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cAMP Signaling Compartmentation: Adenylyl Cyclases as Anchors of Dynamic Signaling Complexes

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ABSTRACT

It is widely accepted that cAMP signaling is compartmentalized within cells. However, our knowledge of how receptors, cAMP signaling enzymes, effectors, and other key proteins form specific signaling complexes to regulate specific cell responses is limited. The multicomponent nature of these systems and the spatiotemporal dynamics involved as proteins interact and move within a cell make cAMP responses highly complex. Adenylyl cyclases, the enzymatic source of cAMP production, are key starting points for understanding cAMP compartments and defining the functional signaling complexes. Three basic elements are required to form a signaling compartment. First, a localized signal is generated by a G protein-coupled receptor paired to one or more of the nine different transmembrane adenylyl cyclase isoforms that generate the cAMP signal in the cytosol. The diffusion of cAMP is subsequently limited by several factors, including expression of any number of phosphodiesterases (of which there are 24 genes plus splice variants). Finally, signal response elements are differentially localized to respond to cAMP produced within each locale. A-kinase-anchoring proteins, of which there are 43 different isoforms, facilitate this by targeting protein kinase A to specific substrates. Thousands of potential combinations of these three elements are possible in any given cell type, making the characterization of cAMP signaling compartments daunting. This review will focus on what is known about how cells organize cAMP signaling components as well as identify the unknowns. We make an argument for adenylyl cyclases being central to the formation and maintenance of these signaling complexes.

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Introduction

The classic model of G protein-coupled receptor (GPCR) signaling, where receptors, G proteins, effector enzymes, and downstream signaling proteins float in a sea of phospholipids and interact with one another stochastically and generate a global signal via a second messenger, does not explain pharmacological reality. If one bathes cells in a catecholamine, such as epinephrine, β-adrenergic receptors are activated and a cAMP signal is generated in the cytosol. However, cAMP is a soluble messenger that can diffuse everywhere, theoretically making the location of its generation irrelevant. Moreover, cAMP can elicit hundreds of different cellular responses based on protein kinase A (PKA) phosphorylation events and activation of exchange protein directly activated by cAMP (Schmidt et al., 2013). Recent efforts using RNA sequencing to define the full range of GPCR expressed in a given cell type reveal that most cells express approximately 100 different heptahelical receptors (Insel et al., 2015). Thus, the pharmacological reality is that at least 20–30 of these receptors couple Gαs, stimulate adenylyl cyclase (AC) activity, and promote cAMP production. The classic model of cAMP signaling does not differentiate Gαs-coupled responses from different GPCR agonists. Ligands, such as prostaglandin E2 (PGE2), that activate Gαs-coupled receptors generate cAMP in the cytosol and should lead to the exact same set of cellular responses as epinephrine, but they do not. Most will readily agree that a model where epinephrine and PGE2 yield the same cellular response is not teleologically supported, and in fact experimental observations debunking this idea date back nearly 40 years (Brunton et al., 1979; Hayes et al., 1979). Indeed, the idea that cAMP signaling is compartmentalized has been readily accepted for decades but has never been incorporated into the general model of cAMP signaling. In fact, the first description of cAMP signaling compartments was made in 1983 by Ian Buxton and Larry Brunton (Buxton and Brunton, 1983).

The arrangement of GPCR signaling proteins to create and maintain strict cAMP compartments is poorly understood,

ABBREVIATIONS: AC, adenylyl cyclase; AKAP, A-kinase-anchoring protein; β1AR, beta-1 adrenergic receptor; β2AR, beta-2 adrenergic receptor; GPCR, G protein-coupled receptor; HASM, human airway smooth muscle; PDE, phosphodiesterase; PGE2, prostaglandin E2; PKA, protein kinase A; PKC, protein kinase C.
Despite the concept of cAMP signaling compartmentation being both palatable and not new. Examples of specific physiologic responses mediated by distinct cAMP signaling compartments are also equally lacking. The three basic elements required to define cAMP signaling compartments are localized signal generation, restricted diffusion of the signal, and localized signal response elements. All cells express a number of different proteins that perform each of these functions in cAMP signaling. This review will discuss these three elements and illustrate the evidence that various signaling proteins contribute to cAMP signaling compartments in different cell models.

**Localized Signal Generation**

GPCRs are the first step in the cascade leading to second messenger generation and are critical elements in determining the nature of the subsequent localized cAMP signal. The initial organization of cell responses to activation of any given GPCR is based on AC partnering with the receptor within discrete signaling microdomains. While GPCRs can be quite mobile with respect to different plasma membrane domains and even intracellular organelles (depending on the nature of the receptor and its level of activity), AC isoforms appear quite steady in their localization. A primary distinction between isoforms of AC is their residence in caveolar/lipid raft microdomains or in nonraft plasma membrane domains (Ostrom and Insel, 2004; Dessauer et al., 2017). For example AC1, AC3, AC5, AC6, and AC8 associate with lipid rafts, which in many cells also contain one or more forms of caveolin. On the other hand, AC2, AC4, AC7, and AC9 are found within nonraft plasma membrane microdomains. ACs within these microdomains appear constant, with no differences among species, cell types, activation states, or even caveolin expression (Ostrom and Insel, 2004; Cooper and Tabbasum, 2014). The availability of $G_{q}$ appears to be in excess of both GPCR and AC and relatively uniform in its distribution across membrane microdomains (Post et al., 1995; Ostrom et al., 2001).

AC localization to lipid raft or nonraft complexes correlates with GPCR coupling with specific ACs, implying that the juxtaposition of receptor and effector with a single microdomain is necessary for efficient coupling. Specific GPCR/AC coupling has been elucidated using knockdown or overexpression of different AC isoforms. For example, AC6 overexpression selectively enhances beta-2 adrenergic receptor ($\beta_2$AR) signaling in airway smooth muscle, lung fibroblasts, and neonatal cardiac myocytes, but not signaling from other G$_{q}$/$\alpha$-coupled receptors (Ostrom et al., 2000, 2002; Liu et al., 2008; Bogard et al., 2011). In contrast, prostaglandin E2 receptors in human airway smooth muscle (HASM) specifically colocalize with and couple to AC2 in nonraft domains based on AC2 overexpression studies (Bogard et al., 2012). Table 1 shows what is known about the localization and GPCR coupling of each of the nine transmembrane forms of AC. Also noted are the cells that have been used for these studies and notes on other key pieces of information. What is clear from Table 1 is that most ACs are significantly understudied in terms of their receptor coupling.

Cardiac myocytes also appear to exhibit some correlation between AC localization and GPCRs. There is evidence that AC5 is preferentially expressed in the T-tubules, while AC6 is found in the peripheral sarcolemma (Timofeyev et al., 2013). Furthermore, beta-1 adrenergic receptors ($\beta_1$ARs) have been reported to couple selectively to AC5 in neonatal cardiac myocytes (Tsunematsu et al., 2015). On the other hand, in adult cardiac myocytes, both $\beta_1$ARs and $\beta_2$ ARs are found in T-tubules, where they couple to AC5, while $\beta_2$ARs alone are found in the peripheral sarcolemma, where they couple specifically to AC6 (Nikolaev et al., 2010; Timofeyev et al., 2013). However, Cros and Brette (2013) found that $\beta_2$AR couples in the peripheral sarcolemma, but not in the T-tubules. Thus, GPCRs display AC-specific coupling that reflects receptor colocalization to AC isoforms in specific plasma membrane environments that can give rise to different cellular responses.

Numerous other GPCRs show differential coupling to AC activity based on localization to specific membrane domains. D1 dopamine receptors are coupled to lipid-raft-dependent AC5, AC6, and AC6 activity, whereas D5 dopamine receptors interact with AC5 in a nonraft localized manner (Yu et al.,

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**TABLE 1**

Microdomain localization and receptor coupling of mammalian AC isoforms

<table>
<thead>
<tr>
<th>AC Isoform</th>
<th>Microdomain Localization</th>
<th>GPCR Coupling</th>
<th>Systems Studied</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>AC1</td>
<td>Lipid raft</td>
<td>$\beta_3$AR</td>
<td>Vascular smooth muscle cells</td>
<td>AC1 overexpression used to demonstrate GPCR coupling</td>
</tr>
<tr>
<td>AC2</td>
<td>Nonraft</td>
<td>EP$_{2\alpha}$R</td>
<td>Airway smooth muscle cells</td>
<td>AC2 overexpression used to demonstrate GPCR coupling</td>
</tr>
<tr>
<td>AC3</td>
<td>Lipid raft</td>
<td>$\beta_2$AR, D$_1$R glucagon</td>
<td>HEK-293</td>
<td>Glucagon receptors were exogenously expressed</td>
</tr>
<tr>
<td>AC4</td>
<td>Nonraft</td>
<td>$\beta_1$AR, $\beta_2$AR, A$_{2a}$R</td>
<td>Unknown</td>
<td>Cardiac myocytes, hepatocytes</td>
</tr>
<tr>
<td>AC5</td>
<td>Lipid raft</td>
<td>Unknown</td>
<td>HEK-293, cardiac myocytes, airway and GI smooth muscle cells, vascular smooth muscle cells, cardiac fibroblasts, lung fibroblasts, hepatocytes, platelets</td>
<td>D$_1$R couple to AC5 in nonrafts of renal tubule cells</td>
</tr>
<tr>
<td>AC6</td>
<td>Lipid raft</td>
<td>$\beta_1$AR, $\beta_2$AR, IPR, D$<em>1$, A$</em>{2a}$R, PAC$_1$</td>
<td>HEK-293, pancreatic cells, hippocampal neurons</td>
<td>AC6 overexpression, knockdown, and antibody inactivation have been used to determine GPCR-AC6 coupling</td>
</tr>
<tr>
<td>AC7</td>
<td>Nonraft</td>
<td>D$_1$R-EP$_1$R</td>
<td>Unknown</td>
<td>Regulation of AC7 by D$<em>1$R is via G$</em>{q}$, and requires EP2R</td>
</tr>
<tr>
<td>AC8</td>
<td>Lipid raft</td>
<td>None</td>
<td>HEK-293, pancreatic cells, hippocampal neurons</td>
<td>Stimulated by store-operated and L-type calcium channels</td>
</tr>
<tr>
<td>AC9</td>
<td>Nonraft</td>
<td>Unknown</td>
<td>Unknown</td>
<td></td>
</tr>
</tbody>
</table>

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GPCR and its cognate AC. For example, PGE2 may regulate coexpressed GPCRs can even influence AC activity of another receptor stimulation is inhibitory in early pregnancy and becomes stimulatory after midterm pregnancy (Zhou et al., 2000). This change in AC activity is probably due to an upregulation of uterine AC2 expression (Zhou et al., 2007). Coexpressed GPCRs can even influence AC activity of another GPCR and its cognate AC. For example, PGE2 may regulate neuronal dopamine signaling through EP1 receptors that facilitate dopamine D1 activation of cAMP production in an AC7/G protein-dependent manner (Ehrlich et al., 2013).

A structural component must exist in all ACs that confers affinity for one domain or the other since AC isoforms are either lipid raft or nonraft localized. Likely mechanisms that stabilize these interactions include sequence-specific protein-protein or protein-lipid interactions, and post-translational modifications. Evidence for the critical role of the intracellular C1 and C2 domains in targeting ACs to lipid rafts come from studies with chimeras of raft and nonraft isoforms (Crosswhait, et al., 2005) and approaches examining fragments of ACs (Thangavel et al., 2009). These studies show AC isoforms have key protein-protein interactions that confer their lipid raft localization. Protein-lipid interactions may also be critical for raft targeting of ACs. Mutation of N-linked glycosylation sites on an extracellular loop of AC8 causes this lipid-raft localized isoform to appear in nonraft membranes (Pagano et al., 2009). Mutation of similar glycosylation sites in AC6 also causes nonraft targeting (unpublished data). Other known components of lipid raft microdomains, such as caveolin, appear to not have a direct functional role in the lipid raft localization of ACs. Cell lines without caveolin-1, whether by knockout approaches or naturally absent, do not show differences in AC expression or localization (Thangavel et al., 2009).

Nine isoforms of AC exist in the human genome and each isoform has long been known to display specific regulatory properties (Dessauer et al., 2017). AC isoform-specific regulation can be mediated by a diverse array of intracellular components including Gβγ proteins, calcium-calmodulin, protein kinases, and nitric oxide (Tang and Gilman, 1992; Sunahara et al., 1996; Hanoune and Defer, 2001; Patel et al., 2001; Cooper, 2003). Any one particular intracellular regulator can differentially influence the direction of AC activity depending on the isoform. For example, Gβγ subunits stimulate AC2 and AC4; inhibit AC1, AC5, and AC6 (Gao and Gilman, 1991; Tang and Gilman, 1991; Taussig et al., 1993; Bayewitch et al., 1998); and produce conditional stimulation of AC5 or AC6 based on differences in the structural homology of each isoform (Thomas and Hoffman, 1996; Gao et al., 2007). Most AC isoforms are stimulated by protein kinase C (PKC) except for AC6, which is inhibited (Jacobowitz et al., 1993; Kawabe et al., 1994; La et al., 1997). Likewise, calcium-calmodulin stimulates AC1, AC3, and AC8 isoforms and AC5 and AC6 are inhibited by the presence of divalent calcium (Tang et al., 1991; Choi et al., 1992; Katsushika et al., 1992; Yoshimura and Cooper, 1992; Cali et al., 1994). The differential effects of other regulatory proteins of AC isoforms continue to be identified including annexin A4, a calcium-dependent phospholipid binding protein that inhibits AC activity in HEK293 and cardiomyocyte cells (Heinick et al., 2015).

The localization of GPCRs and specific ACs to lipid rafts is also responsible for coupling capacitive calcium entry with calcium-dependent AC isoform activity (Cooper et al., 1995; Fagan et al., 1998, 2000; Smith et al., 2002). The importance of calcium-sensitive AC isoforms, such as AC6 or AC8, and their proximity to capacitive calcium entry channels is evident since AC coupling is lost if lipid rafts are disrupted. Thus, regulation of AC isoforms and their downstream cellular functions are also influenced by positioning within different cAMP signaling compartments (Ostrom et al., 2012; Cooper and Tabassum, 2014). Therefore, both AC isoform-specific regulatory properties and cAMP signaling compartments must be considered as additional dimensions that regulate cAMP effects in a spatiotemporal manner specific to cell type and external stimuli conditions.

Restriction of Signal Diffusion

Generating an intracellular signal in a specific locale has little consequence if the signal freely diffuses through the cell. Rich et al. (2000) described some of the first data showing membrane-delimited cAMP signals that were not detected in the cytosol. Ample evidence has emerged since then that cAMP does not freely diffuse inside cells (Agarwal et al., 2016; Richards et al., 2016); however, the manner in which this second messenger’s movement is restricted remains poorly defined. One can readily imagine that both phosphodiesterase (PDE) activity and physical barriers to diffusion could play a role. Experimental data and modeling of cell signaling in cardiac myocytes reveals that PDE activity cannot account for all of the restricted diffusion of cAMP (Saucerman et al., 2014). Thus, long-distance diffusion of cAMP through a cell must reckon with a variety of factors including the buffering effects of PKA, structural components of the cell that are impervious to small molecules in the cytosol, and enzymes that catalyze its breakdown (Yang et al., 2016).

Davare et al. (2001) were among the first to describe that β2AR signal in a preassembled complex to regulate L-type Ca2+ channels. While no PDE (or A-kinase-anchoring protein [AKAP]) was specifically identified in these early studies, it was clear that a membrane-delimited cAMP signaling event occurred due to formation of specific signaling complexes that contained a GPCR and a downstream effector protein. PDEs possess the essential qualities to play roles in maintaining compartmentalized cAMP signaling (Kokkonen and Kass, 2017). In the cAMP degradation realm, most work has focused on PDE4 (Houslay and Adams, 2003). PDE4D binds to β- arrestin, meaning it can be brought into close association with recently activated GPCRs (Bailie et al., 2003). β-arrestin-PDE4 recruitment regulates β-adrenergic receptors switching from Gs to Gi and coupling to the mitogen-activated protein kinase pathway (Bailie et al., 2003; Lynch et al., 2005). PDEs also bind to certain AKAP isoforms to target their enzymatic activity to specific signaling complexes (discussed in more detail subsequently).
Localization of Downstream Effectors

The organization of the GPCR-AC signaling is not limited to just the regional localization of receptor and effector elements based on raft or nonraft compartments. The ramification of cAMP responses also critically depends on the spatial organization of the elements that link PKA activity to a cellular response. The systematic way in which the cAMP signal is sculpted in various cell types to produce differential response has been primarily attributed to the apposition of AC and attendant PDE isoforms (Houslay and Milligan, 1997). However, this view is quickly being expanded with an understanding of the role of AKAPs to direct cAMP actions. AKAPs organize PKA signaling with a wide variety of other molecules via their defining PKA binding motif (Colledge and Scott, 1999). AKAPs distribute regional signals throughout the cell by scaffolding PKA to its intracellular substrates involved in various signaling pathways. These critical functions of AKAPs can be anchored to the site of cAMP generation (Kapiloff et al., 2014). Several diverse members of the AKAP family interact with ACs in an isoform-specific manner and regulate cAMP signaling (Bauman et al., 2006). To date, the specific AKAP-AC interactions have been defined are AKAP9/Yotiao with AC1, AC2, AC3, and AC9 (Piggott et al., 2008); mAKAP with AC2 and AC5 (Kapiloff et al., 2009); and AKAP79/150 with AC2, AC3, AC5, AC6, AC8, and AC9 (Efendiev et al., 2010; Delint-Ramirez et al., 2011; Shen and Cooper, 2013).

The taxonomy of AC-AKAP complexes and their roles in AC regulation continues to be a focus of ongoing research (Dessauer, 2009). The AKAP9/Yotiao complex illustrates the differential role AKAP complexes can have on how associated ACs operate. For example, AKAP9/Yotiao promotes normal functioning of AC1 and AC9, whereas it inhibits activity of AC2 and AC3 upon association (Piggott et al., 2008). Another AKAP, AKAP5 (AKAP79/150), decreases the sensitivity of AC8 to Ca++, while it can also interact with AC5 and AC6 via β3ARs (Bauman et al., 2006; Willoughby et al., 2010). In this configuration PKA associates and phosphorylates β3AR to initiate desensitization, receptor translocation, and even G protein switching (Daaka et al., 1997). PKA can then phosphorylate AC5 and AC6 to inhibit activity or alter AC8 sensitivity to store-operated, AKAP5-mediated calcium entry (Beazely and Watts, 2006; Delint-Ramirez et al., 2011). Therefore, one of the main roles of an AKAP is to provide a substrate for feed forward and backward regulation of the cAMP signaling cascade.

AKAPs also regulate AC activity via PKC phosphorylation. In general, PKC activity will increase activity of certain ACs (AC1, AC2, AC3, AC5, and AC7) but inhibit the activity of others (AC6) (Sunahara et al., 1996; Lai et al., 1997). For example, muscarinic receptor activation of cAMP production occurs via recruitment of PKC to a complex containing the PKC-stimulable AC2, which is organized specifically by AKAP5, with PDE4 and PKA as contributing elements in the complex (Shen and Cooper, 2013). α-Amino-3-hydroxy-5-methyl-4-isoxazolopropionic acid–type glutamate receptors in mouse forebrain slices depend upon AKAP5 anchoring both PKA and AC (Zhang et al., 2013). Mutations of the KCNQ1 subunit of the slow outward potassium channel interrupts its binding to Yotiao, causing loss of PKA regulation of the channel and alteration in the action potential that causes
arrhythmias (Marx et al., 2002). Other AKAP-AC complexes have other clear pathophysiological implications (Efendiev and Dessauer, 2011).

AKAPs also bind PDEs. As discussed previously, AKAP9 brings PDE4D3, but not PDE4D5, to a complex with cardiac IKs channels to create tightly integrated signaling (Terrenoire et al., 2009). Previous studies have shown that mAKAP binds PDE4 to facilitate a negative feedback loop by which local cAMP would activate mAKAP-tethered PKA, which in turn would phosphorylate colocalized PDE4 and upregulate its cAMP hydrolyzing activity (Dodge et al., 2001). This operational model was further expanded by findings that AKAP12 (Gravin) also binds PDE4D to allow it to selectively regulate near-membrane cAMP signaling events (Willoughby et al., 2006). Guinzberg et al. (2017) recently described that adenosine A2A receptors couple to an AC6/AKAP79/PDE3A complex in hepatocytes, while in these same cells A2B receptors couple to an AC5/D-AKAP2 complex. The interplay between ACs, AKAPs, PKA, and PDEs highlights the manifold intracellular signaling circuitry that can be organized to elicit specific, tuned responses throughout different cell types. However, much work remains to fully characterize these complexes and understand how they shape cAMP signaling in differentiated cells.

**From a Signal to a Response**

cAMP signaling compartmentation is of little consequence if the separate signaling pools do not yield distinct cellular responses. We, and others, have described that AC6 has the unique ability to regulate arborization, a form of cytoskeletal reorganization, in HASM cells (Gros et al., 2006; Bogard et al., 2012). AC2 overexpressed in the same cells is unable to mediate the arborization response unless PDE4 activity is inhibited. We have also recently described that overexpression of different AC isoforms yields drastically different gene expression responses. Overexpression of AC2 or AC6 in HASM cells and subsequent stimulation by forskolin leads to gene expression changes that are heterogeneous (Bogard et al., 2013). For example, interleukin 6 production in airway smooth muscle cells is directed by AC2-generated cAMP, whereas somatostatin production was shown to be activated by AC6-derived cAMP (Bogard et al., 2013). In vascular smooth muscle cells and HEK-293 cells, AC1 selectively slowed cell proliferation while AC2, AC5, and AC6 had little effect (Gros et al., 2006). These data suggest the cellular responses depend on the loci of cAMP production within the cell and provide rationale for continuing efforts to define the signaling complexes that lead to cAMP signaling compartmentation in differentiated cells. However, cAMP can elicit hundreds of potential responses in a given cell and we know very little about how many different responses any given localized pool of cAMP can affect.

**Conclusions**

Compartmentation of cAMP signaling allows different GPCRs to use the same second messenger to regulate a unique array of cellular responses. AC isoforms appear to anchor these cAMP signaling compartments since their localization does not change across cell types or activation states. AC isoforms couple to specific GPCRs, synthesize cAMP in distinct subcellular locations, and bind to many other proteins that together can form large signaling complexes. AKAPs are of particular importance because they form specific interactions with GPCRs, ACs, and downstream signaling elements (targets of PKA phosphorylation); cAMP regulatory enzymes (PDEs); and probably other key signaling proteins. However, there are likely other proteins that play critical roles in scaffolding these signaling molecules together. Proteomic and other unbiased approaches are needed to identify the whole range of proteins participating in these signaling complexes. The result of the formation of these multiprotein complexes is that each compartment can have a tailored magnitude, duration, and scope of the cAMP signal generated. These compartmentalized signals are likely dynamic and contextual, as well as cell specific. The proximal elements of two main cAMP signaling compartments in HASM cells have been defined; therefore, we present a diagram that illustrates some of what is known (see Fig. 1). Understanding how these signaling complexes form, what responses they mediate, and how they may be altered in disease should lead to new therapeutic strategies that have far greater efficacy for the desired response with fewer unwanted effects.

**Fig. 1.** Schematic diagram of proposed cAMP signaling compartments in HASM cells. Two main cAMP signaling compartments have been defined, one centered around β2AR-AC6 in lipid raft domains (consisting of sphingolipid, shown in red, and cholesterol) and another around EP3/4R-AC2 in nonraft domains (consisting of phospholipid, shown in blue). Illustrated are the main AKAP and PDE isoforms known to assemble in these signaling complexes or regulate cAMP emanating from each location. Cytoskeletal elements likely also form barriers to cAMP diffusion. Cell responses that have been linked to specific compartments are listed. A number of AKAP and PDE isoforms are expressed in HASM but the locations of some are unknown, and thus are not shown. These include PDE3, AKAP2, AKAP3, ezrin, and Map2B. It is also unknown if PDE4D splice variants have different localizations. PDE8A is located in lipid rafts where it regulates β2AR-AC6 signaling, but it is unknown how it is tethered specifically in this location. AKAP12 (Gravin), AKAP5 (AKAP79), AKAP9 (yotiao).


