

# Assembly of a β<sub>2</sub>-adrenergic receptor—GluR1 signalling complex for localized cAMP signalling

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Central noradrenergic signalling mediates arousal and facilitates learning through unknown molecular mechanisms. Here, we show that the  $\beta_2$ -adrenergic receptor  $(\beta_2AR)$ , the trimeric  $G_s$  protein, adenylyl cyclase, and PKA form a signalling complex with the AMPA-type glutamate receptor subunit GluR1, which is linked to the β<sub>2</sub>AR through stargazin and PSD-95 and their homologues. Only GluR1 associated with the β<sub>2</sub>AR is phosphorylated by PKA on  $\beta_2$ AR stimulation. Peptides that interfere with the  $\beta_2$ AR-GluR1 association prevent this phosphorylation of GluR1. This phosphorylation increases GluR1 surface expression at postsynaptic sites and amplitudes of EPSCs and mEPSCs in prefrontal cortex slices. Assembly of all proteins involved in the classic β<sub>2</sub>AR-cAMP cascade into a supramolecular signalling complex and thus allows highly localized and selective regulation of one of its major target proteins. The EMBO Journal (2010) 29, 482-495. doi:10.1038/ emboj.2009.344; Published online 26 November 2009 Subject Categories: signal transduction; neuroscience Keywords: adrenergic; AMPA; localized; signalling

### Introduction

Although cAMP can in general freely diffuse, it can also be spatially restricted and exert highly localized actions (e.g. Levitzki, 1988; Neubig, 1994; Zaccolo and Pozzan, 2002; Rebois and Hebert, 2003; Fischmeister et al, 2006; Richter et al, 2008; Dai et al, 2009). Evidence for constitutive association of G protein-coupled receptors and their downstream effectors, the trimeric G<sub>s</sub> protein and adenylyl cyclase,

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with each other has been established and interactions of such complexes with downstream targets have been proposed to form the basis for spatially restricted cAMP signalling (Levitzki, 1988; Neubig, 1994; Rebois and Hebert, 2003; Dai et al, 2009). Here, we describe a complex containing the β<sub>2</sub>-adrenergic receptor (β<sub>2</sub>AR), G<sub>s</sub>, adenylyl cyclase, PKA, and the AMPA receptor (AMPAR) GluR1 subunit that allows highly localized signalling by cAMP in neurons.

A brief period of high activity can permanently enhance synaptic transmission between neurons in the hippocampus. This long-term potentiation (LTP) has been a focus of intense study, as a tractable experimental model for learning and memory. Current evidence implicates trafficking of AMPARs to the postsynaptic membrane as the primary mechanism of LTP expression. AMPARs are tetramers formed by GluR1-4 subunits, with GluR1/2 and GluR2/3 heterotetramers accounting for the majority of AMPARs in the hippocampus (Hollmann and Heinemann, 1994; Wenthold et al, 1996; Dingledine et al, 1999) and GluR1/2 constituting the functionally by far prevailing AMPAR component at postsynaptic sites under basal conditions (Lu et al, 2009). The GluR1 subunit is thought to have a special function in initiating the remodelling associated with increased synaptic expression of AMPARs on LTP. Phosphorylation of GluR1 by PKA on serine 845 (S845) is critical for activity-driven accumulation of GluR1 at postsynaptic sites (Esteban et al, 2003) and fosters surface expression of GluR1 (Ehlers, 2000; Swayze et al, 2004; Sun et al, 2005; Oh et al, 2006; Man et al, 2007). Moreover, GluR1 is physically associated with PKA and the counteracting phosphatase calcineurin/PP2B, optimizing the efficacy of this regulation (Colledge et al, 2000; Tavalin et al, 2002).

Norepinephrine, released by an extensive network of fibres originating from the locus coeruleus, supports arousal and learning under novel and emotionally charged situations (Cahill et al, 1994; Nielson and Jensen, 1994; Berman and Dudai, 2001; Strange et al, 2003; Strange and Dolan, 2004; Minzenberg et al, 2008) and facilitates various forms of LTP in the dentate gyrus and CA1 region of the hippocampus (Thomas et al, 1996; Lin et al, 2003; Walling and Harley, 2004; Gelinas and Nguyen, 2005). However, the molecular basis of these noradrenergic actions is unclear. Norepinephrine acts through αAR and βAR. Stimulation of the  $\beta_1AR$  and  $\beta_2AR$  activates  $G_s$ , adenylyl cyclase, and PKA. The  $\beta_2AR$  is concentrated at excitatory postsynaptic sites in pyramidal neurons (Davare et al, 2001). We now show that GluR1 forms a signalling complex with the β<sub>2</sub>AR, G<sub>s</sub>, and adenylyl cyclase for highly localized GluR1 phosphorylation, which promotes surface expression of GluR1 and increases EPSC and mEPSC amplitudes in cortex.

# Results

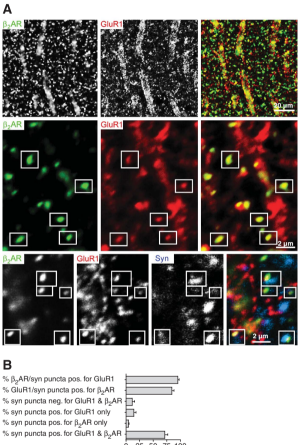
# β<sub>2</sub>AR and GluR1 colocalize in hippocampal neurons

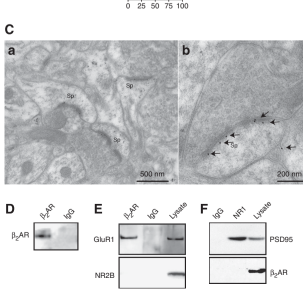
Double labelling for the  $\beta_2AR$  and GluR1 in tissue slices from adult hippocampus showed prominent colocalization of

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immunoreactive puncta (Figure 1A). Most of these puncta were closely associated with puncta immunoreactive for the presynaptic marker synaptophysin, suggesting that they represent bona fide synapses. Quantitative analysis indicates that >90% of the  $\beta_2AR$  puncta associated with synaptophysin were also positive for GluR1 (Figure 1B). Furthermore, >80% of GluR1 puncta associated with synaptophysin were also immunoreactive for  $\beta_2AR$ , implying that the large majority of GluR1-containing synapses also possess the  $\beta_2$ AR.

The precise localization of the  $\beta_2AR$  was determined by post-embedding immunogold labelling in the prefrontal cor-





tex (PFC), which receives especially prominent noradrenergic innervation (e.g. Minzenberg et al, 2008). A large fraction of gold label was synaptic, and a large fraction of asymmetric axospinous synapses were immunopositive (Figure 1C). Gold particles concentrated over the postsynaptic density close to the postsynaptic membrane. Particles could also be seen within spines, in large dendritic shafts, in which they were typically associated with microtubules and in the cytoplasm of neuronal somata (data not shown).

After solubilization with Triton X-100 and removal of nonsolubilized material by ultracentrifugation, immunoprecipitation of the β<sub>2</sub>AR from rat forebrain (Figure 1D) led to coimmunoprecipitation of GluR1 (Figure 1E). This co-precipitation was specific, as the NMDA-type glutamate receptor (NMDAR) subunit NR2B, which has an overall structure and cellular distribution similar to GluR1, did not co-precipitate with the β<sub>2</sub>AR, and control IgG did not immunoprecipitate GluR1 or the β<sub>2</sub>AR. In contrast, immunoprecipitation of the NMDAR-PSD-95 complex (Leonard et al, 1998; Valtschanoff et al, 2000; Lim et al, 2002, 2003) did not result in co-precipitation of the  $\beta_2AR$  (Figure 1F). We conclude that the  $\beta_2AR$  assembles with GluR1-containing AMPARs, but not with NMDAR into a molecular complex at synapses.

# GluR1 interacts with the β<sub>2</sub>AR through PSD-95 and stargazin/y2

PSD-95 co-immunoprecipitated with the β<sub>2</sub>AR from rat brain extracts (Figure 2A). In vitro pull-down experiments indicate that this association is mediated by the C-terminus of the  $\beta_2$ AR. The distal end of the  $\beta_2$ AR C-terminus (DSPL), which conforms to a type 1 PDZ domain ligand, binds to the third PDZ domain of PSD-95 (Figure 2B). Stargazin (or  $\gamma$ 2) and its homologues  $\gamma 3$ ,  $\gamma 4$ ,  $\gamma 5$ ,  $\gamma 7$ , and  $\gamma 8$  (TARPs) associate with AMPARs to promote their surface expression and modulate their biophysical properties. Stargazin binds with its C-terminus to the first two PDZ domains of PSD-95, and this interaction is required for surface expression and postsynaptic

Figure 1 Colocalization and co-immunoprecipitation of  $\beta_2AR$  and GluR1. (A) Rat brain sections were triple-labelled for β<sub>2</sub>AR (left panels; green in overlays), GluR1 (centre panels; red), and synaptophysin (blue in bottom right panel). GluR1 and β<sub>2</sub>AR puncta colocalize with each other (boxes) and associate with synaptophysin puncta (triple overlay, bottom right panel) in the stratum radiatum of the CA1 area. (B) Quantification of the number of  $\beta_2AR$ , GluR1, or synaptophysin (syn)-labelled puncta that shows positive (pos.) or no (neg.) staining for one or both of the other proteins. More than 90% of puncta that are positive for the  $\beta_2AR$ and associated with synaptophysin are also positive for GluR1 (top bar) and >80% of puncta that are positive for GluR1 and associated with synaptophysin are also positive for the  $\beta_2AR$  (second bar; a total of 329 synapses in 15 fields in stratum radiatum 25-100 µm away from the pyramidal cell layer from three adult s.d. rats (five fields per rat) were analysed). (C) Post-embedding immunogold staining for the  $\beta_2AR$  in PFC. (a) Three immunopositive axospinous synapses in a single field. (b) A large strongly immunolabelled synapse. Labelling is mainly over PSD, close to postsynaptic plasma membrane (n=2 brains). (**D-F**) GluR1, but not NMDARs, specifically co-immunoprecipitate with  $\beta_2AR$  from rat brain lysates. Rat brain was solubilized with 1% Triton X-100 and cleared by ultracentrifugation before immunoprecipitation (500 µg protein per lysate sample) with the H-20 antibody against  $\beta_2AR$  or a control antibody and immunoblotting with antibodies against proteins indicated on the left side. The H-20 antibody, but not control IgG, immunoprecipitated the  $\beta_2$ AR (**D**; n=3) and GluR1 (**E**; n=6), but not NR2B (**E**; n = 2). The  $\beta_2 AR$  did not co-precipitate with NMDARs (F; n = 3; 25 µg total lysate protein were loaded where indicated).

targeting of AMPARs (Chen et al, 2000; Schnell et al, 2002). Direct interactions between the β<sub>2</sub>AR with PSD-95 and stargazin with both PSD-95 and GluR1 could establish a physical link between the  $\beta_2AR$  and AMPARs (Figure 2C).

To test whether stargazin links the β<sub>2</sub>AR to GluR1 through PSD-95 in neurons, we evaluated whether co-immunoprecipitation of GluR1 with the β<sub>2</sub>AR depends on the presence of stargazin. Stargazer mice (stg<sup>-/-</sup>) lack functional stargazin, the prevailing TARP in adult cerebellum (Chen et al, 2000; Tomita et al, 2003; Menuz and Nicoll, 2008; Menuz et al, 2008). GluR1 co-immunoprecipitated with the β<sub>2</sub>AR from cerebral cortex and cerebellum from wild-type mice (Figure 2D, top left panel). In stg<sup>-/-</sup> mice, GluR1 co-immunoprecipitated with the β<sub>2</sub>AR from cortex, but not from cerebellum (Figure 2D, top right panel). These results indicate a requirement for stargazin for association of the  $\beta_2AR$ with GluR1 in the cerebellum, but not cortex, consistent with

Deoxycholate Triton X-100 Α 3 8 9 PSD-95 DARIH 20 ARHAO Control В 5 MBPβ<sub>2</sub>AR-CT 50 75 GST-PSD-95 domains β<sub>2</sub>AR IP D WT Stg -/-GluR1 Celapallity Lysate GluR Cerebellur Ε mGluR1 PSD-95 mGluR5 AC Gαs Cav1 NR2A G<sub>B</sub>BN<sub>1</sub> NR2B GBBC1 NR1 PKA RII  $\alpha$ Sta

published evidence that other TARPs substitute for stargazin in cortex

As the β<sub>2</sub>AR forms a complex with GluR1, but not with NMDAR, that depends on stg and PSD-95 and their homologues, the interaction of the β<sub>2</sub>AR with PSD-95 or its homologues is governed by other interactions of PSD-95. In other words, it is controlled by the immediate molecular environment of PSD-95. That only certain combinations of binding partners for complex formation with PSD-95 are realized in vivo seems to be critical to avoid physiologically undesirable assemblies or, worse, chaos by random complex formation.

# β<sub>2</sub>AR-GluR1 complex also contains G<sub>s</sub> and adenylyl cyclase

Association of the β<sub>2</sub>AR with GluR1 could allow selective and spatially restricted signalling. If so, the trimeric G<sub>s</sub> protein, adenylyl cyclase, and PKA must also be localized near GluR1. PKA is structurally and functionally linked to GluR1 through

Figure 2 GluR1 forms a complex with the β<sub>2</sub>AR through PSD-95 and stargazin, which also contains G<sub>s</sub> and adenylyl cyclase. (A) Rat brains were extracted with 1% deoxycholate (left) and 1% Triton X-100 (right) before immunoprecipitation with two different antibodies against β<sub>2</sub>AR (H-20, M-20) or control IgG, followed by immunoblotting with anti-PSD-95 ('JH62092' in Sans et al (2000); n=4). Specificity of the observed co-immunoprecipitation of PSD-95 with  $\beta_2AR$  (lanes 4,5,9,10) is indicated by a lack of PSD-95 signal in the control IgG samples (lanes 3 and 8). (B) GST fusion proteins of the PDZ1-2, PDZ2, PDZ3, SH3 and GK domains of PSD-95 and GST alone were immobilized on glutathione Sepharose and incubated with MBP fusion protein of the full cytosolic C-terminus of the human β<sub>2</sub>AR (MBP-β<sub>2</sub>AR-CT) before washing and immunoblotting with anti-MBP antibodies. MBP-β<sub>2</sub>AR-CT bound strongly and specifically to the third PDZ domain (upper panel lane 3), but not to other domains of PSD-95 or to GST alone (lane 6). Reprobing with anti-GST antibodies shows that the different GST fusion proteins were present in comparable amounts (lower panel; n = 8). (**C**) Schematic illustration of the  $\beta_2$ AR–GluR1 complex. Stg or one of its homologues in the AMPAR complex binds with the C-terminus to the first two PDZ domains of PSD-95 or one of its homologues. PSD-95 interacts with its third PDZ domain with the C-terminus of the  $\beta_2AR$ . (D) Loss of stargazin dissociates GluR1 from β<sub>2</sub>AR. Triton X-100 extracts of cortices and cerebella of wild-type and litter-matched stargazer (Stg<sup>-/-</sup>) mice were cleared by ultracentrifugation before immunoprecipitation of β<sub>2</sub>AR and immunoblotting with anti-GluR1 antibodies. GluR1 co-immunoprecipitated with  $\beta_2AR$  from both cortex and cerebellum of wild-type mice (upper left panel) and with the  $\beta_2AR$  from cortex, but not cerebellum of stg<sup>-/-</sup> mice (upper right panel). Lower panels indicate that similar amounts of GluR1 were present in the various extracts. Similar results were obtained in three independent experiments with three wt and three stg $^{-/-}$  mice. (E) Association of  $G_s$ and adenylyl cyclase with the  $\beta_2AR$ -AMPAR complex. Triton X-100 extracts of crude rat forebrain membrane fractions were cleared by ultracentrifugation before immunoprecipitation with antibodies against GluR1 (lane 1), β<sub>2</sub>AR (lane 2), or a non-specific control IgG (lane 3), followed by immunoblotting for the proteins indicated on the left side (Leonard et al, 1998; Davare et al, 2001). Anti-GluR1 and anti- $\beta_2$ AR precipitated PSD-95 (positive control, n=3), adenylyl cyclase (AC, n=3),  $G_{\alpha s}$  (n=3), and  $G_{\beta}$ . The latter was detected with two different antibodies that recognized the N- (BN1; n = 2) and C-termini (BC1; n = 3) of  $G_{\beta 1-4}$  (left panels). PKA (here the RII $_{\alpha}$ subunit; n = 2) and Stg (n = 4) were also present in GluR1 and  $\beta_2$ AR precipitates. The metabotropic glutamate receptors mGluR1 (n=3)and mGluR5 (n=3), caveolin 1(Cav1; n=3) and the NMDA receptor subunits NR2A (n=1), NR2B (n=3), and NR1 (n=1)did not co-immunoprecipitate with either GluR1 or β2AR (right panels), although all proteins were clearly detectable in lysate (lane 4). All immunoprecipitations were made from lysate samples containing 500 µg total protein; lysate aliquots containing 25 µg protein

were loaded when indicated.

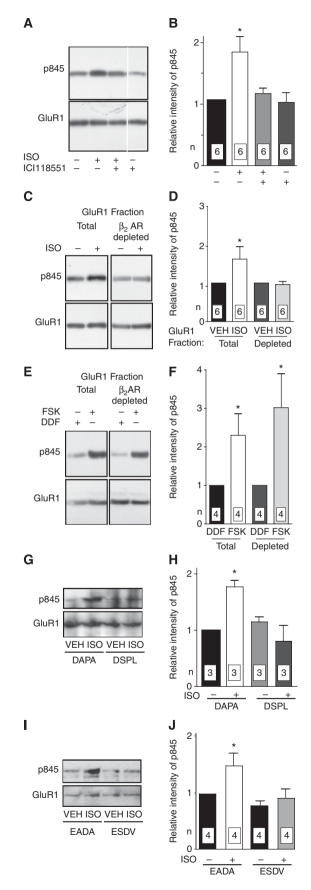
the A kinase anchor protein, AKAP150 (Colledge et al, 2000; Tavalin et al, 2002). AKAP150 associates with SAP97 (Colledge et al, 2000; Tavalin et al, 2002), which in turn binds directly to the C-terminus of GluR1 (Leonard et al., 1998). Figure 2E illustrates that G<sub>s</sub> and adenylyl cyclase are also associated with GluR1: immunoprecipitation of either GluR1 or the  $\beta_2$ AR co-precipitated  $G_{\alpha s}$ ,  $G_{\beta}$ , and adenylyl cyclase in extracts from total forebrain (left upper panels) as well as specifically from PFC and cerebellum (Supplementary Figure 1A). PKA and stargazin also co-precipitated with GluR1 and the  $\beta_2AR$  under these conditions (Figure 2E, left lower panels) as did GluR2 as tested in PFC and cerebellar extracts (Supplementary Figure 1A). As stargazin expression is low in forebrain (Tomita et al, 2003), immunoreactive signals are low in lysate and immunoprecipitates from forebrain. In contrast to these proteins, several other proteins that are likewise concentrated at the postsynaptic site did not co-immunoprecipitate with GluR1 or the  $\beta_2AR$ , including the metabotropic glutamate receptors mGluR1 and mGluR5, and the NMDAR subunits NR1, NR2A, and NR2B (Figure 2E, right panels). The plasma membrane raft marker caveolin 1 was also absent in these immunoprecipitates. Control IgG did not precipitate PSD-95, adenylyl cyclase,  $G_{\alpha s}$ ,  $G_{\beta}$ , PKA, and stargazin. Therefore, we conclude that their co-immunoprecipitation with GluR1 and the  $\beta_2AR$  is specific.

# Regulation of GluR1 S845 phosphorylation by associated β<sub>2</sub>AR

By phosphorylating S845 on the intracellular COOH-terminal region of GluR1 (Roche et al, 1996), PKA promotes activitydriven synaptic targeting of GluR1 (Esteban et al, 2003). To determine whether  $\beta_2AR$ ,  $G_s$ , adenylyl cyclase, PKA, and GluR1 are organized into a functional complex, primary hippocampal cultures were treated with the β-adrenergic agonist isoproterenol (ISO; 3 µM, 15 min). This treatment increased phosphorylation of S845 of GluR1 in hippocampal neurons (Figure 3A and B). The increase was attenuated with the highly selective β<sub>2</sub>AR antagonist ICI118551 (1 μM), which

Figure 3 Localized regulation of GluR1 S845 phosphorylation by the  $\beta_2AR$ . Primary hippocampal cultures (18 DIV) were treated with vehicle (VEH), 3 μM ISO, 10 μM forskolin (FSK), or 10 μM 1,9dideoxyforskolin (DDF; inactive forskolin homologue) for 15 min. Cultures were extracted with Triton X-100 and cleared by ultracentrifugation. GluR1 was immunoprecipitated directly or after pre-immunoprecipitation of β<sub>2</sub>AR complexes (H-20) before immunoblotting with antibodies against phosphorylated S845 (top) and total GluR1 (bottom). Immunosignals were quantified by densitometry. Phospho-S845 (pS845) signals were corrected with respect to total GluR1 signals and normalized to control. (A, B) ISO significantly increased S845 phosphorylation. Pre-treatment for 15 min with 1 μM ICI118551-blocked ISO-induced phosphorylation. All lanes are from same blot and exposure, but non-relevant lanes have been removed between lane 3 and 4. (C-F) After treatments, culture extracts were split into two equal portions. In contrast to the total GluR1 population (left pairs in C, D), GluR1 remaining after depletion of  $GluR1-\beta_2AR$  complexes by pre-immunoprecipitation with H-20 showed no ISO-induced increase in S845 phosphorylation (right pairs in C, D), although forskolin stimulated S845 phosphorylation in both the total and the GluR1-β<sub>2</sub>AR-depleted GluR1 populations (E, F). (G-J) Pre-treatment for 2h with membranepermeant DSPL and ESDV peptides, but not their inactive analogues DAPA and EADA (1 μM each) prevented induction of S845 phosphorylation by ISO. \*P<0.05 compared with control treatments; error bars: s.e.m.; n: number of independent experiments.

by itself reduced basal S845 phosphorylation in some, but not all experiments. We next tested whether the β<sub>2</sub>AR-GluR1 interaction is necessary for S845 phosphorylation, as pre-



dicted if signalling from the β<sub>2</sub>AR to the AMPAR through PKA is localized. After treatment with vehicle or ISO and extraction with Triton X-100, each sample was divided into two equal halves. One half was mock treated before immunoprecipitation of GluR1 to assess the total GluR1 population. The other half was first depleted of GluR1 that was associated with the  $\beta_2AR$  by immunoprecipitation with anti- $\beta_2AR$ antibody. The resulting supernatant lacking  $\beta_2AR$ -GluR1 complexes was then used for immunoprecipitation with anti-GluR1. As before, ISO treatment induced S845 phosphorylation in the total GluR1 pool, but no such increase in phosphorylation was observed for the GluR1 pool from which β<sub>2</sub>AR-associated GluR1 had been removed (Figure 3C and D). An analogous experiment with forskolin (10 µM, 15 min), a direct activator of adenylyl cyclase, led to increased S845 phosphorylation in the total GluR1 as well as the  $\beta_2$ AR-depleted GluR1 populations (Figure 3E and F). These results indicate that a substantial fraction of GluR1 is available for phosphorylation by PKA on S845 on massive stimulation of cAMP production even if not associated with the  $\beta_2AR$ . However, activation of the  $\beta_2AR$  leads to selective phosphorylation of  $\beta_2$ AR-linked GluR1.

To further test whether the complex formation between β<sub>2</sub>AR and GluR1 is necessary for efficient signalling, cultures were pre-treated for 2 h with the membrane-permeable peptide 11R-QGRNSNTNDSPL ('DSPL'). This peptide mimics the extreme C-terminus of the  $\beta_2AR$ , which interacts with the third PDZ domain of PSD-95, and disrupts the co-immunoprecipitation of the β<sub>2</sub>AR with GluR1 (Supplementary Figure 1B). It had little effect on the co-immunoprecipitation of PSD-95 with GluR1, as expected, because the latter engages PDZ domains 1 and 2 of PSD-95 and its homologues. Binding selectivities of the first and second PDZ domains of PSD-95 and related proteins such as SAP97 and SAP102 are very similar to each other, but differ from those of the third PDZ domains (Lim et al, 2002, 2003). Control treatments were performed with its inactive analogue 11R-QGRNSNTNDAPA ('DAPA') altered at the 0 and -2 position (in bold), which are critical for PDZ domain binding. Subsequent incubation with ISO (3 µM, 15 min) led to increased S845 phosphorylation only in samples pre-treated with DAPA, but not in those pre-treated with DSPL (Figure 3G and H). A second pair of peptides gave analogous results: 11R-VYKKMPSIESDV ('ESDV') and its inactive analogue 11R-VYKKMPSIEADA ('EADA') is based on the C-terminus of NR2A. ESDV has optimal binding affinity for PDZ1 and 2 of PSD-95 and its homologues (Lim et al, 2002) and effectively blocks PDZ1 and 2 interactions (Lim et al, 2003). The latter conclusion had been based on immunoprecipitation experiments of PSD-95 with the NMDAR and was now confirmed for the co-immunoprecipitation of PSD-95 with GluR1, which was disrupted as expected (Supplementary Figure 1B). Pre-treatment with ESDV, but not EADA, prevented the ISO-triggered increase in S845 phosphorylation (Figure 3I and J). Collectively, these peptide experiments indicate that PSD-95 and its homologues structurally and functionally connect through PDZ domains the β<sub>2</sub>AR to the GluR1-stargazin complex for S845 phosphorylation.

# Increased surface expression of GluR1 by the associated β<sub>2</sub>AR

The amount of GluR1 is increased at postsynaptic sites on stimulation of S845 phosphorylation by PKA (Swayze et al,

2004; Sun et al, 2005; Man et al, 2007). We asked whether the classic β-adrenergic signalling pathway controls postsynaptic GluR1 accumulation in hippocampal neurons using ectopically expressed GluR1 with superecliptic pHluorin (SEP) at its extracellular N-terminus. The pH-sensitive SEP-GluR1 signal was nearly completely quenched if the extracellular pH was lowered from 7.4 to 6.0 (Supplementary Figure 2), implying that the SEP-GluR1 signal was only detectable if SEP-GluR1 was present at the surface, but not in acidic intracellular organelles. The SEP-GluR1 fluorescence was stable for several hours under our imaging paradigm (data not shown). In initial experiments, 1 µM ISO in combination with the phosphodiesterase inhibitor 3-isobutyl-1-methylxanthine (IBMX; 250 μM) to enhance cAMP levels on β-adrenergic stimulation strongly increased surface expression of SEP-GluR1 at dendritic spines within 15 min, elevating both the intensity for individual puncta and the number of detectable puncta per dendritic length (Supplementary Figure 3). IBMX alone was without effect indicating that signalling was driven by ISO. The ISO effect was completely blocked if the  $\beta_2AR$  antagonist ICI118551 (1 μM) was added immediately before monitoring SEP-GluR1 signals. Subsequent experiments illustrate that a  $5\,\text{min}$  treatment with  $1\,\mu\text{M}$  ISO is sufficient for the full increase in synaptic SEP-GluR1 accumulation (Figure 4).

Similarly, ISO alone (10 µM, 15 min) elevated the number and intensity of GluR1-positive puncta detected by surface labelling of endogenous GluR1 with an antibody against its N-terminus (Figure 5). Double labelling with the synaptic marker synaptophysin confirmed that ISO-induced increases in GluR1 staining occurred at synapses (Figure 5A, B, D, E). This elevation was fully blocked by the  $\beta_2AR$  antagonist ICI118551, but not by the  $\beta_1AR$  antagonist ICI89406 (Figure 5F; Supplementary Figure 4). Disrupting binding of stargazin and its homologues to PDZ1 and 2 of PSD-95 and its homologues with the ESDV peptide or of the  $\beta_2AR$  to PDZ3 of PSD-95 with the DSPL peptide also prevented ISO-induced increases in GluR1 surface expression (Figure 5G; Supplementary Figure 4). The corresponding control peptides EADA and DAPA had no effect (Figure 5G). ICI118551 and more so DSPL (but for unknown reasons not ESDV) showed a clear tendency towards reducing GluR1 surface labelling under control conditions by themselves, which reached statistical significance for changes in synaptic GluR1 puncta frequency (Figure 4D). These results suggest that basal activity of the β<sub>2</sub>AR can enhance GluR1 surface expression under nonstimulated conditions (see also effect of ESDV and DSPL on EPSC in PFC below). At the same time, none of the peptides affected the localization of the  $\beta_2AR$ , PSD-95, or the NMDAR under control conditions (Supplementary Figure 5). These findings exclude a widespread basal effect of these peptides and reassure that the actions of DSPL and ESDV on B2ARmediated effects are due to specific interference with the  $\beta_2$ AR-GluR1 assembly. We conclude that stimulation of the β<sub>2</sub>AR promotes surface expression of GluR1 that resides in a complex with the β<sub>2</sub>AR held together by PSD-95.

## Stimulation of the \$2AR up-regulates postsynaptic AMPAR responses

Activation of BARs can modestly increase synaptic transmission at glutamatergic synapses in hippocampal slices within minutes, although this increase varied and was not observed in all recordings (Dahl and Sarvey, 1989; Thomas et al, 1996;

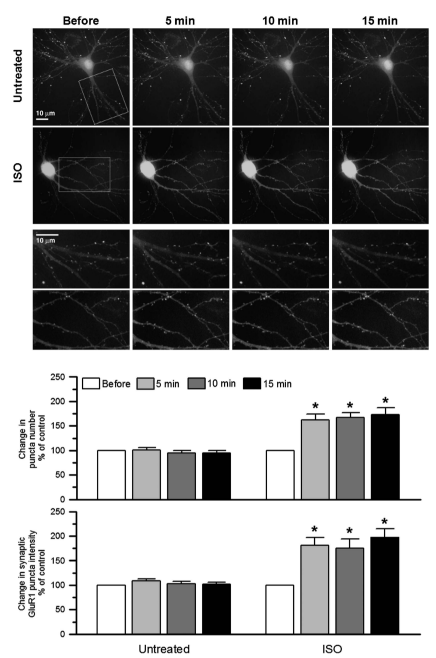


Figure 4 ISO increases SEP-GluR1 surface expression in dendritic spines. Primary hippocampal cultures were transfected with SEP-GluR1 at 5-7 DIV and imaged at 21DIV. ISO (1 μM) increased density and signal intensity of SEP-GluR1 puncta within 5 min, which otherwise remained constant under untreated control conditions. \*P<0.05 compared with control; error bars: SEM. In each case, a minimum of 5 dendrites from 15 neurons in 3 different experiments were analysed.

Katsuki et al, 1997; Winder et al, 1999; Gelinas and Nguyen, 2005). Similarly, we found that 10 µM ISO increased fEPSP initial slope in acute hippocampal slices in 14 out of 23 slices (Supplementary Figure 6A). Averaging all responses gave a statistically significant increase by 24% (Supplementary Figure 6B). Paired-pulse facilitation was unaltered, suggesting that ISO acted postsynaptically (Supplementary Figure 6C).

As ISO-induced up-regulation of fEPSPs was variable in hippocampal slices, we evaluated its effect in the PFC, which receives stronger noradrenergic innervation than the hippocampus (Minzenberg et al, 2008). In acute PFC slices, 10 µM ISO caused a robust and lasting increase of EPSC amplitudes by >50% (Figure 6A), which was completely blocked by the  $\beta_2AR$  antagonist ICI118551 (Figure 6B). ICI118551 itself reduced EPSC amplitude by  $14.4 \pm 2.6\%$  (not illustrated), presumably reflecting basal β<sub>2</sub>AR activity. Paired-pulse facilitation was unaltered in PFC slices (Figure 6C), indicating that the ISO effects were postsynaptic. ISO also significantly increased mEPSC amplitude (Figure 6D and E), but had no effect on mEPSC frequency (Figure 6E), decay time constant (control:  $4.7 \pm 0.5$  ms; n = 6; ISO:  $4.8 \pm 0.32$  ms; n = 5; wash:  $4.7 \pm 0.36$  ms; n = 5), or the 10-90% rise time (control:  $2.7 \pm$ 0.05 ms; n = 6; ISO: 2.7  $\pm$  0.05 ms; n = 5; wash: 2.8  $\pm$  0.07 ms; n = 5). As the ISO effect was more robust and predictable in

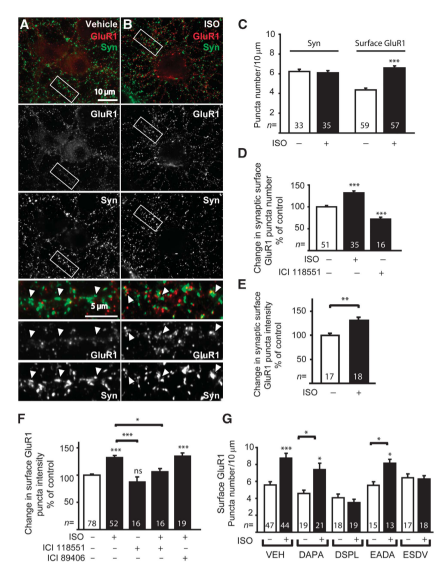


Figure 5 GluR1-associated  $\beta_2$ AR stimulates surface expression of endogenous GluR1. (A, B) Primary hippocampal cultures (18 DIV) were treated with vehicle (VEH) or 10 µM ISO for 15 min, surface labelled with the N-terminal GluR1 antibody (red), fixed, permeabilized, and counter labelled with anti-synaptophysin antibody (green). ISO caused a significant increase in surface expression of GluR1 compared with control cells treated with vehicle as indicated by quantification of total density of GluR1 puncta (C), density of synaptic GluR1 puncta (D; puncta that are colocalized with synaptophysin puncta), and GluR1 staining intensity at synaptic puncta (E). The ISO effect was blocked by 15 min pre-treatment with the  $\beta_2$ AR-selective antagonist ICI118551 (1  $\mu$ M), but not the  $\beta_1$ AR-selective antagonist ICI89406 (1  $\mu$ M; F). (G) Pretreatment for 1 h with membrane-permeant DSPL and ESDV peptides, but not their inactive analogues DAPA and EADA prevented ISO from increasing GluR1 surface expression. \*P < 0.05; \*\*P < 0.001; \*\*\*P < 0.0001 compared with controls treatments or as indicated; error bars: s.e.m.; n: number of neurons analysed in at least three independent experiments for each condition.

PFC than hippocampus slices, we used the PFC for further mechanistic studies.

# Up-regulation of AMPAR responses in PFC by the β<sub>2</sub>AR depends on PKA

To ensure that the ISO response is mediated by PKA, the highly specific PKA inhibitory PKI peptide PKI<sub>5-22</sub> was included in the patch pipette for both evoked EPSC and mEPSC analysis. Dialysis of PKI<sub>5-22</sub> (40 μM) gradually reduced AMPAR-EPSC by  $30.7 \pm 4.4\%$  (n = 9) and blocked the increasing effect of ISO (10.1  $\pm$  3.9%; n = 9; Figure 6F; Supplementary Figure 7A). A scrambled control peptide had no effect on EPSC ( $5.8 \pm 1.5\%$ ; n = 5), and the ISO effect remained intact (51.6  $\pm$  3.5%; n = 5).

Dialysis of  $PKI_{5-22}$ , but not scrambled peptide, also reduced mEPSC amplitude (control:  $14.4 \pm 0.8$ pA; n = 8; PKI<sub>5-22</sub>:  $10.2 \pm 0.8$ pA; n = 8; scrambled peptide:  $13.8 \pm$ 0.6pA; n = 6; Figure 6G; Supplementary Figure 7C and E), but not frequency (control:  $3.9 \pm 0.4 \,\mathrm{Hz}$ ; n = 8;  $PKI_{5-22}$ :  $3.8 \pm 0.4 \,\text{Hz}$ ; n = 8; scrambled peptide:  $3.3 \pm 0.4 \,\text{Hz}$ ; n = 6; Supplementary Figure 7D and F). PKI<sub>5-22</sub> abolished the increasing effect of ISO on mEPSC amplitude ( $PKI_{5-22}$ :  $10.2 \pm 0.8$ pA; n = 8;  $PKI_{5-22} + ISO$ :  $11.6 \pm 0.9$ pA; n = 8; Supplementary Figure 7E), whereas the effect was intact in scrambled peptide-injected neurons (scrambled peptide:  $13.8 \pm 0.6$ pA; n = 6; scrambled peptide + ISO:  $19.3 \pm 0.9$ pA; n = 7, P < 0.001, ANOVA). ISO had also no effect on mEPSC frequency in the presence of  $PKI_{5-22}$  ( $PKI_{5-22}$ : 3.8 ± 0.4 Hz;

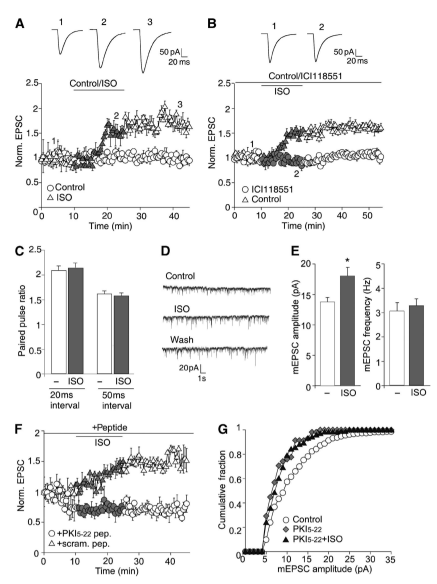


Figure 6 β<sub>2</sub>AR stimulation increases postsynaptic AMPAR-mediated EPSCs in PFC slices. (A) During bath application of ISO (10 μM; grey symbols), the amplitude of evoked AMPAR-EPSC in PFC pyramidal neurons gradually increased by  $53 \pm 7.7\%$  (n = 7), lasting for at least 20 min (after 20 min washout:  $71.4 \pm 23.7\%$ ; n = 7). (B) Bath application of the  $\beta_2$ AR antagonist ICI118551 (1  $\mu$ M) prevented the ISO-triggered increase in AMPAR-EPSC amplitude (109.5  $\pm$  1.8% of control, n = 12), compared with control cells without the antagonist (149.4  $\pm$  4.3%, n = 11). (C) Paired-pulse facilitation (PPF) is unchanged in neurons treated for 15 min with 10 μM ISO (20 ms interval: control: 2.08 ± 0.1, ISO:  $2.14 \pm 0.1$ ; n = 5; 50 ms interval: control:  $1.6 \pm 0.06$ , ISO:  $1.6 \pm 0.06$ ; n = 7; paired-pulse ratio was determined by dividing the amplitude of the second EPSC to the first EPSC). (D, E) Miniature EPSC (mEPSC) recordings from PFC slices. Bath perfusion of ISO (10 µM, 15 min) increased mEPSC amplitude by  $28.6 \pm 2.7\%$  (from  $13.8 \pm 0.7$  pA to  $18.0 \pm 1.4$  pA; n = 6), which remained elevated for at least 20 min (after 20 min washout:  $27 \pm 2.3\%$ ; n = 6). However, there was no significant change for mEPSC frequency (before ISO:  $3.1 \pm 0.35$  Hz; with ISO:  $3.3 \pm 0.3$  Hz; n = 9). (F, G) PKA is required for up-regulation of AMPAR responses by ISO. Injection of PKI<sub>5-22</sub>, but not scrambled peptide (40  $\mu$ M) inhibited EPSC (F) and mEPSC amplitudes (G) and prevented ISO from increasing the amplitudes.

n = 8; PKI<sub>5-22</sub> + ISO:  $3.9 \pm 0.41$  Hz; n = 8) or scrambled peptide (scrambled peptide:  $3.3 \pm 0.4$  Hz; n = 6; scrambled peptide + ISO:  $3.4 \pm 0.16$  Hz; n = 7).

# ISO induces a lasting increase in \$845 phosphorylation through adenylyl cyclase

As the increase in EPSC and mEPSC amplitude by ISO in PFC remained beyond washout of ISO, we tested whether the same would be true for S845 phosphorylation. Incubation of acute PFC slices with 10 µM ISO lead to near maximal phosphorylation within 2 min (Supplementary Figure 8). Importantly, when slices were treated for 15 min with ISO followed by 15 min washout of ISO S845, phosphorylation remained elevated, which explains the continued heightened EPSC and mEPSC responses after removal of ISO.

To corroborate that ISO acts through adenylyl cyclase, which is part of the main signalling pathway downstream of the  $\beta_2AR$  and is present in the GluR1- $\beta_2AR$  complex (Figure 2E; Supplementary Figure 1A), PFC slices were treated with 10 µM ISO for 15 min in the absence and presence of 20 µM SQ22536, an adenylyl cyclase inhibitor. The increase in S845 phosphorylation by ISO was reduced by >50%, indicating that it was largely mediated by adenylyl cyclase (not illustrated).

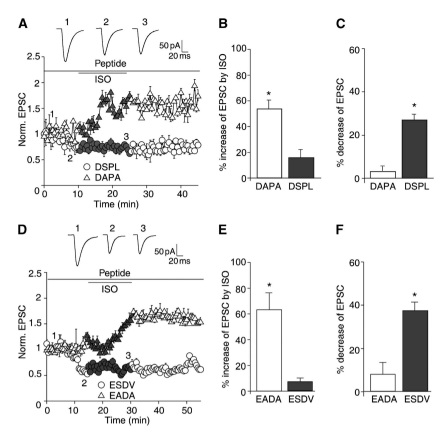


Figure 7 Peptides that disrupt the  $\beta_2$ AR-AMPAR interaction specifically prevent the ISO-induced increase in EPSCs amplitude in PFC slices. (A-C) Basal AMPAR-EPSC amplitudes were significantly reduced by injection of the DSPL peptide ( $50 \mu M$ ) by  $25.2 \pm 4.9 \%$  (n = 5), but not by the control DAPA peptide ( $50 \,\mu\text{M}$ ;  $3.2 \pm 2.2 \,\%$  reduction; n = 5). DSPL diminished the ISO response to an increase of  $15.7 \pm 5.9 \,\%$  (n = 7), but DAPA did not cause any significant reduction (increase was 54.3 ± 6.6%; n=6). (D-F) Basal AMPAR-EPSC amplitudes were significantly reduced by injection of the ESDV peptide ( $50 \mu M$ ) by  $37.5 \pm 4\%$  (n = 10), but not by the control EADA peptide ( $50 \mu M$ ;  $8.8 \pm 4.7\%$  reduction; n = 10). **ESDV** diminished the ISO response to an increase of  $6.8 \pm 2.7\%$  (n = 10), but DAPA did not cause any significant reduction (increase was  $62.7 \pm 12.9\%$ ; n = 10).

# Up-regulation of AMPAR responses by ISO requires β<sub>2</sub>AR-AMPAR association

The DSPL and ESDV peptides were used to test whether upregulation of AMPAR-mediated EPSCs by the β<sub>2</sub>AR depends on PSD-95 binding to TARPs and the β<sub>2</sub>AR. Infusion of either peptide through the whole-cell recording electrode prevented ISO from increasing EPSCs (Figure 7). Similar to ICI118551, both peptides decreased EPSC amplitude by themselves, further supporting the notion that basal activity of GluR1associated β<sub>2</sub>AR enhances AMPAR responses under resting conditions in PFC slices. The corresponding control peptides (DAPA and EADA) showed none of these effects.

# Up-regulation of AMPAR responses by the β<sub>2</sub>AR depends on intact exocytosis

Phosphorylation of S845 up-regulates GluR1 surface expression by promoting its insertion into the plasma membrane (Ehlers, 2000; Swayze et al, 2004; Sun et al, 2005; Oh et al, 2006; Man et al, 2007). It also might increase AMPAR channel activity (Roche et al, 1996). To evaluate whether the β<sub>2</sub>AR up-regulates AMPAR responses mainly by augmenting surface expression, we used two different reagents that inhibit exocytosis. N-ethylmaleimide (NEM) inhibits NEM-sensitive factor, and thereby membrane fusion events required for trafficking of surface proteins to the plasma membrane. Earlier work established that post-

synaptic injection of NEM blocks LTP without affecting basal postsynaptic responses (Lledo et al, 1998). Similarly, NEM infusion did not alter EPSCs in PFC neurons under basal conditions, but prevented up-regulation of EPSCs by ISO (Figure 8A and C). Botulinum neurotoxin B (BoTox) and tetanus toxin (TeTox) block exocytosis by proteolytic cleavage of synaptobrevin/VAMP2, which is critical for exocytosis (Schiavo et al, 2000). Injection of the catalytically active BoTox or TeTox light chain does not reduce basal postsynaptic responses, but abrogates LTP (Lledo et al, 1998; Lu et al, 2001). Similar to NEM, BoTox blocked the ISO-triggered increase in EPSC amplitude (Figure 8B and C). These data show that ISO increases postsynaptic AMPAR responses mainly by promoting insertion of AMPARs into the cell surface.

# Discussion

Our data indicate that the  $\beta_2AR$ ,  $G_s$ , and adenylyl cyclase form a supramolecular signalling complex with GluR1, which is known to associate with PKA and the antagonistic phosphatase PP2B through AKAP150, SAP97, and, alternatively, PSD-95 (Leonard et al, 1998; Colledge et al, 2000; Tavalin et al, 2002). Remarkably, only GluR1-containing complexes that are associated with β<sub>2</sub>ARs are effectively regulated by β-adrenergic stimulation, showing that this assembly

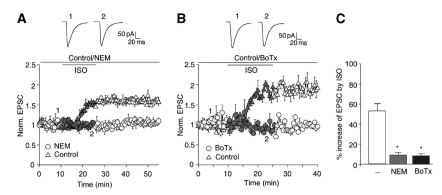


Figure 8 Exocytosis blockage prevents the ISO-induced increase in EPSCs amplitude in PFC slices. (A) Dialysis of NEM (5 mM) itself did not significantly alter AMPAR-EPSC amplitude ( $104.3 \pm 1.4\%$  of control; n = 11), but decreased the effect of ISO to an increase of  $11.8 \pm 2.6\%$ stimulation (n = 11). (B) Dialysis of Botulinum neurotoxin (0.5  $\mu$ M) itself did not significantly alter AMPAR-EPSC amplitude (102.8  $\pm$  1.5%; n=8), but reduced the effect of ISO to an increase of  $10\pm1.9\%$  stimulation (n=10). (C) Averages in changes of EPSC amplitudes from all experiments from panels A and B.

allows for highly localized control of GluR1 phosphorylation by the  $\beta_2AR$ .

Another study suggests that the  $\beta_1AR$ , but not  $\beta_2AR$ , regulates S845 phosphorylation (Vanhoose and Winder, 2003), contrasting our findings. The differences between this and our study that lead to this discrepancy outcome could be related to differences in the exact experimental system, though quite similar, or the high concentrations of the  $\beta_1$  selective blocker betaxolol (10  $\mu$ M) used in the other study at which isotype specificity could have been lost (e.g. Smith and Teitler, 1999; Sharif and Xu, 2004). In addition, there was no evidence presented that the ICI118551 batch was active in the other study. However, we do not want to exclude that the β<sub>1</sub>AR can contribute to S845 phosphorylation under certain conditions.

Phosphorylation of S845 by PKA promotes GluR1 surface expression because of a combination of reduced internalization rate and increased re-insertion rate (Ehlers, 2000; Man et al, 2007). We now show that, upstream of PKA, activation of the β<sub>2</sub>AR specifically augments the synaptic expression of GluR1. S845 phosphorylation might also enhance the activity of GluR1-containing AMPARs by increasing their open probability (Banke et al, 2000), which could have contributed to the increase in AMPAR responses we observed in acute hippocampal and PFC slices. However, ISO-induced increases in the PFC were largely blocked by infusion of BoTox indicating that increased activity of AMPARs that were already present at postsynaptic sites could have made only a modest contribution.

The increase in GluR1 surface expression contrasts with the decrease of  $\beta_2AR$  typically induced by  $\beta_2AR$  ligands. However, agonist-induced internalization typically reduces surface  $\beta_2AR$  by <50%, and in some cases as little as 10% (e.g. Shenoy et al, 2006). Furthermore, association of the β<sub>1</sub>AR with PSD-95 inhibits its ligand-induced internalization (Xu et al, 2001). Analogously, the  $\beta_2$ AR might be resistant to internalization when forming a complex with PSD-95, stargazin, and GluR1.

The β<sub>2</sub>AR-dependent increase in S845 phosphorylation emerges as an important consequence of adrenergic activation throughout the brain. β-adrenergic stimulation can increase synaptic transmission at glutamatergic synapses in hippocampal slices within minutes, although that is not observed in all experiments (Dahl and Sarvey, 1989; Thomas et al, 1996; Katsuki et al, 1997; Winder et al, 1999; Gelinas and Nguyen, 2005). At the same time, β-adrenergic activation fosters induction of LTP, which can transform a reversible increase in synaptic strength into a lasting one (Thomas et al, 1996; Katsuki et al, 1997; Winder et al, 1999; Lin et al, 2003; Gelinas and Nguyen, 2005). Recent work indicates that norepinephrine and emotional stress induce phosphorylation of GluR1 S845 and to a lesser degree on S831, a phosphorylation site for PKC and CaMKII (Hu et al, 2007). Mutating both sites to alanine interferes with norepinephrine-dependent LTP induction by a 10 Hz/90 s tetanus and fear conditioning by a modest conditioning paradigm (Hu et al, 2007). Furthermore, a single mutation of S845 to alanine prevents ISO from priming neurons in the visual cortex for spike time-dependent potentiation (Seol et al, 2007).

In closing, the localized signalling from the  $\beta_2AR$  to GluR1 described here represents a critical regulatory mechanism for postsynaptic functions linking the noradrenergic and glutamatergic systems in the brain. This mechanism is likely to have a major function in mediating noradrenergic effects on the brain, including heightened arousal and facilitation of synaptic plasticity and learning (Cahill et al, 1994; Nielson and Jensen, 1994; Berman and Dudai, 2001; Strange et al, 2003; Strange and Dolan, 2004; Hu et al, 2007). These findings also illustrate the importance of noradrenergic inner vation of the hippocampus, although this innervation is typically eliminated in experiments using hippocampal slices, but must not be neglected. Finally, the characterization of the  $\beta_2AR-G_s$ -adenylyl cyclase-PKA-GluR1 complex reported here illuminates the capability of the  $\beta_2AR$  to drive spatially restricted generation of cAMP. This work firmly establishes the concept of locally restricted and thereby highly selective signalling by cAMP, through formation of supramolecular signalling complexes.

# Materials and methods

### Reagents and antibodies

ISO, ICI118551, forskolin, dideoxyforskolin, and NEM were from Sigma and ICI89406 from Biotrend Chemicals Inc. A peptide derived from the C-terminus of GluR1 (MSHSSGMPLGATGL) was purified by HPLC, cross-linked with glutaraldehyde to bovine serum albumin and injected into rabbits as described (Davare et al, 1999) to obtain anti-GluR1 antiserum. For immunoprecipitation of

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GluR1,  $2\mu l$  of this serum were used in  $0.5\,ml$  extracts. For immunoblotting, this serum was diluted 1:1000. Antibodies against the extracellular N-terminus of GluR1 were from Oncogen Research, Bassoon from Stressgen,  $\beta_2AR$  (H-20 and M-20) from Santa Cruz, and  $G_{\alpha s}$  from Upstate. Non-specific rabbit IgG was from Zymed. Anti-PSD-95 was produced earlier by us and corresponds to 'JH62092' in Leonard and Hell (1997), Leonard et al (1999), Sans et al (2000), and Davare et al (2001). All other antibodies including those against  $G_{\beta}$ , pan-adenylyl cyclase, mGluR1, mGluR5, caveolin-1, NR1, NR2A, and NR2B and the phospho-specific antibody against S845 were as described earlier (Leonard and Hell, 1997; Leonard et al, 1999; Sans et al, 2000; Davare et al, 2001; Lu et al, 2007). The membrane-permeable peptides 11R-QGRNSNTNDSPL (DSPL), 11R-QGRNSNTNDAPA (DAPA), 11R-VYKKMPSIESDV (ESDV), and 11R-VYKKMPSIEADA (EADA) were custom synthesized by the W. M. Keck Biotechnology Resource Center, Yale University and HPLC purified. Botulinum Neurotoxin Type B Light Chain (BoTox) was from Listlabs. All other reagents were from the usual suppliers and of standard quality.

### *Immunocytochemistry*

Adult male Sprague-Dawley rats were deeply anesthetized with sodium pentobarbital and perfused with 4% paraformaldehyde following NIH guidelines as described (Burette et al, 1999) and approved by the University of North Carolina. Fifteen minutes after the onset of fixation, material was flushed with normal saline. Brains were removed, tissue sections cut on a Vibratome and processed for multiple label immunofluorescence. After blocking, rabbit anti-β<sub>2</sub>AR (1:10000; H-20) was added, followed by horseradish peroxidase-conjugated secondary antibody. Fluorophoreconjugated tyramide was then used to amplify the signal followed by PBS washes (Burette et al, 1999). Subsequently, standard double labelling with rabbit anti-GluR1 antibody (1:1000) and mouse-antisynaptophysin (1:5000) was performed. Quantitative data were collected by confocal microscopy from stratum radiatum of hippocampal CA1, at least 25 µm from the pyramidal layer (Melone et al, 2005); 329 synapses from five fields from each of three rats were analysed for quantification of the number of β<sub>2</sub>AR- and GluR1labelled puncta (Melone et al, 2005). We considered only neuropil, avoiding somata and dendritic shafts. To focus specifically on synaptic labelling, we only considered immunopositive puncta clearly associated with puncta stained for synaptophysin.

# Immunogold labelling

Pentobarbital-anesthetized adult rats were perfused with a mixture of 2% depolymerized paraformaldehyde and 2% glutaraldehyde in phosphate buffer, after a brief saline rinse, as per institutionally approved protocols and NIH guidelines. A total of 200 µm Vibratome sections were cryoprotected in 30% glycerol. Chunks of prefrontal/orbital cortex were plunge frozen in isopentane and dehydrated over several days at  $-60^{\circ}$  in ethanol with 1% uranyl acetate, infiltrated with Lowicryl HM-20, and UV-polymerized at low temperature. Thin (~80-100 nm) sections were collected on nickel mesh grids and processed for immunogold labelling, using the M20 antibody at 1:1000 dilution, according to Phend et al (1992, 1995). Grids were examined on a Philips Tecnai transmission electron microscope.

### Immunoprecipitation and immunoblotting

Sprague-Dawley rats were obtained from Harlan and homozygous stargazer mice and litter-matched wild-type control mice from Jackson Laboratories. The Animal Care and Use Committee of the University of Iowa approved all procedures involving animals. Forebrains, neocortices, or cerebella were homogenized in a 10-fold volume of Buffer A (150 mM NaCl, 10 mM EDTA, 10 mM EGTA, 10 mM Tris-HCl, pH 7.4, and protease inhibitors) containing 1% deoxycholate or 1% Triton X-100, and cleared from non-solubilized material by ultracentrifugation (minimally 250 000 g for 30 min) before immunoprecipitation with the C-terminal GluR1 antibody, H20 against the  $\beta_2AR$ , or an equivalent amount of non-specific rabbit IgG and subsequent immunoblotting with the indicated antibodies as described (Leonard and Hell, 1997; Leonard et al,

# GST and MBP fusion proteins and pull-down assay

GST fusion proteins of the PDZ1-2 (residues 1-185), PDZ2 (156-248), PDZ3 (302-402), SH3 (431-500), and GK (534-724) domains

of PSD-95 and an MBP fusion protein of the full cytosolic C-terminus of the human  $\beta_2AR$  (MBP- $\beta_2AR$ -CT; residues 326–413) were expressed in Escherichia coli as described (Seabold et al, 2003). GST fusion proteins were bound to glutathione Sepharose 4B (Amersham), washed, incubated with MBP-B2AR-CT for 90-120 min at 4°C in TBS (10 mM Tris, 150 mM NaCl, 0.1% TX100; pH 7.4), and washed. Immunoblotting was performed with a monoclonal anti-MBP antibody (NEB), stripped with SDS and dithiothreitol and reprobed with anti-GST antibody (Leonard et al, 1998).

#### Hippocampal cultures and phosphorylation studies

Hippocampal cultures were prepared from E18 embryonic Sprague-Dawley rats (Harlan) and maintained as described (Lim et al, 2003) using the B27-derived culture medium supplement NS21 (Chen et al, 2008). Cultures (DIV18) were pre-incubated for 15 min with the  $\beta$ AR antagonists ICI118551 ( $\beta$ <sub>2</sub>AR; 1  $\mu$ M) or ICI 89406 ( $\beta$ <sub>1</sub>AR; 1 μM) if indicated before incubation with vehicle (water) versus ISO  $(1-3 \mu M)$  or forskolin  $(10 \mu M)$ ; from DMSO stock) versus 1.9-dideoxyforskolin (10 uM) for 15 min. For immunoprecipitation and analysis of S845, phosphorylation cultures were extracted with Triton X-100 in Buffer A containing the phosphatase inhibitors microcystin (0.4  $\mu$ M) and para-nitriphenyl phosphate (100  $\mu$ M). GluR1 was either directly immunoprecipitated or immunoprecipitated after initial immunoprecipitation with H-20 to remove GluR1- $\beta_2$ AR complexes before subsequent immunoblotting with antibodies against GluR1 phosphorylated at S845 and, after stripping, with the C-terminal GluR1 antiserum. Relative phosphorylation levels were determined by densitometry of ECL signals on film. After scanning, pixel intensity of bands was quantified using Adobe Photoshop. Signals were in the linear range as shown earlier (Davare and Hell, 2003; Hall et al, 2006). Values were corrected for any variation in total GluR1 loaded and normalized to control treatment.

### GluR1 surface labelling in hippocampal cultures

For surface labelling of GluR1, cultures were incubated after drug treatments with the N-terminal GluR1 antibody for 20 min at 37°C, fixed with 4% paraformaldehyde and 4 % sucrose for 15 min at room temperature, and incubated with Alexa 568 anti-rabbit secondary antibody (Molecular Probes) for 1 h at room temperature under non-permeabilized conditions (Swayze et al, 2004; Lise et al, 2006). For double labelling of GluR1 surface stained cultures, neurons were permeabilized with cold methanol for 5 min and incubated with anti-synaptophysin antibody overnight, followed by incubation with the appropriate secondary antibody. Images were taken using a × 63 objective on a Zeiss Axiovert M200 microscope. To allow comparison between different specimens, pictures were taken under the same settings. To correct for out-of-focus clusters within the field of view, focal plane (z) stacks were acquired and maximum intensity projections performed offline. Images were scaled to 16 bits and analysed in Northern Eclipse (Empix Imaging, Mississauga, Canada), by using user-written software routines (Gerrow et al, 2006; Lise et al, 2006). To measure puncta intensity and number per dendrite length, an experimenter blind to the treatment conditions manually outlined dendrites (at least 300 µM). Puncta were defined as sites of intensity at least twice the background. Average integrated puncta intensity was obtained by subtracting the average background grey from the average puncta grey intensities, which was then multiplied by the average puncta area. GluR1 puncta colocalizing with synaptophysin puncta were considered synaptic. At least eight neurons from two to three independent experiments were analysed. The s.e.m. values were calculated based on number of neurons examined. Two-tailed parametric Student's t-tests were performed to calculate the statistical significance of differences between two groups; oneway ANOVA was used to compare three or more groups, followed by Tukey-b post hoc analysis for multiple comparisons.

# Monitoring SEP-GluR1 surface expression in hippocampal

SEP was kindly provided by Dr Gero Miesenbock (Yale University, New Haven, CT). Its coding sequence was amplified by PCR (QuikChange protocol; Stratagene, La Jolla, CA) and inserted into the N-terminus of rat GluR1 after a signal peptide in pRK5 vector. Rat hippocampal cultures (DIV 5-7) were co-transfected with plasmids encoding DsRed (used to find transfeceted cells not to bleach GFP) and SEP-GluR1 by the calcium phosphate method. At DIV 21, live cell images were captured with a system consisting of a BD CARVII<sup>TM</sup> confocal imager connected to a Leica DMIRE2 fluorescence microscope and a Hamamatsu EM CCD camera with filter sets. The X-Y coordinates of individual neurons on a mechanical X-Y stage were recorded within the IPLab4 software program, so that the same neuron could be located and photographed at different time periods during treatment. To maintain the neuron viability, imaging was carried out with a temperature-controlled stage (37°C) fitted with a CO<sub>2</sub> chamber set at 5% (Leica Microsystems). After capturing an image, the culture dish was immediately put back to a humidified 5% CO2 incubator at 37°C for prolonged treatment.

Stacks of confocal images (not >25 z-planes) at 0.5 μm intervals were merged into one single image before analyses. All digital images were analysed with IPLab4 software (BD Biosciences). SEP-GluR1 puncta were defined as either dendritic protrusions with expanded heads that were 50% wider than their necks or regions of intensity at least twice the dendritic intensity. To quantify the SEP-GluR1 and dendritic intensity, the outline of puncta and the middle line on dendrites were manually drawn and these parameters were measured using the IPlab4 software. Average blank field intensity used as background was subtracted from these measurements to yield the actual intensity of SEP-GluR1. Data are expressed as mean ± s.e.m. of the indicated number of experiments. Statistical significance was determined using a paired t-test (Sigma plot 7.0; Systat software, San Jose, CA) in comparing the puncta density and fluorescence intensity before and after treatment of the same neuron.

### Electrophysiological recording from PFC slices

All experiments were carried out with the approval of State University of New York at Buffalo Animal Care Committee following NIH guidelines. Slices were prepared from 3-5-week-old Sprague-Dawley rats as described before (for details see, Wu et al, 2005). In brief, animals were anesthetized by inhaling 2-bromo-2-chloro-1,1,1-trifluoroethane (1 ml/100g; Sigma) and decapitated. Brains were quickly removed and sliced (300 µm) with a Leica VP1000S Vibratome, whereas bathed in HEPES-buffered salt solution. Slices were then incubated for 1-5 h at room temperature (22-24°C) in a NaHCO<sub>3</sub>-buffered saline (EBSS) bubbled with 95% O<sub>2</sub>, 5% CO<sub>2</sub>.

AMPAR-mediated synaptic transmission was monitored by standard whole-cell patch recordings from pyramidal neurons located in layer V in PFC slices (Yuen et al, 2007). PFC slices were perfused with oxygenated artificial cerebro-spinal fluid (ACSF) containing (mM) NaCl (130), NaHCO<sub>3</sub> (26), CaCl<sub>2</sub> (1), MgCl<sub>2</sub> (5), KCl (3), glucose (10), NaH<sub>2</sub>PO<sub>4</sub> (1.25) plus GABA<sub>A</sub>R antagonist bicuculline (10 µM) and NMDAR antagonist D-aminophosphonovalerate (APV,  $25\,\mu\text{M}$ ). Patch pipettes (5-8 M $\Omega$ ) were filled with internal solution containing (mM) Cs-methanesulfonate (130), CsCl (10), NaCl (4), MgCl<sub>2</sub> (1), HEPES (10), EGTA (5), QX-314 (2.2), phosphocreatine (12), MgATP (5), Na<sub>2</sub>GTP (0.5), and leupeptin (0.1); pH 7.2-7.3, 265-270 mOsm. Neurons were observed with a  $\times$  40 water-immersion lens and illuminated with near infrared (IR) light, and the image was captured with an IR-sensitive CCD camera. Recordings were performed using a Multiclamp 700A amplifier (Axon Instruments). Tight seals  $(2-10 \text{ G}\Omega)$  were obtained by applying negative pressure. The membrane was ruptured with additional suction to obtain whole-cell configuration. Access resistance (13–18 M $\Omega$ ) was compensated by 50–70%. Neurons were held at  $-70 \,\mathrm{mV}$ . EPSCs were evoked by stimulating the neighbouring neurons (50  $\mu s$  pulse) with a bipolar tungsten electrode (FHC, Inc.). To record mEPSC in PFC slices, slices were perfused with a modified ACSF containing a low concentration of MgCl<sub>2</sub> (1 mM) and TTX (1 µM). Data analysis was performed with Clampfit (Axon Instruments). Spontaneous synaptic events were analysed with Mini Analysis Program (Synaptosoft, Leonia, NJ). Statistical comparisons of the amplitude and frequency of mEPSC were made using the Kolmogorov-Smirnov test.

### Supplementary data

Supplementary data are available at The EMBO Journal Online (http://www.embojournal.org).

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