Modulation of Calcium-Dependent Inactivation of L-Type Ca$^{2+}$ Channels via β-Adrenergic Signaling in Thalamocortical Relay Neurons

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Abstract

Neuronal high-voltage-activated (HVA) Ca$^{2+}$ channels are rapidly inactivated by a mechanism that is termed Ca$^{2+}$-dependent inactivation (CDI). In this study we have shown that β-adrenergic receptor (βAR) stimulation inhibits CDI in rat thalamocortical (TC) relay neurons. This effect can be blocked by inhibition of cAMP-dependent protein kinase (PKA) with a cell-permeable inhibitor (myristoylated protein kinase inhibitor-(14–22)-amide) or A-kinase anchor protein (AKAP) St-Ht31 inhibitory peptide, suggesting a critical role of these molecules downstream of the receptor. Moreover, inhibition of protein phosphatases (PP) with okadaic acid revealed the involvement of phosphorylation events in modulation of CDI after βAR stimulation. Double fluorescence immunocytochemistry and pull down experiments further support the idea that modulation of CDI in TC neurons via βAR stimulation requires a protein complex consisting of CaV1.2, PKA and proteins from the AKAP family. All together our data suggest that AKAPs mediate targeting of PKA to L-type Ca$^{2+}$ channels allowing their phosphorylation and thereby modulation of CDI.

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Introduction

Voltage-gated Ca$^{2+}$ channels of the plasma membrane consist of three subfamilies (CaV1, CaV2 and CaV3) [1]. They are composed of 10 pore-forming α1 channel subunits and are important components of a universal cellular Ca$^{2+}$ signaling tool kit [2]. Voltage-dependent Ca$^{2+}$ channels are one of the main routes of cellular Ca$^{2+}$ entry. Intracellular Ca$^{2+}$ ions control processes as diverse as cell proliferation, neuronal development and transmitter release [2]. All of these functions have to be accomplished within a narrow range of Ca$^{2+}$ concentrations. CDI of voltage-dependent Ca$^{2+}$ channels is an auto-inhibitory feedback mechanism controlling Ca$^{2+}$-influx [3,4]. Previously we have shown that in TC neurons of the dorsal part of the lateral geniculate nucleus (dLGN), Ca$^{2+}$-induced Ca$^{2+}$ release (CICR) contributes to intracellular Ca$^{2+}$ transients [5], leads to the activation of Ca$^{2+}$-dependent K$^+$ channels and thereby supports regular tonic firing [6]. Furthermore, CDI, which is under the control of multiple biochemical and activity-dependent mechanisms, has been shown to limit Ca$^{2+}$ entry into TC neurons [7,8,9,10]. Stimulation of the βAR/adenyl cyclase (AC)/PKA-dependent pathway in TC neurons mediates behavioural state-dependent shifts in thalamic activity modes by modulating pacemaker channels, L-type Ca$^{2+}$ channels, and Ca$^{2+}$-dependent K$^+$ channels [3,11]. Direct application of cAMP and the catalytic subunit of PKA reduced the degree of CDI in TC neurons [9]. Although cAMP-dependent signaling and CDI represent prominent mechanisms in TC neuron function, their possible functional coupling by direct stimulation of βAR has not been investigated in these neurons yet. Recent studies in cardiac [12] as well as in hippocampal cells have shown a functional link between βAR and one type of L-type Ca$^{2+}$ channels, namely CaV1.2, via PKA [13,14], however a possible link to CDI has not been addressed. Furthermore, AKAP has been shown to be an important element in organizing βAR-dependent pathways in neurons [13,15,16]. Although dephosphorylation of Ca$^{2+}$ channels by calcineurin (PP2B) has originally been proposed to be the crucial mechanism of CDI [17], calmodulin closely tethered to the channel has been identified as the Ca$^{2+}$ sensor and central mediator of this process [18,19,20]. The role of phosphorylation/dephosphorylation in CDI attracted less attention, although the close association of CaV1.2 channels, phosphorylating PKA, and dephosphorylating calcineurin has been shown [17]. Based on the finding that βARs are directly associated with one of their main effector channels, namely CaV1.2, via a protein complex also containing G-proteins, AC, PKA, and a counterbalancing protein phosphatase [13], it was also suggested that this protein complex might be the basis for β-adrenergic modulation of CDI [3] (Figure S1). Here we provide
evidence for this hypothesis and show that stimulation of βARs reduces CDI in TC neurons via a protein complex including AKAP and PKA.

Materials and Methods

Ethics Statement

All experiments were carried out in accordance with the European Communities Council Directive (86/609/EEC) and approved by the local animal care committee (Landesverwaltungs- samt Sachen-Anhalt, AZ: 42502/5.19 UNI MD).

Reverse transcription-polymerase chain reaction (RT-PCR) assays

Poly A+ mRNA was prepared from freshly dissected thalamic tissue (the ventrobasal thalamic complex (VB) and dLGN were visually identified) or whole brain by extraction with Trizol reagent according to the manufacturer's instruction (Qiagen, Hilden, Germany). First-strand cDNA synthesis was primed with oligo (dT) from 0.5–1 μg of mRNA, using the SuperScript II enzyme (Invitrogen Life Technologies). Specific primers for calcium channels α1C-α1E, G-proteins, adenylcyclase, and β-receptors were used according to manufacturer's description (Figure 1A–1F). Accession numbers for used primers were: Alpha1S (nucleotides 445–828) No. L04684; Alpha1C (nucleotides 2624–3033) No. NM012517; Alpha1D (nucleotides 3691–4200) No. NM017298; Alpha1F (nucleotides 3196–3660) No. NM0053701; Alpha1A (nucleotides 5115–5387) No. NM012918; Alpha1B (nucleotides 1742–2051) No. NM147114; Alpha1E (nucleotides 3031–6377) No. NM012924; G alpha i–1 (nucleotides 669–1198) No. AF234260; G alpha i–2 (nucleotides 139–554) No. AF239674; G alpha i–3 (nucleotides 1993–2367) No. NM013106; β1–AR (nucleotides 742–1104) No. NM012701; β2–AR (nucleotides 676–964) No. NM012492; β3–AR (nucleotides 390–797) No. NM013108; AC1 (nucleotides 1361–1863) No. XM223616; AC6 (nucleotides 1531–1940) No. NM012812; AC8 (nucleotides 3001–3480) No. NM017142; AKAP5 (nucleotides 122–389) No. NM 133515; AKAP7 (nucleotides 494–1059) No. NM 00101801; Gs (nucleotides 332–756) No. NM 019132.

Cell type-specific RT - PCR

Brain tissue from P14–P24 Long-Evans rats consisting of dLGN was sliced using a vibratome to 500 μm. After trypan blue staining, the slices were transferred into 3 ml carrier RNA buffer (RNeasy Micro Kit, Qiagen) by breaking the tip of the pipette and expelling 3 μl of solution with positive pressure. The pipette solution (6 μl) contained 5% sucrose, 10% glucose, 10. Slices were kept for 20 min at 34 °C, then oxygenated slicing solution (pH 7.35, with 95% O2-5% CO2) was added to the bath solution in order to prevent oxidation of the drugs. Slices were transferred to the oxygenated slicing solution (pH 7.35, with NaOH) containing the extracellular solution (ACSF; pH 7.35, with 95% O2–5% CO2). Complete cells were sucked into the pipette and depolarized with a double-pulse protocol in which conditioning pulses were used to a fixed potential of V. Series resistance compensation was routinely used (30%). Voltage-clamp experiments were governed by Pulse software (HEKA, Germany). Typical electrode resistances amounted to 2–4 MΩ, while access resistance was 5–15 MΩ. Series resistance compensation was routinely used (30%). Voltage-clamp experiments were conducted with Pulse software (HEKA, Germany). For standard recordings the following solutions were used: (i) extracellular solution (in mM): NaCl, 125; KCl, 2.5; NaH2PO4, 1.25; NaHCO3, 22–26; MgSO4, 2; CaCl2, 2; and glucose, 10; 1 min at 60 °C before being cooled to room temperature, and allowed to rest for 60 to 90 min.

Quantitative Real – time PCR

Real-time PCR was performed using the Real PCR Master Mix 2.5X (Eppendorf) and the ABI Prism 7000 Sequence Detection System (Applied Biosystems); PCR program was: 2 min at 50 °C, 10 min at 95 °C, 50 cycles: 15 s at 95 °C and 1 min at 60 °C. Analysed probes were AKAP 5, AKAP 7, and β-2 microglobulin as the internal reference gene used for normalisation (Figure 2B). All primers were purchased from Applied Biosystems. Results were analysed with the ABI Prism 7000 SDS software. The efficiency of real-time primer/probes was nearly identical. Quantification was done using the comparative Ct method as described earlier [21]. In a standard PCR GAPDH (nucleotides 799–1028, accession No. P28291) expression was checked as a positive control to confirm the integrity of transcribed cDNA. The PCR protocol for GAPDH amplification was: 3 min 94 °C; 50 cycles (30 s 94 °C, 1 min 61 °C, 1 min 72°C); 7 min 72°C.

Tissue Preparation

Thalamic slices were prepared from juvenile postnatal day (P) 12–21 Long-Evans rats. After anaesthesia with isoflurane, animals were decapitated and a block of tissue containing the dLGN was rapidly removed and placed in chilled (2–4 °C), oxygenated slicing solution (pH 7.35, with NaOH) containing the following (in mM): sucrose, 195; glucose, 11; Pipes, 10; MgCl2, 5; MgSO4, 10; and CaCl2, 0.5. Coronal slices of the thalamus were cut at 300 μm on a vibratome and kept submerged in artificial cerebrospinal fluid (ACSF; pH 7.35, with 95% O2–5% CO2) containing the following (in mM): NaCl, 125; KCl, 2.5; NaH2PO4, 1.25; NaHCO3, 22–26; MgSO4, 2; CaCl2, 2; and glucose, 10. Slices were kept for 20 min at 34 °C before being cooled to room temperature, and allowed to rest for 60 to 90 min.

Patch clamp recordings

Whole-cell recordings under voltage clamp condition were performed on visually identified TC neurons of the dLGN at room temperature (21–23 °C) using borosilicate glass pipettes (GC150TF-10, Clark Electromedical Instruments, Pangbourne, UK) connected to an EPC-9/2 amplifier (HEKA Electronics, Lamprecht, Germany). Typical electrode resistances amounted to 2–4 MΩ, while access resistance was 5–15 MΩ. Series resistance compensation was routinely used (30%). Voltage-clamp experiments were governed by Pulse software (HEKA, Germany). For standard recordings the following solutions were used: (i) extracellular solution (in mM): NaCl, 125; KCl, 2.5; NaH2PO4, 1.25; NaHCO3, 22–26; MgSO4, 2; CaCl2, 2; and glucose, 10; TTX, 0.001; 4-AP, 4; pH 7.35 with NaOH. (ii) Intracellular solution: Cs-glucuronate, 85; Cs3-citrate, 10; NaCl, 10; KC1, 1; EGTA, 1.1; CaCl2, 0.1; MgCl2, 0.25; HEPES, 10; TEA-Cl, 15; Mg-ATP, 3; Na2-GTP, 0.5; pH 7.25 with CsOH. In some experiments equimolar concentrations of ascorbic acid were added to the bath solution in order to prevent oxidation of the drugs. HVA Ca2+ currents were evoked from a holding potential of −40 mV by a double-pulse protocol in which conditioning pulses to varying potentials (−40 to +60 mV, 200 ms duration) were followed by a brief gap (−40 mV, 50 ms) and a subsequent analyzing test pulse to a fixed potential of +10 mV (200 ms) (Figure 3A). For standard recordings Ca2+ was used as a charge carrier and 1.1 mM EGTA was included to the intracellular solution. The degree of inactivation (Dmax) was determined by dividing the current amplitudes elicited by a test pulse to +10 mV with maximal and minimal preceding conditioning current amplitude (Figure 3B–3C). Dmax = \[1 - \frac{I_{test}}{I_{max}}\] x 100(%). A2 (I_{max} Figure 3C) represents the minimal current amplitude after a preceding conditioning pulse, as measured at the beginning of
the voltage step to +10 mV (upper arrowhead in Figure 3B), and $I_{\text{max}}$ (Figure 3C) representing the maximal current amplitude (lower arrowhead, Figure 3B), as measured without a preceding conditioning pulse. GraphPad Prism 5.0 and Microcal Origin 6.0 software were used for data analysis and figure plotting.

**Drugs**

The following drugs were used: isoproterenol hydrochloride, propranolol hydrochloride, xamoterol hemifumarate, salmeterol, BRL 37344, sodium salt, CGP 20712 dihydrochloride, ICI 118,551 hydrochloride, SR 59230A hydrochloride, okadaic acid, PKA inhibitor myristoylated PKI 14–22 amide (all from Tocris, USA). Equimolar concentrations of ascorbic acid were added to the extracellular bath solution in order to prevent oxidation of the drugs. The selective AKAP inhibitor InCELLect™ AKAP St-Hi31 and the corresponding control peptide were purchased from Promega (Germany).

**Cell culture**

After incubation for 5 minutes at room temperature with 2 ml Trypsin solution, immortalized kidney cells of the African green monkey (COS-7 cells) were collected with 10 ml cell culture medium and transferred into a 15 ml Falcon tube. Cell pellets obtained after centrifugation at 500 x g for 5 minutes at 4°C were resuspended in 10 ml of Dulbecco’s modified Eagle medium (DMEM; Gibco, Eggestain, Germany). Transfection was done according to the manufacturer’s instructions (Invitrogen, Germany). DNA-Lipofectamine complexes were added into the flask and cells were left in incubator for the next two days. 24 hours after transfection sodium-butyrate was added in order to enhance expression of transfected proteins. 48 hours after transfection, the cells were lysed and obtained lysates were either used immediately or stored at −20°C. For GFP constructs, transfection efficiency was analyzed under a fluorescence microscope (Zeiss axioplan microscope, Carl Zeiss, Jena, Germany).
Preparation and culturing of dissociated cell cultures from the dorsal thalamus and hippocampus

Dorsal thalami were prepared from embryos (Long-Evans rats) at stage E19 and subsequently transferred into ice-cold HBSS without Ca²⁺/Mg²⁺. After triple washing with HBSS (5 ml), 2.0 ml HBSS containing 0.5% trypsin was added to the tissue, followed by incubation at 37°C for 20 min. Thereafter tissue was washed again five times with 5 ml HBSS and finally transferred into tubes (2 ml) with HBSS containing 0.01% DNaseI. For dissociation thalamic tissue was pressed slowly three times through a 0.9 mm gauge needle followed by three passages through a 0.45 mm gauge needle. The remaining cell suspension was poured through a nylon tissue into a 50 ml tube and filled up with 18 ml DMEM. After estimating cell quantity, the suspension was diluted with DMEM in order to achieve a cell density of 30,000 cells/ml. A 500 µl aliquot of this suspension was placed on each well of a
24-well plate, containing defatted, baked, and poly-D-lysine-coated coverslips. The cell cultures were incubated at 37°C and 5% CO₂ up to the appropriate time points. Between 3rd and 4th day of incubation, AraC (final concentration: 6 μM) was added to prevent further growth of glial cells.

Primary hippocampal cells were prepared in the same way as described for thalamic neurons and grown in culture until day 10 to 14. Some of the cover slips were taken out and immediately fixed with 4% paraformaldehyde (PFA), stained with PKARIß antibody (1:500, BD Bioscience; see below) and used as control. For comparison cells were treated with βAR agonist isoproterenol hydrochloride (50 μM) for 5 minutes before fixation and PKARIß antibody staining. All cells were then incubated with the appropriate secondary antibody, mounted on microscope slides, and analysed under the microscope. Finally, distances (in pixels) between the dendritic localizations of PKA and the centre of the soma were quantified using MetaMorph software (Visitron Systems GmbH, Puchheim, Germany).

Immunocytochemistry

After 10–14 days in vitro (DIV 10–14), PFA-fixed thalamic cells were washed two times with 25 mM glycine in phosphate buffered saline (PBS, 10 mM), one time with PBS, and subsequently preincubated at room temperature in PBS-blocking solution containing 10% horse normal serum, 2% bovine serum albumin, 5% sucrose, and 0.3% Triton X-100. After 1 h, the following primary antibodies were added in different combinations to the blocking solution and incubated for 90 min at room temperature: rabbit anti-CaV1.2 (1:200, Alomone Labs, Israel); mouse anti-cAMP-dependent protein kinase type II beta regulatory subunit (PKARIß, 1:500, BD Bioscience, USA); rabbit anti-ß2-AR (H-73 1:400, Santa Cruz, USA); mouse anti-microtubule-associated protein 2 (MAP2, HM-2, 1:1000, Sigma, Germany); goat anti-AKAP-150 (N-19, Santa Cruz, USA); sheep anti-protein phosphatase 2A (PP2A, Acris, Germany). Cultures were then treated with PBS including 0.3% Triton X-100, incubated with appropriate specific fluorescent secondary antibodies: Alexa 488 donkey anti-sheep IgG; Alexa 488 goat anti-mouse IgG; Alexa 488 goat anti-rabbit IgG; Alexa 488 rabbit anti-goat IgG; Alexa 647 donkey anti-mouse IgG; Alexa 647 donkey anti-goat IgG; Alexa 647 donkey anti-rabbit IgG, (all from Molecular Probes, Invitrogen, Germany); Cy3-conjugated goat anti-rabbit IgG; Cy3-conjugated goat anti-mouse IgG; Cy3-conjugated rabbit anti-goat IgG, (all from Dianova, Germany) for 90 min, washed, and mounted with Mowiol. Omission of primary and secondary antibodies resulted in lack of fluorescent signals.

Immunohistochemistry

Long-Evans rats P25–27 were deeply anesthetized using pentobarbital (50 mg/kg body weight) and transcardially perfused with PBS, followed by an ice-cold 4% PFA/PBS for 35–40 min. Brains were removed, postfixed for 4 h in 4% PFA/PBS, and cryoprotected with 30% sucrose. Coronal sections (20 μm) were cut at the level of the dLGN, mounted onto polylysine slide glass (Menzel, Germany), and air dried. For detection of CaV1.2, fresh-frozen sections were used. In this case, brains from isoflurane-anesthetized rats were removed and frozen in −50°C isopentane. Cryostat coronal sections of 20 μm thickness were cut at the level of the dLGN, thaw-mounted onto Polylysine slide glass, air dried, and fixed in 4% PFA/PBS for 10 min. After permeabilization with 0.1% Triton X-100 in PBS for 10 min and several washes with PBS, sections were blocked with 10% normal horse serum (NHS), 2% BSA in PBS for 3 h to minimize nonspecific binding before incubation of slices with primary antibodies (rabbit anti-CaV1.2, l:200; mouse anti-PKARIß, 1:500) in 2% NHS, 2% BSA in PBS at 4°C for 16–18 h. After washing, sections were incubated with Cy3-conjugated donkey anti-rabbit IgG (1:400 in 2% NHS, 2% BSA in PBS, Dianova, Germany) or Alexa488 goat anti-mouse IgG (1:750 in 2% NHS, 2% BSA in PBS, Molecular probes, Invitrogen, Germany) for 1.5 h, washed again, and mounted with Immumount.

Culturing of COS-7 cells and pull down assays

Dulbecco’s modified Eagle medium (DMEM; Gibco, Eggstein, Germany) and washing buffer PBS were warmed to 37°C and frozen trypsin aliquots were thawed at room temperature. COS-7 cells (grown to confluence) were washed once with PBS. After incubation for 5 minutes at room temperature with trypsin (2 ml), cells were collected with 10 ml cell culture medium and transferred into a 15 ml Falcon tube. Cell pellets obtained after centrifugation at 500 x g for 5 minutes at 4°C were resuspended in 10 ml of Dulbecco’s modified Eagle medium (DMEM; Gibco, Eggstein, Germany). Transfection of the cells was done according to the manufacturer’s instruction (Invitrogen, Germany). Transfection was conducted for 2 days and after 24 hours sodium butyrate was added in order to enhance expression of transfected proteins. In the following cells were lysed and lysates were used immediately or stored at −20°C. For GFP constructs, transfection efficiency was analyzed under a fluorescence microscope (Zeiss Axioplan microscope, Carl Zeiss, Jena, Germany).

Transfected COS-7 cells expressing proteins of interest tagged to GFP or c-myc were scraped in medium and transferred to a 15 ml Falcon tube. Cell flask was washed with 2 ml of Trisbuffered saline (TBS) and the solution was pulled with scraped cells. Content of the cell was made accessible by repeating two times centrifugation (1000 x g, 5 min, 4°C), discarding the supernatant, resuspension of cells in 1 ml of TBS, and transfer to a 1.5 ml Eppendorf tube. In order to disrupt cells, the pellet was frozen and thawed using liquid nitrogen. After careful resuspension in 200 μl of TBS/Triton X-100 and vortexing, the lysate was rotated for 1 h at 4°C. A cleared lysate which was obtained after 20 min centrifugation (14000 rpm, at 4°C) was diluted with TBS containing 0.2% Triton X-100 and either used directly for pull down assay or stored at −20°C for further use.

Next, 50 μl matrix beads coupled to one of the proposed binding partners with GST or MBP tag were washed 2x and equilibrated in TBS containing 0.1% Triton X-100. Appropriate lysates from COS-7 cells (200–300 μl) were added and beads were gently shaken overnight at 4°C. On the next day matrices were centrifuged (5 min, 600 x g, 4°C) washed three times (TBS/0.2% Triton X-100, 10 min) to remove unbound proteins. Bound proteins were eluted by 5 min boiling with 4 x SDS sample buffer and either loaded on SDS-PAGE gels directly or stored at −20°C for further use.

Immunoprecipitation

AKAP5-GFP, co-expressed with PKARIß-c-myc in COS-7 cells, was immunoprecipitated using μMACSTM Epitope Tag Protein Isolation Kit (Miltenyi Biotec, Germany) according to manufacturer’s instructions. Eluted probes were loaded on SDS-PAGE and presence of proteins interacting with AKAP5 was detected after Western blot analysis using a PKARIß specific antibody (BD Bioscience, USA).

Western blotting procedures

For immuno-blot experiments solubilized protein fractions were separated on 5–20% SDS-polyacrylamide gradient gels and subsequently transferred to nitrocellulose membranes (90 min,
200 mA). The transfer buffer contained 25 mM Tris, 192 mM glycine, 0.02% SDS and 20% methanol. After blotting, membranes were blocked with 5% dry milk and 0.1% Tween 20 in TBS for 2 h and subsequently incubated at 4°C overnight with a specific dilution of antibodies in TBSA containing 0.1% Tween 20. After final washing steps, the blots were incubated with HRP-conjugated secondary antibodies (1:5000) for 2 h, washed again and finally developed using ECL films.

Microscopy

Immunofluorescence analysis of cultured neurons was done using a computer-controlled inverted laser scanning microscope (Leica, Bensheim, Germany) allowing recording of Z-stacks and enabling 3D deconvolution of the obtained images. Image analysis was done using MetaMorph, ImageJ (NIH), and Adobe Photoshop CS (version 9.0 CS2) software.

Data analysis

Statistical data analysis was done by Student’s t test or one way ANOVA as indicated using GraphPad Prism 5 and Microcal Origin 6. All values were presented as mean ± SEM. As we were able to demonstrate a Gaussian distribution for the three main parameters (current amplitude, Dinact and ratio of inactivation) analyzed in the present study under control conditions, statistical significance was evaluated by Student’s t-test. Where applicable, control values were compared with corresponding values obtained during drug application for the same cells. When intracellular substance application had to be used, an appropriate number of cells during drug application for the same cells. When intracellular substance application had to be used, an appropriate number of cells. Statistical data analysis was done by Student’s t test or one way ANOVA as indicated using GraphPad Prism 5 and Microcal Origin 6. All values were presented as mean ± SEM. As we were able to demonstrate a Gaussian distribution for the three main parameters (current amplitude, Dinact and ratio of inactivation) analyzed in the present study under control conditions, statistical significance was evaluated by Student’s t-test. Where applicable, control values were compared with corresponding values obtained during drug application for the same cells. When intracellular substance application had to be used, an appropriate number of cells during drug application for the same cells. When intracellular substance application had to be used, an appropriate number of cells.

Results

PCR expression patterns of the main components of the βAR signaling cascade in dLGN

In a first attempt to investigate the modulation of CDI via βARs, we analyzed the expression patterns of the main components of the proposed β-AR signaling pathway in dLGN TC neurons by performing RT-PCR analyses on a tissue and single cell level. Expression of RNA of the main neuronal L-type Ca2+ channels (CaV1.2/α1C, CaV1.3/α1D) were found in whole brain samples (Figure 1A), the VB (Figure 1B), and dLGN (Figure 1F). The skeletal muscle type CaV1.1/α1S and the retinaspecific CaV1.4/α1F were not detected in thalamic tissue (Figure 1), indicating a specific RT-PCR amplification. Furthermore, the specific expression of main components of the βAR signaling cascade supposed to be involved in CDI modulation in TC neurons was confirmed in dLGN tissue (Figure 1C–1E).

The expression of AKAP and the stimulatory G protein (Gs) was probed in a cell type-specific manner using single identified cells obtained from dLGN tissue. Acutely dissociated cells were observed under an inverted microscope and small bipolar interneurons and larger multipolar TC neurons were visually identified by using established criteria [22,23]. Single cells were collected by means of a suction pipette and used for standard PCR or quantitative real-time PCR (qRT-PCR). Because of very low amounts of the mRNA species targeted here, in some sets of experiments it was necessary to pool up to 10 cells. As Figure 2A shows, in three independent experiments (n = 3) the Gs subunit was expressed in both types of neurons, AKAP5 was only expressed in TC neurons, and AKAP7 was not detected at the single cell level. To obtain larger amounts of mRNA, we also performed qRT-PCR with samples from dLGN tissue. Using specific primers for AKAP5 and AKAP7, we were able to detect expression signals for these two genes in dLGN with almost identical expression levels (Figure 2B). Of note, that while AKAP7 is nearly equally expressed in the brain, AKAP5 exhibited a slightly higher expression level in the primary somatosensory cortex and hippocampus (data not shown). In view of positive signals for both AKAP subtypes from dLGN tissue and AKAP5 in single TC neurons, the absence of positive PCR bands for AKAP7 in extracts from single cells indicted that the amount of mRNA was below the detection limit of the method used here or pointed to an expression in other cell types present in intact tissue (glia cells, endothelial cells, blood cells).

CDI is active in TC neurons in brain slices

Next, we demonstrated the occurrence of CDI in TC neurons in brain slices. Total HVA Ca2+ current, which is composed of about 40% current through L-type channels, was measured in the presence of TTX (1 μM) [24]. We have shown before that blocking of L-type calcium channels using nifedipine (1 μM), (n = 8) indeed significantly reduced CDI [5]. HVA Ca2+ currents were recorded in 160 cells and a double pulse voltage protocol (Figure 3A) was used to effectively disclose CDI [3]. If CDI is operative, the current evoked by the test pulse should exhibit a U-shaped dependence on the conditioning pulse potential, with maximal inactivation occurring at the peak of the conditioning pulse current-voltage (I-V) relationship. Under standard conditions the I-V relationship of the conditioning pulse demonstrated HVA Ca2+ currents with an activation threshold of ~40 mV, maximal inward current at around +10 mV, and an apparent reversal potential at around +60 mV (data not shown). The test pulse I-V (peak amplitude of the test pulse current plotted vs. conditioning pulse voltage) revealed minimal current occurring at +10 mV (Figure 3C) as expected for a CDI mechanism. With respect to the amplitude of the test pulse current elicited from the holding potential of ~40 mV, the degree of inactivation (Dinact) was 39.5 ± 0.5% (n = 100; Figure 3C). By default, the test pulse I-V was obtained with the conditioning pulses altered in 10 mV increments from ~40 mV (Figure 3A). I-V relationships were obtained as described before and all experiments were done using a protocol resulting in maximal CDI [5]. Taken together, these findings demonstrate a CDI mechanism in TC neurons in the slice preparation that is very similar to that described in these cells after acute isolation [9].

The role of β-AR in CDI modulation of L-type Ca2+ channels

Since activation of the cAMP/PKA pathway is able to reduce CDI in isolated TC neurons [9,10], activation of βARs is expected to modulate CDI of L-type calcium channels in the dLGN. βARs consist of three similar receptor subtypes (β1, β2, β3). Using commercially available pharmacological substances, we specifically stimulated and blocked receptor subtypes and evaluated their possible relevance for CDI modulation. First, we used general agonists and antagonists for βARs, namely isoproterenol and propranolol (data not shown). Experiments testing βAR activation alone using 10 μM isoproterenol revealed an inhibiting influence of βAR stimulation on CDI. Dinact was decreased to 33.7 ± 2.2% (n = 4, p<0.05) as compared to control (Dinact = 40.1 ± 3%). The effect of isoproterenol on CDI was reversed by co-application of the βAR antagonist propranolol (100 μM, Dinact = 43.7 ± 2.1, n = 5, p<0.05; control Dinact = 46.2 ± 1.6%, n = 5; data not shown). These experiments demonstrate a general role of βAR in modulation of CDI in TC neurons. The rather moderate effect
of βAR on CDI may be limited by basal dephosphorylation processes in TC neurons [9] (see below).

Next, we tested agonists which bind preferentially to one of the three βAR subtypes. Therefore xamoterol (β2AR agonist), salmeterol (β2AR agonist), and BRL 37344 (β3AR agonist) were used. Challenging TC neurons with xamoterol (10 μM), reduced the degree of inactivation from 43.8 ± 2.3% to 37.4 ± 1.8% (n = 4, p < 0.05). This decrease was similar to that induced by salmeterol (10 μM) application (reduction of D\text{inact} from 39.0 ± 2.2% to 34.5 ± 0.3%, n = 4, p < 0.05; data not shown) and BRL 37344 (10 μM) application (reduction of D\text{inact} from 43.9 ± 0.6% to 38.3 ± 0.8%, n = 4, p < 0.05; data not shown). To minimize side effects of each drug on non-preferred βAR subtypes and to further enhance the specificity of our pharmacological approach, we used agonists that prefers one type of receptor (in 10 μM concentration) in combination with antagonists for the other two receptor types (in 100 μM concentration), thereby allowing to more selectively investigate the role of each type of receptor in CDI modulation. The following antagonists were applied: CGP 20712 (β2AR antagonist), ICI 118,351 (β3AR antagonist), and SR 59230A (β3AR antagonist). Using this approach, effects on CDI were detected only when the β2AR agonist salmeterol was used in combination with CGP 20712 and SR 59230A (reduction of D\text{inact} from 33.5 ± 1.2% to 28.4 ± 1%, n = 4; Figure 4A). For the other two βAR subtypes, no significant modulation of CDI was obtained in this set of experiments (data not shown). These results clearly demonstrate that β2AR specifically contribute to CDI modulation in dLGN TC neurons.

Localization of β2ARs in cultured TC neurons

Next, we analyzed the specific expression and localization of β2AR in 10 days old cultured thalamic neurons and performed double immunostaining using an antibody against β2AR (Figure 4B) in combination with an antibody against the dendritic marker protein microtubule associated protein 2 (Map2; Figure 4B). The expression of both proteins was detected with β2AR mainly localized in somatic and proximal dendrites of TC neurons (Figure 4B, merge image), cellular compartments that are involved in modulation of CDI in TC neurons.

Blocking of PKA suppresses βAR-dependent modulation of CDI in TC neurons

Next, we assessed the possible contribution of PKA in CDI modulation of TC neurons by intracellular application of a PKA inhibitor (PKI, 10 μM) in whole cell patch clamp experiments. As shown in Figure 5A, the degree of inactivation was significantly reduced from 44.6 ± 2.3% (n = 5) under control conditions to 33.1 ± 2.1% (n = 4, p < 0.001; Figure 5A) when TC neurons were challenged with salmeterol (10 μM), while in combination with PKI the inhibitory effect of βAR stimulation was absent (D\text{inact} = 43.2 ± 1%; n = 5). These findings indicate that the modulation of CDI in TC neurons depends on the activity of PKA.

To address the possible association between PKA and CaV1.2, we performed double immunostainings of cultured TC neurons treated with the general βAR agonist isoproterenol. In this set of experiments we used an antibody directed against the regulatory subunit IIβ of PKA (PKARIIβ), which is highly expressed in thalamus and hippocampus (Figure 5C). Image analysis pointed to a close spatial proximity of the two proteins at the somatodendritic junction following βAR stimulation (Figure 5B). On the other hand untreated cells showed more somatic expression of the PKARIIβ (data not shown).

AKAPs play a crucial role in the modulation of CDI in TC neurons

Next, we investigated the possible contribution of AKAP in CDI modulation in TC neurons. AKAP is assumed to simultaneously bind to PKA, Ca2⁺ channels, and protein phosphatases, including calcineurin [17]. Therefore a selective AKAP inhibitory peptide which binds to the PKA binding sites of AKAP thereby blocking the binding of intracellular PKA and a corresponding control peptide which allows normal PKA binding were used. When salmeterol (10 μM) was applied in the presence of intracellular AKAP inhibitory peptide (10 μM), the degree of inactivation (37.6 ± 1.5%, n = 9; Figure 6A and 6B) was comparable to control conditions (39.5 ± 0.5, n = 4, p > 0.05) but different from conditions where salmeterol was applied alone (34.5 ± 0.3%, n = 4, p < 0.01; Figure 6A and 6B). Using the same experimental protocol, application of salmeterol in the presence of intracellular control peptide did not change the reduction in CDI (data not shown). In summary, these experiments point to a role of AKAP in the modulation of CDI in TC neurons.

Complex formation by the main components of the β-AR signaling cascade in TC neurons

After stimulation of the β-AR signaling cascade, it is assumed that a ternary complex is formed between PKA, AKAP, and CaV1.2, the formation of which is important for CDI modulation in TC neurons. In order to find evidence corroborating this assumption, we next performed immunocytochemical staining using antibodies specific for CaV1.2 (Figure 7A, upper left panel), AKAP 5 (Figure 7A, middle left panel), and PKARIIβ (Figure 7A, lower left panel), in 10 days old cultured thalamic neurons. Fluorescence imaging revealed that all three proteins are densely expressed in somatic regions and proximal dendrites of TC neurons. Merged images pointed to rather close spatial association of the three proteins (Figure 7A, right panel). In contrary to CaV1.2, another calcium channel CaV2.1 is similarly expressed at somatic regions and proximal dendrites but also additionally in distal dendrites (Figure S2).

Protein-protein interactions between components of the ternary inactivation complex

Since immunocytochemical stainings pointed to the possibility that PKA, AKAP, and CaV1.2 are located close to each other, thereby contributing to the CDI process, we performed pull down assays which confirmed an interaction between PKARIIβ and AKAP7 (Figure 7B) as well as PKARIIβ and PKAcsβ (Figure 7D). Control incubations of samples of interest with the appropriate fusion-protein partner (empty vector) showed no signal in Western blots. Moreover, indications for an interaction of PKARIIβ with another member of the AKAP family, namely AKAP5, were obtained by co-immunoprecipitation experiments. Therefore, magnetic beads coupled to GFP antibody and Western blotting using a PKARIIβ-specific antibody were used to verify the binding of PKARIIβ to GFP-tagged AKAP5 (Figure 7C). Control incubations of samples of interest (c- myc PKARIIβ) with the appropriate fusion-protein partner (empty GFP vector) showed no signal in Western blots. These results provided further evidence for a close coupling between PKA and AKAP in the thalamus.

Phosphorylation and dephosphorylation processes in modulation of CDI

It is well documented that phosphorylation/dephosphorylation processes play an important role in the regulation of calcium channel activity [3,9]. In an additional series of patch clamp experiments we focused on the role of dephosphorylation in the
modulation of CDI in TC neurons after βAR stimulation. The double-pulse protocol was applied using the phosphatase inhibitor okadaic acid (OA; 10 μM alone; p > 0.05; Figure 8A), or in combination with isoproterenol (10 μM; Figure 8B). After blocking dephosphorylation processes, the degree of inactivation is reduced to 29.3 ± 1.1% (n = 5) in comparison to βAR stimulation alone (35.7 ± 2%, n = 5; p < 0.01), and non-stimulated control cells which showed D_{inact} of 40 ± 3.3% (n = 5; p < 0.001) in these experiments.

Next, the expression of protein phosphatases PP2A was investigated in 10 days old cultured thalamic neurons by using antibodies for PP2A (Figure 8C, upper right panel) and Cav1.2

Figure 4. β2AR specifically modulates CDI in TC neurons of dLGN. (A) Representative current traces recorded under control conditions (upper left panel) and during extracellular application of specific β2AR agonist salmeterol (10 μM) plus antagonists for the other two types of receptor (CGP 20712 dihydrochloride-β1 antagonist and SR 59230A hydrochloride-β3-antagonist, 100 μM each; lower left panel) elicited by the indicated pulse protocol (middle left panel). The bar graph (right panel) represents D_{inact} under different recording conditions (as indicated). The mean value of four cells recorded under control conditions was taken for comparison with four cells recorded under 10 μM salmeterol plus 100 μM β1+β3 antagonists. Data are presented as means ± SEM of several independent experiments. **p < 0.01. Significance of salmeterol plus antagonists versus control was calculated by Student’s t test. The degree of inactivation is given by the normalized current amplitude of the mean postpulse I/V at +10 mV. (B) Immunocytochemical analysis of primary cultures of the dorsal thalamus using MAP2 specific antibody (left upper panel, red) and β2AR specific antibody (left down panel, green). The merged picture revealed the co-expression of these two proteins in somatic regions and proximal dendrites. Enlarged inlay represents magnification of the area indicated by the rectangle. Yellow dots represent places were these two proteins are in close proximity. Data shown are representative pictures from several independent immunostainings and preparations of neurons. In all cases, omission of primary antibodies resulted in no fluorescence signal above background (negative control).

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Both proteins were mainly localized in somatic regions and proximal dendrites of TC neurons (Figure 8C, lower panel). Merged pictures pointed to a close spatial arrangement.

**Discussion**

The present study provides clear evidence that the CDI of HVA Ca\(^{2+}\) channels is decreased by \(\beta\)AR signaling in central neurons. Moreover, we present findings that are in agreement with the existence of a possible protein complex including PKA, AKAP, and CaV1.2 which underlies the modulation of CDI. The most important findings of the present study are: (i) Classical double pulse protocols reveal the occurrence of CDI in TC neurons in a slice preparation; (ii) Activation of \(\beta\)2AR induces the down-regulation of CDI; (iii) Blocking of PKA signaling completely suppresses the effect of \(\beta\)AR stimulation on CDI; (iv) AKAP might play a crucial role in CDI modulation and the docking of PKA to sites close to CaV1.2 [see Text S1]. Moreover, blocking of the interaction between AKAP and PKA significantly reduces CDI and translocation of PKA. (v) Phosphorylation and dephosphorylation processes represent the basis for bidirectional up- and down-regulation of CDI in TC neurons, respectively.

**\(\beta\)AR stimulation and modulation of CDI of CaV1.2 via phosphorylation and dephosphorylation processes**

We have previously demonstrated the existence of CDI of HVA Ca\(^{2+}\) currents in TC neurons in different (acutelyisolated cells, brain slices) thalamic preparations [5,8,9,10]. Based on these findings, the basic features of CDI in TC neurons can be defined as follows: (i) Under control conditions the degree of inactivation varies between about 35–40%. (ii) In addition to L-type Ca\(^{2+}\) channels, Q-type channels are also governed by CDI. (iii) CDI is influenced by a number of cellular mechanisms including repetitive neuronal activity, phosphorylation and dephosphorylation, Ca\(^{2+}\)-binding proteins, the cytoskeleton, and intracellular Ca\(^{2+}\) release. The present study adds to these findings by demonstrating the specific influence of \(\beta\)2AR stimulation via PKA and scaffolding proteins of the AKAP family on the degree of CDI.
In the present study, we demonstrated that stimulation of βAR, which leads to phosphorylation of CaV_{1.2} channels in a cAMP-dependent manner via PKA activation, significantly reduced the degree of inactivation of L-type CaV_{1.2} channels. The same results were obtained when channel dephosphorylation was inhibited. This indicates that phosphorylation keeps L-type CaV_{1.2} channels in a state of high open probability, ready to open during depolarization [9,25]. The modulation of L-type CaV_{1.2} channels through phosphorylation via different second messenger systems is well established, and includes phosphorylation by PKA and CaM kinase II (for review, see [26,27]). Both types of modulation result in an increase in peak current amplitude [28,29]. Furthermore, CDI has been shown to be reduced by activation of cAMP-dependent phosphorylation [9,25,30]. Therefore, both the increment in HVA CaV_{2+} current amplitude and the reduction of CDI after βAR stimulation observed in the present study are consistent with a phosphorylation of L-type CaV_{1.2} channels by PKA.

The serine residue at position 1928 (Ser1928) of CaV_{1.2} channels is one important target of PKA activity in heart and brain [31,32,33]. However, it has been recently shown that mutation of Ser1928 of cardiac L-type CaV_{1.2} channels has only a small effect on channel phosphorylation after βAR stimulation [34], pointing to the existence of additional phosphorylation sites [34,35,36]. Moreover, meaningful regulation of channel activity by PKA phosphorylation requires a proper balance with dephosphorylation processes. Several studies in hippocampal neurons have shown that protein phosphatases including PP1, PP2A, and calcineurin directly bind to the C-terminal region of CaV_{1.2} [17,31,33]. Moreover, the signaling pathway from β2AR to the CaV_{1.2} channel, including G-proteins, adenylyl cyclase, PKA and the counterbalancing protein phosphatases, forms a closely associated protein complex in the forebrain [13]. However, the role of this complex in context of CDI modulation has not been addressed yet. CaV_{1.2} channels are expressed in TC neurons [8] and the existence of βARs, coupled positively to adenylyl cyclase in these neurons, has been shown [37]. Moreover, we previously demonstrated effects of calyculin A and ascomycin, which are blockers of PP1, PP2A, and calcineurin [38] on CDI and CaV_{1.2} current amplitude in TC neurons [9]. The present study on brain slices confirmed and extended previous findings obtained in acutely isolated TC neurons by identifying βAR as the receptor subtype involved in CDI modulation and showing that blocking protein phosphatases by okadaic acid, has a significant effect on CDI during β-AR stimulation. From our data, it is therefore reasonable to conclude that PKA and protein phosphatases antagonistically modulate CDI of CaV_{1.2} channels in TC neurons via phosphorylation and dephosphorylation processes.

Although our experiments with okadaic acid demonstrated the role of dephosphorylation processes in modulation of CDI after βAR stimulation in TC neurons, the specific type of phosphatase involved in TC neurons is not clear. The original model of CDI in Helix aspersa included a dephosphorylation cycle by calcineurin as the fundamental step leading to channel closure [39]. Later, evidence for and against an involvement of calcineurin was found [17,40,41,42,43,44,45]. In TC neurons, application of the calcineurin blocker ascomycin boosts CDI [9,46]. Although these observations clearly indicate a modulation of CDI by calcineurin in TC neurons, the inactivation process itself is probably not a dephosphorylation reaction. As shown above, PKA is the main enzyme which phosphorylates CaV_{1.2} channels, thereby keeping them in an open state (increasing their open probability) and restraining the effects of CDI. Besides this phosphorylation processes that occur after βAR stimulation, there might be a constant dephosphorylation driven by PP1 and PP2.

AKAP mediates the modulation of CaV_{1.2} channel during βAR stimulation

CaV_{1.2} channels can physically associate with either AKAP5 or AKAP7 through a leucine zipper interaction [17,47]. Both, the modulation of channels and its downstream signaling depend upon the identity of the associated AKAPs. Although both AKAPs...
subtypes target PKA to the channel, AKAP5 also targets calcineurin and thereby confers unique characteristics upon AKAP5-complexed L-type channels in neurons [17]. AKAP5 is the major AKAP protein in neurons, where it is widely distributed and has been shown to anchor protein kinases and other signaling proteins to multiple receptors and ion channels [48]. On the one side, AKAP5 recruits PKA and calcineurin to the AMPA receptor [49], associates with CaV1.2 [14,50] in neurons, recruits PKA and calcineurin to the channel and is necessary for the β2AR stimulation of L-type calcium currents [13,14,48] as well as for the L-type calcium current-mediated activation of the transcriptional regulator NFATc4 [17]. On the otherside, AKAP5 binds also to β2AR and facilitates receptor phosphorylation and signaling [51]. Moreover, colocalization with CaV1.2 and postsynaptic density (PSD) proteins in dendritic spines of hippocampal neurons has been shown [52]. Three different binding sites for AKAP5 were described in the N terminus, the cytoplasmic loop connecting repeats I and II, and in the C terminus of CaV1.2 [14]. The C-terminal leucine zipper was shown to be essential for AKAP binding and for βAR stimulation and reversible phosphorylation of CaV1.2 in heart muscle and in neurons [17,47]. Mutation of the three basic residues of this motif blocked AKAP5 and PKA binding and phosphorylation of the Ca2+ channel in response to βAR stimulation [17]. Moreover, mutation of known binding sites

![Figure 7. Interaction partners important for CDI modulation in TC neurons.](https://www.plosone.org/figure/image/10.1371/journal.pone.0027474.g007)

**Figure 7. Interaction partners important for CDI modulation in TC neurons.** (A) Immunocytochemical analysis of primary cultures of the dorsal thalamus using CaV1.2- (red), AKAP150- (green) and PKARIIβ- (blue) specific antibodies. Merged picture shows the close connection of the components of the proposed ternary complex, especially in somatic regions and proximal dendrites. Enlarged inlay represents a magnification of the area indicated by the rectangle. Data shown are representative pictures from several independent immunostainings and preparations of neurons. In all cases, omission of primary antibodies resulted without signal (negative control). Western blot analysis and pull down assays were done as described in "Methods". (B) Interaction of AKAP7-MBP and PKARIIβ-c-myc was detected using antibodies against c-myc. (C) IP of PKARIIβ-c-myc and AKAP5-GFP detected after incubation with GFP-coupled magnetic beads using antibodies derived against PKARIIβ. (D) Existence of PKA holoenzyme consisting of PKARIIβ-GST and PKAcsβ-GFP was detected with antibodies against GFP protein.

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![Figure 8. The role of dephosphorylation processes in CDI modulation of TC neurons.](https://www.plosone.org/figure/image/10.1371/journal.pone.0027474.g008)

**Figure 8. The role of dephosphorylation processes in CDI modulation of TC neurons.** (A) Representative current traces recorded under control conditions (upper left panel) and in the presence of okadaic acid (OA) (10 μM; left down panel). The bar graph (right panel) represents D_inact under different recording conditions (as indicated). The mean value of three cells recorded under control conditions was taken for comparison with three cells recorded under 10 μM OA. (B) Representative current traces recorded under control conditions (upper left panel) and in the presence of the isoproterenol alone (10 μM; left middle panel) or in combination with OA (10 μM; lower left panel). The bar graph (right panel) represents D_inact under different recording conditions (as indicated). The mean value of five cells recorded under control conditions was taken for comparison with five cells recorded under 10 μM isoproterenol and five cells recorded under isoproterenol plus OA. Data are presented as means ± SEM of several independent experiments. ***P<0.001. Significance of isoproterenol plus OA (n=5) versus controls (n=5) was calculated by Student’s t test. **P<0.01. Significance of isoproterenol (n=5) versus isoproterenol plus OA (n=5) was calculated by Student’s t test. The degree of inactivation is given by the normalized current amplitude of the mean postpulse IV at +10 mV. (C) Immunocytochemical analysis of primary cultures of the dorsal thalamus using CaV1.2- (left panel, green) and PP2A-specific antibodies (right panel, red). Merge picture showed close association of the two proteins, especially in somatic regions and proximal dendrites. Data shown are representative pictures from several independent immunostainings and thalamic neurons preparations. In all cases, omission of primary antibodies resulted in no fluorescence signal above background (negative control).

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on Ca$_{v}$1.2 for the scaffold proteins, like AKAP and PSD proteins did not change membrane expression of Ca$_{v}$1.2 [32]. The above findings indicate that binding of these proteins is necessary for the regulation of Ca$_{v}$1.2 after βAR stimulation but does not have an influence on the membrane localization of Ca$_{v}$1.2.

Based on the previous findings discussed above and the results presented here, we propose the following role for AKAP in the regulation of CDI in TC neurons. Following βAR stimulation, AC gets activated via G-proteins and produces cAMP which then activates PKA. With support of AKAPs, PKA targets its final effector, the CaV1.2 channels and phosphorylates them (see Figure S1). In the present study we demonstrated that blocking the binding of AKAP to PKA significantly reduces CDI. Moreover, under similar conditions we have shown that PKA phosphorylation of Ca$_{v}$1.2 and translocation of this enzyme close to the channel depends on binding of PKA and AKAPs in hippocampal neurons (see Text S1 & Figure S3). Most recent studies demonstrated that AKAP5 is required to localize both RIIs and RIIb containing holoenzymes to the dendritic regions of neurons in the hippocampus and striatum. PKA is dramatically delocalized within dendrites in both the KO and D36 (mutant that lacks the PKA binding domain of AKAP5) mice indicating that no other AKAP subtype is able to compensate and maintain normal PKA localization [33]. Our study confirmed an interaction between AKAP and PKA in pull down assays and demonstrated that after stimulation of the βAR signaling cascade a ternary complex is formed between PKA, AKAP, and Ca$_{v}$1.2 and that the formation of this complex is important for CDI modulation in TC neurons. Moreover, as mentioned before, AKAP5 also targets calcineurin which is able to activate protein phosphatase PP1 [54] and therefore might have multiple function in regulation of CDI by influencing both phosphorylation by PKA and dephosphorylation processes.

Functional significance of Ca$^{2+}$ channel phosphorylation after β-AR stimulation

Release of transmitters from a number of brainstem terminals modulates the behavioural states of an individual by depolarizing TC relay neurons [37]. During states of slow-wave sleep, thalamic relay neurons are hyperpolarized and display rhythmic burst activity [11]. During states of wakefulness, these cells are depolarized and display tonic single spike activity, resulting in the faithful transmission of sensory signals through the dorsal thalamus. The shift from burst activity to tonic activity is mediated by increased activity of ascending brainstem fibres that are thought to increasingly release acetylcholine (ACh), noradrenaline (NA) and serotonin (5-HT) during wakefulness. Both NA via β-receptors and 5-HT via an unknown 5-HT receptor subtype, activate adenylate cyclase [37] in TC relay neurons and are thus able to positively modulate HVA Ca$^{2+}$ currents.

When attempting to integrate the findings of the present study into the known framework of thalamic physiology it can be assumed that HVA Ca$^{2+}$ currents are especially activated during tonic firing. Furthermore, following release of NA, HVA Ca$^{2+}$ current amplitudes will be increased while CDI is decreased. Another consequence of βAR stimulation may be the AC/cAMP-dependent inhibition of high conductance Ca$^{2+}$-dependent K$^+$ (BK$_{Ca}$) channels in sensory TC neurons [55]. In addition, tonic sequences of action potentials are coupled to CICR from intracellular stores, thereby further increasing Ca$^{2+}$ entry into TC neurons during wakefulness [6]. It has been shown that intracellular Ca$^{2+}$ release provides Ca$^{2+}$ that contributes to CDI and activates BK$_{Ca}$ channels in TC neurons [3,6]. Computer modelling indicated that activation of BK$_{Ca}$ channels leads to the occurrence of spike frequency adaptation (P. Meuth & T. Budde, unpublished observations), a condition that would impair the faithful 1:1 relay of incoming trains of sensory action potentials. These data indicate a fine-tuned interplay between activity dependent Ca$^{2+}$ influx, phosphorylation/dephosphorylation processes and the mode of activity, possibly to enable faithful signal integration and transfer during wakefulness.

Future studies will have to unravel the different modulatory pathways that act upstream of the multiple CDI mechanisms thereby pointing to additional functions of CDI and unraveling further the elusive role of HVA Ca$^{2+}$ channels in thalamic physiology.

**Supporting Information**

**Text S1** AKAP5 assist in PKA translocation from somatic regions to the plasma membrane. Translocation of PKAIIβ after βAR stimulation with isoproterenol in hippocampal neurons. (DOC)

**Figure S1** Main components of the βAR signaling cascade in dLGN. (TIF)

**Figure S2** Differential expression of Ca$_{v}$2.1 from Ca$_{v}$1.2 in TC neurons of thalamus. Immunocytochemical analysis of primary cultures of the dorsal thalamus using Ca$_{v}$2.1-(green) and MAP2-specific (neuronal marker, red) antibodies. Merge image showed expression of these two proteins in somatic, proximal and in distal regions of TC neurons. Data shown are representative pictures from several independent experiments. In all cases, omission of primary antibodies resulted in staining without signal (negative control). (TIF)

**Figure S3** PKA translocation from somatic regions to the plasma membrane. (A) Cultured hippocampal cells were treated with 50 μM isoproterenol or kept under control conditions (control/no treatment), fixed and immunostained with PKAIIβ antibody. Enlarged inlays represent magnification of the area indicated by the rectangle. Note translocation of PKA from somatic perinuclear region in control cells to more distal regions close to the plasma membrane and probably Ca$^{2+}$ channels in cells treated with isoproterenol. (B) Ten days old hippocampal cultures were treated for 5 minutes with 50 μM isoproterenol in combination with 50 μM control peptide AKAP St-Ht31 (left panel), 50 μM isoproterenol in combination with 50 μM AKAP St-Ht31 inhibitory peptide (right panel) and 50 μM isoproterenol in combination with 7.5 μM Anisomycin (down panel). The cells were then fixed and immunostained with PKAIIβ antibody. Note that PKA is still translocated to proximal dendrites, after blocking the association of PKA with AKAPs, translocation is almost completely inhibited and anisomycin treatment did not block PKA translocation. (C) Quantification of translocation experiments using MetaMorph was done by measurement of PKA distance from centre of the soma to dendrites in pixels after different treatments. Data are presented as means ± SEM of several independent experiments. ***p<0.001, Anova test. (TIF)

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Author Contributions

Conceived and designed the experiments: VR TM PL. Performed the experiments: VR TK PE. Analyzed the data: VR TM PL. Contributed reagents/materials/analysis tools: PL MK SM. Wrote the paper: VR TB TM. Edited the manuscript: VR TM TB SM MK.

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