Cross Talk between Metabotropic and Ionotropic Glutamate Receptor-Mediated Signaling in Parallel Fiber-Induced Inositol 1,4,5-Trisphosphate Production in Cerebellar Purkinje Cells

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In many excitatory glutamatergic synapses, both ionotropic glutamate receptors (iGluRs) and metabotropic glutamate receptors (mGluRs) are closely distributed on the postsynaptic membrane. However, the functional significance of the close distribution of the two types of glutamate receptors has not been fully clarified. In this study, we examined the functional interaction between iGluR and mGluR at parallel fiber (PF)→Purkinje cell synapses in the generation of inositol 1,4,5-trisphosphate (IP$_3$), a key second messenger that regulates many important cellular functions. We visualized local IP$_3$ dynamics in Purkinje cells using the green fluorescent protein-tagged pleckstrin homology domain (GFP-PHD) as a fluorescent IP$_3$ probe. Purkinje cells were transduced with Sindbis virus encoding GFP-PHD and imaged with a two-photon laser scanning microscope. Translocation of GFP-PHD from the plasma membrane to the cytoplasm attributable to an increase in IP$_3$ concentration was observed on PF stimulation in fine dendrites of Purkinje cells. Surprisingly, this PF-induced IP$_3$ production was blocked not only by the group I mGluR antagonist but also by the AMPA receptor (AMPAR) activation and Ca$^{2+}$ buffering by BAPTA. These results show that IP$_3$ production is mediated cooperatively by group I mGluR and AMPAR through G-protein coupling to phospholipase C (PLC) via the heterotrimeric G-protein. We have shown that intracellular IP$_3$ dynamics can be visualized within intact cells using the green fluorescent protein-tagged pleckstrin homology domain of PLC-δ1 (GFP-PHD) (Hirose et al., 1999). GFP-PHD binds to PIP$_2$ within the plasma membrane and translocates to the cytoplasm on an increase in the concentration of IP$_3$ because of its 20-fold higher affinity for IP$_3$ than for PIP$_2$. Therefore, by imaging the translocation of GFP-PHD, we can measure the change in intracellular IP$_3$ concentration. We have also shown that GFP-PHD can be used to measure intracellular IP$_3$ dynamics in single Purkinje cells (Okubo et al., 2001). In this report, the IP$_3$ imaging method was combined with a high-
spatiotemporal resolution imaging technique using a two-photon laser scanning microscope (TPLSM) (Denk and Svoboda, 1997) to visualize the local IP₃ dynamics in response to PF inputs in Purkinje cells within cerebellar slices. This imaging technique now allows us to visualize the local IP₃ dynamics on PF inputs in the fine dendrites of Purkinje cells. Unexpectedly, IP₃ production on PF stimulation was found to be dependent on the simultaneous activation of group 1 mGlur and AMPAR. In addition, this PF-induced IP₃ production was blocked by the inhibition of the G-protein function and by buffering of intracellular Ca²⁺ concentration. These results show that the PF-induced IP₃ production is mediated cooperatively by group 1 mGlur and AMPAR through G-protein activation and Ca²⁺ influx in Purkinje cells.

Materials and Methods

Sindbis viruses. The enhanced GFP (EGFP) gene in pIRE2-EGFP (Clontech, Palo Alto, CA) was replaced by the cDNA encoding internal ribosomal entry site (IRES)-GFP-PHD. The cDNA encoding IRES-GFP-PHD was cloned into pSinRep5 (Invitrogen, Carlsbad, CA). This vector was then used as the template for in vitro transcription using SP6 RNA polymerase (Invitrogen or Ambion, Austin, TX). The RNA transcript and the helper RNA from a D/H(26S) cDNA template (Invitrogen) were co-transfected into baby hamster kidney cells by electroporation. Twenty-four hours after transfection, the culture medium was harvested because this contained the infectious particles of Sindbis virus (SIN)-IRES-GFP-PHD. After transfection, the culture medium was harvested because this contained the infectious particles of Sindbis virus (SIN)-IRES-GFP-PHD. IRES-dependent translations were shown to be less efficient than cap-tailed the infectious particles of Sindbis virus (SIN)-IRES-GFP-PHD. After transfection, the culture medium was harvested because this contained the infectious particles of Sindbis virus (SIN)-IRES-GFP-PHD. The cDNA encoding IRES-GFP-PHD was cloned into pSinRep5 with IRES-entry site (IRES)-GFP-PHD. The cDNA encoding IRES-GFP-PHD was then inserted into the cerebellum so that the tips of the pipettes were exposed by removing the cranium and the dura. The tips of glass pipettes were backfilled with virus in the medium (HEK cell, Palo Alto, CA) was replaced by the cDNA encoding GFP-PHD (Hirose et al., 1999) to generate the cDNA encoding 5-phosphatase [wild type (WT)]-IRES-GFP-PHD. Arginine 343 of IP₃ 5-phosphatase was mutated to alanine by the PCR technique using a mutated primer. The cDNA fragment including the mutated site was used to replace the corresponding site in 5-phosphatase (WT) to generate the cDNA of 5-phosphatase (R343A)-IRES-GFP-PHD cloned into pSinRep5. The infectious particles were acquired by the same procedure described above.

Infection of Purkinje cells and preparation of cerebellar slices. C57BL/6 mice [postnatal day 18 (P18)–P29] were deeply anesthetized with pentobarbital, and the surface of cerebellar lobule 6 beside the midline was exposed by removing the cranium and the dura. The tips of glass pipettes were backfilled with virus in the medium (0.5 μl). Glass pipettes were then inserted into the cerebellum so that the tips of the pipettes were placed at the molecular layer of cerebellar lobule 6 facing lobule 4/5. The virus-containing medium was delivered over 5 min by air pressure applied through the microsyringe. The glass pipettes were then left in place for 5 min to prevent virus leakage. Twenty to twenty-six hours after the infection, parasagittal cerebellar slices (250 μm thickness) were described as prepared previously (Edwards et al., 1989; Alba et al., 1994; Kakizawa et al., 2000). Slices were incubated in a holding chamber containing artificial CSF (ACSF) bubbled with 95% O₂ and 5% CO₂ at 35°C for 1 hr and then returned to 23°C before recording. ACSF for slicing and incubation contained (in mM): 125 NaCl, 2.5 KCl, 2 CaCl₂, 1 MgSO₄, 1.25 NaH₂PO₄, 26 NaHCO₃, and 20 glucose.

Electrophysiology. For recording, slices were transfected to a recording chamber that was continuously perfused with ACSF bubbled with 95% O₂ and 5% CO₂ and containing 10 μM bicuculline to block spontaneous IPSCs. Whole-cell recordings were made from Purkinje cells using an upright microscope (BX61WI; Olympus Optical, Tokyo, Japan). The resistances of patch pipettes were 2–5 MΩ when filled with an intracellular solution containing (in mM): 148 K gluconate, 2 NaCl, 4 MgCl₂, 4 ATP (Na salt), 0.4 GTP (Na salt), and 10 HEPES, pH 7.3 (adjusted with KOH). An intracellular solution also contained 0.2 mM X-rhod-1 (Molecular Probes, Eugene, OR) or 0.2 mM Calcium Green-5N (Molecular Probes). EPSCs were recorded with an EPC-9 patch-clamp amplifier (HEKA Elektronik, Lambrecht/Pfalz, Germany) using the voltage-clamp mode. The signals were filtered at 2 kHz and digitized at 10 kHz. The on-line data acquisition and off-line analysis of data were performed using PULSE software (HEKA). A stimulation pipette (3–7 μm tip diameter) was filled with ACSF and placed near the site where the PF-induced response was to be imaged (within 15 μm in the lateral direction and 10–50 μm above in the vertical direction) to apply square pulses for the focal stimulation of PFs (duration, 0.1 msec; amplitude, 4–36 V). Low-amplitude Ca²⁺ transients attributable to direct stimulation of Purkinje cells were observed when high-frequency stimulus pulses (10 at 50 Hz) with large amplitudes (25–36 V) were applied via the stimulation electrode placed very close to the dendrites (10–15 μm above, 12 runs from 12 cells). Such a stimulation configuration was avoided in our experiments. Experiments were performed at room temperature (23–25°C).

Imaging. Imaging was performed with a custom-made TPLSM (Nemoto et al., 2001; Takahashi et al., 2002) consisting of an upright microscope (BX61WI; Olympus Optical) equipped with a scanning unit (FV300; Olympus Optical) and a Ti:sapphire laser providing a mode-locked femtosecond-pulse laser (Tsunami; Spectra-Physics, Mountain View, CA). Purkinje cells that expressed a sufficient amount of GFP-PHD were selected based on morphological criteria (large, round soma and prominent dendritic arbor) and then viewed under a water immersion objective (40×; numerical aperture, 0.80). The distal dendrite that showed a clear membranous distribution of GFP-PHD was selected for imaging. For the imaging of Ca²⁺ transients, X-rhod-1 or Calcium Green-5N was applied to Purkinje cells through the patch pipette. Imaging was started at least 20 min after the establishment of the whole-cell configuration to allow the diffusion of X-rhod-1 or Calcium Green-5N. Both GFP-PHD and X-rhod-1 were excited at 900 nm (pulse width, 70–100 fsec). Calcium Green-5N was excited at 810 nm (pulse width, 70–100 fsec). Emitted fluorescence was filtered using an infrared ray cutting filter and then separated by a 565 nm dichroic mirror (DM565; Olympus Optical). A 495–540 nm barrier filter (BA495–540; Olympus Optical) was placed in the shorter-wavelength pathway to isolate GFP or Calcium Green-5N fluorescence and, a 590 nm long-pass filter (BA590; Olympus Optical) was placed in the longer-wavelength pathway to isolate X-rhod-1 fluorescence. External photomultiplier tubes (Hamamatsu Photonics, Hamamatsu, Japan) were used to detect fluorescence. Data were acquired using FluoView 4.2 software (Olympus Optical). Stimulus-induced changes in fluorescence were recorded using the line scan mode with a 5 msec time resolution. Fluorescence signals were corrected for background fluorescence by measuring a nonfluorescent area.

Drugs. (S)-4- Carboxyphenylglycine (4-CPG), 6-nitro-7-sulfamoylbenzo[f]quinoline-2,3-dione (NBQX; disodium salt), and bicuculline were obtained from Tocris Cookson (Ballwin, MO); GDP-βS was from Sigma (St. Louis, MO); and BAPTA was from Dojindo (Kumamoto, Japan). 4-CPG, NBQX, and bicuculline were dissolved in ACSF and administered to the cells through the perfusion system of the recording chamber. GDP-βS and BAPTA were dissolved in the intracellular solution and introduced into the cytoplasm through the patch pipette. For the intracellular delivery of BAPTA, the intracellular solution was used, which contained (in mM): 114 K glutonate, 2 NaCl, 4 MgCl₂, 6 CaCl₂, 4 ATP (Na salt), 0.4 GTP (Na salt), 10 HEPES, 20 BAPTA, and 0.2 X-rhod-1, pH 7.3 (adjusted with KOH).

Results

Sindbis-mediated expression of GFP-PHD in Purkinje cells in cerebellar slices

We expressed GFP-PHD in Purkinje cells using Sindbis virus carrying a gene encoding GFP-PHD (SIN-IRE5-GFP-PHD). Twenty to twenty-six hours after the injection of infectious particles of the Sindbis virus into the cerebellum of P18–P29 mice, acute cerebellar slices were prepared. Figure 1A shows a Purkinje cell transduced with SIN-IRE5-GFP-PHD in the cerebellar slice imaged using TPLSM. GFP-PHD was localized mainly in the narrow region of cell borders corresponding to the plasma membrane. This is in accordance with its affinity for PIP₂ on the plasma membrane.
PF-induced GFP-PHD translocation reflects IP₃ dynamics

To confirm that PF-induced GFP-PHD translocation depends on the increase in the intracellular IP₃ concentration, we examined the effect of IP₃ 5-phosphatase overexpression (Fig. 2). Because IP₃ 5-phosphatase hydrolyzes IP₃ but not PIP₂ (Majerus, 1992; Takechi et al., 1998), the overexpression of IP₃ 5-phosphatase should specifically inhibit an increase in cytoplasmic IP₃ concentration. We therefore constructed a Sindbis virus carrying the gene encoding IP₃ 5-phosphatase followed by GFP-PHD under the control of the internal ribosomal entry site [SIN-5-ppase (WT)-IRES-GFP-PHD] to coexpress IP3 5-phosphatase and GFP-PHD. In addition, we also generated a similar virus in which arginine 343 of the IP₃ 5-phosphatase was replaced by alanine [SIN-5-ppase (R343A)-IRES-GFP-PHD]. This mutation (R343A) resulted in the loss of enzymatic activity (Communi et al., 1996) and was used here as a negative control. As shown in Figure 2, most of the PF-induced GFP-PHD translocation was blocked in the IP₃ 5-phosphatase (WT)-expressing cells, whereas the IP₃-induced translocation was observed in cells expressing the R343A-mutant IP₃ 5-phosphatase. These results show the requirement of the increase in IP₃ concentration for the GFP-PHD translocation during PF stimulation, indicating that PF-induced GFP-PHD translocation reports the intracellular IP₃ concentration.

We observed a delay in the appearance of GFP-PHD translocation after PF stimulation. The duration of delay time shows a strong correlation with the peak amplitude of PF-induced GFP-PHD translocation during PF stimulation (Fig. 1B). Averaged time course of fractional changes in fluorescent intensity (ΔF/F0) on PF stimulation showed a transient increase in ΔF/F0 in the plasma membrane and a concomitant decrease in the cytoplasm (Fig. 1C). The changes in the fluorescent intensity indicate the translocation of GFP-PHD from the plasma membrane to the cytoplasm (Hirose et al., 1999; Okubo et al., 2001). The difference in the peak amplitude of PF-induced ΔF/F0 transients between the cytoplasm and the plasma membrane is mainly attributable to a high surface-to-volume ratio in fine dendrites (i.e., the thinner the dendrites, the greater the cytoplasmic increase in the concentration of GFP-PHD for the same amount of translocation from the membrane to the cytoplasm). It is also possible that the definition of the plasma membrane region included the adjacent cytoplasmic region to a small extent, resulting in a slight exaggeration of the difference. Because neuronal activity is known to cause changes in intracellular pH (Chesler and Kaila, 1992) that could affect the fluorescence intensity of GFP (Knee et al., 1998) and thus the GFP-PHD signal, we evaluated PF stimulation-induced changes in GFP fluorescence. PF stimulation (10 stimuli at 50 Hz) was applied to Purkinje cells transduced with Sindbis virus encoding GFP (Okubo et al., 2001). Only a small reduction in ΔF/F0 was observed (∼0.018 ± 0.004, mean of peak change ± SEM, during 1 sec from onset of stimulation, 10 runs from 10 cells), indicating that there is very little effect of PF-induced pH changes on GFP-PHD signals.

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Figure 1. PF-induced translocation of GFP-PHD in Purkinje cells. A, Purkinje cells infected with SIN-IRES-GFP-PHD were imaged with TPLSM. GFP-PHD was excited at 900 nm. The right panel shows the high-magnification image of a part of the dendrite in the left panel (indicated by the white box). Twenty images acquired at 2 μm intervals were projected onto a plane for the left panel, and seven images were projected for the right panel. Fragments of adjacent cells made during the slicing procedure are seen as small dots near and over the Purkinje cells. B, Representative line scan image during PF stimulation. The left panel shows the fluorescence image of a Purkinje cell dendrite. The dashed line indicates the location of time-lapse line scanning. The right panel shows a line scan image on repetitive PF stimulations (10 stimuli at 50 Hz, indicated by the horizontal bar). C, Averaged time courses (14 runs from 14 cells) of fractional changes in the fluorescence intensity (ΔF/F0, normalized by the average of 10 line scans taken immediately before PF stimulation) in the cytoplasm and the plasma membrane. PF stimulation (10 stimuli at 50 Hz) is indicated by the gray bar. Averaged time courses of the reduction in fluorescence intensity attributable to photobleaching were subtracted from the traces.
time, although the finite dissociation rate constant (>5 sec⁻¹) of GFP-PHD from PIP2 (Hirose et al., 1999) may account for a fraction of the observed delay.

**Simultaneous imaging of IP₃ production and Ca²⁺ transients**

Simultaneous imaging of IP₃ and Ca²⁺ transients offers valuable information for understanding the relationship between IP₃ and Ca²⁺ signaling (Hirose et al., 1999) and is indispensable for the evaluation of the involvement of AMPAR in PF-induced IP₃ production via Ca²⁺ influx. We applied the whole-cell patch-clamp method to GFP-PHD-expressing Purkinje cells and introduced into the cells a fluorescent Ca²⁺ indicator, X-rhod-1, through the patch pipette. Both GFP-PHD and X-rhod-1 were excited at 900 nm. Because X-rhod-1 emits a red fluorescence, its signal can be resolved from the fluorescence of GFP-PHD. This enabled us to simultaneously measure the concentrations of Ca²⁺ and IP₃, as well as EPSC, on PF stimulation (Fig. 3A). We voltage clamped Purkinje cells to eliminate the possible Ca²⁺ influx via voltage-gated Ca²⁺ channels triggered by somatic action potentials (Denk et al., 1995) and to evaluate the local involvement of AMPAR in PF-induced IP₃ production. It has been reported that AMPAR-triggered Ca²⁺ influx via voltage-gated Ca²⁺ channels could be observed under voltage clamping because of insufficient clamping in dendritic regions of Purkinje cells (Denk et al., 1995; Eilers et al., 1995; Finch and Augustine, 1998; Takechi et al., 1998).

It was reported that the Ca²⁺ transient induced by a brief train of PF inputs has two distinct components. The initial component coincides with the EPSC and depends on AMPAR-triggered Ca²⁺ influx via voltage-gated Ca²⁺ channels (Denk et al., 1995; Eilers et al., 1995; Finch and Augustine, 1998; Takechi et al., 1998). The delayed component is observed after the end of PF stimulation and is mediated by mGluR-triggered Ca²⁺ release via IP₃R (Finch and Augustine, 1998; Takechi et al., 1998). Thus, we first stimulated PFs with 10 stimuli at 50 Hz with a minimal intensity sufficient for inducing a substantial delayed Ca²⁺ transient (left traces) was followed by PF stimulation at a higher intensity for inducing a substantial initial component in the Ca²⁺ transient (right traces) in the same preparations. Stimulus artifacts in the EPSC traces were omitted for clarity in this and the following figures. B, Time integrals of IP₃ production (during 1.5 sec from the onset of PF stimulation; middle graph), and amplitudes of EPSC (during 200 msec from the onset of PF stimulation; right graph), time integrals of the initial component of Ca²⁺ transients (during 500 ms from the onset of PF stimulation; middle graph), and amplitudes of EPSC evoked by the first stimulus pulse of 10 stimuli at 50 Hz (left graph) are shown (mean ± SEM). Results from seven experiments are compiled.
To investigate the involvement of AMPAR-triggered Ca$^{2+}$ influx in PF-induced IP$_3$ production, we then increased the stimulus intensity to stimulate a larger number of PFs, enhancing AMPAR-triggered Ca$^{2+}$ influx. EPSC showed an increased amplitude by a factor of ~2.7 and a prolonged decay time course (Fig. 3A, bottom traces, B, left graph). There was a substantial increase in the Ca$^{2+}$ concentration, the onset of which coincided with the EPSC (Fig. 3A, middle traces, B, middle graph). There was also a significant delayed component in the Ca$^{2+}$ signal attributable to Ca$^{2+}$ release, although an apparent plateau phase was reached because of the saturation of X-rhod-1 with Ca$^{2+}$ (Fig. 3A, middle traces). IP$_3$ production was considerably increased (Fig. 3A, top traces, B, right graph) when Ca$^{2+}$ transient showed a substantial AMPAR-triggered Ca$^{2+}$ influx.

**PF-induced IP$_3$ production and Ca$^{2+}$ transient were mediated cooperatively by mGluR and AMPAR**

There are two possibilities for the cause of enhanced IP$_3$ production with the increased intensity of PF stimulation as shown in Figure 3. It may be simply attributable to an increased glutamate concentration at the synaptic site inducing an enhanced activation of mGluR. Alternatively, AMPAR-mediated Ca$^{2+}$ influx may have induced IP$_3$ production, as has been shown previously (Okubo et al., 2001). To test these possibilities, we studied the contribution of mGluR and AMPAR to PF-induced IP$_3$ production using specific antagonists for these receptors.

We first studied the effect of a group I mGluR antagonist, (4-CPG, 1 mM). The antagonist reduced PF-induced IP$_3$ production to a level that could not induce a significant GFP-PHD translocation (Fig. 4A, top traces) and blocked the later component of PF-induced Ca$^{2+}$ transient (Fig. 4A, middle traces). The residual Ca$^{2+}$ transient should be attributable mostly to AMPAR-triggered Ca$^{2+}$ influx via the voltage-gated Ca$^{2+}$ channel. Thus, AMPAR-triggered Ca$^{2+}$ influx itself could not induce IP$_3$ production that could induce a significant GFP-PHD translocation under this condition (10 stimuli at 50 Hz) (also see Fig. 5). The peak amplitude of EPSC was unaffected by 4-CPG, although the decay phase of EPSC was partially decreased (Fig. 4A, bottom traces). This 4-CPG-sensitive EPSC component presumably corresponds to the current via the TRPC1 channel triggered by the activation of group I mGluRs (mGluR-EPSC) in Purkinje cells (Templia et al., 1998; Kim et al., 2003). Because it was reported that mGluR-EPSC could mediate Ca$^{2+}$ influx (Templia et al., 2001), it is possible that the 4-CPG-sensitive Ca$^{2+}$ component includes not only Ca$^{2+}$ release from the intracellular store but also Ca$^{2+}$ influx mediated by mGluR-EPSC.

We then studied the effect of an AMPAR antagonist (NBQX, 10 μM). Most of the EPSC was blocked by NBQX (Fig. 4A, bottom traces). The residual EPSC component is probably mediated by mGluR-EPSC. Quite unexpectedly, NBQX blocked not only the first component but also the later component of the Ca$^{2+}$ transient (Fig. 4A, middle traces). The Ca$^{2+}$ release via IP$_3$R from the intracellular store and perhaps Ca$^{2+}$ influx mediated by mGluR-EPSC seems to underlie the residual Ca$^{2+}$ transient. A remarkable feature of the effect of NBQX was that it significantly reduced the PF-induced IP$_3$ production (Fig. 4A, top traces). This result indicates that the activation of mGluR is not sufficient for but the coactivation of mGluR and AMPAR is necessary for the full IP$_3$ production on PF stimulation, although the activation of AMPAR alone cannot induce a significant amount of IP$_3$ production. Indeed, the sum of the IP$_3$ transients in the presence of either 4-CPG or NBQX was significantly smaller than the IP$_3$ transient in the absence of antagonists (Fig. 4B) (also see Fig. 5).

Taken together, the effects of 4-CPG and NBQX on the PF-induced IP$_3$ production indicate that PF-induced IP$_3$ production is dependent on the simultaneous activation of group I mGluR and AMPAR. Thus, there is a robust crosstalk between mGluR and iGluR for IP$_3$ production at the PF–Purkinje cell synapse.

We then examined how the crosstalk between mGluR and AMPAR affects the PF-induced Ca$^{2+}$ transients. Because the Ca$^{2+}$ transients measured by X-rhod-1 (K$_p$ 700 nM) show the sign of saturation (Fig. 4A, Control), we imaged the PF-induced Ca$^{2+}$ response using a low-affinity Ca$^{2+}$ indicator, Calcium Green-5N (K$_p$ 14 μM) (Fig. 4C). Calcium Green-5N was applied through the patch pipette to Purkinje cells and was excited at 810 nm. In the presence of 4-CPG, a greater part of the Ca$^{2+}$ transient was blocked, leaving the small response coinciding with PF stimulation. NBQX almost completely blocked the PF-induced Ca$^{2+}$

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**Figure 4.** Effect of glutamate receptor antagonists on PF-induced response. A. Effects of the group I mGluR antagonist 4-CPG (1 mM) and AMPAR antagonist NBQX (10 μM). Averaged traces (7 runs from 7 cells for the 4-CPG experiment, 6 runs from 6 cells for the NBQX experiment) of IP$_3$ production (top traces), Ca$^{2+}$ transient (middle traces), and EPSC (bottom traces) during PF stimulation (10 stimuli at 50 Hz, indicated by the gray bar). After the measurement under the control condition, 4-CPG or NBQX was perfused for 4–7 min, and then the measurement was performed at the same location. B, Cooperativity in IP$_3$ production during PF stimulation. The linear sum of the averaged traces in the presence of 4-CPG or NBQX shown in A is compared with an averaged trace under the control condition (13 runs from 13 cells), C. Cooperativity in Ca$^{2+}$ transient detected using Calcium Green-5N. Averaged time courses (9 runs from 9 cells for the 4-CPG experiment, 8 runs from 8 cells for the NBQX experiment) of the Ca$^{2+}$ transient on PF stimulation are shown (top traces). The linear sum of averaged traces in the presence of 4-CPG or NBQX is significantly smaller than that of the averaged trace under the control condition (bottom trace).
response. The Ca\(^{2+}\) transient under the control condition was significantly larger than the sum of the Ca\(^{2+}\) transients in the presence of either antagonist.

**Dependence on the number of PF inputs of the cooperative IP\(_3\) production**

To study how the cooperativity of mGluR and AMPAR in IP\(_3\) signaling depends on the intensity of synaptic inputs, we examined the magnitude of IP\(_3\) production on the number of PF stimulations while maintaining the frequency at 50 Hz. In the absence of antagonists, trains of as few as three pulses induced detectable IP\(_3\) production, and the size of the response showed a steep dependence on the number of pulses between 3 and 20 pulses (Fig. 5A, black circles). The amplitude and the duration of PF-induced IP\(_3\) production saturated at 20 pulses. When mGluR was blocked in the presence of 1 mM 4-CPG, no significant IP\(_3\) production was observed in response to 10 or fewer pulses, showing that the activation of AMPAR alone is not sufficient for IP\(_3\) production in this range of PF inputs (Fig. 5A) (also see Fig. 4A). However, significant IP\(_3\) production was observed on 20 and 40 pulses of PF inputs, although the magnitude was approximately half that of control experiments (Fig. 5A, red circles). Thus, strong AMPAR activation could induce IP\(_3\) production in Purkinje cells, in accordance with our previous report (Okubo et al., 2001). When AMPAR was blocked by 10 \(\mu\)M NBQX, the amplitude of PF-induced IP\(_3\) production increased with the number of pulses more than 3 (Fig. 5A, blue circles). The mGluR-triggered IP\(_3\) production showed a less steep dependence on the number of pulses than the AMPAR-mediated IP\(_3\) production. The sum of the IP\(_3\) transients in the presence of either 4-CPG or NBQX was significantly smaller than the IP\(_3\) transient in the absence of antagonists at 5 or 10 pulses, but the two values were not significantly different at greater number of pulses (Fig. 5C). Thus, the cooperativity of mGluR and AMPAR in IP\(_3\) signaling is dependent on the number of pulses and is prominent in the window of 5–10 pulses at 50 Hz, which is within the physiological firing patterns of PFs (Merrill et al., 1978; Chadderton et al., 2004).

**PF-induced IP\(_3\) production required both G-protein activation and an increase in intracellular Ca\(^{2+}\) concentration**

Having clarified the cross talk between mGluR and AMPAR in IP\(_3\) production, we then examined the intracellular mechanisms involved in the cross talk. Because mGluR is one of the G-protein-coupled receptors (Masu et al., 1991; Aramori and Nakanishi, 1992), we investigated the involvement of the G-protein in PF-induced IP\(_3\) production using GDP-\(\beta\)S to inhibit G-protein activity. GDP-\(\beta\)S (2 \(\mu\)M) was introduced into the cytoplasm through the patch pipette. In the presence of GDP-\(\beta\)S, PF-mediated IP\(_3\) production was almost completely inhibited (Fig. 6, left traces). The PF-induced Ca\(^{2+}\) transients in the presence of GDP-\(\beta\)S were similar to those in the presence of 4-CPG (Fig. 4A, middle traces), showing that the AMPAR-dependent Ca\(^{2+}\) transient is normally activated in the presence of GDP-\(\beta\)S. This result shows that the signaling pathway via the G-protein is critical for the cross talk between mGluR and AMPAR in IP\(_3\) production.

Next, we evaluated the signaling event triggered by the activation of AMPAR. In Purkinje cells, the activation of AMPAR is followed by depolarization, triggering Ca\(^{2+}\) influx via the P-type voltage-gated Ca\(^{2+}\) channel (Gruol et al., 1996). Thus, there is a possibility that the AMPAR-triggered Ca\(^{2+}\) influx enhances PF-induced IP\(_3\) production when mGluR is activated. To evaluate the involvement of Ca\(^{2+}\), we applied BAPTA through the patch pipette to buffer intracellular Ca\(^{2+}\) (Fig. 6, right traces). We observed no significant IP\(_3\) production in the presence of BAPTA. This observation was made possible by the direct measurement of IP\(_3\) generation and indicates that an increase in intracellular Ca\(^{2+}\) concentration is necessary for PF-induced IP\(_3\) production.

These results indicate that the coincidental increase in G-protein activity and intracellular Ca\(^{2+}\) concentration is necessary for PF-induced IP\(_3\) production. This strongly suggests that the cross talk between mGluR and AMPAR in IP\(_3\) production is mediated through mGluR-triggered G-protein activation and AMPAR-triggered Ca\(^{2+}\) influx.
G-protein (Rhee and Bae, 1997; Rebecchi and Pentyala, 2000), explain the supralinear increase in IP3 concentration.

Possible physiological roles of the mGlur–AMPAR cross talk

The group I mGlurRs (including mGlur1) induce IP3 production through the activation of PLC-β isozymes that are coupled to the G-protein (Masu et al., 1991; Aramori and Nakanishi, 1992). Because Purkinje cells highly express mGlur1 (Baude et al., 1993) and PLC-β isozymes (Watanabe et al., 1998), it is likely that the PF input-induced IP3 production (Fig. 4B, top trace) is mediated by the G-protein–PLC-β signaling pathway. In accordance with this notion, PF-mediated IP3 production was completely blocked by the inhibition of G-protein activity by GDP-βS (Fig. 6). Although the activity of PLC-β is primarily regulated by the G-protein (Rhee and Bae, 1997; Rebecchi and Pentyala, 2000), the PLC-β activity is enhanced by an increase in intracellular Ca2+ concentration (Taylor et al., 1991; Hirose et al., 1999). Indeed, IP3 production was abrogated when the intracellular Ca2+ concentration was buffered by BAPTA (Fig. 6). Taken together, PLC-β is the most plausible candidate protein that mediates the cross talk between mGlur-mediated G-protein activation and AMPAR-triggered Ca2+ influx, although saturation of IP3-metabolizing enzymes and/or IP3-binding proteins may partially explain the supralinear increase in IP3 concentration.

Figure 6. PF-induced IP3 production depends on G-protein and intracellular Ca2+. Averaged traces (6 runs from 5 cells for the GDP-βS experiment, 7 runs from 6 cells for the BAPTA experiment) of IP3 production, Ca2+ transient, and EPSC during PF stimulation (10 stimuli at 50 Hz, indicated by the gray bar) are shown. PF-induced IP3 production was completely blocked by the intracellular delivery of 2 mM GDP-βS (left traces) or 20 mM BAPTA (right traces).

Discussion

We visualized and analyzed the PF-induced IP3 dynamics in Purkinje cells. The present study showed that PF-induced IP3 production is cooperatively mediated by mGlur and AMPAR via G-protein activation and Ca2+ influx. The present results may shed light on the functional significance of the close distribution of mGlur and iGlur at excitatory synapses in the CNS.

Candidate components that mediate the mGlur-AMPAR cross talk

The group I mGlurRs (including mGlur1) induce IP3 production through the activation of PLC-β isozymes that are coupled to the G-protein (Masu et al., 1991; Aramori and Nakanishi, 1992). Because Purkinje cells highly express mGlur1 (Baude et al., 1993) and PLC-β isozymes (Watanabe et al., 1998), it is likely that the PF input-induced IP3 production (Fig. 4B, top trace) is mediated by the G-protein–PLC-β signaling pathway. In accordance with this notion, PF-mediated IP3 production was completely blocked by the inhibition of G-protein activity by GDP-βS (Fig. 6). Although the activity of PLC-β is primarily regulated by the G-protein (Rhee and Bae, 1997; Rebecchi and Pentyala, 2000), the PLC-β activity is enhanced by an increase in intracellular Ca2+ concentration (Taylor et al., 1991; Hirose et al., 1999). Indeed, IP3 production was abrogated when the intracellular Ca2+ concentration was buffered by BAPTA (Fig. 6). Taken together, PLC-β is the most plausible candidate protein that mediates the cross talk between mGlur-mediated G-protein activation and AMPAR-triggered Ca2+ influx, although saturation of IP3-metabolizing enzymes and/or IP3-binding proteins may partially explain the supralinear increase in IP3 concentration.

Possible physiological roles of the mGlur–AMPAR cross talk

PF→Purkinje cell synapses show synaptic plasticity, such as long-term depression (LTD), which requires the concomitant activation of PF and climbing fiber (CF) synapses, and may underlie certain forms of motor learning (Ito, 2001). It was reported that, at the PF→Purkinje cell synapse, the simultaneous activation of AMPAR and mGlur is necessary for the induction of LTD (Kano and Kato, 1988; Linden et al., 1991; Hartell, 1994; Hemart et al., 1995). Indeed, LTD is absent in mGlur1-deficient mice (Aiba et al., 1994; Conquet et al., 1994). However, the role of AMPAR remains unclear. The present results indicate that AMPAR at the PF synapses cooperates with mGlur to activate PLC-β, which is involved in the induction of LTD (Miyata et al., 2001). This cooperative activation of PLC-β may be the reason why the simultaneous activation of AMPAR and mGlur at PF synapses is necessary for the induction of LTD.

It was reported that PF inputs in combination with a CF-triggered Ca2+ influx, the stimulus used for LTD induction, induce a supralinear increase in Ca2+ release from the intracellular store within spines (Wang et al., 2000). The supralinear Ca2+ release probably involves the cooperative activation of IP3, which requires both IP3 and Ca2+ for full activation (Iino, 1990; Bezprozvanny et al., 1991). However, possible cooperative production of IP3 by simultaneous inputs from PFs and a CF may also contribute to the generation of supralinear Ca2+ release. Future development of a technique that allows the imaging of IP3 dynamics within spines may clarify this issue.

The activation of PLC enzymes produces not only IP3, but also diacylglycerol, which is a precursor of endocannabinoids (Stella et al., 1997) as well as an activator of PKC. Endocannabinoids are retrograde signaling molecules that modulate the release rate of neurotransmitters in many brain regions and are produced on two distinct signals, an increase in Ca2+ concentration (Di Marzo et al., 1994; Stella et al., 1997) and G-protein activation via group I mGlur (Maejima et al., 2001). The present results, therefore, suggest that mGlur and AMPAR may cooperatively mediate the production of endocannabinoids at PF synapses. In general agreement with this notion, mGlur activation and depolarization-induced Ca2+ influx cooperatively generated endocannabinoids in cultured hippocampal neurons (Ohno-Shosaku et al., 2002).

In summary, the functional interaction between mGlur and iGlur was identified in PF→Purkinje cell synapses. IP3 production was cooperatively induced by the activation of mGlur and iGlur, and the molecular coincidence detector is most likely PLC-β. This cooperation between mGlur and iGlur may play important roles in the regulation of various synaptic functions.

References


