

NIH Public Access

Author Manuscript

FEBS Lett. Author manuscript; available in PMC 2012 October 11.

Published in final edited form as: *FEBS Lett.* 1990 February 12; 261(1): 67–70.

Characterization of the locomotor depression produced by an A₂-selective adenosine agonist

Olga Nikodijević, John W. Daly, and Kenneth A. Jacobson

Laboratories of Chemistry and Bioorganic Chemistry, National Inst. of Diabetes, Digestive and Kidney Diseases, National Institutes of Health, Bethesda, MD 20892, USA

Abstract

Adenosine analogs, such as N^6 -cyclohexyladenosine (CHA) that are selective for A₁-adenosine receptors, and analogs, such as 5'-N-ethylcarboxamidoadenosine (NECA) that are active at both A₁ and A₂ receptors, cause a profound depression of locomotor activity in mice via a central mechanism. The depression is effectively reversed by non-selective adenosine antagonists such as theophylline. We report that 2 - [(2-aminoethyl-amino)carbonylethylphenylethylamino] - 5' - Nethylcarboxamidoadenosine (APEC), an amine derivative of the A₂-selective agonist, CGS21680, is a potent locomotor depressant in mice. The in vivo pharmacology is consistent with A₂selectivity at a central site of action. Two parameters indicative of locomotor activity, horizontal activity and total distance travelled, were measured using a computerized activity monitor. From dose-response curves it was found that APEC (ED₅₀ 16 μ g/kg) is more potent than CHA (ED₅₀ 60 μ g/kg) and less potent than NECA (ED₅₀ 2 μ g/kg). The locomotor depression by APEC was reversible by theophylline, but not by the A₁-selective antagonists 8-cyclopentyltheophylline (CPT) and 8-cyclopentyl-1,3-dipropyl-2-thioxanthine, nor by the peripheral antagonists 8-psulfophenyltheophylline (8-PST) and 1,3-dipropyl-8-p-sulfophenylxanthine. The locomotor activity depression elicited by NECA and CHA was reversed by A1-selective antagonists. These results suggest that the effects of APEC are due to stimulation of A₂ adenosine receptors in the brain.

Keywords

Adenosine analog; Locomotor depression; Adenosine receptor

1. INTRODUCTION

Adenosine, as a neuromodulator, inhibits the firing of neurons and the release of neurotransmitters in the central nervous system [1]. In behavioral models, adenosine agonists, acting via a central mechanism, cause a dramatic depression of locomotor activity, which is effectively reversed by the non-selective adenosine antagonists, caffeine and theophylline [2]. This effect has been demonstrated in rodents using a number of potent adenosine analogs, including the non-selective agonist, *N*-ethylcarboxamidoadenosine (NECA), and the A₁ selective agonists, N^6 -cyclohexyl-and N^6 -cyclopentyladenosine [3–5]. The potencies in producing hypomobility, as measured by head dipping and locomotor assays, of a series of adenosine agonists were recently found to correlate to the potencies of

^{© 1990} Federation of European Biochemical Societies

Correspondence address: O. Nikodijevi, Laboratories of Chemistry and Bioorganic Chemistry, National Inst. of Diabetes, Digestive and Kidney Diseases, National Institutes of Health, Bethesda, MD 20892, USA.

the analogs at A_2 receptors [6], suggesting that primarily A_2 receptors are involved in these effects. The lack of a truly A_2 -selective agonist has hampered these studies.

Recently, several classes of A₂-selective adenosine agonists have been reported. N^{6} -[2-(3,5-dimethoxyphenyl)-2-(2-methylphenyl)ethyl] adenosine [7] and 2-(carboxyethylphenylethylamino)adenosine-5'-carboxamide (CGS21680) [8,9] are A₂-selective in competitive binding experiments at central A₁- (measured in cortex) and A₂- (measured in striatum) adenosine receptors by factors of 32 and 140, respectively. CGS21680 was also shown to be A₂-selective in the cardiovascular system [9]. CGS21680 contains a carboxylic acid functionality, which is expected to limit its passage across the blood/brain barrier [9]. Using a functionalized congener approach, a series of long-chain derivatives of CGS21680 that retain A₂ potency and selectivity and do not contain the carboxylic functionality, was synthesized [10]. An amine derivative, 2-[(2 -aminoethyl-amino)carbonylethylphenylethylamino]-5'-carboxamidoadenosine (APEC; 1), served as a synthetic intermediate for molecular probes for A₂-adenosine receptors, including the first photo-affinity ligand, ¹²⁵I-PAPA-APEC [11]. We report that APEC, which is 17-fold A₂-selective in vitro [10], is a potent locomotor depressant in mice. The in vivo pharmacology is consistent with A₂-selectivity at a central site of action.



2. MATERIALS AND METHODS

2.1. Chemicals

NECA, CHA, 8-PST, CPT, DPSPX and XAC were obtained from Research Biochemicals, Inc. (Natick, MA). 2-Thio-CPX [13] was the generous gift of Professor W. Pfleiderer (Univ. of Konstanz, FRG and Dr J. Neumeyer, RBI). CGS 21680C (Na salt) was the generous gift of Dr A. Hutchison (CIBA-Geigy Corp.). APEC was synthesized as described [10].

2.2. Animal studies

2.2.1. Subjects—Adult male mice of the NIH (Swiss) strain weighing 25–30 g were housed in groups of 12 animals per cage with a light-dark cycle of 12:12 h. The animals were given free access to standard pellet food and water and were habituated for 24 h in laboratory conditions prior to testing. Each animal was used only once in the activity monitor.

2.2.2. Locomotor activity—Individual animals were studied in a Digiscan activity monitor (Omnitech Electronics Inc., Columbus, OH) equipped with an IBM-compatible computer. Data was collected in the morning, for 3 consecutive intervals of 10 min each and analyzed as a group for 30 min sampling period. Two non-equivalent parameters [15] were analyzed: (i) horizontal activity, which represents the total number of beam interruptions in the horizontal direction; and (ii) total distance travelled, which indicates the distance in cm travelled by the animal. The latter is dependent on the path taken.

2.2.3. Drug administration—All drugs were dissolved in a 1:4 v/v mixture of Emulphor EL-620 (GAF Chemicals Corp., Wayne, NJ) and phosphate-buffered saline and administered i.p. in a vol. of 5 ml/kg b. wt. Warming and sonication aided in dissolving the drugs. When appropriate, an adenosine antagonist was injected first followed by an agonist

FEBS Lett. Author manuscript; available in PMC 2012 October 11.

after 10 min. Immediately after the final injection, the mouse was placed in the activity monitor cage, and data collection was begun after a delay of 10 min. Statistical analysis was performed using the Student's *t*-test. Each value reported represents the mean \pm SE for 6–10 animals, except for the control points (vehicle injected) for which n = 22.

3. RESULTS

The locomotor effects at different doses of APEC and the adenosine agonists NECA and CHA, administered intraperitoneally in mice, were measured. The dose-response curves are given in fig. 1. APEC was found to have an ED_{50} value for horizontal activity of 14 μ g/kg b. wt. Thus, APEC is more potent than CHA ($ED_{50} = 70 \ \mu$ g/kg) and less potent than NECA ($ED_{50} = 2 \ \mu$ g/kg). CGS21680 was also tested as a locomotor depressant at several doses. CGS21680 at a dose of 16 μ g/kg^a was nearly inactive with 3 ± 0.2% and 13 ± 1% depression of horizontal activity (h.a.) and total distance travelled (t.d.), respectively. At a dose of 1 μ mol/kg, CGS21680 caused decreases of 64 ± 4.5% (h.a.) and 62 ± 5% (t.d.) in locomotor activity, and at 3 μ mol/kg the locomotor depression was 94 ± 9% (h.a.) and 96 ± 20% (t.d.).

The locomotor depressant activity of APEC was not reversed by the peripheral adenosine antagonist, 8-p-sulfophenyltheophylline (8-PST; fig. 2). This is consistent with a central mechanism for the locomotor depression by APEC. Similarly, the more potent 1,3dipropyl-8-(p-sulfophenyl)xanthine (DPSPX) at 5 mg/kg did not antagonize APEC (fig. 2). Since 8-PST and DPSPX are relatively non-selective, a peripheral action at either A_1 or A_2 subtypes is precluded as the mechanism for the locomotor depression by APEC. Curiously, at high doses, these peripheral antagonists both elicited some locomotor depression. This depression is particularly evident in the effect of 10 mg/kg 8-PST on total distance travelled $(24 \pm 2\%$ decrease). At 5 mg/kg the depressant effect on total distance travelled was 6 ± 0.5% and 12 \pm 1.8% for 8-PST and DPSPX, respectively. The A₁-selective antagonist, 2thioCPX (see below), depressed locomotor activity only at a dose of 10 mg/kg, with decreases of $23 \pm 3\%$ (h.a.) and $28 \pm 3.6\%$ (t.d.). The mechanisms underlying such depressant effects are unclear. At a dose of 10 mg/kg 8-(p-sulfophenyl)caffeine, which is structurally related to 8-PST but inactive or weakly active, respectively, as an adenosine antagonist at A₁ and A₂ receptors [16], stimulated locomotor activity slightly by $6 \pm 0.2\%$ (h.a.) and $12 \pm 1.8\%$ (t.d.).

The A₁-selective antagonist, 8-cyclopentyltheophylline (CPT), has been reported to antagonize the central depressant activities of adenosine agonists, such as N^{6_-} cyclopentyladenosine [4]. Similarly, we found that CPT could reverse the depression by an ED₅₀ dose of CHA (fig. 3), an agonist that is A₁-selective by a factor of 390 [12]. The depression evoked by an ED₅₀ dose of NECA, an agonist that has marked activity at both adenosine receptor subtypes, was also completely reversed by this A₁-selective antagonist (fig. 4). However, the locomotor depression evoked by APEC was not reversible by a comparable dose of CPT (fig. 4). This suggests that the depressant effect of APEC is due to stimulation of A₂ adenosine receptors in the brain. The depressant effects of APEC were reversed by the non-selective antagonist theophylline. At the ED₅₀ dose of APEC, 10 mg/kg theophylline restore the total distance travelled to 95 ± 12% of control.

In contrast to PST, DPSPX and 2-thio-CPX at high doses, CPT alone at 10 mg/kg was found to be a weak central stimulant (fig. 3) causing increases of $41 \pm 8\%$ and $24 \pm 7\%$ for total distance travelled and horizontal activity, respectively. This finding is in contrast to a

^aThe dose equal to the average of ED₅₀ values for APEC for horizontal activity and total distance (corresponds to 0.03 μ mol/kg APEC).

previous report [4] in which no stimulation by CPT (up to 30 mg/kg) was seen. To further study the effects of A₁-selective adenosine antagonists on the locomotor depression by APEC, we searched for another centrally active A₁-selective xanthine. The potent ($K_i = 0.66$ nM) and 480-fold A₁-selective antagonist, 8-cyclopentyl-1,3-dipropyl-2-thioxanthine (2thio-CPX) [13,14], was shown to reverse the locomotor depression elicited by CHA (data not shown). At a dose of 5 mg/kg, 2-thio-CPX alone, unlike CPT, did not appreciably stimulate locomotor activity (increases of $1 \pm 0.1\%$ in both h.a. and t.d.). Like CPT, this xanthine failed to antagonize the locomotor effects of APEC, further supporting the conclusion of in vivo A₂ selectivity of APEC. A combination of 5 mg/kg 2-thio-CPX and the ED₅₀ dose of APEC depressed locomotor activity by 50 ± 10% (h.a.) and 55 ± 13% (t.d.).

4. DISCUSSION

The results show that APEC, previously determined to be A2-selective in binding assays at rat brain adenosine receptors [10], is a potent locomotor depressant in mice. The doseresponse curves and the ED_{50} values for the effects of adenosine agonists on two parameters indicative of locomotor activity (horizontal activity and total distance travelled; fig. 1) show an order of potency of NECA > APEC > CHA. CHA (K_i 5 = 10 nM) is also less potent than NECA and APEC ($K_i = 10.3$ and 5.73 nM, respectively) in competitive binding experiments at A₂ adenosine receptors [10,12]. The locomotor effects of APEC were not reversed by the A1-selective adenosine antagonists, CPT and 2-thio-CPX. Both of these xanthines are centrally active antagonists and both reverse the locomotor depression elicited by the A_1 agonist, CHA. Moreover, the effect of the non-selective antagonist, theophylline, and the inactivity of peripheral, non-selective antagonists, PST and DPSPX, in reversing locomotor depression elicited by APEC suggests central action at adenosine receptors. The lack of antagonism of APEC-elicited depression by the A1-selective xanthines, CPT and 2-thio-CPX, strongly suggests that activation of A₂ receptors are involved in the behavioral depression. Prior studies have suggested that activation of either A_1 or A_2 receptors can elicit dramatic depressant effects [2-6]. APEC now provides an important tool for investigation of the role of central A2 receptors, just as the A1-selective agonists, CHA and CPA, provide tools for investigation of the role of central A₁ receptors.

CGS21680, the structural precursor of APEC, has been evaluated as a potentially therapeutic, hypotensive agent [9]. The charged carboxylate group is predicted to tend to restrict the action of this potent and highly selective A_2 adenosine agonist ($K_i = 14 \text{ nM}$) to the periphery [8]. It is possible that the reason for the behavioral inactivity of CGS21680 at a low dose, at which APEC is active, is due to diminished passage across the blood/brain barrier. At higher doses, CGS21680 acts as a locomotor activity depressant, but less potent than CHA, The aliphatic amino group of APEC is predominantly but not completely charged at physiological pH, which obviously does not prevent its entry into the CNS.

Yet to be resolved is why NECA is much more potent than APEC despite similar affinity at A_2 -receptors. In addition to pharmacokinetic factors, there remains the possibility that dual activation of A_1 and A_2 receptors by NECA is acting synergistically on locomotor depression. The blockade by CPT of NECA-elicited behavioral depression [4] suggests a key role for A_1 receptors, yet other data [3] suggests that A_2 receptors are involved. Preliminary results (unpublished) suggest that the effects of APEC and CHA are more than additive, supporting possible synergistic interactions of A_1 and A_2 receptors in eliciting behavioral depression.

References

- 1. Snyder SH. Annu Rev Neurosci. 1985; 8:103-124. [PubMed: 2858998]
- Snyder SH, Katims JJ, Annau Z, Bruns RF, Daly JW. Proc Natl Acad Sci USA. 1981; 78:3260– 3264. [PubMed: 6265942]
- 3. Seale T, Abla KA, Shamim MT, Carney JM, Daly JW. Life Sci. 1988; 43:1671–1684. [PubMed: 3193854]
- Bruns, RF.; Davis, RE.; Ninteman, FW.; Poschel, BPH.; Wiley, JN.; Heffner, TG. Physiology and Pharmacology of Adenosine and Adenine Nucleotides. Paton, DM., editor. Taylor and Francis; London: 1988. p. 39-49.
- Heffner TG, Wiley JN, Williams AE, Bruns RF, Coughenour LL, Downs DA. Psychopharmacology. 1989; 98:31–37. [PubMed: 2498959]
- 6. Durcan MJ, Morgan PF. Eur J Pharmacol. 1989; 168:285-290. [PubMed: 2583238]
- Bridges AJ, Bruns RF, Ortwine DF, Priebe SR, Szotek DL, Trivedi BK. J Med Chem. 1988; 31:1282–1285. [PubMed: 3385722]
- Hutchison AJ, Williams M, DeJesus R, Oei HH, Ghai GR, Webb RL, Zoganas HC, Stone GA, Jarvis MF. J Med Chem. 1989 in press.
- 9. Hutchison AJ, Webb RL, Oei HH, Ghai GR, Williams M. J Pharm Exp Ther. 1989; 251:47-55.
- Jacobson KA, Barrington WW, Pannell LK, Jarvis MF, Ji X-D, Hutchison AJ, Stiles GL. J Mol Recognition. 1990 in press.
- Barrington WW, Jacobson KA, Williams M, Hutchison AJ, Stiles GL. Proc Natl Acad Sci USA. 1989; 86:6572–6576. [PubMed: 2771944]
- 12. Bruns RF, Lu GH, Pugsley TA. Mol Pharmacol. 1986; 29:331–346. [PubMed: 3010074]
- Jacobson KA, Kiriasis L, Barone S, Bradbury BJ, Kammula U, Campagne JM, Secunda S, Daly JW, Pfleiderer W. J Med Chem. 1989; 32:1873–1879. [PubMed: 2754711]
- Neumeyer, JL.; De la Cruz, D.; Kiriasis, L.; Barone, S.; Bradbury, BJ.; Kammula, U.; Campagne, JM.; Secunda, S.; Daly, JW.; Pfleiderer, W.; Jacobson, KA. Abstract B-16, Purine Nucleosides and Nucleotides in Cell Signalling: Targets for New Drugs (meeting); Sept. 1989; 1989.
- Sandberg PR, Hagenmeyer SH, Henault MA. Neurobehav Toxicol Teratol. 1985; 7:87–94. [PubMed: 3839052]
- Shamim MT, Ukena D, Padgett WL, Daly JW. J Med Chem. 1989; 32:1231–1237. [PubMed: 2724296]



Fig. 1.

Dose-response curves for locomotor depression in mice by the adenosine agonists, NECA (squares), APEC (triangles) and CHA (circles). For each analog, percent decrease compared to vehicle control is shown for horizontal activity (open symbols) and total distance (closed symbols).





The effects of the peripheral adenosine antagonists, 8-PST and DPSPX, on locomotor depression induced by APEC (16 μ g/kg). Changes in horizontal activity (solid bars) and total distance travelled (hatched bars), relative to vehicle control, are shown.

Nikodijevi et al.



Fig. 3.

The effects of the A₁-selective adenosine antagonist, CPT, on locomotor depression induced by CHA (60 μ g/kg). Percent changes in total distance travelled (A) and horizontal activity (B), relative to vehicle control, are shown.

Nikodijevi et al.



Fig. 4.

The effects of an A₁-selective antagonist on locomotor depression induced by adenosine 5'carboxamide analogs. The percent change in locomotor activity, relative to vehicle control, induced by NECA (2 μ g/kg) and APEC (16 μ g/kg) at the ED₅₀ doses, in the presence (hatched bars) and absence (solid bars) of 10 mg/kg CPT, is shown.