Biosensor-Based Approach Identifies Four Distinct Calmodulin-Binding Domains in the G Protein-Coupled Estrogen Receptor 1

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Abstract

The G protein-coupled estrogen receptor 1 (GPER) has been demonstrated to participate in many cellular functions, but its regulatory inputs are not clearly understood. Here we describe a new approach that identifies GPER as a calmodulin-binding protein, locates interaction sites, and characterizes their binding properties. GPER coimmunoprecipitates with calmodulin in primary vascular smooth muscle cells under resting conditions, which is enhanced upon acute treatment with either specific ligands or a Ca\(^{2+}\)-elevating agent. To confirm direct interaction and locate the calmodulin-binding domain(s), we designed a series of FRET biosensors that consist of enhanced cyan and yellow fluorescent proteins flanking each of GPER’s submembrane domains (SMDs). Responses of these biosensors showed that all four submembrane domains directly bind calmodulin. Modifications of biosensor linker identified domains that display the strongest calmodulin-binding affinities and largest biosensor dynamics, including a.a. 83–93, 150–175, 242–259, 330–351, corresponding respectively to SMDs 1, 2, 3, and the juxta-membranous section of SMD4. These biosensors bind calmodulin in a strictly Ca\(^{2+}\)-dependent fashion and with disparate affinities in the order SMD2>SMD4>SMD3>SMD1, apparent K\(_d\) values being 0.44±0.03, 1.40±0.16, 8.01±0.29, and 136.62±6.56 μM, respectively. Interestingly, simultaneous determinations of biosensor responses and suitable Ca\(^{2+}\) indicators identified separate Ca\(^{2+}\) sensitivities for their interactions with calmodulin. SMD1-CaM complexes display a biphasic Ca\(^{2+}\) response, representing two distinct species (SMD1 sp1 and SMD1 sp2) with drastically different Ca\(^{2+}\) sensitivities. The Ca\(^{2+}\) sensitivities of CaM-SMDs interactions follow the order SMD1sp1>SMD4>SMD2>SMD1sp2>SMD3, EC\(_{50}(\text{Ca}^{2+})\) values being 0.13±0.02, 0.75±0.05, 2.38±0.13, 3.71±0.13, and 5.15±0.25 μM, respectively. These data indicate that calmodulin may regulate GPER-dependent signaling at the receptor level through multiple interaction sites. FRET biosensors represent a simple method to identify unknown calmodulin-binding domains in G protein-coupled receptors and to quantitatively assess binding properties.

Introduction

Plasma estrogen concentrations are closely related to cardiovascular health. Along with menopause comes a substantial increase in the risk of cardiovascular diseases [1,2]. Estrogen modulates gene expression, growth, development and immune responses, and has many cardiovascular protective effects. The actions of estrogen are extensive and include effects that are both dependent and independent of transcriptional activities [3,4]. The mechanisms underlying these effects are still far from being completely understood [5,6]. Indeed, two classical estrogen receptors, ER\(_{\alpha}\) and ER\(_{\beta}\), which function as transcriptional factors that bind estrogen responsive elements in promoters of target genes [3], were thought to be totally responsible for estrogen’s effects. However, a novel G protein-coupled receptor, GPR30, was cloned around 1997 as an orphan receptor [7–11], and was demonstrated to be an estrogen receptor in 2005 [12,13]. It was subsequently termed G protein-coupled estrogen receptor 1 (GPER) by International Union of Basic and Clinical Pharmacology (IUPHAR). GPER has since been shown to be involved in many cellular activities, including Ca\(^{2+}\) mobilization [13,14], cAMP production [12,14], activation of protein kinases [13,15], and activation of transcription [16–19]. Clarifying the regulation of GPER and related pathways will enhance our knowledge of how estrogen works and provide grounds to target estrogen receptor subtypes for preventive and therapeutic purposes.

Calmodulin (CaM) is a highly conserved 148-a.a. protein that contains four EF-hand Ca\(^{2+}\)-binding motifs. Ca\(^{2+}\)-bound CaM has a dumbbell conformation with two EF-hand motifs on either end connected by a central helix. This central helix functions as a tether that is bent upon target interaction while the two lobes exert regulatory inputs in a significant number of cell types, including vascular endothelial cells [25,26], smooth muscle cells [27], and...
cardiomyocytes [28,29]. This insufficiency of CaM has been demonstrated to generate functional coupling among its targets due simply to competition for CaM [25,26], and suggests that factors controlling CaM expression and dynamics can vastly alter cellular functions.

G protein-coupled receptors (GPCRs) represent a superfamily of cell surface proteins that convey extracellular inputs to vast changes in cellular functions via dynamic associations with heterotrimeric G proteins and numerous other partners at their submembrane domains. Recently, CaM has been demonstrated to interact with a number of G protein-coupled receptors (GPCRs), such as the metabotropic glutamate receptors mGluR1 and mGluR5 [30,31], the opioid µ receptor [32], the parathyroid hormone receptor 1 [33], the 5-HT2C and 5-HT1A receptors [34,35], the D2 dopamine receptor [36], and the angiotensin II receptor type 1A [37,38]. In the case of the 5HT1A receptor, CaM binding to the third submembrane domain alters the receptor’s phosphorylation or interaction with G protein subunit(s) [35], while for the 5-HT2C receptor, CaM binding to the C terminal tail has been demonstrated to be important for β-arrestin recruitment and for receptor-operated extracellular signal-regulated kinase [39]. Nevertheless, the roles of CaM in GPCR biology at the receptor level is still not entirely clear. Part of the reason for this is the lack of an approach to exhaustively identify all interaction sites for CaM on a GPCR. For example, a CaM-binding domain in the juxtamembrane region of the cytoplasmic tail of the angiotensin II receptor type 1A was first identified in 1999 using a peptide deduced from a sequence-based comparison with known CaM-binding motifs [37]. More than a decade later, another CaM-binding domain in the third submembrane domain in the same receptor was recently identified using a similar approach [30]. In addition, while the interactions between CaM and these GPCRs are Ca²⁺-dependent, information regarding the specific Ca²⁺ sensitivities of these interactions is lacking. Knowledge of the Ca²⁺ sensitivities of GPCR-CaM interactions will be of value in determining the roles of CaM interaction with each submembrane domain at different physiological scenarios in cells.

With respect to estrogen biology, it is completely unknown if CaM plays a role GPER-dependent signaling. We have begun to test the idea that CaM is directly involved in GPER-dependent signaling in the vasculature at both the receptor level and downstream effectors. In this paper, we describe a new approach to identify GPER as a novel CaM-binding protein, locate all of its CaM-binding domains, and determine their binding characteristics including K_d and Ca²⁺ sensitivities. Using FRET-based biosensors designed to scan all submembrane domains of GPER, we identified up to four binding sites located each in submembrane domains 1, 2, 3 and 4 of GPER that have distinct binding affinities for CaM. In addition, we simultaneously incorporated these biosensors and suitable Ca²⁺ indicators in single assays to precisely determine the Ca²⁺ sensitivities of the interactions between CaM and these different domains in GPER. This approach in turn revealed quite distinct sensitivities to Ca²⁺ of the interactions between CaM and GPER’s submembrane domains. Interestingly, submembrane domain 1 interacts with CaM with biphasic Ca²⁺ dependency, representing two distinct populations of complexes formed at drastically different free Ca²⁺ values that range from cytoplasmic resting levels to those that occur during Ca²⁺ entry. The results demonstrate that FRET biosensor-based screening is an effective and efficient approach to identify unknown CaM-binding domains in GPCRs that allows for highly quantitative assessment of their binding properties.

### Materials and Methods

#### Ethics Statement

The protocol to isolate primary vascular smooth muscle cells from porcine aortas (obtained commercially from a local slaughter house) was approved by the Des Moines University Institutional Biosafety Committee.

#### Cell Isolation and Culture

Descending segments of porcine thoracic aortas were freshly collected from a local slaughter house. Primary vascular smooth muscle cells (VSMCs) were isolated from these aortas using a modification of the methods published by Bolzon [40], Ulrich-Merzenich [41], and Leik [42]. Briefly, endothelial cells were initially completely removed by a sequence of mechanical scraping, short enzyme incubation (0.02% collagenase, 0.1% papain, and 4 mM dithiothreitol) and re-scraping. After rinsing in sterile PBS, aortas were cut into rectangular strips and the luminal surface was incubated with the same enzyme mix for 90 minutes. The luminal surface was then scraped again, and cells from this enzyme treatment and scraping were grown in M-199 containing 10% fetal bovine serum and 2% penicillin/streptomycin in 90% humidified condition with 5% CO₂ at 37°C for one week. This procedure consistently yields highly homologous populations VSMCs, verified by both morphological properties and smooth muscle α-actin immunofluorescence staining. This protocol was approved by Des Moines University Institutional Biosafety Committee.

#### Co-immunoprecipitation

Co-immunoprecipitation was performed using a Pierce immunoprecipitation kit (Rockford, IL) as per manufacturer’s instructions. The use of an amino-reactive resin in this approach prevents the co-elution of the antibody heavy and light chains that may co-migrate with the bands of the proteins of interest. Typically, 10 µL of resin was conjugated with appropriate amounts of primary antibody. After appropriate treatment, cell lysis was performed on monolayer cells without trypsinization to avoid complications having to do with shear during the process. Cells were lysed on ice for 15 minutes and the lysate was centrifuged at 21,000 x g for 5 min at 4°C. Following pre-clearing, the protein content of the eluant was determined using the BCA assay (Pierce). One milligram of total cellular proteins was then rocked with the resin-antibody conjugates for 3 hours in 1-mL columns at 4°C prior to eluting. Following electrophoresis and transfer, membranes were cut between the levels of the bait and the prey proteins prior to incubation with separate primary antibodies against CaM (EMD Millipore Corp., MA) and GPER (R&D System, MN; GeneTex Inc., Irvine, CA, or Santa Cruz Biotech, CA).

#### Construction of Biosensors to Identify GPER’s CaM-binding Domains

Biosensors to screen for CaM-binding domains in GPER were constructed based on the principle of fluorescent resonance energy transfer (FRET). Sub-membrane domains (SMDs) 1–4 from human GPER (Fig. 1A), corresponding to a.a. 83–93, 150–175, 242–259, 330–375, and shorter fragments of these domains were PCR amplified from the human GPER cDNA (GeneCopoeia Inc., Rockville, MD) and inserted between enhanced variants of cyan and yellow fluorescent proteins as a donor–acceptor FRET pair. We termed these biosensors BSGPER_x, where x denotes the amino acid numbering of the GPER insert between the fluorophores. In total, the BSGPER_x made were BSGPER_R3-93.
BSGPERGGG83–93GGG, BSGPER150–170, BSGPER150–175, BSGPER242–259, BSGPER330–345, BSGPER330–351, BSGPER330–361, BSGPER330–375, and BSGPER 362–375 (Fig. 1B). The BSGPERGGG83–93GGG construct consists of three additional glycine residues on the N- and C-terminal ends of SMD1 (see Results section). Biosensors were introduced into a bacterial expression pET vector with a 6-histidine tag at the C terminus downstream of ECFP (a gift from Dr. Anthony Persechini, University of Missouri-Kansas City).

Expression and Purification of Biosensors

BSGPERx were expressed in BL21(DE3) competent cells. An overnight culture of 5 mL was inoculated into 300 mL of 2XYT culture medium and shaken at 37°C until OD 0.6–0.8. Expression was then induced with 0.5 mM isopropyl 1-thiogalactopyranoside (IPTG) and the culture was grown at 22°C for an additional 24 hrs. Expressed biosensors were then purified using standard affinity chromatography with Cb2+ affinity resin (Pierce). Eluant was dialyzed twice against large volumes of 50 mM Tris, 100 mM KCl, pH 7.5 at 4°C. Purified BSGPERx were stored at –80°C.

Expression and Purification of CaM

CaM was purified with a slight modification of previously published approach. A bacterial expression pET vector encoding the rat CaMl gene was transformed into BL21(DE3) E. Coli. An overnight culture of 10 mL was added to 1L of culture media and grown at 37°C until O.D. reached 0.6–0.8. Expression was induced by 1 mM IPTG. The culture was then grown for 4 hrs at 37°C. After centrifugation, pellets were resuspended in 50 mM Tris, 100 mM KCl, pH 7.5 at room temperature. Resuspension was treated with 200 μg/mL lysozyme, protease inhibitor cocktail (0.06 units/mL) and 0.01 μL/mL PMSF for 1 hr. Cells were lysed by sonication. After centrifugation at 100,000 × g for 30 min, 10 mM CaCl2 was added to the supernatant and heated to 65°C for 15 minutes. Supernatant was brought to room temperature and centrifuged again at 100,000 × g for 30 min in the presence of 4 μL/mL DNAse and 3 mM MgCl2. Supernatant was loaded onto a CL-4B Phenyl Sepharose column. Columns were subjected to three long successive washes with buffer A (50 mM Tris, 1 mM CaCl2, pH 7.5), B (50 mM Tris, 1 mM CaCl2, 5 mM imidazole and 300 mM NaCl), followed by a brief wash with buffer A. CaM was eluted with 50 mM Tris, 100 mM NaCl, and 5 mM EGTA. Eluant was dialyzed twice against 50 mM Tris, 100 mM KCl, pH 7.5 and concentrated using Centriprep (Millipore) prior to use. This protocol consistently produces high yields of pure CaM.

Binding Assay to Screen for CaM-binding Domains in GPER

BSGPERx were tested in vitro for direct binding with purified CaM. BSGPERx were mixed in titration buffer (25 mM Tris, 100 mM KCl, pH 7.5) containing 0.1 mg/mL bovine serum albumin and 1 mM CaCl2 in a quartz cuvette (Hellma Analytics). Small aliquots of purified CaM were then added to this mixture as biosensor responses were monitored in a QuantaMaster™-40 spectrofluometer (Photon Technology International Inc.). All titrations were performed at 22°C. Direct interactions between the specified GPER fragments and CaM were identified by disruption of FRET between donor ECFP and acceptor EYFP. Biosensor fractional responses were determined by the formula

\[ BS_{frac} = \frac{R - R_{min}}{R_{max} - R_{min}} \]

where Rmin and Rmax are the ratios between emission intensities at \( \lambda_{475} \) and \( \lambda_{535} \) nm (F475/F535) when the biosensor is in unbound and maximally bound state, respectively. Fractional biosensor responses could also be determined using the emission intensity at \( \lambda_{475} \) nm (F475) or \( \lambda_{535} \) nm (F535), as follow.

\[ BS_{frac} = \frac{F - F_{min}}{F_{max} - F_{min}} \]

where F, Fmax and Fmin represent the observed, maximal and minimal intensities at \( \lambda_{475} \) nm.

Figure 1. Schematic topography of human GPER (A) and sub-membrane fragments used to generate biosensors (B). SMD, sub-membrane domain. Hydrophobic residues are in bold-face type. Positively charged residues are underlined. doi:10.1371/journal.pone.0089669.g001
Or,

\[ BS_{\text{frac}} = \frac{F_{\text{max}} - F}{F_{\text{max}} - F_{\text{min}}} \]  

where \( F, F_{\text{max}} \) and \( F_{\text{min}} \) represent the observed, maximal and minimal intensities at \( \lambda = 535 \) nm, as described previously [43]. Dilution factors were tightly calculated for each addition of CaM using a detailed excel spreadsheet algorithm and applied for intensity correction when the F475 or F535 values were used to derive BS_{frac}; however, we determined that the use of ratiometric responses (formula (1)) negated this necessity. Precise fractional biosensor responses were plotted against titrated CaM concentrations. Apparent \( K_d \) values for BSGPERx-CaM interactions were obtained by fitting biosensor’s fractional responses as a function of concentration.
CaM concentration to hyperbolic or quadratic binding equations [43]:

\[
BS_{\text{frac}} = \frac{[\text{CaM}]_{\text{tot}}}{[\text{CaM}]_{\text{tot}} - K_d}
\]

(4)

where \(BS_{\text{frac}}\), \([\text{BS}]\), and \([\text{CaM}]\) are fractional response of BSGPER\(_x\) (equation (1)) and the total concentration of biosensor and CaM in the mixture, respectively.

**Determination of \(Ca^{2+}\) Sensitivity of the Interaction between BSGPER\(_x\) and CaM**

\(Ca^{2+}\) sensitivities of the interactions between CaM and the different submembrane domains of GPER were determined by simultaneous spectrofluorometric measurements of the responses of a suitable \(Ca^{2+}\) indicator and BSGPER\(_x\) in the presence of saturating CaM concentration and incremental increases in free \(Ca^{2+}\). Saturating CaM concentrations were obtained from determinations of apparent \(K_d\) values of interactions between BSGPER\(_x\) and \(Ca^{2+}\)-CaM described above. Free \(Ca^{2+}\)

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**Table 1. BSGPER\(_x\) dynamic ranges and affinities for CaM.**

| BSGPER\(_x\)     | SMD | Dynamic Range | Apparent \(K_d\) (nM) |
|------------------|-----|---------------|-----------------------|
| BSGPER33–93      | 1   | 1.31±0.01     | 136.62±6.56           |
| BSGPER150–175    | 2   | 1.61±0.03     | 0.44±0.03             |
| BSGPER150–170    | 2   | 1.34±0.02     | 3.41±0.09             |
| BSGPER242–259    | 3   | 2.65±0.05     | 8.01±0.49             |
| BSGPER330–351    | 4   | 2.50±0.03     | 1.40±0.16             |
| BSGPER330–345    | 4   | 2.45±0.04     | 1.60±0.22             |

Data are means ± SD of at least three independent determinations.

*All \(K_d\) values are significantly different (p<0.05) from one another.

\*, these values are different (p<0.05) from the remaining dynamic range values.

**Figure 3. Design of biosensors to screen for CaM binding sequences in GPER.** Biosensor is composed of a FRET donor ECFP and a FRET acceptor citrine EYFPc flanking each of the four submembrane domains in GPER and their truncated segments corresponding to our predicted CaM-binding sequences. The proximity between the donor and acceptor in the unbound state facilitate robust FRET when the donor ECFP is excited at 430 nm giving emission light at 475 nm, which in turn excites the acceptor EYFPc, which emits at 535 nm (A). Upon addition of CaM, if there is specific interaction with the GPER insert fragment, the conformational changes that occur upon specific interaction with CaM will disrupt FRET, decreasing emission from EYFPc while increasing that from donor ECFP. This signature spectral change identifies CaM binding (B). C and D, Typical spectrofluorometric response of biosensor in the absence of specific CaM binding to its linker (C), which corresponds to the biosensor configuration in (A), and in the presence of CaM and saturating concentration of \(Ca^{2+}\) (D), which corresponds to the biosensor configuration in (B).

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concentrations were determined by the responses of X-Rhod5F emission intensity at \(\lambda 600\) nm with excitation at \(\lambda 580\) nm, which were simultaneously monitored with biosensor response. Biosensor FRET emission ratio (F475/F535) following excitation at \(\lambda 430\) nm were concurrently monitored in a QuantaMaster-40 spectrofluorometer. Typically, starting reaction mixture contained 0.5 mM GPER biosensor, 2 mM X-Rhod5F, 0.1 mg/mL BSA, 0.25 mM Br2BAPTA and saturating CaM concentrations (obtained from \(K_d\) titrations) in 25 mM Tris, 100 mM KCl, pH 7.5 at 22°C. In some assays with high saturating CaM concentrations, the concentration of Br2BAPTA was increased to guarantee chelation of Ca\(^{2+}\) prior to the start of Ca\(^{2+}\) titrations (see Results). In this nominally Ca\(^{2+}\)-free condition, there was no interaction between CaM and BSGPERx. Incremental Ca\(^{2+}\) was then titrated into the reaction mix. Free Ca\(^{2+}\) values were calculated using the formula

\[
[Ca^{2+}]_{\text{free}} = \left[ \frac{[Ca^{2+}]_{\text{max}}}{[Ca^{2+}]_{\text{max}} + [EC_{50}(Ca^{2+})]} \right]^{n}
\]

where \([Ca^{2+}]_{\text{free}}\) is obtained from equation (1); \(n\) is the Hill coefficient.

Statistical Analysis

Data are expressed as means ± SD of at least three independent experiments. Statistical analysis was performed using Student’s \(t\)-test, assuming unequal variances between control and treated groups. Statistical significance was determined as \(P<0.05\).

Results

Complex Formation between GPER and CaM in Primary Vascular Smooth Muscle Cells

Estrogen activates many CaM-dependent processes. We considered the possibility that CaM might be involved in estrogen actions through GPER at the receptor level. We asked whether GPER interacts with CaM and whether this interaction requires GPER activation by specific ligand binding. To test this, primary VSMCs were acutely treated with vehicle or the endogenous ligand 17β-estradiol (E2, 5 nM), the GPER agonist G-1 (100 nM), or the SERCA pump inhibitor thapsigargin (2 µM) for 3 minutes in the presence of 1 mM extracellular Ca\(^{2+}\). Both E2 and G-1 have been demonstrated to trigger intracellular Ca\(^{2+}\) signals [13,14,44–48], and thapsigargin is a well-known stimulator of store-operated Ca\(^{2+}\) entry. GPER was immunoprecipitated from the lysate and the pull-down fractions were probed for CaM (Fig. 2A). Reciprocally, CaM was immunoprecipitated from the lysate and the pulled down fractions were subsequently probed for GPER.
In both co-IP directions, CaM appears to associate with GPER under resting conditions. Both 17β-estradiol and GPER agonist G-1 apparently enhanced the association between GPER and CaM. Interestingly, thapsigargin, which stimulates Ca\textsuperscript{2+} entry independently of GPER activation, also produced a similar effect (Fig. 2A and 2B). To confirm specificity of the co-IPs, total cell lysate from a set of samples used in the co-IP reactions in Fig. 2A and 2B was first probed with non-immune IgGs from both mouse (Fig. 2C, left panel, species compatible with that of the anti-CaM antibody) and rabbit (Fig. 2D, left panel, species compatible with that of the anti-GPER antibody). These panels, predictably, showed no detectible bands. Following probing with the non-immune IgGs, the same membranes were stripped and re-probed with mouse anti-CaM antibody (Fig. 2C, right panel) or rabbit anti-GPER antibody (Fig. 2D, right panel). These panels now showed clear GPER and CaM bands, confirming the specificity of the antibodies used for GPER and CaM. To further confirm the specificity of the co-IPs in Fig. 2A and 2B, a control co-IP paradigm was tested in which GPER and CaM were immuno-precipitated from lysate of cells treated acutely with 5 nM E\textsubscript{2}, followed by immunoblotting with non-immune rabbit antibody (Fig. 2E, left upper panel) and mouse non-immune IgG (Fig. 2E, left lower panel). As predicted, these panels showed no detectible immunoprecipitates. The same membranes were then stripped and immunoblotted with non-immune rabbit antibody (Fig. 2E, right upper panel) and mouse non-immune IgG (Fig. 2E, right lower panel). Additionally, immunoprecipitation of lysate from E\textsubscript{2}-treated cells using the non-immune IgGs showed no detectible GPER or CaM upon immunoblotting with their specific antibodies (Fig. 2F, upper and lower panels, respectively).

**Identification of GPER's CaM-binding Domains**

The co-immunoprecipitation of GPER and CaM in VSMCs suggested that GPER might directly interact with CaM. In general, CaM-binding sequences do not belong to a common motif, so it is challenging to define CaM-binding domains in different proteins using sequence alignment [49]. For a GPCR, sub-membrane domains represent logical site(s) for these interactions. To confirm direct GPER-CaM interaction and locate the CaM-binding domains, we decided to use FRET technology as a new approach to confirm direct interaction and identify CaM-binding sites. The typical length of CaM-binding sequences, ~20 amino acids, and the large conformational change produced upon CaM binding are ideal for FRET technology. FRET biosensors based on known CaM-binding sequences have been developed, modified and used successfully to monitor free Ca\textsuperscript{2+}-CaM signals in cells [25,26,28,43,50]. However, FRET technique has not been used to locate unknown CaM-binding sequences. We believe that this technique could also be used to cost-effectively and precisely locate unknown CaM-binding domains in GPCRs and quantitatively determine their binding properties. Figure 3 recapitulates the principle of this approach. Given sufficient proximity of the energy donor ECFP and acceptor EYFP, FRET is formed between these moieties upon excitation of ECFP at 430 nm (Fig. 3A). Specific binding of CaM to the linker causes a conformational change between the two fluorescent proteins and

![Figure 5. Reversibility and Ca\textsuperscript{2+} dependency of BSGPER\textsubscript{CaM} interactions.](image-url)
affinity of BSGPER150–170 (closed diamonds) and BSGPER150–175 (open squares) revealed the role of the basic patch (a.a. 170–175) in CaM binding to SMD2. To confirm Ca2+-dependency of these interactions, saturating concentrations of CaM determined from titrations similar to those shown in Fig. 4 were added to a mixture of 500 nM BSGPER150–175 and 250 μM Ca2+(3 mM Ca2+ for BSGPER330–375). After maximal responses were obtained for BSGPER83–93, BSGPER150–175, BSGPER242–259, BSGPER330–375 (solid blue traces, Fig. 5A-D), 2 mM BAPTA (5 mM for BSGPER83–93) was added to chelate Ca2+ in the mixture. This addition completely reversed the responses for these biosensors (dotted traces, Fig. 5A-D). These data indicate direct Ca2+-dependent interaction between CaM and GPER’s submembrane domains 1–4.

To narrow down the precise CaM-binding domains in GPER, our strategy was to adjust the length of the linker sequences in keeping if possible with common CaM-binding motifs [51] and then screen for biosensors that yield the lowest Kd values for CaM and largest dynamic ranges. BSGPER83–93 (SMD1) represents a fairy short sequence (11 residues) that does not follow any known CaM-binding motifs and yet apparently binds CaM in a strictly Ca2+-dependent fashion, albeit with relatively low affinity. BSGPER150–175 (SMD2) contains a 20-a.a. segment (a.a. 150–170) of hydrophobic residues interspersed among positively charged residues, while the remaining part (a.a. 170–175) is a patch of dense basic, positively charged residues (Fig. 1B). Although the positions of bulky hydrophobic residues in the 150–170 segment do not fit any common CaM-binding motifs [51], the composition of hydrophobic and positively charged residues does suggest potential of a CaM-binding domain. BSGPER150–170 was thus constructed for comparison with BSGPER150–175. This comparison would also reveal the role of the basic patch (a.a. 170–175) in CaM binding to SMD2. Figure 6A–C shows a typical experiment to determine the correct CaM-binding sequence in GPER’s SMDs 2–4, or whether biosensor responses, operationally defined as the fold difference between the observed Rmin and Rmax (equation 1). This value reflects the conformational change of the linker that occurs upon binding and was used as an additive factor in determining the precise location of the CaM-binding domains.

As an initial screen, we inserted the sub-membrane domains (SMDs) of GPER as linkers between the FRET donor ECFP and acceptor EYFPc and screened for direct interactions with purified CaM. Fig. 4 shows the emission spectral changes of these biosensors upon incremental additions of Ca2+-saturated CaM (Ca2+-CaM), BSGPER83–93, whose linker corresponds to GPER’s SMD1, showed reciprocal responses in its emission at λ475 nm and λ535 nm only to fairly high CaM concentrations (Fig. 4A), and was saturated at 500 μM BSGPER150–175, BSGPER242–259 and BSGPER330–375, whose linker sequences correspond to GPER’s SMDs 2–4, also display reciprocal changes in its emission intensities at λ475 nm and λ535 nm upon additions of Ca2+-CaM (Fig. 4B–D), albeit with different dynamic ranges (Table 1). The lower dynamic range of BSGPER150–175 (Fig. 4D) is likely due to its longer linker sequence. To confirm Ca2+-dependency of these interactions, saturating concentrations of CaM determined from titrations similar to those shown in Fig. 4 were added to a mixture of 500 nM BSGPER150–170 and 250 μM Ca2+(3 mM Ca2+ for BSGPER330–375). After maximal responses were obtained for BSGPER83–93, BSGPER150–175, BSGPER242–259, BSGPER330–375 (solid blue traces, Fig. 5A–D), 2 mM BAPTA (5 mM for BSGPER83–93) was added to chelate Ca2+ in the mixture. This addition completely reversed the responses for these biosensors (dotted traces, Fig. 5A-D). These data indicate direct Ca2+-dependent interaction between CaM and GPER’s submembrane domains 1–4.

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As an initial screen, we inserted the sub-membrane domains (SMDs) of GPER as linkers between the FRET donor ECFP and acceptor EYFPc and screened for direct interactions with purified CaM. Fig. 4 shows the emission spectral changes of these biosensors upon incremental additions of Ca2+-saturated CaM (Ca2+-CaM), BSGPER83–93, whose linker corresponds to GPER’s SMD1, showed reciprocal responses in its emission at λ475 nm and λ535 nm only to fairly high CaM concentrations (Fig. 4A), and was saturated at 500 μM BSGPER150–175, BSGPER242–259 and BSGPER330–375, whose linker sequences correspond to GPER’s SMDs 2–4, also display reciprocal changes in its emission intensities at λ475 nm and λ535 nm upon additions of Ca2+-CaM (Fig. 4B–D), albeit with different dynamic ranges (Table 1). The lower dynamic range of BSGPER150–175 (Fig. 4D) is likely due to its longer linker sequence. To confirm Ca2+-dependency of these interactions, saturating concentrations of CaM determined from titrations similar to those shown in Fig. 4 were added to a mixture of 500 nM BSGPER150–170 and 250 μM Ca2+(3 mM Ca2+ for BSGPER330–375). After maximal responses were obtained for BSGPER83–93, BSGPER150–175, BSGPER242–259, BSGPER330–375 (solid blue traces, Fig. 5A–D), 2 mM BAPTA (5 mM for BSGPER83–93) was added to chelate Ca2+ in the mixture. This addition completely reversed the responses for these biosensors (dotted traces, Fig. 5A-D). These data indicate direct Ca2+-dependent interaction between CaM and GPER’s submembrane domains 1–4.

To narrow down the precise CaM-binding domains in GPER, our strategy was to adjust the length of the linker sequences in keeping if possible with common CaM-binding motifs [51] and then screen for biosensors that yield the lowest Kd values for CaM and largest dynamic ranges. BSGPER83–93 (SMD1) represents a fairy short sequence (11 residues) that does not follow any known CaM-binding motifs and yet apparently binds CaM in a strictly Ca2+-dependent fashion, albeit with relatively low affinity. BSGPER150–175 (SMD2) contains a 20-a.a. segment (a.a. 150–170) of hydrophobic residues interspersed among positively charged residues, while the remaining part (a.a. 170–175) is a patch of dense basic, positively charged residues (Fig. 1B). Although the positions of bulky hydrophobic residues in the 150–170 segment do not fit any common CaM-binding motifs [51], the composition of hydrophobic and positively charged residues does suggest potential of a CaM-binding domain. BSGPER150–170 was thus constructed for comparison with BSGPER150–175. This comparison would also reveal the role of the basic patch (a.a. 170–175) in CaM binding to SMD2. Figure 6A–C shows a typical experiment to determine the correct CaM-binding sequence in GPER’s SMDs 2–4, or whether biosensor responses, operationally defined as the fold difference between the observed Rmin and Rmax (equation 1). This value reflects the conformational change of the linker that occurs upon binding and was used as an additive factor in determining the precise location of the CaM-binding domains.
indicates that the entire SMD2 participates in CaM interaction. It also demonstrates the importance of the patch of basic, positively charged residues (170–175) in CaM binding to SMD2.

The 18-residue SMD3 (a.a. 242–259) contains several hydrophobic residues interspersed among a very dense group of basic residues. The hydrophobic residues here again do not appear to follow any of the common motifs of CaM-binding sequences. Given the short length of this submembrane domain, it is likely that the entire domain interacts with CaM (Fig. 4C). As for SMD4, we noted that BSGPER330–375 showed limited dynamic range, due probably to its long linker sequence (Fig. 4D). The SMD4 sequence shows in its juxtamembrane region a potential CaM-binding domain. Additional BSGPERs were thus constructed with linkers corresponding to fragments a.a. 330–345, 330–351, and 330–361. To rule out a role of the C-terminal end of SMD4 in its interaction with CaM, a biosensor was also generated based on a.a. 362–375. As shown in Fig. 7A–C, BSGPER330–345, BSGPER330–351 and BSGPER330–361 all display robust responses to Ca2+-CaM, in agreement with the prediction that the CaM-binding domain in SMD4 is located in its juxtamembrane segment. Consistently, BSGPER362–375 does not respond to Ca2+-CaM, even at such high concentration as 700 μM CaM (Fig. 7D); this serves as a negative control that CaM does not affect ECFP and EYFP fluorescence. BSGPER330–351 displays slightly larger dynamic range and higher affinity than BSGPER330–345, while BSGPER330–361 shows a limited dynamic range. These data indicate that a.a. 330–351 represents the CaM-binding sequence in SMD4. The data presented so far clearly indicate that GPER possess four distinct CaM-binding sites in all of its submembrane domains with disparate affinities. An aggregate plot representing these remarkable differences is shown in Fig. 8. The apparent Kd values for these interaction are significantly different from one another (p<0.03) and are summarized in Table 1 alongside their dynamic ranges.

Ca2+ Sensitivities of the Interactions between CaM and GPER’s CaM-binding Domains

The disparate affinities for CaM binding of the four domains identified above suggest that they may interact with CaM under different physiological scenarios. Given the complete Ca2+-dependent nature of these interactions (Fig. 5), knowledge of
their Ca\textsuperscript{2+} sensitivities would be very helpful in determining the roles of CaM binding to each submembrane domain when studying GPER's physiological functions. The best approach was to simultaneously monitor BSGPER\textsubscript{x} fractional responses and free Ca\textsuperscript{2+} concentrations in the presence of saturating CaM and increasing amounts of free Ca\textsuperscript{2+} measured using an appropriate Ca\textsuperscript{2+} indicator. Based on the \( K_d \) values determined (Table 1), saturating CaM concentrations for the biosensors were known. The choice of a suitable Ca\textsuperscript{2+} indicator to do this depends on two factors. First, its excitation and emission spectra should not interfere with those of the biosensors. Second, it must have an apparent \( K_d \) for Ca\textsuperscript{2+} that is close enough to the Ca\textsuperscript{2+} sensitivity of the biosensor being tested, which could be quickly assessed by simultaneous measurement of both biosensor and the Ca\textsuperscript{2+} indicator responses as Ca\textsuperscript{2+} is titrated to the mixture. A wide range of commercially available Ca\textsuperscript{2+} indicators made this task relatively easy. A few Ca\textsuperscript{2+} indicators were tested, including indo-1, Mag-indo1, and XRhod-5F. XRhod-5F represented an ideal Ca\textsuperscript{2+} indicator for use with these biosensors, due to the closeness of its \( K_d \) values to the estimated Ca\textsuperscript{2+} sensitivities of the biosensors and the fact that both excitation (3580 nm) and emission (3605 nm) peaks are far to the right of the emission peak of the EVFP; moity of the biosensors. Figure 9A and B show changes in XRhod-5F (A) and biosensor ratio (B) simultaneously monitored as Ca\textsuperscript{2+} was titrated to a mixture of biosensor, saturating concentration of CaM. Starting mixture contained 0.25–3 mM BAPTA to guarantee nominally Ca\textsuperscript{2+}-free condition at the beginning of the titration.

Figure 9C shows plots of the fractional responses of BSGPER\textsubscript{83–93}, BSGPER\textsubscript{150–175}, BSGPER\textsubscript{242–259}, and BSGPER\textsubscript{350–351} to saturating CaM concentrations as a function of free Ca\textsuperscript{2+}. The four CaM-binding domains display significantly different Ca\textsuperscript{2+} sensitivities in their interaction with CaM. BSGPER\textsubscript{150–175} has the strongest affinity among the four (Fig. 8), only displays an intermediate Ca\textsuperscript{2+} sensitivity compared to the others (Fig. 9C).

Interestingly, BSGPER\textsubscript{83–93}, which binds CaM with the lowest affinity among the four domains identified (Fig. 8 and Table 1), displays a clearly biphasic Ca\textsuperscript{2+} response curve (Fig. 9C), the majority of which with a much higher Hill coefficient. To rule out a potential complication due to the fairly short distance between the FRET donor and acceptor of this biosensor (11 residues), which might limit access of CaM to the linker, we decided to lengthen the linker by adding three glycine residues to each end of SMD1 in this biosensor configuration, generating BSGPER\textsubscript{GGG83–93GGG}. BSGPER\textsubscript{GGG83–93GGG} binds CaM with essentially identical \( K_d \) value with BSGPER\textsubscript{83–93} (not shown), despite a slightly increased dynamic range due to a longer linker (Fig. 10B vs 10A). Interestingly, simultaneous determinations of BSGPER\textsubscript{GGG83–93GGG} response and free Ca\textsuperscript{2+} values still demonstrated the biphasic behavior (Fig. 10D) identical to that of BSGPER\textsubscript{83–93} (Fig. 10C). These results ruled out the short distance between the FRET pair as an obstacle for CaM access and suggested that the biphasic nature of the Ca\textsuperscript{2+} titration curves is due perhaps to the composition of the SMD1 sequence. Examinations of the response curves in Fig. 10C and 10D show two distinct species of SMD1-CaM complexes with respect to their Ca\textsuperscript{2+} sensitivity. BSGPER\textsubscript{83–93}CaM species 1 are complexes formed at low free Ca\textsuperscript{2+} concentrations (below 1 \( \mu \)M free Ca\textsuperscript{2+}). Fits of this species formation against free Ca\textsuperscript{2+} (blue curve, Fig. 10C) yielded an average EC\textsubscript{50}(Ca\textsuperscript{2+}) value of 0.13 ± 0.02 \( \mu \)M. Analyses of BSGPER\textsubscript{GGG83–93GGG} CaM species 1 (blue curve, Fig. 10D) responses yielded essentially identical value. It is interesting to note that these EC\textsubscript{50}(Ca\textsuperscript{2+}) values are well in the range of resting free cytoplasmic Ca\textsuperscript{2+} concentrations in cells. BSGPER\textsubscript{83–93}CaM species 2 are complexes formed at drastically higher free Ca\textsuperscript{2+} concentrations than species 1. Fits of the entire population formed as a function of free Ca\textsuperscript{2+} values follow the behavior of this more abundant species (black dotted lines, Fig. 10C), which yielded an average EC\textsubscript{50}(Ca\textsuperscript{2+}) value of 3.71 ± 0.13 \( \mu \)M. This value is well within cytoplasmic Ca\textsuperscript{2+} values measured within the sustained phase of vasopressin-induced Ca\textsuperscript{2+} entry in smooth muscle cells [52]. Fits of species 2 formation also yielded significantly higher Hill coefficients than those of species 1. Table 2 summarizes the different EC\textsubscript{50}(Ca\textsuperscript{2+}) values and the Hill coefficients for the four CaM binding domains in their Ca\textsuperscript{2+}-dependent interactions with CaM.
Discussion

Identification of CaM-binding domains in CaM-dependent proteins has traditionally involved synthesis of peptides that correspond to the predicted binding sequence and study of CaM-peptide interaction or of the interactions between CaM and the wild-type protein or a mutant with deletion of the predicted CaM binding sequence. For a G protein-coupled receptor, purification of the entire protein remains a challenge. The lack of an exhaustive approach to identify all CaM-binding domains in a GPCR and determine the Ca\(^{2+}\) sensitivity of their interactions with CaM has made it difficult to fully assess the potential roles of CaM in GPCR biology.

FRET biosensors have been developed based on known CaM-binding sequences to measure free Ca\(^{2+}\)-CaM levels in cells [25,26,28,50,53]. In our opinion, FRET technology offers a simple alternative approach to identify unknown CaM-binding sequences in GPCRs that provides several advantages. First, typical CaM-binding domains are 20 a.a. in length and represent ideal linkers for outstanding FRET efficiency. Using apparent K\(_d\) values and dynamic range of biosensors, a parameter introduced in this paper, the length of linker based on a predicted domain can be easily adjusted until the accurate domain responsible for CaM binding is identified. For GPCRs, the submembrane domains are short fragments that allow for easy screening using this approach. Second, biosensors can be expressed and purified with relative ease and cost effectiveness. This is particularly helpful when there are multiple CaM-binding sites in the same protein, as in the case of GPER. Third, since FRET is proportionate to the inverse 6th power of the molecular distance between the donor and acceptor fluorophores, the technique is very sensitive and therefore allows highly quantitative characterization of the interaction between CaM and its binding domains. Fourth, specific CaM binding to the linker sequence only involves changes in the fluorescence intensities of the donor and acceptor fluorophores and not any shifts in their spectral peaks, facilitating precise quantitation. Finally, the Ca\(^{2+}\) sensitivity of CaM-target interactions can be measured with high precision using the approach described in this paper.

Table 2. EC\(_{50}\)(Ca\(^{2+}\)) values and Hill coefficients for interactions between CaM and GPER’s submembrane domains.*

| Complexes | SMD | EC\(_{50}\)(Ca\(^{2+}\)) (µM) | Hill Coefficient |
|-----------|-----|----------------------------|-----------------|
| CaM-GPER\(_{83-93}\) (Species 1) | 1   | 0.13±0.02                  | 1.99±0.14\*     |
| CaM-GPER\(_{83-93}\) (Species 2) | 2   | 3.71±0.13                  | 2.53±0.15\*     |
| CaM-GPER\(_{150-175}\)          | 2   | 2.38±0.13                  | 1.21±0.09       |
| CaM-GPER\(_{242-259}\)          | 3   | 5.15±0.55                  | 1.43±0.11       |
| CaM-GPER\(_{330-351}\)          | 4   | 0.75±0.05                  | 1.18±0.08       |

Data are means ± SD of at least 3 independent determinations.

*All EC\(_{50}\)(Ca\(^{2+}\)) values are significantly different (p<0.05) from one another; Hill coefficients for GPER\(_{83-93}\) (Species 1) and GPER\(_{83-93}\) (Species 2) are different (p>0.05) from the rest.

\* Significantly different (p<0.05) from GPER\(_{83-93}\) (Species 1).

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Figure 10. CaM titrations and Ca\(^{2+}\) titrations for BSGPER\(_{83-93}\) and BSGPER\(_{GGG83-93GGG}\). (A & B) Typical CaM titrations for BSGPER\(_{83-93}\) (A) and BSGPER\(_{GGG83-93GGG}\) (B). Starting reaction mix contained 0.5 µM biosensor, 3 mM Ca\(^{2+}\), 0.1 mg/mL BSA. (C & D) Aggregate data from Ca\(^{2+}\) titrations for the interactions between CaM and BSGPER\(_{83-93}\) (C) and BSGPER\(_{GGG83-93GGG}\) (D). Starting reactions contained 0.5 µM biosensor, 700 µM CaM, 3 mM BAPTA, 0.5 mM BSA, 2 µM XRhod-5F. Fits for the entire set of data were shown in black dotted lines. Fits for the separate first phase of data (up to measured free Ca\(^{2+}\) values of 1 µM) were shown in blue lines.

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Data are means ± SD of at least 3 independent determinations.

*All EC\(_{50}\)(Ca\(^{2+}\)) values are significantly different (p<0.05) from one another; Hill coefficients for GPER\(_{83-93}\) (Species 1) and GPER\(_{83-93}\) (Species 2) are different (p>0.05) from the rest.

\* Significantly different (p<0.05) from GPER\(_{83-93}\) (Species 1).

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paper. Knowledge of this sensitivity allows prediction of the physiological scenarios that facilitate interactions between CaM and the individual sub-membrane domains of a GPCR. This is particularly true given the constantly trafficking nature of GPCRs through different cellular locales, where the free Ca$^{2+}$ concentrations may vary substantially.

In this study, we have utilized FRET biosensor technique for the first time to demonstrate direct interaction between GPER and CaM and identify four distinct CaM-binding sequences in the submembrane domains of this receptor. The majority of CaM binding domains identified so far in GPCRs are located in either the third submembrane domain or the juxtamembrane section of the fourth. To our knowledge, GPER is the first GPCR identified to possess four CaM-binding domains. Detailed characterizations of biosensor responses indicate that these domains bind Ca$^{2+}$-CaM with significantly different affinities in the sub- to high micromolar ranges. The sensitive nature of the biosensor technique has enabled identification of the role of different fragments in the sequence under study in the interaction with CaM. For example, fragment 150–170 in GPER’s SMD2 binds CaM, but with an apparent affinity 8-fold lower than that of fragment 150–175. This clearly indicates an important role of the basic patch 170–175 in CaM binding to SMD2. The apparent K$_d$ values are obtained in vitro on isolated purified biosensors and, like values obtained using synthetic peptides, are not necessarily the same as in the entire receptor, whose expression and purification remains a challenge. Nevertheless, they provide a useful guide to predict interactions that might occur where GPER resides in its cycle in the cell. With the order in affinity of SMD2 > SMD1 > SMD3 > SMD1, one would predict that SMD2 would be the first to bind CaM in response to a Ca$^{2+}$ signal produced at the receptor’s location in cells. In addition, the identification of these domains provides a basis for future investigations of the roles of CaM in the various interactions at these locations between GPER and other proteins in cells.

As all of these interactions are Ca$^{2+}$-dependent, knowledge of the Ca$^{2+}$ sensitivity of each domain’s interaction with CaM becomes helpful in predicting the physiological scenarios in which each might occur. The concurrent determination of free Ca$^{2+}$ concentrations and CaM-SMD interactions using responses of a suitable Ca$^{2+}$ indicator and FRET biosensor allows highly precise determination of the Ca$^{2+}$ sensitivities of these interactions. We believe that this approach avoids some potential difficulties associated with using Ca$^{2+}$ buffers. One such would be uncertainty in the amount of free Ca$^{2+}$ contained in biosensor stocks or CaM stocks, which would require complete Ca$^{2+}$ chelation of these stocks prior to mixing with buffers, or uncertainly of the effect of dilution from additions of stock solutions. Based on the EC$_{50}$ (Ca$^{2+}$) values determined, SMD2, while apparently being the most avid binder of CaM among the four SMDs, only possesses an intermediate Ca$^{2+}$ sensitivity (EC$_{50}$ (Ca$^{2+}$) ~ 2.3 μM) for its interaction with CaM. Knowledge of the free Ca$^{2+}$ concentrations produced in different physiological scenarios then can be used together with these values in predicting what interactions will take place in cells. Indeed, from our co-immunoprecipitation data, it is obvious that GPER and CaM are in the same complexes even at resting conditions. Based on the condition of the co-IP reaction, this would at first sight seem to be a Ca$^{2+}$-independent interaction; however, based on the Ca$^{2+}$ sensitivity data presented in Fig. 10 and Table 2, it is apparent that SMD1-CaM complexes can be formed at very low free Ca$^{2+}$ concentration (species 1), with an EC$_{50}$ (Ca$^{2+}$) value of ~ 130 nM, a value in the range of resting cytoplasmic Ca$^{2+}$ concentrations in many cell types. So from the standpoint of free Ca$^{2+}$ value, this interaction can certainly occur in unstimulated cells. From the standpoint of available CaM, it is noteworthy that the interaction between CaM and SMD1 is of low affinity. So the question arises as to whether this interaction can occur in cells. It has been demonstrated that CaM is a limiting factor in cells and CaM targets compete for Ca$^{2+}$-dependent interactions with CaM [25,26,28,54]. However, at resting Ca$^{2+}$ levels, the competition among targets for Ca$^{2+}$-dependent interaction with CaM must be much reduced, as only those interactions that are highly sensitive to Ca$^{2+}$, such as that of SMD1-CaM species 1, can take place. A similar case was recently demonstrated, where the interaction between endothelial nitric oxide synthase and CaM, and hence nitric oxide production, can occur at fairly low free Ca$^{2+}$ concentrations (EC$_{50}$ (Ca$^{2+}$) ~ 150 nM) and that phosphorylation of the synthase substantially reduces, albeit by no means negate, the Ca$^{2+}$ dependency of the interactions (EC$_{50}$ (Ca$^{2+}$) ~ 50 nM) [55,56]. These observations have contributed to the explanation of the effects of eNOS phosphorylation or shear stress to increase basal nitric oxide production that previously was thought to be Ca$^{2+}$-independent. In the case of GPER83–93-CaM complex species 1, the biochemical results obtained here lend a nice plausible interpretation of the co-IP data obtained in control cells. In addition, it is noteworthy that GPER has been found in a number of different locations in cells in addition to the plasma membrane [57], including the ER [30] and the trans-Golgi-proteasome network [59]. Under resting conditions, Ca$^{2+}$ values in the Golgi network have been shown to be in the range of 100–250 μM, while that in the ER can reach above 450 μM [60,61]. Given these values and the potential locations of GPER, it is not out of the question that interactions between CaM and the other SMDs of GPER can also occur in unstimulated cells, since the EC$_{50}$ (Ca$^{2+}$) values for these interactions are all in the sub- or low micromolar range (Table 2). Of course, we cannot rule out an alternative possibility that other interaction partner(s) with GPER rather than CaM may also bind CaM with very high Ca$^{2+}$ sensitivity or in a Ca$^{2+}$-independent fashion, and showed up in the GPER pull-down fractions probed for CaM and vice-versa, under resting conditions. Nevertheless, our in-vitro data clearly demonstrate absolute Ca$^{2+}$ dependence of the interactions between CaM and GPER and provide parameters that can sufficiently explain the co-IP data under resting condition.

Under stimulated conditions, it has been shown that vasopressin-induced SR Ca$^{2+}$ release from VSMCs can raise cytoplasmic Ca$^{2+}$ level above 1 μM, while sub-plasma membrane Ca$^{2+}$ can reach 45 μM and remain above 5 μM for a sustained period during subsequent Ca$^{2+}$ entry [32]. Data from Fig. 9C, 10C–D and Table 2 then allow prediction that in addition to SMD1-CaM species 1, SMD1-CaM complexes can easily form in the cytoplasm in response to SR release Ca$^{2+}$ signal, while SMD3 and 2 can interact with Ca$^{2+}$-CaM during the peak and sustained phase of agonist-induced Ca$^{2+}$ entry in VSMCs. In this regard, it is interesting to consider the observation that thapsigargin, which stimulates store-operated Ca$^{2+}$ entry independently of estrogen activity, enhanced the association between CaM and GPER independently of specific ligand binding in VSMCs (Fig. 1). An interesting possibility to investigate from this result is whether a generic cytoplasmic Ca$^{2+}$ signal can activate GPER or alter its function as a G protein-coupled receptor by promoting its multilevel interactions with CaM. Consistent with this idea, it has been demonstrated that experimental shear stress, which triggers Ca$^{2+}$ signals in vascular endothelial cells, causes conformational changes in the B2 bradykinin receptor independently of ligand binding [62].
At the moment we do not have a complete explanation for the biphasic nature of the Ca\(^{2+}\) response curve of SMD1-CaM complex formation. It is possible that sequential binding of the C-terminal lobe and N-terminal lobes of CaM to SMD1 is responsible for this behavior and that the short half-life of GPER’s submembrane domain in intact cells may be the stage such property to be discerned. In a speculative scheme, at low Ca\(^{2+}\) increases, such as during SR Ca\(^{2+}\) release or Ca\(^{2+}\) entry, the N-terminal lobe may now contribute to the interaction, causing a much larger conformational change. Consistent with this possibility, Ca\(^{2+}\) binding to separate N- and C-terminal halves of CaM in the presence of a peptide from the CaM-binding domain of the plasma membrane Ca\(^{2+}\)-ATPase has been shown to be of biphasic nature [63]. These possibilities represent interesting potential for future studies to delineate the biochemical basis for interactions between CaM and SMD1 and their roles in GPER function, especially at basal conditions in cells.

In summary, we have used a novel approach to identify four CaM-binding domains in GPER and characterize properties of their interactions with CaM. Our data demonstrate that this approach can be used with ease to locate CaM binding sequences in GPCRs in particular and perhaps in other proteins in general.

The results of our study suggest that CaM may play a regulatory role in actions of estrogen mediated by GPER. It is interesting to note that the nuclear estrogen receptor ER\(\alpha\) has been demonstrated to be a Ca\(^{2+}\)-dependent CaM-binding protein, whose interaction with CaM stabilizes its dimerization, modulates the interaction with estrogen responsive element and activates transcription [64–69]. Together with these earlier findings, the presence of multiple CaM-binding domains in GPER suggests that CaM is important for estrogen signaling at multiple levels. With the large differences in affinity and Ca\(^{2+}\) sensitivity, the effects of CaM binding to each of the sub-membrane domains on GPER’s functions present an interesting and necessary area of investigations.

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Author Contributions

Conceived and designed the experiments: Q-KT. Performed the experiments: Q-KT MVM. Analyzed the data: Q-KT. Wrote the paper: Q-KT.

References

1. Stampfer MJ, Colditz GA (1991) Estrogen replacement therapy and coronary heart disease: a quantitative assessment of the epidemiologic evidence. Prev Med 20: 47–63.
2. Grady D, Rubin SM, Pettiti DB, Fox CS, Black D, et al. (1992) Hormone therapy to prevent disease and prolong life in postmenopausal women. Ann Intern Med 117: 1016–1037.
3. Edwards DP (2005) Regulation of signal transduction pathways by estrogen and progesterone. Ann Rev Physiol 67: 335–376.
4. Mendelsohn ME, Karas RH (2010) Rapid progress for non-nuclear estrogen receptor signaling. J Clin Invest 120: 2277–2279.
5. Murphy E, Lagahana G, Deschamps A, Kehr M, Nguyen T, et al. (2011) Mechanism of Cardioprotection: What Can We Learn from Females? Pediatr Cardiol.
6. Prossnitz ER, Barton M (2009) Signaling, physiological functions and clinical relevance of the G-protein-coupled estrogen receptor GPER. Prostaglandins Other Lipid Mediat 89: 89–97.
7. Carmeci C, Thompson DA, Ring HZ, Francke U, Weigel RJ (1997) Identification of a gene (GPR30) with homology to the G-protein-coupled receptor superfamily and associated with estrogen receptor expression in breast cancer. Genomics 45: 607–617.
8. Takaya Y, Kato C, Kondo S, Koenaga R, Ando J (1997) Cloning of cDNAs encoding G protein-coupled receptor expressed in human endothelial cells exposed to fluid shear stress. Biochem Biophys Res Commun 240: 737–741.
9. Kvingedal AM, Smeland EB (1997) A novel putative G-protein-coupled receptor expressed in lung, heart and lymphoid tissue. FEBS Lett 407: 59–62.
10. Owman C, Blay P, Nilsson C, Lohal SJ (1996) Cloning of human cDNA encoding a novel heptahelix receptor expressed in Burkitt’s lymphoma and widely distributed in brain and peripheral tissues. Biochem Biophys Res Commun 228: 285–292.
11. O’Dowd BF, Nguyen T, Marchese A, Cheng R, Lynch KR, et al. (1998) Discovery of three novel G-protein-coupled receptor genes. Genomics 47: 310–313.
12. Thomas P, Pang Y, Filardo EJ, Dong J (2005) Identity of an estrogen membrane receptor coupled to a G protein in human breast cancer cells. J Biol Chem 280: 661–662.
13. Levitan IB (1999) It is calmodulin after all! Mediator of the calcium modulation of the parathyroid hormone 1 receptor and a sub-set of class b G-protein-coupled receptor superfamily. Endocrinology 140: 3236–3245.
14. Minakami R, Jinnai N, Sugiyama H (1997) Phosphorylation and calmodulin binding of the cytoplasmic tails of the parathyroid hormone 1 receptor and a sub-set of class b G-protein coupled receptors. FEBS Lett 579: 805–807.
15. Mahon MJ, Shimada M (2005) Calmodulin interacts with the cytoplasmic tails of the parathyroid hormone 1 receptor and a sub-set of class b G-protein coupled receptors. FEBS Lett 579: 805–807.

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