Carboxyl-Terminal Receptor Domains Control the Differential Dephosphorylation of Somatostatin Receptors by Protein Phosphatase 1 Isoforms

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Abstract

We have recently identified protein phosphatase 1β (PP1β) as G protein-coupled receptor (GPCR) phosphatase for the sst2 somatostatin receptor using siRNA knockdown screening. By contrast, for the sst5 somatostatin receptor we identified protein phosphatase 1γ (PP1γ) as GPCR phosphatase using the same approach. We have also shown that sst2 and sst5 receptors differ substantially in the temporal dynamics of their dephosphorylation and trafficking patterns. Whereas dephosphorylation and recycling of the sst2 receptor requires extended time periods of ~30 min, dephosphorylation and recycling of the sst5 receptor is completed in less than 10 min. Here, we examined which receptor domains determine the selection of phosphatases for receptor dephosphorylation. We found that generation of tail-swap mutants between sst2 and sst5 was required and sufficient to reverse the patterns of dephosphorylation and trafficking of these two receptors. In fact, siRNA knockdown confirmed that the sst2 receptor carrying the sst5 tail is predominantly dephosphorylated by PP1β, whereas the sst5 receptor carrying the sst2 tail is predominantly dephosphorylated by PP1γ. Thus, the GPCR phosphatase responsible for dephosphorylation of individual somatostatin receptor subtypes is primarily determined by their different carboxyl-terminal receptor domains. This phosphatase specificity has in turn profound consequences for the dephosphorylation dynamics and trafficking patterns of GPCRs.

Introduction

The signaling output of G protein-coupled receptors (GPCRs) is desensitized by mechanisms involving phosphorylation, β-arrestin binding and internalization. GPCR signaling is re-sensitized by mechanisms involving dephosphorylation, but details about the phosphatases responsible are generally lacking. We and others have recently succeeded in identifying bona fide GPCR phosphatases for a number of receptors using a combined approach of phosphosite-specific antibodies and siRNA screening in HEK293 cells. First, we identified protein phosphatase 1β (PP1β) as GPCR phosphatase for the sst2 somatostatin receptor [1]. Second, we identified PP1γ as GPCR phosphatase for the μ-opioid receptor and the sst5 somatostatin receptor [2] [3]. Third, more recently Gehret and Hinkle identified PP1ζ as GPCR phosphatase for the thyrotropin-releasing hormone receptor [4]. All of the above observations were made in a similar cellular background. This suggests that a given GPCR may recruit its specific PP1 isoform for rapid dephosphorylation with remarkable selectivity. However, it is not known which GPCR domain directs the engagement of specific PP1 isoforms to the receptor.

Here, we have addressed this question using the closely-related sst2 and sst5 somatostatin receptors. The sst2 and sst5 receptors exhibit a high degree of homology in their transmembrane domains but exhibit divergent carboxyl-terminal tails. Both the sst2 and the sst5 receptor are pharmacological relevant targets for clinically-used drugs [5] [6] [7] [8] [9] but the two receptors exhibit strikingly different phosphorylation and trafficking patterns. The sst2 receptor is a prototypical class B receptor that is phosphorylated at a cluster of at least six carboxyl-terminal serine and threonine residues upon agonist exposure. The sst5 receptor then forms stable complexes with β-arrestin that co-internalize into the same endocytic vesicles. Consequently, the sst2 receptor recycles slowly [1] [10] [11]. By contrast, the sst5 receptor is a prototypical class A receptor in that its endocytosis is regulated by a single phosphorylation at T333. The sst5 receptor then forms relatively unstable β-arrestin complexes that dissociate at or near the plasma membrane. The receptor internalizes without β-arrestin and recycles rapidly [2] [12]. Here, we show that a tail-swap mutation of sst2 and sst5 receptors is required and sufficient to reverse the patterns of dephosphorylation and trafficking of these two receptors.

Materials and Methods

Reagents, plasmids and antibodies

SS-14 was obtained from Bachem (Weil am Rhein, Germany). DNA for HA-tagged human sst2 and sst5 receptor, 2-5- and 5-2-chimaera were generated via artificial gene synthesis and cloned into pcDNA3.1 by imaGenes (Berlin, Germany). The human HA-tagged sst2 receptor was obtained from UMR cDNA Resource
Center (Rolla, MO). The phosphorylation-independent rabbit monoclonal anti-sst2 antibody {UMB-1} and anti-sst5 antibody {UMB-4} were obtained from Epitomics (Burlingame, CA). The phosphosite-specific sst2A antibodies anti-pS341/pS343 (3157), anti-pT353/pT354 (0521), anti-pT356/pT359 (0522) and phosphosite-specific sst5 antibodies anti-pT333 (3567) as well as the rabbit polyclonal anti-HA antibodies were generated and extensively characterized as previously described [1] [2].

Cell culture and transfection

Human embryonic kidney HEK293 cells were obtained from the German Resource Centre for Biological Material (DSMZ, Braunschweig, Germany). HEK293 cells were grown in DMEM supplemented with 10% fetal calf serum. Cells were transfected with plasmids using Lipofectamine 2000 according to the instructions of the manufacturer (Invitrogen, Carlsbad, CA). Stable transfectants were selected in the presence of 400 μg/ml G418. Stable cells were characterized using radioligand-binding assays, Western blot analysis, and immunocytochemistry as described previously. All receptors and chimeras tested were present at the cell surface, expressed similar amounts of receptor protein and had similar affinities for SS-14 as the wild-type receptors and had similar affinities for SS-14 as the wild-type receptors.

Analysis of receptor internalization by confocal microscopy

Cells were grown on poly-L-lysine-coated coverslips overnight. After treatment with 1 μM SS-14 for 0, 15 or 30 min at 37°C, cells were fixed with 4% paraformaldehyde and 0.2% picric acid in phosphate buffer (pH 6.9) for 30 min at room temperature and washed several times. Specimens were permeabilized and then incubated with anti-sst2A {UMB-1} or anti-sst5 antibody {UMB-4} antibodies followed by Alexa488-conjugated secondary antibodies. Specimens were mounted and examined using a Zeiss LSM510 META laser scanning confocal microscope.

Quantification of receptor internalization by ELISA

Stably transfected HEK293 cells were seeded onto poly-L-lysine-treated 24-well plates. The next day, cells were preincubated with 1 μg/ml anti-HA antibody for 2 h at 4°C. After the appropriate treatment with SS-14 (1 μM) for 30 min at 37°C, cells were fixed and incubated with peroxidase-conjugated anti-rabbit antibody overnight. After washing, plates were developed with ABTS solution and analyzed at 405 nm using a microplate reader.

Western blot analysis

Cells were plated onto 60-mm dishes and grown to 80% confluence. After treatment with SS-14 (1 μM), cells were lysed in detergent buffer (50 mM Tris-HCl, pH 7.4, 150 mM NaCl, 5 mM EDTA, 10 mM NaF, 10 mM disodium pyrophosphate, 1% Nonidet P-40, 0.5% sodium deoxycholate, 0.1% SDS, 0.2 mM phenylmethylsulfonyl fluoride, 10 μg/ml leupeptin, 1 μg/ml pepstatin A, 1 μg/ml aprotinin, and 10 μg/ml bacitracin). All phosphorylation and dephosphorylation assays were performed at both physiological temperature (37°C) and at room temperature (22°C) for the indicated time periods. Glycosylated proteins were partially enriched using wheat germ lectin-agarose beads as described. Proteins were eluted from the beads using SDS-sample buffer for 20 min at 65°C and then resolved on 8% SDS-polyacrylamide gels. After electroblotting, membranes were incubated with phosphosite-specific antibodies anti-pS341/pS343 {3157}, anti-pT353/pT354 {0521}, anti-pT356/pT359 {0522}, anti-pT333 {3567} at a concentration of 0.1 μg/ml followed by detection using enhanced chemiluminescence. Blots were subsequently stripped and reprobed with anti-sst2A antibody {UMB-1} or anti-sst5 antibody {UMB-4} to confirm equal loading of the gels.

β-Arrestin-EGFP mobilization assay

HEK293 cells were seeded onto 35-mm glass-bottom culture dishes (Mattek, Ashland, MA). The next day, cells were transiently cotransfected with 0.2 μg β-arrestin-2-EGFP and 2 μg human or chimeric somatostatin receptor or with a mixture of 0.2 μg β-arrestin-2-EGFP, 0.8 μg GRK2 and 1.2 μg human/chimeric sst2 receptor per dish containing 200,000 cells using TurboFectTM (Fermentas) according to the instructions of the manufacturer. After 24 h, cells were transferred onto a temperature-controlled microscope stage set at 37°C of a Zeiss LSM510 META laser scanning confocal microscope. Images were collected sequentially using single line excitation at 488 nm with 513-540-nm band pass emission filters. Saturating concentrations of SS-14 (1 μM) were applied directly into the culture medium immediately after the initial image was taken.

Small Interfering RNA Silencing of Gene Expression

Chemically synthesized double-stranded siRNA duplexes (with 5′dTdT overhangs) were purchased from Qiagen (Hilden, Germany) for the following targets: PP1α catalytic subunit (5′- AACAGACCGTACAACATCCAA-3′), PP1β catalytic subunit (5′- ACAGGAGATGTGTCAGGAA-3′ and 5′-GTTCGAG-GCTTATGTATCA-3′), PP1y catalytic subunit (5′-ACATCGACGAGGATTATCCAA-3′ and 5′-AGAGGCGTTGGTCACT-3′), and a nonsilencing RNA duplex (5′- GCTTAGGAG-GATTAGTGAAA-3′ and 5′- AAA CTC TAT CTG CAC GGT GAC-3′). HEK293 cells were transfected with 150 nM siRNA for single transfection or with 100 nM of each siRNA for double transfection using HiPerFect (Qiagen). Silencing was quantified by immunoblotting. All experiments showed protein levels reduced by ≥80%.

Data Analysis

Data were analyzed using GraphPad Prism 4.0 software. Statistical analysis was carried out with Student’s t-test as well as with one-way or two-way ANOVA followed by the Bonferroni post-test. p-Values of <0.05 were considered statistically significant.

Results

The sst2 and sst5 receptors exhibit a high degree of homology, yet these somatostatin receptors are dephosphorylated by different PP1 isoforms. To elucidate which receptor domains determine this remarkable phosphatase specificity, we first constructed tail-swap mutants of these two receptors. In initial studies, we confirmed that all receptors were expressed at similar levels on the cell surface, and exhibited similar binding properties. All four receptors also exhibited similar signaling properties determined as their ability to activate ERK in a pertussis toxin-sensitive manner (not shown). We then compared agonist-induced phosphorylation of the wild-type sst2 receptor with that of the sst2-5 receptor using phosphosite-specific antibodies for pS341/343, pT353/354 and pT356/359 (Figure 1, left panel). Phosphorylation at the three sites was not detectable in untreated cells. In the presence of SS-14 phosphorylation at all three sites became detectable within seconds of agonist exposure in both the sst2 and the sst2-5 receptor. Next, we compared agonist-induced phosphorylation of the wild-type sst2 receptor with that of the sst2-5 receptor using phosphosite-specific antibodies for pT333 and pT347.
Phosphorylation at T347 was already detectable in untreated cells for both receptors (not shown). By contrast, phosphorylation at T333 was not detectable in untreated cells. However, upon addition of SS-14 T333 phosphorylation occurred within a few seconds in both sst5 and sst2-5 receptors (Figure 1, right panel).

The sst2 receptor and the sst5 receptor dramatically differ in the extent of their agonist-induced internalization. In the presence of SS-14, nearly all cell surface sst2 receptors are removed from the plasma membrane resulting in a 80% loss of surface receptors after 30 min agonist exposure (Figure 2A, B). By contrast, the sst5 shows only partial receptor internalization with a large proportion of receptors remaining at the plasma membrane resulting in a maximal internalization of 25% after 30 min SS-14 treatment (Figure 2A, B). Interestingly, swapping the cytoplasmic tails completely reversed this trafficking pattern in that the sst5-2 receptor revealed nearly complete (70%) and the sst2-5 receptor partial (25%) endocytosis (Figure 2).

We then employed functional β-arrestin-2 conjugated to enhanced green fluorescent protein (EGFP) to visualize the patterns of dephosphorylation and recycling. Interestingly, dephosphorylation of individual sst2 phosphate acceptor sites occurs with distinct temporal dynamics. Whereas T353/T354 dephosphorylation occurred rapidly (~5 min), T356/T359 dephosphorylation was delayed (~20 min) and S341/S343 dephosphorylation is only observed after extended SS-14 washout (~60 min) (Figure 4). When sst5-expressing cells were exposed to SS-14 for 5 min, washed and then incubated in agonist-free medium, T333 dephosphorylation occurred very rapidly (~2 min) (Figure 4). In contrast, T347 phosphorylation was although to a lesser extent still detectable even after prolonged incubation in the absence of agonist (not shown). Analysis of chimeric receptors under identical conditions showed that transplantation of the sst2 tail to the sst5 receptor led to an sst2-like dephosphorylation profile and β-arrestin-2 that appeared as punctuate staining within the cytoplasm at later time points (Figure 3B). The addition of 1 μM SS-14 to the human sst5 receptor induced a redistribution of β-arrestin-2 from the cytoplasm to the plasma membrane that was less pronounced compared to that seen in sst2-expressing cells (Figure 3A). Although over-expression of GRK2 clearly facilitated β-arrestin-2 recruitment in sst2-expressing cells at early time points, it did not lead to a redistribution of β-arrestin-2-EGFP into the cytosol at later time points (Figure 3B). Again, swapping the cytoplasmic tails led to a complete reversal of the β-arrestin trafficking patterns of these two receptors (Figure 3A, B).

The sst2 receptor and the sst5 receptor also dramatically differ in their patterns of dephosphorylation and recycling. Interestingly, dephosphorylation of individual sst2 phosphate acceptor sites occurs with distinct temporal dynamics. Whereas T353/T354 dephosphorylation occurred rapidly (~5 min), T356/T359 dephosphorylation was delayed (~20 min) and S341/S343 dephosphorylation is only observed after extended SS-14 washout (~60 min) (Figure 4). When sst5-expressing cells were exposed to SS-14 for 5 min, washed and then incubated in agonist-free medium, T333 dephosphorylation occurred very rapidly (~2 min) (Figure 4). In contrast, T347 phosphorylation was although to a lesser extent still detectable even after prolonged incubation in the absence of agonist (not shown). Analysis of chimeric receptors under identical conditions showed that transplantation of the sst2 tail to the sst5 receptor led to an sst2-like dephosphorylation profile.
Conversely, transplantation of the sst5 tail to the sst2 receptor led to an sst5-like dephosphorylation profile (Figure 4). Finally, we examined the PP1 specificity of the sst2 receptor and the sst5 receptor as well as their respective tail-swap mutants. To date, three distinct catalytic subunits \( \alpha, \beta \) and \( \gamma \) are known for PP1 [13]. To elucidate which of these PP1 isoforms is involved in sst5 dephosphorylation, we performed siRNA knockdown experiments. As depicted in Figure 5, PP1\( \beta \) knockdown resulted in a robust inhibition of sst2 dephosphorylation. In contrast, transfection of PP1\( \alpha \) or PP1\( \beta \) siRNA did not result in a significant inhibition of sst2 dephosphorylation (Figure 5). For the sst5 receptor only PP1\( \gamma \) knockdown resulted in a detectable inhibition of its dephosphorylation, while transfection of PP1\( \alpha \) or PP1\( \beta \) siRNA had no effect (Figure 5). Interestingly, swapping the cytoplasmic tails conferred PP1\( \gamma \) specificity to the sst2 receptor and PP1\( \beta \) specificity to the sst5 receptor (Figure 5). These results suggest that PP1 specificity of individual somatostatin receptor subtypes is primarily determined by their different carboxyl-terminal receptor domains.

**Discussion**

Although the regulation of agonist-induced phosphorylation and internalization has been studied in detail for many GPCRs, the molecular mechanisms and functional consequences of receptor dephosphorylation are far from understood. We have recently observed that closely related somatostatin receptor subtypes can be dephosphorylated by distinct PP1 isoforms. However, it is not known which GPCR domain directs the engagement of specific PP1 isoforms to the receptor. The major
finding of this study is that carboxyl-terminal regions of different somatostatin receptor subtypes is a major determinants for their PP1 selectivity. This conclusion is based on the observation that transplantation of the sst2 tail to the sst5 receptor led to a predominant dephosphorylation by PP1β, whereas transplantation of the sst5 tail to the sst2 receptor led to a predominant dephosphorylation by PP1γ. Moreover, swapping the cytoplasmic...

Figure 4. Dephosphorylation of sst2 and sst5 tail-swap mutants. Stably transfected HEK 293 cells were exposed to 1 μM SS-14 for 5 min, washed and incubated at room temperature in the absence of agonist for the indicated time periods. Cells were lysed and immunoblotted with the indicated phosphosite-specific antibodies. Blots were then stripped and reprobed with the phosphorylation-independent anti-sst2 antibody (UMB-4) or anti-sst5 antibody (UMB-1) to confirm equal loading of the gels. Shown are representative results from one of three independent experiments. The position of molecular mass markers is indicated on the left (in kDa). doi:10.1371/journal.pone.0091526.g004

Figure 5. PP1 specificity of sst2 and sst5 tail-swap mutants. (A) HEK 293 cells stably expressing the sst2 receptor or the sst5-2 tail-swap mutant were transfected with siRNA targeted to PP1α, PP1β, PP1γ or non-silencing siRNA control (SCR) for 72 h and then exposed to 1 μM SS-14 for 5 min. Cells were washed three times and then incubated for 0, 15 or 30 min in the absence of agonist. Cells were lysed and immunoblotted with anti-pT356/T359 antibody (0522). Blots were stripped and reprobed with the phosphorylation-independent anti-sst2 antibody (UMB-1) to confirm equal loading of the gels. (B) HEK 293 cells stably expressing the sst5 receptor or the sst2-5 tail-swap mutant were transfected with siRNA targeted to PP1α, PP1β, PP1γ or non-silencing siRNA control (SCR) for 72 h and then exposed to 1 μM SS-14 for 5 min. Cells were washed three times and then incubated for 0, 2 or 5 min in the absence of agonist. Cells were lysed and immunoblotted with anti-pT333 (3567) antibody. Blots were stripped and reprobed with the phosphorylation-independent anti-sst5 antibody (UMB-4) to confirm equal loading of the gels. Receptor phosphorylation was quantified and expressed as percentage of maximal phosphorylation in SCR-transfected cells, which was set at 100%. Data correspond to the mean ± SEM from three independent experiments. Results were analyzed by two-way ANOVA. (C) siRNA knockdown of PP1 was confirmed by Western blot using isoform-specific PP1 antibodies. The positions of molecular mass markers are indicated on the left (in kDa). doi:10.1371/journal.pone.0091526.g005
tails led to a complete reversal of the trafficking profiles of these two receptors.

The remarkable selectivity in the recruitment of specific PP1 catalytic subunits to individual somatostatin receptor subtypes is surprising. PP1 catalytic subunits bind to their regulatory subunits and some substrates in a mutually exclusive manner through a conserved RvxF motif. The three isoforms of the PP1 catalytic subunit share greater than 90% sequence identity, including the regions that interact with the RvxF sequence [13]. However, neither the human sst2 nor the human sst5 receptor has a potential PP1-binding motif in its carboxyl-terminal tail suggesting that somatostatin receptors do not bind to PP1 exclusively by the canonical RvxF motif. Instead, association of PP1 may occur directly through a noncanonical interaction or multiple weak interactions or indirectly via one or more regulatory subunits of PP1. Such targeting PP1 subunits are prime candidates to bring phosphatases in proximity to phosphorylated GPCRs. Nevertheless, the identity of the targeting PP1 subunits remains to be elucidated for both sst2 and sst5.

Somatostatin receptor subtypes exhibit strikingly different β-arrestin trafficking patterns. The sst2 receptor is a prototypical class B receptor that is phosphorylated at clusters of carboxyl-terminal serine and threonine residues. In turn the sst2 receptor forms stable β-arrestin complexes, co-internalizes with β-arrestin and recycles slowly. By contrast, sst5 is a prototypical class A receptor in that its endocytosis is driven by phosphorylation of a single threonine residue. The sst5 receptor then forms unstable β-arrestin complexes, internalizes without β-arrestin and recycles rapidly. Thus, our finding that swapping the cytoplasmic tails led not only to reversal of the PP1 specificity but also to a reversal of the β-arrestin trafficking profiles of somatostatin receptors suggests a simple model in which fast recycling class A receptors are preferentially dephosphorylated by PP1γ, whereas slow recycling class B receptors are preferentially dephosphorylated by PP1β. So far only few bona fide GPCR phosphatases have been identified [14] [15] [1] [2] [4]. However, it should be noted that this hypothesis is supported by our recent observation that the μ-opioid receptor, which is a prototypical fast recycling class A receptor, is rapidly dephosphorylated by PP1γ [3]. However, it should be noted that PP1γ was identified as GPCR phosphatase for the thyrotropin-releasing hormone receptor [4], suggesting that different phosphatases can interact with different GPCRs to mediate their dephosphorylation. It is also possible that distinct phosphatase activities mediate dephosphorylation of plasma membrane receptors versus internalized receptors.

GPCR dephosphorylation has long been viewed as an unregulated process with little or no functional implications. Nevertheless, more recent evidence suggests that PP1β-mediated dephosphorylation is involved in fine-tuning unconventional β-arrestin-dependent GPCR signaling. Indeed, inhibition of PP1β expression results in a specific enhancement of sst2-driven ERK activation [1]. Given that arrestin-dependent signaling is initiated by binding to phosphorylated receptors, this finding suggests that PP1β-mediated GPCR dephosphorylation limits β-arrestin-dependent signaling by disrupting the β-arrestin-GPCR complex.

In conclusion, different GPCRs can recruit specific PP1 isoforms for their rapid dephosphorylation with remarkable selectivity. This GPCR phosphatase specificity is primarily determined by carboxyl-terminal receptor domains. Recruitment of different GPCR phosphatases has in turn profound consequences for the dephosphorylation dynamics and trafficking patterns of GPCRs.

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Author Contributions

Conceived and designed the experiments: SS AL FN AK. Performed the experiments: AL AK JCM FN. Wrote the paper: SS AL.

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