α_{1D}-Adrenoceptors are responsible for the high sensitivity and the slow time-course of noradrenaline-mediated contraction in conductance arteries

Nicla Flacco¹, Jaime Parés¹, Eva Serna¹, Vanessa Segura¹, Diana Vicente¹, Miguel Pérez-Aso¹, María Antonia Noguera¹, María Dolores Ivorra¹, John C. McGrath² & Pilar D'Ocon^{1,*}

¹Departamento de Farmacología, Facultad de Farmacia, Universitat de València, Valencia, Spain

²Autonomic Physiology Unit, School of Life Sciences, College of Medical, Veterinary and Life Sciences, University of Glasgow, Glasgow, U.K.

Keywords

 $\alpha_{1\text{A}}\text{-}\text{adrenoceptors, conductance and}$ resistance vessels, contraction time-course

Correspondence

Pilar D'Ocon, Departament de Farmacologia, Universitat de València Facultad de Farmàcia, Avda. Vicent Andres Estelles s/n 46100 Burjassot, València, Spain. Tel: +34963544828; Fax: +34963544943; E-mail: doconp@uv.es

Funding Information

This study was supported by research grants to the University of Glasgow from the UK Medical Research Council (G0000042, I.D.51,240) and to the University of Valencia from the Instituto de Salud Carlos III Fondo de Investigaciones Sanitarias (FIS PI070509), Ministerio de Ciencia e Innovacion, SAF (2007-62,120), Fondos FEDER, Generalitat Valenciana (GVACOMP2009/261) and Research funds of the Universitat de Valencia (UV-INV-AE11-42,176).

Received: 2 May 2013; Revised: 8 May 2013; Accepted: 17 May 2013

Pharma Res Per, 1 (1), 2013, e00001, doi: 10.1002/prp2.1

doi: 10.1002/prp2.1

Abstract

The objective of this study was to determine whether the different time-course characteristics of α_1 -adrenoceptor-mediated contraction in arteries can be related to the subtypes involved. Contractile responses to noradrenaline (NA) were compared with inositol phosphate accumulation and extracellular signalregulated kinase (ERK)1/2 phosphorylation after α_1 -agonist stimuli in the same vessels in the presence or absence of α_1 -antagonists in rat or in α_1 -subtype knockout (KO) mice. Aorta, where α_{1D} -AR is the main functional subtype, had higher sensitivity to NA (in respect of inositol phosphate [IP], pERK1/2, and contractile response) than tail artery, where the α_{1A} -adrenoceptor subtype is predominant. Furthermore, the contraction in aorta exhibited a slower decay after agonist removal and this was consistent in all strains harboring α_{1D} adrenoceptors (from rat, α_{1B} -KO, and wild-type [WT] mice) but was not observed in the absence of the α_{1D} -adrenoceptor signal (α_{1D} -adrenoceptor blocked rat aorta or aorta from α_{1D} -KO). IP formation paralleled α_1 -adrenoceptor-mediated contraction (agonist present or postagonist) in aorta and tail artery. High sensitivity to agonist and persistence of response after agonist removal is a property of α_{1D} -adrenoceptors. Therefore, the preponderance of this subtype in noninnervated conductance arteries such as aorta allows responsiveness to circulating catecholamines and prevents abrupt changes in vessel caliber when the stimulus fluctuates. Conversely, in innervated distributing arteries, high local concentrations of NA are required to activate α_{1A} -adrenoceptors for a response that is rapid but short lived allowing fine adjustment of the contractile tone by perivascular sympathetic nerves.

Abbreviations

AR, α_1 -adrenoceptors; CRC, Concentration–response curves; EDTA, ethylenediaminetetraacetic acid; IP, inositol phosphate; KO, knockout; MAPK, mitogen-activated protein kinase; NA, noradrenaline; PBS, phosphate-buffered saline; PDZ, PSD95/DlgA/Zo-1; PVDF, polyvinyldiene fluoride; WT, wild-type.

Introduction

The α_1 -adrenoceptors (ARs) are responsible for the contractile response to catecholamines in blood vessels and, classically, three different subtypes have been characterized, α_{1A} , α_{1B} , and α_{1D} -ARs. The presence of mRNA or receptor protein is not well correlated with contractile function; in several examples, mRNA and protein for all three α_1 -AR subtypes are expressed in a vessel, yet pharmacological analysis shows that a single subtype is mainly responsible for mediating contraction. There is, however, some correlation between the subtype involved in mediating vascular contraction and the type of vessel. For example, α_{1A} -ARs mediate contraction of well-innervated distributing arteries such as renal (Hrometz et al. 1999), tail (Lachnit et al. 1997; Tanaka et al. 2004), and distal

© 2013 The Authors. *Pharmacology Research & Perspectives* published by John Wiley & Sons Ltd, British Pharmacological Society and American Society for Pharmacology and Experimental Therapeutics. This is an open access article under the terms of the Creative Commons Attribution License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited. mesenteric and resistance arteries such as small mesenteric branches (Philipp and Hein 2004; Martí et al. 2005; Methven et al. 2009a). On the other hand the α_{1D} -AR has been shown to regulate the contraction of poorly innervated conductance arteries such as the aorta, femoral, iliac, carotid, pulmonary, and superior mesenteric artery (Piascik et al. 1995; Hussain and Marshall 1997; Rudner et al. 1999; Gisbert et al. 2000; Arévalo-León et al. 2003; Martí et al. 2005; Methven et al. 2009b) and there is only limited direct evidence that the α_{1B} -AR is a mediator of contractile function in blood vessels (Cavalli et al. 1997; Daly et al. 2002; Tanoue et al. 2003; Cotecchia 2010; Docherty 2010). Thus, vascular α_1 -AR subtypes may correlate with the different functions of smooth muscle in these different vascular types, that is, compliance of large arteries (α_{1D}) and redistribution of blood flow between different organ systems (α_{1A}) as we have previously discussed (Daly et al. 2002; Ziani et al. 2002).

In general, a1-ARs manifest different sensitivity to agonists, the α_{1D} -subtype being the most sensitive (Theroux et al. 1996; Taguchi et al. 1998; Gisbert et al. 2000; Piascik and Perez 2001; Daly et al. 2002). Once activated, the three α_1 -AR subtypes interact with the Gq protein but can also activate a variety of other signaling pathways such as Gi and Go proteins or mitogen-activated protein kinases (MAPKs) (Hawrylyshyn et al. 2004; Hein and Michel 2007; Cotecchia 2010) that are less well explored in native tissues. Nevertheless, there are marked differences in the ability of each subtype to generate intracellular second messengers (García-Sainz et al. 1999b; Zhong and Minneman 1999; Keffel et al. 2000; Piascik and Perez 2001). The α_{1A} -AR is most efficiently coupled to inositol phosphate production, increases in cytosolic calcium concentrations, and MAPKs pathway, whereas the α_{1D} -AR is poorly coupled to intracellular signaling cascades (Schwinn et al. 1991; Theroux et al. 1996; Taguchi et al. 1998; Zhong and Minneman 1999; García-Sainz and Villalobos-Molina 2004; Hein and Michel 2007; García-Cazarín et al. 2008). This points to the possibility that potential differences in their efficacy result in different functional outcomes for each α_1 -AR subtype.

There are more observations that add complexity to this scenario. As previous results obtained by our research group indicate, native α_{1D} -AR remains active after removing the agonist (Noguera and D'Ocon 1993; Noguera et al. 1996; Gisbert et al. 2000, 2002, 2003b; Ziani et al. 2002) in vessels where this subtype play a functional role. They act as "*constitutively active*" receptors which maintain an increased vascular tone for some time after the adrenoceptor-mediated stimulus is removed (Ziani et al. 2002). This constitutive activity of α_{1D} -ARs has also been found in stably transfected rat fibroblasts and HEK293 cells where a α_{1D} -mediated pERK1/2 signal was observed

in the absence of an adrenoceptor-mediated stimulus (García-Sainz and Torres-Padilla 1999a; McCune et al. 2000; Chalothorn et al. 2002; Pérez-Aso et al. 2013).

We propose that the characteristic behavior of the α_{1D} subtype: higher sensitivity, sustained activity after removal of the agonist, and its presence in the poorly innervated conductance vessels, could determine a distinctive timecourse of the adrenoceptor-mediated contraction in these vessels, and permit them to respond to the circulating levels of catecholamines (rarely above 10 nmol/L) (Goldstein et al. 2003).

In the present work, we confirm this hypothesis by analyzing the characteristics of the response elicited by α_1 -ARs in two different vessels, aorta, a territory where the α_{1D} -AR subtype plays the main functional role, and tail artery as a vessel where the α_{1A} -AR subtype is the main one responsible of the adrenoceptor-mediated contractile response. Involvement of subtypes was manipulated by the use of selective antagonists in the rat and subtype knockouts in the mouse (α_{1B} -KO, α_{1D} -KO, and $\alpha_{1B/D}$ -KO). Signaling pathways were investigated alongside contractility studies by analyzing inositol phosphate accumulation and extracellular signal-regulated kinase (ERK)1/2 phosphorylation after adrenoceptor stimulus in the same vessels.

Materials and Methods

Animals

Thoracic aorta and tail artery were obtained as previously described (Gisbert et al. 2000) from male Wistar rats (200–250 g) from colonies of wild-type (WT) and α_{1D} -KO mice, kindly supplied by Professor Gozoh Tsujimoto (Department of Molecular Cell Pharmacology, National Research Institute for Child Health and Development, Tokyo), α_{1B} -KO mice, kindly supplied by Professor Susanna Cotecchia (Départment de Pharmacologie et de Toxicologie, Université de Lausanne, Switzerland), and $\alpha_{1B/D}$ -KO mice generated by crossing these α_{1D} -KO and α_{1B} -KO strains at University of Glasgow (see Methven et al. 2009a). All protocols complied with European Community guidelines for experimental animals and were approved by the Ethics Committee of the University of Valencia.

Functional studies

Rings obtained from rat vessels and mouse aorta, were denuded of endothelium by gentle rubbing and suspended in an organ bath. Tension was recorded isometrically according to the protocol previously described (Martí et al. 2005). Arterial rings from the mouse tail were mounted on an isometric wire myograph (J.P. Trading, Aarhus, Denmark) according to the procedure previously described (Martinez-Rivelles et al. 2012). All vessels were maintained in Krebs buffer, at 37° C and gassed with 95% O₂ and 5% CO₂.

An initial load of 9.81 mN was applied to each preparation and maintained throughout a 75–90 minutes equilibration period. The rings were stimulated with noradrenaline (NA) (10 μ mol/L in tail artery or 1 μ mol/L in aorta) which produced a maximal contraction. The lack (<10%) of a relaxant response to acetyl-choline (100 μ mol/L) in these precontracted preparations indicated the absence of a functional endothelium. After 30 minutes washout, contractile responses to NA were elicited according to different experimental procedures:

(a) Concentration-response curves (CRC) to NA. This experimental procedure was performed in each vessel by addition of cumulative concentrations of NA (0.0001–10 μ mol/L) until a maximal response was obtained. From these curves, pD₂ and Emax were calculated using a nonlinear regression plot (Graph Pad Software; San Diego, CA).

(b) Sustained contractile response to NA. The experimental procedure was performed according to previous studies (Noguera and D'Ocon 1993; Gisbert et al. 2000; Ziani et al. 2002). A maximal contractile response to NA (1 µmol/L in aorta, 10 µmol/L in tail artery) was obtained in Ca2+-containing medium; this concentration was maintained until a stable tone was reached and then washed. In some experiments, BMY 7378 or 5methylurapidil were added 15 minutes prior to NA addition. The washing procedure was carried out with a total replacement of the bathing solution by three repeated washes within the first 30 seconds and by two other repeated washes every 5 minutes in all cases. The tone was measured at different times during the development of the contractile response and after washing until total recovery of the basal tone. The results were expressed as percentages of maximal contractile responses.

(c) Increase in vascular tone after removal of the agonist. Vessels were incubated in a Ca²⁺-free solution (containing 0.1 mmol/L ethylenediaminetetraacetic acid, EDTA) for 20 minutes, which led to a small loss in tension (<10–15%); then, vessels were exposed to NA (1 μ mol/L in aorta, 10 μ mol/L in tail artery) twice, 10 minutes each time, the tissues being carefully washed between the two exposures, following the same procedure described above. A spontaneous increase in vascular tone was observed when the Ca²⁺-free solution was substituted by a Ca²⁺-containing medium. The effects of prazosin, 5-methylurapidil, and BMY 7378 were assessed on this spontaneous increase in tone. One micromolar of each antagonist was added during incubation in Ca^{2+} -free medium, 10 minutes before addition of Ca^{2+} -containing solution and was maintained during the Ca^{2+} -loading period. Contraction was expressed in mN.

Accumulation of [³H]-inositol phosphates

The determination of the accumulation of inositol phosphates (IPs) has been previously described (Gisbert et al. 2003b). Briefly, rat tail arteries or thoracic aortas were cut into rings, pooled, and submitted to different experimental procedures:

(a) Incubation for 30 minutes with increasing concentrations of NA (0.01 μ mol/L–0.1 mmol/L) in the presence of LiCl (10 mmol/L) in order to inhibit the metabolism of inositol monophosphates.

(b) Incubation for 30 minutes with 1 μ mol/L of prazosin, 5-methylurapidil, or BMY 7378, in the presence of LiCl (10 mmol/L) and in presence or absence of NA (1 μ mol/L in aorta, 10 μ mol/L in tail artery).

(c) Incubation for 30 minutes with NA (1 μ mol/L in aorta, 10 μ mol/L in tail artery) in Ca²⁺-free medium in the absence of LiCl to avoid accumulation of inositol phosphates, followed by removal of the agonist by careful washing, and incubation for 30 minutes in Ca²⁺-containing medium in the presence of LiCl (10 mmol/L).

At the end of the functional experiments all vessels were immediately frozen and processed as previously published (Monto et al. 2012) to obtain total proteins.

Accumulation of $[{}^{3}H]$ -IPs was routinely calculated as dpm of total $[{}^{3}H]$ -inositol labeled lipids/µg of protein in each individual sample. CRC for NA-induced $[{}^{3}H]$ -IPs accumulation were fitted by nonlinear regression plot (Graph Pad Software; San Diego, CA) and the pEC₅₀ and Emax values were obtained.

Determination by immunoblotting of NAmediated ERK1/2 activation

Rings of rat aorta or tail artery were loaded in tubes containing 5 mL of Krebs solution gassed with 95% O_2 and 5% CO_2 , at 37°C. After 30 minutes of stabilization in Ca^{2+} -containing medium, selective antagonists were added when indicated, and maintained for 15 minutes. Aorta and tail artery segments were then stimulated or not with NA for 5 minutes and then the tissues were immediately frozen by liquid N_2 immersion.

Protein extracts (50 μ g) were loaded onto 10% Sodium dodecyl sulphate-Polyacrylamide gels, and electrophoresed proteins were transferred to polyvinyldiene fluoride (PVDF) membranes 2 hours at 375 mA, using a liquid

Mini Trans-Blot[®] Electrophoretic Transfer Cell system (Bio-Rad Laboratories, Inc., S. A. Madrid, Spain). Membranes were blocked in albumin from bovine serum 3% in phosphate-buffered saline (PBS) containing 0.1% Tween 20 for 1 hour at room temperature with gentle agitation. Membranes were incubated overnight at 4°C with anti-phospho-p42/44 ERK MAPK (Thr202/Thr204) and anti-p42/44 ERK MAPK (1:500; Cell Signaling Technology, Beverly, MA). Membranes were then washed three times with PBS with 0.1% Tween 20, incubated with antirabbit immunoglobulin G horseradish peroxidase-linked whole antibody (1:2500; GE Healthcare, Buckinghamshire, U.K.) for 45 minutes at room temperature and washed extensively with phosphate buffered saline with tween before chemiluminescent detection was performed using the the ECL[™] Prime Western Blotting Detection Reagents (GE Healthcare, Buckinghamshire, U.K.). The image was captured with the AutoChemi System (Ultra-Violet Products Bioimaging Systems, Cambridge, U.K.) and band intensity was measured using LabWorks 4.6 Image acquisition and Analysis (Ultra-Violet Products Bioimaging Systems, Cambridge, U.K.).

Drugs and solutions

The following drugs were obtained from SIGMA (St. Louis, MO): (-)-NA, prazosin, BMY 7378 (8-[2-[4-(2-Methoxyphenyl)-1-piperazynil]-8-azaspiro [4,5]decane-7,9-dione dihydrochloride) and 5-methylurapidil. Other reagents were of analytical grade. All compounds were dissolved in distilled water. The composition of Krebs solution was (mmol/L) NaCl 118, KCl 4.75, CaCl₂ 1.8, MgCl₂ 1.2, KH₂PO₄ 1.2, NaHCO₃ 25, and glucose 11. Ca²⁺-free solution had the same composition except that CaCl₂ was omitted and EDTA (0.1 mmol/L) was added. The terminology for receptors employed is as recommended in Alexander et al. (2011).

Results

NA exhibits higher potency but lower efficacy in aorta than in tail artery

NA elicited a concentration-dependent contraction, ERK 1/2 phosphorylation, and [³H]-IPs accumulation in both rat aorta and tail artery (Fig. 1). The potency (pEC₅₀) of NA was higher in aorta than in tail artery, for all three measures; conversely, this increase in potency was accompanied by an apparently lower efficacy in IPs formation and ERK1/2 phosphorylation in aorta when normalized for protein content (Fig. 1A and B). It is not practicable to compare contractile efficacy between different vessels.

CRC of NA were also performed in aortic and tail artery rings from α_{1B} -KO, α_{1D} -KO, $\alpha_{1B/D}$ -KO, or WT mice and the results are shown in Figure 2. As in rat vessels, in WT mice the pEC₅₀ of NA was significantly higher in aorta than in tail artery (8.28 \pm 0.02 and 7.23 \pm 0.05, respectively, P < 0.001). Comparing α_{1D} -KO with WT, the pEC₅₀ of NA was reduced in aorta and tail artery, although the maximal response (Emax) was not different (Fig. 2) in either case. No significant difference in potency of NA was observed between aortic rings from WT and α_{1B} -KO mice although the Emax was significantly reduced. In $\alpha_{1B/D}$ -KO, a contractile response of aorta was detectable only with the three highest concentrations of NA and was so small that the pEC₅₀ of the CRC could not be calculated (Fig. 2A).

NA-induced contraction exhibits a slower time-course in aorta than in tail artery

The concentration of NA needed to obtain the maximal response (1 μ mol/L in aorta, 10 μ mol/L in tail artery from rat and WT mice) evoked a contraction with different time-course profile in each vessel (Figs. 3 and 4).

In order to clarify the role that each α_1 -AR subtype plays in the distinctive time-course of the adrenoceptormediated response observed in each vessel, we analyzed changes in this response in the presence of subtype selective α_1 -antagonists in rat vessels or in knockout mice vessels (α_{1B} -KO, α_{1D} -KO, $\alpha_{1B/D}$ -KO, or WT mice) with the objective of isolating the response to each receptor subtype.

We produced responses to single concentrations of NA (1 μ mol/L in rat aorta and 10 μ mol/L in tail artery) in the presence and absence of two selective antagonists, 5-methylurapidil, selective for α_{1A} -ARs and BMY 7378 selective for α_{1D} -AR (Koshimizu et al. 2002). The concentration of each antagonist was of the same order as its pA₂ and/or pK_B in each vessel (Gisbert et al. 2003a).

In rat aorta, the kinetic of the NA-induced contraction was not affected by 5-methylurapidil (Fig. 3A), nor was the recovery of the basal tone after removal of the agonist (Fig. 3B). In the presence of BMY 7378 a slightly faster contractile response was observed (Fig. 3C) followed by a faster recovery of basal tone after washing the tissue. As Figure 3D shows, without antagonist, 5 minutes after agonist removal 50% of the maximal response to NA remained. However, in the presence of BMY, 5 minutes after NA removal the vascular tone was only 20% of the maximal contraction, and reached the basal levels around 10 minutes versus 20 minutes in the absence of BMY 7378.

In aorta from α_{1D} -KO mice, the contractile response elicited by NA was faster than in WT and not sustained,



Figure 1. Concentration–response curves of noradrenaline (NA) on (A) p-ERK signaling pathway determined by immunoblotting. A representative immunoblot was also included, (B) inositol phosphates accumulation (IPs), (C) vascular tone expressed as force units (mN). Experiments were performed in aorta (white circles) and tail artery (black circles) from rat. Emax and pEC₅₀ of the concentration–response curves were included in each case. Values are represented as mean \pm SEM of n = 3–6 experiments. Statistical signification was calculated by Student's *t* test. **P* < 0.05, ****P* < 0.001.

with a slow decay after reaching its maximal response (Fig. 3E). In α_{1B} -KO and $\alpha_{1B/D}$ -KO mice, NA-induced contraction reached the maximal value more slowly than the contraction observed in aorta from WT (Fig. 3E). After removal of the agonist by washing, the return to the baseline was markedly slower in aorta from α_{1B} -KO or WT mouse than in aorta from α_{1D} -KO and $\alpha_{1B/D}$ -KO mouse (Fig. 3F).

In rat tail artery, no significant changes were observed in the time-course profile of the NA-induced contraction in presence of any drug (Fig. 4A–D). In tail artery from the α_{1D} -KO mouse the profile of the contractile response was similar to WT (Fig. 4E). The return to the baseline was only slightly slower in WT than in α_{1D} -KO mouse (Fig. 4F) confirming the minor role of the α_{1D} subtype in this vessel.



		Emax (mN)	pEC ₅₀
AORTA	WT	3.88 ± 0.04	8.28 ± 0.02
	α _{1B} -KO	2.88 ± 0.01*	8.00 ± 0.37
	α _{1D} -KO	3.64 ± 0.06	6.89 ± 0.06***
	α _{1B/D} ΚΟ	0.17 ± 0.01***	n.d.
TAIL	WT	8.24 ± 0.20	7.23 ± 0.05
	α _{1D} -KO	7.52 ± 0.13	6.70 ± 0.04***

Figure 2. Contractile responses to cumulative concentrations of noradrenaline NA in aorta (A) or tail artery (B) of wild-type (WT), α_{1D} -adrenoceptor knockout (α_{1D} -KO), α_{1B} -adrenoceptor knockout ($\alpha_{1B/C}$ -KO) and $\alpha_{1B/D}$ -adrenoceptor knockout ($\alpha_{1B/D}$ -KO) mice. Emax (expressed as mN) and pEC₅₀ of the concentration–response curves were included in each case. Values are represented as mean \pm SEM of n = 3–6 experiments. Statistical significance was calculated by Student's *t* test. **P* < 0.05, ****P* < 0.001, n.d.= not determined.

α₁-adrenoceptors exhibit activity after removal of the agonist in aorta but not in tail artery

Previous evidence indicates that cells expressing α_{1D} -ARs exhibit elevated basal levels of calcium [32] and of pERK (McCune et al. 2000; Chalothorn et al. 2002) which could be decreased by α_1 -AR antagonists including the nonsub-type-selective prazosin or the α_{1D} -AR-selective BMY 7378. In rat aorta, the basal level of pERK1/2 (in the absence of an adrenoceptor-mediated stimulus) was not significantly changed by BMY 7378 (1 μ mol/L) or prazosin (1 μ mol/

L). No changes in basal phosphorylation of ERK1/2 were observed in rat tail artery incubated with 5-methylurapidil (1 μ mol/L) or prazosin (1 μ mol/L) (Fig. 5, white bars). As expected, NA induced an increase in the phosphorylation of ERK1/2 in both vessels, which was inhibited by prazosin and BMY 7378 in aorta and by prazosin and 5-methylurapidil in tail artery (Fig. 5, black bars).

The incubation with selective antagonists did not change the basal IPs levels in rat aorta, nor in rat tail artery (Fig. 6, white bars). Addition of NA (1 μ mol/L in aorta, 10 μ mol/L in tail artery) induced a marked increase in the IPs accumulation which was completely inhibited by prazosin (1 μ mol/L) and by BMY 7378 (1 μ mol/L) or 5-methylurapidil (1 μ mol/L) in aorta or tail artery, respectively (Fig. 6, black bars).

Following the experimental procedure described in the methods section, we had previously observed in aorta but not in tail artery a population of active α_{1D} -ARs that increases IPs accumulation after removal of the agonist (Gisbert et al. 2003b). The same results were obtained in the present work. Figure 6 (gray bars) quantifies the magnitude of the IPs accumulation in aorta observed after agonist removal, and shows that incubation with prazosin (1 μ mol/L) or BMY 7378 (1 μ mol/L) inhibits it. Different results were observed in tail artery. No increase in IPs was observed after removal of the agonist and incubation with prazosin (1 μ mol/L) did not modify IPs levels (Fig. 6, gray bars).

None of the antagonists assayed modified the basal tone of the rat aorta or tail artery (Fig. 7, white bars) and, as expected, addition of NA to the bath chamber promoted a sustained increase in tone that was almost completely inhibited in presence of prazosin or either BMY 7378 or 5-methylurapidil (Fig. 7, black bars). After careful removal of the agonist and following the experimental procedure previously described, a spontaneous increase in the vascular tone was observed in aorta but not in tail artery (Fig. 7, gray bars). Incubation with prazosin (1 μ mol/L) or BMY 7378 (1 μ mol/L) inhibits this increase in tone observed after agonist removal, as has been previously shown (Gisbert et al. 2000, 2003b).

In order to elucidate if an inadequate washing or tissue peculiarity rather than a special activity of one α_1 -AR subtype could explain the increase in tone observed after NA removal, we performed the same experimental procedure in vessels from knockout mice. Experiments in aorta from WT and α_{1B} -KO mice show a spontaneous increase in tone after NA-removal similar to that found in rat aorta. However, a similar increase in tone was not observed in aorta from α_{1D} -KO or $\alpha_{1D}/_{1B}$ -KO mice after NA-removal (Fig. 8). The spontaneous increase in tone



Figure 3. Time-course of the contractile response to noradrenaline in aorta from rat or transgenic mice. The magnitude of the contraction was determined at different times after addition of NA (1 μ mol/L) to the bath chamber (A, C, and E) or after removal of the agonist (B, D, and F), in absence (control) or presence of selective antagonists (5-methylurapidil 0.1 μ mol/L and BMY7378 0.01 μ mol/L) or in different mouse strains: wild type (WT), α_{1D} -adrenoceptor knockout (α_{1D} -KO), α_{1B} -adrenoceptor knockout (α_{1B} -KO), and $\alpha_{1B/D}$ -adrenoceptor knockout ($\alpha_{1B/D}$ -KO). Values represent mean \pm SEM of n = (3–6) experiments.

observed in WT mouse was completely inhibited by prazosin (1 μ mol/L) and BMY 7378 (1 μ mol/L), but not by 5-methylurapidil (1 μ mol/L) (Fig. 8B).

Discussion and Conclusions

The major findings of the present study are that the α_{1D} subtype is responsible of the greater contractile sensitivity, slower time-course and postactivation contraction to an adrenoceptor-mediated stimulus in conductance vessels. This conclusion can be drawn from differences observed in the contractile response to NA between a conductance artery (aorta) and distributing artery (tail artery) in two species, the receptor subtype being isolated pharmacologically in the rat and by receptor subtype knockout in the mouse.

Conducting arteries respond to adrenoceptor-mediated stimulus with higher sensitivity than distributing vessels

Rat aorta exhibited a higher sensitivity for NA than tail artery and this difference was observed independently of the signaling pathway analyzed: IPs accumulation, ERK1/ 2 phosphorylation, or contractile response. Interestingly when a measure of maximal response representing efficacy was calculated this was not greater in aorta so a generally "larger" response signal per se is not implicated.

It is well known that rat, mouse, and human aorta express protein of the three α_1 -ARs subtypes but, in functional terms, the vasoconstrictor role of α_{1D} is predominant (Kenny et al. 1995; Hussain and Marshall 1997;



Figure 4. Time-course of the contractile response to NA in tail artery from rat or transgenic mice. The magnitude of the contraction was determined at different times after addition of NA (1 μ mol/L) to the bath chamber (A, C and E) or after removal of the agonist (B, D, and F), in absence (control) or presence of selective antagonists (5-methylurapidil 0.01 μ mol/L and BMY7378 0.1 μ mol/L) or in different mouse strains: wild type (WT), α_{1D} -adrenoceptor knockout (α_{1D} -KO), α_{1B} -adrenoceptor knockout ($\alpha_{1B/D}$ -adrenoceptor knockout ($\alpha_{1B/D}$ -KO). Values represent mean \pm SEM of n = (3–6) experiments.

Gisbert et al. 2000, 2002, 2003a; Yamamoto and Koike 2001; Hosoda et al. 2005), whereas in rat tail artery the α_{1A} -subtype is mainly implicated in adrenoceptor-mediated contraction (Gisbert et al. 2003a; Martí et al. 2005; Docherty 2010). Thus, the greater sensitivity to NA observed in aorta could be attributed to the main functional role played by α_{1D} -AR in this vessel. This proposal was based on previous reports showing a higher potency of NA and adrenaline on cloned α_{1D} -ARs expressed in different cell lines (Theroux et al. 1996; Pérez-Aso et al. 2013), as well as aorta and other conducting arteries where the α_{1D} subtype plays a main functional role (Daly et al. 2002; Tanoue et al. 2002; Deighan et al. 2005; Hosoda et al. 2005; Methven et al. 2009a,b).

However, in native tissues, changes in potency of the agonists could be also explained by structural or cellular characteristics of vessels independent of the α_1 subtype involved. The present results obtained with gene-targeted mice confirmed the higher sensitivity of the native α_{1D} subtype to NA as responsible for the higher potency exhibited by the agonist in aorta since, in the mouse model lacking the α_{1D} -AR (α_{1D} -KO), the pEC₅₀ of NA





Figure 5. Basal and noradrenaline (NA)-induced phosphorylation of ERK1/2 in rat aorta and tail artery. When indicated, vessels were incubated or not with NA (10 µmol/L in aorta and tail artery) for 5 minutes, in absence or presence of selective ligands as prazosin (PRAZ), BMY 7378 (BMY), and 5-methylurapidil (5-MeU) at 1 µmol/L. After stimulation, cellular extracts were prepared as described under the methods section. Equal amounts (50 µg) of each sample were used to visualize the ERK1/2 expression (upper panels). The lower panels show equal amounts of ERK1/2 loaded on each sample. Bar graphics represents the quantification of basal (white bars) or NA-induced (black bars) ERK1/2 phosphorylation. Values represent means \pm SEM of 3–4 independent experiments. Statistics was performed by the Dunnett's test **P* < 0.05 versus NA.

was significantly lower than that observed in aorta from α_{1B} -KO or WT mice; $\alpha_{1B/D}$ -KO compared with α_{1B} -KO gave a similar result. Moreover, the much smaller difference in potency and efficacy of CRC to NA observed in tail artery between α_{1D} -KO and WT mice suggests a lesser role for α_{1D} -ARs in this vessel; an earlier study using α_{1B} -KO showed also a lesser role for α_{1B} -AR in this artery (Daly et al. 2002).

Figure 6. Inositol phosphates accumulation determined in rat aorta and tail artery in basal conditions (white bars), after addition of noradrenaline (NA) (black bars) and after addition and careful removal of NA (gray bars), according to the experimental procedure described in Methods. The experiments were performed in absence or presence of the selective ligands prazosin (PRAZ), BMY 7378 (1 µmol/L) (BMY), and 5-methylurapidil (5-MeU) at 1 µmol/L. Values represent means \pm SEM of 3–4 independent experiments. Statistics was performed by the Dunnett's test **P* < 0.05, ***P* < 0.01, ****P* < 0.001 versus noradrenaline or noradrenaline removal. NA, noradrenaline 1 µmol/L in aorta, and 10 µmol/L in tail artery.

After removal of the adrenoceptormediated stimulus, the contractile response disappears more slowly in conducting than in distributing arteries

Aorta from rat or mouse exhibits a distinctive timecourse in the contractile response to an adrenoceptormediated stimulus. The time-course profile of this response is characterized by a slow decay in the contractile tone when the agonist was removed. We have previously described a similar time-course in other conducting vessels such as iliac and proximal mesenteric arteries, where α_{1D} -AR plays a functional role (Ziani et al. 2002).





Figure 7. Changes in the vascular tone observed in rat aorta and tail artery in basal conditions (white bars), after addition of noradrenaline (black bars) and after addition and careful removal of noradrenaline (gray bars) according to the experimental procedure described in the methods section. The experiments were performed in absence or presence of the selective ligands prazosin (PRAZ), BMY 7378 (BMY), and 5-methylurapidil (5-MeU) at 1 μ mol/L. Values represent means \pm SEM of 4–5 independent experiments. Statistical analysis was performed by the Dunnett's test: ****P*<0.001 to test the effects of antagonists versus noradrenaline or noradrenaline removal. NA, noradrenaline 1 μ mol/L in aorta and 10 μ mol/L in tail artery.

On the contrary, in distributing vessels such as tail artery, or resistance vessels such as small mesenteric branches, where the α_{1D} -AR has not a predominant role, and the α_{1A} -AR is the main subtype involved, this slow time-course is not observed, and a fast decay in the contractile tone after agonist removal was observed (Ziani et al. 2002). Therefore, we can attribute the slower timecourse profile to the presence of the α_{1D} -subtype in a vessel but also it could be due to structural differences between arteries. The use of selective antagonists in rat aorta as well as studies in vessels from mice lacking the α_{1D} -subtype confirms the involvement of this receptor in the distinctive time-course observed in conducting vessels.

Figure 8. (A) Representative tracings of the changes in tone observed in aorta of wild-type (WT) and α_{1D} -knockout (α_{1D} -KO) mice after addition of NA (1 μ mol/L) in a calcium-free medium, subsequent removal of the agonist and washing (W), and subsequent incubation in a calcium-containing solution. Arrows show the spontaneous increase in tone observed in aorta from WT but not from α_{1D} -KO mouse. (B) Quantification of the spontaneous increase in tone observed in aorta from WT mice incubated with or without the selective ligands prazosin (PRAZ), BMY 7378 (BMY), and 5-methylurapidil (5-MeU) at 1 μ mol/L (black bars), and in α_{1B} -KO, α_{1D} -KO, and $\alpha_{1B/D}$ -KO mice (gray bars). Values represent means \pm SEM of 4–5 independent experiments. Statistics was performed by the Dunnett's test, ***P < 0.001 to test the effect of antagonists versus WT

A selective antagonist of the α_{1D} -ARs, BMY 7378, but not the α_{1A} selective antagonist 5-methylurapidil (Michelotti et al. 2000; Koshimizu et al. 2002), affects to a great extent the recovery of basal tone after agonist removal. This difference was even more evident in knockout mice. In strains where the α_{1D} -AR was not expressed (α_{1D} -KO and $\alpha_{1B/D}$ -KO), the recovery of basal tone was almost complete 5 minutes after agonist washing whereas it takes up 30 minutes in WT and α_{1B} -KO mice. In tail artery from all strains, the decay in this maximal response to NA was faster than in aorta from rats or WT mice, and similar to aorta from α_{1D} -KO and $\alpha_{1B/D}$ -KO mouse. In conclusion, after removal of the agonist, a faster decay in the contractile tone was observed in aorta from α_{1D} -KO and $\alpha_{1B/D}$ -KO mice versus WT and α_{1B} -KO, and this time-course profile of the adrenoceptor-mediated contraction is similar to that observed in distributing vessels such as tail artery. Therefore, a consequence of α_{1D} -ARs activation in conducting vessels is a sustained contractile response when the stimulus disappears.

The next question that arises was the possible involvement of the constitutive activity of α_{1D} -ARs in this sustained response. It has been reported that cloned α_{1D} -ARs exhibit constitutive activity, evidenced by increased levels of calcium (García-Sainz and Torres-Padilla 1999a) or pERK1/2 (McCune et al. 2000) that were selectively inhibited by prazosin or BMY 7378, acting as inverse agonists.

After incubation with prazosin or BMY 7378, no change was registered in p-ERK1/2 IPs or contractile tone in aorta, which suggests that constitutive activity observed in cloned α_{1D} -ARs is not so evident in native receptors and has not a relevant impact on the signaling pathway or in the vascular tone. Therefore, there is no evidence for the presence of a population of constitutively active α_{1D} -AR, coupled to p-ERK signal with a modulator role on the basal vascular tone in aorta.

Interestingly, after activation by NA and removal of the agonist, the α_{1D} -ARs continue actively coupled to the IP pathway and, at the same time, we observed a temporary increase in vascular tone when NA was no longer present in the bath. The experiments performed in knockout mice confirmed this peculiarity of α_{1D} -ARs as the increased tone which appears after agonist removal was observed only in WT and α_{1B} -KO mouse, but not in α_{1D} -KO or $\alpha_{1B/D}$ -KO mice.

Recent evidences indicate that α_{1D} -ARs are expressed as a multiprotein complex at the plasma membrane (Lyssand et al. 2008, 2010) interacting with the syntrophin family through a PSD95/DlgA/Zo-1 (PDZ)-domain (Chen et al. 2006). Syntrophin isoforms play selective roles in the α_{1D} -AR/dystrophin-associated protein complex signalosome as α -syntrophin increases α_{1D} -AR binding site density while β_2 -syntrophin enhances α_{1D} -AR coupling to downstream signaling effectors (Lyssand et al. 2011). In addition, this signaling complex is not mimicked by the α_{1A} or α_{1B} -AR subtypes which suggest that it could be involved in the peculiar activity exhibited by the α_{1D} subtype.

The characteristic behavior of the α_{1D} -AR has been previously reported (Gisbert et al. 2000, 2003b; Ziani et al. 2002) as constitutive activity which manifests only after agonist stimulation and removal. Thus, in native vascular smooth muscle, the α_{1D} -ARs remain constitutively active after agonist activation, and maintain the adrenoceptormediated response when the agonist is removed; finally, α_{1D} -ARs are internalized and the vessel recovers the basal tone. However, this activity observed after removal of the agonist could be also attributed to a prolonged binding of NA to α_{1D} -ARs which activates them for a while. As we have discussed in previous papers, this explanation does not hold (Gisbert et al. 2000, 2003b) but in any case, the more interesting result is not related to the fact that the activity showed by α_{1D} -ARs after removal of the agonist was "truly" or only "apparent" constitutive activity. The more interesting question is the physiological consequence of this activity which explains the slower decay observed in the adrenoceptor-mediated response when the agonist is removed.



Figure 9. Schematic picture showing that preponderance of the most sensitive α_{1D} -AR subtype in noninnervated conductance arteries such as aorta, allows responsiveness to physiological levels of circulating catecholamines. The activity showed by this subtype after agonist removal sustains the contractile tone and prevents abrupt changes in vessel caliber when the stimulus fluctuates. In innervated distributing arteries, high local concentrations of NA are required to activate the less sensitive subtype of α_{1A} -adrenoceptors which elicit a response that is rapid but short lived, allowing fine adjustment of the contractile tone by perivascular sympathetic nerves.

In fact, aortas from α_{1D} -KO and $\alpha_{1B/D}$ -KO mice, which did not exhibit a spontaneous increase in tone after agonist removal, had a time-course profile of recovery of the basal tone faster than aorta from WT and α_{1B} -KO mouse, and similar to tail artery from any strain.

Physiological and therapeutic relevance of α_{1D}-ARs in conductance vessels

The present results show that, in response to a systemic adrenoceptor-mediated stimulus, a poorly innervated conductance vessel such as aorta, where α_{1D} -AR is the main functionally relevant subtype, responds with higher sensitivity than a vessel where the α_{1A} -AR subtype is dominant, as occurs in tail artery. Thus, plasma levels of catecholamines, which rarely exceed 10 nmol/L (Goldstein et al. 2003) could induce a moderate adrenoceptor-mediated response in conductance vessels. This contractile response can be temporarily sustained when the agonist is removed, due to the constitutive coupling of the α_{1D} -subtype to IPs/contraction pathway after agonist removal, and this mechanism would prevent abrupt changes in the caliber of conductance arteries when the adrenoceptor-mediated stimulus fluctuates.

However, the higher threshold for α_{1A} -adrenoceptors present in tail artery might take them out of the reach of circulating levels of catecholamines. Thus the α_{1A} -ARs might require the high local concentrations produced only by release of noradrenaline from perivascular nerves. In this case, the response is fast and intense, and disappears when the stimulus does. This mechanism would permit the fine adjustment of the contractile tone of distributing vessels by the local nervous stimulus and consequently the precise adjustment of blood flow. This concept is consistent with the hypothesis of Stassen et al. (1998) that α_{1A} -adrenoceptors are present in blood vessels only when adrenergic nerves are present and might add the further idea that α_{1A} -ARs are activated physiologically only or mainly by nerves (Daly et al. 2002).

The differential role exhibited by the α_{1D} and the α_{1A} subtypes, present in conductance or distributing and resistance vessels, respectively, opens new lines of pharmacological research looking for the selective modulation of a given subtype as a more vessel selective, accurate, and safe strategy to control vascular tone.

In conclusion, as Figure 9 depicts, high sensitivity to agonist and persistence of response after agonist removal is a property of α_{1D} -adrenoceptors. Therefore, the preponderance of this subtype in noninnervated conductance arteries such as aorta allows responsiveness to circulating catecholamines and prevents abrupt changes in vessel caliber when the stimulus fluctuates. Conversely, in innervated distributing arteries, high local concentrations of

NA are required to activate α_{1A} -adrenoceptors for a response that is rapid but short lived allowing fine adjustment of the contractile tone by perivascular sympathetic nerves.

Acknowledgements

This study was supported by research grants to the University of Glasgow from the UK Medical Research Council (G0000042, I.D.51,240) and to the University of Valencia from the Instituto de Salud Carlos III Fondo de Investigaciones Sanitarias (FIS PI070509), Ministerio de Ciencia e Innovacion, SAF(2007-62,120), Fondos FEDER, Generalitat Valenciana (GVACOMP2009/261) and Research funds of the Universitat de Valencia (UV-INV-AE11-42,176).

Disclosures

None declared.

References

Alexander S, Mathie A, Peters J (2011). Guide to receptors and channels (GRAC), 5th edn. Br. J. Pharmacol. 164: S1–S324.

Arévalo-León LE, Gallardo-Ortiz IA, Urquiza-Marin H, Villalobos-Molina R (2003). Evidence for the role of alpha1Dand alpha1A-adrenoceptors in contraction of the rat mesenteric artery. Vascul. Pharmacol. 40: 91–96.

Cavalli A, Lattion AL, Hummler E, Nenniger M, Pedrazzini T, Aubert JF, et al. (1997). Decreased blood pressure response in mice deficient of the alpha1b-adrenergic receptor. Proc. Natl. Acad. Sci. USA 94: 11589–11594.

Chalothorn D, McCune DF, Edelmann SE, Garcia-Cazarin M, Tsujimoto G, Piascik M (2002). Differences in the cellular localization and agonist-mediated internalization properties of the alpha(1)-adrenoceptor subtypes. Mol. Pharmacol. 61: 1008–1016.

Chen Z, Hague C, Hall RA, Minneman KP (2006). Syntrophins regulate α_{1D} adrenergic receptors through a PDZ domain-mediated interaction. J. Biol. Chem. 281: 12414–12420.

Cotecchia S (2010). The alpha1-adrenergic receptors: diversity of signaling networks and regulation. J. Recept. Signal Transduct. Res. 30: 410–419.

Daly CJ, Deighan C, McGee A, Mennie D, Ali Z, McBride M, et al. (2002). A knockout approach indicates a minor vasoconstrictor role for vascular alpha1B-adrenoceptors in mouse. Physiol. Genomics 9: 85–91.

Deighan C, Methven L, Naghadeh MM, Wokoma A, Macmillan J, Daly CJ, et al. (2005). Insights into the functional roles of alpha(1)-adrenoceptor subtypes in mouse

carotid arteries using knockout mice. Br. J. Pharmacol. 144: 558–565.

Docherty JR (2010). Subtypes of functional alpha1-adrenoceptor. Cell. Mol. Life Sci. 67: 405–417.

García-Cazarín ML, Smith JL, Olszewski KA, McCune DF, Simmerman LA, Hadley RW, et al. (2008). The alpha1D-adrenergic receptor is expressed intracellularly and coupled to increases in intracellular calcium and reactive oxygen species in human aortic smooth muscle cells. J. Mol. Signal. 3: 6.

García-Sainz JA, Torres-Padilla ME (1999a). Modulation of basal intracellular calcium by inverse agonists and phorbol myristate acetate in rat-1 fibroblasts stably expressing alpha1d-adrenoceptors. FEBS Lett. 443: 277–281.

García-Sainz JA, Villalobos-Molina R (2004). The elusive alpha (1D)-adrenoceptor: molecular and cellular characteristics and integrative roles. Eur. J. Pharmacol. 500: 113–120.

García-Sainz JA, Vázquez-Prado J, Villalobos-Molina R (1999b). Alpha 1-adrenoceptors: subtypes, signaling, and roles in health and disease. Arch. Med. Res. 30: 449–458.

Gisbert R, Noguera MA, Ivorra MD, D'Ocon P (2000). Functional evidence of a constitutively active population of alpha(1D)-adrenoceptors in rat aorta. J. Pharmacol. Exp. Ther. 295: 810–817.

Gisbert R, Ziani K, Miquel R, Noguera MA, Ivorra MD, Anselmi E, et al. (2002). Pathological role of a constitutively active population of alpha(1D)-adrenoceptors in arteries of spontaneously hypertensive rats. Br. J. Pharmacol. 135: 206– 216.

Gisbert R, Madrero Y, Sabino V, Noguera MA, Ivorra MD, D'Ocon P (2003a). Functional characterization of alpha 1-adrenoceptor subtypes in vascular tissues using different experimental approaches: a comparative study. Br. J. Pharmacol. 138: 359–368.

Gisbert R, Pérez-Vizcaino F, Cogolludo AL, Noguera MA, Ivorra MD, Tamargo J, et al. (2003b). Cytosolic Ca2+ and phosphoinositide hydrolysis linked to constitutively active alpha 1D-adrenoceptors in vascular smooth muscle. J. Pharmacol. Exp. Ther. 305: 1006–1014.

Goldstein DS, Eisenhofer G, Kopin IJ (2003). Sources and significance of plasma levels of catechols and their metabolites in humans J. Pharmacol. Exp. Ther. 305: 800–811.

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.

Hein P, Michel M (2007). Signal transduction and regulation: are all alpha1-adrenergic receptor subtypes created equal? Biochem. Pharmacol. 73: 1097–1106.

Hosoda C, Tanoue A, Shibano M, Tanaka Y, Hiroyama M, Koshimizu TA, et al. (2005). Correlation between

vasoconstrictor roles and mRNA expression of alpha1-adrenoceptor subtypes in blood vessels of genetically engineered mice. Br. J. Pharmacol. 146: 456–466.

Hrometz SL, Edelmann SE, McCune DF, Olges JR, Hadley RW, Perez DM, et al. (1999). Expression of multiple alpha1-adrenoceptors on vascular smooth muscle: correlation with the regulation of contraction. J. Pharmacol. Exp. Ther. 290: 452–463.

Hussain MB, Marshall I (1997). Characterization of alpha1-adrenoceptor subtypes mediating contractions to phenylephrine in rat thoracic aorta, mesenteric artery and pulmonary artery. Br. J. Pharmacol. 122: 849–858.

Keffel S, Alexandrov A, Goepel M, Michel MC (2000). alpha (1)-adrenoceptor subtypes differentially couple to growth promotion and inhibition in Chinese hamster ovary cells. Biochem. Biophys. Res. Commun. 272: 906–911.

Kenny BA, Chalmers DH, Philpott PC, Naylor AM (1995). Characterization of an alpha 1D-adrenoceptor mediating the contractile response of rat aorta to noradrenaline. Br. J. Pharmacol. 115: 981–986.

Koshimizu TA, Yamauchi J, Hirasawa A, Tanoue A, Tsujimoto G (2002). Recent progress in alpha 1-adrenoceptor pharmacology. Biol. Pharm. Bull. 25: 401–408.

Lachnit WG, Tran AM, Clarke DE, Ford AP (1997). Pharmacological characterization of an alpha 1A-adrenoceptor mediating contractile responses to noradrenaline in isolated caudal artery of rat. Br. J. Pharmacol. 120: 819–826.

Lyssand JS, DeFino MC, Tang XB, Hertz AL, Feller DB, Wacker JL, et al. (2008). Blood pressure is regulated by an alpha1D-adrenergic receptor/dystrophin signalosome. J. Biol. Chem. 283: 18792–18800.

Lyssand JS, Whiting JL, Lee KS, Kastl R, Wacker JL, Bruchas MR, et al. (2010). Alpha-dystrobrevin-1 recruits alpha-catulin to the alpha1D-adrenergic receptor/dystrophin-associated protein complex signalosome. Proc. Natl Acad. Sci. USA 107: 21854–21859.

Lyssand JS, Lee KS, DeFino M, Adams ME, Hague C (2011). Syntrophin isoforms play specific functional roles in the alpha1D-adrenergicreceptor/DAPC signalosome. Biochem. Biophys. Res. Commun. 412: 596–601.

Martí D, Miquel R, Ziani K, Gisbert R, Ivorra MD, Anselmi E, et al. (2005). Correlation between mRNA levels and functional role of alpha1-adrenoceptor subtypes in arteries: evidence of alpha1L as a functional isoform of the alpha1A-adrenoceptor. Am. J. Physiol. Heart Circ. Physiol. 289: H1923–H1932.

Martinez-Rivelles S, Caracuel L, Marquez-Martin A, Dantas AP, Oliver E, D'Ocon P, et al. (2012). Increased endothelin-1 vasoconstriction in mesenteric resistance arteries after superior mesenteric ischemia-reperfusion. Br. J. Pharmacol. 165: 937–950.

McCune DF, Edelmann SE, Olges JR, Post GR, Waldrop BA, Waugh DJ, et al. (2000). Regulation of the cellular localization and signaling properties of the alpha(1B)- and alpha (1D)-adrenoceptors by agonists and inverse agonists. Mol. Pharmacol. 57: 659–666.

Methven L, McBride M, Wallace GA, McGrath JC (2009a). The alpha 1B/D-adrenoceptor knockout mouse permits isolation of the vascular alpha 1A-adrenoceptor and elucidates its relationship to the other subtypes. Br. J. Pharmacol. 158: 209–224.

Methven L, Simpson PC, McGrath JC (2009b). Alpha1A/ B-knockout mice explain the native alpha1D-adrenoceptor's role in vasoconstriction and show that its location is independent of the other alpha1-subtypes. Br. J. Pharmacol. 158: 1663–1675.

Michelotti GA, Price DT, Schwinn DA (2000). Alpha 1-adrenergic receptor regulation: basic science and clinical implications. Pharmacol. Ther. 88: 281–309.

Monto F, Oliver E, Vicente D, Rueda J, Aguero J, Almenar L, et al. (2012). Different expressions of adrenoceptors and GRKs in the human myocardium depend on heart failure etiology and correlate with clinical variables. Am. J. Physiol. Heart Circ. Physiol. 303: H368–H376. PMID:22685168

Noguera MA, D'Ocon P (1993). Evidence that depletion of internal stores sensitive to noradrenaline elicits a contractile response dependent on extracellular calcium in rat aorta. Br. J. Pharmacol. 110: 861–867.

Noguera MA, Ivorra MD, D'Ocon P (1996). Functional evidence of inverse agonism in vascular smooth muscle. Br. J. Pharmacol. 119: 158–164.

Pérez-Aso M, Segura V, Montó F, Barettino D, Noguera MA, Milligan G, et al. (2013). The three α_1 -adrenoceptor subtypes show different spatio-temporal mechanisms of internalization and ERK1/2 phosphorylation. Biochim. Biophys. Acta 1833: 2322–2333.

Philipp M, Hein L (2004). Adrenergic receptor knockout mice: distinct functions of 9 receptor subtypes. Pharmacol. Ther. 101: 65–74.

Piascik MT, Perez DM (2001). Alpha1-adrenergic receptors: new insights and directions. J. Pharmacol. Exp. Ther. 298: 403–410.

Piascik MT, Guarino RD, Smith MS, Soltis EE, Saussy DL Jr, Perez DM (1995). The specific contribution of the novel alpha-1D adrenoceptor to the contraction of vascular smooth muscle. J. Pharmacol. Exp. Ther. 275: 1583–1589. Rudner XL, Berkowitz DE, Booth JV, Funk BL, Cozart KL, D'Amico EB, et al. (1999). Subtype specific regulation of human vascular alpha(1)-adrenergic receptors by vessel bed and age. Circulation 100: 2336–2343.

Schwinn DA, Page SO, Middleton JP, Lorenz W, Liggett SB, Yamamoto K, et al. (1991). The alpha 1C-adrenergic receptor: characterization of signal transduction pathways and mammalian tissue heterogeneity. Mol. Pharmacol. 40: 619–626.

Stassen FR, Maas RG, Schiffers PM, Janssen GM, De Mey JG (1998). A positive and reversible relationship between adrenergic nerves and alpha-1A adrenoceptors in rat arteries. J. Pharmacol. Exp. Ther. 284: 399–405.

Taguchi K, Yang M, Goepel M, Michel MC (1998). Comparison of human alpha1-adrenoceptor subtype coupling to protein kinase C activation and related signalling pathways. Naunyn Schmiedebergs Arch. Pharmacol. 357: 100–110.

Tanaka T, Zhang L, Suzuki F, Muramatsu I (2004). Alpha-1 adrenoceptors: evaluation of receptor subtype-binding kinetics in intact arterial tissues and comparison with membrane binding. Br. J. Pharmacol. 141: 468–476.

Tanoue A, Koshimizu TA, Tsujimoto G (2002). Transgenic studies of alpha(1)-adrenergic receptor subtype function. Life Sci. 71: 2207–2215.

Tanoue A, Koshimizu TA, Shibata K, Nasa Y, Takeo S, Tsujimoto G (2003). Insights into alpha1 adrenoceptor function in health and disease from transgenic animal studies. Trends Endocrinol. Metab. 14: 107–113.

Theroux TL, Esbenshade TA, Peavy RD, Minneman KP (1996). Coupling efficiencies of human alpha 1-adrenergic receptor subtypes: titration of receptor density and responsiveness with inducible and repressible expression vectors. Mol. Pharmacol. 50: 1376–1387.

Yamamoto Y, Koike K (2001). alpha(1)-Adrenoceptor subtypes in the mouse mesenteric artery and abdominal aorta. Br. J. Pharmacol. 134: 1045–1054.

Zhong H, Minneman KP (1999). Differential activation of mitogen-activated protein kinase pathways in PC12 cells by closely related alpha1-adrenergic receptor subtypes. J. Neurochem. 72: 2388–2396.

Ziani K, Gisbert R, Noguera MA, Ivorra MD, D'Ocon P (2002). Modulatory role of a constitutively active population of alpha(1D)-adrenoceptors in conductance arteries. Am. J. Physiol. Heart Circ. Physiol. 282: H475–H481.