Proprotein Convertases Process and Thereby Inactivate Formylglycine-generating Enzyme

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This article containssupplemental Table S1.

Formylglycine-generating enzyme (FGE) post-translationally converts a specific cysteine in newly synthesized sulfatases to formylglycine (FGly). FGly is the key catalytic residue of the sulfatase family, comprising 17 nonredundant enzymes in human that play essential roles in development and homeostasis. FGE, a resident protein of the endoplasmic reticulum, is also secreted. A major fraction of secreted FGE is N-terminally truncated, lacking residues 34–72. Here we demonstrate that this truncated form is generated intracellularly by limited proteolysis mediated by proprotein convertase(s) (PCs) along the secretory pathway. The cleavage site is represented by the sequence RYSR72↓, a motif that is conserved in higher eukaryotic FGEs, implying important functionality. Residues Arg-69 and Arg-72 are critical because their mutation abolishes FGE processing. Furthermore, residues Tyr-70 and Ser-71 confer an unusual property to the cleavage motif such that endogenous as well as overexpressed FGE is only partially processed. FGE is cleaved by furin, PACE4, and PC5a. Processing is disabled in furin-deficient cells but fully restored upon transient furin expression, indicating that furin is the major protease cleaving FGE. Processing by endogenous furin occurs mostly intracellularly, although also extracellular processing is observed in HEK293 cells. Interestingly, the truncated form of secreted FGE no longer possesses FGly-generating activity, whereas the unprocessed form of secreted FGE is active. As always both forms are secreted, we postulate that furin-mediated processing of FGE during secretion is a physiological means of higher eukaryotic cells to regulate FGE activity upon exit from the endoplasmic reticulum.

Background: The ER-resident formylglycine-generating enzyme (FGE), essential for post-translational activation of all eukaryotic sulfatase enzymes, is N-terminally processed during secretion.

Results: Processing is mediated by furin at a conserved R/K-YS-R/K↓ cleavage site and leads to FGE inactivation.

Conclusion: Processing serves to regulate the amount of FGE activity following ER exit.

Significance: Processing is never complete, thus suggesting an extra-ER function of FGE.

Sulfatases form a family of enzymes that catalyze the hydrolysis of sulfate esters and sulfamates in a wide variety of substrates like glycosaminoglycans, sulfolipids, and steroid sulfates (1, 2). In pro- and eukaryotic sulfatases, post-translational modification of the crucial cysteine residue in the conserved CXPXRX motif to formylglycine (FGly) is a hallmark for their activation and sulfatases devoid of this modification are catalytically inactive. Multiple sulfatase deficiency (MSD), a rare but fatal lysosomal storage disorder in humans, is characterized by the production of all 17 human sulfatases with almost no FGly formation in their active sites (3). The formylglycine-generating enzyme (FGE) catalyzes this unique and critical modification in nascent sulfatase polypeptides in the endoplasmic reticulum (ER), and mutations in the FGE encoding gene SLUMF1 were discovered as the basis of MSD (4–8). Recently, a role for FGE in regulating cell lineage commitment was reported. FGE, via activation of sulfatases Sulfl and Sulfl2, was shown to control hematopoietic lineage development through FGE and Wnt signaling (9).

In eukaryotes, FGE is localized in the lumen of the ER. The mature 41-kDa protein in humans, lacking the signal peptide (amino acids 1–33), is an N-glycosylated monomer containing 8 cysteines. The core domain containing the active site exhibits a novel fold with remarkably low secondary structural elements that is stabilized by two Ca2+ ions and two intramolecular disulfide bridges (10). The presence of two catalytic cysteines (residues 336 and 341) in the active site, which are involved in binding and oxidation of the cysteine in the sulfatase polypeptide, is highly conserved in FGE homologues from prokaryotes to eukaryotes (11). One of the defining features unique to eukaryotic FGE is the presence of a 55-residue N-terminal extension (amino acid positions 34–88 in the mature form of the protein). We have previously shown that this N-terminal extension of human FGE, for which the structure is unknown, is required for activation of sulfatases in cultured cells. Especially a conserved pair of cysteines (residues 50 and 52) within this

4 The abbreviations used are: FGly, formylglycine; CHO, Chinese hamster ovary; CMK, chloromethylketone; EGF, enhanced GFP; ER, endoplasmic reticulum; FD, furin-deficient; FGE, formylglycine-generating enzyme; fl, full-length; MSD, multiple sulfatase deficiency; PC, proprotein convertase; rFurin, recombinant furin; STS, steroid sulfatase.
extension was shown to be involved, with Cys-52 being critical for this activation (12). Moreover, the N-terminal extension has been shown to confer efficient retention of FGE in the ER by interaction with ERP44, a redox sensor and retention factor for Ero1α and adiponectin (13, 14). FGE that escapes the ER retention machinery is secreted (15). Recently, the reuptake of secreted FGE has also been reported; and interestingly, the endocytosed FGE has been shown to activate sulfatases after it reaches the ER by an unknown mechanism (16).

However, in our previous studies we have shown that a large fraction of secreted FGE is in an N-terminally truncated form starting at glutamate 73 (15). The nature of this proteolytic truncation and the identity of the protease(s) involved have not been defined so far. It is well known that proteolytic processing mediated by proprotein convertases (PCs) along the secretory pathway activates and thereby regulates the function of several secreted proteins (17). In case of FGE, however, the mechanism and relevance of processing for controlling its function as a master regulator in sulfatase biogenesis have not been investigated so far, and truncation of a functionally indispensable N-terminal fragment of FGE during secretion validates a detailed analysis of this process.

In this study, we show that FGE is cleaved off its N-terminal extension during secretion by furin and other furin-like PCs. The N-terminal trimming of secreted FGE is abolished in furin-deficient cells, suggesting that furin is the major protease involved in this processing. The cleavage site is represented by a RYSR72 motif, which is conserved among higher eukaryotes and wherein residues Arg-69 and Arg-72 are absolutely required for cleavage of human FGE. Interestingly, residues Tyr-70 and Ser-71 are highly conserved and play a role in lowering the cleavage efficiency. Therefore, FGE is inefficiently cleaved during secretion. Furthermore, the N-terminally truncated FGE is devoid of its formyglycine-generating activity, suggesting that the furin-mediated processing during secretion leads to inactivation of FGE. As always full-length FGE is secreted as well, regulation by processing may in fact address a role of FGE executed outside the ER.

**EXPERIMENTAL PROCEDURES**

**Expression Plasmids**—The construction of plasmids used to express FGE-tagged C-terminally with either -RGS-His6 (pSB-FGE-His) or -HA (pBI-FGE-HA), FGE with an appended KDEL motif at the C terminus (pBI-STS), and myc-ERp44 (pBI-myc-ERp44) was described earlier (12). All variants of the cleavage site motif of FGE based on a ClustalW alignment in case of the sequences of group III that do not comprise a R/K/-motif motif RYSR (group I) or R/K/-/X-S-R/K/X (group II; X denotes any amino acid) or no common sequence (group III) at position P1-P4 of the cleavage site in most species of the given classifications (Fig. 3). The sequences were centered at P1 (Arg-72) or based on a ClustalW alignment in case of the sequences of group III that do not comprise a R/K/-/X-S-R/K/X motif. 16 amino acids corresponding to positions 65–80 in human FGE are displayed. All 88 sequences are shown in supplemental Table S1.

**Cell Culture and Transfection**—HT1080, HeLa, HEK293, and baby hamster kidney cells (American Type Culture Collection) were cultured at 37 °C under 5% CO2 in Dulbecco’s modified Eagle’s medium (DMEM; Invitrogen) containing 10% fetal calf serum (FCS) (Lonza). CHO-K1 and furin-deficient CHO-FD11 cells (kindly provided by Steven Leppla (21)) were cultured in DMEM supplemented with 40 μg/ml proline (Fisher Scientific). HT1080 Tet-On and MSDi Tet-On cells, for doxycycline-inducible protein expression were maintained as described earlier (12). All transfections were performed with Lipofectamine LTX as recommended by the manufacturer (Invitrogen). A stable CHO-FD11 Tet-On cell line was generated by transfection of CHO-FD11 cells with pUHrT62 (kindly provided by Nadja Jung) encoding the reverse tetracycline controlled transactivator and neomycin resistance vector pSB4.7pA in a 10:1 ratio (15). The stable clones were selected with medium containing 0.8 mg/ml neomycin (Invitrogen) and screened through Western blotting for doxycycline-dependent FGE expression after transient transfection with pBI-FGE-HA plasmid. CHO-FD11 cells stably expressing FGE-RGS-His6 were generated by transfecting CHO-FD11 cells with pSB-FGE-RGS-His6, and transfectedants were selected with 0.8 mg/ml G-418 sulfate (PAA). The stable clone was selected through FGE expression analysis by Western blotting.

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The phylogenetic tree (see Fig. 3, left) is based on the Newick format of 13 representative species of a total of 88 SUMF1 sequences from different species. For sequence logo generation through the WebLogo 3.0 program (19, 20) species could be divided into three subgroups, based on the presence of the motif RYSR (group I) or R/K/-/X-S-R/K/X (group II; X denotes any amino acid) or no common sequence (group III) at position P1-P4 of the cleavage site in most species of the given classifications (Fig. 3). The sequences were centered at P1 (Arg-72) or based on a ClustalW alignment in case of the sequences of group III that do not comprise a R/K/-/X-S-R/K/X motif. 16 amino acids corresponding to positions 65–80 in human FGE are displayed. All 88 sequences are shown in supplemental Table S1.
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Preparation of Cell and Medium Samples for SDS-PAGE and Western Blot Analysis—5 μg of pSB-FGE-RGS-His<sub>s</sub> vector was used for single transfections of HT1080, HeLa, HEK293, baby hamster kidney, and CHO cells. Conditioned medium was collected after 24 h and centrifuged (500 × g, 5 min). Cells were washed once with PBS, treated with trypsin (Lonza), and pelleted by 250 × g to remove the medium. Cell pellets were resuspended in PBS (pH 7.4) containing protease inhibitor mixture (Sigma) and lyzed by sonication 3 × 10 s on ice. For co-expression of FGE with one of the PCs in CHO-FD11 Tet-On cells 2 μg of pBl-FGE-HA and 2 μg of the plasmids encoding PCs (see above) were used for transient transfection. In case of transient transfections using pBl vector constructs the protein expression was induced by replenishing the medium with medium containing 2 μg/ml doxycycline (BD Biosciences) at 5 h post-transfection. After induction for 24 h, the cells and media were collected and processed as described above for further analysis by SDS-PAGE and Western blotting.

Western blot analyses were carried out using rabbit polyclonal antiserum against FGE and rabbit polyclonal anti-GFP antibody (Living Colors<sup>®</sup> A.v. Peptide Antibody, Clontech) as primary antibodies and the peroxidase-conjugated goat anti-rabbit secondary antibody (Invitrogen). Western blot signal was performed using AIDA 2.1 software package (Raytest), and calculations of FGE amounts were based on 10- or 20-ng FGE standard signals, present on the same blot.

**Immunoprecipitation—**HT1080 cells were cultured in four 14-cm plates with 5% FCS-containing medium for 48 h. Medium was collected and centrifuged 10 min at 1000 × g, to remove cell debris. Cells of one plate were harvested by trypsinization, pelleted at 1000 × g for 5 min, and lysed by sonication in PBS (pH 7.4) containing 0.1% Triton X-100 and protease inhibitor mixture, followed by centrifugation at 20,000 × g for 15 min. As a negative control fresh medium containing 5% FCS was used, and 0.01% Triton X-100 and protease inhibitor mixture were added to it and the cleared medium. All supernatants were preincubated with rabbit pre-immune serum for 30 min at 4 °C, and after addition of protein A-Sepharose CL-4B (Sigma-Aldrich) the bound material was pelleted down by centrifugation at 7000 × g for 10 min. The supernatants of the conditioned medium and the cell lysate were split into two parts, and either rabbit preimmune serum or rabbit FGE antiserum was added. After incubation at 4 °C and addition of protein A-Sepharose CL-4B the bound material was pelleted down by centrifugation at 7000 × g for 10 min. The pellets were washed stepwise as described earlier (22). The pellets were boiled in 1× Laemmlı buffer for 5 min at 95 °C and centrifuged at maximum speed for 5 min. 100% of the medium and 30% of the cell lysate supernatants were loaded for 12.5% SDS-PAGE followed by Western blotting and detection with FGE antiserum.

**Steroid Sulfatase (STS) Activity Assay and Western Blotting—**Activity assays of STS were performed as described earlier (15, 23). For Western blot analysis, polyclonal antisera against FGE and STS were used as primary antibodies. Horseradish peroxidase-conjugated goat anti-rabbit antibody was used as secondary antibodies. Signals of STS are given as relative amounts, i.e. related to signal intensities detected in cells expressing STS only. Relative specific sulfatase activities were calculated, i.e. catalytic activity divided by the Western blot signal (arbitrary units) and referred to that of cells expressing the sulfatase only (relative specific sulfatase activity = 1). Absolute values for this reference are given in the legends.

**In Vitro Furin Cleavage Assay and Treatment of Cells with RVKR-Chloromethylketone (CMK) Inhibitor—**CHO-FD11 cells stably expressing FGE-His<sub>s</sub> or HT1080 cells were grown for 24 h, and cells and medium were harvested. For treatment with furin in vitro cell pellets were resuspended in HEPES (pH 7.5) buffer containing 1 mM CaCl<sub>2</sub> and 0.5% Triton X-100 and lyzed in the absence of protease inhibitors by sonication on ice. Appropriate amounts of cell lysate and medium were incubated for 3 h at 25 °C with 4 units of furin (NEB) with or without 25 μM decanoyl-RVKR-CMK (Alexis Biochemicals) as indicated in the figures. For in vivo treatment, HT1080 Tet-On cells were transiently transfected with pBl-FGE-HA, and 4 h post-transfection, the medium was replenished with medium containing 20 ng/ml doxycycline and various concentrations of decanoyl-RVKR-CMK or only dimethyl sulfoxide as a carrier control. The cells and medium were collected after 16 h of induction/treatment for further analysis. The samples were boiled in Laemmli buffer and analyzed by SDS-PAGE and Western blotting with FGE antiserum.

**Extracellular Processing of FGE-containing Conditioned Medium—**CHO-FD cells stably expressing FGE-His<sub>s</sub> were cultured on 25-cm plates in 15 ml 5% FCS-containing medium for 24 h. Medium was centrifuged at 500 × g, and supernatants were sterile filtered to remove any floating producer cells. The conditioned medium was added to confluent 10-cm plates of CHO-FD11 (negative control), MDSi, HT1080, HeLa, and HEK293 cells for the indicated incubation times. Medium was collected and centrifuged at 500 × g. Equal amounts of medium were used for Western blot analysis.

**In Vitro FGE Activity Assay—**FGE activity was assayed using conditioned media that contained either secreted full-length (fl)-FGE (obtained from CHO-FD FGE-His<sub>s</sub> cells) or Δ72-FGE (obtained from CHO-FD Tet-On cells after transient co-transfection with pBl-FGE-HA and pIRES-Furin constructs). After expression for 48 h at 1% or 2.5% FCS-containing DMEM, the medium was collected and centrifuged at 500 × g for 5 min. The supernatant was analyzed by SDS-PAGE and Western blotting, and aliquots were used directly for activity testing.

The activity assay was performed in triplicate with three sets of conditioned media at pH 9.3 under standard conditions as described earlier (4) with some modifications; no BSA was added to the reaction mixture, and 0.01% Triton X-100 and 0.01% Triton X-100 and GSH was used as a reducing agent.

**RESULTS**

**FGE Is N-terminally Truncated during Secretion by a Non-saturable Mechanism in a Post-ER compartment—**FGE, an ER-localized enzyme, lacks the canonical (KDEL-like) ER retention signal and is retained in the ER via interactions with ERP44 by a saturable mechanism (13). It is known that recombiantly expressed FGE (41 kDa) in HT1080 cells is released into the medium, and the majority of the secreted protein represents an N-terminally truncated form of 37 kDa in size lacking residues
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![Diagram](image)

**FIGURE 1. FGE is N-terminally truncated in a post-ER compartment.** A, FGE is N-terminally processed only upon secretion. HT1080 Tet-On cells were transiently transfected with cDNA encoding FGE-HA or FGE with an appended KDEL signal (FGE-HA-KDEL). 6 h post-transfection, FGE expression was induced with 20 ng/ml doxycycline. After 22 h of induction, cells (C) and medium (M) were analyzed by Western blotting with FGE antiserum. B, FGE truncation is observed in various cell lines. FGE was transiently expressed in the indicated cell lines for 24 h, and cells and medium were analyzed by Western blotting with FGE antiserum. C, N-terminal processing of FGE is independent of its expression level. FGE was transiently expressed in HT1080 Tet-On cells, and expression was induced with the indicated concentrations of doxycycline. After 22 h of induction, cells and medium (in a ratio of 2:1) were analyzed by Western blotting. The amount of FGE in cells and medium was determined by calibration of the Western blot with known amounts of purified FGE (data not shown). D, endogenous FGE is also secreted and proteolytically processed. HT1080 cells were cultured for 48 h, and cell lysates and conditioned media were subjected to immunoprecipitation (IP) with FGE antiserum or preimmune serum (PIS). 100% of IP fractions were analyzed by Western blotting. Nonconditioned (non-c) cell culture medium served as a control.

34–72 (Δ72-FGE) (15). To examine whether this proteolytic processing occurs in a compartment along the secretory route, we expressed FGE and FGE with an appended KDEL sequence at the C terminus (FGE+KDEL) in HT1080 cells (Fig. 1A). Analysis of the cell homogenate and medium clearly shows that the FGE population escaping the ER retention machinery is eventually secreted (Fig. 1 lane 2 and thereby proteolytically processed, whereas retaining FGE in the ER through an appended KDEL sequence prevents the secretion and in turn the proteolytic processing (Fig. 1A, lanes 3 and 4). These data indicate that the transport of FGE to a post-cis-Golgi compartment during secretion is a prerequisite for cleavage. It should be noted that the unprocessed FGE in the medium (see also Fig. 1C) has a lower electrophoretic mobility in SDS-PAGE compared with the intracellular FGE due to modification of the attached N-glycan in the secretory pathway (see Ref. 15).

To analyze the proteolytic processing in detail and to exclude any cell type-specific effect, we studied the secretion profile of FGE in various cell lines. We transiently expressed FGE in HT1080, HeLa, HEK293, baby hamster kidney, and CHO cells. The analysis of the cell homogenate and medium revealed that FGE is secreted in all cell lines tested (Fig. 1B) and that usually the majority of the secreted FGE is in the truncated form. Of note, the extent of processing was comparable across cell lines except in CHO cells, wherein the truncation was less pronounced. The secretion of the unprocessed fl-form suggests that the processing mechanism could be saturated due to high FGE expression levels. To test this hypothesis, FGE was expressed under the control of a doxycycline-inducible promoter in HT1080 Tet-On cells. FGE expression was induced with increasing concentrations of doxycycline, and quantification of the amount of fl- and Δ72-FGE in the secretions shows that even a 7-fold increased FGE expression does not increase the proportion of the fl-form in the medium (Fig. 1C). Approximately 25–35% of secreted FGE is in the unprocessed form independent of the expression level, indicating that incomplete processing is not due to saturation of the processing machinery. To determine whether endogenous FGE is also secreted and whether it is secreted in a processed form, we immunoprecipitated FGE from untransfected HT1080 and HeLa cell homogenate and medium samples. In both cell lines FGE was traceable in the secretions, as shown in Fig. 1D for HT1080 cells. The majority of secreted FGE was in the truncated form, and surprisingly the unprocessed form was also detected to similar relative levels as observed under overexpression conditions.

The RYSR Motif in the N Terminal of FGE Is Indispensable for Proteolytic Processing of Secreted FGE but Not Required for FGE Activity in Vivo—Because many secreted glycoproteins that are processed along the conventional secretory pathway contain an RXXR-like motif, we speculated that the RYSR motif in FGE (residues 69–72, Fig. 2A) represents a potential PC cleavage site. This is supported by our previous finding that the N-terminally truncated FGE starts at glutamate 73 (15). Surprisingly, an in silico analysis using a widely used PC cleavage site prediction program (24) did not yield any potential cleavage motif in FGE, and only a recently published program PiTou predicts a cleavage at the RYSR motif, albeit with a low score (+0.93) (25).

To analyze biochemically the role of the RXXR-like motif in the N terminus of FGE, we generated the following alanine variants: FGE-R69A, -R72A, -R69A/R72A, -Y70A, and -S71A. Expression of FGE-R69A, -R72A, or -R69A/R72A led to a clear resistance to truncation, and all secreted FGE was in the unprocessed fl-form (Fig. 2B, lanes 4, 6, 8) indicating that both arginine residues (Arg-69 and Arg-72) are equally essential for pro-
Promote cleavage. Expression of FGE-S71A led to a nearly complete processing of secreted FGE (Fig. 2B, lane 12), whereas the Y70A mutation conferred partial resistance to cleavage, with 1.5-fold more FGE secreted in the fl-form compared with wild-type (wt) FGE (Fig. 2B, compare lanes 2 and 10). These data indicate that the RYSR motif is a bona fide cleavage motif and that residues Tyr-70 and Ser-71 determine cleavage efficiency.

Expression of sulfatases in immortalized MSD cells that lack endogenous FGE activity leads to the production of inactive sulfatases whereas co-expression of a sulfatase with FGE in these cells yields active sulfatases, thus providing a reliable system to investigate FGE-mediated FGly generation in vivo (12). Using this method we examined whether the RYSR motif is required for the activation of sulfatases. We transfected cDNAs encoding STS alone or co-transfected along with cDNAs encoding RYSR motif alanine variants of FGE in immortalized MSD Tet-On cells (Fig. 2C). Upon doxycycline induction, STS was barely active when expressed alone, whereas co-expression with FGE-wt increased the activity of STS approximately 50-fold, as observed previously (12). Co-expressing the alanine variants of the RYSR motif led to an increase in STS activity to similar levels observed for FGE-wt, indicating that the integrity of the RYSR motif is not required for the activity of FGE in vivo. In conclusion, the RYSR motif in the N terminus of FGE serves as an authentic cleavage motif rather than a role in the activity of cellular FGE.

The RYSR Motif Is Highly Conserved among Higher Eukaryotes—Because the RYSR motif was only recognized as a weak potential cleavage site by the prediction program, we analyzed the conservation of the cleavage site flanking regions across species in the animal kingdom. The whole N terminus including the cleavage site as well as the important Cys-Gly-Cys-motif that has been studied earlier (12, 13) is encoded by exon 1 of the SUMF1 gene. Notably, this region has no counterpart in homologous prokaryotic genes (26) and in fact arose as an extension in early eukaryotes and persisted to their present day descendants.

A phylogenetic tree with 13 representative species (Fig. 3, left) provides an overview for our set of 88 available eukaryotic sequences, which were arranged according to modern molecular taxonomy (27; see supplemental Table S1 for the full set of sequences). Within this taxonomy-based list of species we could identify three groups. Group III consisted of the 30 earlier
diverging eukaryote species from sponge (basal metazoan) to sea pineapple (urochordates), among which no significant sequence conservation was detected in the relevant region (Fig. 3, WebLogo III). By contrast, in the 58 later diverging eukaryote species from northern pike (ray-finned fish) to human the sequence conservation was detected in the relevant region (Fig. 3, WebLogo IV). The high degree of conservation of these seven consecutive amino acids (residues 65–80 of human FGE) and were generated as described under “Experimental Procedures.” A high degree of conservation of a single amino acid at a particular position is represented by a large size (in units of bits) of the amino acid.

FIGURE 3. The RYSR motif is conserved in later diverging eukaryotes. The phylogenetic tree (left) of 13 representative species, chosen out of a total of 88 analyzed species (see supplemental Table S1), was generated from Newick format according to modern molecular consensus taxonomy (27) and visualized with the program Phylodendron. The species were divided into three groups based on their taxonomy and the presence of RYSR or YS at the cleavage site position. Group I represents 36 species from euarchontoglires, laurasiatheres, and atlantogenata; group II consists of 22 sequences from marsupials to ray-finned fish; and group III includes 30 species of urochordates to basal metazoan. WebLogo 3.0 was used to create logos (I–III) of the three groups as well as a combined logo (IV) of all 58 later diverging eukaryote sequences (19, 20). The four sequence logos display the degree of conservation of amino acids at positions P8 to P8 of each group (representing residues 65–80 of human FGE) and were generated as described under “Experimental Procedures.” A high degree of conservation of a single amino acid at a particular position is represented by a large size (in units of bits) of the amino acid.

Notably, all 88 species from human to sponge contain the essential Cys-Gly-Cys sequence mentioned above, which is required for sulfatase activation (12). Therefore, the cleavage site motif arose later in the molecular evolution of the N-terminus and gained a function that may serve as a tool to regulate FGE activity in later diverging eukaryotes.

RYSR72 ↓E Represents a Unique Cleavage Motif That Imparts Suboptimal Cleavage Efficiency—Our observation that under any condition tested approximately 20–30% of secreted FGE-wt is in the unprocessed form (Figs. 1 and 2) suggested that the processing of the N-terminal extension of secreted FGE is suboptimal. The cleavage efficiency of mammalian PCs has been shown to be directly dependent on ~20 amino acid residues surrounding the cleavage site, and especially the positions P4 to P1’ (RYSR72 ↓E in FGE) are important. To determine biochemically the functional advantage of the conserved residues in the RYSR72 ↓E motif, we transiently expressed motif variants in HT1080 cells and quantified the extent of processing in the secretions by SDS-PAGE and Western blotting. Expression of the motif variants KYSR72 ↓E and RYSK72 ↓E, representing motifs found in group II species that contain Lys at position P4 to P1’ (RYSR72 ↓E in FGE) are important. To determine biochemically the functional advantage of the conserved residues in the RYSR72 ↓E motif, we transiently expressed motif variants in HT1080 cells and quantified the extent of processing in the secretions by SDS-PAGE and Western blotting. Expression of the motif variants KYSR72 ↓E and RYSK72 ↓E, representing motifs found in group II species that contain Lys at position P4 to P1’, respectively, led to truncation of the secreted protein, but the cleavage efficiency was lowered, as indicated by a doubling of the fl-FGE/Δ72-FGE ratio compared with that of FGE-wt (Fig. 4A). The occurrence of proline at position P1’ is known to compromise the cleavage efficiency of the PCs, and as expected, substitution of Glu-73 with proline (RYSR72 ↓P) also led to a reduction in processing, indicating that Glu-73 is conducive to cleavage.

Mutating the highly conserved Tyr-70 to position P3 to Phe (RFSR ↓E) or Lys (RKSR ↓E) led to a clear increase in cleavage efficiency, whereas substitution with Ser (RSSR ↓E) only had a minor effect (Fig. 4B). A dramatic increase in the cleavage efficiency was observed when Ser-71 at position P2 was mutated to a positively charged arginine residue (RYRR ↓E or RARR ↓E) to a level where virtually all of the secreted FGE was processed.

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From these data we conclude that RYSR↓E is a unique cleavage motif with the highly conserved Tyr (at position P3) and Ser (P2) conferring an inefficient cleavage property to FGE. Whether these residues, apart from probably being unfavorable for PC recognition, make contact to the FGE core domain (or to other proteins), which may restrict access of the cleavage site, remains to be determined.

Furin Mediates the Proteolytic Processing of Secreted FGE—To verify that the N terminus of FGE containing the RYSR motif is subject to processing by furin or furin-like convertases, we studied the secretion profile of FGE in cells in the presence of decanoyl-RVKR-CMK, an inhibitor for PCs (Fig. 5A). HT1080 Tet-On cells expressing FGE under the control of doxycycline were treated with only dimethyl sulfoxide (as a carrier control) or increasing concentrations of the inhibitor for 6 h after transfection. After 20 h of induction, cells (C) and medium (M) were analyzed by Western blotting with FGE antiserum (upper panels). The cleavage efficiency was quantified from these blots and expressed as signal ratio of fl-FGE/Δ72-FGE in the medium, as indicated below each medium lane in a bar graph. These data are representative of two independent experiments.

FIGURE 4. Analysis of the conserved RYSR↓E cleavage motif by alanine scanning mutagenesis. HT1080 Tet-On cells were transiently transfected with pBI plasmids encoding FGE-wt (with RYSR↓E motif) or mutants thereof (with the mutated residues in the RYSR↓E motif indicated in bold). FGE expression was induced with 20 ng/ml doxycycline, 6 h after transfection. After 20 h of induction, cells (C) and medium (M) (at a ratio of 2:1) were analyzed by Western blotting with FGE antiserum (upper panels). The cleavage efficiency was quantified from these blots and expressed as signal ratio of fl-FGE/Δ72-FGE in the medium, as indicated below each medium lane in a bar graph. These data are representative of two independent experiments.
face (30). In addition, a soluble form of furin can be generated by sheddases (31). To investigate the possibility of post-secretion processing, we incubated fl-FGE containing conditioned medium obtained from CHO-FD11 cells stably expressing FGE with different cell lines (Fig. 6B). In fact, FGE was processed after incubation with HEK293 cells but not with HeLa, HT1080, or immortalized MSD patient cells. In parallel, conditioned medium was incubated with CHO-FD11 cells to show that truncation of FGE does not occur due to the cultivation conditions used.

Further, increased incubation times with HEK293 cells led to increased level of Δ72-FGE in the conditioned medium (Fig. 6C). These data clearly show that FGE can be processed extracellularly by surface-exposed proteases and this cleavage is cell type-specific and time-dependent. The cell type specificity may represent the furin expression level of the cell line or, at least, the amount of cell surface-exposed and/or soluble furin.

**Processing of the N Terminus Leads to Inactivation of FGE**—We have recently shown that the N-terminal part of FGE (residues 34–68) is essential to activate sulfatases in vitro (12). By contrast, the N-terminal extension was not required for in vitro FGly-generating activity of purified secreted Δ72-FGE. However, this in vitro activity is dependent on the presence of the reductant DTT (26, 32). To assess the physiological consequence of the N-terminal truncation by furin on the function of FGE, we analyzed the FGly-generating activity of the processed and the unprocessed forms of secreted FGE in the presence of GSH, a physiological reductant, or DTT serving as a control. The in vitro FGE activity assay (based on mass spectrometry; see “Experimental Procedures”) was performed with conditioned media obtained from CHO-FD11 cells expressing either FGE alone or co-expressing FGE plus furin leading to secretion of unprocessed FGE or processed Δ72-FGE, respectively (Fig. 7B, Western blot panels). Reaction conditions (amount of FGE and incubation time) in presence of DTT were set to 50% turnover of the cysteine-containing substrate peptide (2526.3 m/z) to the FGly-containing product peptide (2508.3 m/z). The unprocessed FGE is active in the presence of both DTT and GSH as shown by the appearance of the product (2508.3 m/z) in the representative spectra (Fig. 7A, panels a and b), whereas processed FGE (Δ72-FGE) showed activity only in the presence of DTT but not when GSH was used as reductant (Fig. 7A, panels c and d). Quantitative analysis of the FGly-generating activity (percentage substrate turnover) of FGE and Δ72-FGE in the presence of GSH normalized to that of DTT (100%) revealed that the unprocessed form is as active with GSH as with DTT, but Δ72-FGE is barely active in the presence of GSH. These data clearly indicate that the FGly-generating activity of FGE with a physiological reductant is exclusively dependent on the presence of the N-terminal extension, and furin-mediated processing functionally inactivates FGE during secretion. This agrees with the earlier observation that the forced expression of truncated FGE in MSD patient cells does not lead to intracellular sulfatase activation in vivo (12).
Accessibility of the N Terminus of FGE for Furin-mediated Cleavage Is Abolished When FGE Is in Complex with ERp44—Although FGE is an ER resident protein, it lacks a canonical ER retention motif. We recently showed that FGE is retained in the ER by ERp44 via a thiol-independent mechanism but nevertheless forms a disulfide-linked covalent complex with ERp44 through its N-terminal Cys-50 and Cys-52 (13). Interestingly, when co-expressed together with ERp44 lacking the RDEL sequence (ERp44/H9004 RDEL), a larger fraction of FGE was secreted, mainly in the fl-form. This indicates that FGE when secreted along with ERp44/H9004 RDEL as a complex escapes N-terminal trimming. In this study, using the in vitro furin cleavage assay, we assessed furin-mediated processing of the FGE-ERp44 covalent complex (Fig. 8). Homogenates, pretreated with N-ethylmaleimide (to prevent post-lysis disulfide shuffling), from HT1080 cells expressing either FGE alone or myc-ERp44 together from a bi-directional promoter were incubated with rFurin and analyzed by SDS-PAGE under nonreducing conditions. Upon overexpression, intracellular FGE via its N-terminal cysteines (Cys-50 and Cys-52) forms covalently cross-bridged homodimers (Fig. 8, lanes 1 and 2) and, additionally, FGE-ERp44 heterodimers when co-expressed with ERp44 (Fig. 8, lanes 4 and 6) as previously observed (13). Removal of the N terminus by furin cleavage should lead to a disappearance of the homo-/heterodimeric FGE forms, which serves as an indicator for accessibility of the furin cleavage site in FGE. Upon treatment with rFurin, the signals for monomeric as well as the homodimeric fraction of FGE decrease with a concomitant increase in the signal corresponding to the N-terminally truncated form (Δ72-FGE), indicating that in both monomer and homodimer the furin-cleavage site is accessible to furin (Fig. 8, compare lanes 2 and 3; lanes 4 and 5). However, rFurin treatment does not abolish the covalent interaction between FGE and ERp44 as evidenced by the presence of an intact FGE-ERp44 heterodimer observed in untreated samples (Fig. 8, compare lanes 2 and 3; lanes 4 and 5) and a quantitative increase of the unprocessed FGE form due to heterodimer formation when analyzed under reducing (+SH) conditions (Fig. 8, lanes 3 and 5). These data show that the cleavage motif in the N terminus of FGE is inaccessible when in complex with ERp44 and is thereby protected from furin cleavage.

DISCUSSION

FGE as a Noncanonical Substrate for Furin and Other PCs—In this study we could show that furin-like PCs cleave the N terminus of secreted FGE, with furin itself being the most effective and primary protease that performs this N-terminal processing. Several lines of evidence support our conclusion. FGE, like other proteins that are processed by PCs, bears a conserved cleavage site sequence containing the minimal consensus motif [R/K]Xn[R/K]. Our phylogenetic analysis revealed that the RYSR motif of FGE is conserved in later diverging eukaryote species (Fig. 3); experimentally we show that this RYSR sequence represents an authentic processing motif, as alanine variants of the conserved arginines conferred resistance to
Moreover, the processing of FGE was abolished either when treated with RVKR-CMK, a peptide-based inhibitor of PCs, or when expressed in CHO-FD11 cells that are deficient for furin; replenishing furin in these cells by transient expression led to a complete processing of secreted FGE. Expression of PACE4 or PC5α also led to FGE processing, albeit to a lesser extent compared with furin. Further, both intracellular and secreted FGE was processed by recombinant furin under in vitro conditions. All of these data unambiguously lead to the conclusion that furin is the primary PC that mediates the processing of FGE during secretion.

The observation that this N-terminal processing occurs for endogenous secreted FGE signifies the physiological relevance of this processing event. Interestingly, a fraction (~20–30%) of secreted FGE, independent of its expression level, is found in the unprocessed form. The escape from complete processing even at the very low amounts of endogenous FGE exiting the ER may reveal a kinetic limitation during the processing step and, at the same time, the importance of secreted fl-FGE. Because we observed an intracellular accumulation of FGE upon treatment with brefeldin A indicating that an intact Golgi is necessary for secretion of FGE,5 it is very unlikely that the fraction of the unprocessed form represents a population of FGE secreted via unconventional trafficking routes. Rather, we find that the processing is inefficient due to properties of the cleavage motif.

A classical furin cleavage motif is defined by the presence of arginines at P1 and P4, but residues in close vicinity of the cleavage site play an important role in the recognition and cleavage efficiency of PCs (33). The presence of a positively charged amino acid at P2 (mostly lysine or arginine) has been shown to improve the processing efficiency of furin, whereas no clear consensus has been found for residues at position P3. Interestingly, the presence of Ser at position P2 in some substrates has been suggested to be unfavorable for cleavage by mammalian furin and other PCs (29). Our mutational analysis of Tyr-70 and Ser-71 residues indicates that this holds true also for FGE. Modifying the residues at positions P2 and P3 to positively charged residues, thus representing a more favorable

5 E. C. Ennemann, K. Radhakrishnan, M. Mariappan, M. Wachs, T. H. Pringle, B. Schmidt, and T. Dierks, unpublished results.
motif, led to a strongly increased efficiency of processing with a maximum effect observed by mutating Ser-71. Thus, the presence of Ser at position P2 is the major cause for inefficient processing of FGE. Interestingly, Tyr-70 and Ser-71 are absolutely conserved throughout evolution to a degree even higher than the essential basic (arginine) residues, supporting our conclusion that the RYSR motif in FGE is a noncanonical cleavage motif that evolved to confer suboptimal cleavage efficiency.

Why Should FGE Be Processed at All?—Both phylogenetic comparison and experimental data clearly demonstrate that the FGE protein during evolution gained a function by addition of the N-terminal extension harboring the Cys-Gly-Cys sequence. This motif is critical for the biological activity of eukaryotic FGE (12) and fully conserved from human to sponge. The N-terminal extension is found only in eukaryotes (encoded by exon 1), and the fact that eukaryotic Fgly generation is a co-translational event occurring in a specialized compartment, the ER, suggests that the N-terminal extension serves to adapt FGE to ER-based functioning in eukaryotes. One of the gained properties is ER retention through interaction of ERP44 with the N terminus (13); another seems to be related to inter- or intramolecular activation of the catalytic domain of FGE by the Cys-Gly-Cys motif (12). Further ER-specific aspects, such as competence to act on nascent sulfatase polypeptides emerging at ER import sites, might be associated with the N-terminal extension.

Interestingly, prokaryotic FGE from Streptomyces coelicolor, which lacks the N-terminal extension, when expressed in the cytoplasm of eukaryotic cells was shown to possess Fgly-generating activity acting on engineered cytosolic model substrates containing the Fgly modification signature (34, 35). On the other hand, we have shown that N-terminally truncated human FGE, when expressed in the ER, did not possess Fgly-generating activity, which agrees with the hypothesis that N-terminal processing irreversibly abrogates ER-based FGE functioning. On the basis of these two observations one can speculate that furin-mediated removal of the N terminus during secretion could serve a possible mechanism in later diverging eukaryotes to generate FGE that is active under extracellular conditions. However, an extracellular formylglycine generation has not been reported so far.

Processing by PCs as a Means of Enzyme Inactivation—Our data indicate that a direct consequence of N-terminal processing of secreted human FGE is its inactivation. The N-terminal extension was found to be essential for in vitro activity in the presence of glutathione, a physiological reductant, which does not sustain activity of N-terminally truncated FGE. This corroborates our previous observation that the N terminus is essential for biological activity in cultured cells. Because removal of this part renders secreted FGE inactive, we propose that the observed furin-mediated processing during secretion is a physiological means for regulation of FGE function/activity. The need for inactivation of the enzyme is not clear, but it might be a mechanism necessary to avoid generation of toxic aldehydes in extracellular or cell surface proteins in case that FGE evades its indirect ERP44-mediated ER retention.

Proteolytic processing mediated by PCs along the secretory pathway is a widely used necessary step in the activation or maturation of many proteins that are involved in various cellular processes. In contrast, inactivation of a protein function by furin-mediated processing only recently has been recognized as a novel mode of regulation, as shown for PCSK9 (36). The function of PCSK9, in mediating LDL receptor internalization and degradation, was shown to be inactivated by furin-catalyzed cleavage of PCSK9. FGE could represent another protein that is inactivated by proteolytic processing mediated by PCs. Nevertheless, one cannot exclude the possibility that the secreted N-terminally truncated FGE could perform as yet unknown extracellular functions other than Fgly generation.

Incomplete Processing as a Means to Provide Active FGE to the Extracellular Space?—On the other hand, it is tempting to assume that the inefficient processing could represent a functional tool for regulated release of FGE in the unprocessed active form. The low but significant secretion of fl-FGE even under endogenous expression levels might indicate such a regulated release of active FGE. In fact, it has been shown that secreted FGE can enter other cells by mannose receptor-mediated internalization and activate sulfatases in a paracrine manner after reaching the ER of the recipient cells (16). To exert this function, FGE should escape furin-mediated inactivation during secretion. As a proof of principle our data show that FGE, when in complex with ERP44, is barred from furin processing. This indicates that masking the cleavage site by an interacting protein during secretion could significantly hinder recognition by furin and could lead to the secretion of FGE in the unprocessed form. Alternatively, a population of FGE destined to be secreted in the active form could have an altered conformation wherein the cleavage site is masked due to intramolecular interactions of the N terminus to the core of the protein, thus preventing truncation. In combination with the suboptimal cleavage motif, an impaired processing by furin due to an inaccessible cleavage site (either by inter- or intramolecular interactions) could further increase the fraction of unprocessed secreted FGE. The secretion of FGE (14, 15) and possible reuptake by other cells (16) has been shown to be a multistep-regulated process, and our findings extend this complex regulation of FGE function to an additional level.

Perspective—The finding that FGE is secreted in the unprocessed form by furin-deficient CHO cells should pave the way for production and detailed in vitro analysis of fl-FGE, which hopefully will lead to a more complete understanding of the structure-function relationship of FGE. It should further allow studying the cell biology of FGE in more detail to address the questions and concepts put forward in this study. One of the concepts to be tested involves uptake of recombinant fl-FGE by recipient cells, which if true, might be developed into strategies for enzyme-replacement therapy in MSD patients.

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