mediated transformation, and its expression was confirmed by flow cytometry with the anti-mouse Fas mAb (Jo2) (40). Human PLB-985 and mouse WR19L cells were transformed by retrovirus infection with amphotropic retrovirus envelope or VSVγ envelope. In brief, retrovirus was generated by transfecting 293T cells with the pMXs retrovirus vector, pGP (Takara Bio) for
Gag and pol-fusion protein, and pE-ampho (Takara Bio) or pCMV-VSV-G-RSV-Rev (provided by Dr. H. Miyoshi, Riken). Virus in the culture supernatant was concentrated by centrifugation and used to transform cell lines. To express Xkr8-GFP, 293T cells were transfected with pMXs puroXkr8-GFP with Fugene 6 (Promega), and transformants were selected in medium containing 1.0 μg/ml puromycin.

**Induction of Apoptosis, Treatment with Ca²⁺ ionophore, and Flow Cytometry**

Apoptosis was induced with FasL, staurosporine, or UV. In brief, 5 x 10⁵ cells in 500 μl of medium were incubated at 37°C with 10-400 units/ml hFasL for 1.2-2.0 h or with 10 μM staurosporine for 1.5-8.0 h. For UV exposure, 1 x 10⁶ cells in 2 ml of PBS were exposed to 500-2000 J/m² UV radiation (254 nm) in a StrataLinker (Stratagene), and incubated at 37°C for 1.5-2.0 h in 4 ml of RPMI1640-10% FCS. The cell viability was assayed by WST-1 assay with 2-(4-Iodophenyl)-3- (4-nitrophenyl)-5-(2,4-disulfophenyl)-2H-tetrazolium, monosodium salt (WST-1; Dojin Laboratories) and 1-Methoxy-5-methylphenazinium methylsulfate as described (41). To detect the apoptotic DNA fragmentation, DNA extracted from cells was analyzed by electrophoresis on 1.5% agarose gel. To detect active caspase-3, 1 x 10⁶ cells were fixed at 37°C for 10 min in PBS containing 1% paraformaldehyde, washed with chilled PBS-0.5% BSA, and permeabilized by overnight incubation at -20°C in 90% methanol. Cells in 100 μl of PBS-0.5% BSA were then incubated with 200-fold-diluted rabbit anti-active caspase-3 at room temperature for 30 min, followed by incubation for 30 min with 1,000-fold-diluted Alexa 488-labeled goat anti-rabbit IgG. The cells were washed with PBS-0.5% BSA, filtered into the FACS tube, and analyzed by a FACSAria. To monitor A23187-induced PtdSer exposure, 5 x 10⁵ cells in 500 μl of Annexin V staining buffer (10 mM Hepes-NaOH buffer [pH7.4] containing 140 mM NaCl and 2.5 mM CaCl₂) were treated with 3.0-10 μM A23187, and analyzed with a FACSAria at 20°C.

**Assay for Phospholipid Scrambling Activity**

To detect phosphatidylserine (PtdSer) and phosphatidylethanolamine (PtdEtn) on the cell surface, cells were stained on ice for 15 min with 2500-5000-fold diluted Cy5-Annexin V (Biovision) or 800-fold diluted biotin-Ro09-0198 (42) followed by 1.0 μg/ml APC-streptavidin in Annexin V staining buffer in the presence of 5 μg/ml propidium iodide, and analyzed by a FACSAria or FACSCalibur (BD Biosciences). To assay the internalization of phosphatidylcholine (PtdCho) and sphingomyelin (SM), 1 x 10⁶ cells in 0.5 ml of Hank’s Balanced Salt Solution (HBSS) containing 1 mM CaCl₂ (HBSS-Ca) were incubated on ice for 7 min. An equal volume of 200 nM 1-oleoyl-2-[(6-[7-nitro-2,1,3-benzo[d]oxadiazol-4-yl]aminohexanoyl]-sn-glycero-3-phosphocholine (NBD-PC)(Avanti Polar Lipids), or N-[6-{(7-nitro-2,1,3-benzo[d]oxadiazol-4-yl]amino]hexanoyl}sphingosine-1-phosphocholine (NBD-SM) (Avanti Polar Lipids) in HBSS-Ca was added, and incubated at 20°C. Aliquots (150 μl) were mixed with 150 μl HBSS containing 5 μg/ml fatty-acid free BSA (Sigma-Aldrich) and 500 nM Sytoxblue (Molecular Probes), and analyzed by FACSAria.

**Engulfment of Apoptotic Cells by Macrophages**

Engulfment of apoptotic cells by macrophages was assayed essentially as described previously (43). In brief, cells were treated with UV to induce apoptosis and labeled with 0.1 μg/ml pHrodo succinimidyl ester (pHrodo, Invitrogen). Thioglycollate-elicited peritoneal macrophages were prepared from mouse peritoneal cavity 4 days after injection of 2 ml of 3% thioglycollate, and
incubated with apoptotic cells at 37°C for 2 h. The pHrodo-positive population in CD11b+ macrophages was regarded as the cells engulfing apoptotic cells.

**Treatment with 5-aza-2’-deoxycytidine, and Bisulfite Genomic Sequencing**

To treat human PLB-985 cells with 5-aza-2’-deoxycytidine (DAC, Sigma-Aldrich), 1.0 x 10^6 cells in 10 ml of RPMI-10% FCS were incubated with 0.5 µM DAC for up to 7 days. Since DAC is an unstable compound, the medium was changed every 24 h. After DAC treatment, the cells were divided into three portions: one for FACS to analyze the PtdSer exposure, one for real-time RT-PCR for Xkr8 expression, and one for methylation-specific PCR analysis (12). For the bisulfite genomic sequencing, DNA was modified with bisulfite using a kit (MethyEasy Xceed, Human Genetic Signatures). In brief, 3 µg DNA was denatured at 37°C for 15 min in 0.3 M NaOH, and treated with sodium bisulfite according to the protocol provided by the supplier except that the incubation time was changed to 90 min. The modified DNA was denatured at 95°C for 20 min, and amplified by PCR using primers (TTAGGGATTAGAATGTGTTT and CCTATACAAATAACCCAAC). PCR was carried out with EpiTaq HS polymerase (Takara Bio), and the product was cloned into the pGEM-Teasy vector for sequencing.

**Real-time RT-PCR**

Total RNA was reverse-transcribed using Superscript III reverse-transcriptase (Invitrogen) or High Capacity RNA-to-cDNA™ kit (Applied Biosystems). Specific cDNA was amplified in a reaction mixture containing LightCycler®480 SYBR Green I Master (Roche Diagnostics), and the mRNA was quantified at the point where the LightCycler System detected the upstroke of the exponential phase of PCR accumulation with the linearized plasmid DNA as reference. Primers for real-time RT-PCR were: mXkr8, 5’-GCGACGCCACAGCTCACACT-3’ and 5’-CCCCAGCAGCAGAGTTCCC-3’; mGapdh, 5’-AGCAGGCATCTGAGGGCCCA-3’ and 5’-GAGAGCAATGCCAGCCCCGG-3’; hXkr8, 5’-AGGCCGGGCCCATCATCCACT-3’ and 5’-TGCCGCCTGTTCAGGGCAGC-3’; and hβ-actin, 5’-GCATCCTCACCTGAAAGTAC-3’ and 5’-CTTAATGTCACGCAGGTTTC-3’.

**Caspase Treatment of Cell Lysates**

The membrane fraction was prepared from PLB-985 transformants expressing hXkr8-GFP or hXkr8 2DA-GFP as described (28) and solubilized in 20 mM Tris-HCl buffer (pH 7.2)-140 mM NaCl, 1% Triton X-100, 10% glycerol, and 1 mM (p-aminophenyl) methanesulfonyl fluoride (APMSF). After removing insoluble materials by centrifugation, the membrane proteins (20 µg) were incubated at 37°C for 1 h with 3 units of various recombinant human caspases (Biovision) in 100 µl of 50 mM Hepes-NaOH (pH 7.4), 50 mM NaCl, 5% glycerol, 5 mM DTT, 10 mM EDTA, 0.1 mM APMSF and 0.1% CHAPS, and analyzed by Western blotting.

**Western Blotting**

Cells were lysed in RIPA buffer (50 mM Hepes-NaOH buffer [pH 8.0], 1% NP-40, 0.1% SDS, 0.5% sodium deoxycholate, 150 mM NaCl, and 10% protease inhibitor cocktail). The lysates were mixed with 5 x SDS sample buffer (200 mM Tris-HCl [pH 6.8], 10% SDS, 25% glycerol, 5% β-mercaptoethanol, and 0.05% bromophenolblue), and incubated at room temperature for 1 h to detect Xkr8-GFP, or boiled for 5 min to detect other proteins. Proteins were separated by SDS-PAGE on a 10-20% gradient gel (Bio Craft), and transferred to a PVDF membrane (Millipore). The membranes were probed with 3000-fold-diluted mouse anti-GFP mAb, 3000-fold-diluted mouse
anti-human ICAD mAb, or 3000-fold-diluted rabbit anti-active caspase-3 mAb followed by incubation with 1,000-fold-diluted HRP-conjugated goat anti-mouse or rabbit Igs (Dako). The peroxidase activity was detected by the Western Lightning®-ECL system (PerkinElmer).

C. elegans Strains, Plasmids and Experimental Procedures
C. elegans strains were cultured as described (44) and maintained at 20°C. The Bristol strain N2 was used as the wild-type strain. Mutations used are listed below:

**LGI:** *ced-1(e1735, n2091).
LGI1: *ced-6(n1813), ced-7(n1892), ced-12(n3261).
LGV: *ced-2(e1752), ced-3(n3692, n2424), ced-5(n1812), ced-10(n1993).
LGVX: *ced-8(n1891, n1999, n2090), nIs106[*lin-11::gfp].
unknown linkage: *nIs398[*dyn-1::mfg-e8::Venus, *myo-2::dsRed].

Shed cells or floaters were counted in eggs between the 2-fold and 3.5-fold stages of development (approximately 450–600 min after the first cell division) using a Zeiss Axioskop wide-field microscope equipped with a 100x objective equipped and Nomarski differential interference contrast (DIC) optics as described (17). PtdSer exposure on the corpses was detected using secreted MFG-e8::Venus expressed from the transgene *nIs398* (18). All images were acquired using OpenLab software (PerkinElmer) and modified for publication using ImageJ.
**Fig. S1.** An alignment of the amino acid sequences of six vertebrate Xkr8 orthologs. The putative transmembrane regions are shaded blue, perfectly conserved amino acids are red, and the caspase-recognition sites are yellow.
Fig. S2. An alignment of amino acid sequences of mouse Xkr8, C. elegans CED-8, and Drosophila CG32579. Identical amino acids are red, and similar amino acids are pink. Similar amino acids are defined as residues belonging to one of the following groups: S, T, P, A, and G; N, D, E, and Q; H, R, and K; M, I, L and V; F, Y, and W. Putative transmembrane regions are underlined, and the caspase recognition site in mouse Xkr8 is indicated in green. The number at the end of sequence indicates the number of amino acids of each protein.
Fig. S3.
Xkr8 and its GFP-fusion proteins. (A) The predicted topology of mammalian Xkr8 proteins. Putative transmembrane regions are numbered. (B) Representations of wild-type and caspase-resistant (2DA) mutant forms of hXkr8 and mXkr8 fused to GFP. TM, transmembrane.
Fig. S4
FasL-induced activation of caspase-3. WR-Fas transformants expressing GFP, mXkr8-GFP or mTMEM16F-GFP were treated with FasL. The cell lysates were analyzed by western blotting with anti-active caspase-3 and anti-α-tubulin antibodies.
Fig. S5

Real-time PCR quantification of mXkr8 mRNA abundance (relative to GAPDH mRNA) in WR-Fas clones transformed with plasmid DNA carrying mXkr8 shRNA or scrambled (Scr) shRNA. Two clones for each transformant were analyzed. Right, effect of mXkr8-shRNA on caspase-3 activation. For each WR-Fas transformation (Scr shRNA-expressing, mXkr8 shRNA-expressing, and mXkr8 shRNA- plus hXkr8-expressing transformants), two independent samples were treated with FasL, lysed, analyzed by western blotting with anti-active caspase-3 and anti-α-tubulin antibodies.
**Fig. S6**

Xkr8 does not affect most aspects of apoptotic cell death. Human PLB985 cells (parental) and their human Xkr8-transformants (+hXkr8) were treated with 10 mM staurosporine (STS) for 4 hr or left untreated (control). (A) Activation of caspase-3. Paraformaldehyde-fixed cells were permeabilized by treating with methanol and stained with rabbit anti-active caspase-3. The FACS profiles of control and STS-treated cells are represented by green and open areas, respectively. (B) DNA fragmentation. DNA was prepared from control and STS-treated cells, submitted to agarose gel electrophoresis, and stained with ethidium bromide. (C) Cell death. Cell death was assayed by the WST-1 method as described in Materials and Methods. (D) Cell shrinkage. The cells were analyzed by FACS Aria. The FSC profiles represent cell size.
Fig. S7.
Methylation of the CpG islands in hXkr8 gene. (A) CpG islands in hXkr8 gene promoter. Open and filled boxes, 5’ non-coding and coding regions, respectively. Arrow, transcription start site. Each CpG is represented by a vertical bar. (B) The nucleotide sequence from -232 to +4 of the hXkr8 gene. The 23 CpGs are shown in red, and predicted transcription-factor recognition sites are underlined. (C) Methylation status of the CpG islands. Each circle represents a CpG between positions -232 and +4, and the extent of methylation is represented by shading: black, 76-100%; grey, 26-75%; white, 0-25%.
Fig. S8.
Characterization of the hXkr8-mediated scrambling of phospholipids. (A) Apoptotic cleavage of ICAD. PLB-985 cells and hXkr8-expressing transformants were treated with STS, and cell lysates were analyzed by western blotting with anti-ICAD. (B) Apoptotic exposure of PtdEtn. PLB-985 and hXkr8-expressing transformants were treated with STS, stained with biotin-RO peptide and streptavidin-APC and PI, and analyzed by FACS. (C and D) Scrambling of PtdCho and SM in apoptotic cells. PLB-985 and hXkr8-expressing transformants were treated with STS and incubated with NBD-PC (C) or NBD-SM (D). Unincorporated lipids were extracted and analyzed by FACSAria at two minute time intervals. The fluorescence intensity in the SytoxBlue-negative fraction is shown in arbitrary units as the internalized NBD-PC or NBD-SM.
**Fig. S9.**
hXkr8 does not alter Ca^{2+}-dependent PtdSer exposure. PLB-985 and its hXkr8 transformants were treated at 20°C with 10 μM A23187 and stained with Cy5-labeled Annexin V. Annexin V-binding to the cells was monitored by flow cytometry at two minute intervals for 10 min.
Fig. S10

*Xkr8* mRNA abundance in different mouse tissues. Tissue-specific *Xkr8* mRNA levels were determined by real-time RT-PCR and expressed relative to *Gapdh* mRNA.
Fig. S11.
Gene structures of wild-type, floxed and deleted alleles of mXkr8 and the targeting vector.
mXkr8 is not required for apoptosis-induced caspase-3 activation. (A) MEFs from Xkr8<sup>+/−</sup> and Xkr8<sup>−/−</sup> embryos were treated with (+) or without (−) staurosporine (STS) for 8 h. Cell lysates were analyzed by western blotting using anti-active caspase-3 or anti-α-tubulin antibodies. (B) The Xkr8<sup>−/−</sup>, Xkr8<sup>+/+</sup>, mXkr8-transformed Xkr8<sup>−/−</sup>, and TMEM16F<sup>−/−</sup> IFETs expressing Fas were treated with FasL, stained with anti-active caspase-3, and analyzed with flow cytometry. The FACS profiles for the untreated (open) and FasL-treated cells (filled) are shown. The percentage of cells carrying the active caspase-3 is indicated.
Fig. S13.
ced-8, the *C. elegans* homolog of mammalian Xkr8, promotes the engulfment of apoptotic cells. Shown are Nomarski DIC micrographs of (A) *ced-1(e1735)*, (B) *ced-8(n1891)*, and (C) *ced-1(e1735); ced-8(n1891)* eggs that contain unengulfed apoptotic cells (termed “floaters” and indicated by arrowheads), which have detached from the developing embryo.
The pro-apoptotic caspase CED-3 is required for the generation of floater cells in the eggs of ced-8 mutants and other mutants defective in cell-corpse engulfment. Shown are the combined numbers of embryonic floater and shed cells that were counted in the eggs of each genotype. ced-3 mutations cause the extrusion of shed cells (17). Shed cells are morphologically different from floater cells generated by mutants defective in engulfment. We observed that ced-1 and ced-8 mutations synergistically cause the appearance of floater cells. By contrast, ced-3 mutations caused the appearance of shed cells and suppressed the generation of floaters in all tested genetic backgrounds, including ced-1; ced-8. Thus, ced-3 is epistatic to ced-1 and ced-8 with respect to the generation of shed cells and the suppression of floater cells, indicating that the floater cells of ced-1; ced-8 mutants are dependent on apoptosis. For ease of presentation, we have indicated that ced-3 mutants generate shed cells (black bars) and that ced-1 and ced-8 mutants generate floaters (white bars). Error bars, standard deviations.
Fig. S15.

*ced-8* promotes PtdSer exposure on the cell surface of floaters, the unengulfed apoptotic cell corpses that detach from engulfment-defective mutant embryos. Shown are Nomarski DIC and fluorescent micrographs of floaters (indicated by arrowheads) generated by (A) *ced-1(e1735)*, (B) *ced-8(n1891)*, and (C) *ced-1(e1735); ced-8(n1891)* mutants. PtdSer was detected using secreted MFG-e8::Venus expressed from the transgene *nIs398*. All floaters imaged were generated by embryos at the 1.5-fold stage of development or earlier. At later stages of embryonic development, *ced-8* mutant floaters exhibit some PtdSer exposure, indicating that *ced-8* mutations delay the process of PtdSer exposure and that *C. elegans* contains additional mechanisms of PtdSer exposure.