Signaling Microdomains Define the Specificity of Receptor-Mediated InsP₃ Pathways in Neurons

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Summary

M₁ muscarinic (M₁AChRs) and B₂ bradykinin (B₂Rs) receptors are two PLC_β-coupled receptors that mobilize Ca²⁺ in nonexcitable cells. In many neurons, however, B₂Rs but not M₁AChRs mobilize intracellular Ca²⁺. We have studied the membrane organization and dynamics underlying this coupling specificity by using Trp channels as biosensors for real-time detection of PLC_β products. We found that, in sympathetic neurons, although both receptors rapidly produced DAG and InsP₃ as messengers, only InsP₃ formed by B₂Rs has the ability to activate IP₃Rs. This exclusive coupling results from spatially restricted complexes linking B₂Rs to IP₃Rs, a missing partnership for M₁AChRs. These complexes allow fast and localized rises of InsP₃, necessary to activate the low-affinity neuronal IP₃R. Thus, these signaling microdomains are of critical importance for the induction of selective responses, discriminating proinflammatory information associated with B₂Rs from cholinergic neurotransmission.

Introduction

Coordination of cellular functions resides in the ability of a cell to convert extracellular stimuli into appropriate cellular responses. These cellular responses are dictated by the nature of the signal transduction pathway initiated by the receptor. However, in any individual cell, the number of signaling proteins appears limited, and often these proteins are molecular targets for multiple plasma membrane receptors. A persistent dilemma therefore is how receptors that are coupled to identical transducing proteins and second messenger systems can produce different output signals. One possibility is that specificity results from differences in spatial organization and dynamics in the receptor-transduction systems. Structural mechanisms for the spatial organization of some G protein-coupled receptors (GPCRs) are known (e.g., Drosophila photoreceptors and metabotropic glutamate receptors) (Hardie and Raghu, 2001; Fagni et al., 2000; Xiao et al., 2000), but there is no comparable information about the differential organization and dynamics of GPCRs activating the same effector system within a single neuron.

Here, we explore the question of how different neurotransmitter receptors that utilize one common pathway-the phospholipase C (PLC) pathway-can induce different responses in the same neuron. The PLC pathway provides a ubiquitous mode of transduction in eukaryotic cells that regulates fundamendal cell functions as diverse as gene expression, exocytosis, neuronal signaling, cell growth, and differentiation and is the focal point for signal transduction triggered by G proteincoupled receptors and tyrosine kinase receptors (Berridge, 1993, 1998). Active PLC catalyses the breakdown of phosphatidylinositol biphosphate into the second messengers diacylglycerol (DAG) and inositol-1,4,5-triphosphate (InsP₃). DAG is the physiological activator of the different members of the protein kinase C family. while InsP₃ mobilizes Ca²⁺ from internal IP₃R-stores.

M₁ muscarinic (M₁AChR) and B₂ bradykinin (B₂R) receptors are two G protein-coupled receptors that activate PLC. M₁AChRs play a key role in regulating neuronal excitability (Caulfield, 1993; Brown et al., 1997; Hamilton et al., 1997), whereas B₂Rs mediate inflammation and hyperalgesia (Dray and Perkins, 1993; Dray, 1997). In nonexcitable cells, stimulation of either of these two receptors leads to PLC-mediated production of InsP₃, which triggers release of Ca2+ through endoplasmic reticulum-associated InsP₃ receptors (IP₃R) (Felder, 1995; Lee and Rhee, 1995). Surprisingly, however, in neurons, M1AChRs appear to be relatively weak mobilizers of intracellular Ca2+. A typical example is found in rat sympathetic neurons, which express both M1AChRs and B₂Rs (Marrion et al., 1989; Bernheim et al., 1992; Jones et al., 1995; Hamilton et al., 1997). While stimulation of either receptor can activate PLC sufficiently to increase the formation of InsP₃ (del Río et al., 1999; Bofill-Cardona et al., 2000) and DAG, with consequent mobilization of protein kinase C (Marsh et al., 1995), only B₂Rs produce a significant rise in intracellular Ca²⁺ (Cruzblanca et al., 1998; del Río et al., 1999). This implies that only the B₂R efficiently couples to the endoplasmic IP₃R, and raises the question, what membrane organization of receptors and transducing molecules might underly this differential coupling?

In the present study, we have tried to answer this question by dynamically tracing receptor-mediated PLC signals in individual neurons. We have developed the use of exogenously expressed mammalian homologs of the *Drosophila* transient receptor potential (Trp) proteins as biosensors for PLC signaling. Trp homologs form cation channels (TrpC) that are activated by signaling events downstream from receptor activation and PLC stimulation (Zhu et al., 1996; Harteneck et al., 2000; Clapham et al., 2001). In particular, we used mouse TrpC6 and human TrpC1, which are activated by DAG (Hofmann et al., 1999) and via IP₃Rs (Lockwich et al., 2000; Rosado and Sage, 2000), respectively. The activity of these channels was found to be a high-fidelity indicator



Figure 1. Mechanism of Activation of hTrpC1 and mTrpC6 in SCG Neurons

(A–C) Effects of 1-oleoyl-2-acetyl-glycerol, OAG (50 μ M), and thapsigargin (500 nM) recorded in control neurons (A) and in neurons expressing either mTrpC6 (B) or hTrpC1 (C). Cells were voltage clamped at -70 mV using the perforated patch method.

(D) mTrpC6 and hTrpC1 I-V relationships. TrpC I-V relationships are difference-currents obtained by subtracting I-V curves before and after application of OAG (mTrpC6 current) or thapsigargin (hTrpC1 current). Points are mean \pm SEM (n = 5–7). Insets, confocal sections showing plasma membrane expression of myc-tagged mTrpC6 and myc-tagged hTrpC1. Scale bars, 10 μ m.

of PLC stimulation when expressed in neurons, providing a real-time detection device for submembrane production of DAG and InsP₃. These experiments were paralleled by testing the ability of M1AChRs and B2Rs to translocate GFP-tagged DAG- or Ca2+-sensing domains of PKCy. Using these approaches in combination with patch-clamp recording of macroscopic and microscopic membrane domains, we show that although both M_1 AChR and B_2 R are indeed robust activators of PLC β , only InsP₃ produced by B₂R had the ability to activate reticular IP₃Rs. This peculiarity results from a membrane arrangement that links IP₃Rs to B₂Rs, but not M₁AChRs, allowing InsP₃ to be produced at the required concentration and the precise location inside the cell. Thus, membrane-transducing microdomains play a key role in spatio-temporal coding of InsP₃ signals, enabling the cell to discriminate between identical signaling pathways that are triggered by different membrane receptors.

Results

Trp Channels as Biosensors for Detecting Submembrane PLC Products in Neurons

We first determined the properties of activation of Trp channels expressed in superior cervical ganglion (SCG) neurons. Nuclei of SCG neurons were microinjected with cDNA constructs encoding hTrpC1, hTrpC3, and mTrpC6 and perforated-patch somatic recordings were made 12–24 hr later. Expression of these TrpC minigenes in SCG neurons did not produce constitutive activity.

In cells expressing mTrpC6, 1-oleoyl-2-acetyl-snglycerol (OAG, 50 μ M) produced inward currents that were absent in control cells (Figures 1A and 1B). Mouse TrpC6 inward currents were not activated by activators of PKC (pdBU 500 nM, n = 5) or by thapsigargin (TG), an inhibitor of endoplasmic reticulum Ca²⁺-ATPase (Figure 1B, n = 6). OAG-induced mTrpC6 currents displayed properties of nonselective cation currents with an apparent reversal potential of +4 ± 3 mV (Figure 1D, n = 7). These currents were blocked by 90%–97% by 100 μ M La³⁺, as expected for this family of Trp channels (Boulay et al., 1997).

Cytoplasmic injection of $InsP_3$ at a final intracellular concentration of ${\sim}15\,\mu$ M (see Experimental Procedures) activated hTrpC1 but not mTrpC6 (Figure 3; n = 4–6). Lower [InsP_3]_i were largely ineffective in stimulating hTrpC1. hTrpC1-expressing cells also responded to TG by large inward currents (Figure 1C) that reversed at -1 \pm 4 mV (Figure 1D, n = 5) and were blocked by La^{3+}. Bath application of OAG had no effect on hTrpC1 (Figure 1C, n = 9), whereas hTrpC3 could be activated by either OAG (n = 6 of 7) or microinjection of InsP_3 ([InsP_3]_i ~15 ~\muM, n = 3 of 4) (data not shown).

Taken together, these data indicate that mTrpC6 expressed in SCG neurons is activated by OAG, independently of PKC, whereas hTrpC1 is activated via an InsP₃-dependent mechanism. This accords with previous studies in cell lines showing gating of TrpC6 by DAG analogs and activation of TrpC1 by InsP₃ or TG (Boulay et al., 1997; Hofmann et al., 1999; Rosado and Sage, 2000). In the subsequent experiments, we therefore used mTrpC6 and hTrpC1 as real-time sensors of the



Figure 2. Differential Activation of TrpC by M_1AChRs and B_2Rs

(A) Shown are inward currents induced by oxotremorine-M (Oxo-M, 5 μ M, top traces) and bradykinin (BK, 100 nM, bottom traces) in control cells and in cells expressing either mTrpC6 or hTrpC1 (as indicated). Holding potential, -80 mV.

(B) Tonic activation of mTrpC6 and hTrpC1 cation currents in neurons coexpressing GTPase-deficient forms of $G\alpha_q$ but not $G\alpha_{\alpha a}$ subunits ($G\alpha_q$ -QL and $G\alpha_{\alpha a}$ -QL, respectively). Horizontal bars indicate isoosmotic substitution of external Na⁺ by N-methyl-D-glucosamine, NMDG⁺. Holding potential, -70 mV.

PLC β products DAG and InsP₃, respectively. Human TrpC3 was not used further as at least two distinct routes (DAG and InsP₃) appeared to activate this channel in our system (in agreement with Ma et al., 2000).

We confirmed the plasma membrane distribution of TrpC in sympathetic neurons by expressing cDNA constructs encoding mTrpC6 and hTrpC1 C-terminally fused to myc-tag. Both fusion proteins formed functional channels with properties indistinguishable from the channels lacking the c-*myc* tags (n = 4–5). mTrpC6 and hTrpC1 were prominently targeted to the plasma membrane (insets in Figure 1D).

Different Ability of B₂R and M₁AChR to Activate Trp Channels

To trace receptor signals, we tested the ability of constitutively expressed B₂Rs and M₁AChRs to couple to TrpC. The concentration of agonists were chosen to be saturating (see below). In control neurons held at -70 mV, neither bradykinin (BK, 100 nM) nor the muscarinic agonist oxotremorine-M (Oxo-M, 5 μ M) produced inward currents (Figure 2A). Reversible cation currents developed in response to both BK and Oxo-M in cells expressing mTrpC6 (Figure 2A and Table 1). By contrast, in neurons expressing hTrpC1, only BK was able to generate large inward currents while Oxo-M was ineffective (n = 11 of 11; Figure 2A). Stimulation of the G_{i/o} protein-coupled α_2 -adrenoceptor by norepinephrine had no effect in cells expressing either hTrpC1 or mTrpC6 (n = 4–6, data not shown). The EC₅₀ value for activation of mTrpC6 by Oxo-M was 450 ± 35 nM (n = 5). Higher concentrations of Oxo-M (>10 μ M) did not produce activation of hTrpC1. However, in some cases, rapid inward currents were observed with these high concentrations of Oxo-M. They were fully blocked by the nicotinic receptor antagonist, hexamethonium (100 μ M), implying a contribution of nicotinic receptors. EC₅₀ for activation of mTrpC6 and hTrpC1 by BK were 6 ± 3.5 nM (n = 5) and 12 ± 5 nM (n = 6), respectively.

Activation of TrpCs by Oxo-M and/or BK was resistant to overnight treatment with *pertussis* toxin (PTX), consistent with the coupling specificity of M₁AChR and B₂R to PTX-insensitive G proteins (Delmas et al., 1998b; Haley et al., 1998; Haley et al., 2000a). In order to identify further the G protein species involved, GTPase-deficient forms of G protein α subunits (G α Q-L) were expressed together with Trp channels. Tonic activation of TrpC currents was then detected as a standing inward current

Table 1. Receptor/G Protein Coupling to Trp Channels in SCG Neurons									
Treatment	Oxo-M	BK	NE	G _q QL	G ₁₁ QL	G _{oA} QL	G _z QL	G ₁₂ QL	-
Current density ^a	0.040					0.05	o o -		
(pA/pF) Control	0-0.18	0-0.5	0-0.3	0-0.3	0-0.2	0-0.5	0-0.7	0-0.8	
mTrpC6	13 ± 2°	6 ± 1°	$\textbf{0.1}~\pm~\textbf{0.2}$	11.8 ± 3°	9 ± 2°	1.4 ± 0.7	$\textbf{0.3}~\pm~\textbf{0.3}$	0.7 ± 0.3	
hTrpC1	$\textbf{0.9}\pm\textbf{0.4}$	17 ± 3°	0.5 ± 0.3	$8.4 \pm 2^{\circ}$	$6 \pm 2^{\circ}$	$\textbf{0.6}~\pm~\textbf{0.2}$	$\textbf{0.2}~\pm~\textbf{0.1}$	1.1 ± 0.4	
hTrpC3	$\textbf{4.6}~\pm~\textbf{2^{b}}$	$3.5\pm1^{ m b}$	0.4 \pm 0.2	5 ± 2^{b}	NT	NT	0.4 \pm 0.3	NT	

Saturative concentrations were used: [Oxo-M] 5µM, [BK] 100 nM, and [NE] 1µM. NT, not tested.

^aAs estimated from the NMDG⁺-sensitive inward current.

 ${}^{\scriptscriptstyle b}p < 0.05$ versus control (one-way ANOVA and Dunnest's multiple comparison test).

 $^{\circ}p < 0.01$ versus control. (one-way ANOVA and Dunnest's multiple comparison test.)



Figure 3. The B₂R, but Not the M₁AChR, Couples to hTrpC1 via IP₃R (A) Top left panel, schematic diagram of intracellular microinjection of InsP₃ during perforated patch-clamp recording. Cytoplasmic microinjection of InsP₃ (0.5 pl at 100 μ M, arrows) activated hTrpC1 but not mTrpC6. Note that InsP₃ occluded the activation of hTrpC1 by BK (100 nM) but not activation of mTrpC6.

(B) BK (100 nM) but not Oxo-M (5 $\mu M)$ mobilized intracellular Ca^{2+} in control neurons. All cells were voltage clamped at -60 mV.

sensitive to either La³⁺ or external Na⁺ substitution. Human TrpC1 and mTrpC6 currents (revealed by substituting N-methyl-D-glucosamine, NMDG, for external Na⁺) were seen with G α_q Q-L and G α_{11} Q-L mutants but not with G α_{oA} Q-L, G α_z Q-L, and G α_{12} Q-L mutants (Figure 2B and Table 1).

Both B₂R and M₁AChR Stimulate PLC β , but Only B₂R Mobilizes IP₃R

Activation of hTrpC1 by BK and of mTrpC6 by Oxo-M were blocked by application of U73122 (10 μ M for 10 min, n = 4-6), an inhibitor of PLC, but not by calphostin C (500 nM for 10 min, n = 3-5), a specific inhibitor of PKC. This is consistent with the involvement of PLC β in the modulation of both hTrpC1 and mTrpC6. The stimulation of hTrpC1 by BK was blocked by the membranepermeable inhibitors of IP₃R, xestospongin C (50 μ M, n = 6) and diphenylboric acid 2-amino-ethyl ester (2APB, 80 μ M, n = 5; see Figures 7 and 8). These agents had no significant effect on mTrpC6 currents activated by either BK or Oxo-M (paired t tests, p > 0.1, n = 5-6). Consistent with this, preactivation of hTrpC1 by cytosolic $InsP_3$ occluded by 91% (n = 6) further activation of hTrpC1 by BK (Figure 3A). Thus, although both M₁AChRs and B₂Rs stimulate PLC_β, activation of IP₃R-operated hTrpC1 was strictly restricted to B₂Rs. This preferential coupling was correlated with the ability of B₂Rs to mobilize Ca²⁺ from intracellular IP₃R-stores in both the somata (Figure 3B; see also Cruzblanca et al., 1998) and dendrites of control neurons. In contrast, Ca²⁺ mobilization was not detectable either in somata or dendrites in



Figure 4. Kinetics of Receptor-Mediated Production of DAG and \mbox{InsP}_3 in Individual Neurons

Time course of activation of mTrpC6 (A) and hTrpC1 (B) in response to bath application of BK, Oxo-M, and/or OAG. Inward currents induced by a high external K⁺ solution (40 mM) have been rescaled for comparison. Agonists have been applied for 5 s and high K⁺ for 2 s. Same concentrations as in Figures 1 and 2.

(C) Rate of production of DAG and InsP₃ by M₁AChR and B₂R as detected with Trp channels. Dashed, broken lines are rising phases of the currents averaged from three to four cells, and smooth solid lines are polynomial fits to the data. T = 0 (s) indicates the onset of mTrpC6 activation by Oxo-M.

response to M₁AChR stimulation (Figure 3B; in agreement with Cruzblanca et al., 1998; del Río et al., 1999). Addition of the protein phosphatase inhibitors, okadaic acid (PP1/PP2A, 1 μ M, n = 6), calyculin A (PP1, PP2A, 50 nM, n = 5), and FK506 (PP2B, 100 nM, n = 4) to the bath had no significant effect on the lack of coupling of M₁AChRs with either Ca²⁺ organelles or hTrpC1.

Kinetics of Receptor-Mediated Production of DAG and $InsP_3$ in Individual Neurons

We examined the kinetics of production of DAG and $InsP_3$ in response to B_2R and M_1AChR stimulation. In these experiments, a fast perfusion system was used and the responses to high external K⁺ and OAG were taken as estimates for solution exchange and "direct" modulation, respectively (Figures 4A and 4B). When cor-

rection is made for the time course of solution exchange (Figure 4C), mTrpC6 currents developed with a similar rate of rise (10–12 \pm 2% s⁻¹) in response to either Oxo-M or BK. However, responses to BK were consistently delayed by 2 \pm 1 s (n = 7), perhaps indicating that binding of BK to B₂Rs is rate limiting (see also Jones et al., 1995). This later inference is substantiated by the faster activation of mTrpC6 by OAG (Figure 4A). BK activated hTrpC1 with a slower rising rate of 6.7 \pm 2% s⁻¹ (Figure 4C), probably reflecting additional downstream steps including the activation of IP₃Rs.

DAG- and Ca²⁺-Sensing Domains of PKC γ Are Differentially Translocated by M₁AChR and B₂Rs

The above data indicate that both B₂Rs and M₁AChRs activate PLC_B, but only B₂Rs mobilize IP₃R-stores. We sought an alternative assay to verify this selective coupling, by testing the ability of B₂Rs and M₁AChRs to translocate DAG- and Ca²⁺- binding domains of PKC_V (C1₂ and C2, respectively). A previous study in basophilic leukemia cells has shown that the translocation of C1₂ to the plasma membrane is induced by DAG, whereas the translocation of C2 depends on Ca2+ binding (Oancea and Meyer, 1998). cDNAs encoding GFPtagged C1₂ and C2 domains of PKC_Y were microinjected into the nuclei of neurons, and translocation was assessed 8-16 hr later in the absence of external Ca²⁺. Figure 5 shows images of neurons expressing C1₂-GFP and C2-GFP under control conditions and after receptor stimulation. In the controls, C12-GFP and C2-GFP were seen in greatest density in the cytosol (Figure 5B). DAGsensing C1₂ domain translocated to the plasma membrane in response to OAG application and to stimulation of both M_1AChRs and B_2Rs (n = 14 of 19 and n = 12 of 18, respectively). Other receptors such as the Gi/ocoupled α_2 -adrenoceptor and somatostatin receptors were ineffective (n = 6–11). The Ca^{2+} -sensing C2 peptide translocated to the plasma membrane in response to TG and the Ca²⁺ ionophore, ionomycin (in the presence of 0.5 mM external Ca²⁺). Addition of BK led to a marked translocation of C2-GFP to the plasma membrane (n = 11 of 17) (Figure 5B). This change in distribution can be clearly seen using a line scan analysis of the fluorescence across the cell (Figure 5C). BK-induced translocation was fully suppressed by preincubation of cells with xestospongin C (20 μ M for 10 min; n = 6 of 6). In contrast, no plasma membrane translocation of C2-GFP was observed after Oxo-M addition (n = 17 of 17) (Figures 5B and 5C).

Cell Dependence of M₁AChR Coupling to IP₃R

We also tested the ability of M_1AChRs to activate IP_3R in nonneuronal cells and neuroblastoma hybrid cells. We used CHO cells and NG108-15 mouse neuroblastoma × rat glioma hybrid cells (clone BM-8) stably transfected with the human M_1 muscarinic receptor. NG108-15 cells also constitutively express B_2Rs (Fukuda et al., 1988). In these two cell lines, M_1AChR (as well as the B_2R , data not shown) activated transfected hTrpC1 (70%–85% of responsive cells). A representative experiment obtained in CHO cells is shown in Figure 6B. This property of the M_1AChR was associated with its ability to mobilize intracellular Ca²⁺ from IP_3R -stores in control



Figure 5. Receptor-Mediated Translocation of DAG- and Ca^{2+}- Sensing Domains of PKC $\!\gamma$

(B) Distribution of C1₂-GFP (left panels) or C2-GFP (right panels) fusion proteins in neurons before and after addition of the drugs indicated (see Experimental Procedures for more details on drug application).

(C) Line scan profiles of the fluorescence intensity for the C2-GFPexpressing cells shown in (B).

cells (Figure 6A; see also Robbins et al., 1993). M₁AChR stimulation also translocated the initially cytosolic C2-GFP fusion protein to the plasma membrane in CHO cells (Figure 6C, n = 14 of 16). All these responses were abolished by 10 min pretreatment with xestospongin C (n = 3-8).

In clear contrast to primary sympathetic neurons, microinjection of low concentrations of InsP₃ were able to activate hTrpC1 in both CHO and NG108-15 cells. [InsP₃]_i of less than 1 μ M resulted in consistent activation of hTrpC1 in NG108-15 cells (n = 6 of 7), whereas even lower concentrations (~250 nM) were sufficient in CHO cells (n = 2, due to difficulties in impaling these flat cells).

Block of Calmodulin Reveals Remote Production of InsP₃ by M₁AChR

Calmodulin (CaM) has been shown to be responsible for the Ca²⁺-dependent inactivation of type-1 IP₃R (Michikawa et al., 1999). We therefore tested whether CaM can act as a negative feedback regulator of receptor-mediated activation of IP₃Rs in our neuronal system. To do this we used mutants of rat CaM (CaM₁₋₄) in which the aspartate residue found in the first position of each of the four Ca²⁺ binding E-F hand motifs have been



Figure 6. Activation of hTrpC1 and Mobilization of Ca $^{2+}$ by M1AChRs in CHO Cells

(A and B) Oxo-M (5 μ M) mobilizes [Ca²⁺], (A) and activates hTrpC1 (B) in CHO cells stably transfected with M₁AChR. Holding potential, -70 mV.

(C) Oxo-M addition leads to the plasma membrane translocation of the Ca²⁺-sensing C2-GFP in a living CHO cell. Right panel, line scan of the fluorescence intensity across the cell before and after Oxo-M addition.

mutated to an alanine. Such mutations reduced or abolished Ca²⁺ binding (Xia et al., 1998). Mutant CaMs were expressed in neurons and activation of hTrpC1 by B₂Rs or M₁AChRs was examined (Figure 7). In cells expressing CaM₁₋₄, BK produced hTrpC1 currents qualitatively similar to those recorded in control cells or cells expressing wt CaM. However, the peak amplitude of these currents was slightly increased (by 0.95 \pm 0.2 pA/pF, n = 7), and their deactivation rate after removal of agonist was markedly slowed (Figure 7A). The slow deactivation and



Figure 7. $\mbox{IP}_3\mbox{R}$ Activation by $\mbox{M}_1\mbox{AChRs}$ Is Unmasked by Dominant-Negative Calmodulin

(A) hTrpC1 currents evoked by BK (100 nM) in neurons expressing hTrpC1 alone (control trace) or coexpressing hTrpC1 and dominant-negative CaM₁₋₄. Note the slow deactivation of the hTrpC1 current following agonist removal in the presence of CaM₁₋₄. Currents were normalized to cell capacitance for comparison.

(B) Slow developing hTrpC1 current in response to Oxo-M (5 μ M) in a neuron expressing dominant-negative CaM₁₋₄. Note the different scale bars in (A) and (B). 2 APB, 80 μ M.

the slight increase of hTrpC1 currents resulted from the reduced inactivation of IP₃Rs in the presence of CaM₁₋₄, as evidenced by the effects of 2-APB (Figure 7A). Interestingly, in the presence of CaM₁₋₄, M₁AChR stimulation was now also capable of activating hTrpC1 (n = 6 of 10, Figure 7B). Activation of hTrpC1 by M₁AChRs however remained weak and 4- to 7-fold slower in onset than BK responses, suggesting that the M₁AChRs were rather remote from the IP₃R-stores, leading to a diffusional delay in the effect of InsP₃. As in control cells, Oxo-M had no effect in cells expressing wt CaM.

We also tested the effects of Oxo-M (5 μ M) before and after treatment with the calmodulin inhibitor, calmidazolium (1 μ M for > 5 min), in neurons expressing hTrpC1. Oxo-M activated a modest TrpC1 current (-35 ± 5 pA) in the presence of calmidazolium in five out of nine cells tested, whereas no current (-2 ± 1.2 pA) could be observed before treatment (data not shown). This current could be fully blocked by xestospongin-C treatment (20 μ M, n = 4) and by bath application of La³⁺ (100 μ M, n = 3). In control cells (uninjected) and cells expressing mTrpC6 (n = 5 and 4, respectively), the combination of Oxo-M and calmidazolium had no effect.

Actin Cytoskeleton Is Essential for B_2R Coupling to IP_3R

The actin cytoskeleton provides a molecular scaffold upon which signaling complexes may be assembled (Lin et al., 2001). We therefore tested whether the actin cytoskeleton may serve as a structural link between IP_3R -stores and B_2Rs . Neurons were preincubated with the cell-permeant toxin cytochalasin D, an inhibitor of actin filament polymerization. As indicated by staining with Alexa-red-labeled phalloidin, actin filaments in neurons are located mainly in the cell periphery, forming a submembrane cytoskeleton. Incubation with cytochalasin D for 20 min induced reorganization of the membrane cytoskeleton into diffuse foci near the plasma membrane. This treatment abolished activation of hTrpC1 by BK (n = 6 of 8), whereas activation of mTrpC6 by both BK and Oxo-M remained largely unchanged (n = 4–6) (data not shown). The loss in coupling between B_2R and hTrpC1 resulted primarily from the disruption of the B_2R -IP₃R link since hTrpC1 cation currents could still be activated by cytosolic injection of InsP₃ (n = 3 of 3). Disruption of cytoskeletal tubulin by exposure to the microtubule inhibitor nocodozole had no effect on membrane receptor coupling to either hTrpC1 or mTrpC6 (n = 3–4, data not shown).

Tight Functional Association of B_2R to IP_3R in Perforated Microvesicles

The strong functional coupling of B₂Rs to IP₃Rs suggests that they may be associated within a tight signaling complex. We tested this by recording from perforated microvesicles excised from neural membranes (Figure 8A). First, an amphotericin-perforated patch was obtained on a neuron expressing hTrpC1, and BK was tested on the whole-cell macroscopic currents. If the cell was found responsive (e.g., Figure 8B, left panel) the patch was excised and a perforated microvesicle of \sim 3 μ m diameter (see Experimental Procedure and Sakmann and Neher, 1995) formed in an outside-out configuration (Figures 8A and 8B). BK was then reapplied on the voltage-clamped microvesicle, and hTrpC1 currents were monitored (Figure 8B, right panel). Using this method we could observe activation of microscopic hTrpC1 currents in 6 out of 11 microvesicles tested. As shown in Figure 8B, hTrpC1 activation was reversibly blocked by local superfusion of 2APB (as well as by xestospongin C, n = 3), indicating that interaction between B₂R, PLCβ, IP₃R, and hTrpC1 was conserved in these microvesicles. In contrast, out of nine microvesicles tested with mTrpC6, none responded to BK, while six of them generated mTrpC6 inward currents in response to OAG application (data not shown). Consistent with this, only 2 out of 12 microvesicles expressing mTrpC6 (tested with subsequent application of OAG), responded to Oxo-M. This confirms the random distribution of mTrpC6 versus B₂Rs and M₁AChRs.

Coimmunoprecipitation of B₂R with IP₃R

To assess possible physical interactions between B₂R and IP₃R, we tested the ability of IP₃R₁ antibody to coimmunoprecipitate myc-tagged B₂Rs overexpressed in NG108-15 cells. As shown in Figure 8C, myc-tagged B₂Rs, but not myc-tagged M₁AChRs, were detected in IP₃R₁ immunoprecipitates, whereas both proteins were easily detected by Western blot (Figure 8D, bottom panel). In addition, IP₃R₁ antibody also coimmunoprecipitates endogenous B₂Rs but not M₁AChRs from SCG lysate (Figure 8D). Immunoprecipitates were also reprobed with antibodies against G $\alpha_{q/11}$ or G_{oA/B} G protein subunits. G $\alpha_{q/11}$ subunits were most clearly part of the immunoprecipitated complex, whereas G $\alpha_{oA/B}$ subunits were not detected (data not shown).

Colocalization of IP₃R with B₂R but Not M₁AChR

Triple-label immunoconfocal microscopy was performed in sympathetic neurons maintained 3–4 days in culture to determine the cellular localization of endogenous B_2Rs and M_1AChRs versus reticular IP₃Rs. B_2R and



Figure 8. Docking of IP₃Rs with Plasma Membrane B₂Rs

(A and B) Same cell. (A) Schematic diagram of the manipulations performed to obtain perforated microvesicles. After cell-attached patch formation and permeabilization, the pipette was withdrawn to form an outside-out vesicle. The superimposed traces on the right show the change in membrane capacitance transients that results from the formation of the perforated vesicle. (B) hTrpC1 currents induced by BK in the whole-cell recording configuration (left panel) and in the perforated vesicle (right panel). Note the block of microscopic hTrpC1 currents by 2APB (80 μ M).

(C and D) B₂R coprecipitates with IP₃R₁. (C) Detergent lysates of NG108-15 cells transfected with myc-tagged B₂R (left lane) or myc-tagged M₁AChR (middle and right) were immunoprecipitated with rabbit anti-IP₃R₁ antibody (IP) and blotted (IB) with anti-myc antibody (mouse) or rabbit anti-IP₃R₁ antibody (reprobe, right lane). Control IPs using rabbit serum or IgG fraction were negative for mouse antimyc antibody (data not shown). Multiple high mw bands in IP₃R₁ are presumed proteolysis products. The band at ~50 kDa obtained with the anti-IP₃R₁ antibody represents reduced heavy chains of the precipitated with rabbit anti-IP₃R₁ antibody and blotted with mouse anti-B₂R, goat anti-M₁AchR, or rabbit anti-IP₃R₁ antibodies. Bottom panel, Western blot for myc-tagged B₂Rs and M₁AChRs.

 M_1 AChR labelings were heterogeneous in distribution, with patches of punctate labeling separated by areas in which little labeling was apparent (Figure 9A). IP₃R staining also showed punctate fluorescence within the cytoplasm and extended to the plasma membrane forming discrete patches of concentrated fluorescence, outlining the cell periphery. Simultaneous viewing of the staining patterns observed in 33 neurons revealed a close relationship between IP₃R puncta and B₂R staining



Figure 9. Colocalization of Endogenous B_2Rs with IP_3Rs

(A) Fluorescence patterns for endogenous B₂Rs, M₁AchRs, and IP₃Rs in a cultured sympathetic neuron. Cells were triple labeled with mouse monoclonal B₂R antibody (red), rabbit anti-IP₃R₁₋₃ polyclonal antibody (green), and goat polyclonal M₁AChR (blue). All images are reconstructions from serial optical sections.

(B) Panel showing a combined visualization of B_2R/IP_3R and M_1AChR/IP_3R labelings (Cy5-M_1AChR fluorescence was converted to red for comparison). Note that where B_2R staining occurs its distribution overlaps that of IP_3R (arrows and yellow spots).

(Figure 9B). Of 437 B₂R-positive structures, 369 (84.5%) showed juxtaposition or partial overlap with an IP₃R positive punctum. By contrast, of 511 M₁AChR-positive structures, only 89 (17.5%) were coincident or juxtaposed with IP₃R-fluorescent spots. Taken together, these data show that IP₃R-positive structures predominated adjacent to B₂R labeling and are consistent with the hypothesis that IP₃R and B₂R form functional complexes.

Discussion

The present study shows that the different Ca²⁺ signaling capabilities of B₂Rs and M₁AChRs in these (sympathetic) neurons reside in the spatio-temporal mobilization of IP₃Rs. In contrast to M₁AChRs, B₂Rs—together with G_{q/11} proteins and PLCβ—are clustered with IP₃Rs to form discrete signaling complexes. These complexes enhance the efficiency of signal propagation and ensure specificity of the signaling pathway.

Transfected Trp Cation Channels as PLC Biosensors in Neurons

In our experiments, we have developed the use of Trp cation channels as sensors of PLC signaling, taking advantage of several of their unique features, including selective activation by PLC products (Harteneck et al., 2000). Perforated whole-cell recordings and immunostaining established that the homomeric hTrp1 and mTrp6 channels are expressed, inserted into the plasma membrane, and respond selectively to either DAG or InsP₃. Thus, by comparing the activation of hTrpC1 and mTrpC6, we could monitor the formation of InsP₃ and DAG in submembrane compartments, allowing real-time

detection of PLC products in living neurons. Although there is good agreement that TrpC is activated downstream to PLC, the identity of the final regulatory element(s) for the multiple TrpC isoforms remains controversial. When expressed in our system, mTrpC6 was activated by DAG (or possibly one of its metabolites) in accordance with recent reports (Hofmann et al., 1999), whereas hTrpC1 was activated via the mobilization of reticular IP₃R, possibly by direct conformational coupling (Lockwich et al., 2000; Rosado and Sage, 2000; see also Boulay et al., 1999; Ma et al., 2000). In this respect, we found that myc-tagged hTrpC1 expressed in SCG neurons colocalized with native IP₃R in discrete regions of the surface membrane (our unpublished data), which makes them ideally suited for studying IP₃R signals.

This approach has clear advantages over previous methods of measuring PLC in living neurons. The most commonly used method is by monitoring inositol polyphosphate formation in large population of cells, but this affords poor spatial and temporal resolution. Other methods for tracing $InsP_3$ dynamics in single cells rely on an indirect measure with Ca^{2+} -sensing dyes (Tsien, 1998) or $InsP_3$ -mediated translocation of GFP-tagged pleckstrin homology domain of PLC- δ 1 (Hirose et al., 1999). Our method has the advantage in that it incorporates the PLC sensors in the surface membrane where PLC is known to reside and is therefore most suited for real-time monitoring of membrane-localized PLC signaling.

Signaling Microdomains for InsP₃

We have provided ample evidence that the B₂R, but not the M₁AChR, activates IP₃Rs in sympathetic neurons. Our data confirm and complement previous observations using intracellular Ca2+ measurements (Cruzblanca et al., 1998; del Río et al., 1999). Two arguments however demonstrate that the lack of coupling of M1AChR to IP3R does not result from its inability or relative inefficiency in stimulating PLCβ. First, M₁AChRs strongly activated mTrpC6 channels and translocated the DAG-sensing C1₂ domain of PKC γ to the plasma membrane. Second, the kinetics of M₁AChR- and B₂Rmediated production of DAG were similar. Taken together, these data indicate that both receptors are equally effective in stimulating PLC, in agreement with population-based analysis of InsP₃ production in SCG neurons (Bofill-Cardona et al., 2000) and with previous observations that demonstrate that both Oxo-M and BK can activate protein kinase C to open chloride channels in these cells (Marsh et al., 1995).

Instead, we suggest that the differential abilities of B_2Rs and M_1AChRs to stimulate IP_3Rs results from a more intimate relationship between the B_2Rs and the IP_3Rs . B_2Rs are not evenly distributed along the plasma membrane but instead often appear to be clustered with IP_3Rs . The endoplasmic reticulum (ER) is well known to come in close contact with the plasma membrane (Berridge, 1998; Patterson et al., 1999). Close apposition of the ER to the plasma membrane has been previously reported in rat sympathetic neurons (Henkart, 1980) where it appears to follow the contours of the somatic plasma membrane in the cell body of rat SCG neurons

(S.J. Marsh, personal communication). Clusters of B₂Rs have also been observed in neurites of differentiated neuroblastoma cells where they play a key role in initiating bradykinin-induced Ca²⁺ waves (Fink et al., 2000). The finding that the B₂R-IP₃R link was preserved in microvesicles excised from neural membrane strongly suggests that their association resembles a physical docking. This is supported by the finding that B₂R can be immunoprecipitated by IP₃R₁ antibodies. A key question therefore that arises out of these findings is how reticular IP₃Rs interact with B₂Rs. We have provided evidence that a cortical actin skeleton might play a role in stabilizing B₂R-IP₃R association, perhaps forming a bridge between the B₂R and the IP₃R by its ability to anchor both PLC β and IP₃R. Linkage of IP₃Rs to the plasma membrane through an actin bridge has previously been reported in liver cells (Rossier et al., 1991), and Bourgignon et al. (1993) have reported that cytochalasin D also inhibits InsP₃-induced Ca²⁺ release in platelets. More recently, Homer proteins have been shown to form a physical tether cross-linking mGluRs with IP₃Rs (Tu et al., 1998). Although the proline-rich Homer ligand motif is not obviously present in the B₂R, it remains to be tested whether Homer proteins or related proteins may link B₂R to IP₃R.

The fact that IP₃Rs are in close vicinity to the PLCcoupled B₂R allows them to be flooded by high InsP₃ concentration, which results in effective IP₃R stimulation. These signaling microdomains do not appear to have restricted InsP₃ diffusion, as IP₃R stimulation diminished smoothly after the stimulus had ceased; this would not be the case if InsP₃ was confined to specialized compartments with restricted diffusion (such as demonstrated for cAMP in HEK-293 cells; Rich et al., 2000). Thus, B₂R and M₁AChR signals do not operate in isolated subcellular compartments. As demonstrated with dominant-negative CaM, InsP₃ produced by the M₁AChR is not physically excluded from B2R-IP3R-hTrpC1 microdomains but simply does not reach a sufficient concentration with appropriate speed. One reason for this is the random distribution of M₁AChRs versus IP₃Rs. Longer and probably more intense muscarinic stimulation may therefore be required to reach InsP₃ levels high enough to activate IP₃Rs. It is worth noting that the level of resting cytosolic Ca²⁺ can influence M₁AChR coupling to IP₃Rs, as demonstrated by preconditioning the cell to depolarized voltages (del Río et al., 1999).

Two other important factors also contribute to attenuate Ca^{2+} signals produced by muscarinic receptor stimulation. First, the negative feedback played by CaM on IP₃R activity serves as a filter that is more efficient in inhibiting slow rather than rapid rises in InsP₃. Second, IP₃Rs in SCG neurons—and in neurons in general—have low affinities for InsP₃ in comparison with those in NG108-15 or CHO cells. These differences in sensitivities between neurons, neuroblastoma, and nonneuronal cells have been reported previously (Khodakhah and Ogden, 1993; see also Fink et al., 2000). They may explain why M₁AChRs are able to stimulate IP₃Rs in NG108-15 and CHO cells, but not in SCG neurons, and emphasize the need in neurons for high local InsP₃ concentrations.

A simplified view of InsP₃ signaling emerging from our



Figure 10. Model for Spatio-Temporal Coding of \mbox{InsP}_3 Signaling in SCG Neurons

Agonist activation of M₁AChRs and B₂Rs via heterotrimeric G proteins stimulates PLC β , leading to the prodution of DAG and InsP₃ second messengers. Complexes of B₂Rs with reticular IP₃Rs create high local [InsP₃] in close proximity to the IP₃Rs, releasing Ca²⁺ from endoplasmic reticulum stores. The activation of IP₃Rs and/or the fall in luminal Ca²⁺ in the store (see Putney, 1999) induce a conformational change of the IP₃R, causing hTrpC1 to open. The signaling complexes containing B₂Rs, G proteins, PLC β , and IP₃Rs (along with exogenous hTrpC1) requires an actin-provided scaffold to support the IP₃Rs in a position close to the plasma membrane. Although M₁AChRs activate PLC β and consequently open mTrpC6, they fail to mobilize IP₃Rs domains. Activation of IP₃Rs by diffusional InsP₃ is prevented further by the low sensitivity of IP₃R for InsP₃ and the inhibitory forg leadable by calmodulin.

study is depicted in Figure 10. Extracellular signals such as ACh or BK bind their respective receptors in the cell plasma membrane, which then activate the membraneassociated G proteins and PLC β with comparable efficiencies. B₂R/PLC β are clustered with (some) IP₃Rs, thus creating spatially compact signaling complexes. (This complex may also contain a specific isoform of PLC, PLC β 4, linked to the B₂R; see Haley et al., 2000b). This causes the Ca²⁺-mobilizing messenger InsP₃ to build up in discrete domains and activate IP₃R. IP₃Rs do not form such signaling complexes with M₁AChRs, which make them "diffusionally isolated" from these receptors.

Physiological Implication of Signaling Microdomains

Localizing B_2Rs and IP_3Rs within signaling microdomains provides a mechanism for achieving specificity and efficiency by ensuring that IP_3Rs located remotely from receptors are not exposed to the same level of $InsP_3$ as those in the vicinity of the complex. This results in the coordinated gating of IP_3Rs sequestered to the

microdomain and thereby removes the need for InsP₃ to accumulate throughout the cytosol. This is particularly important in view of the low sensitivity of neuronal IP₃R for InsP₃. An example of the differential functional effects of B₂Rs and M₁AChRs in sympathetic neurons resulting from this localization concerns the mechanisms whereby they inhibit voltage-gated M channels. While activation of either receptor inhibits these channels, and thereby increases neuronal excitability (see Marrion et al., 1989; Jones et al., 1995), the mechanisms whereby they do so differ. Inhibition by B₂Rs results from the rise in Ca²⁺ produced by the tight coupling to IP₃Rs, whereas inhibition by M₁AChRs results from some other, non-PLC/ Ca²⁺-dependent mechanism (Cruzblanca et al., 1998; Haley et al., 2000a; Bofill-Cardona et al., 2000). In contrast, since both receptors stimulate the production of DAG, both activate the same PKC-dependent chloride current through a common mechanism (Marsh et al., 1995).

Nevertheless, it is clear that InsP₃ production is enhanced by M₁AChR stimulation in these neurons (see del Río et al., 1999, and references therein). What could be the function of this? Since the most effective stimulation of IP₃Rs is achieved when both intracellular Ca²⁺ and InsP₃ are presented together (Berridge, 1998), this dual activation system might act as a coincidence detector: small elevations of InsP₃ by M₁AChRs that are not able to cause Ca²⁺ mobilization may enhance the Ca²⁺ sensitivity of the IP₃R, thereby facilitating further PLC-mediated receptor signals.

In conclusion, the extensive and intricate network of signaling pathways requires mechanisms to achieve specificity. Here, we have described the presence of spatially restricted signaling complexes between the B_2R and the IP₃R. These signaling microdomains allow localized formation of InsP₃ and coordinated activation of sequestered IP₃Rs. Our data provide a framework for understanding the multiple and differential cellular functions regulated by PLC-coupled membrane receptors.

Experimental Procedures

cDNA Constructs

cDNA for hTrpC1, hTrpC3, and mTrpC6 have been kindly provided by Dr. Christian Harteneck (Institut für Pharmakologie, Freie Universität Berlin, Germany) and were subcloned into pcDNA3 or pMT vectors (Stratagene). cDNA encoding GTPase-deficient forms of G α_q (Q209L), G α_{11} (Q209L), G α_{0A} (Q205L), and G α_{12} (Q229L) were subcloned either in pcDNA1 or pcDNA3 vectors and have been described previously (Delmas et al., 1998b; Haley et al., 1998). C1₂ and C2 domains of PKC γ were subcloned in pEGFP-N1 vector. cDNA encoding a Ca²⁺-insensitive mutant of calmodulin (CaM₁₋₄) (first D in E-F hand 1,2,3,4 mutated to A) and wt CaM were subcloned in pBK-CMV or pMT. D to A mutation in amino acids 21, 57, 94, and 130 was verified by sequencing. Mouse B₂ bradykinin receptor was in pMT. Plasmids were propagated in DH5 α *Escherichia coli* and purified using maxiprep columns (Qiagen, Hilden, Germany). All plasmids were verified by sequencing.

Cell Cultures and cDNA Delivery

Sympathetic neurons were isolated from Sprague Dawley rats (2 weeks old) and cultured on glass coverslips as described previously (Delmas et al., 1998a). DNA plasmids were diluted to 100 μ g/ml in Ca²⁺-free solutions (pH 7.3) containing 0.2% FITC-dextran (70,000 MW) or rhodamine (10,000 MW) and were pressure injected into the nucleus of SCG neurons as largely described (Delmas et al., 1999;

Delmas et al., 2000). Cells were maintained in culture after injection and identified for patch clamping by fluorescence microscopy. CHO hm1 and differentiated NG108-15 cells were cultured on glass coverslips in α -MEM medium supplemented with 10% fetal calf serum, 1% L-glutamine, and 1% penicilin/streptomycin. Cells were transfected 1–3 days after plating using LipofectAmine Plus (Life Technologies, Gaithersburg, MD). The plasmids of interest were cotransfected with cDNA for green fluorescent protein as marker (10:1 ratio). Recordings were made 12–24 hr after transfection.

Cytosolic Microinjection of InsP₃

Microinjection was made using a method similar to that used for cytoplasmic injection of antibodies (Delmas et al., 1999). Briefly, micropipettes with resistance of about 60–70 MΩ were tip filled with standard intracellular solution (300 mOsmol/l) for 1 min and back filled with the same solution containing either 1, 10, 100, or 300 μ M of InsP₃ plus 0.2% TRITC-dextran (10 kDa). The cell of interest was first voltage clamped using the perforated patch-clamp method and then impaled with the InsP₃-containing micropipette. A slight negative pressure was applied to the interior of the micropipette in order to avoid leakage of InsP₃. Five minutes was allowed after impalement to allow the holding current to recover if any leak current developed. Cells that did not recover within 5 min were not used. The injected volume was \sim 0.5 pl, giving a mean dilution factor of 8, 10.4, and 4 in SCG neurons, NG108-15, and CHO cells, respectively. Final concentrations of InsP₃ were calculated from these dilution factors.

Translocation of GFP-Tagged PKCγ Domains

cDNA of GFP-tagged minimal C1₂ and C2 domains of PKC_V have been described (Oancea and Meyer, 1998). As previously shown in basophilic leukemia 2H3 cells, the C-terminal GFP tags do not affect cofactor dependence and translocation of these peptides to the plasma membrane. Experiments in SCG neurons were carried out 7-18 hr after nuclear microinjection of cDNA to obtain relatively low levels of expression. Treatment with agonists or drugs were as follows: oxotremorine-M (Oxo-M, 5 µM) for 1 min; bradykinin (BK, 500 nM) for 30 s; 1,oleoyl-2-acetyl-glycerol (OAG, 50 μM) for 5 min; and ionomycin (10 µM) for 5 min. These experiments were made in the absence of extracellular Ca2+ (except for ionomycin, 0.5 mM Ca2+) in order to prevent Ca2+ influx eventually caused by druginduced depolarization. Cells were then fixed with 4% paraformaldehyde and 0.15 M sucrose in phosphate-buffered saline (PBS) (25 min at room temperature). Images were acquired on a Zeiss Axiophot microscope or a confocal microscope and analyzed using NIH image 1.62f software.

Measurement of Intracellular Calcium

Intracellular free Ca²⁺ concentration was estimated from indo-1 fluorescence using the ratiometric method (Grynkiewicz et al., 1985). Briefly, cells were incubated with 5 mM indo-1-AM for 30–45 min and then superfused with standard bath solution for 10–20 min to allow ester hydrolysis. Indo-1 was excited at 360 nm (330–380 nm bandwidth) and the emitted light was split by a dichroic mirror to two photomultipliers (Thorn EMI QL 30) with input filter at 408 and 480 nm. The 408/480 emission ratio was digitized using a Digidata 1200 interface driven by pClamp software. Sample rate was 100 Hz. Calibration was achieved by whole-cell dialysis of SCG neurons with intracellular solutions containing known concentrations of free Ca²⁺ (Calcium Calibration Kit, adjusted for pH and osmolarity).

Patch-Clamp Recording

Whole-cell macroscopic currents were recorded using the perforated patch method as described (Haley et al., 2000a) on neurons kept 3–4 days in culture (e.g., microinjected at day 2). Briefly, amphotericin B (0.07–0.1 mg/ml in dimethylsulfoxide) was dissolved in the intracellular solution consisting of 90 mM K⁺ acetate, 30 mM KCl, 3 mM MgCl₂, and 40 mM Hepes (adjusted to pH 7.3 with KOH, 290 mOsmol/l). When filled with this internal solution, pipette resistance was 2–3 M Ω . Access resistances after permeabilization ranged from 8–15 M Ω . The external solution consisted of 130 mM NaCl, 3 mM KCl, 1 mM MgCl₂, 10 mM HEPES, 0.0005 mM tetrodotoxin (TTX), 0.2–2 mM CaCl₂, and 11 mM glucose (adjusted to pH 7.3 with NaOH,

292 mOsmol/l). Recordings were obtained with an Axopatch 200A amplifier (Axon Instruments) and filtered at 2 kHz.

For recording of perforated microvesicles, patch pipettes were pulled from thin-walled borosilicate glass capillaries (Harvard Apparatus, UK), polished, and coated with Sylgard. They had a resistance of ${\sim}2~M\Omega~(1{-}1.25~\mu{\rm m}$ tip diameter). After permeabilization (incomplete), the pipette was withdrawn from the cell to form a perforated vesicle. During this process, the patch-clamp amplifier was switched to current-clamp mode with zero resting current. Assuming an Ω -shaped geometry of these vesicles (Sakmann and Neher, 1995), the total membrane area was estimated to be 30–40 $\mu{\rm m}^2$, from which the sealed membrane portion was 21 $\mu{\rm m}^2$, the perforated membrane portion was 13 $\mu{\rm m}^2$ and the patch area in contact with the bath was 0.8–3 $\mu{\rm m}^2$.

Coimmunoprecipitation

NG108-15 cells were transfected with myc-tagged mB₂R or myctagged rM₁AChR cDNAs using Tfx-50 reagent. Cells were scraped in TE buffer (50 mM Tris and 1 mM EDTA, pH 7.4) containing 1% CHAPS and protease inhibitor cocktail (10 μl per 10 6 cells). Isolated SCGs were cleansed of connective tissue, cut into small pieces, and sonicated in TE buffer containing 1% CHAPS and protease inhibitor cocktail (100 mg wet weight/ml). NG108-15 and SCG homogenates were then centrifuged at 60,000-90,000 rpm at 4°C for 20 min. 1 ml of the NG108-15 cellular extract and 100 μl of SCG cellular extract were used in the immunoprecipitation assay with 1.3–3 μ l anti-IP₃R₁ antibody (rabbit, Alomone labs). Antibodies and extract were then incubated for 1 hr at 4°C before adding protein G sepharose. Immunoprecipitates were washed three times in TE-CHAPS buffer, separated by SDS-PAGE, and analyzed by immunoblot. Blots of immunoprecipitated complexes were probed (or reprobed) with anti-myc antibody (mouse, 1/1000, oncogene, Research products), anti-M1AChR antibody (goat, 1/500, Santa Cruz Biotechnology, Inc.), anti-B2R antibody (mouse, 1/1000, Research Diagnostics, Inc), anti-IP₃R₁ antibody (rabbit, 1/1000), and anti-G $\alpha_{q/11}$ or anti-GaoA/B antibodies (1/800, gift from Graeme Milligan) and visualized using ECL-Western blotting detection reagents (Amersham pharmacia).

Immunofluorescence

SCG neurons were fixed with 4% paraformadehyde, permeabilized with 0.1% triton, and blocked by 1% BSA (PBS buffered) for 30 min before incubation for triple labeling. The optimal procedure for triple labeling was treatment with a mixture of mouse monoclonal B2R antibody (1/800, Research Diagnostics, Inc.) and goat polyclonal M1AChR (1/800, Santa Cruz Biotechnology, Inc.) overnight at room temperature, followed by a rabbit polyclonal anti-IP₃R₁₋₃ antibody (2–4 μ g/ml, Calbiochem) for 2–3 hr. Cells were washed four times and incubated with a mixture of the FITC- (sheep anti-rabbit, 1/100), Cy3- (donkey anti-mouse, 1/250), and Cy5- (donkey anti-goat, 1/250) conjugated secondary antibodies (all Chemicon, Euromedex, France) at room temperature for 1 hr. Controls involved secondary antibodies crossover experiments and competitive peptide inhibition. Images were obtained by confocal laser scanning microscopy using a Leica TCS SP equipped with krypton, argon, and helium neons lasers. Series of images were sequentially acquired to avoid signal crossover, and quantification of puncta was performed on 10 µm confocal z-axis stacks from 33 selected neurons.

Solutions

Solutions were made just before the experiments and applied using a gravity-fed perfusion system at 10–15 ml/min. For the experiments in Figure 3, test agents were applied to neurons through a glass micropipette (30–50 μ m i.d.) placed 50 μ m away from the neuron under study. Dose-response curves for Oxo-M were constructed using cumulative application of increasing concentrations of the muscarinic agonist, whereas noncumulative dose-response curves were made for BK (to avoid desensitization). Data were expressed as the mean \pm SE of the mean. ANOVA and Student's t test were applied to determine the statistical significance, and differences were considered significant if p < 0.05. All experiments were performed at 31°C–33°C.

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