A natural inactivating mutant of human glucagon receptor exhibits multiple abnormalities in processing and signaling

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Abstract
Background and aim: To elucidate the pathogenetic mechanisms of a mutant P86S glucagon receptor (GCGR) in causing a novel human disease (Mahvash disease).

Material and method: Enhanced green fluorescent protein (EGFP)-tagged WT and P86S GCGR were expressed in HEK 293 or H1299 cells either transiently or stably. Receptor localization and internalization, and cell apoptosis were studied by fluorescence microscopy, and calcium signaling by Rhod-3 labeling. Gene expression was assayed by RT-PCR or Western blot. Cell fate was determined by live cell imaging.

Results: Unlike WT GCGR, P86S was partially localized to the plasma membrane and partially in the cytoplasm as previously reported and did not undergo internalization upon glucagon treatment. P86S did not elicit calcium response after treatment with 1\textsuperscript{\textmu}M glucagon. Cells transiently expressing P86S exhibited more apoptosis than those expressing WT GCGR (18.3\% vs 2.1\%, \(P<0.05\)) but the X-box binding protein 1 mRNA cleavage, a marker of endoplasmic reticulum (ER) stress, was not evident, suggesting that the apoptosis did not result from ER stress. Cells stably expressing P86S did not exhibit apoptosis and a quarter of them harbored a novel inclusion body-like circular structure that was marked by P86S and ER residential proteins. These circular ER bodies were not seen in cells expressing WT GCGR or transiently expressing P86S and were not affected by treatment with proteasome inhibitor or microtubule depolymerizer, suggesting that they do not represent aggresome structures. The circular ER bodies could fuse and split to form new bodies.

Conclusion: The naturally-occurring P86S mutant GCGR exhibits abnormal receptor internalization and calcium mobilization, and causes apoptosis. The novel dynamic circular ER bodies may be adaptive in nature to nullify the toxic effects on P86S. These findings provide further insights into the pathogenetic mechanisms of Mahvash disease.

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Introduction

Naturally-occurring mutations of G protein-coupled receptors are uncommon but cause many human diseases.\(^1\,2\) Inactivating mutations of G protein-coupled receptors result in loss of function of the normal signaling pathway mediated by the receptors. The mechanisms for loss of function by the inactivating mutations can be categorized into 4 classes of defective receptor function: receptor biosynthesis, trafficking to the cell surface, ligand binding, and receptor activation.\(^2\) The glucagon receptor (GCGR) belongs to the secretin receptor family of G protein-coupled receptors.\(^3\)-\(^5\) GCGR is mainly expressed in liver and kidney. GCGR couples to Gs and Gq and increases cAMP and intra-cellular calcium levels, and undergoes glucagon-induced internalization.\(^4\)-\(^9\) We have recently identified a novel human disease of hyperglucagonemia without glucagonoma syndrome, hypoglycemia, pancreatic α cell hyperplasia, and pancreatic neuroendocrine tumors associated with a homozygous inactivating mutation (P86S) of the GCGR (Mahvash disease).\(^10\,11\) The mechanisms for pancreatic α cell hyperplasia and neuroendocrine tumors are not clear but appear to be loss of negative feedback inhibition on α cells by glucagon signaling. The P86S mutation is in the N terminal extracellular domain of GCGR. The homologous proline residue in the receptors of gastric inhibitory polypeptide (GIP) and glucagon-like peptide 1 (GLP-1), both highly homologous to GCGR, is critical for the binding with their respective ligands.\(^12\,13\) The P86S mutant localizes partially to the plasma membrane and partially in the cytoplasm, binds glucagon with much lower affinity, and produces lower cAMP levels at physiological concentrations of glucagon but was able to achieve maximal cAMP levels at 100 nM of glucagon. As the patient’s phenotype is almost identical to that of mice lacking GCGR,\(^14\,15\) it is likely that the P86S mutation causes the patient’s unique condition.

To further characterize the P86S GCGR, we studied its internalization, calcium mobilization, effects on cell survival, and detailed subcellular localization by expressing enhanced green fluorescent protein (EGFP)-tagged WT or P86S GCGR in HEK 293 or H1299 cells, and we demonstrated that the P86S mutant has multiple abnormalities in receptor function. Moreover, the P86S mutant induces formation of a novel subcellular structure, dynamic circular endoplasmic reticulum bodies, which may be an important mechanism for cellular processing of the mutant GCGR.
Figure 1  Internalization of WT GCGR and P86S. A) GCGR internalization by increasing concentrations of glucagon. Two days after transfection with plasmids encoding WT GCGR-EGFP or P86S-EGFP, HEK 293 cells were treated with glucagon from 1 nM to 1 μM at 37°C for 30 minutes. Cells were fixed and representative cells shown. B) Time course of glucagon-induced GCGR internalization. HEK 293 cells were treated with 1 μM glucagon for 10-60 minutes. Note that untreated WT GCGR was always localized to plasma membrane while P86S was localized to cytoplasm in some cells (shown in A) or to plasma membrane in others (shown in B). P86S did not change localization upon glucagon stimulation. Bar, 10 μm.

EGFP: enhanced green fluorescent protein; GCGR: glucagon receptor.

Materials and methods

Cell Culture, plasmids, transfection, and reagents

HEK 293 (from human embryonic kidney) and H1299 (from human non-small cell lung carcinoma) cells were grown in Dulbecco’s modified Eagle’s medium with 10% fetal bovine serum in a 37°C, humidified incubator with 5% CO2. HEK 293 cells were used in most experiments as they are derived from the kidney which normally expresses GCGR. H1299 cells were only used in calcium measurements because they did not apparently express GCGR intrinsically through preliminary experiments and HEK 293 cells did not load the calcium dye Rhod-3 AM well. Plasmids encoding enhanced green fluorescent protein (EGFP)-tagged human WT and P86S glucagon receptor (GCGR) and the parental plasmid pEGFP-N3 were described before.11 Cells were transfected with Lipofectamine 2000 (Invitrogen, Carlsbad, CA, EE.UU.). Stable HEK 293 cell lines expressing EGFP, WT GCGR-EGFP, or P86S-EGFP were established by incubating cells with 0.6 μg/ml G418 after transfection. Thapsigargin was from Sigma (St Louis, IL, EE.UU.), LLnL from Calbiochem (La Jolla, CA, EE.UU.), and colcemid from Invitrogen.

Calcium analysis

H1299 cells were incubated with 10 μM Rhod-3 AM, PowerLoad, and 2.5 mM probenecid (Molecular Probes, Eugene, OR, EE.UU.) in Hank’s buffered salt solution (HBSS) for 45 minutes, followed by incubation in HBSS with probenecid for another 45 minutes. Loaded cells were placed in a confocal microscope incubator chamber and images were acquired with a TCSP confocal microscope (Leica Microsystems, Heidelberg, Germany) using excitation/emission 550/580 nm. Scanning rate was set to 7 frames per second for 200 seconds.

Apoptosis assay

HEK 293 cells were grown on coverslips for 2-5 days after transfection. Cells were fixed with 4% paraformaldehyde, permeabilized with 0.6% Tween 20, stained with Hoechst 33258 (1:10,000; Molecular Probes), and observed with a
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Figure 2  Calcium signaling by WT GCGR and P86S. H1299 cells were transfected with plasmids encoding WT GCGR-EGFP or P86S-EGFP, loaded with Rhod-3 AM and stimulated with 1 µM glucagon. Shown in upper panels were average Rhod-3 fluorescence levels ± standard error of the mean before and after glucagon stimulation. Shown in lower panels were traces of 2 representative individual cells. P86S did not respond to glucagon with an increase in intracellular calcium.

EGFP: enhanced green fluorescent protein; GCGR: glucagon receptor.

RT PCR and Western blot

Total RNA was extracted with RNeasy kit (Qiagen, Valencia, CA, EE.UU.) and treated with DNase. RNA (2 µg) was reverse-transcribed using SuperScript II Reverse Transcriptase (Invitrogen). Human XBP1 (X-box binding protein 1) was amplified by PCR using 100 ng cDNA and primer set GAAGCCAAGGGGAA TGAAGT (forward) and ACTGGGTC-CAAGTTGTCCAG (reverse) for XBP-1 and primer set for actin B (Real Time Primers, Elkins Park, PA). Western blot of EGFP-tagged WT or P86S GCGR was described previously.

Immunofluorescent staining and fluorescence microscopy

These were performed as described. Briefly, HEK 293 cells expressing WT GCGR-EGFP or P86S-EGFP were fixed with 4% paraformaldehyde, and permeabilized with blocking buffer. Primary antibody, rabbit polyclonal anti-calregulin or anti-glucosidase II, goat polyclonal anti-ubiquitin (Santa Cruz Biotechnology, Santa Cruz, CA, EE.UU.), or mouse monoclonal anti-alpha tubulin (Calbiochem) was added at 1:50 in the blocking buffer and incubated for 2 h. Cells were washed and incubated with appropriate rhodamine-labeled secondary antibody for 30 min. EGFP fluorescence was directly visualized without staining. Immunostained cells were visualized on a TE200 inverted epifluorescence microscope (Nikon, Melville, NY, EE.UU.) equipped with relevant fluorescence filters. For live cell observation, HEK 293 cells stably expressing WT GCGR-EGFP or P86S-EGFP were grown in gridded flasks and the same groups of cells were directly visualized over time with green fluorescence filters.

Results

Receptor internalization

In HEK 293 cells, WT GCGR was predominantly localized on the plasma membrane and underwent glucagon-induced internalization into intracellular vesicles in a glucagon dose-dependent manner (fig. 1A), in agreement with the results from Sachdev et al. Internalization of WT GCGR appeared to be complete after 30 minutes of maximal glucagon stimulation (fig. 1B). P86S was partially localized to the plasma membrane and partially in the cytoplasm as previously reported. Cells with more cytoplasmic P86S localization were shown in Figure 1A while those with more plasma membrane localization were shown in Figure 1B. Regardless of the basal receptor localization, P86S failed to accumulate in intracellular vesicles in response to glucagon stimulation, even at 1 µM (which stimulates maximal cAMP production by...
P86S for 1 hour. Thus, the P86S is defective in receptor

Calcium mobilization

We used red calcium sensor Rhod-3 to study glucagon-stimulated calcium response of EGFP-tagged WT or P86S GCGR. In cells expressing WT GCGR, 1 μM glucagon elicited a clear increase in intracellular calcium levels (fig. 2). Most cells exhibited a monophasic calcium response that largely tapered in 100 seconds after glucagon stimulation. In a few cells, calcium levels began to oscillate after glucagon. In cells expressing P86S, no calcium increase was observed in response to glucagon stimulation. A few cells exhibited spontaneous low-amplitude oscillation of calcium levels which was not changed by glucagon.

Apoptosis

We measured apoptotic rate of HEK 293 cells expressing EGFP, WT GCGR-EGFP, and P86S-EGFP. The apoptotic rates of cells expressing P86S were consistently higher than those of cells expressing EGFP or WT GCGR, starting 2 days after transient transfection (fig. 3A). The difference in apoptotic rates between cells expressing P86S and WT GCGR was largest 4 days after transfection (18.3 ± 0.2% vs 2.1 ± 0.2%, P < 0.05). As P86S was significantly localized to endoplasmic reticulum (ER) (data not shown for cells transiently expressing P86S), we tested cleavage of XBP-1 mRNA, a marker of ER stress. Treatment with thapsigargin, a known inducer of ER stress, clearly showed XBP-1 cleavage, but no such cleavage was seen in cells expressing any of the 3 constructs (fig. 3B), suggesting that P86S-induced apoptosis was caused by mechanisms other than ER stress.

Circular ER bodies

To further examine the effects of P86S expression on cell survival, we established stable HEK 293 cells expressing WT GCGR-EGFP or P86S-EGFP. Apoptotic rates were similar in cells expressing the two constructs: 1.2 ± 0.1% for WT GCGR and 0.9 ± 0.4% for P86S. The expression levels of WT GCGR and P86S were similar (fig. 4). Unlike in cells transiently transfected, P86S was present in cells stably expressing P86S (24.9 ± 1.9%) but not in those expressing WT GCGR (0 ± 0%, P < 0.001) (fig. 4A and B). These novel structures measured 2-3 μm in diameter, and were mostly perinuclear and present in all 7 stable clones of HEK 293 cells expressing P86S-EGFP but rarely seen in cells transiently expressing it. Of 70 cells that contain that novel structure, 52 (74%) had 1, 10 (14%) had 2, 5 (7%) had 3, while 3 (4%) had 4 or more. In WT GCGR-expressing cells, WT GCGR and ER marker calregulin were localized distinctly without overlap (fig. 4A). In P86S-expressing cells, P86S and calregulin and glucosidase II, another ER marker, were almost completely colocalized (fig. 4B a-h). The circular, inclusion body-like structures containing P86S were perfectly colocalized with both ER markers. We hereafter call these novel structures circular ER bodies.

Characterization of circular ER bodies

As mutant G protein-coupled receptors such as mutant rhodopsin can localize to a structure called aggresome, we examined whether the circular ER bodies are aggresomes. Aggresome formation is microtubule-dependent and increases with proteasome inhibition; aggresomes are localized around microtubule organizing centers and contain ubiquitinated proteins. In both interphase (fig. 4B i-l) and metaphase (fig. 4B m-p) cells, the circular ER bodies were distinct from the microtubule organizing centers and they did not colocalize. The circular ER bodies did not colocalize with ubiquitin (fig. 4B q-t). Treatment with proteasome inhibitor LmL or microtubule depolymerizer colcemid or both did not increase the number or size of the circular ER bodies (fig. 4B u-w). Our results thus demonstrated that the circular ER bodies are not aggresomes.

To study the fates of the circular ER bodies and the cells harboring them, we followed 791 live cells over 9-13 hours. None of the cells harboring the circular ER bodies died over the observation period and all cells divided
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Discussion

The P86S mutant is a naturally-occurring, inactive mutant human GCGR which causes a novel human disease (Mahvash disease) when expressed homozygously. The Mahvash disease is a rare genetic disorder characterized by severe hyperglycemia, hypoglycemia, and growth retardation. It is caused by a mutation in the glucagon receptor gene (GCGR) that results in a loss of function. This study further elucidates the cellular and molecular mechanisms underlying the pathogenesis of Mahvash disease by characterizing the endoplasmic reticulum (ER) bodies induced by the P86S mutation. The presence of ER bodies suggests that the mutant receptor is retained in the ER, potentially interfering with cellular function. Further studies are needed to understand the specific molecular mechanisms and to develop therapeutic strategies for this disease.
Our previous studies demonstrate that GCGR increases intracellular calcium in response to cellular stress. It has been suggested that gluconeogenesis is a main function of GCGR. As gluconeogenesis is a main function of GCGR, it appears that besides defective glucagon binding and abnormal receptor localization, failure of P86S internalization may also cause other abnormalities as well. To understand the pathogenesis of the Mahvash disease, we have used transiently or stably expressed EGFP-tagged WT and P86S human GCGR to further study their differences.

The current data clearly demonstrated that unlike the WT GCGR which internalizes upon glucagon binding, the P86S does not internalize even after incubation in very high concentrations of glucagon for an extended period of time. The WT GCGR internalizes with Gsα and GCGR-Gs signal transduction may continue after GCGR internalization at the endo-lysosomal apparatus. It appears that besides defective glucagon binding and abnormal receptor localization, failure of P86S internalization may also cause decreased cAMP production.

It is interesting to note that P86S does not mobilize calcium as we have shown previously that high concentrations of glucagon (1 μM) produce maximal cAMP signals in cells expressing P86S. GCGR increases intracellular calcium levels by releasing calcium from intracellular stores through activating phospholipase C. It has been suggested that that the increase in intracellular calcium stimulates gluconeogenesis by upregulating the Krebs cycle and the electron-transfer chain and by activating gluconeogenic enzymes. As gluconeogenesis is a main function of GCGR in vivo, the absence of calcium signaling by P86S would contribute to decreased neogenesis, explaining in part the hypoglycemia in the patient. How a point mutation in the extracellular domain of GCGR abrogates the coupling between GCGR and Gq is not clear.

Our study also unravels the deleterious effects of transiently expressed P86S on cell survival, adding one more abnormality to P86S. As P86S is significantly localized to the ER and may cause ER stress, we tested cleavage of XBP-1 mRNA, a marker of ER stress, induced by P86S expression. Since ER stress is not detected, other unknown mechanisms should underlie the apoptosis induced by transiently expressed P86S. In contrast to cells transiently expressing P86S, those stably expressing P86S do not exhibit excess apoptosis, consistent with the normal liver function of the patient harboring homozygous P86S mutation. The cells stably expressing P86S must develop mechanisms to alleviate the toxic effects of P86S, one of which may be the novel, dynamic inclusion body-like structures we termed "circular ER bodies".

Our study has clearly demonstrated that the circular ER bodies are not aggresomes (solid cytoplasmic inclusion bodies) because the former did not change with proteasome inhibition or microtubule disruption, exhibited circular configurations, and were localized away from the microtubule organizing centers. Morphologically the circular ER bodies resemble another inclusion body-like structure called "concentric membranous bodies of the ER" formed by mutant CFTR (cystic fibrosis transmembrane conductance regulator) or wildtype D1 and D2 dopamine receptors, both G protein-coupled, in response to cellular stress. The formation of concentric membranous bodies by mutant CFTR and D1 and D2 receptors requires overexpression of calnexin, an ER chaperone that interacts with glycosylated proteins, and intact microbubules. In contrast, the circular ER bodies formed by P86S as shown in this study are independent of calnexin overexpression or microtubules, indicating they are different structures from the concentric membranous bodies of the ER. The functions of the circular ER bodies are still unclear but could be inferred from this study as an adaptive response to P86S expression. The circular ER bodies are only present in the stable clones of cells expressing P86S but not in cells transiently expressing it, suggesting that they are required for the stable P86S expression. The absence of significant apoptosis in cells stably expressing P86S also supports the notion that the circular ER bodies are protective against the P86S toxicity. We do not have access to the liver or kidney of the patient with Mahvash disease, therefore cannot ascertain whether the...
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In summary, we have described further differences between WT GCGR and P86S in glucagon-induced receptor internalization, calcium mobilization, and effects on cell survival. The P86S mutant induces the formation of dynamic circular ER bodies are also present in these organs. The circular ER bodies are unlikely an artifact of overexpression as the WT GCGR, expressed at similar levels, did not form them.

Figure 6  Dynamic circular endoplasmic reticulum (ER) bodies revealed by live cell imaging. Live HEK 293 cells stably expressing P86S-EGFP were observed hourly and the same groups of cells were followed over time. A) Disappearance of circular ER bodies from 2 cells, 1 in 3 hours, the other in 5 hours. B) Emergence of new ER bodies in 2 cells. C) The life cycle of one circular ER body (in the cell in the center of field). Note that an adjacent cell on the left has a dynamic circular EB body that disappeared in 4 hours. D) Fusion and disappearance of 3 ER bodies. Time of image acquisition is indicated at the left low corner. The circular ER bodies were dynamic structures. Bar, 10 μm.

EGFP: enhanced green fluorescent protein.
circular ER bodies, which may be adaptive in nature to nullify the P86S toxicity to cells. These differences provide further insights into the pathogenetic mechanisms of the Mahvash disease.

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Conflict of interest

The authors declare that they have no conflict of interest.

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