

RESEARCH PAPER

Acute dilation to α_2 -adrenoceptor antagonists uncovers dual constriction and dilation mediated by arterial α_2 -adrenoceptors

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Background and purpose: In mouse tail arteries, selective α_2 -adrenoceptor antagonism with rauwolscline caused powerful dilation during constriction to the α_1 -adrenoceptor agonist phenylephrine. This study therefore assessed phenylephrine's selectivity at vascular α -adrenoceptors and the mechanism(s) underlying dilation to rauwolscline.

Experimental approach: Mouse isolated tail arteries were assessed using a pressure myograph.

Key results: The α_2 -adrenoceptor agonist UK14,304 caused low-maximum constriction that was inhibited by rauwolscline (3×10^{-8} M) but not by the selective α_1 -adrenoceptor antagonist prazosin (10^{-7} M). Concentration–effect curves to phenylephrine, cirazoline or noradrenaline were unaffected by rauwolscline but were inhibited by prazosin, which was more effective at high compared with low levels of constriction. In the presence of prazosin, rauwolscline inhibited the curves and was more effective at low compared with high levels of constriction. Although rauwolscline alone did not affect concentration–effect curves to phenylephrine, noradrenaline or cirazoline, it caused marked transient dilation when administered during constriction to these agonists. Dilation was mimicked by another α_2 -adrenoceptor antagonist (RX821002, 3×10^{-8} M), was dependent on agonist selectivity, and did not occur during adrenoceptor-independent constriction (U46619). During constriction to UK14,304 plus U46619, rauwolscline or rapid removal of UK14,304 caused transient dilation that virtually abolished the combined constriction. Endothelial denudation reduced these dilator responses.

Conclusions and implications: Inhibition of α_2 -adrenoceptors caused transient dilation that was substantially greater than the contribution of α_2 -adrenoceptors to the constriction. This reflects a slowly reversing α_2 -adrenoceptor-mediated endothelium-dependent dilation and provides a rapid, sensitive test of α_2 -adrenoceptor activity. This approach also clearly emphasizes the poor selectivity of phenylephrine at vascular α -adrenoceptors.

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Introduction

α_1 - and α_2 -adrenoceptors (nomenclature follows Alexander *et al.*, 2008) are expressed on vascular smooth muscle cells and initiate constriction in response to the endogenous catecholamines, adrenaline and noradrenaline (Flavahan and McGrath, 1980; Flavahan *et al.*, 1984; Guimaraes and Moura, 2001; Philipp *et al.*, 2002). Although α_1 -adrenoceptors are expressed by most blood vessels, functional constrictor α_2 -adrenoceptors have a more restricted distribution in the vascular system. Because of their role in mediating cold-induced cutaneous vasoconstriction, α_2 -adrenoceptors are more active in cutaneous compared with deep blood vessels

(Flavahan *et al.*, 1985; Flavahan, 2005). Furthermore, in contrast to α_1 -adrenoceptor activity, α_2 -adrenoceptor-mediated constriction increases in distal compared with proximal arteries; a pattern that continues in the venous system where α_2 -adrenoceptor activity is widely distributed (Flavahan *et al.*, 1984; 1987; Flavahan and Vanhoutte, 1986b; Rajanayagam and Medgett, 1987; Guimaraes and Moura, 2001). However, even when the α_2 -adrenoceptor constrictor system is at its most powerful, it generally provides a lower maximum response compared with other receptor systems, including α_1 -adrenoceptors (Flavahan and McGrath, 1984; Flavahan *et al.*, 1984; Chotani *et al.*, 2000).

Phenylephrine is a commonly used nasal decongestant (Eccles, 2007; Kollar *et al.*, 2007). It is generally considered to be a selective α_1 -adrenoceptor agonist and is widely used in animal and clinical studies to assess the functional activity of α_1 -adrenoceptors (Inoue *et al.*, 2001; Arain *et al.*, 2002; Mueed *et al.*, 2004; Lewis *et al.*, 2007; Kirby *et al.*, 2008). While

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studying the agonist in mouse isolated tail arteries, we observed that selective α_2 -adrenoceptor antagonism with rauwolsine (3×10^{-8} M) caused powerful transient dilation of phenylephrine-induced constriction, suggesting that the agonist might cause constriction by activating α_1 - and α_2 -adrenoceptors. The present study was therefore performed to characterize the selectivity of phenylephrine at mouse vascular α_1 - and α_2 -adrenoceptors, and to investigate the mechanism underlying the relaxation to rauwolsine (using mouse isolated tail and mesenteric arteries). The results demonstrate that phenylephrine has limited selectivity at α -adrenoceptor subtypes and does indeed initiate vasoconstriction by activating α_1 - and α_2 -adrenoceptors. However, the transient dilator response evoked by α_2 -adrenoceptor antagonists is much greater than the contribution of α_2 -adrenoceptors to the constrictor response. This reflects the presence of a slowly reversing α_2 -adrenoceptor-mediated endothelium-dependent dilation, which transiently amplifies the dilator response to α_2 -adrenoceptor antagonism. Indeed, this provides a rapid and sensitive method of assessing agonist selectivity at vascular α -adrenoceptors.

Methods

Animals

All animal care and experimental procedures were approved by the Johns Hopkins University Institutional Animal Care and Use Committee. We used (C57BL6) mice from The Jackson Laboratory, Bar Harbor, ME, USA.

Blood vessel chamber. Eight- to fourteen-week-old male mice were killed by CO₂ asphyxiation. Segments of mesenteric and proximal tail arteries were then rapidly and carefully isolated and placed in cold Krebs-Ringer-bicarbonate solution: 118.3 mM NaCl, 4.7 mM KCl, 1.2 mM MgSO₄, 1.2 mM KH₂PO₄, 2.5 mM CaCl₂, 25.0 mM NaHCO₃ and 11.1 mM glucose (control solution). Arteries were cannulated at both ends with glass micropipettes, secured using 12-0 nylon monofilament suture, and placed in a microvascular chamber (Living Systems, Burlington, VT, USA) (Flavahan, 2005). They were studied in the absence of flow and maintained at a constant transmural pressure (P_{TM}) of 60 mmHg (Flavahan, 2005). Under these conditions, the internal diameter of the arteries was approximately 300 μ m. The chamber was placed on the stage of an inverted microscope (Nikon TMS-F; Nikon, Tokyo, Japan) connected to a video camera, and superfused with control solution (maintained at 37°C, gassed with 16% O₂, 5% CO₂, balance N₂; pH 7.4). In general, as performed in our previous studies, the control solution filling the chamber and superfusing the arterial segments was re-circulated in a continuous fashion (total volume 90 mL, flow rate 25 mL·min⁻¹). However, in one series of experiments, continuous perfusion of the chamber was achieved in an open, non-re-circulating, manner (rate 25 mL·min⁻¹). The blood vessel image was projected onto a video monitor and internal diameters continuously monitored by a video dimension analyser (Living Systems) and BIOPAC data acquisition system (Santa Barbara, CA, USA).

Experimental protocols

Arterial segments were allowed to equilibrate for at least 30 min before starting experiments. Concentration–effect curves to the adrenergic agonists phenylephrine (10^{-8} to 3×10^{-5} M), cirazoline (10^{-9} to 10^{-6} M), UK14,304 [5-bromo-N-(4,5-dihydro-1H-imidazol-2-yl)-6-quinoxalinamine] (10^{-9} to 10^{-7} M), or noradrenaline (10^{-8} to 10^{-5} M), or to the thromboxane receptor agonist U46619 (10^{-9} to 10^{-7} M) were generated by increasing the concentration of the agonists in half-log increments, once the constriction to the previous concentration had peaked. Following completion of the concentration–effect curve, the response to the agonists was terminated by repeatedly exchanging the buffer solution and allowing the blood vessels to return to their stable baseline levels. In some experiments, concentration–effect curves to agonists were determined under control conditions or in the presence of the selective α_2 -adrenoceptor antagonists, rauwolsine (3×10^{-8} M) or RX821002 (3×10^{-8} M) (Flavahan *et al.*, 1984; Devedjian *et al.*, 1994), and/or the selective α_1 -adrenoceptor antagonist, prazosin (10^{-7} M) (Flavahan *et al.*, 1984). At the concentrations used, these antagonists are potent and selective antagonists at their respective receptors. Antagonists were present 30 min before and during the concentration–effect curves to the agonists. In some experiments, arteries were stably constricted with phenylephrine (3×10^{-7} M), cirazoline (3×10^{-8} M), or noradrenaline (3×10^{-7} M), or to a combination of U46619 (to ~60% of baseline diameter, 10^{-8} to 3×10^{-8} M) plus UK14,304 (3×10^{-8} M), and then vasodilator responses to rauwolsine (3×10^{-8} M) or RX821002 (3×10^{-8} M) determined. Time control studies were performed using paired arterial segments that were not exposed to the receptor antagonists. In some experiments, endothelium denudation was performed by inserting a 70 μ m wire into the lumen of the artery and was confirmed by loss of the dilator response to acetylcholine (10^{-6} M, assessed during constriction to U46619 to approximately 60% of baseline diameter).

Data analysis and statistics

Vasoconstriction was expressed as a percentage of the internal diameter of the blood vessel prior to administering the agonist(s). α_2 -Adrenoceptor constriction is generally a low-maximum response compared with other constrictor systems (including α_1 -adrenoceptors), and generates only ~30% constriction of mouse tail arteries. This difference in maximal response can facilitate identification of α -adrenoceptor subtypes activated by non-selective agonists. This is achieved by analysing the effect of antagonists on agonist concentration–effect curves at response levels attainable by α_2 -adrenoceptor activation and at levels attainable only by α_1 -adrenoceptor activation (Flavahan and McGrath, 1984; Flavahan *et al.*, 1984; 1987; Flavahan and Vanhoutte, 1987). Therefore, the effect of α -adrenoceptor antagonists was analysed at agonist concentrations causing 15% and 60% constriction (CC₁₅ and CC₆₀ respectively). The 15% level of constriction (i.e. to 85% of baseline diameter) was selected because it approximates 50% of the maximal response to α_2 -adrenoceptor stimulation in tail arteries whereas the 60% level of constriction (i.e. to 40% of baseline diameter) exceeds the contractile activity of

α_2 -adrenoceptors (Flavahan, 2005). Statistical evaluation of the data was performed by Student's paired *t*-test. When more than two means were compared, analysis of variance was used. If a significant *F* value was found, Tukey's test for multiple comparisons was employed to identify differences among groups. Data are expressed as means $\pm$ SEM for *n* number of experiments, where *n* equals the number of animals from which blood vessels were studied. Values were considered to be statistically different when *P* < 0.05.

Materials

All drugs were obtained from Sigma-Aldrich (St Louis, MO, USA) and were dissolved in distilled water with the exception of UK14,304, which was dissolved in dimethyl sulphoxide (highest chamber concentration of 0.001%). Drug concentrations are described as final molar concentration (M) in the chamber superfusate.

Results

Characterization of α -adrenoceptors involved in constriction to phenylephrine, noradrenaline or cirazoline

The selective α_1 -adrenoceptor antagonist prazosin (10^{-7} M) did not affect constrictor responses to the prototypic α_2 -adrenoceptor agonist UK14,304 (10^{-9} to 10^{-7} M) confirming prazosin's selectivity for α_1 -adrenoceptors in the mouse tail artery (Figure 1). In contrast, the selective α_2 -adrenoceptor antagonist rauwolscine (3×10^{-8} M) markedly depressed responses to UK14,304 (Figure 1).

Rauwolscine (3×10^{-8} M) did not significantly affect the concentration–effect curve to phenylephrine (Figure 2). Prazosin (10^{-7} M) caused a non-parallel shift in the concentration–effect curve to phenylephrine, having a greater

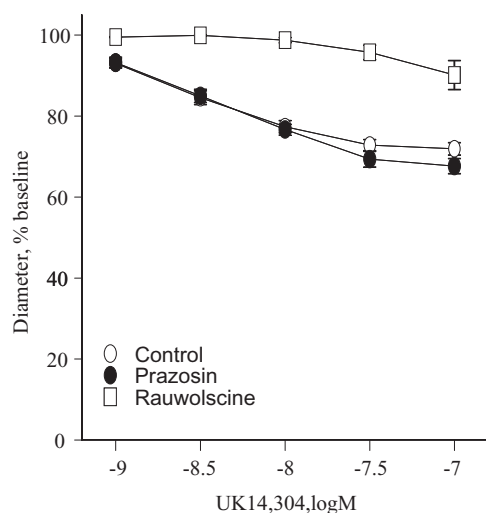


Figure 1 Effects of the selective α_1 -adrenoceptor antagonist prazosin (10^{-7} M) or the selective α_2 -adrenoceptor antagonist rauwolscine (3×10^{-8} M) on constrictor responses to the selective α_2 -adrenoceptor agonist, UK14,304 (10^{-9} to 10^{-7} M) in mouse isolated tail arteries. Constriction was expressed as a percentage of the stable baseline diameter of the arteries (prior to agonist administration) and is presented as means $\pm$ SEM for *n* = 4–5.

inhibitory effect at a high (CC_{60}) compared with a low level of constriction (CC_{15}) (Table 1, Figure 2). In the presence of prazosin (10^{-7} M), rauwolscine (3×10^{-8} M) caused a significant rightward shift in the concentration–effect curve to phenylephrine, having a significantly greater inhibitory effect at a low (CC_{15}) compared with a high level of constriction (CC_{60}) (Table 1, Figure 2). Indeed, the combination of prazosin plus rauwolscine caused a parallel shift in the concentration–effect curve to phenylephrine (Table 1, Figure 2). Similar results were obtained with RX821002 (3×10^{-8} M), another selective α_2 -adrenoceptor antagonist (data not shown).

The effects of antagonists on concentration–effect curves to noradrenaline or cirazoline were qualitatively similar to those observed with phenylephrine. Prazosin (10^{-7} M) caused non-parallel shifts in the concentration–effect curve to the agonists, with greater inhibitory effects at a high (CC_{60}) compared with a low level of constriction (CC_{15}) (Table 1, Figures 3 and 4). Rauwolscine (3×10^{-8} M), when administered alone, did not significantly affect the concentration–effect curves to noradrenaline or cirazoline. However, in the presence of prazosin (10^{-7} M), rauwolscine (3×10^{-8} M) inhibited the concentration–effect curves, having a greater inhibitory effect

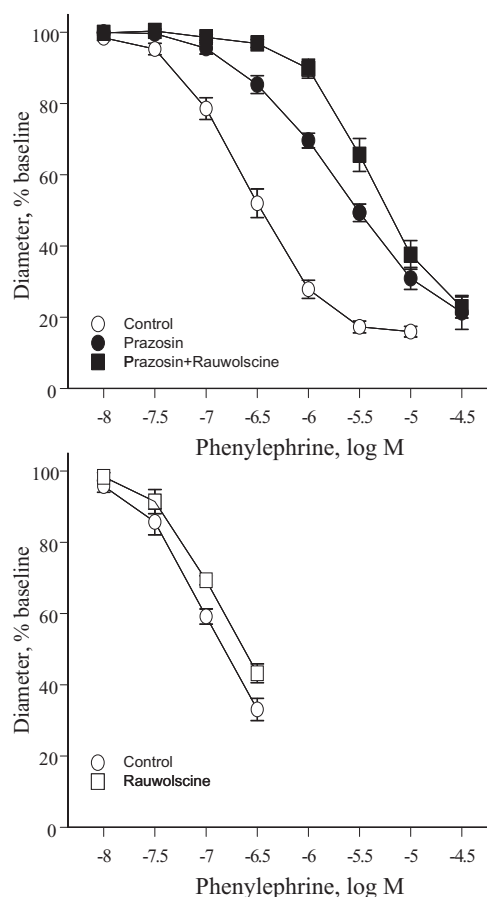


Figure 2 Effects of the selective α_1 -adrenoceptor antagonist prazosin (10^{-7} M), the selective α_2 -adrenoceptor antagonist rauwolscine (3×10^{-8} M) or the combination of prazosin plus rauwolscine on constrictor responses to phenylephrine in mouse isolated tail arteries. Constriction was expressed as a percentage of the stable baseline diameter of the arteries (prior to agonist administration) and is presented as means $\pm$ SEM for *n* = 10 (upper panel) or 5 (lower panel).

Table 1 Effects of antagonists on the concentration–effect curves to adrenoceptor agonists^a

Agonist	Level	Prazosin		Rauwolsine ^b (after Prazosin)		Prazosin + Rauwolsine	
		Log shift ^c	shift	Log shift ^c	shift	Log shift ^c	shift
Phenylephrine (10)	CC ₁₅	0.54 ± 0.13**	3.5	0.66 ± 0.09***	4.5	1.31 ± 0.14***	20.4
	CC ₆₀	1.07 ± 0.15***##	11.6	0.18 ± 0.03*** ##	1.5	1.28 ± 0.14***	19.3
Noradrenaline (7)	CC ₁₅	0.52 ± 0.16*	3.3	0.40 ± 0.14*	2.5	1.06 ± 0.16***	11.5
	CC ₆₀	1.11 ± 0.08***##	12.8	No effect	–	1.14 ± 0.16***	13.8
Cirazoline (4)	CC ₁₅	1.03 ± 0.06***	10.7	0.51 ± 0.06**	3.2	1.60 ± 0.11***	40.2
	CC ₆₀	1.56 ± 0.18**	36.4	No effect	–	1.59 ± 0.23**	39.4

^aShifts in the concentration–effect curves were determined at the CC₁₅ and CC₆₀ levels of constriction. Data are expressed as means ± SEM. The numbers of experiments are indicated in parentheses after the name of the agonist.

^bEffect of rauwolsine refers to the effect of the antagonist in arteries in which α_1 -adrenoceptors have already been blocked with prazosin (10^{–7} M), i.e. the difference between the concentration–effect curves performed in the presence of ‘prazosin’ and ‘prazosin + rauwolsine’. The other comparisons were made to the control concentration–effect curves.

^c*, **, *** indicates the log shift is significant at $P < 0.05$, $P < 0.01$ or $P < 0.001$ respectively. ##, ###, indicates the log shift is significantly different between the CC₁₅ and CC₆₀ levels of constriction at $P < 0.01$ or $P < 0.001$ respectively.

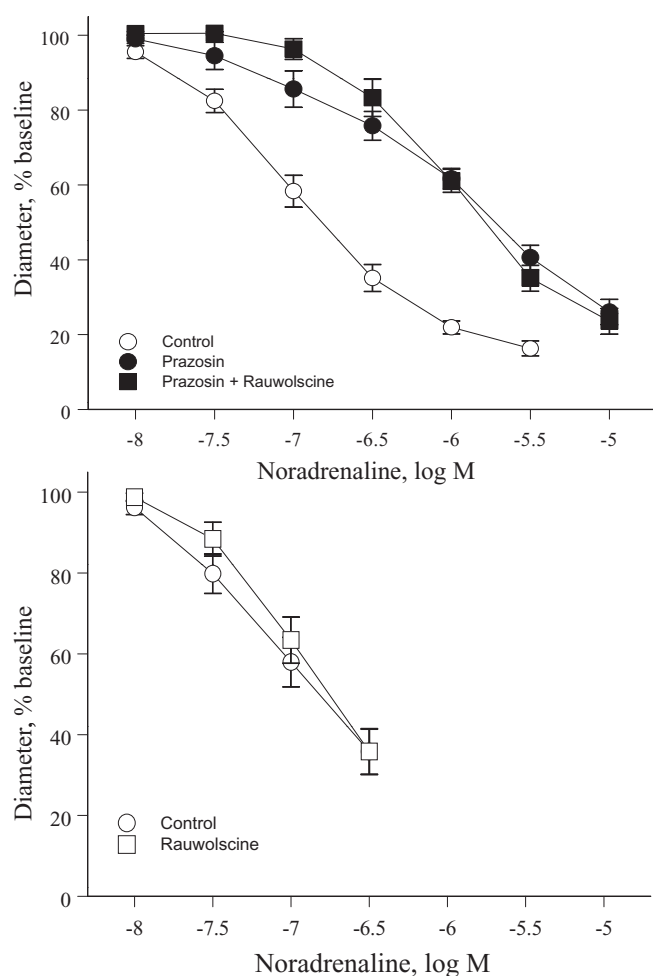


Figure 3 Effects of the selective α_1 -adrenoceptor antagonist prazosin (10^{–7} M), the selective α_2 -adrenoceptor antagonist rauwolsine (3 × 10^{–8} M) or the combination of prazosin plus rauwolsine on constrictor responses to noradrenaline in mouse isolated tail arteries. Constriction was expressed as a percentage of the stable baseline diameter of the arteries (prior to agonist administration) and is presented as means ± SEM for $n = 7$ (upper panel) or 6 (lower panel).

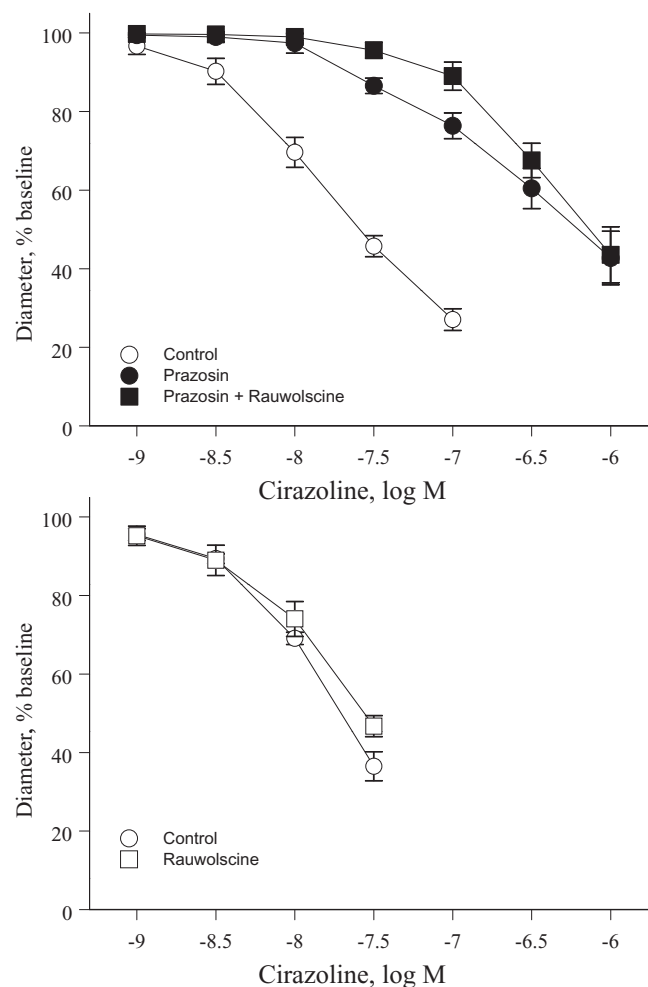


Figure 4 Effects of the selective α_1 -adrenoceptor antagonist prazosin (10^{–7} M), the selective α_2 -adrenoceptor antagonist rauwolsine (3 × 10^{–8} M) or the combination of prazosin plus rauwolsine on constrictor responses to cirazoline in mouse isolated tail arteries. Constriction was expressed as a percentage of the stable baseline diameter of the arteries (prior to agonist administration) and is presented as means ± SEM for $n = 4$ (upper panel) or 5 (lower panel).

Table 2 Antagonist-induced dilation during responses to constrictor agonists^a

Agonist	Treatment	Constriction ^b	Dilation ^{b,c}	
			Peak	After 15 min
Phenylephrine (12)	Rauwolscine	49.5 ± 2.5%	76.2 ± 2.6%***	64.5 ± 2.5%*** ###
Phenylephrine (5)	RX821002	56.2 ± 5.4%	83.4 ± 3.9%***	68.0 ± 5.0%* ##
Phenylephrine (12)	Time Control	55.6 ± 3.0%	55.5 ± 3.1%	54.5 ± 3.2%
Noradrenaline (4)	Rauwolscine	37.5 ± 5.4%	76.9 ± 2.7%***	64.4 ± 4.1%*** #
Noradrenaline (4)	Time Control	45.3 ± 3.9%	47.4 ± 3.2%	52.6 ± 3.7%*** ##
Cirazoline (5)	Rauwolscine	50.6 ± 4.5%	67.5 ± 2.8%***	63.4 ± 4.6%***
Cirazoline (5)	Time Control	56.5 ± 3.7%	57.1 ± 3.5%	57.6 ± 4.6%
U46619 (3)	Rauwolscine	62.4 ± 1.0%	63.8 ± 1.8%	63.0 ± 1.9%

^aDilations to rauwolscine (3×10^{-8} M) or RX821002 (3×10^{-8} M) were evaluated during constrictor responses to adrenoceptor agonists phenylephrine (3×10^{-7} M), noradrenaline (3×10^{-7} M), or cirazoline (3×10^{-8} M), or to the non-adrenoceptor agonist U46619 (3×10^{-9} M). Time-dependent changes in constriction to adrenoceptor agonists (in the absence of antagonists) were evaluated in paired arterial segments (time control).

^bDiameters associated with constriction to the agonists and dilation to the antagonists (or time-dependent changes in agonist constrictions) are all presented as a percentage of the baseline diameter prior to administration of the agonist, and expressed as means ± SEM. The numbers of experiments are provided in parenthesis after the name of the agonist.

***, **, indicates the level of constriction is significant different from the initial response to the constrictor agonist at $P < 0.01$ or $P < 0.001$ respectively. #, ##, ###, indicates the diameter is significantly different from that occurring during the peak response to the antagonist at $P < 0.05$, $P < 0.01$ or $P < 0.001$, respectively.

at a low (CC_{15}) compared with high level of constriction (CC_{60}) (Table 1, Figures 3 and 4). As with phenylephrine, the combination of prazosin plus rauwolscine caused parallel shifts in the concentration–effect curve to noradrenaline and cirazoline (Table 1, Figures 3 and 4).

Reversal of agonist-induced constriction by α_2 -adrenoceptor antagonists

α -Adrenoceptor activation in quiescent arteries. During stable constriction to phenylephrine (3×10^{-7} M), noradrenaline (3×10^{-7} M) or cirazoline (3×10^{-8} M), rauwolscine (3×10^{-8} M) caused an impressive transient dilation that diminished over time and was significantly decreased 15 min after the peak relaxation (Table 2, Figure 5). RX821002 (3×10^{-8} M) had a similar dilator effect to rauwolscine (Table 2, Figure 5). In paired arterial segments not treated with α_2 -adrenoceptor antagonists, there was no time-dependent change in constriction to phenylephrine or cirazoline during the exposure period to the antagonists (Table 2, Figure 5). There was however a small time-dependent decrease in constriction to noradrenaline (Table 2, Figure 5), which may reflect oxidation and inactivation of this agonist.

Therefore, although pretreatment with rauwolscine (when administered alone) did not inhibit the concentration–effect curves to noradrenaline, phenylephrine or cirazoline (Figures 2B, 3B and 4B), it caused impressive transient dilation when administered during sustained constriction to these agonists (Figure 5). Concentration–effect curves were generated using initial peak constrictions, whereas the antagonist dilator responses were assessed during more prolonged constrictor responses to the agonists. Therefore, the difference in antagonist activity could reflect distinct temporal characteristics of α_1 -adrenoceptor and α_2 -adrenoceptor constrictor components, with α_1 -adrenoceptors contributing predominantly to the initial response and α_2 -adrenoceptors contributing to maintained constriction. However, pretreatment with rauwolscine (3×10^{-8} M) did not affect the time course of constriction to phenylephrine (3×10^{-7} M) (Figure 6).

The dilator response to α_2 -adrenoceptor antagonists was dependent on α_2 -adrenoceptor activity. Mouse mesenteric arteries do not express functional constrictor α_2 -adrenoceptors, and do not constrict in response to UK14,304 (Figure 7) (Flavahan, 2005). When administered during a sustained constriction to phenylephrine in mouse mesenteric artery, rauwolscine (3×10^{-8} M) did not cause dilation (Figure 7). Similarly, during constriction of tail arteries to the thromboxane receptor agonist U46619, rauwolscine (3×10^{-8} M) did not cause dilation (Table 2, Figure 8).

α_2 -Adrenoceptor activation during constriction with U46619. The ability of rauwolscine (3×10^{-8} M) to reverse (or dilate) an α_2 -adrenoceptor-mediated constriction to UK14,304 was evaluated during simultaneous constriction with U46619. In tail arteries constricted with U46619, the subsequent addition of the α_2 -adrenoceptor agonist UK14,304 (3×10^{-8} M) caused a significant further constriction (Figure 8). When administered during this combined constriction, the α_2 -adrenoceptor antagonist rauwolscine (3×10^{-8} M) caused an initial dilation that completely reversed the α_2 -adrenoceptor component of the constriction and also almost abolished the U46619-dependent component of constriction (Figure 8). After 15 min, the dilation to rauwolscine was significantly attenuated and the new stable level of constriction was not significantly different from the original constriction to U46619, although it was still significantly less than the combined constrictor response to U46619 plus UK14,304 (Figure 8). In the absence of UK14,304, rauwolscine (3×10^{-8} M) did not cause dilation during constriction to U46619 (Figure 8).

The dilation associated with antagonist-induced termination of α_2 -adrenoceptor activity was also observed following rapid removal of UK14,304 during a combined constriction to U46619 plus UK14,304. For these studies, constriction of tail arteries was analysed in an open perfusion system rather than the standard re-circulating perfusion system used in our experiments. In tail arteries constricted with U46619, the α_2 -adrenoceptor agonist UK14,304 (3×10^{-8} M) caused a

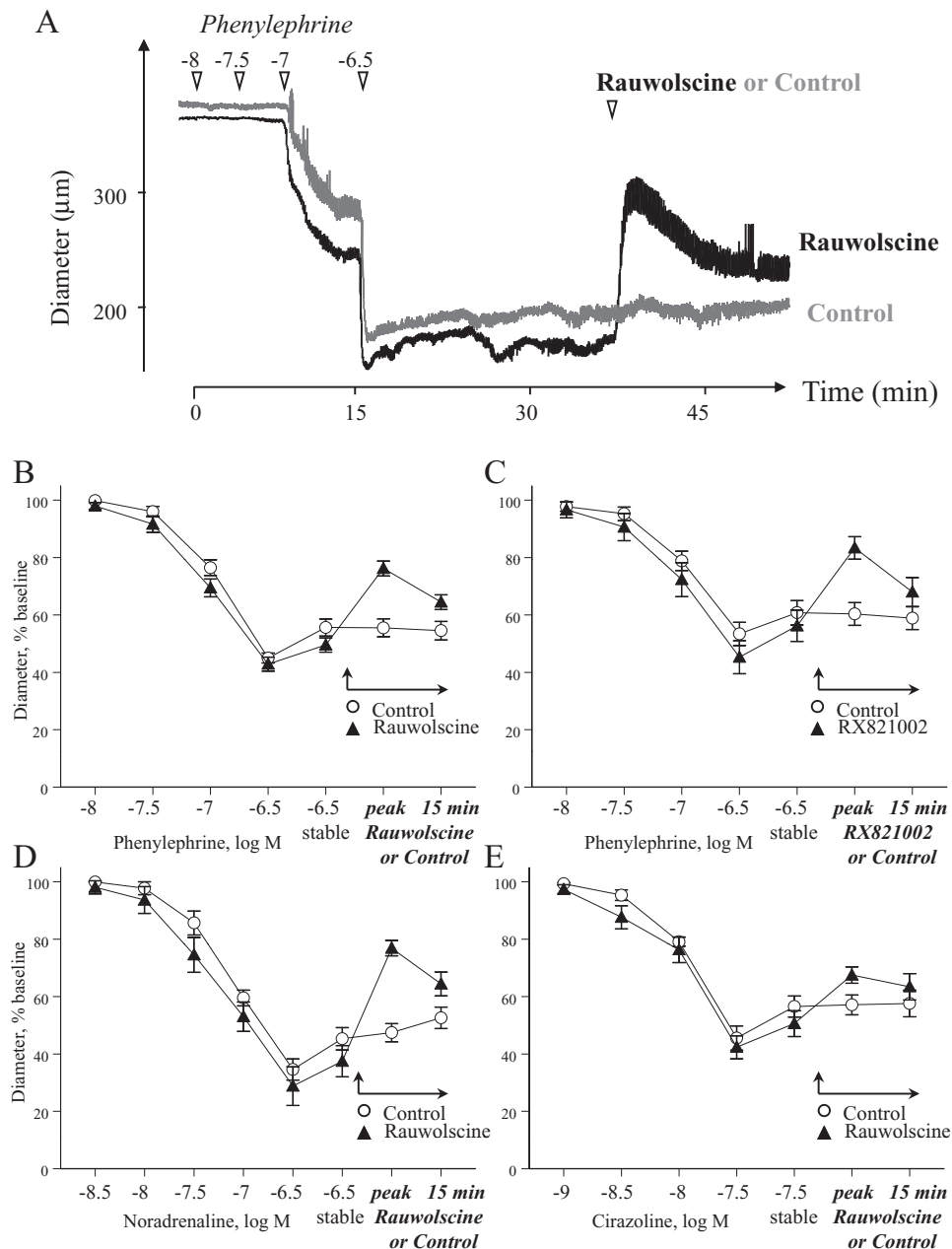


Figure 5 The acute dilator effect of selective α_2 -adrenoceptor antagonists rauwolscline (3×10^{-8} M) or RX821009 (3×10^{-8} M) during stable constriction of mouse isolated tail arteries with phenylephrine, cirazoline or noradrenaline. Paired arterial segments were exposed to increasing concentrations of an α -adrenoceptor agonist and constriction allowed to stabilize ('stable', at 3×10^{-7} M for noradrenaline or phenylephrine, and 3×10^{-8} M for cirazoline). Administration of the antagonists then caused dilation, which was assessed at the peak response and 15 min later. In all experiments, one of each paired arterial segments served as a time control and was not exposed to the antagonists (Control). The upper trace (A) presents a typical recording from one of the experiments with phenylephrine and rauwolscline. The lower graphs represent the means $\pm$ SEM for $n = 12$ (B), $n = 5$ (C), $n = 4$ (D) and $n = 5$ (E). All responses were expressed as a percentage of the stable baseline diameter of the arteries prior to agonist administration.

significant further constriction (Figure 9). Abrupt removal of UK14,304 from the perfusate (in the continued presence of U46619) caused an initial dilation that completely reversed the α_2 -adrenoceptor component of the constriction and almost abolished the U46619-dependent component of constriction (Figure 9). After 15 min, the dilation was significantly diminished and the new stable level of constriction was not significantly different from the original constriction

to U46619, although it was still significantly less than the combined constrictor response to UK14,304 plus U46619 (Figure 9). The constriction and dilation associated with introduction and removal, respectively, of UK14,304 (3×10^{-8} M) in the open perfusion system were dramatically reduced when the experiment was repeated in the presence of the α_2 -adrenoceptor antagonist, rauwolscline (3×10^{-8} M). Following pretreatment with rauwolscline, the introduction of

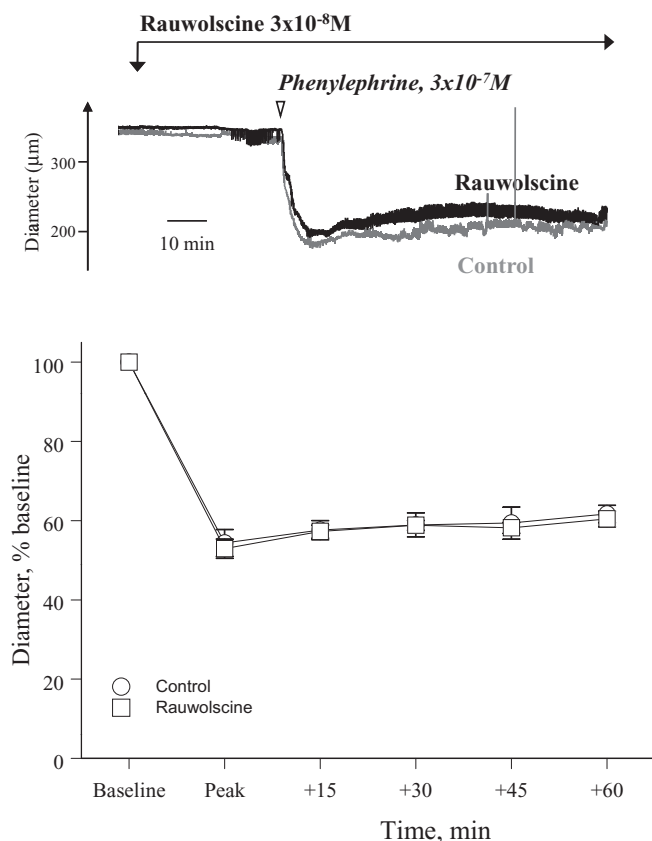


Figure 6 Effect of the selective α_2 -adrenoceptor antagonist rauwolscline (3×10^{-8} M) on the time course of the constrictor response to phenylephrine (3×10^{-7} M) in mouse isolated tail arteries. The upper panel presents a recording from a typical experiment, whereas the lower panel presents the means $\pm$ SEM for $n = 4$. In the recording, internal diameter is presented in microns, whereas in the graph, internal diameter is expressed as a percentage of the baseline prior to agonist administration. The response to phenylephrine was assessed at the peak of constriction and then 15, 30, 45 and 60 min later.

UK14,304 (3×10^{-8} M) to U46619-constricted arteries caused constriction equivalent to $6.4 \pm 1.5\%$ of baseline diameter ($n = 4$) (was $26.5 \pm 0.9\%$ under control conditions, $n = 4$, Figure 9), whereas removal of UK14,304 was associated with simple reversal of this constriction and a dilation equivalent to $5.4 \pm 1.6\%$ of baseline diameter ($n = 4$) (was $51.2 \pm 2.8\%$ under control conditions, $n = 4$, Figure 9).

Role of the endothelium. Endothelial denudation significantly reduced the dilator response evoked by administration of rauwolscline (3×10^{-8} M) (Figure 8) or abrupt removal of UK14,304 (Figure 9) in tail arteries constricted with U46619 plus UK14,304. In endothelium-denuded arteries, the dilation in each case was sufficient to abolish the α_2 -adrenoceptor component of the combined constriction but it did not significantly affect the component of constriction evoked by U46619 (Figures 8 and 9).

Endothelium denudation did not significantly affect the constrictor response to activation of α_2 -adrenoceptors with UK14,304 either during combined constriction with U46619 (e.g. Figures 8 and 9) or under basal conditions (i.e. absence of U46619). Under basal conditions, concentration–effect curves to UK14,304 were characterized by log CC₁₅ values of -8.07 ± 0.21 and -7.88 ± 0.13 ($n = 5$; $P = \text{NS}$, not significant) and maximal constriction to $74.9 \pm 3.5\%$ and $77.6 \pm 1.3\%$ of baseline diameter ($n = 5$, $P = \text{NS}$) for endothelium-containing and endothelium-denuded arteries respectively. Likewise, endothelium denudation did not significantly affect constriction to U46619 (log CC₁₅ values of -8.17 ± 0.04 and -8.12 ± 0.07 , $n = 5$, $P = \text{NS}$ and maximal-attained constriction, at 10^{-7} M, to $25.7 \pm 2.6\%$ and $17.7 \pm 3.6\%$ of baseline diameter, $n = 5$, $P = \text{NS}$, for endothelium-containing and endothelium-denuded arteries respectively).

Mesenteric Arteries

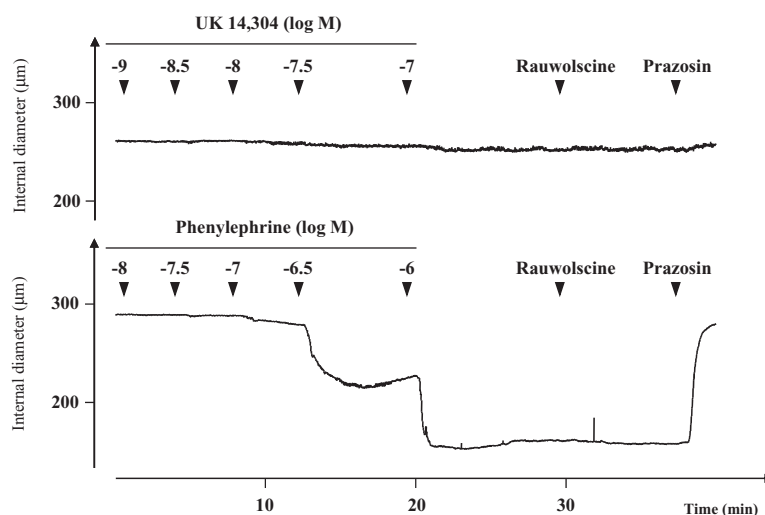


Figure 7 Representative recording demonstrating the activity of the α -adrenoceptor agonists UK14,304 (UK, 10^{-9} to 10^{-7} M) or phenylephrine (PE, 10^{-8} to 10^{-6} M) in mouse mesenteric arteries. UK14,304 failed to constrict the artery. Constriction to phenylephrine was not affected by acute administration of the selective α_2 -adrenoceptor antagonist rauwolscline (3×10^{-8} M), but was reversed by the selective α_1 -adrenoceptor antagonist prazosin (10^{-7} M). The recording is representative of four experiments.

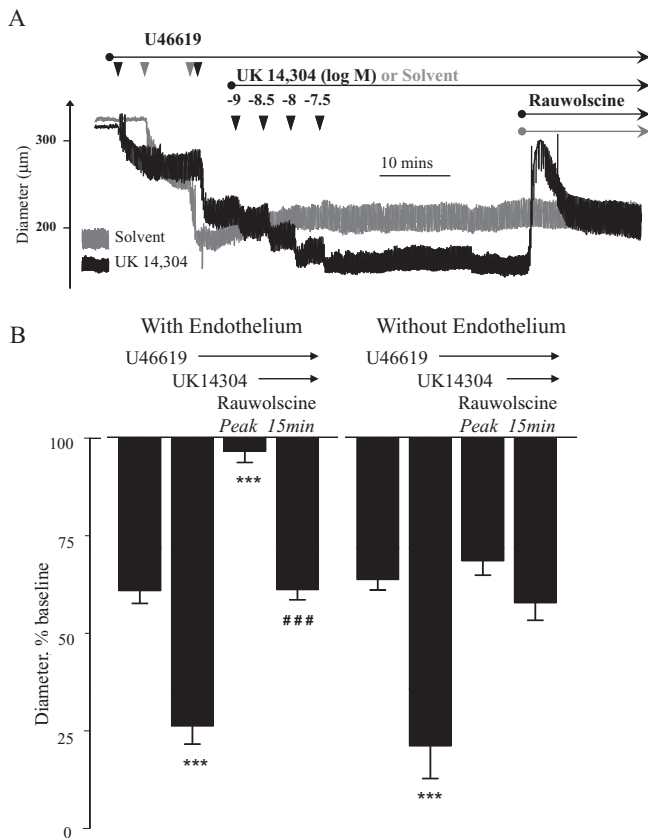


Figure 8 Effects of acute administration of rauwolscine (3×10^{-8} M) on combined constriction to the thromboxane receptor agonist U46619 and the α_2 -adrenoceptor agonist UK14,304 in mouse isolated tail arteries. Arteries (with and without endothelium) were initially constricted with U46619 to approximately 60% of resting baseline diameter, and then further constricted with UK14,304 (up to 3×10^{-8} M). Once the constriction stabilized, the dilator effect of the selective α_2 -adrenoceptor antagonist rauwolscine (3×10^{-8} M) was determined. (A) Representative recording from a typical experiment. In arteries constricted only with U46619 (grey trace), rauwolscine did not cause dilation (Table 2). (B) Means $\pm$ SEM for $n = 5$. Responses were expressed as a percentage of the baseline internal diameter prior to administration of the agonists. The influence of rauwolscine was assessed at the peak of dilation and 15 min later. * indicates that the diameter is significantly different from the initial constriction to U46619 ($***P < 0.001$). # indicates the effect of rauwolscine after 15 min is significantly different from the peak effect (### $P < 0.001$).

Discussion

Because of its role in thermoregulation, the mouse tail artery is somewhat unusual by possessing a prominent smooth muscle α_2 -adrenoceptor constrictor response (Chotani *et al.*, 2000; Flavahan, 2005). However, even at its most powerful, α_2 -adrenoceptor constriction evoked by high-efficacy agonists such as UK14,304 is a low-maximum response compared with α_1 -adrenoceptor activation (Flavahan and McGrath, 1984; Flavahan *et al.*, 1984), causing only ~30% constriction of the tail artery (Chotani *et al.*, 2000; Flavahan, 2005). Because of this disparity in maximal responses to activation of α_1 - and α_2 -adrenoceptor subtypes, the selective α_1 -adrenoceptor antagonist prazosin causes non-parallel shifts in the concentration–effect curve to non-selective agonists (e.g.

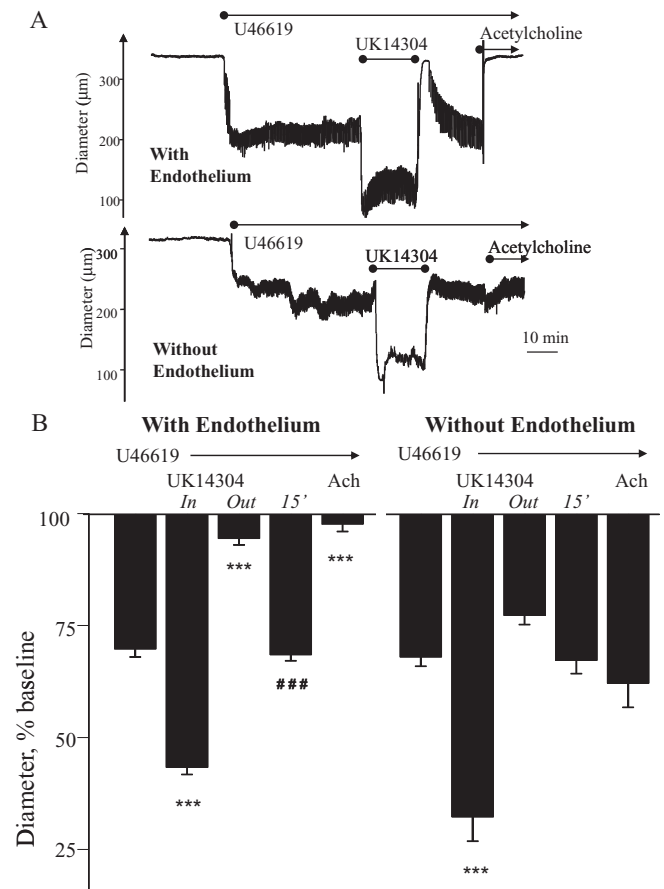


Figure 9 Effects of rapid withdrawal of the selective α_2 -adrenoceptor agonist UK14,304 (3×10^{-8} M) during combined constriction of mouse isolated tail arteries to UK14,304 and the thromboxane receptor agonist U46619. In an open perfusion system, arteries (with and without endothelium) were initially constricted with U46619 to approximately 60% of resting baseline diameter, and then further constricted with UK14,304 (3×10^{-8} M). Once the constriction had stabilized, UK14,304 was removed from the perfusate and constriction was maintained only by the continued presence of U46619. (A) Representative recording from a typical experiment in arteries with and without endothelium. (B) Means $\pm$ SEM for $n = 4$. Responses were expressed as a percentage of the baseline internal diameter prior to administration of the agonists. The effect of removing UK14,304 was assessed at the peak of dilation ('out') and 15 min later (15'). * indicates that the diameter is significantly different from the initial constriction to U46619 ($***P < 0.001$). # indicates the effect of removing UK14,304 is significantly different between the peak effect ('out') and 15 min later (15') (### $P < 0.001$).

noradrenaline) being more potent at high compared with low levels of constriction (Flavahan and McGrath, 1984; Flavahan *et al.*, 1984; 1987; Flavahan and Vanhoutte, 1986b). At high levels of constriction, the inhibitory potency of prazosin is consistent with its antagonism of α_1 -adrenoceptors whereas at low levels of constriction, the presence of the low-maximum α_2 -adrenoceptor response restricts the inhibitory effect of prazosin resulting in 'prazosin-resistance' (Flavahan and McGrath, 1984; Flavahan *et al.*, 1984; 1987; Flavahan and Vanhoutte, 1986b). Indeed, in the present study, prazosin caused a non-parallel shift in the concentration–effect curve to phenylephrine, being more potent at a high level of constriction (60%, which is outside the range of the

low-maximum α_2 -adrenoceptor response) compared with a lower intensity of constriction (15%, which is approximately 50% of the maximal α_2 -adrenoceptor response). The potency of prazosin at the high level of constriction (11.6-fold shift, CC_{50}) is consistent with an antagonist dissociation constant (K_B) of 9×10^{-9} M and α_1 -adrenoceptor activity in cutaneous blood vessels (Flavahan *et al.*, 1984). However, at the low level of constriction, the potency of prazosin (3.5-fold shift, CC_{15}) was significantly lower and not consistent with simple α_1 -adrenoceptor activity (Flavahan *et al.*, 1984; Flavahan and Vanhoutte, 1986a). In the presence of prazosin, the selective α_2 -adrenoceptor antagonist rauwolscline caused inhibition of the concentration–effect curve at the low but not at the high level of constriction, and therefore removed this prazosin-resistant component. The demonstration that the prazosin-resistant component is rauwolscline-sensitive confirms that it represents α_2 -adrenoceptor constriction to the agonist. In the absence of prazosin, rauwolscline did not significantly affect the concentration–effect curve to phenylephrine, which confirms that rauwolscline acts selectively to inhibit α_2 -adrenoceptors and indicates that phenylephrine acts predominantly through α_1 -adrenoceptors. If one compares the concentration–effect curve to phenylephrine after rauwolscline with the curve after prazosin plus rauwolscline, the inclusion of prazosin caused a parallel shift in the curve to phenylephrine and there was no indication of resistance to prazosin. This again demonstrates that rauwolscline inhibits the prazosin-resistant, α_2 -adrenoceptor-mediated constriction to phenylephrine. However, loss of the α_2 -adrenoceptor component did not influence the control concentration–effect curve to phenylephrine and its influence was only evident following inhibition of the α_1 -adrenoceptor response. We have previously demonstrated that there is minimal interaction between α_1 - and α_2 -adrenoceptor components of vasoconstriction in cutaneous blood vessels (Flavahan *et al.*, 1984; Flavahan and Vanhoutte, 1987). The control concentration–effect curve to phenylephrine therefore approximates the α_1 -adrenoceptor component of the response, and the curve after prazosin (i.e. the prazosin-resistant component at the CC_{15} level of constriction) identifies the α_2 -adrenoceptor component. The magnitude of the difference between these components characterizes the functional selectivity of phenylephrine at α_1 - and α_2 -adrenoceptors in the tail artery, which is 3.5-fold (Table 1).

The effects of prazosin and rauwolscline on responses to cirazoline or the physiological agonist noradrenaline were qualitatively similar to those observed with phenylephrine and were also consistent with activation of α_1 -adrenoceptors and α_2 -adrenoceptors by these agonists. As with phenylephrine, rauwolscline did not inhibit the concentration–effect curves indicating that the control curve approximates the α_1 -adrenoceptor component of the response. The difference between the control curve and the prazosin-resistant component of the response (at the CC_{15} level of constriction), which provides an indication of the functional selectivity of the agonists at α_1 - and α_2 -adrenoceptors, was 3.3-fold and 10.7 for noradrenaline and cirazoline respectively. Therefore, in the mouse tail artery, all three agonists act as preferential α_1 -adrenoceptor agonists. However, phenylephrine and noradrenaline exhibit similar low selectivity at α_1 -adrenoceptor

and α_2 -adrenoceptors, and cirazoline was the most selective α_1 -adrenoceptor agonist tested.

Phenylephrine is generally considered to be a highly selective α_1 -adrenoceptor agonist. Although previous studies have challenged this assumption, reporting selectivity ratios at α_1 - and α_2 -adrenoceptors of ≥ 10 -fold (Flavahan and McGrath, 1981; Guimaraes *et al.*, 1987; Brown *et al.*, 1988), the relative lack of selectivity for phenylephrine observed in the present study was still surprising. There are theoretical conditions when selective activation of one receptor could provide an erroneous appearance of non-selectivity. For example, constitutively active α_2 -adrenoceptors (Jansson *et al.*, 1998; Murrin *et al.*, 2000) could theoretically amplify an α_1 -adrenoceptor response to phenylephrine. Likewise, the possible existence of α_1/α_2 -adrenoceptor oligomers could theoretically enable α_1 -adrenoceptor-dependent activation of α_2 -adrenoceptor function. Under these conditions, phenylephrine could function as a selective α_1 -adrenoceptor agonist, but the resulting vasoconstriction would be inhibited by selective α_2 -adrenoceptor antagonists acting as inverse agonists (Jansson *et al.*, 1998; Murrin *et al.*, 2000; Vilardaga *et al.*, 2005). However, these scenarios are unlikely to have occurred in the mouse tail artery. There was no evidence of constitutively active α_2 -adrenoceptors: rauwolscline did not inhibit constriction to U46619 or to the α -adrenoceptor agonists (when administered in the absence of prazosin). Furthermore, these scenarios require α_1 -adrenoceptor activation to enable visualization of the α_2 -adrenoceptor constrictor response, which would cause the α_2 -adrenoceptor response to be sensitive to inhibition of α_1 -adrenoceptors, i.e. prazosin-sensitive. The presence of a prazosin-resistant, rauwolscline-sensitive constriction to phenylephrine suggests that the α_2 -adrenoceptor constrictor component of the response to the agonist is not dependent on α_1 -adrenoceptor activation. One caveat to this interpretation is that α_1/α_2 -adrenoceptor oligomerization might change the binding affinity of prazosin for α_1 -adrenoceptors. However, at least for α_1/α_1 -adrenoceptor dimers, the oligomerization process does not alter the ability of prazosin to inhibit receptor activation (Uberti *et al.*, 2003). Furthermore, oligomerization of α_1 - and α_2 -adrenoceptors has never been documented. Therefore, the pharmacological analysis indicates that phenylephrine is acting as a poorly selective agonist at α_1 -adrenoceptors in this blood vessel.

Although rauwolscline did not affect the control concentration–effect curves to the α -adrenoceptor agonists, it caused a rapid and marked relaxation when administered during stable constriction to the agonists. The magnitude of the dilation to rauwolscline was dependent on the selectivity of the agonists (most powerful with phenylephrine and noradrenaline, less effective with cirazoline) suggesting that the effect was mediated by inhibition of α_2 -adrenoceptors. Indeed, rauwolscline did not cause relaxation during constriction of mouse tail artery to U46619 or during phenylephrine-induced constriction of mouse mesenteric artery, which does not express functional smooth muscle constrictor α_2 -adrenoceptors (Flavahan, 2005). Furthermore, the dilator response to rauwolscline in tail arteries was mimicked by another α_2 -adrenoceptor antagonist, RX821002. Indeed, the acute administration of α_2 -adrenoceptor antagonists appears to be an efficient and reliable approach to evaluate agonist

selectivity. However, the ability of α_2 -adrenoceptor antagonists to cause powerful relaxation was perplexing. For example, with phenylephrine or noradrenaline, the magnitude of the peak dilation to rauwolscine or RX821002 would be equivalent to an approximate threefold shift in the concentration–effect curve to the agonist (Figure 5). Surprisingly, this inhibitory effect was not evident from directly analysing the effects of the α_2 -adrenoceptor antagonists (when given alone) on the concentration–effect curves. Because these curves were constructed using the initial peak response to the agonists, we considered that the dilator responses might result from temporal differences in the α_1 -adrenoceptor and α_2 -adrenoceptor components of the response: the initial peak response could be dominated by the α_1 -adrenoceptor response, whereas α_2 -adrenoceptors may play a more important role to maintain vasoconstriction (McGrath *et al.*, 1982; Fukui *et al.*, 2005). However, there was no difference in the time course to phenylephrine when the tissues were pretreated with the α_2 -adrenoceptor antagonist. Indeed, we found no evidence for temporal differences in α_1 -adrenoceptor and α_2 -adrenoceptor vasoconstriction.

The basis for the surprising dilation to α_2 -adrenoceptor antagonists became evident when rauwolscine's ability to reverse an α_2 -adrenoceptor-dependent constriction to UK14,304 was assessed during simultaneous constriction to U46619. Rauwolscine caused dilation that not only completely reversed the α_2 -adrenoceptor-mediated component of constriction; it virtually abolished the constriction to U46619. This profound dilation, however, was only transient: after 15 min the dilator effect of rauwolscine was restricted to elimination of the α_2 -adrenoceptor constriction without any effect on the U46619 component of the constriction. The large transient dilation did not result from non-selective activity of the antagonist because rauwolscine did not affect constriction to U46619 in the absence of the α_2 -adrenoceptor agonist. Therefore, the dilation appeared to be dependent on the termination of α_2 -adrenoceptor agonist activity. Indeed, abrupt removal of UK14,304 (during combined constriction to UK14,304 and U46619) also caused marked dilation of the U46619 component of constriction. As with rauwolscine, this dilation was transient and after 15 min, the original U46619 constriction was regained. These results indicate that α_2 -adrenoceptor activation initiates two responses in this blood vessel: constriction that normally predominates and reverses quickly following interruption of receptor stimulation, and dilation, which is slower to reverse than the constrictor response. By rapidly terminating agonist–receptor interaction, acute administration of an α_2 -adrenoceptor antagonist leads to immediate cessation of the constrictor response and a slower reversal of the dilator response. The peak transient dilation therefore reflects the combined loss of the α_2 -adrenoceptor constrictor component and the uncovering of the α_2 -adrenoceptor dilator response. With the slower decay of the α_2 -adrenoceptor-mediated dilator response, the peak dilation gradually recedes (within 15 min) to reveal the true magnitude of the underlying α_2 -adrenoceptor-independent constriction. Endothelial denudation abolished the transient dilator responses associated with rapid termination of the α_2 -adrenoceptor constrictor response (caused either by administration of an antagonist or abrupt removal

of the agonist) indicating that the slowly reversing dilation is mediated by activation of endothelial α_2 -adrenoceptors (Cocks and Angus, 1983; Bockman *et al.*, 1993; Shafaroudi *et al.*, 2005). Endothelial α_2 -adrenoceptors were considered to be most prominent in the coronary system and to play a limited role in blood vessels (including cutaneous arteries) where smooth muscle α_2 -adrenoceptors play a prominent constrictor role (Miller and Vanhoutte, 1985; Angus *et al.*, 1986; Flavahan *et al.*, 1989; Ohgushi *et al.*, 1993; Flavahan, 2005). The results of the present study indicate that α_2 -adrenoceptors can initiate powerful endothelium-dependent dilation and smooth muscle constriction in cutaneous arteries. Endothelium denudation did not significantly affect the vasoconstrictor response to α_2 -adrenoceptor activation in the mouse tail artery. This is consistent with our previous studies (Flavahan, 2005) and indicates that the α_2 -adrenoceptor endothelium-dependent dilator response is normally dominated by the smooth muscle constrictor response.

At high concentrations, rauwolscine and other α_2 -adrenoceptor antagonists have been reported to relax contractions evoked by phenylephrine in the rat isolated aorta and mesenteric arteries (Kim *et al.*, 1999; Artigues-Varin *et al.*, 2002). These rat arteries do not express functional constrictor α_2 -adrenoceptors, and the contraction to phenylephrine is mediated by α_1 -adrenoceptors (Macia *et al.*, 1984; Flavahan and Vanhoutte, 1986a; Nielsen *et al.*, 1991; Flavahan, 2005). Indeed, because of the high concentrations of rauwolscine ($IC_{50} \sim 10^{-6}$ M) and other α_2 -adrenoceptor antagonists employed by Kim *et al.* (1999) and Artigues-Varin *et al.* (2002), these agents no longer function as selective α_2 -adrenoceptor antagonists and inhibit α_1 -adrenoceptors (Digges and Summers, 1983). Therefore, in contrast to the results of the present study, the sustained relaxation to rauwolscine in rat aorta and mesenteric artery is mediated by antagonism of α_1 -adrenoceptors (Artigues-Varin *et al.*, 2002).

In conclusion, the results of the present study demonstrate that the prototypic α_1 -adrenoceptor agonist phenylephrine has limited selectivity at α -adrenoceptor subtypes and is a powerful agonist at vascular smooth muscle α_2 -adrenoceptors. However, the transient dilator response evoked by α_2 -adrenoceptor antagonists is much greater than the contribution of α_2 -adrenoceptors to the constrictor response. This reflects the presence of a slowly-reversing α_2 -adrenoceptor-mediated endothelium-dependent dilation, which transiently amplifies the dilator response to α_2 -adrenoceptor antagonism. Indeed, this provides a novel, rapid and sensitive approach to assess agonist activity at vascular α -adrenoceptors.

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Statement of conflicts of interest

None.

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