Dimerization Domain of Retinal Membrane Guanylyl Cyclase 1 (RetGC1) Is an Essential Part of Guanylyl Cyclase-activating Protein (GCAP) Binding Interface*

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Background: GCAPs regulate photoreceptor guanylyl cyclase RetGC1 but not hormone receptor guanylyl cyclase NPRA.

Results: Mutations in RetGC1 dimerization domain disrupt GCAP1 and GCAP2 binding.

Conclusion: Met823 in dimerization domain strongly contributes to its specificity in forming GCAP binding interface.

Significance: Congenital blindness-causing mutation in the neighboring residue prohibits GCAP binding.

The photoreceptor-specific proteins guanylyl cyclase-activating proteins (GCAPs) bind and regulate retinal membrane guanylyl cyclase 1 (RetGC1) but not natriuretic peptide receptor A (NPRA). Study of RetGC1 regulation in vitro and its association with fluorescently tagged GCAP in transfected cells showed that R822P substitution in the cyclase dimerization domain causing congenital early onset blindness disrupted RetGC1 ability to bind GCAP but did not eliminate its affinity for another photoreceptor-specific protein, retinal degeneration 3 (RD3). Likewise, the presence of the NPRA dimerization domain in RetGC1/NPRA chimera specifically disabled binding of GCAPs but not of RD3. In subsequent mapping using hybrid dimerization domains in RetGC1/NPRA chimera, multiple RetGC1-specific residues contributed to GCAP binding by the cyclase, but the region around Met823 was the most crucial. Either positively or negatively charged residues in that position completely blocked GCAP1 and GCAP2 but not RD3 binding similarly to the disease-causing mutation in the neighboring Arg822. The specificity of GCAP binding imparted by RetGC1 dimerization domain was not directly related to promoting dimerization of the cyclase. The probability of coiled coil dimer formation computed for RetGC1/NPRA chimeras, even those incapable of binding GCAP, remained high, and functional complementation tests showed that the RetGC1 active site, which requires dimerization of the cyclase, was formed even when Met823 or Arg822 was mutated. These results directly demonstrate that the interface for GCAP binding on RetGC1 requires not only the kinase homology region but also directly involves the dimerization domain and especially its portion containing Arg822 and Met823.

Photoreceptor retinal membrane guanylyl cyclase 1 and 2 (RetGC1 and RetGC2)2 (1–3) produce a secondary messenger of phototransduction, cGMP. RetGCs make complexes with two types of photoreceptor-specific proteins, guanylyl cyclase-activating proteins (GCAPs) (4–8) and retinal degeneration 3 protein (RD3) (9), that bind with high affinity and inhibit RetGC (10, 11). GCAPs are Mg2+/Ca2+-binding proteins (12) that control the shape of the photoreceptor (13–16) through acceleration of cGMP synthesis by RetGC in the light when Ca2+ concentrations in the photoreceptor outer segment decline (17, 18). RD3 protein, which is linked to rare forms of congenital blindness (9), is required for normal accumulation of RetGC in photoreceptors and can form a very tight complex with RetGC (10), effectively blocking both RetGC catalytic activity and activation by GCAPs (11). Mutations in RetGC1 isozyme cause different forms of inherited blindness: loss-of-function mutations cause recessive Leber congenital amaurosis (LCA) (19–21), and gain-of function mutations cause dominant cone-rod degeneration 6 (CORD6) (22–25). The disease-causing mutations are found in various RetGC1 domains defined by their function and homology to other membrane guanylyl cyclases (19, 22). RetGC1 has a single transmembrane region that connects the extracellular domain (ECD) and the cytoplasmic portion that consists of the kinase homology (KHD), dimerization, and catalytic domains (2, 26). The region that includes the KHD and dimerization domain forms the binding interface for both GCAP1 and GCAP2 (27), but the precise location of the side chains that enable the docking site(s) for GCAPs on RetGC1 remains to be identified.

RetGC1 can function only as a homodimer (28, 29) in which the two catalytic subunits complement each other by supplying residues coordinating two Mg2+/GTP molecules in the active site (28). The complex is also likely to contain up to two GCAPs.

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2 The abbreviations used are: RetGC, retinal membrane guanylyl cyclase; CORD6, cone-rod degeneration 6; GCAP, guanylyl cyclase-activating protein; KHD, kinase homology domain; LCA, Leber congenital amaurosis; PCC, Pearson’s correlation coefficient; RD3, retinal degeneration 3 protein; NPR, natriuretic peptide receptor; ECD, extracellular domain; DD, dimerization domain.
molecules, in the case of RetGC1 primarily GCAP1 and in the case of RetGC2 primarily GCAP2 (27, 30, 31). However, when compared with a hormone receptor membrane guanylyl cyclase, such as NPRA (GUCY2A) (26), the role of the dimerization domain in RetGC function appears to be rather intriguing. On the one hand, mutations affecting the coiled coil interactions of the dimerization dimer change the Ca$$^{2+}$$ sensitivity of RetGC1 regulation by GCAP1 (23–25). On the other hand, the role of this domain as being essential for the cyclase activity (which requires the cyclase dimerization) and for binding GCAPs has been disputed (32).

Contrary to the models hypothesized elsewhere (32–34), we recently demonstrated that the docking interface(s) for both GCAP1 and GCAP2 is encoded by a region, Arg$$^{488}$$–Arg$$^{851}$$, containing the KHD and dimerization domain of RetGC1 (27) rather than its C terminus and transmembrane-proximal portions, making two independent docking sites for different GCAPs (33, 35). In the present study, we tested how mutations (site-directed or naturally occurring) specifically in the RetGC1 dimerization domain affected binding of GCAPs. We found that (i) RetGC1-specific residues in the dimerization domain are essential for the ability of cyclase to bind GCAP albeit not essential for the RD3 binding interface; (ii) the ability of RetGC1 to bind GCAP lacking in hormone receptor cyclase, such as NPRA, requires RetGC1-specific residues in the dimerization domain, especially Met$$^{822}$$, the residue that does not constitute a direct contact between the RetGC1 coiled coil domains; and (iii) mutation of Met$$^{822}$$ and LCA-linked mutation of the neighboring Arg$$^{822}$$ destabilize the binding of GCAP1 and -2 but not the binding of RD3.

**Experimental Procedures**

**Guanylyl Cyclase Expression and Mutagenesis—**Human RetGC1 (GUCY2D) and NPRA (GUCY2A) guanylyl cyclases, either tagged with mOrange red fluorescent protein (Clontech) or untagged, were expressed in HEK293 cells as described (27). Mutagenesis was performed utilizing Thermo Scientific Phusion Flash DNA polymerase in a “splicing by overlap extension” approach (36) to produce a series of single amino acid substitutions or replacing portions of RetGC1 sequence with that of NPRA. The amplified fragments were inserted into the DraIII/Clal sites of the modified plasmid, mOrangeRetGC1 plasmid, encoding new restriction sites in the RetGC1 without changing the amino acid sequence (27). The resultant constructs contained mOrange fluorescent tag in the extracellular segment as described in detail previously (27). Oligonucleotide primers were synthesized by Integrated DNA Technologies. Restriction endonucleases were purchased from New England Biolabs.

**Immunostaining—**Following 7% polyacrylamide–SDS gel electrophoresis, the samples were transferred on PVDF membrane using an Invitrogen/Life Technologies iBlot apparatus, stained with Ponceau S to mark positions of the molecular mass standards, and probed with rabbit polyclonal antibodies (37) raised against the Arg$$^{540}$$–Asn$$^{815}$$ human RetGC1 KHD fragment or the Met$$^{797}$$–Ser$$^{1103}$$ catalytic domain fragment (residues are numbered from Met$$^1$$ of the leader peptide encoded by GUCY2D gene). The blots were developed using a Pierce SuperSignal Femto chemiluminescence substrate kit, and the images were taken using a Luminous FX imaging system.

**GCAP and RD3 Expression—**Myristoylated GCAP1 and GCAP2 were expressed from pET11d vector in BLR(DE3) Escherichia coli strain harboring N-myristoyltransferase, extracted from inclusion bodies, and purified using hydrophobic and size exclusion chromatography as described in detail previously (38–40). For the in cyto binding analysis, GCAP1 and GCAP2 were tagged at the C terminus with the SuperGlo enhanced green fluorescent protein (Clontech) and expressed from pQB-IfN3 vector (Clontech) using a Promega FuGENE protocol as described previously (27, 41, 42).

Human RD3 (9) cDNA was amplified with HindIII/EcoRI restriction sites at the ends using as a template MH5101-9206149 Thermo Scientific/Open Biosystems cDNA clone (catalog number 6140075) in high fidelity Phusion Flash polymerase chain reaction mixture and inserted into the HindIII/EcoRI sites of the pQBIfN3 expression vectors to tag RD3 with GFP at the C terminus. RD3-GFP was expressed in HEK293 cells using a FuGENE transfection reagent protocol.

**Transfection for Confocal Imaging—**Unless specified otherwise, HEK293 cells were transfected in a Lab-Tek 4-well cover glass chamber with 1 μg of mOrangeRetGC1 DNA/well using 3 μl/μg DNA of the FuGENE reagent as described (27) at an ~1:100 molar ratio of GCAP-GFP- or RD3-GFP-coding plasmid versus mOrangeRetGC1-coding plasmid. Confocal images were taken after 24 h of incubation utilizing an Olympus FV1000 Spectral instrument using the respective 543- and 488-nm excitation for the red and green fluorochromes and processed using Olympus Fluoview FV10-ASW software as described previously (27, 41, 42). No changes to the original images were made except for occasional minor γ correction applied to the whole image for more clear presentation in print. Quantitative analysis was performed using only original images without γ corrections. Where applicable, Pearson’s correlation coefficient (PCC) for testing co-localization of RD3-GFP with mOrange-tagged RetGC1 variants was calculated using Olympus Fluoview FV10-ASW software as described previously (27, 41, 42), and the statistical difference between the PCC values was tested using the analysis of variance function in Synergy KaleidaGraph 4 software applying Bonferroni post hoc processing.

**RetGC1 Activity Assay—**Human RetGC1 cDNA was expressed in HEK293 cells from a modified pRCCMV vector (Invitrogen) using calcium phosphate precipitation for the transfection, and the membrane fraction containing expressed RetGC1 was isolated as described in detail previously (27, 43). The activity of the cyclase was assayed using [α$$^{32}$$P]GTP (PerkinElmer Life Sciences) as a substrate, and the [$$^{32}$$P]cGMP product was quantified using TLC as described previously (27, 39, 43). Briefly, the assay mixture (25 μl) incubated at 30 °C contained 30 mM MOPS-KOH (pH 7.2), 60 mM KCl, 4 mM NaCl, 1 mM DTT, 2 mM Ca$$^{2+}$/EGTA buffer (<10 mM [Ca$$^{2+}$/free]), 6 mM free Mg$$^{2+}$$, as indicated in the text. 0.3 mM ATP, 4 mM cGMP, 1 mM GTP, and 1 μCi of [α$$^{32}$$P]GTP. The resultant [$$^{32}$$P]cGMP product was separated by TLC on fluorescently backed polyethyleneimine cellulose plates (Merck), developed in 0.2 M LiCl, and eluted with 2 M LiCl. For the
Regulation of Retinal Guanylyl Cyclase

mutants of RetGC1 converted into adenylyl cyclase by mutations in the active site (44), the assay was performed using the same protocol except that 2 mm \([\alpha-32P]ATP\) (PerkinElmer Life Sciences) was used as a substrate instead of GTP, and 3 mm cAMP was the internal standard to identify the product of the reaction on the TLC plate.

Computation of Coiled Coil Probability—The coiled coil structure probability for the whole intracellular segment of RetGC1 and the RetGC/NPRA chimeric protein was computed using conventional MARCOIL software, a hidden Markov model-based program (45) that predicts the existence and location of potential coiled coil domains in protein sequences based on a standard MTIDK matrix, an implementation of the method of Lupas et al. (46).

Results

LCA Mutation R822P in Dimerization Domain of RetGC1 Suppresses GCAP Binding—LCA mutation R822P (20) (residues here and further are numbered from the Met1 of the leader peptide as coded by the wild type RetGC1 cDNA; Ref. 2) disabled RetGC1 activity in vitro by blocking its binding with GCAP1 and GCAP2 (Fig. 1). GCAP complex with the cyclase is known to disintegrate upon extraction of RetGC1 from the membranes with detergents (47). Therefore, the RetGC-GCAP complex cannot be detected using conventional co-precipitation or pulldown assays. However, it can be detected using a cell-based assay in which GCAP1-GFP or GCAP2-GFP expressed in HEK293 cells displays a uniform distribution through the cytoplasm and the nucleus of the cell but becomes anchored to the membranes by the cyclase (primarily to the endoplasmic reticulum; Ref. 41) when co-expressed with either untagged or mOrange-tagged RetGC1 (27, 40 – 42). The tagged cyclase carrying the R822P substitution was neither able to become effectively stimulated by GCAP1 and GCAP2 (similarly to the untagged mutants; see Ref. 20) nor to alter GCAP distribution. A PCC (mean ± S.E.) value of 0.25 ± 0.04 (n = 23), which is below the co-localization criterion threshold (~0.5; Ref. 48), effectively confirmed the lack of strong GCAP1 binding to the mutated cyclase in a sharp contrast to the normal cyclase (PCC = 0.9 ± 0.05, n = 33) (Fig. 1, B and C, and Table 1).

Unlike GCAP1-GFP, RD3-GFP binding to RetGC1 in HEK293 cells remained despite the R822P substitution (Fig. 2). We found that in a fashion similar to that of GCAP1-GFP (41), RD3-GFP displayed a mostly diffuse distribution throughout the cytoplasm and the nucleus when expressed in HEK293 cells (Fig. 2A, top panel) but became anchored to the membranes when co-expressed with the cyclase and void from the nucleus (Fig. 2B). It needs to be pointed, however, that the overall pattern of expressed RD3-GFP in the conditions of our experiments was different from the punctate pattern of RD3 produced in the HEK293 nuclei (9) and/or the cytoplasm (10). Because recombinant RD3 was found to be very prone to self-aggregation (11), we tested whether the variations in the protein expression levels could explain the variability between its distribution patterns reported by different groups. We found that the dose of RD3-GFP DNA used for transfection indeed critically affected the RD3 pattern in HEK293 cells (Fig. 2A). At a low dosage of the RD3-GFP-expressing plasmid (≤0.02 μg of DNA cm\(^{-2}\)), the distribution was mostly uniform throughout the cell (top panel) similar to that of GCAP1-GFP expressed in the absence of RetGC1 (41), whereas at the increased dosage (up to 2 μg of DNA cm\(^{-2}\)), a distinct bright punctate staining of various sizes appeared in both the cytoplasm and the nucleus (Fig. 2A). The granules of the aggregated protein were also observed when a non-tagged RD3 was expressed and probed with anti-RD3 antibody (data not shown). Therefore, it needs to be underscored that to avoid the potential artifacts caused by RD3 self-aggregation, such as shown in the bottom three panels in Fig. 2A, we used a very low dose (≤0.02 μg of DNA cm\(^{-2}\)) (Fig. 2A, top panel) of the RD3-GFP-expressing vector in all subsequent co-transfection experiments designed to probe for the RD3-GFP co-localization with the mOrangeRetGC1. That was the concentration at which RD3 expressed in the absence of
the cyclase was distributed fairly uniformly through the cell (Fig. 2A, top panel) but acquired a sharp membrane-bound, rather than punctate, pattern through its association with the cyclase (Fig. 2B and C) similar to that of GCAP1 (41). The R822P RetGC1 co-localized with RD3-GFP very strongly (PCC = 0.9 ± 0.01, n = 35), thus arguing that the disruption of GCAP binding by the R822P mutation was not a result of a nonspecific unfolding of the cyclase. Hence, altering RetGC1 amino acid sequence in the dimerization domain specifically disrupted the GCAP binding interface.

Replacing RetGC1 Dimerization Domain with That of NPRA Incapacitates Both GCAP1 and GCAP2 Binding—In contrast to RetGC1, the peptide hormone receptor cyclase NPRA does not bind GCAPs (27). However, a chimeric protein containing the KHD and dimerization domain from RetGC1 and catalytic domain of NPRA (construct shown in Fig. 3A and referred to hereafter as “Cat1”) binds GCAP1 and GCAP2 with high apparent affinity, even exceeding that of wild type RetGC1 (Ref. 27 and Fig. 4, A and B). To test whether or not the dimerization domain was essential for creating the GCAP binding interface on the cyclase, we used a chimera with the NPRA sequence extended to include the dimerization domain (“Cat1DD I” chimera in Fig. 3A). The Cat1DD I chimera completely failed to bind GCAP1 and GCAP2 (Fig. 3, B and C, and Table 1). However, Cat1 and Cat1DD I were both able to bind RD3 (Fig. 3D). These results indicated that the dimerization domain of NPRA cyclase did not cause unfolding of the Cat1DD I chimera but rather specifically disrupted its GCAP binding interface.

Consistent with the GCAP binding pattern in cyto, the hybrid cyclase containing the dimerization domain of NPRA in a striking contrast to the Cat1 chimera containing the RetGC1 dimerization domain (Fig. 4, A–C) also completely failed to undergo activation by GCAP1 and GCAP2 in vitro despite the comparable levels of expression in HEK293 membranes for both chimeras (Fig. 4B, inset). Hence, not only the LCA-related mutation but also substitution with a homologous sequence derived from a receptor membrane guanylyl cyclase dimerization domain failed to support RetGC ability to bind either GCAP.

### Table 1

| Co-transfection                | PCC (mean ± S.E.; n cells) |
|-------------------------------|----------------------------|
| GCAP1-GFP + mOrangeRetGC1     | 0.90 ± 0.05; 33            |
| GCAP1-GFP + mOrangeRetGC1 R822P | 0.25 ± 0.04; 23             |
| GCAP1-GFP + Cat1DD I          | 0.30 ± 0.03; 41             |
| GCAP1-GFP + DD II             | 0.39 ± 0.04; 40             |
| GCAP1-GFP + DD III            | 0.91 ± 0.01; 41             |
| GCAP1-GFP + DD IV             | 0.66 ± 0.03; 42             |
| GCAP1-GFP + DD V              | 0.87 ± 0.01; 53             |
| GCAP1-GFP + DD VI             | 0.46 ± 0.02; 27             |
| GCAP1-GFP + DD VII            | 0.52 ± 0.04; 21             |
| GCAP1-GFP + DD VIII           | 0.83 ± 0.01; 40             |
| GCAP1-GFP + DD IX             | 0.49 ± 0.03; 21             |
| GCAP1-GFP mOrangeRetGC1 M823R | 0.55 ± 0.03; 31             |
| GCAP1-GFP mOrangeRetGC1 M823E | 0.52 ± 0.03; 32             |
| GCAP1-GFP mOrangeRetGC1 M823S | 0.89 ± 0.01; 41             |
| GCAP2-GFP mOrangeRetGC1 M823R | 0.40 ± 0.03; 28             |
| GCAP2-GFP mOrangeRetGC1 M823S | 0.87 ± 0.01; 43             |
| RD3-GFP + mOrangeRetGC1 R822P | 0.90 ± 0.01; 35             |
| RD3-GFP + DD hybrid IV        | 0.88 ± 0.01; 21             |
| RD3-GFP + DD hybrid V         | 0.90 ± 0.01; 23             |
| RD3-GFP + DD hybrid VI        | 0.91 ± 0.01; 15             |
| RD3-GFP + mOrangeRetGC1 M823R | 0.91 ± 0.01; 44             |

* The mOrangeRetGC1 and GCAP-GFP or RD3-GFP were co-expressed in HEK293 cells and confocal microscopy was performed as described under “Experimental Procedures.” PCC values indicating strong co-localization are high-lighted in bold (note that PCC ≤ 0.5 generally means no co-localization, whereas PCC = 1.0 means co-localization of all red and green pixels in the image (40)).

b p < 0.0001 when compared with the mOrangeRetGC1; from one-way analysis of variance/Bonferroni all-pairs comparison test (n = 0.01; confidence level, 99%) processed using Synergy KaleidaGraph 4 software.

p < 0.0001, unpaired t test.

### Figure 2

**A** hRD3GFP + DIC

**B** hRD3GFP + mOrangeR822P

**C** hRD3GFP + mOrangeR822P

Consistent with the GCAP binding pattern in cyto, the hybrid cyclase containing the dimerization domain of NPRA in a striking contrast to the Cat1 chimera containing the RetGC1 dimerization domain (Fig. 4, A–C) also completely failed to undergo activation by GCAP1 and GCAP2 in vitro despite the comparable levels of expression in HEK293 membranes for both chimeras (Fig. 4B, inset). Hence, not only the LCA-related mutation but also substitution with a homologous sequence derived from a receptor membrane guanylyl cyclase dimerization domain failed to support RetGC ability to bind either GCAP.

The Part of RetGC1 Dimerization Domain Containing Met823 Has the Strongest Effect in Enabling GCAP1 and GCAP2 Binding.
Regulation of Retinal Guanylyl Cyclase

Binding—The prominent difference between the affinities of Cat1 and Cat1 DDI chimeric cyclases for GCAPs (Figs. 3 and 4, B and C) made it possible to probe for the regions in the dimerization domain that strongly contributed to the unique ability of RetGC1 to associate with GCAPs. The amino acid residues encoded by RetGC1 and NPRA cDNAs (Fig. 4A) were arranged in various hybrid combinations (“DD I–DD VI”) and tested for their activation by GCAP1 (Fig. 4D). Replacement of the C-proximal half of the NPRA dimerization domain (Cat1DD I chimera) was replaced with the catalytic domain of RetGC1 (Cat1) and in the other both the catalytic domain and dimerization domain (Cat1DD I chimera) were replaced with the homologous regions from NPRA; the “+” and the “−” indicate that the construct binds or fails to bind GCAPs in the co-transfection assays, respectively. LP, leader peptide; TM, transmembrane region; CAT, catalytic domain. B and C, unlike WT or Cat1 chimera, the Cat1DD I chimera fails to anchor either GCAP1 (B) or GCAP2 (C) to the membranes in cyto. Note the diffuse pattern of GCAP-GFP spread through the cytoplasm and the nucleus. D, in contrast to GCAPs, the Cat1DDI chimera continues to anchor RD3-GFP to the membranes and prevents its diffusion to the nuclei.

FIGURE 3. Binding of GCAPs but not RD3 by RetGC1/NPRA chimera strongly depends on RetGC1-specific amino acid sequence in dimerization domain. A, schematics representing the domain structures of RetGC1 (blue), NPRA (black), and two chimeric constructs: in one of them (27) the catalytic domain of RetGC1 (Cat1) and in the other both the catalytic domain and dimerization domain (Cat1DD I chimera) were replaced with the homologous regions from NPRA; the “+” and the “−” indicate that the construct binds or fails to bind GCAPs in the co-transfection assays, respectively. LP, leader peptide; TM, transmembrane region; CAT, catalytic domain. B and C, unlike WT or Cat1 chimera, the Cat1DD I chimera fails to anchor either GCAP1 (B) or GCAP2 (C) to the membranes in cyto. Note the diffuse pattern of GCAP-GFP spread through the cytoplasm and the nucleus. D, in contrast to GCAPs, the Cat1DDI chimera continues to anchor RD3-GFP to the membranes and prevents its diffusion to the nuclei.
Subsequent single amino acid substitutions restoring RetGC1-specific residues in DD II, such as S822R (hybrid “DD VII”), R822M (hybrid “DD VIII”), and M824L (hybrid “DD IX”) pointed to Met823 as the residue enhancing the affinity for GCAP1 in the N-proximal portion of the RetGC1 dimerization domain (Fig. 4, A and E). Co-localization assay (Figs. 5 and 6 and Table 1) also demonstrated that the GCAP1 binding pattern was the sharpest with hybrids DD III, V, and VIII (respective PCC values were 0.91, 0.87, and 0.83, which are well above the co-localization threshold of 0.5 (48)). Co-localization was poorly defined for the rest of the tested hybrids (respective PCC values for hybrids DD II, IV, V, and VI were 0.39, 0.66, 0.46, 0.52, and 0.49; Table 1). Importantly, even the hybrids demonstrating a weaker GCAP1 binding pattern (DD hybrids II, IV, V, and VI) retain a normal membrane-anchored pattern regardless of the effectiveness of the hybrid in anchoring GCAP1-GFP. The PCC values for the co-localization are presented in Table 1.

The importance of Met823 in contributing to the strength of the GCAP1 binding interface created by the RetGC1-specific residues was further tested in wild type RetGC1 by a series of single amino acid substitutions at this position. The abilities of the non-chimeric cyclase to become activated by GCAPs (Fig. 7) and to relocate GCAPs to the membranes in the co-transfection assay (Fig. 8) both tolerated substitution with a non-charged polar residue (Ser) but became compromised by the replacement of the Met823 with either negatively charged Glu or positively charged Arg (data for M823E RetGC1/GCAP2-GFP co-expression is not shown due to the space constraint in the figure). Most interestingly, the Arg residue prohibiting effective GCAP1 binding is present in the corresponding position of the dimerization domains from all peptide hormone receptor guanylyl cyclases: NPRA (GUCY2D), NPRB (GUCY2B), and NPRC (GUCY2C) (Fig. 7, top). At the same time, neither Arg823 nor Glu823 (not shown) disrupted the cyclase co-localization with RD3-GFP (Fig. 8, C).

The Ability of the Mutated Dimerization Domains to Promote Coiled Coil Dimerization of the Cyclase—The probabilities of creating the coiled coil structure for the hybrid RetGC1/NPRA chimeras and for RetGC1 harboring point mutations were evaluated using a conventional MARCOIL computation (45, 46) in which the primary structure of the entire intracellular segment of the enzyme downstream of the transmembrane domain was tested for the likelihood of a coiled coil dimerization between two identical protein molecules (Fig. 9). All constructs tested in the present study yielded a high probability of a coiled coil dimer. The lack of the apparent correlation between the probability to create a coiled coil dimer and the extent to which they maintained GCAP binding was quite remarkable for the hybrids DD I–VI (Fig. 9A). For example, DD hybrids I, II, and V, although strikingly different in binding GCAP (see Figs. 3–5 and Table 1), all exceeded 99% probability of forming coiled coil dimer in the dimerization domain between residues 812 and 846.

It needs to be noted that based on the computational analysis in the heptad I helix, which is the most N-proximal of the four heptads in the RetGC1 dimerization domain (44), neither

FIGURE 5. Localization of GCAP1-GFP co-expressed in HEK293 cells with RetGC1/NPRA dimerization domain hybrids II–VI. A, GCAP1-GFP was co-expressed with mOrange Cat1 chimera containing hybrids (top to bottom) DD II–VI. The DD II and DD V hybrids that harbor the RetGC1-specific 822RML824 peptide sequence (bold) retain an obvious GCAP binding pattern, whereas in the other hybrids it has been compromised. B, RD3-GFP co-expressed with (top to bottom) hybrids DD IV, V, and VI retains a normal membrane-anchored pattern regardless of the effectiveness of the hybrid in anchoring GCAP1-GFP. The PCC values for the co-localization are presented in Table 1.

FIGURE 6. Met823 in dimerization domain enhances GCAP1-GFP co-localization with RetGC1/NPRA chimera. GCAP1-GFP was co-expressed with mOrange Cat1 chimera containing hybrids (left to right) DD VII–IX. Note that hybrid DD VIII retains a clear GCAP binding pattern, which is compromised in the hybrids lacking the Met823. The PCC values for the co-localization are presented in Table 1.
Arg822 nor Met823 create the immediate contacts between two coils (positions “a” and “d” in the heptad shown in Fig. 9B). Instead, Met823 occupies position “g” and Arg822 occupies position “f” opposite from the points of contact between the dimerizing coils. The effect of point mutations introduced into wild type RetGC1 was tested and is shown in Fig. 9C. The point mutations failed to negate the coiled coil dimerization by the RetGC1. No evidence of the coiled coil structure being disrupted was found for the Met823 mutants, but for the R822P, the N-proximal portion of the coiled coil appeared partially destabilized: the >90% coiled coil probability region in the R822P RetGC1 spanned the residues 828–855, shorter than the 818–856 region in wild type or Met823 mutants.

Testing Dimerization of RetGC Mutants by Functional Complementation within the Cyclase Homodimer—The ability of the M823R and R822P RetGC1 to dimerize was directly verified in a biochemical assay using a method originally introduced by Ramamurthy et al. (24) by functional complementation between two RetGC molecules whose substrate binding was altered (Fig. 10). RetGC1 is a homodimer (28) whose active site is formed by two complementary subunits converting two GTP substrate molecules into two cGMP molecules (Fig. 10A). Each subunit coordinates the purine base of one GTP molecule and the ribose ring of another GTP molecule (28). Replacement of two residues, Glu925 and Cys997, coordinating the purine base (shown in Fig. 10A, yellow) with Lys and Asp, respectively, changes the substrate specificity of RetGC1 from GTP to ATP (44), thus converting RetGC1 from a guanylyl cyclase into an adenylyl cyclase (construct “GC1AC” in Fig. 10B). A different point mutation, Asp929 (shown in Fig. 10A, red) to Ala, prevents proper ribose coordination and the GTP → cGMP (or ATP → cAMP in GC1AC) catalysis and thus completely inactivates the homodimeric enzyme (24) (Fig. 10B, “GC1AC rib(-)”). Because the ECD portion of RetGC1 is not required for either
catalytic activity or regulation by GCAPs (27, 37, 42), we deleted most of the ECD in both GC1AC and GC1AC rib(−) variants to distinguish them from the full-size RetGC1 by immunoblotting in the subsequent co-transfection experiments. It is worth re-emphasizing that GCAP binding to RetGC1 did not require preservation of the guanylyl cyclase active site. GCAP1-GFP became anchored to the membranes and void from the nuclei when co-expressed in with the GC1AC rib(−) cyclase mutant HEK293 cells (Fig. 10C) similarly to the wild type RetGC1 (41).

The ability of various mutants to utilize GTP versus ATP as a substrate as summarized in Table 2 predicted that GC1AC rib(−) could not display adenylyl cyclase activity unless dimerized with a subunit harboring unchanged Asp929. Indeed, when expressed alone, neither wild type nor GC1AC rib(−) RetGC1 or M823R RetGC1 was able to produce cAMP in the presence of a high concentration of GCAP1 (Fig. 10D). However, when co-expressed with the GC1AC rib(−), the M823R and the wild type both promoted well detectable adenylyl cyclase activity, which could only become possible if they dimerized with GC1AC rib(−) and complemented its ATP-binding active site with the Asp929, the essential residue (Fig. 10A) that GC1AC rib(−) was lacking. In a similar experiment, the R822P RetGC1 was also able to rescue GC1AC rib(−) adenylyl cyclase activity (Fig. 10E).

In a reciprocal experiment, we co-expressed GC1AC with the M823R RetGC1. Membranes isolated from the transfected HEK293 cells were then reconstituted with purified GCAP1 but this time to measure GCAP-stimulated guanylyl cyclase activity (Fig. 10F). Although GC1AC alone could not utilize GTP as a substrate (Table 2 and Ref. 44) and the M823R RetGC1 alone had very low guanylyl cyclase activity due to its poor binding of GCAP1 (see Figs. 7 and 8), the two mutants when co-expressed displayed markedly enhanced cGMP synthesis. That would only be possible if the two mutated cyclases were able to form a dimer in which a portion of the active site (from the M823R RetGC1 subunit) was able to catalyze conversion of GTP, whereas the other subunit, GC1AC, contributed its unabated GCAP1 binding to the cyclase dimer.

**Discussion**

RetGC remains one of the most important but least understood photoreceptor enzymes in rod and cone physiology and disease in part because the data on its three-dimensional structure and organization of the quaternary complex with GCAPs are scarce. A model of the three-dimensional structure for the RetGC1 catalytic domain based on the homology with the known structure of adenylyl cyclase (28) gives a reliable description of its active site and has been directly verified by biochemical experiments (24, 44). However, no other structural data about the rest of the molecule, including the KHD and dimerization domain, is presently available. Likewise, the structure of the Ca\(^{2+}\)-regulated RetGC1-GCAP complex has never been established primarily because the complex could not be stabilized in the presence of a detergent (47). Evidently, regulation of RetGC1 by GCAPs is accompanied by a change in the tightness of the dimer (49), and one can reasonably suggest that changes affecting the RetGC1 homodimer could make the catalysis in the active site more or less efficient by even subtle conformational shifts imparted by the Mg\(^{2+}\) versus Ca\(^{2+}\)-bound GCAPs (50, 51). However, even the exact binding site(s) for GCAPs on RetGC1 has not been unequivocally established despite several prior attempts to assign them to a particular region(s) in the RetGC1 primary structure (32, 35, 52–55). The difficulties in mapping the binding interface for GCAPs exacerbated by the lack of reliable detection of the complex in detergent solution can be effectively mitigated by applying the *in cyto* assay in which the formation of the complex becomes visualized by confocal microscopy using fluorescently labeled GCAPs and RetGC1 (41). This approach was effective in the recent mapping of the RetGC1 binding interface in GCAP1 (42) and in the partial mapping of the GCAP binding interface in RetGC1 (27). The latter revealed that the binding interface for both GCAPs is located in the portion of the RetGC1 molecule that includes both the KHD and the dimerization domain (27). As the first step toward more detailed mapping of the interface, we found here that the dimerization domain strongly contributes...
to the interface. Although some of the earlier studies pointed at the KHD as the possible part of the interface (52, 54, 55), the dimerization domain as a regulatory element in RetGC-GCAP interactions was deemed irrelevant for GCAP2 in a publication describing that GCAP2, but not GCAP1, activated recombinant RetGC1 lacking the dimerization domain (32). We find it difficult to reconcile our findings with the results described in the cited publication because in our study GCAP1 and GCAP2 regulate RetGC in a similar fashion (56), and we show here that both lost the ability to activate the cyclase when its dimerization domain was affected. Our recent study rules out a possibility that the primary binding site for GCAP1 and GCAP2 operates outside the KHD and dimerization domain of the cyclase (27). Furthermore, our experiments presented here clearly demonstrate that GCAP1 and GCAP2 both failed to activate recombinant RetGC1 not only when the entire dimerization domain was replaced by that of a peptide receptor cyclase but even when just a single amino acid residue, Arg$_{922}$ or Met$_{923}$, was altered. Results of RetGC1 mutagenesis combined with the GCAP-dependent activation and in cyto GCAP binding assays conducted in our study strongly argue that the dimerization domain participates in creating and/or directly regulating the GCAP1- and -2 binding interface on RetGC1. The selectivity for GCAP binding enabled by the dimerization domain of RetGC1 in comparison with that of NPRA is attributable to the RetGC1-specific amino acid residues in different portions of the domain but most strongly depends on the presence the N-proximal portion of the domain containing Met$_{923}$ (Figs. 5–8). This residue is evolutionarily conserved between different species (e.g. human, mouse, and bovine) in both RetGC1 and RetGC2 (GUCY2F) in contrast to hormone receptor cyclases GUCY2A (NPRA), GUCY2B, and GUCY2C where it becomes replaced by
TABLE 2

Substrate(s) in active site formed by the cyclase dimer

Conversion of the substrate highlighted in bold to the respective cyclic monophosphate was tested (Fig. 10) to probe for the presence of the corresponding functional cyclase dimer.

| RetGC1 dimer (subunit A/subunit B) | Substrate in active site (subunit A/subunit B) |
|-----------------------------------|-----------------------------------------------|
| mOrangeRetGC1/mOrangeRetGC1<sup>+</sup> | GTP/GTP<sup>+</sup> |
| mOrangeRetGC1/mOrangeRetGC1<sup>+</sup> | APT/ATP<sup>+</sup> |
| GC1AC/GC1AC<sup>+</sup> | Inactive/inactive<sup>a</sup> |
| GC1AC rib(–)/GC1AC rib(–) | GTP/GTP<sup>+</sup> |
| mOrangeM823R/mOrangeM823R<sup>+</sup> | Inactive/ATP<sup>+/-</sup> |
| mOrangeM823R/mOrangeM823R<sup>+</sup> | GTP/ATP<sup>+</sup> |
| mOrangeM823R/mG1AC rib(–)/<sup>+</sup> | Inactive/ATP<sup>+</sup> |
| mOrangeM823R/mG1AC rib(–)/<sup>+</sup> | GTP/ATP<sup>+</sup> |
| mOrangeM823R/mG1AC rib(–)/<sup>+</sup> | GTP/ATP<sup>+</sup> |
| mOrangeM823R/mG1AC rib(–)/<sup>+</sup> | GTP/ATP<sup>+</sup> |
| mOrangeR822P/mOrangeR822P<sup>+</sup> | GTP/GTP<sup>+</sup> |
| mOrangeR822P/mOrangeR822P<sup>+</sup> | Inactive/ATP<sup>+</sup> |
| mOrangeR822P/mG1AC rib(–)/<sup>+</sup> | GTP/ATP<sup>+</sup> |
| mOrangeR822P/mG1AC rib(–)/<sup>+</sup> | Inactive/ATP<sup>+</sup> |
| mOrangeR822P/mG1AC rib(–)/<sup>+</sup> | GTP/ATP<sup>+</sup> |
| mOrangeR822P/mG1AC rib(–)/<sup>+</sup> | Inactive/ATP<sup>+</sup> |

<sup>a</sup> Each subunit of RetGC1 in a dimer coordinates the purine base of GTP and provides coordination of ribose and Mg<sup>2+</sup> for the second Mg<sup>2+</sup> GTP molecule in the active site (Fig. 10A) (28).

<sup>b</sup> RetGC1 with short ECD (37) was converted to adenyl cyclase by the E925K/C997D substitutions as in Ref. 44.

<sup>c</sup> In addition to the E925K/C997D substitutions, the GC1AC also had the D929A mutation (24), disabling coordination of ribose and eliminating catalytic activity.

<sup>d</sup> The purine base of GTP is recognized by the subunit A, but complete subunit B fails to coordinate ribose of the GTP, whereas ATP bound through the purine base by subunit B is held in a normal fashion by the Asp<sup>929</sup> of subunit A (28).

<sup>e</sup> Subunit A cannot utilize Mg<sup>2+</sup> GTP as a substrate because subunit B cannot coordinate the ribose and Mg<sup>2+</sup> of GTP; subunit B coordinates the purine base of ATP, whereas subunit A provides coordination of ribose and Mg<sup>2+</sup> of the ATP (24, 28, 44).

<sup>f</sup> Subunit A coordinates purine base of GTP, and subunit B coordinates that of ATP; each subunit provides coordination of ribose and Mg<sup>2+</sup> for the NTP whose purine base is recognized by the opposite subunit (28, 44).

Arg, a residue that does not support binding of GCAPs (Figs. 7 and 8). This would be consistent with the unique regulatory properties of the photoreceptor RetGC isozymes as Ca<sup>2+</sup>/GCAP-regulated enzymes in contrast to the peptide hormone receptor guanylyl cyclase (27). The hydrophilic non-charged residue in this position of the dimerization domain is required for the cyclase to enable its contact with GCAPs (Figs. 7 and 8).

The RETGC1 DD and especially Met<sup>823</sup> are more likely to be a part of GCAP binding interface than to affect it through the dimerization of the cyclase. First, MARCOIL prediction disfavors Met<sup>823</sup> making a direct contact between the coiled coil structures in heptad I (Fig. 9). Second, computational analysis (Fig. 9) and functional complementation assays (Fig. 10) both demonstrated that the M823R mutation, although strongly disabling GCAP binding, did not preclude RetGC1 from dimerization.

Our finding that the dimerization domain is a part of the GCAP binding interface on the cyclase agrees with earlier observations that mutations of Arg<sup>838</sup> (23–25) or Glu<sup>447</sup> (57) causing dominant CORD6 shift Ca<sup>2+</sup> sensitivity of RetGC1 regulation by GCAPs. There is, however, an important difference. In contrast to Arg<sup>822</sup> and Met<sup>823</sup>, Arg<sup>838</sup> is one of the residues in the corresponding heptad of the RetGC1 homodimer that make immediate contact between the two coils; hence, the CORD6 mutations alter the coiled coil structure itself (24, 57). Therefore, unlike the recessive LCA mutation in Arg<sup>822</sup> (20), which breaks the interface for GCAP and blocks its binding to the cyclase, CORD6 mutations of Arg<sup>838</sup> only alter the relative...
affinities for the Mg\(^{2+}\)-liganded versus Ca\(^{2+}\)-liganded GCAP1 (24, 25). One can speculate that the CORD6-related change of the coiled coil structure alters the accessibility of those residues in the upstream portion of the dimerization domain that contribute to the binding interface for GCAPs and/or directly affect it. The precise mechanism of RetGC1 activation by GCAPs presently remains unclear, but considering that (i) the dimerization domain evidently contributes to the GCAP binding interface, (ii) the overall packing of the cyclase dimer is sensitive to the presence of GCAP (49), (iii) GCAPs drastically increase \( V_{\text{max}} \) of the GTP conversion to cGMP in the active site (1, 56), and (iv) the active site of the cyclase requires dimerization of two catalytic domains (24, 28, 44), it is conceivable that Mg\(^{2+}\)-liganded GCAP bound to RetGC1 via a part of the dimerization domain increases the efficiency of catalysis in the active site by altering the coiled coil structure of the dimer and hence adjusting the orientation of the two catalytic subunits relative to each other.

The R822P mutation linked to LCA (20) also implicates the RetGC1 dimerization domain as an important regulatory region. Unlike Met\(^{823}\), the neighboring residue Arg\(^{822}\) does not define the specificity of GCAP1 binding in RetGC1 versus NPRA, but its replacement with Pro evidently prevents binding (Fig. 1) and subsequent activation of the cyclase by both GCAP1 and GCAP2. A likely explanation for this is that Pro\(^{822}\) despite its not being at the coiled coil contact partially destabilizes the structure of the α-helix at the first heptad of the domain (Fig. 9C).

As yet another intriguing observation from our experiments, M823R RetGC1 and R822P RetGC1, which both poorly bind GCAPs, were both able to sustain the GC1AC rib (−) activation by GCAP in the functional complementation experiments (Fig. 10, D and E). At present, it remains unclear whether one or two GCAP molecules are required to fully activate the cyclase. On the one hand, at saturation by GCAP, the complex is likely to contain two GCAP molecules per two cyclase subunits (30). However, it is presently impossible to conclude whether or not the ratio could be lowered to just one GCAP per complex in the complementation assays and still remain sufficient for activation of the cyclase dimer. On the other hand, because GCAPs themselves are able to dimerize (58), it cannot be excluded that the GCAP dimer is able to regulate the cyclase homodimer even when firmly anchored by only one of the RetGC1 subunits (provided that secondary interaction required for the cyclase regulation is preserved).

The R822P is an LCA-causing mutation (20) that blocks GCAP1 binding but does not disrupt binding of RD3 (Figs. 1 and 2). This argues that (i) R822P does not cause nonspecific unfolding of the cyclase but affects specifically the GCAP binding interface on the cyclase and (ii) the interfaces for the two types of regulator proteins on RetGC1 are not identical. However, there is a strong possibility that the two interfaces can overlap in a tertiary structure of the enzyme based on the following evidence. (i) RD3 and GCAPs have opposite and mutually exclusive effects on RetGC1 activity in recombinant HEK293 cells and in native photoreceptor membranes (11), and (ii) some LCA mutations, such as R768W, evidently block binding of both GCAP1 (30) and RD3 (59) altogether. It needs to be reiterated that the R822P substitution per se does not block RD3 binding. Hence, LCA in that case is unlikely to be a result of an impaired RetGC1 maturation or trafficking to the outer segment membranes, processes in which RD3 has been implicated (10, 59). Hence, in the case of R822P, the LCA it causes is likely a result of impaired activation and/or regulation of the mutated cyclase per se.

It also needs to be emphasized that although our study demonstrates that the cyclase dimerization domain is essential for the GCAP binding, this is only a part of the binding interface formed by the dimerization domain and KHD together (27). The KHD is also an indispensable part of that interface because relative affinities of GCAPs for RetGC1 and RetGC2 depend on the KHD (52) and because deletion and point mutations in KHD can completely disable GCAP binding by RetGC1 (20, 27, 30, 54, 55) even when the dimerization domain remains unaltered. We have verified in a separate experiment that RetGC1-specific sequence of the dimerization domain alone without the KHD could not turn NPRA into a GCAP-activated enzyme (data not shown).

To conclude, our study directly demonstrates that the dimerization domain of the photoreceptor guanylyl cyclase, especially its portion surrounding Met\(^{823}\), is a part of the GCAP binding interface and is therefore essential for the physiological regulation of the cyclase by Ca\(^{2+}\). When mutated, the portion of the interface containing the cyclase dimerization domain can cause LCA by altering normal activation of the enzyme by GCAPs (20). Another portion of the GCAP binding interface that includes the KHD represents a much larger portion of the RetGC1 primary structure than the dimerization domain. Therefore, detailed functional mapping of the residues in the KHD to identify their roles in forming the binding interface for GCAPs remains a major challenge for future studies.

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