Role of different voltage-gated Ca$^{2+}$ channels in cortical spreading depression

Specific requirement of P/Q-type Ca$^{2+}$ channels

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Gain-of-function mutations in Cav2.1 (P/Q-type) Ca$^{2+}$ channels cause familial hemiplegic migraine type 1 (FHM1), a subtype of migraine with aura. Knockin (KI) mice carrying FHM1 mutations show increased neuronal P/Q-type Ca$^{2+}$ current and facilitation of induction and propagation of cortical spreading depression (CSD), the phenomenon that underlies migraine aura and may activate migraine headache mechanisms. We recently studied cortical neurotransmission in neuronal microcultures and brain slices of FHM1 KI mice, and showed (1) gain-of-function of excitatory neurotransmission, due to increased action potential-evoked Ca$^{2+}$ influx and increased probability of glutamate release at pyramidal cell synapses, but unaltered inhibitory neurotransmission at fast-spiking interneuron synapses and (2) a causative link between enhanced glutamate release and facilitation of CSD induced by brief pulses of high K$^+$ in cortical slices. Here, we show that blockage of either the P/Q-type Ca$^{2+}$ channels or the NMDA receptors, CSD cannot be induced in wild-type mouse cortical slices. In contrast, blockade of N- or R-type Ca$^{2+}$ channels has only a small inhibitory effect on CSD threshold and velocity of propagation. Our findings support a model in which Ca$^{2+}$ influx through presynaptic P/Q-type Ca$^{2+}$ channels with consequent release of glutamate from recurrent cortical pyramidal cell synapses and activation of NMDA receptors are required for initiation and propagation of the CSD involved in migraine.

Introduction

Missense mutations in the gene that encodes the pore-forming α subunit of voltage-gated Ca$^{2+}$ channels cause a rare autosomal dominant subtype of migraine with aura: familial hemiplegic migraine type 1 (FHM1). Cav2.1 channels are located in somatodendritic membranes and presynaptic terminals throughout the brain, and play a dominant role in initiating action potential-evoked neurotransmitter release at central nervous system synapses. FHM1 mutations produce gain-of-function of human recombinant Cav2.1 channels, mainly due to a shift to lower voltages of channel activation. Knockin (KI) mice carrying FHM1 mutations show increased P/Q-type Ca$^{2+}$ current density in central neurons including cortical pyramidal cells. Interestingly, the FHM1 KI mice also show a reduced threshold for induction of cortical spreading depression (CSD) and an increased velocity of propagation of CSD. CSD can be induced in animals by focal stimulation of the cerebral cortex and consists in a slowly propagating wave of cortical neuronal and glial depolarization, whose mechanisms remain unclear and controversial. Neuroimaging studies in humans indicate that CSD underlies the migraine aura; animal studies indicate that CSD may also trigger the migraine headache mechanisms. To study the cortical mechanisms that produce facilitation of CSD in FHM1 mutant mice, we investigated cortical neurotransmission in neuronal microcultures and in brain slices from FHM1 KI mice.
mice. The results show increased strength of excitatory synaptic transmission due to increased action potential-evoked Ca\textsuperscript{2+} influx through presynaptic P/Q-type Ca\textsuperscript{2+} channels and increased probability of glutamate release at cortical pyramidal cell synapses of mutant mice. In striking contrast, inhibitory neurotransmission at connected pairs of fast-spiking interneurons and pyramidal cells was unaltered in FHM1 mice, despite being initiated by P/Q-type Ca\textsuperscript{2+} channels. To test the hypothesis that the gain-of-function of glutamate release at synapses onto pyramidal cells may explain the facilitation of induction and propagation of experimental CSD in FHM1 KI mice, we measured the threshold for CSD induction and the velocity of CSD propagation in acute slices of somatosensory cortex of R192Q KI mice before and after perfusion with a concentration of ωAgIVA that reduced glutamate release at KI pyramidal cell synapses to wild-type (WT) levels. Strikingly, restoration of glutamate release to WT levels completely rescued CSD facilitation, as both CSD triggering threshold and CSD propagation rate in mutant mice became similar to those in WT mice. This finding provides direct evidence of a causative link between enhanced glutamate release at pyramidal cell synapses and facilitation of experimental CSD, thus giving insights into the controversial mechanisms of CSD initiation and propagation. The initiation of the positive feedback cycle that ignites CSD and almost zeroes the neuronal membrane potential depends on the local increase of the extracellular concentration of K\textsuperscript{+} ions [K\textsuperscript{+]\textsubscript{e}} above a critical value and on the activation of a net inward current at the pyramidal cell dendrites; the nature of the cationic channels mediating this inward current remains unclear and controversial, although there is strong pharmacological support for a key role of NMDA receptors (cf also Discussion and references therein). The findings of Tottene et al. support a model of CSD initiation in which activation of presynaptic P/Q-type Ca\textsuperscript{2+} channels with consequent release of glutamate from recurrent cortical pyramidal cell synapses and activation of NMDA receptors (and possibly postsynaptic P/Q-type Ca\textsuperscript{2+} channels) are key components of the positive feedback cycle that ignites CSD in normally metabolizing cortical tissue. Regarding CSD propagation, the findings are consistent with a model based on interstitial K\textsuperscript{+} diffusion initiating in adjacent dendrites this positive feedback cycle.

Since it is controversial whether activation of NMDA receptors and in particular Ca\textsuperscript{2+} influx and synaptic transmission play only a modulatory role or are required for CSD induction and propagation,\textsuperscript{9,10} we further investigated the role of NMDA receptors and of the different voltage-gated Ca\textsuperscript{2+} channel by studying the effect of specific blockers of these channels on the threshold for CSD induction and the velocity of CSD propagation in acute slices of mouse cerebral cortex.

## Results

To investigate the role of the NMDA receptors and the P/Q-, N-, R- and L-type Ca\textsuperscript{2+} channels in initiation and propagation of experimental CSD, we measured the threshold for CSD induction and the velocity of CSD propagation, induced by brief pulses of high K\textsuperscript{+} in acute slices of mouse somatosensory cortex, before and after application of saturating concentrations of D-AP5 (50 μM), ωAgIVA (300 or 400 nM), ωCgTxGVIA (1 μM), SNX-482 (300 or 500 nM) and nimodipine (10 μM), respectively, as described in Tottene et al.\textsuperscript{1} Pressure pulses of increasing duration were applied to a 3 M KCl-containing pipette positioned on layer 2/3 until a CSD was observed. CSD was revealed by both the associated changes in light transmittance and the typical depolarization to almost zero mV recorded in a pyramidal cell located 600 μm apart from the pressure-ejection pipette tip (to ensure that a true propagating event was studied). The duration of the first pulse eliciting a CSD was taken as CSD threshold, and the rate of horizontal spread of the change in intrinsic optical signal as velocity of CSD propagation.\textsuperscript{2}

CSD could not be induced after blocking either the NMDA receptors or the P/Q-type Ca\textsuperscript{2+} channels; in fact, in the presence of either D-AP5 or ωAgIVA, neither a depolarization of the patch-clamped pyramidal cell nor a propagating change in intrinsic optic signal were measured even with K\textsuperscript{+} pulses 30 times longer than the threshold pulses triggering a CSD in control (Table 1). After the inhibition of P/Q-type Ca\textsuperscript{2+} channels, K\textsuperscript{+} pulses of 10 s duration were unable to elicit a CSD in slices in which the average threshold duration in control was 0.29 ± 0.01 s (n = 7). Similarly, after the inhibition of NMDA receptors with D-AP5, K\textsuperscript{+} pulses of 10 s duration were unable to elicit a CSD in slices in which the average threshold duration in control was 0.34 ± 0.03 s (n = 5). Moreover, CSD was also not elicited in nine (in the presence of ωAgIVA) and nine (in the presence of D-AP5) additional slices in which the maximal duration of the K\textsuperscript{+} pulse was 1 s (about three times larger than the threshold in control) (Table 1).

In contrast, inhibition of the other types of presynaptic voltage gated Ca\textsuperscript{2+} channels had only a relatively small effect on CSD initiation and propagation. In the presence of either ωCgTxGVIA (n = 13) or SNX-482 (n = 10), the CSD threshold was about 10% higher and the CSD rate of propagation 15% lower than in control (Fig. 1 and Table 1). Neither threshold nor velocity of propagation of CSD was significantly affected by inhibition of L-type Ca\textsuperscript{2+} channels with nimodipine (n = 5) (Table 1).

## Discussion

Our pharmacological study supports the conclusions that (1) Ca\textsuperscript{2+} influx through P/Q-type Ca\textsuperscript{2+} channels and activation of NMDA receptors are required for CSD induction by K\textsuperscript{+} pressure pulses in acute slices of mouse sensory cortex; (2) Ca\textsuperscript{2+} influx through N- and, probably, R-type Ca\textsuperscript{2+} channels may play a modulatory role on CSD threshold and velocity of propagation. In agreement with our findings, most previous studies investigating the effect of NMDA receptor antagonists on CSD, induced by brief K\textsuperscript{+} pulses in cortical slices, reported complete blockade of CSD recorded at ≥500 μm from the local K\textsuperscript{+} ejection (but see ref. 18 for an exception in hippocampal slices); moreover CSD could not be recorded after perfusing the slices in Ca\textsuperscript{2+}-free medium or after blocking the Ca\textsuperscript{2+} channels.
higher electrical stimulation intensity to evoke a CSD than wild-type mice. However, if one considers previous in vitro and in vivo pharmacological studies of CSD induced by perfusing the cortical slices or the cortex with a high K+ solution (rather than with brief K+ pulses or electrical stimulation) a completely different picture emerges: NMDA antagonists only slightly increase the CSD threshold without affecting its velocity; similarly, blockade of the P/Q-type (or the N-type) Ca2+ channels hardly affects the CSD threshold obtained by perfusing cortical slices or the NMDA receptors, stimuli 30 times larger than the CSD triggering threshold were unable to induce a CSD. Our findings in cortical slices are consistent with in vivo studies of CSD induced by electrical stimulation of the cortex, showing that, after i.p. injection of NMDA antagonists, even stimulation currents ten times longer and eight times larger than the CSD triggering threshold were unable to induce a CSD, and spontaneous cacna1a mouse mutants, with mutations that produce partial loss-of-function of the P/Q-type Ca2+ channel, required an approximately 10 fold higher electrical stimulation intensity to evoke a CSD than wild-type mice.

However, if one considers previous in vitro and in vivo pharmacological studies of CSD induced by perfusing the cortical slices or the cortex with a high K+ solution (rather than with brief K+ pulses or electrical stimulation) a completely different picture emerges: NMDA antagonists only slightly increase the CSD threshold without affecting its velocity; similarly, blockade of the P/Q-type (or the N-type) Ca2+ channels hardly affects the CSD threshold obtained by perfusing cortical slices or the NMDA receptors, stimuli 30 times larger than the CSD threshold are unable to induce a CSD.

Figure 1. Relative values of CSD threshold and velocity before and after inhibition of N-type or R-type Ca2+ channels. The threshold for CSD induction and the velocity of CSD propagation were measured before (control) and after application of saturating concentrations of ω-AgaIVA, ω-CgTxGVIA, SNX-482 and nimodipine to inhibit NMDA receptors and P/Q-, N-, R- and L-type Ca2+ channels, respectively. CSD was induced in acute cortical slices by high K+ pressure pulses of increasing duration: the duration of the first pulse eliciting a CSD was taken as CSD threshold and the rate of horizontal spread of the change in intrinsic optical signal as velocity of CSD propagation. In the presence of ω-AgaIVA or D-AP5, the CSD could not be induced even with very long pulses, about 3 (1 s) or 30 (10 s) times larger than the control CSD triggering threshold.

Table 1. Effect of inhibition of the different types of voltage-gated Ca2+ channels or the NMDA receptors on CSD threshold and CSD velocity

| Channel inhibition | Control | Channel inhibition |
|--------------------|--------|--------------------|
|                     | n      | Threshold (ms)     | Velocity (mm/min) |
| ω-AgaIVA            | 7      | 293 ± 12           | 2.3 ± 0.2         |
| D-AP5               | 9      | 292 ± 7            | 2.1 ± 0.1         |
|                     | 5      | 338 ± 27           | 1.8 ± 0.1         |
|                     | 9      | 286 ± 10           | 2.4 ± 0.1         |
| ω-CgTxGVIA          | 13     | 261 ± 9            | 2.1 ± 0.1         |
| SNX-482             | 10     | 288 ± 9            | 2.4 ± 0.2         |
| Nimodipine          | 5      | 310 ± 21           | 2.2 ± 0.1         |

The threshold for CSD induction and the velocity of CSD propagation were measured before (control) and after (channel inhibition) application of saturating concentrations of D-AP5, ω-AgaIVA, ω-CgTxVIA, SNX-482 and nimodipine to inhibit NMDA receptors and P/Q-, N-, R- and L-type Ca2+ channels, respectively. CSD was induced in acute cortical slices by high K+ pressure pulses of increasing duration: the duration of the first pulse eliciting a CSD was taken as CSD threshold and the rate of horizontal spread of the change in intrinsic optical signal as velocity of CSD propagation.
slices with progressively increasing K+ concentrations, and removal of extracellular Ca2+ ions does not block CSD induced by perfusing the slices with a high K+ solution for 80–90 s (but reduces to about half the rate of propagation). Moreover, multiple CSDs induced in vivo by continuous K+ microdialysis or topical application of a KCl crystal are strongly reduced in frequency but not completely suppressed by P/Q-type (or N-type) Ca2+ channel blockers and by NMDA antagonists (that also reduce their amplitude and duration). The Ca2+ channel blockers do not affect CSD induced by pinprick in vivo.

Thus, even limiting our discussion to the methods that are used to elicit CSD in normally metabolizing cortical tissues (because the pharmacological profile of hypoxia and/or ouabain-induced CSD is again different), it is clear that different methods lead to different pharmacological profiles regarding the role of NMDA receptors and Ca2+ influx through P/Q-type Ca2+ channels in CSD induction and propagation. The different pharmacological profiles likely reflect the fact that CSD is a complex phenomenon and there may be sequential mechanisms with different pharmacology leading to the final common downstream CSD event; some of the upstream mechanisms may be variably bypassed and/or occluded by the different CSD-inducing methods.

It becomes then crucial to understand which experimental CSD-inducing method is more relevant (and eventually most predictive of drug efficacy) for the CSDs that arise “spontaneously” in a given brain pathology. In the case of migraine, insights into how “spontaneous” CSDs in migraineurs. The specificity of the P/Q-type Ca2+ channel requirement compared to that of the other presynaptic N- and R-type Ca2+ channels may reflect the fact that excitatory synaptic transmission at pyramidal cell synapses in different areas of the cerebral cortex depends predominantly on P/Q-type Ca2+ channels (reviewed in ref. 28), and/or may point to a specific role of postsynaptic P/Q-type Ca2+ channels in CSD induction.

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References

1. Tootene A, Conti R, Fabbro A, Vecchia D, Shapovalova M, Santello M, et al. Enhanced excitatory transmission at cortical synapses as the basis for facilitated spreading depression in Caja1 knockin migraine mice. Neuron 2009; 61:762-73.
2. Ophoff RA, Terwindt GM, Verdure MN, van Eijk R, Offner PJ, Hoffman SMG, et al. Familial hemiplegic migraine and episodic ataxia type-2 are caused by mutations in the Ca2+ channel gene CACNL1A4. Cell 1996; 87:543-52.
3. Pietrobon D. Function and dysfunction of synaptic calcium channels: insights from mouse models. Curr Opin Neurobiol 2005; 15:257-65.
4. Pietrobon D. Ca2+ channelopathies. Pflugers Arch 2010; 460:375-93.
5. Tootene A, Fellin T, Pagnutti S, Lisovette S, Striessnig J,letcher C, et al. Familial hemiplegic migraine mutations increase Ca2+ influx through single human Caja1.2 channels and decrease maximal Ca2+1 current density in neurons. PNAS 2002; 99:13284-9.
6. van den Maagdenberg AM, Pietrobon D, Pizzorusso T, Kaja S, Broos LA, Ceserti T, et al. A Caca1a knockin migraine mouse model with increased susceptibility to cortical spreading depression. Neuron 2004; 41:701-10.
7. Inchauste CG, Urbano FJ, Di Giuli MN, Forsythe JD, Ferrari MD, van den Maagdenberg AM, et al. Gain of function in FHM-1 Caja1 knock-in mice is related to the shape of the action potential. J Neurophysiol 2010; 104:291-9.
8. van den Maagdenberg AM, Pizzorusso T, Kaja S, Tepoloni N, Shapovalova M, Hoebeek FE, et al. High cortical spreading depression susceptibility and migraine-associated symptoms in Caja1.2 S1218L mice. Ann Neurol 2010; 67:85-98.
9. Sonjen GC. Mechanisms of spreading depression and hypoxic spreading depression-like depolarization. Physiol Rev 2001; 81:1065-96.
10. Charles A, Brenan K. Cortical spreading depression—new insights and persistent questions. Cephalalgia 2009; 29:1115-24.
11. Pietrobon D, Striessnig J. Neurobiology of migraine. Nat Rev Neurosci 2003; 4:386-98.
12. Ayata C. Spreading depression: from serendipity to targeted therapy in migraine prophylaxis. Cephalalgia 2009; 29:1095-114.
13. Zhang X, Levy D, Nosea R, Kainz V, Jakubowski M, Burstein R. Activation of meningeal nociceptors by cortical spreading depression: implications for migraine with aura. J Neurosci 2010; 30:8807-14.
14. Footitt DR, Newberry NR. Cortical spreading depression induces an LTP-like effect in rat neocortex in vitro. Brain Res 1998; 781:359-42.
15. Vilaj I, Klupa N, Luhmann JH. Optical recording of spreading depression in rat neocortical slices. Brain Res 2001; 898:288-96.
16. Peters O, Schipke CG, Hashimoto Y, Kettenmann H. Different mechanisms promote astrocyte Ca2+ waves and spreading depression in the mouse neocortex. J Neurosci 2003; 23:9888-96.
17. Perlzold GC, Windmoller O, Haack S, Major S, Buchheim K, Megow D, et al. Increased extracellular K+ concentration reduces the efficacy of N-methyl-D-aspartate receptor antagonists to block spreading depression-like depolarizations and spreading ischemia. Stroke 2005; 36:1270-7.
18. Larrosa B, Pastor J, Lopez-Aguado L, Herreraza O. A role for glutamate and glia in the fast network oscillations preceding spreading depression. Neuroscience 2006; 141:1057-68.
19. Jing J, Artken PG, Sonjen GC. Role of calcium channels in spreading depression in rat hippocampal slices. Brain Res 1993; 604:251-9.
20. Marrannes R, Willems R, De Prins E, Wauquier A. Evidence for a role of the N-methyl-D-aspartate (NMDA) receptor in cortical spreading depression in the rat. Brain Res 1988; 457:226-40.
21. Ayata C, Shimizu-Sasamata M, Lo EH, Noebels JL, Moskovitz. Impaired neurotransmitter release and elevated threshold for cortical spreading depression in mice with mutations in the α1 subunit of P/Q type calcium channels. Neuroscience 2000; 95:639-45.
22. Perlzold GC, Haack S, von Bohlen Und Halbach O, Priller J, Lehmann TN, Heinemann U, et al. Nitric oxide modulates spreading depolarization threshold in the human and rodent cortex. Stroke 2008; 39:1292-9.
23. Zhou N, Gordon GR, Frighan D, MacVicar BA. Transient swelling, acidification, and mitochondrial depolarization occurs in neurons but not astrocytes during spreading depression. Cereb Cortex 2010; 20:2614-24.
24. Richter F, Ebersberger A, Schaible HG. Blockade of voltage-gated calcium channels in rat inhibits repetitive cortical spreading depression. Neurosci Lett 2002; 34:123-6.
25. Herreras O, Somjen GG. Analysis of potential shifts associated with recurrent spreading depression and prolonged unstable SD induced by microdialysis of elevated K+ in hippocampus of anesthetized rats. Brain Res 1993; 610:283-94.
26. Peeters M, Gunthorpe MJ, Strijbos LM, Goldsmith P, Upton N, James MF. Effects of pan-and subtype-selective N-Methyl-D-aspartate receptor antagonists on cortical spreading depression in the rat: therapeutic potential for migraine. J Pharmacol Exp Ther 2007; 321:564-72.
27. Akerman S, Holland PR, Goadsby PJ. Mechanically-induced cortical spreading depression associated regional cerebral blood flow changes are blocked by Na+ ion channel blockade. Brain Res 2008; 1229:27-36.
28. Pietrobon D. Insights into migraine mechanisms and Ca2.1 calcium channel function from mouse models of familial hemiplegic migraine. J Physiol 2010; 588:1871-8.