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The subcellular dynamics of GPCR signaling

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Abstract

G protein-coupled receptors (GPCRs) are the largest family of membrane receptors and mediate the effects of a multitude of extracellular cues, such as hormones, neurotransmitters, odorants and light. Because of their involvement in numerous physiological and pathological processes and their accessibility, they are extensively exploited as pharmacological targets. Biochemical and structural biology investigations have clarified the molecular basis of GPCR signaling to a high level of detail. In spite of this, how GPCRs can efficiently and precisely translate extracellular signals into specific and well-orchestrated biological responses in the complexity of a living cell or organism remains insufficiently understood. To explain the high efficiency and specificity observed in GPCR signaling, it has been suggested that GPCR might signal in discrete nanodomains on the plasma membrane or even form stable complexes with G proteins and effectors. However, directly testing these hypotheses has proven a major challenge. Recent studies taking advantage of innovative optical methods such as fluorescence resonance energy transfer (FRET) and single-molecule microscopy have begun to dig into the organization of GPCR signaling in living cells on the spatial (nm) and temporal (ms) scales on which cell signaling events are taking place. The results of these studies are revealing a complex and highly dynamic picture, whereby GPCRs undergo transient interaction with their signaling partners, membrane lipids and the cytoskeleton to form short-lived signaling nanodomains both on the plasma membrane and at intracellular sites. Continuous exchanges among such nanodomains via later diffusion as well as via membrane trafficking might provide a highly sophisticated way of controlling the timing and location of GPCR signaling. Here, we will review the most recent advances in our understanding of the organization of GPCR signaling in living cells, with a particular focus on its dynamics.

Keywords

GPCR, signal compartmentalization, nanodomains, FRET, single-molecule microscopy

1 1. Introduction

2 During evolution from simple unicellular to complex multicellular organisms, cells have developed 3 increasingly sophisticated strategies to sense the extracellular environment and communicate with 4 each other. The large superfamily of G protein-coupled receptors (GPCRs) arguably represent the most successful result of this amazing evolutionary endeavor, which enabled our cells to sense and 5 6 decode a large number of extracellular cues, encompassing light, odorants, hormones and 7 neurotransmitters (Pierce et al., 2002; Lefkowitz, 2004). Given their accessibility, diversity and well-8 defined pharmacology, GPCRs have also served as major drug targets. As a result, at least one third 9 of all drugs currently on the market target these receptors (Hauser et al., 2017). Moreover, there is 10 large scope for further development since only a fraction of all potentially targetable GPCRs are 11 currently exploited for pharmacological purposes.

12 Given the fundamental biological role and importance of GPCRs as drug targets, all major steps in GPCR signaling have been intensively investigated. By as early as the late 70's, pioneering work on 13 14 the mechanisms of hormone action initiated by Earl Sutherland and Ted Rall in the late 50's and later taken over by Alfred G. Gilman and Martin Rodbell had already clarified that hormones like 15 16 adrenaline and glucagon act via binding to a specific receptor located on the plasma membrane, 17 triggering the activation of G proteins in a process that requires GTP, ultimately leading to the 18 production of cyclic AMP (cAMP) by adenylyl cyclase – for a historical perspective see (Beavo and 19 Brunton, 2002). In 1986, the sequence of the β_2 -adrenergic receptor was elucidated by the group of 20 Robert Lefkowitz, revealing an unexpected similarity with rhodopsin (Dixon et al., 1986). More 21 recently, there has been enormous progress in the clarification of the structural basis of GPCR 22 signaling, with the successful elucidation of the three dimensional structures of several GPCRs in 23 different conformational states (Cherezov et al., 2007; Rasmussen et al., 2007) as well as in complex 24 with both G proteins (Rasmussen et al., 2011; Koehl et al., 2018; Liang et al., 2017; Zhang et al., 25 2017) and arrestins (Kang et al., 2015; Zhou et al., 2017).

26 Despite these tremendous advances, we are just beginning to understand how GPCRs function and 27 are regulated within the complexity of an intact cell or organism to produce specific effects. Indeed, 28 recent data, mainly obtained with innovative microscopy approaches, indicate that GPCRs are far 29 more complex and dynamic than previously thought, being able to signal at distinct signaling 30 nanodomains both at the cell surface and on intracellular membranes. Understanding this complexity is going to be crucial to answer fundamental and still unresolved questions such as how 31 32 GPCRs can produce specific effects and might pave the way to innovative pharmacological 33 approaches.

34

35 **2. Early evidence for signal compartmentalization**

Whereas the highly successful model of GPCR signaling derived from the early biochemical studies 36 37 described well the general mechanisms of GPCR signaling, it soon emerged that such a model was 38 insufficient to fully explain the effects observed in intact cells and tissues - for a comprehensive 39 review see (Beavo and Brunton, 2002; Steinberg and Brunton, 2001). For example, already in early 40 studies in perfused hearts it was noted that whereas both epinephrine and prostaglandin E1 induce 41 similar increases of cAMP and activate protein kinase A (PKA), only stimulation of adrenergic 42 receptors with epinephrine was able to significantly activate glycogen phosphorylase, increase heart 43 contractility and induce troponin I phosphorylation (Keely, 1979; Brunton et al., 1979). Subsequently 44 it was found that the β -adrenergic agonist isoproterenol but not PGE1 increases the amount of 45 cAMP and PKA activity in the particulate fraction of rabbit heart lysates, which mainly contain type-II 46 isoforms of PKA (Hayes et al., 1980). These and similar findings obtained in isolated cardiomyocytes 47 led Buxton and Brunton to hypothesize that β -adrenergic and PGE1 receptors might induce cAMP 48 accumulation and PKA activation in distinct subcellular microdomains, leading to different biological 49 effects (Buxton and Brunton, 1983).

50 In parallel, experiments suggested that adenylyl cyclases and other signaling proteins might not be 51 randomly distributed on biological membranes. For instance, pioneering work by Tolkovsky and 52 Levitzki provided indirect evidence that on turkey erythrocyte membranes adenosine receptors 53 might be pre-coupled to adenylyl cyclases (note that the role of G proteins was not known at that 54 time), whereas β -adrenergic receptors would activate adenylyl cyclases by random collision 55 (Tolkovsky and Levitzki, 1978; Tolkovsky and Levitzki, 1978).

Several early studies concentrated on the role of lipid domains on the plasma membrane. These 56 57 studies suggested that GPCR signaling might preferentially occur in lipid rafts and caveolae (Insel et 58 al., 2005). Lipid rafts were initially identified biochemically as small (micrometer-sized) sphingolipid-59 and cholesterol-rich membrane domains that are resistant to detergent extraction at low temperature (Simons and Ikonen, 1997). A fraction of these domains contains also caveolins and, 60 61 morphologically, corresponds to small invaginations of the plasma membrane, known as caveolae, 62 which can be visualized by electron microscopy (Simons and Toomre, 2000). Based on biochemical 63 evidence, several groups proposed that receptors, G proteins and adenylyl cyclases preferentially 64 accumulate within lipid rafts and caveolae, suggesting a possible functional role for the resulting 65 inhomogeneous distribution of GPCRs and their signaling partners on the plasma membrane (Insel et 66 al., 2005). For instance, it has been proposed that, in cardiomyocytes, β_2 -adrenergic receptors are 67 preferentially localized in caveolae and T-tubules, which have a membrane composition similar to 68 caveolae, whereas β_1 -adrenergic receptors would be mainly excluded from these structures (Xiang et al., 2002; Nikolaev et al., 2010). This has been suggested to play an important role in determining the specificity of signaling downstream of β_{1^-} vs. β_{2^-} adrenergic receptors, which have distinct biological effects on cardiomyocytes. In particular, the spatial proximity of β_{2^-} adrenergic receptors, G_s proteins and adenylyl cyclases in caveolae has been suggested to be required for β_{2^-} adrenergic receptors to produce physiological responses (Xiang et al., 2002; MacDougall et al., 2012; Wright et al., 2014).

75

76 **3.** New imaging approaches to address an old problem

77 Although the concept of signal compartmentalization is now widely accepted and supported by 78 growing evidence, there has initially been considerable resistance to accept a non-random 79 distribution of signaling molecules, possibly due to the popularity of the Singer and Nicolson's fluid mosaic model of the plasma membrane (Singer and Nicolson, 1972). Moreover, directly 80 81 demonstrating the existence of signaling domains in living cells has proven challenging. A major 82 problem was the lack of adequate tools to localize signaling events in living cells, as the classical 83 biochemical and pharmacological methods typically used in these studies require cell disruption and 84 have no spatial and very low temporal resolution. These limitations have been at least partially 85 overcome by the introduction of innovative microscopy methods that allow scientists to directly 86 visualize receptor signaling in living cells. A first major advance was represented by the introduction 87 of reporters based on fluorescence resonance energy transfer (FRET) (Milligan and Bouvier, 2005; 88 Lohse et al., 2012; Lefkimmiatis and Zaccolo, 2014; Calebiro and Maiellaro, 2014). These methods 89 were crucial to provide direct evidence for the existence of signaling domains on the plasma 90 membrane as well as inside cells (Calebiro et al., 2010; Irannejad et al., 2013; Irannejad et al., 2017; 91 Godbole et al., 2017; Surdo et al., 2017; Castro et al., 2010; Maiellaro et al., 2016). Among other 92 findings, this revealed that despite cAMP being a small water-soluble molecule, cAMP and PKA 93 signals can be highly confined on the plasma membrane as well as at other intracellular 94 compartments (Surdo et al., 2017; Castro et al., 2010; Maiellaro et al., 2016). This organization is 95 likely particularly relevant in highly specialized cells such as cardiomyocytes and neurons, where 96 neurotransmitter and hormone signals have to be rapidly converted into coordinated cellular 97 responses such as synaptic plasticity or heart contraction.

More recently, the rapid development of innovative methods based on single-molecule microscopy has allowed probing the organization and dynamics of GPCR signaling nanodomains with unprecedented spatiotemporal resolution – for a detailed review see (Calebiro and Sungkaworn, 2017)). These methods hold great promise to directly study the dynamic organization of GPCR signaling on the spatial and temporal scales where GPCR signaling events are taking place.

103

104 **4.** The importance of the cytoskeleton

105 The cytoskeleton that is closely associated with the plasma membrane is often termed membrane 106 skeleton. It consists of actin filaments, microtubules, and associated proteins. The membrane 107 skeleton is somewhat different from the bulk cytoskeleton: it interacts with the plasma membrane 108 and proteins that are located at the plasma membrane, playing a role in numerous cellular functions, 109 such as endocytosis and exocytosis. Moreover, it provides anchors for the localization of 110 transmembrane proteins. The membrane skeleton covers almost the entire cytoplasmic surface of 111 the plasma membrane, and is intimately associated with clathrin-coated pits and caveolae (Morone 112 et al., 2006). Based on results of early single-particle tracking (SPT) experiments with gold 113 nanoparticles (Sako and Kusumi, 1994) and optical tweezers (Edidin et al., 1991; Sako and Kusumi, 114 1995), it was proposed that the membrane skeleton partitions the plasma membrane, influencing the diffusion of membrane-associated molecules (Jacobson et al., 1995; Kusumi et al., 2005). These 115 116 and later measurements with fluorescently labelled proteins indicated that membrane molecules 117 are temporarily confined into membrane compartments of 40-300 nm, consistent with the size of the membrane skeleton mesh (Murase et al., 2004). By observing membrane proteins and lipids 118 119 embedded in the plasma membrane, it was found that they both undergo so called 'hop' diffusion, 120 characterized by alternating phases of free diffusion and transient confinement (Suzuki et al., 2005; 121 Fujiwara et al., 2002; Hiramoto-Yamaki et al., 2014). 'Hop' diffusion has been demonstrated for 122 different types of membrane receptors, including GPCRs, which were investigated in a pioneering single-particle tracking study by the group of Akihiro Kusumi (Suzuki et al., 2005). In this study, 123 124 tracking of μ -opiod receptors labelled with gold nanoparticles at the impressive temporal resolution 125 of 25 µs revealed that these receptors jump between adjacent membrane compartments, in which 126 they are temporally trapped. These findings led to the formulation of the 'fence-and-picket' model 127 of the plasma membrane. According to this model, the membrane skeleton ('fences') and integral 128 membrane proteins associated with it ('pickets') provide physical barriers to the diffusion of both 129 membrane proteins and lipids, leading to compartmentalization of the plasma membrane in small 130 nanodomains (Fujiwara et al., 2002; Kusumi et al., 2011). Importantly, the confinement of 131 interacting molecules in such nanodomains has been suggested to increase the probability of their 132 encounters, thus potentially increasing the rate of biochemical reactions (Saxton, 2002).

The cytoskeleton does not only provide barriers to receptor diffusion, but also provides anchor points for receptors and other membrane proteins, further contributing to the formation of signaling nanodomains and controlling their spatial arrangement on the plasma membrane. This has been probably best studied for ionotropic receptors at chemical synapses. It has been shown that

137 ionotropic glutamate receptors are constantly diffusing within the plasma membrane, which 138 promotes the exchange of receptors between synaptic and extrasynaptic sites (Triller and Choquet, 139 2003). However, the local entrapment of receptors at postsynaptic densities is essential for synaptic 140 function. This is achieved via a network of molecular interactions with the membrane skeleton and 141 associated transmembrane proteins (Sheng and Sala, 2001; Garner et al., 2000). For instance, single-142 particle tracking in living cells and super-resolution imaging in fixed cells have revealed that the 143 nanoscale localization and lateral mobility of α -amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid 144 (AMPA) receptors can greatly influence synaptic transmission (Compans et al., 2016). In the case of 145 AMPA receptors, the interactions with the cytoskeleton have been shown to be mediated by 146 stargazin (also known as TARP γ 2), an auxiliary subunit of the AMPA receptor, which, in turn, 147 interacts with the postsynaptic density protein 95 (PSD95). PSD95 is a scaffold that plays a crucial 148 role in the organization of post-synaptic densities. This involves interactions with the actin 149 cytoskeleton mediated by binding of PDS95 to α -actinin (Matt et al., 2018). Since the interactions 150 between AMPA receptor and stargazin are transient, this allows a dynamic exchange of AMPA 151 receptors between synaptic and extrasynaptic sites (Bats et al., 2007). Moreover, it has been shown 152 that glutamate stimulation decreases the stargazin-mediated immobilization of AMPA receptors at 153 post-synaptic sites, allowing a faster replacement of desensitized receptors with new ones. Thus, 154 this might provide a mechanism to fine tune synaptic sensitivity to repeated stimulation.

155 Although the mechanisms responsible for the organization of GPCRs on the plasma membrane are 156 less understood than in the case of ionotropic receptors, there is evidence that also GPCRs might 157 interact with the actin cytoskeleton. For example, single-molecule experiments in simple cell models 158 have shown that $GABA_{B}$ receptors undergo dynamic interactions with the actin cytoskeleton, likely 159 mediated by an as yet unknown scaffold, causing their preferential arrangement along actin fibers 160 (Calebiro et al., 2013). As in the case of AMPA receptors, agonist stimulation with GABA was also 161 found to weaken the interaction of GABA_B receptors with the cytoskeleton, leading to an increase of 162 their lateral mobility. Whereas the occurrence and functional relevance of this organization in 163 neurons is presently unknown, it is tempting to speculate that it might be involved in controlling the 164 precise localization and/or function of $GABA_{B}$ receptors at synapses (Calebiro et al., 2013). Another 165 receptor that has been recently investigated in this respect is the somatostatin receptor type 2 166 (SSTR2), which had been shown to interact with the acting-binding scaffold filamin A (Peverelli et al., 167 2014). Recent single-molecule data by our group indicate that SSTR2 undergoes transient 168 interactions with filamin A, which lead to a preferential localization of SSTR2 along actin fibers and 169 participate in restraining SSTR2 diffusion on the plasma membrane. These interactions are increased

- ¹⁷⁰ by agonist stimulation and are apparently required for efficient SSTR2 recruitment to clathrin-coated
- ¹⁷¹ pits and internalization in response to agonist stimulation (Treppiedi et al., 2018).
- 172

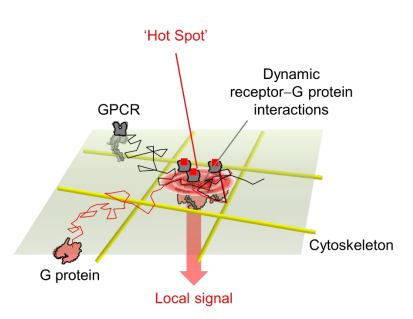
173 **5. Hot-spots for GPCR signaling on the plasma membrane**

174 As mentioned above, despite several lines of evidence suggesting the possible existence of GPCR 175 signaling nanodomains on the plasma membrane, their demonstration has proven extremely 176 challenging. In fact, it has been only recently, with the further development of methods based on 177 single-molecule and super-resolution microscopy that their direct visualization has become possible. 178 These methods are not only able to resolve the organization of receptors and their signaling partners 179 but can also localize downstream signaling events. For instance, in a recent elegant study, Mo et al. 180 developed a new type of sensors - based on changes in the fluorescence fluctuation of the 181 fluorescent protein TagRFP-T when in close proximity to another fluorescent protein called Dronpa -182 that allowed them to resolve PKA activity on the plasma membrane of living cells with a resolution 183 up to three times better than the diffraction limit (Mo et al., 2017). This revealed the existence of 184 PKA signaling nanodomains, which likely result from PKA clustering at the plasma membrane 185 mediated via interaction with PKA anchoring proteins such as AKAP79.

186 More recently, our group succeeded for the first time in directly visualizing individual receptors and 187 G proteins as they diffuse, interact and signal on the surface of intact cells (Sungkaworn et al., 2017). 188 This study – which focused on α_{2A} -adrenergic receptor/G_i and β_2 -adrenergic receptors/G_s as model 189 receptor/G protein pairs – has led to a number of important observations. Notably, we could directly 190 measure the duration of receptor-G protein interactions in intact cells, showing that they are 191 transient and last approximately 1-2 seconds. Whereas we observed the occurrence of transient 192 receptor-G protein interactions also in the absence of ligands, which were linked to the basal 193 constitutive activity of the receptors, we did not observe preformed, stable receptor-G protein 194 complexes, as proposed by some previous studies. However, differences might exist in the degree 195 and stability of pre-association among different receptors and/or G proteins. For instance, using 196 bioluminescence resonance energy transfer (BRET) and time-resolved fluorescence resonance 197 energy transfer (FRET), the protease-activated receptor 1 (PAR1) was found in a previous study to 198 pre-associate with G_i but not with G_{12} (Ayoub et al., 2010). Second, we found that agonists mainly act 199 by increasing the association rate (k_{on}) between receptors and G proteins, as expected in the case of 200 protein interactions that proceed through major conformational changes. Third, we discovered that 201 receptor-G protein interactions and G protein activation do not occur randomly on the plasma 202 membrane, but rather at dynamic nanodomains that we termed 'hot spots' (see Figure 1). Based on 203 our recent results and simulations, we hypothesize that, by increasing the local effective

204 concentration of receptors and G proteins, these hot spots increase the speed and efficiency of their 205 interactions, while allowing GPCR signals to occur locally. These data help to clarify one of the most 206 fundamental and debated aspect of GPCR signaling, i.e. whether receptors are pre-coupled to G 207 proteins or interact with them via random collisions. At least in principle, stable receptor complexes 208 could permit fast and local signaling, but at the expense of signal amplification. In contrast, pure 209 random coupling would favor signal amplification but would also bring low speed and efficiency. By 210 establishing a sort of 'dynamic pre-coupling' - i.e. via allowing transient receptor-G protein 211 interactions in the basal state, and keeping the involved receptors and G proteins near to each other 212 thanks to the barriers provided by the cytoskeleton – nature seems to have found the ideal balance 213 between signal amplification and speed. These and similar mechanisms might play a crucial role in 214 determining the high efficiency and specificity observed among GPCRs and might provide a means of 215 controlling GPCR signaling in space and time.

216



217

Figure 1. 'Hot spots' for GPCR signaling on the plasma membrane. Dynamic interactions between receptors, G proteins and barriers provided by the cytoskeleton lead to the formation of dynamic nanodomains on the plasma membrane that increase the efficiency of G protein activation, while allowing GPCRs to induce local signals.

222

6. Lipid nanodomains revisited

In spite of the biochemical evidence supporting the existence of lipid rafts, the size, dynamics and even existence of lipid domains on the plasma membrane has proven hard to demonstrate in living cells. This has sparkled an intense debate on the exact nature, role and functional relevance of lipid rafts (Munro, 2003; Eggeling et al., 2009; Eggeling, 2015). Traditionally, it was assumed that lipid rafts are rather stable plasma membrane compartments containing sphingolipids, cholesterol and a unique set of resident (mainly GPI-anchored) proteins which float as 'rafts' on the plasma membrane
and might provide specialized platforms for receptor signaling (Simons and Ikonen, 1997). This idea
was mainly based on the fact that rafts could be isolated as detergent-resistant lipid patches and the
observation of phase separation in model membranes (Eggeling, 2015). However, the most recent
imaging studies that attempted to directly visualize lipid rafts in living cells failed to detect stable
lipid domains on the plasma membrane (Eggeling et al., 2009).

235 Early attempts to directly visualize the spatial arrangement of fluorescently labelled GPI-anchored 236 proteins by fluorescence microscopy in living cells showed fairly homogenous membrane staining, 237 suggesting that either lipid rafts did not exist in vivo or were smaller than the lateral resolution of 238 conventional fluorescence microscopy, which is about 200 nm (Mayor and Maxfield, 1995). The 239 latter hypothesis was supported by FRET measurements suggesting that GPI-anchored proteins may 240 associate in clusters smaller than 70 nm (Varma and Mayor, 1998). A subsequent study by the same 241 group further narrowed down the estimated size of such clusters to less than 5 nm and a maximum 242 of 4 GPI-anchored proteins per cluster (Sharma et al., 2004).

As a complementary approach, several groups have measured the diffusion of fluorescently labelled 243 244 lipids and GPI-anchored proteins by either single-particle tracking or fluorescence correlation 245 spectroscopy (FCS) in an attempt to understand their spatiotemporal organization at the plasma 246 membrane. The results have shown that both GPI-anchored proteins and lipids undergo transient 247 confinement in small nanodomains on the plasma membrane (Fujiwara et al., 2002; Lenne et al., 248 2006). However, multiple factors could contribute to such behavior, complicating the interpretation 249 of the results. A first important factor is the cytoskeleton, which, as mentioned above, has been 250 shown to provide barriers to both protein and lipid diffusion in the plasma membrane. However, 251 there is some evidence that the confinement of membrane lipids and GPI-anchored proteins is not 252 always dependent on the presence of an intact cytoskeleton and could be altered by manipulating 253 the lipid composition of the plasma membrane, supporting a role for lipid-protein interactions in the 254 transient confinement of GPI-anchored proteins (Lenne et al., 2006).

255 To more precisely measure the residency time of fluorescent lipids in membrane nanodomains, the 256 groups of Stefan Hell and Christian Eggeling have developed novel approaches based on stimulated 257 emission depletion (STED) microscopy, which enable them to control and narrow down the size of 258 the excited volume in FCS measurements (Eggeling et al., 2009; Honigmann et al., 2014). This 259 allowed them to investigate the diffusion of membrane molecules on different spatial scales. 260 Overall, their results indicate that sphingolipids exhibit transient arrests as they diffuse on the 261 plasma membrane, which are most likely caused by transient interactions with immobile or slow-262 diffusing membrane proteins. These trapping events last for approximately 10 ms and occur in areas

that are smaller than the resolution of the employed method, which is about 20 nm. These interaction sites are stable over a few seconds, during which they do not seem to diffuse within the plasma membrane. This behavior seems to be specific for sphingolipids, and to a much lesser extent phosphoethanolamine, with no correlation with the preference of the investigated lipid analogs for liquid-ordered membrane environments. Whereas these observations further support the occurrence of short-lived lipid–protein complexes, they seem to rule out the classical model of lipid rafts as stable and ordered lipid patches floating within the plasma membrane.

Another point to consider is that whereas lipid phase separation might affect protein dynamics on the plasma membrane, protein–protein interactions, such as those leading to clustering of GPIanchored proteins during their activation, might also favor the formation of larger and possibly more stable lipid–protein complexes or nanodomains (Kusumi et al., 2012).

Overall, these recent data point to a much more dynamic picture than previously imagined, whereby membrane proteins and lipids undergo transient interactions that might still be relevant for the spatiotemporal organization of receptor signaling, but which do not necessarily lead to the formation of stable lipid–protein domains.

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279 7. GPCR signaling at intracellular domains

280 Whereas signaling by GPCRs has long been thought to be restricted to the plasma membrane, a 281 growing body of evidence indicates that GPCRs can also signal on intracellular membranes. The use 282 of advanced optical methods such as FRET and conformation-sensitive biosensors was not only 283 instrumental for these recent discoveries but also allowed scientists to identify the intracellular 284 compartments where GPCR signaling is taking place. A first study by our group on endogenous 285 thyroid stimulating hormone (TSH) receptors in thyroid cells revealed that these prototypical GPCRs 286 for glycoprotein hormones remain active after internalization, leading to persistent cAMP signaling 287 (Calebiro et al., 2009). Similar results were obtained by the group of Jean-Pierre Vilardaga studying the parathyroid hormone (PTH) receptor (Ferrandon et al., 2009). These early studies provided the 288 289 first demonstration that GPCRs can continue signaling via cAMP at intracellular sites after 290 internalization. Moreover, they suggested that TSH and PTH receptors were probably signaling in 291 distinct compartments, i.e. a perinuclear compartment associated with the Golgi complex and early endosomes, respectively. However, it was only with the introduction of biosensors based on 292 293 conformation-sensitive nanobodies that it was possible to directly visualize the subcellular sites of 294 receptor and G protein activation in living cells. Using this elegant approach, the group of Mark von 295 Zastrow was able to show that β_2 -adrenergic receptors remain active in early endosomes, where 296 they induce local G₅ protein activation (Irannejad et al., 2013). More recently, our group further

297 investigated the nature and dynamics of the intracellular compartment where TSH receptors are 298 signaling. For this purpose, we used a combination of FRET sensors measuring cAMP levels and PKA 299 activity, which we tethered to different subcellular compartments, together with the previously 300 developed conformation-sensitive biosensor for G_s protein activation (Godbole et al., 2017). The 301 results of this study demonstrated that upon TSH stimulation, the TSH receptor and its ligand traffic 302 retrogradely to the trans-Golgi network, where they induce local Gs protein activation, cAMP 303 production and PKA activation. Importantly, this leads to a delayed phase of cAMP/PKA signaling at 304 the Golgi/trans-Golgi network, which is required for TSH to efficiently induce phosphorylation of the 305 cAMP response element-binding protein (CREB) and transcription of early genes (Godbole et al., 306 2017). Interestingly, signaling within the Golgi complex has also been recently demonstrated for the 307 β_1 -adrenergic receptor, even though in this case it has been proposed that it is the ligand (adrenalin) 308 to reach the receptors, which are already located on membranes of the Golgi complex, via facilitated 309 transport across cellular membranes (Irannejad et al., 2017). In the meantime, studies form several 310 groups have demonstrated signaling at intracellular sites for a number of receptors (Kotowski et al., 311 2011; Feinstein et al., 2013; Kuna et al., 2013; Merriam et al., 2013; Ismail et al., 2016; Lyga et al., 312 2016), suggesting that this might be a rather common feature among GPCRs. Moreover, there is 313 some evidence that GPCRs might also signal on the nuclear envelope (Tadevosyan et al., 2012) as 314 well as in mitochondria (Hebert-Chatelain et al., 2016; Suofu et al., 2017).

315 These novel and exciting findings point to a previously unsuspected level of complexity in GPCR 316 signaling. The fact that individual GPCRs can induce local signals in distinct subcellular compartments likely plays a major role in determining their biological effects. This provides a new basis to explain 317 318 the high diversity found within the GPCR superfamily, in spite of the fact that all these receptors 319 converge on just a few common signaling pathways. At the same time, they reveal an extremely 320 dynamic picture. Indeed, data on both TSH (Godbole et al., 2017) and β_2 -adrenergic receptors 321 (Irannejad et al., 2013) indicate that GPCR signaling is highly controlled in space and time during 322 receptor internalization and intracellular trafficking, apparently occurring in short 'bursts' once 323 receptors enter well-defined membrane sub-domains of early endosomes or the trans-Golgi 324 network. This high degree of integration between intracellular trafficking and signaling likely 325 provides a key mechanism to fine tune GPCR signaling, which, once understood in its complexity, 326 might also offer novel opportunities to modulate GPCRs for therapeutic purposes.

327

328 8. Concluding remarks

Altogether, the new findings obtained with advanced optical methods in living cells are deeply changing our views on the spatiotemporal organization of GPCR signaling cascades. Above all, they

331 have revealed a highly complex and dynamic picture, whereby GPCRs can rapidly form transient signaling nanodomains on the plasma membrane as well as at intracellular sites (Figure 2). We are 332 333 only beginning to characterize these nanodomains, investigate which factors lead to their formation and understand their impact on GPCR signaling. As it is often the case, the development of 334 335 innovative methods with increased power and resolution, has been instrumental for these 336 discoveries. The current rapid progress in the fields of single-molecule and super-resolution 337 microscopy combined with a growing awareness of the need for large multidisciplinary efforts to 338 tackle biological complexity is likely to offer novel, exciting opportunities in the near future to 339 further investigate the mechanisms and relevance of the spatiotemporal dynamics found in GPCR 340 signaling.

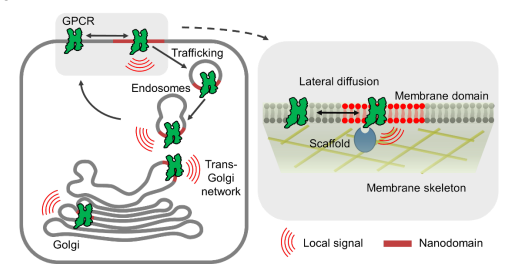




Figure 2. Dynamic nanodomains for GPCR signaling. Recent studies have revealed a highly complex and dynamic picture, whereby GPCRs can signal at dynamic nanodomains located both on the plasma membrane and on membranes of intracellular compartments such as early endosomes, the trans-Golgi network or the Golgi complex. Dynamic interactions of receptors and other signaling proteins with membrane lipids, the cytoskeleton and protein scaffolds likely play important role in the formation of these nanodomains. Lateral diffusion and trafficking control the localization of GPCRs within the nanodomains, which might provide a means to modulate the timing and location of GPCR signaling.

349

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