Retention in the Golgi apparatus and expression on the cell surface of Cfr/Esl-1/Glg-1/MG-160 are regulated by two distinct mechanisms

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Cfr (cysteine-rich fibroblast growth factor receptor) is an Fgf (fibroblast growth factor)-binding protein without a tyrosine kinase. We have shown previously that Cfr is involved in Fgf18 signalling via Fgf receptor 3c. However, as Cfr is also known as Glg (Golgi apparatus protein)-1 or MG-160 and occurs in the Golgi apparatus, it remains unknown how the distribution of Cfr is regulated. In the present study, we performed a mutagenic analysis of Cfr to show that two distinct regions contribute to its distribution and stability. First, the C-terminal region retains Cfr in the Golgi apparatus. Secondly, the Cfr repeats in the extracellular juxtamembrane region destabilizes Cfr passed through the Golgi apparatus. This destabilization does not depend on the cleavage and secretion of the extracellular domain of Cfr. Furthermore, we found that Cfr with a GPI (glycosylphosphatidylinositol) anchor was predominantly expressed on the cell surface in Ba/F3 cells and affected Fgf18 signalling in a similar manner to the full-length Cfr, indicating that the interaction of Cfr with Fgfs on the cell surface is important for its function in Fgf signalling. These results suggest that the expression of Cfr in the Golgi apparatus and on the plasma membrane is finely tuned through two distinct mechanisms for exhibiting different functions.

Key words: cysteine-rich fibroblast growth factor receptor (Cfr), E-selectin ligand (Esl), fibroblast growth factor (Fgf), Golgi apparatus, mutagenesis.

INTRODUCTION

Cfr (cysteine-rich fibroblast growth factor receptor) was identified originally as a transmembrane protein with affinity for Fgf (fibroblast growth factor) 1 and Fgf2 by biochemical screening [1]. It binds to Fgfs via a large extracellular domain comprising 16 repeats of a unique motif called the Cfr repeat, but has a short intracellular domain consisting of only 13 amino acid residues [2]. Cfr has no homology with known Fgf-binding molecules including Fgfrs (Fgf receptors) with a tyrosine kinase, and its physiological function had remained unknown. Recently, we generated Cfr-deficient mice, and found that they died in the perinatal period and show growth retardation and skeletal phenotypes. Because these phenotypes were similar to those of Fgf18-deficient mice, we examined the interaction between Cfr and Fgf18 biochemically and genetically, and demonstrated that Cfr binds to Fgf18 to positively regulate Fgf18 signalling through Fgfr3c [3].

However, Cfr is also known as Glg (Golgi apparatus protein)-1 or MG-160, and accumulating evidence indicates that it occurs predominantly in the Golgi apparatus in various cells [4–6]. Furthermore, Cfr has been shown to bind at least two other molecules, E-selectin and TGFβ (transforming growth factor β), and is also known as Esl (E-selectin ligand)-1 and Ltbp (latent TGFβ-binding protein)-1 [7,8]. It was reported recently that Cfr binds to the TGFβ precursor in the Golgi apparatus to modify its maturation and secretion, regulating TGFβ function [9]. Because Cfr acts as an Fgf-binding protein, a ligand of E-selectin and a TGFβ-binding protein, its intracellular position is critical for its function. To interact with extracellular cytokines Cfr must be on the cell surface, but it can also interact with cytokines produced in the cell in the Golgi apparatus as reported for TGFβ. To bind E-selectin, Cfr must be on the cell surface. Thus Cfr is a unique multifunctional molecule and its intracellular distribution must be regulated.

Previous studies have suggested that the C-terminal cytoplasmic tail is important for the intracellular distribution of Cfr. Truncation of the cytoplasmic tail caused partial translocation of rat Cfr on to the cell surface in CHO (Chinese hamster ovary) cells [10]. Moreover, there exists an alternatively spliced variant of Cfr with an extension of 24 amino acid residues to the cytoplasmic tail in primates and humans. Human Cfr with or without the extension is located in the Golgi apparatus and on the cell surface respectively [11]. However, the involvement of other regions of Cfr in the regulation of its intracellular localization has not been examined.

To address this issue, we first confirmed that the cytoplasmic tail is important for the expression of Cfr in the Golgi apparatus, but noticed that without the tail the protein was extremely unstable. A GPI (glycosylphosphatidylinositol)-anchored Cfr mutant was directed to the cell surface and enhanced Fgf18 signalling via Fgfr3c, suggesting that Cfr positively regulates Fgf signalling on the cell surface rather than in the Golgi apparatus. However, it was also unstable and deletion mutants revealed that the Cfr repeats in the juxtamembrane domain contribute to the instability. Although it is known that Cfr is proteolytically cleaved at the juxtamembrane region and its extracellular domain is secreted [8,12], this cleavage is not involved in the instability. Furthermore, we demonstrated that the insertion of the Cfr repeats into the juxtamembrane region of EpCAM (epithelial cell-adhesion molecule) destabilizes Cfr passed through the Golgi apparatus. This destabilization does not depend on the cleavage and secretion of the extracellular domain of Cfr. Furthermore, we found that Cfr with a GPI anchor was predominantly expressed on the cell surface in Ba/F3 cells and affected Fgf18 signalling in a similar manner to the full-length Cfr, indicating that the interaction of Cfr with Fgfs on the cell surface is important for its function in Fgf signalling. These results suggest that the expression of Cfr in the Golgi apparatus and on the plasma membrane is finely tuned through two distinct mechanisms for exhibiting different functions.

Key words: cysteine-rich fibroblast growth factor receptor (Cfr), E-selectin ligand (Esl), fibroblast growth factor (Fgf), Golgi apparatus, mutagenesis.
molecule) also destabilized it, suggesting that the repeats are a module for protein destabilization. These results indicate that the intracellular distribution of Cfr is regulated by two distinct mechanisms. First, the transmembrane region and cytoplasmic tail retain Cfr in the Golgi apparatus, and secondly the extracellular juxtamembrane Cfr repeats destabilize Cfr passed through the Golgi apparatus, allowing Cfr to exhibit its unique multiple functions.

MATERIALS AND METHODS

cDNAs

In all of the mutants of Cfr used in the present study, the signal sequence of mouse Cfr (amino acids 1–27) was replaced by the signal sequence of mouse CD2 and a FLAG tag. FLAG–CfrΔ1169, corresponding to the reported rat Δ1165 mutant [10], was generated by deleting the C-terminal six amino acid residues (RELKD). In all of the GPl-anchored mutants, the extracellular domain of Cfr (amino acids 1–1141) was fused to the signal sequence for the GPl-anchor of human CD58 [also known as LFA (leucocyte function-associated antigen) 3, amino acids 204–237] [13]. In FLAG–Cfr–KARA–GPI, KR (amino acids 1096–1097) and KKR (amino acids 1105–1107) were replaced with AA and AAA respectively. In FLAG–CfrΔ13–16–GPI and FLAG–CfrΔ13–16, Cfr repeats 13–16 (amino acids 870–1141) were deleted. Similarly, in FLAG–CfrΔ15–16–GPI and FLAG–CfrΔ16–GPI, repeats 15 and 16 (amino acids 995–1141) and repeat 16 (amino acids 1057–1141) were deleted. In EpCAM–CfrRepeat, Cfr repeats 13–16 were inserted between the C-terminus of the extracellular domain and the N-terminus of the transmembrane region of mouse EpCAM (between amino acids 266 and 267).

Antibodies

The antibodies used were as follows: anti-FLAG M2 antibody (F3165) from Sigma, anti-actin antibody (sc-1616) from Santa Cruz Biotechnology, anti-TGOLN2 (F3165) from Sigma, anti-actin antibody (sc-1616) from Santa Cruz Biotechnology, anti-actin antibody (sc-1616) from Santa Cruz Biotechnology, anti-actin antibody (sc-1616) from Santa Cruz Biotechnology, anti-actin antibody (sc-1616) from Santa Cruz Biotechnology, and anti-EpCAM antibody (for immunofluorescent staining). The antibodies used were as follows: anti-FLAG M2 antibody (F3165) from Sigma, anti-actin antibody (sc-1616) from Santa Cruz Biotechnology, anti-actin antibody (sc-1616) from Santa Cruz Biotechnology, anti-actin antibody (sc-1616) from Santa Cruz Biotechnology, and anti-EpCAM antibody (for immunofluorescent staining).

Inhibition of lysosomes and proteasomes

Ba/F3 cells were incubated in medium containing 10 mM NH₄Cl (Wako) as a lysosomal enzyme inhibitor, or 10 mM MG-132 (Sigma) as a proteasome inhibitor at 37 °C for 12 h. In parallel with these treatments, Ba/F3 cells were also incubated in medium with PBS and DMSO as a control. The cells were then either harvested for Western blot analysis or analysed by flow cytometry for cell-surface expression of the FLAG-tagged Cfr proteins.

Retroviral infection and proliferation of Ba/F3 cells

Ba/F3 cells and NIH 3T3 cells were infected with retroviruses containing cDNA for a Cfr mutant with an IRES (internal ribosomal entry site)–EGFP (enhanced green fluorescent protein) sequence. The methods used for the proliferation assay have been described previously [3]. Briefly, we sorted EGFP-positive Ba/F3 cells by FACSVantage (BD), and performed the assay in 96-well plates with WST-1 (water-soluble tetrazolium salt 1; Roche Applied Science). GST (glutathione transferase) and GST–Fgf18 used in the present study were described previously [3].

Protein stability assay

Ba/F3 cells were incubated with 50 μg/ml cycloheximide for 4 % paraformaldehyde in PBS at room temperature (25 °C) for 10 min. The cells were incubated with primary antibodies in 5 % skimmed milk in PBS at 4 °C overnight. They were then incubated with secondary antibodies in 5 % skimmed milk in PBS at room temperature for 2 h, and embedded in GEL/MOUNT (Biomega) containing Hoechst 33342. Signals were observed under a confocal microscope.

Biotin-labelling assay

Ba/F3 cells were washed three times with ice-cold PBS with 0.7 mM CaCl₂ and 0.5 mM MgCl₂. Biotinylation of cell-surface proteins was performed with 500 μl of 300 μg/ml EZ-Link-Sulfo-NHS-SS-biotin (Pierce) in PBS for 30 min at 4 °C. Unbound biotin was removed by three washes with ice-cold PBS. Biotinylated cells were incubated with 500 μl of lysis buffer [50 mM Tris/HCl (pH 7.5), 150 mM NaCl, 1 % Nonidet P40, 1 mM EDTA, 1 μg/ml leupeptin (Roche Applied Science), 1 mM PMSF (Wako) and 500 μg/ml Pefabloc (Roche Applied Science)] for 1 h at 4 °C. Immuno-Pure immobilized streptavidin (Pierce) was then added to biotinylated cell lysate and incubated for 2 h at 4 °C to pull down the biotin-labelled proteins. After washing three times with the lysis buffer, the resins were boiled and the pulled down proteins were subjected to Western blot analysis.

Western and Northern blot analyses

The methods used for the Western and Northern blot analyses have been described previously [15].

RESULTS

Intracellular distribution of Cfr in Ba/F3 cells

We reported previously that forced expression of Cfr enhanced Fgf18 signalling via Fgfr3c in Ba/F3 cells [3]. Cfr, however, has been known as a protein of the Golgi apparatus, so we investigated the intracellular distribution of Cfr by expressing the protein with a FLAG tag at its N-terminus (FLAG–Cfr) in Ba/F3 cells (Figure 1A). Flow cytometric analysis with an anti-FLAG antibody showed that FLAG–Cfr was significantly expressed on the cell surface (Figure 1B). By contrast, immunofluorescent microscopic observation with the anti-FLAG antibody revealed that FLAG–Cfr occurred predominantly in the Golgi apparatus, whereas its expression on the cell surface was barely detectable (Figure 1C).

Cfr lacking the cytoplasmic tail enhances Fgf18 signalling on the cell surface

Because FLAG–Cfr was located both on the cell surface and in the Golgi apparatus, it was not clear where Cfr functions in Fgf18 signalling. Therefore we attempted to restrict Cfr to the...
Intracellular distribution of Cfr

Figure 1 Intracellular distribution of the C-terminally modified Cfr mutants in Ba/F3 cells

(A) Schematic representation of the Cfr mutants. All of the mutant proteins have a FLAG tag at their N-terminus. The GPI-anchored mutants have a GPI signal sequence by which they are recruited to the cell surface via GPI. Deleted regions and point-mutated sites are also indicated. SS, signal sequence; TM, transmembrane domain. (B) Expression of FLAG–Cfr, FLAG–CfrΔ1169 and FLAG–Cfr–GPI on the Ba/F3 cell surface monitored by flow cytometry. Ba/F3 cells with only EGFP were used as a control. Note that FLAG–Cfr was significantly expressed on the cell surface compared with the control cells, and the C-terminal modification enhanced the surface expression. The filled plot is the isotype control and open plot is the anti-FLAG. (C) Immunofluorescent staining of FLAG (red) in Ba/F3 cells. The Golgi apparatus was also visualized by staining of TGOLN2 (Tgn46, green). In the merged photographs, signals of FLAG, TGOLN2 and nuclei (blue) are merged. Scale bar, 20 μm.

We then established Ba/F3 cells expressing Fgfr3c with either FLAG–Cfr or FLAG–Cfr–GPI to determine their response to Fgf18. Consistent with our previous report, FLAG–Cfr enhanced Fgf18 signalling via Fgfr3c in Ba/F3 cells (Figure 2). Furthermore, Ba/F3 cells with FLAG–Cfr–GPI, like those with FLAG–Cfr, more strongly responded to Fgf18 than control cells (Figure 2). These results suggest that the positive effect of Cfr on Fgf signalling is determined by its level of expression on the cell surface, not in the Golgi apparatus. Furthermore, the cytoplasmic tail is unnecessary for the function of Cfr in Fgf signalling.

Figure 2 Effects of FLAG–Cfr–GPI on Fgf18 signalling

Ba/F3 cells expressing both Fgfr3c and FLAG–Cfr or FLAG–Cfr–GPI, as well as those with Fgfr3c and FLAG–Cfr were stimulated with 500 ng/ml GST (glutathione transferase)–Fgf18. The same amount of GST was added as a control. The extent of proliferation was monitored by WST-1 assay, and absorbance at 450 nm minus basal absorbance at 650 nm ± S.D. is shown. We prepared eight wells for each condition. FLAG–Cfr–GPI enhanced Fgf18 signalling like FLAG–Cfr. P value was calculated with Student’s t test: * P < 0.05; ** P < 0.01; NS, P > 0.2.

Instability of the C-terminally modified Cfr mutants

As Cfr expressed on the cell surface enhances Fgf18 signalling, the regulation of its transport from the Golgi apparatus to the cell surface is important for its functions. Therefore we investigated further the regulatory mechanisms governing the intracellular distribution of Cfr.

We noticed that the cell-surface expression of the C-terminally modified mutants FLAG–CfrΔ1169 and FLAG–Cfr–GPI was rather weak and that it was under the detection limit of the biotin-labelling assay (Figure 1B and results not shown). Therefore we checked mRNA and protein levels and found that the amount of protein was greatly reduced for FLAG–CfrΔ1169 and FLAG–Cfr–GPI compared with FLAG–Cfr, whereas the amount of mRNA was comparable with that for FLAG–Cfr (Figure 3A). The results indicate that these mutant proteins are unstable, raising the possibility that lysosomal or proteasomal degradation may occur. To address this possibility, a lysosomal enzyme inhibitor NH4Cl or a proteasome inhibitor MG-132 were added to the cells. However, neither inhibitor rescued the total amount of FLAG–Cfr–GPI protein (Figure 3B). Moreover, these inhibitors failed to recover the cell-surface expression of FLAG–Cfr–GPI (Figure 3C). The results suggest that these Cfr mutants were destabilized by another mechanism.

Shedding of Cfr is not involved in the instability of the Cfr mutants

Because previous reports have shown that the extracellular domain of Cfr is cleaved in the juxtamembrane region to be secreted into the culture supernatant [8,12], we considered the possibility that the instability of Cfr is caused by shedding rather than degradation through the lysosome or proteasome. Therefore...
we investigated the amount of the shed extracellular domain of Cfr in the culture supernatants of Ba/F3 cells expressing FLAG–Cfr or FLAG–Cfr–GPI. We immunoprecipitated the extracellular domain from the culture supernatants by the anti-FLAG antibody. As shown in Figure 4(A), the amounts of proteins precipitated were comparable between FLAG–Cfr and FLAG–Cfr–GPI, whereas the amount of FLAG–Cfr–GPI in the cell lysate was extremely low compared with that of FLAG–Cfr, indicating that the shedding of the extracellular domain was not a main cause of the instability. As there are two potential basic proteolytic cleavage sites in the juxtamembrane region of Cfr [8], we investigated further the involvement of shedding in the protein instability by introducing point mutations into FLAG–Cfr–GPI to replace the basic sites KR and KKR with an alanine residue (FLAG–Cfr–KARA–GPI) (Figure 1A). We expressed the mutant in Ba/F3 cells and monitored its expression. Flow cytometric analysis showed that there was no difference in cell-surface expression between FLAG–Cfr–GPI and FLAG–Cfr–KARA–GPI (Figure 4B). Although Northern blot analysis showed that the mRNA levels of FLAG–Cfr–KARA–GPI and FLAG–Cfr–GPI were comparable, and greater than that of FLAG–Cfr, the protein levels were not increased by the mutation (Figure 4C). These results clearly indicate that proteolytic cleavage in the juxtamembrane region is not involved in the instability of the Cfr mutants with the C-terminal modification.

Cfr repeats in the juxtamembrane region destabilize Cfr

Because the basic amino acid sites in the juxtamembrane region do not appear to destabilize Cfr, we examined a possibility that the stability of Cfr is regulated by the Cfr repeats. For this purpose we generated deletion mutants. As shown in Figure 1(A), the extracellular domain of Cfr is composed of 16 repeats of a unique motif, the Cfr repeat. First, we deleted repeats 13–16 from FLAG–Cfr–GPI (FLAG–CfrΔ13–16–GPI) (Figure 1A), and expressed the mutant in Ba/F3 cells. Surprisingly, FLAG–CfrΔ13–16–GPI was expressed more strongly on the cell surface than FLAG–Cfr–GPI, and FLAG–CfrΔ13–16–GPI was clearly detected by Western blotting unlike FLAG–Cfr–GPI (Figures 4A, 5A and 5B). Therefore the deleted juxtamembrane domain is critical for the stability of Cfr. To narrow down the region, we generated two additional mutants FLAG–CfrΔ15–16–GPI and FLAG–CfrΔ16–GPI in which the last two repeats and last repeat were deleted from FLAG–Cfr–GPI respectively (Figure 1A), and expressed them in Ba/F3 cells. Unexpectedly, FLAG–CfrΔ15–16–GPI and FLAG–CfrΔ16–GPI were expressed on the cell surface at levels in between those of FLAG–Cfr–GPI and FLAG–CfrΔ13–16–GPI (Figure 5A). Overlaying the flow cytometric profiles clearly showed that the GPI-anchored mutants with fewer Cfr repeats exhibited stronger expression on the cell surface (Figure 5A). We also detected FLAG–CfrΔ15–16–GPI and FLAG–CfrΔ16–GPI by Western blotting, indicating that these mutants were more stable than FLAG–Cfr–GPI (Figure 5B).

Furthermore, immunofluorescent staining revealed increased cell-surface expression of FLAG–CfrΔ13–16–GPI compared with FLAG–Cfr–GPI (Figure 5C). On the basis of these results, we concluded that without the C-terminal domain, more Cfr is recruited to the cell surface from the Golgi apparatus, but the juxtamembrane Cfr repeats destabilize the Cfr protein, reducing its cell-surface expression.

Cfr protein is unstable outside the Golgi apparatus

To investigate the instability of Cfr protein more directly, we determined the half-lives of the Cfr mutants. For this purpose, we added cycloheximide to Ba/F3 cells expressing the Cfr mutants and monitored their amounts by Western blot analysis. The full-length Cfr was stable and the protein level was not changed significantly, even after 5 h in this assay (Figure 6A). By contrast, the protein level of FLAG–Cfr–GPI was too low to determine the half-life in this assay (Figures 3A and 3B, and results not shown). Therefore we determined the half-lives of the Cfr mutants with intermediate stability, FLAG–CfrΔ13–16–GPI and FLAG–CfrΔ15–16–GPI. In contrast with the full-length Cfr, these mutants were unstable and their amounts declined by the addition of cycloheximide (Figure 6A). On the basis of three independent experiments, the half-lives of FLAG–CfrΔ13–16–GPI and FLAG–CfrΔ15–16–GPI were estimated to be 102.6 and 92.7 min respectively (Figure 6B). These results clearly show that Cfr is stable in the Golgi apparatus, whereas it is highly unstable outside the Golgi apparatus.

Retention of Cfr in the Golgi apparatus through its C-terminus prior to destabilization by the Cfr repeats

We then deleted the Cfr repeats 13–16 from FLAG–Cfr (FLAG–CfrΔ13–16) (Figure 1A), and expressed the mutant in Ba/F3 cells. The level of FLAG–CfrΔ13–16 on the cell surface was comparable with that of FLAG–Cfr in the flow cytometric analysis (Figure 7A). Western blot analysis revealed that the amount of protein was also comparable with that of FLAG–Cfr, and robustly increased compared with that of FLAG–CfrΔ13–16–GPI (Figure 7B). Moreover, we observed that FLAG–CfrΔ13–16 was mostly present in the Golgi apparatus like FLAG–Cfr,
Figure 4  Expression of the Cfr with mutations in the juxtamembrane basic sites

(A) Secretion of the extracellular domain of Cfr into the culture supernatants of Ba/F3 cells. The extracellular domain was immunoprecipitated from the culture supernatants of Ba/F3 cells expressing FLAG–Cfr or FLAG–Cfr–GPI, and then subjected to Western blot analysis (IP). Expression of Cfr in Ba/F3 cells and the total amount of the cell lysate loaded was monitored by blotting with anti-Cfr serum and anti-actin antibody respectively (Lysate). The amount of the extracellular domain secreted was comparable between FLAG–Cfr and FLAG–Cfr–GPI. (B) Cell-surface expression of FLAG–Cfr–KARA–GPI. Filled, isotype control; open, anti-FLAG. No difference in the expression level was observed between FLAG–Cfr–KARA–GPI and FLAG–Cfr–GPI. (C) The total amount of FLAG–Cfr–KARA–GPI protein and mRNA in Ba/F3 cells. Mutations in the two basic sites did not alter the amount of total protein expression from FLAG–Cfr–GPI.

Figure 5  Expression of the mutants deleted of the Cfr repeats in Ba/F3 cells

(A) Flow cytometric analysis of the GPI-anchored deletion mutants. Cell-surface expression of the mutants was monitored using the FLAG tag. For each sample: filled, isotype control; open, anti-FLAG. For the panel showing GPI-anchored deletion mutants: filled, isotype control of FLAG–Cfr–GPI; overlaid, anti-FLAG of FLAG–Cfr–GPI (black), FLAG–CfrΔ15–16–GPI (red), FLAG–CfrΔ13–16–GPI (blue) and FLAG–Cfr–GPI (green). Cell-surface expression of the GPI-anchored Cfr mutants increased as the number of Cfr repeats decreased. (B) Western and Northern blot analyses of total protein and mRNA of the deletion mutants. The mutants were clearly detected in the Western blot analysis. (C) Immunofluorescent staining of Ba/F3 cells expressing the deletion mutants. In the merged photographs, signals of FLAG (red), TGOLN2 (Tgn46, green) and nuclei (blue) are merged. Scale bar, 20 μm. The FLAG signals on the cell surface were slightly stronger in FLAG–CfrΔ13–16–GPI than FLAG–Cfr–GPI.

Figure 6  Protein stability assay of the Cfr mutants

(A) Ba/F3 cells expressing the Cfr mutants were incubated with 50 μg/ml cycloheximide for the indicated periods, and then harvested for Western blot analysis. As a control the solvent DMSO was added. The GPI-anchored mutants showed decreased stability. By contrast, FLAG–Cfr and FLAG–CfrΔ13–16 was stable and showed no decline in the amount of protein. (B) The half-life curves for the GPI-anchored mutants. The results were derived from three independent experiments. The half-lives of FLAG–CfrΔ13–16–GPI and FLAG–CfrΔ15–16–GPI are 102.6 and 92.7 min respectively.
The intracellular distribution and protein stability of the Cfr mutants were monitored by flow cytometry and Western analysis. The results are summarized in Table 1.

Table 1 Summary of intracellular distribution and protein stability of the Cfr mutants

<table>
<thead>
<tr>
<th>Cfr mutant</th>
<th>Intracellular distribution</th>
<th>Protein stability</th>
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<tbody>
<tr>
<td>FLAG-Cfr</td>
<td>Golgi and weakly cell surface</td>
<td>Very stable</td>
</tr>
<tr>
<td>FLAG-CfrΔ1169</td>
<td>Cell surface</td>
<td>Unstable</td>
</tr>
<tr>
<td>FLAG-Cfr-GPI</td>
<td>Cell surface</td>
<td>Unstable</td>
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<tr>
<td>FLAG-CfrΔ13–16–GPI</td>
<td>Cell surface (&gt;FLAG–Cfr-GPI)</td>
<td>Stable (&gt;FLAG–Cfr-GPI)</td>
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<tr>
<td>FLAG-CfrΔ15–16–GPI</td>
<td>Cell surface (&gt;FLAG–Cfr-GPI)</td>
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<tr>
<td>FLAG-CfrΔ16–GPI</td>
<td>Cell surface (&gt;FLAG–Cfr-GPI)</td>
<td>Stable (&gt;FLAG–Cfr-GPI)</td>
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<tr>
<td>FLAG-Cfr-KARA–GPI</td>
<td>Cell surface (= FLAG–Cfr-GPI)</td>
<td>Unstable (= FLAG–Cfr-GPI)</td>
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<tr>
<td>FLAG-CfrΔ16</td>
<td>Golgi and weakly cell surface</td>
<td>Very stable</td>
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indicating that destabilization of Cfr by the Cfr repeats does not occur in the Golgi apparatus (Figure 7C). This possibility was further supported by the stability of FLAG–CfrΔ13–16, which was comparable with that of the full-length Cfr (Figure 6A).

Regulatory mechanisms for the intracellular distribution of Cfr in other cells and proteins

Finally, we addressed whether the regulatory mechanisms for the intracellular distribution of Cfr are general. First, we expressed the Cfr mutants in NIH 3T3 cells to confirm our results in Ba/F3 cells, and found that the mutants were expressed in NIH 3T3 cells with patterns basically the same as those in Ba/F3 cells (Figure 8A). In addition, we also observed that the Cfr mutants were expressed with the same patterns in COS7 cells (results not shown). We have summarized the intracellular distribution and protein stability of the Cfr mutants in Table 1.

Next, we investigated whether the mechanisms work in other proteins. It has been already shown that fusion of the C-terminal region of Cfr to the extracellular domain of CD8 directs the fusion protein to the Golgi apparatus [11]. Therefore we generated a fusion protein of another cell-surface protein EpCAM and the Cfr repeats (EpCAM-CfrRepeat; Figure 8B). We expressed this fusion protein in Ba/F3 cells in parallel with wild-type EpCAM, and monitored their expression on the cell surface by flow cytometry and a biotin-labelling assay. Interestingly, the insertion of the Cfr repeats decreased the surface expression of EpCAM (Figures 8C and 8D). We then monitored mRNA and protein expression of EpCAM–CfrRepeat and found that whereas the amount of mRNA was comparable between EpCAM and EpCAM–CfrRepeat, the amount of EpCAM–CfrRepeat protein was extremely low (Figure 8E). We monitored further the intracellular distribution of EpCAM–CfrRepeat by immunofluorescent staining. EpCAM was expressed strongly on the cell surface, but the strength of the EpCAM–CfrRepeat signal was lower than that of EpCAM (Figure 8F).

These results clearly show that Cfr is at first retained in the Golgi apparatus through its transmembrane domain and cytoplasmic tail, and then subjected to stability control through the Cfr repeats during its recruitment to the plasma membrane or on the cell surface (Figure 8G).

DISCUSSION

Cfr is unique in that it has been identified independently as an Fgfbinding protein, a latent TGFβ-binding protein, a ligand for E-selectin and a protein of the Golgi apparatus [1,4,7,8]. However, its physiological functions had been unknown. Recently, we [3] and Yang et al. [9] independently generated Cfr-deficient mice and reported that they show skeletal phenotypes. We reported that Cfr is a positive regulator of the Fgf18 signalling pathway, and revealed in the present study that the interaction between Cfr and Fgf18 on the cell surface is important for Cfr’s actions (Figure 2), whereas Yang et al. [9] indicated that it also contributes to TGFβ signalling by binding the precursor of TGFβ in the Golgi apparatus. Furthermore, the function of Cfr as a ligand for E-selectin has been well confirmed by knockdown experiments, although no obvious phenotype in cell adhesion via selectins, such as lymphocyte rolling, has been observed in Cfr-deficient mice [16]. Cfr is ubiquitously expressed in almost all cells and tissues from very early in embryonic development to adulthood [3]. Therefore, considering the pleiotropic roles of Fgf and TGFβ signalling in development, the phenotypes of Cfr-deficient mice would be much more severe if Cfr functions the same way in all cells. Rather, Cfr seems to perform different roles according to its cellular context. We speculate that a key determinant is its intracellular distribution. When located in the Golgi apparatus,
Cfr can act as a regulator of the TGFβ signalling pathway, but it needs to be on the cell surface to function as a regulator of the Fgf signalling pathway or a ligand for E-selectin. We also observed that wild-type Cfr expressed in Ba/F3 cells and NIH 3T3 cells occurred not only in the Golgi apparatus, but also on the cell surface (Figures 1B and 8A), and it has been reported that endogenous Cfr is also expressed on the cell surface in 32Dcl3 cells and neutrophils [17]. Therefore we investigated the mechanisms regulating the intracellular distribution of Cfr.

Previous studies have shown that the cytoplasmic tail is important for the localization of Cfr to the Golgi apparatus in rats and humans [10,11]. We also confirmed that it is required for the retention of Cfr in the Golgi apparatus in mice (Figures 1B and 1C). In addition, we revealed that replacement of the transmembrane and cytoplasmic domains with the GPI anchor sequence further enhanced the cell-surface expression (Figures 1B and 1C). From these results, not only the cytoplasmic tail but also the transmembrane domain could contribute to the retention of Cfr in the Golgi apparatus, like many other transmembrane proteins in the Golgi apparatus [18]. Moreover, these C-terminally modified mutants were extremely unstable compared with full-length Cfr, indicating that having left the
Golgi apparatus Cfr is destabilized (Figure 3A). This possibility was further directly confirmed by the protein stability assay. The Cfr mutants that were no longer localized in the Golgi apparatus exhibited much shorter half-lives compared with full-length Cfr (Figure 6). Unexpectedly, this destabilization is not due to the lysosome or proteasome (Figures 3B and 3C). Therefore we searched by mutagenic analysis for regions responsible for the instability.

It has been reported that the extracellular domain of Cfr is cleaved and secreted into the culture supernatant. The molecular mass of the secreted Cfr fragment almost corresponded with that of the extracellular domain, indicating that the cleavage occurs very near the transmembrane domain of Cfr [8,12]. Indeed, we also observed the shed extracellular domain of Cfr in the culture supernatant, but its amount was comparable between FLAG–Cfr and FLAG–Cfr–GPI. These results indicate that the shedding is not a main cause of the instability of Cfr outside the Golgi apparatus. As it was suggested previously that basic amino acid residues in the juxtamembrane region are the targets of the cleavage [8], we examined further the possibility that the cleavage contributes to the instability by making substitutions with an alanine residue. However, the expression of FLAG–Cfr–GPI protein was not affected by the mutations (Figures 4B and 4C). Instead, we found that deletion of the last four Cfr repeats dramatically enhanced the cell-surface expression of FLAG–Cfr–GPI (Figure 5). To find the specific site for the instability, we deleted the last repeat or two repeats from FLAG–Cfr–GPI and found that the surface expression of these GPI-anchored mutants reduced gradually depending on the number of Cfr repeats (Figure 5A). These results suggest that the Cfr repeats rather than a specific sequence for cleavage in the juxtamembrane region are responsible for the instability of Cfr. Interestingly, the repeats also reduced the amount of EpCAM protein when they were inserted in its juxtamembrane region (Figure 8). These results also support that the Cfr repeats function as a general module for protein destabilization.

To examine the function of the Cfr repeat further, we deleted the last four repeats from the full-length Cfr (Figure 1A). However, FLAG–CfrΔ13–16 was still found in the Golgi apparatus, indicating that the presence of the transmembrane domain and cytoplasmic tail is sufficient for the retention of Cfr in the Golgi apparatus (Figures 7A and 7C). Moreover, the stability control via the Cfr repeats does not take place in the Golgi apparatus because the stability control via the Cfr repeats also reduced the amount of EpCAM protein when they were inserted in its juxtamembrane region (Figure 8). These results clearly demonstrate that Cfr is destabilized after passing through the Golgi apparatus.

On the basis of our findings, we propose a two-step model for regulation of the intracellular distribution of Cfr (Figure 8G). First, Cfr is retained via its transmembrane domain and cytoplasmic tail in the Golgi apparatus in a manner similar to that of many other transmembrane proteins. It has been accepted that transmembrane proteins located in the Golgi apparatus have a relatively short transmembrane domain (~15 amino acids) compared with that of the plasma membrane proteins (~20 amino acids). This short transmembrane domain fits the properties of lipids constituting the membrane of the Golgi apparatus and retains proteins with this domain in the Golgi apparatus [19]. However, the transmembrane domain of Cfr is composed of 21 amino acids, indicating that the localization of Cfr in the Golgi apparatus is independent of the length of the transmembrane domain. Furthermore, the tyrosine-based motif in the cytoplasmic domain functions as a retrieval signal from the plasma membrane to the Golgi apparatus in some transmembrane proteins located in the Golgi apparatus such as TGOLN2 [20–22]. However, Cfr has no tyrosine residues in its cytoplasmic tail, indicating that Cfr occurs in the Golgi apparatus by other mechanisms. It has been reported that the transmembrane domains and cytoplasmic regions interact with each other to form a large complex, preventing transport to other destinations (the ‘kin recognition’ mechanism) [18,23]. This is a possible mechanism regulating the distribution of Cfr. If Cfr is also retained in the Golgi apparatus by ‘kin recognition’, modification of the transmembrane region and cytoplasmic tail should alter the intracellular localization of Cfr. Secondly, after leaving the Golgi apparatus, Cfr is destabilized via its Cfr repeats in the juxtamembrane region, resulting in reduced cell-surface expression. This step seems to be specific to Cfr, and we speculate that there may be a factor recognizing the Cfr repeats to destabilize Cfr. The fact that EpCAM–CfrRepeat is also further destabilized supports the existence of a trans-acting factor on the Cfr repeats (Figure 8). Therefore expression of this factor, or modification of the recognized sites in the Cfr repeat should affect the localization of Cfr. Alternatively, a portion of Cfr may be transported to the surface by simply escaping from this two-step regulation when expressed abundantly. Thus the intracellular distribution of Cfr can be modified at multiple points to direct it into the Golgi apparatus, or on to the cell surface, determining its functions.

**AUTHOR CONTRIBUTION**

Yuichiro Miyaoka, Hidenori Kato and Kazuki Ebato performed the experiments and analysed the data. Shigeru Saito and Naoko Miyata performed the cell sorting. Toru Imamura contributed the Ba/F3 cell line expressing Fgfr3c and provided advice on experimental design. Yuichiro Miyaoka designed the research project. Yuichiro Miyaoka and Atsushi Miyajima wrote the paper.

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