SHORT COMMUNICATION

A Mechanism for Nerve Cell Excitation by Norepinephrine via Alpha-1 Adrenoceptors: Inhibition of Potassium M-Current

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Abstract Some of the excitatory effects of norepinephrine on central neurons are mediated by alpha-1 (\(α_1\)) adrenoceptors. These receptors are coupled to the Gq family of G proteins, and hence stimulate hydrolysis of the membrane phospholipid phosphatidylinositol-4,5-bisphosphate. Other receptors of this type can excite neurons by inhibiting the subthreshold voltage-gated potassium M-current. We tested this possibility using rat sympathetic neurons transformed to express \(α_1α\) receptors. The \(α_1\) agonist phenylephrine strongly inhibited the M-current recorded under voltage-clamp by 72 ± 11 % (\(n = 4\)) and in an unclamped neuron dramatically increased the number of action potentials produced by a 2 s depolarizing current step from 2 to 40, without effect on control neurons devoid of \(α_1\) receptors. We suggest that this might be a potential cause of the increased excitability produced by norepinephrine in some central neurons.

Keywords \(α_1\) Adrenoceptors · Norepinephrine · Phenylephrine · M-current · Potassium current · Sympathetic neurons

Introduction

Alpha-1 (\(α_1\)) adrenoceptors are widely distributed in the central nervous system (Jones et al. 1985; Day et al. 1997). There they mediate some of the effects of norepinephrine, including a membrane depolarization and enhanced action potential firing (e.g., supra-optic nucleus: Ogata and Matsuo 1986; cerebral cortex and thalamus: McCormick et al. 1991; lateral geniculate nucleus: McCormick 1992).

These receptors are G protein-coupled receptors (GPCRs) that link primarily to the Gq family of G proteins and thereby stimulate hydrolysis of the membrane phospholipid phosphatidylinositol-4,5-bisphosphate (PIP\(_2\)) (Hawrylyshyn et al. 2004). Other receptors of this type can excite neurons by inhibiting the subthreshold voltage-gated K\(^+\) current termed the “M-current” (Delmas and Brown 2005), which normally acts as a brake on firing activity. However, this action does not appear to have been hitherto detected in mammalian central neurons following \(α_1\)-stimulation. This may be because most of the cells studied so far did not possess prominent M-currents, or perhaps because the current is not very responsive to \(α_1\) receptor stimulation (though tests on peripheral parasympathetic neurons suggest otherwise: Shibata and Taketani 2001).

We have recently had an opportunity to assess the response of M-currents to specific \(α_1\)-adrenoceptor stimulation using neurons from the rat superior cervical sympathetic ganglion. These have large M-currents (Constanti and Brown 1981) but do not seem to possess an appreciable number of endogenous \(α_1\) receptors (Grayson et al. 1998; Dawson et al. 2010). Hence we inserted \(α_1α\) adrenoceptors by intranuclear cDNA injection, as used for other receptors (see Ikeda 1997; Filippov et al. 2000). We show that this receptor is indeed well capable of inhibiting the M-current, and can thereby strikingly enhance excitability.

Materials and Methods

Superior cervical sympathetic ganglia were isolated from 18-day-old rats (killed by UK Home Office approved methods), neurons dissociated, plated and cultured in vitro for
1–2 days as described by Filippov et al. (1998). To express z1a receptors, a cDNA plasmid encoding the receptor (provided by Dr. John Pediani, Glasgow University; see Pediani et al. 2000) was injected into individual cell nuclei 4 h after initial plating, together with a cDNA plasmid encoding the jellyfish enhanced green fluorescent protein (eGFP) to mark the injected neurons. For electrophysiological recording, cells were patched with electrodes containing (in mM) 90 potassium acetate, 20 KCl, 3 MgCl₂, 40 HEPES, and 0.1 BAPTA (pH-adjusted to 7.4 by KOH), and back-filled with the same solution containing 0.125 mg/ml amphotericin B as a membrane permeabilizing agent, to provide “perforated-patch” recording. (This is essential to retain the normal cytosolic constituents and maintain normal M-currents and excitability.) Cells were maintained in flowing Krebs’ solution at 20–22°C containing (in mM) 120 NaCl, 3 KCl, 1.5 MgCl₂, 2.5 CaCl₂, 10 HEPES, and 11.1 glucose (pH-adjusted to 7.3 with NaOH). Recordings were made, data collected and analyzed as described by Filippov et al. (1998).

Results

M-Current

When these sympathetic neurons are clamped at a depolarized membrane potential, potassium M-channels open and generate a steady outward current (they do not inactive). Then when the cells are briefly hyperpolarized M-channels slowly close, producing a slow inward tail-current (see Fig. 1a). The amount of M-current lost during the hyperpolarization can be measured from the initial amplitude of this deactivation tail-current at the point of hyperpolarization. This procedure isolates M-current from other membrane currents (Adams et al. 1982a).

In neurons that had not been transformed to express z1a adrenoceptors, the z1-agonist phenylephrine (10 μM) had no effect on either the amount of steady outward current or the amplitude of the deactivation tail-current (Fig. 1a, top trace; Fig. 1c). In contrast, in the same neuron the muscarinic acetylcholine-receptor agonist oxotremorine-M (oxo-M, 10 μM) clearly reduced both steady outward current and deactivation current-tail (Fig. 1a, lower trace), signaling M-current inhibition (see Adams et al. 1982b). Mean inhibition in three such neurons measured from the extrapolated initial amplitude of the deactivation tail-current (Adams et al. 1982a) was 59 ± 14 % (Fig. 1c).

In contrast to the negative effect of phenylephrine in Fig. 1a, this z1-agonist clearly did reduce the M-current if a neuron had been pre-injected with z1a receptor cDNA (Fig. 1b), to a mean extent of 72 ± 11 % (n = 4; Fig. 1c). Thus, in an z1a-expressing neuron, phenylephrine inhibits the M-current just like a muscarinic agonist.

Excitability

M-current confers strong spike-frequency adaptation on these neurons, so one effect of M-current inhibition is to facilitate repetitive firing during sustained depolarization (Brown 1983). Figure 2a shows such an effect of oxotremorine-M. A 2 s depolarizing current injection initially generated only two action potentials at the beginning of the pulse but a sustained train of action potentials after adding oxotremorine-M, rising to 50 action potentials (25 Hz) with increasing current injections (Fig. 2c). Phenylephrine had no effect on induced action potential firing in normal cells but precisely replicated the effect of oxotremorine-M in a neuron pre-injected with the z1a cDNA (Fig. 2b, c). Thus, M-current inhibition by z1a-adrenoceptors would be expected to increase neuronal excitability.

Discussion

These experiments show that, when present in a neuron, z1-adrenoceptors are well capable of strongly inhibiting the M-current and greatly increasing neuronal excitability,
just like an endogenous Gq-coupled GPCR such as the muscarinic M1-receptor.

It could be argued that this is an artifact of receptor overexpression, and that any endogenous α1-receptors are somehow insulated from exerting such an effect. For example, muscarinic receptor overexpression in these neurons can overcome endogenous barriers that otherwise restrict downstream Ca$^{2+}$-signaling pathways (Zaika et al. 2011). However, we think this type of exclusion zone unlikely to confer resistance of M-channels to inhibition, since, in previous tests, we found that overexpression of the small number of endogenous P2Y1 purinoceptors, while amplifying the signals, did not qualitatively alter their actions on M-channels and Ca$^{2+}$-channels (Filippov et al. 2010). Further, endogenous α1-adrenoceptors have been reported to inhibit M-currents in other peripheral neurons (Shibata and Taketani 2001).

In previous experiments on central neurons (see “Introduction”), the depolarization produced by α1-receptor activation was usually accompanied by a reduced K$^+$ conductance but in only one case (cultured embryonic spinal neurons: Legendre et al. 1988) did the depolarization show some, though not all, of the properties expected for M-current inhibition. Probably M-currents were insufficiently prominent in the other cells tested. Notwithstanding, bearing in mind the wide distribution of both M-channels and α1 receptors in the brain, it seems likely that more of the α1-mediated effects of norepinephrine on central neurons will prove to be caused by M-channel inhibition when examined in detail. There are only a few thousand noradrenergic neurons in their main site of origin in the locus coeruleus, but their axons ramify widely throughout the c.n.s. (Berridge and Waterhouse 2003), and their activation of α1-receptors are thought to be involved in many aspects of central noradrenergic function, including memory and cognition (Gibbs and Summers 2002; Ramos and Arnsten 2007), sleep (Berridge et al. 2012), depression (Stone et al. 2003), and pain (Pertovaara 2006). Hence, further test for M-current inhibition would seem worthwhile. From the data of Legendre et al. (1988), a plausible start-point might be the spinal cord.

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Fig. 2 Effects of a oxotremorine-M (oxo-M, 10 μM) and b phenylephrine (Phe, 10 μM) on action potential firing in two neurons induced by 2 s depolarizing current injections (120 pA in a, 160 pA in b). Neuron A: wild-type; neuron B: pre-injected with α1a cDNA.

c Plots of the number of action potentials (“spikes”) recorded in 2 s (ordinates) against the amplitude of the depolarizing current injection (abscissae) for the two cells illustrated in a and b. Discharges reverted to control after drug washout.
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References

Adams PR, Brown DA, Constanti A (1982a) M-currents and other potassium currents in bullfrog sympathetic neurones. J Physiol 330:537–572

Adams PR, Brown DA, Constanti A (1982b) Pharmacological inhibition of the M-current. J Physiol 332:223–262

Berridge CW, Waterhouse BD (2003) The locus coeruleus–noradrenergic system: modulation of behavioral state and state-dependent cognitive processes. Brain Res Brain Res Rev 42:33–84

Berridge CW, Schmeichel BE, España RA (2012) Noradrenergic modulation of wakefulness/arousal. Sleep Med Rev 16:187–197

Brown DA (1983) Slow cholinergic excitation—a mechanism for increasing neuronal excitability. Trends Neurosci 6:302–306

Constanti A, Brown DA (1981) M-currents in voltage-clamped mammalian sympathetic neurones. Neurosci Lett 24:289–294

Dawson LF, Phillips JK, Finch PM, Inglis JJ, Drummond PD (2010) Expression of α1-adrenoceptors on peripheral nociceptive neurons. Neuroscience 175:300–314

Day HE, Campeau S, Watson SJ Jr, Akil H (1997) Distribution of alpha 1a-, alpha 1b- and alpha 1d-adrenergic receptor mRNA in the rat brain and spinal cord. J Chem Neuroanat 13:115–139

Delmas P, Brown DA (2005) Pathways modulating neutral KCNQ/M (Kv7) potassium channels. Nat Rev Neurosci 6:850–862

Filippov AK, Webb TE, Barnard EA, Brown DA (1998) P2Y2 nucleotide receptors expressed heterologously in sympathetic neurons inhibit both N-type Ca2+ and M-type K+ currents. J Neurosci 18:5170–5179

Filippov AK, Brown DA, Barnard EA (2000) The P2Y(1) receptor closes the N-type Ca(2+) channel in neurones, with both adenosine triphosphates and diphosphates as potent agonists. Br J Pharmacol 129:1063–1066

Filippov AK, Simon J, Barnard EA, Brown DA (2010) The scaffold protein NHERF2 determines the coupling of P2Y1 nucleotide and mGluR5 glutamate receptor to different ion channels in neurons. J Neurosci 30:11068–11072

Gibbs ME, Summers RJ (2002) Role of adrenoceptor subtypes in memory consolidation. Prog Neurobiol 67:345–391

Grayson TH, Ellis JM, Chen S, Graham RM, Brown RD, Hill CE (1998) Immunohistochemical localisation of alpha1B-adrenergic receptors in the rat iris. Cell Tissue Res 293:435–444

Hawrylyshyn KA, Michelotti GA, Cogé F, Guénin SP, Schwinn DA (2004) Update on human alpha1-adrenoceptor subtype signaling and genomic organization. Trends Pharmacol Sci 25:449–455

Ikeda SR (1997) Heterologous expression of receptors and signaling proteins in adult mammalian sympathetic neurons by microinjection. Methods Mol Biol 83:191–202

Jones LS, Gauger LL, Davis JN (1985) Anatomy of brain alpha 1-adrenergic receptors: in vitro autoradiography with [125I]-hep. J Comp Neurol 231:190–208

Legendre P, Dupouy B, Vincent JD (1988) Excitatory effect of noradrenaline on pacemaker cells in spinal cord primary cultures. Neuroscience 24:647–658

McCormick DA (1992) Cellular mechanisms underlying cholinergic and noradrenergic modulation of neuronal firing mode in the cat and guinea pig dorsal lateral geniculate nucleus. J Neurosci 12:278–289

McCormick DA, Pape HC, Williamson A (1991) Actions of norepinephrine in the cerebral cortex and thalamus: implications for function of the central noradrenergic system. Prog Brain Res 88:293–305

Ogata N, Matsuo T (1986) The effects of catecholamines on electrical activity of neurons in the guinea pig supraoptic nucleus in vitro. Brain Res 385:122–135

Pediani JD, MacKenzie JF, Heeley RP, Daly CJ, McGrath JC (2000) Single-cell recombinant pharmacology: bovine alpha(1a)-adrenoceptors in rat-1 fibroblasts release intracellular Ca(2+), display subtype-characteristic agonism and antagonism, and exhibit an antagonist-reversible inverse concentration-response phase. J Pharmacol Exp Ther 293:887–895

Pertovaara A (2006) Noradrenergic pain modulation. Prog Neurobiol 80:53–83

Ramos BP, Arnsten AF (2007) Adrenergic pharmacology and cognition: focus on the prefrontal cortex. Pharmacol Ther 113:523–536

Shibata K, Taketani K (2001) Excitatory effect of noradrenaline on rat airway parasympathetic ganglion neurons. Fukuoka Igaku Zasshi 92:377–383 (In Japanese.)

Stone EA, Lin Y, Rosengarten H, Kramer HK, Quatermain D (2003) Emerging evidence for a central epinephrine-inervated alpha 1-adrenergic system that regulates behavioral activation and is impaired in depression. Neuropsychopharmacology 28:1387–1399

Zaika O, Zhang J, Shapiro MS (2011) Combined phosphoinositide and Ca2+ signals mediating receptor specificity toward neuronal Ca2+ channels. J Biol Chem 286:830–841