Proteolysis in *Caenorhabditis elegans* sex determination: cleavage of TRA-2A by TRA-3

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The *Caenorhabditis elegans* tra-3 gene promotes female development in XX hermaphrodites and encodes an atypical calpain regulatory protease lacking calcium-binding EF hands. We report that despite the absence of EF hands, TRA-3 has calcium-dependent proteolytic activity and its proteolytic domain is essential for in vivo function. We show that the membrane protein TRA-2A, which promotes XX female development by repressing the masculinizing protein FEM-3, is a TRA-3 substrate. Cleavage of TRA-2A by TRA-3 generates a peptide predicted to have feminizing activity. These results indicate that proteolysis regulated by calcium may control some aspects of sexual cell fate in *C. elegans.*

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Proteolysis is an essential feature of all living cells and is responsible for eliminating proteins during normal degradative turnover and also when their activities are no longer required during processes such as cell cycle regulation and apoptosis [Nunez et al. 1998; Koeppl et al. 1999]. There is increasing evidence indicating that regulated or specialized proteolysis plays an important role in controlling gene expression, cell fate, and cytoskeletal architecture during development. For example, proteolysis allows an intracellular region of the Notch/LIN-12 receptor to translocate to the nucleus and to activate transcription [Kimble et al. 1998]. In addition, during the establishment of segment polarity in *Drosophila*, the transcriptional regulator Ci is activated in response to Hedgehog signaling, and is proteolytically inactivated when signaling is repressed [Aza-Blanc and Kornberg 1999]. We became interested in understanding how the activities of one family of regulatory proteases, the calpains, modulate cell fate when the *Caenorhabditis elegans* tra-3 sex-determining gene was shown to encode a member of this family [Barnes and Hodgkin 1996].

The calpains are calcium-regulated thiol-proteases that selectively cleave proteins to modulate their activities, and are found universally in eukaryotes [Sorimachi et al. 1997; Mykles 1998]. Typical mammalian calpains are heterodimeric, consisting of a large catalytic and a small regulatory subunit, and are ubiquitous in expression. Mammalian μ-, m-, and muscle-specific p94 calpains represent the most extensively studied large subunit isoforms. The large subunit has been divided into four domains (D-I–D-IV). D-II contains the catalytic triad of Cys, His, and Asn. D-IV contains four calmodulin-like calcium-binding EF hand motifs, which confer calcium dependence on proteolytic activity, and a fifth EF hand that may dimerize with the small subunit [Blanchard et al. 1997; Lin et al. 1997]. The recently solved crystal structure of m-calpain indicates that D-III forms a C2-like fold that may bind calcium [Hosfield et al. 1999]. The genome of *C. elegans* encodes many atypical calpains, but a small regulatory subunit has not yet been found [Consortium 1998].

The importance of calpains is emphasized by the finding that a defective p94 gene leads to human limb–girdle muscular dystrophy 2A [Richard et al. 1995]. However, substrates for p94 are unknown, and in general, little is known about the physiological activities of calpains, although they affect cell division, apoptosis, and cytoskeletal remodeling [Sorimachi et al. 1997]. Alterations in calpain activity are also associated with degenerative pathologies such as Alzheimer disease, cataract, and arthritis [Wang and Yuen 1999]. Therefore, the finding that tra-3 encodes a predicted calpain protease provided an excellent opportunity to adopt a genetic strategy for investigating calpain activity. Moreover, TRA-3 is the founding member of a new calpain subfamily that includes human and mouse homologs [Dear et al. 1997; Mugita et al. 1997]; therefore, our studies should also provide insights about these vertebrate homologs.

The genetic basis of sex determination has been analyzed extensively in the nematode *C. elegans,* which has two naturally occurring sexes, an XX hermaphrodite and an XO male [Meyer 1997; Kuwabara 1999; and references therein]. The genes controlling sexual fate decisions have been ordered in a hierarchical pathway involving negative regulation [Fig. 1A]. Somatic sex is determined by tra-1: High tra-1 activity leads to female development and low tra-1 activity leads to male development. At the molecular level, protein–protein interactions control many aspects of somatic sex determination [Fig. 1B]. During XX hermaphrodite development, it is postulated that the membrane protein TRA-2A inactivates one or more of the FEM proteins. It was shown that a cytoplasmic carboxy-terminal region of TRA-2A binds FEM-3 and inhibits its ability to promote male development [Mehra et al. 1999; Fig. 1B, left]. In turn, the transcriptional regulator TRA-1 promotes female development. During XO male development, TRA-2A is inactivated by its repressive ligand HER-1, thus allowing the FEM proteins to inactivate TRA-1 [Fig. 1B, right].

The tra-3 gene promotes female development in XX hermaphrodites [Hodgkin 1986]. The absence of tra-3 transforms XX hermaphrodites into pseudomales, but
has no effect on XO male development. The predicted TRA-3 protein was shown to share sequence similarity with members of the calpain protease family (Barnes and Hodgkin 1996); however, it was unknown whether TRA-3 had proteolytic activity or whether any of the sex-determining proteins were TRA-3 substrates. In this study, we have established that the \textit{C. elegans} TRA-3 protein functions as a protease. We have found that TRA-3 undergoes calcium-dependent autolysis, and that its proteolytic domain is essential for in vivo function. We have also shown that TRA-2A is a substrate for TRA-3 and that the feminizing activity of \textit{tra-3} is dependent on \textit{tra-2} in vivo.

**Results and Discussion**

\textit{TRA-3 requires an intact proteolytic active site to promote female sexual development}

The \textit{C. elegans} sex-determining gene \textit{tra-3} encodes a predicted atypical calpain protease; TRA-3 lacks the calcium-binding EF hands found in ubiquitous calpains and instead carries an unrelated domain T [Fig. 2A] (Barnes and Hodgkin 1996). In general, little is known about the regulation of atypical calpains or their physiological roles in development (Sorimachi et al. 1997). Thus, it was not clear that TRA-3 would display calcium-dependent proteolytic activity. As a first step toward understanding the role of TRA-3 in \textit{C. elegans} sex determination, we showed that a full-length \textit{tra-3} cDNA driven by the heat shock promoter (HS–TRA-3) rescued all aspects of the XX \textit{tra-3} mutant phenotype; 25/30 XX \textit{tra-3}; \textit{crEx10} animals expressing HS–TRA-3 developed as fertile hermaphrodites and not as pseudomales.

To test the role of the predicted catalytic domain, we mutated the catalytic cysteine to serine (C83S) and attempted to rescue XX \textit{tra-3} mutant animals with HS–TRA-3(C83S). However, we failed to detect any fertile XX \textit{tra-3}; \textit{crEx58} hermaphrodites after heat shock \((n = 55)\). This experiment was repeated using the native \textit{tra-3} promoter because we had obtained evidence that some forms of TRA-3 might be less stable when induced by heat shock than when expressed from the native promoter (S. Sokol and P. Kuwabara, unpubl.). However, the TRA-3(C83S) construct driven from the native \textit{tra-3} promoter was again unable to rescue the fertility of XX \textit{tra-3}; \textit{crEx66} or \textit{crEx67} mutants \((n = 35 \text{ and } n = 55)\), indicating that an intact active site is required for TRA-3 activity.

**The \textit{C. elegans} sex-determining \textit{TRA-3} protein is a protease**

To test more directly whether TRA-3 has proteolytic activity, we asked whether TRA-3 was capable of autolysis (Goll et al. 1992; Mykles 1998). This was achieved by programming calcium-depleted rabbit reticulocyte ly-
sates to synthesize \(^{35}\)S-labeled TRA-3 (Fig. 2A). In the absence of exogenous calcium, only full-length wild-type TRA-3 was detected (76 kD) (Fig. 2B, lane 1). However, after adding calcium, a TRA-3 cleavage product (~50 kD) was also detected (Fig. 2B, lane 2); this cleavage was consistently observed using independent batches of lysate. The generation of this cleavage product was dependent on active TRA-3. The addition of calcium failed to produce a cleavage product from a catalytically inactive TRA-3(C83S) (Fig. 2B, lanes 3,4), although wild-type and inactive TRA-3 were expressed at comparable levels (Fig. 2B, lanes 1,3).

How do we explain the basis for this calcium-dependent autolysis when TRA-3 is known to lack calmodulin-like EF hands? The recently solved crystal structure of m-calpain provides a possible explanation. This structure reveals that D-III unexpectedly possesses a C2-like fold (Hosfield et al. 1999). C2 domains participate variously in calcium binding, phospholipid binding, and in mediating protein–protein interactions (Nalefski and Falke 1996). Sequence similarities between the D-III regions of m-calpain and TRA-3 suggest that TRA-3 may also contain a C2 domain. We have also detected by sequence analysis a second C2 site in domain T of TRA-3. Because most known C2 domains bind calcium, these C2 domains may be responsible for the calcium dependence of TRA-3 activity, and perhaps for localizing TRA-3 to the membrane. However, because TRA-3 can promote female development when domain T is absent (Barnes and Hodgkin 1996), this indicates that the C2 domain associated with D-III is primarily responsible for the calcium dependence of TRA-3. Therefore, we further suggest that D-III also participates in the calcium-mediated regulation of other calpains. The purpose of the calcium-dependent autolysis of TRA-3 is unclear. Although some studies indicate that limited amino-terminal autolysis of calpains can lower the calcium concentration required to reach maximal proteolytic activation (Goll et al. 1992, Mykles 1998), the autolysis of TRA-3 observed by us is not similarly amino-terminal (Fig. 2C). Therefore, we speculate that autolysis probably inactivates TRA-3 and may serve to limit its activity.

**C. elegans sex-determining proteins are stably expressed and properly localized in Sf9 cells**

Physiological substrates for calpain proteases have yet to be identified. However, potential TRA-3 substrates are probably confined in function to sex determination and are unlikely to be involved in any essential cellular processes because the absence of TRA-3 does not affect XO male development. On the basis of our knowledge of the pathway controlling sex determination (Fig. 1), potential TRA-3 targets include TRA-2A, the FEM proteins, and the not yet cloned laf-1 product (Goodwin et al. 1997). However, we concentrated on evaluating the transmembrane TRA-2A protein as a candidate substrate for three reasons. First, it had been suggested that tra-3 might act as a positive cofactor for tra-2 because it displayed a weaker loss-of-function phenotype than tra-2 and because strong hypermorphic tra-2 dominant alleles could partially suppress the requirement for tra-3 activity (Doniach 1986; Hodgkin 1986). Second, the activity and substrate specificity of calpains may be promoted by their localization to the membrane (Suzuki and Sorimachi 1998), which is also the location of TRA-2A. Third, TRA-2A has a number of carboxy-terminal PEST sequences, which sometimes target proteins for proteolysis (Kuwabara et al. 1992; Shumway et al. 1999).

To initiate these studies, we used a baculovirus system to express TRA-3 and TRA-2A, because a *C. elegans* tissue culture cell line does not exist and because this system is suited to the expression of integral membrane proteins (Tate and Grisshammer 1996). We first established that a TRA-3 protein carrying both myc and 6xHis tags could be stably expressed in Sf9 cells without requiring exogenous cofactors, such as a small regulatory subunit, although we could not eliminate the possibility that such factors might be present in Sf9 cells (Fig. 3A,B, lane 1). As a control, we expressed an inactive, but otherwise identical, TRA-3(C83S) (Fig. 3A,B, lane 2). We found that both wild-type TRA-3 and TRA-3(C83S) were robustly expressed, migrating with a mobility of 76 kD. To demonstrate that the addition of myc and 6xHis tags did not affect wild-type TRA-3 activity, we showed that an equivalent epitope-tagged TRA-3 construct driven from the *C. elegans* heat shock promoter retained the ability to rescue completely tra-3 mutants (data not shown).

We next expressed the *C. elegans* sex-determining TRA-2A protein carrying a carboxy-terminal myc-tag in Sf9 cells. As shown by the immunoblot probed with anti-TRA-2 (C-term), TRA-2A is present in membrane-enriched extracts, and migrates as a doublet with a relative molecular weight of 112 kD (Fig. 3C, lane 2). The generation of this cleavage product was dependent on active TRA-3. The addition of calcium failed to produce a cleavage product from a catalytically inactive TRA-3(C83S) proteins expressed in Sf9 cells (see legend to Fig. 2A). (B) Immunoblot of Sf9 extracts expressing TRA-3 or TRA-3(C83S) probed with anti-myc (lanes 1,2). (C) Immunoblot of membrane-enriched Sf9 extracts expressing myc-tagged TRA-2A probed with anti-TRA-2. (D) Localization of TRA-2A to the plasma membrane of Sf9 cells using anti-TRA-2. Note that cell at right expresses TRA-2A more strongly than cell at left because of variability in infection efficiency.
molecular mass of 170 kD [Fig. 3C]. Only the upper band of the doublet is consistently detected with anti-myc or anti-TRA-2A [N-term, data not shown], suggesting that the lower band represents a partially degraded TRA-2A. Immunofluorescence analysis of infected Sf9 cells using anti-TRA-2A [C-term] verified that TRA-2A is associated with the plasma membrane and also with a perinuclear compartment that may be the endoplasmic reticulum [Fig. 3D]. An identical pattern of fluorescence was also observed using anti-myc [data not shown]. We also found that the ability to detect TRA-2A with anti-TRA-2A [C-term] or anti-myc was dependent first on permeabilizing the cell membrane with detergent, supporting the prediction that the carboxyl terminus of TRA-2A is cytoplasmic (Kuwabara et al. 1992; Mehra et al. 1999). Given that we could detect robust expression of TRA-3 and TRA-2A in Sf9 cells, and that the cellular localization and topology of TRA-2A appeared to be correct, we next tested whether TRA-2A might be a proteolytic substrate for TRA-3.

**The C. elegans sex-determining protein TRA-2A is a substrate for TRA-3 calpain**

As shown by the immunoblot probed with anti-TRA-2, coexpression of TRA-2A and TRA-3 in Sf9 cells led to the production of a doublet corresponding to full-length TRA-2A and a smaller 55-kD carboxy-terminal TRA-2A band. [Fig. 4A, lane 2]. Because the antibodies are directed against the carboxyl terminus of TRA-2A and detect a discrete 55-kD band, we suggest that the 55-kD band results from endoproteolytic cleavage [Fig. 4C]. We further found that the 55-kD TRA-2A peptide was only present when TRA-2A was coexpressed with TRA-3; we did not detect the 55-kD TRA-2A peptide when TRA-2A was expressed alone [Fig. 4A, lane 1] or when it was coexpressed with a catalytically inactive TRA-3[C83S] [Fig. 4A, lane 3]. Expression levels of TRA-3 and TRA-3[C83S] were comparable on the basis of Ponceau S staining [Fig. 4B]; the identity of these bands as TRA-3 was confirmed by Western analysis using anti-TRA-3 [data not shown]. These data strongly indicate that TRA-3 can cleave TRA-2A to generate a carboxy-terminal TRA-2A peptide, hence, TRA-2A is a TRA-3 substrate.

**The in vivo activity of tr-3 is dependent on tr-2**

If TRA-2A is a substrate for TRA-3 in vivo, then we would predict that the ability of tr-3 to promote female development should be dependent on wild-type tr-2. To test this hypothesis, we generated the transgenic array *crEx65*, which expresses both HS–TRA-3 and a *vit-2:gfp* reporter. The vit-2:gfp reporter was included in this array because it is a sensitive indicator of feminization: normally vit-2:gfp expression is only detected in the intestine of XX hermaphrodites (Yi and Zarkower 1999). As expected, GFP was expressed in the intestines of all XX *him-8; crEx65* hermaphrodites (*n* = 34), but never in the XO males (*n* = 36) [Fig. 5A,B]. However, after applying multiple heat shocks to provide additional TRA-3 activity, we found that many (18/44) XO males were partially feminized because they expressed GFP in their intestines [Fig. 5C,D]. These results show that excess TR-3 can feminize XO males. We next asked whether excess TR-3 could still feminize when wild-type tr-2 was absent. For this experiment, we expressed the *crEx65* transgene in XX tr-2 pseudomales; pseudomales might be expected to be more readily femi-
nized than XO males, if feminization by tra-3 is not tra-2 dependent. After providing multiple heat shocks, no GFP fluorescence or other signs of feminization were detected in XX tra-2; crEx65 pseudomales \((n = 60)\). These results show that (1) tra-3 overexpression causes limited XO intestinal feminization, (2) wild-type tra-2 is required for tra-3 function, and (3) tra-2 is epistatic to tra-3 in the sex determination pathway.

The role of TRA-3 and proteolysis in controlling sexual cell fate in C. elegans

How might cleavage of TRA-2A affect sex determination? First, we propose that the TRA-2A carboxy-terminal cleavage peptide has feminizing potential, because it is likely to encompass at least part of the region of TRA-2A shown previously to promote feminization, and may be comparable in sequence to the germ-line-specific TRA-2B (Fig. 4C) [Kuwabara and Kimble 1995; Kuwabara et al. 1998; Mehra et al. 1999]. Because TRA-2B is identical in sequence to a predicted carboxy-terminal intracellular feminizing domain of TRA-2A, it was proposed that TRA-2A/B would be subject to the same shared regulatory controls [Kuwabara et al. 1998]. We now extend this logic to include the TRA-2A cleavage peptide; more specifically, we propose that TRA-2A/B and the TRA-2A cleavage peptide share two regulatory sites, the FEM-3-binding region and the MX region.

The FEM-3-binding region was defined by the ability of non-membrane-bound TRA-2A peptides to bind FEM-3 and to feminize XO males [Kuwabara and Kimble 1995; Mehra et al. 1999]. The balance between the activities of TRA-2A and FEM-3 influences the decision to adopt either a female or male fate. The MX region was identified by analyzing self-sterile tra-2(mz) mutants that failed to make sperm [Doniaich 1986]. It was proposed that the sites of these mutations delineated a binding domain for a potential tra-2 germ-line repressor that helped to promote hermaphrodite spermatogenesis by inhibiting tra-2. Subsequently, the germ-line-specific TRA-2B might titrate such a repressor to promote the switch from spermatogenesis to oogenesis [Kuwabara et al. 1998]. Therefore, the TRA-2A cleavage peptide has the potential to promote female development by binding and inactivating FEM-3, as discussed previously [Kuwabara and Kimble 1995; Mehra et al. 1999], or perhaps to carry out functions similar to those ascribed to the germ-line-specific TRA-2B [Kuwabara et al. 1998].

Second, we propose that the TRA-2A peptide generated by tra-3 may potentiate the activity of TRA-2A, but not replace the need for full-length TRA-2A. It was shown previously that overexpression of TRA-2A completely transformed an XO animal from male to hermaphrodite. Under similar conditions, the carboxy-terminal region of TRA-2A partially feminized the intestine and tail of XO animals [Kuwabara and Kimble 1995], similar in effect to HS–TRA-3 overexpression in XO animals. In conclusion, TRA-3 probably acts as a potentiatot of tra-2 by proteolytically cleaving TRA-2A to generate a feminizing fragment.

Materials and methods

Nematode strains and handling

Standard methods for culturing and genetic manipulation of nematodes have been described [Sulston and Hodgkin 1988]. Because tra-3 has a maternal effect, all rescue experiments refer to tra-3(mz-) animals. Genes referred to in this manuscript: tra-3(e1767)IV; him-8(e1489) IV; unc-4(e1200II); and tra-2(e1059)III. tra-3 homoygotes were generated from the strain tra-3(n101)(dm); him-8 increases the frequency of XO male progeny from 0.2% to 37%. unc-4 is closely linked to tra-2 and displays a recessive uncoordinated phenotype, which facilitates identification of tra-2 unc-4 homozygotes. tra-2 strains were maintained as tra-2unc-4/immC1.

Plasmid construction

Transgenic expression plasmids, pPK247 carries an RT–PCR-generated full-length tra-3 cDNA cloned in pBluescript KSII (Stratagene) and verified by sequencing. pPK251 encodes HS–TRA-3 and carries the insert from pPK247 cloned in pID49.83 [A. Fire, Carnegie Institute, Baltimore, MD]. pSS33 encodes HS–TRA-3(C83S) and was derived from pPK251. pPK297 encodes HS–TRA-3 with carboxy-terminal myc and 6xHis tags and was derived from pPK251. pSS41 carries the native tra-3 promoter [S. Sokol and P. Kuwabara, unpubl.] fused to the insert from pSS23. Plasmids for in vitro transcription/translation, pSS65 encodes tra-3 and carries the insert from pPK247 cloned in pF7TAG, which carries the T7 promoter, β-globin 5’ UTR, and encodes an amino-terminal myc tag. pSS66 encodes TRA-3(C83S) and was derived from pSS65. Baculoviral expression constructs, pPK296 encodes TRA-3 with carboxy-terminal myc and 6xHis tags and was derived by cloning the insert from pPK247 into pBlue-BacA4 [Invitrogen] and ligating a sequence carrying the myc and 6xHis tags into a unique KpnI site of tra-3. pSS62 encodes TRA-3(C83S) and was derived from pPK296. pPK198 encodes the myc-tagged C. elegans sex-determining TRA-2A protein, and was constructed by ligating a 4.7-kb tra-2 DNA with a carboxy-terminal myc-tag into the baculovirus transfer vector pVL1392 [Invitrogen]. For details regarding plasmid constructions, contact author.

Generation of transgenic animals and heat shock experiments

Transgenic nematodes were generated by standard methods [Mello and Fire 1995] and carry a dominant Rol marker, rol-6(su1006) [pRF4 or pCes1943]. pCR2 carries the viti-2-gfp reporter [Yi and Zarkower 1999]. Extrachromosomal arrays were verified by PCR and are described below: crEx10, HS–TRA-3 [pPK251 + pRF4]; crEx58, HS–TRA–3(C83S) [pSS23 + pRF4]; crEx65, HS–TRA–3(myc,6xHis) + viti-2-gfp [pPK297 + pCR2 + pCes1943]; crEx66, crEx67 tra-3:TRA–3(C83S) [pSS41 + pCes1943]. Heat shock experiments consisted of incubating animals of the appropriate genotype daily for 1 hr at 33°C in an incubator or for 20–40 min in a 33°C water bath. XX tra-3(mz-) pseudomales are variably masculinized, but are rarely self-fertile except when raised at 15°C (Hodgkin 1986). Because of this variability in somatic phenotype, we scored a transgenic construct as capable of rescuing tra-3 mutants only if it restored self-fertility.

Expression and detection of in vitro-synthesized TRA-3

pSS65 and pSS67 were used as templates to synthesize 35S-labeled TRA-3 using the Reticulocyte Translation kit (Promega); samples were prepared in duplicate. After translation, one sample was adjusted to a CaCl2 concentration of 5 mM then all samples were incubated for 1 hr at 30°C. Samples were separated on a 10%–20% gradient polyacrylamide gel and processed for autoradiography.

Insect cell culture and production of recombinant baculovirus

Spodoptera frugiperda (Si9) cells were grown in T8N-FH medium (Sigma) with 10% FCS, 50 µg/ml gentamycin and lipids (GIBCO BRL), at 27°C, with shaking at 150 rpm. Recombinant Autographica californica nuclear polyhedrosis virus (AcMNPV) was generated with either the Max Bac or Bac-n-Blue system (Invitrogen) and propagated as instructed.

Immunofluorescence

Recombinant protein was obtained from bacteria expressing the carboxy-terminal 543 amino acids common to TRA-2A and TRA-2B (Kuwabara et al. 1998) and used to raise antisera in rats. Si9 cells were grown on sterile ring slides for 24–36 hr at 27°C. Cells
were fixed with 4% paraformaldehyde for 10 min, washed with PBS and permeabilized with 0.2% Triton X-100, then incubated overnight at 4°C with rat anti-TRA-2 serum (1:1000). After washing, cells were incubated for 30 min with FITC-conjugated anti-rat IgG (1:200). Slides were mounted for microscopy using a medium containing DABCO.

**Immunoblot detection of recombinant proteins**

Recombinant TRA-3 proteins were prepared for immunoblot analysis as follows. Sf9 cells were harvested, washed with PBS, and solubilized in buffer A (10 mM HEPES at pH 7.9, 2 mM EDTA, 2% Triton X-100, 2 mM DTT, 100 µM PMSF, 5 µM pepstatin, 10 µg/ml aprotinin) for 1 hr on ice. Extracts were electrophoresed under denaturing conditions and the proteins transferred to nitrocellulose; the efficiency of transfer was assessed by Ponceau S staining. Blots were probed with either 9E10 (anti-myc; 1:50) or anti-TRA-2 (1:5000) and visualized using ECL (Amersham). Membrane-enriched extracts were prepared by homogenizing Sf9 cells in buffer A, removing nuclei by centrifugation at 2K for 15 min at 4°C and pelleting membranes by centrifugation at 55K for 1 hr at 4°C. The membrane pellet was resuspended on ice in 25 mM HEPES (pH 7.9), 0.1 M NaCl, 2 mM DTT, 2 mM EDTA, 100 µM PMSF, 5 µM pepstatin, and 1% Triton X-100.

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