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Chen, Xiaofeng; Rateri, Debra L.; Howatt, Deborah A.; Balakrishnan, Anju; Moorleghen, Jessica J.; Morris, Andrew J.; Charnigo, Richard; Cassis, Lisa A.; and Daugherty, Alan, "Amlodipine Reduces AngII-Induced Aortic Aneurysms and Atherosclerosis in Hypercholesterolemic Mice" (2013). Saha Cardiovascular Research Center Faculty Publications. 9. https://uknowledge.uky.edu/cvrc_facpub/9

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Notes/Citation Information

Published in PLOS One, v. 8, issue. 11, e81743.

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Digital Object Identifier (DOI)

http://dx.doi.org/10.1371/journal.pone.0081743

Amlodipine Reduces Angll-Induced Aortic Aneurysms and Atherosclerosis in Hypercholesterolemic Mice

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Abstract

Background: The purpose of this study was to determine effects of amlodipine, a dihydropyridine calcium channel blocker, on development of angiotensin II (AngII)-induced vascular pathologies.

Methods and Results: Male LDL receptor -/- mice were infused with vehicle, amlodipine (5 mg/kg/d), AngII (1,000 ng/kg/min), or AngII + amlodipine for 4 weeks through osmotic pumps (n=10/group). Mice were fed a saturated fatenriched diet for 1 week prior to pump implantation and during 4 weeks of infusion. Infusion of amlodipine resulted in plasma concentrations of 32 ± 2 ng/ml and 27 ± 2 ng/ml for mice in saline + amlodipine and AngII + amlodipine groups, respectively. This infusion rate of amlodipine did not affect AngII-induced increases in systolic blood pressure. Three of 10 (30%) mice infused with AngII died of aortic rupture, while aortic rupture did not occur in mice co-infused with AngII + amlodipine. Suprarenal aortic width and intimal area of ascending aortas were measured to define aortic aneurysms. In the absence of AngII infusion, amlodipine did not change suprarenal aortic width and ascending aortic area. Infusion of AngII led to profound increases of suprarenal aortic width (saline + vehicle versus AngII + vehicle: 0.86 ± 0.02 versus 1.72 ± 0.26 mm; P=0.0006), whereas co-infusion of AngII and amlodipine diminished abdominal dilation (1.02 ± 0.14 mm; P=0.003). As expected, AngII infusion increased mean intimal area of ascending aortas (saline + vehicle versus AngII + vehicle: 8.5 ± 0.3 versus 12.5 ± 1.1 mm²; P=0.001), while co-infusion of AngII and amlodipine ablated dilation of the ascending aorta (8.6 ± 0.2 mm²; P=0.03). Co-administration of amlodipine also significantly attenuated AngII-induced atherosclerosis in the thoracic region as quantified by percent lesion area (AngII + vehicle versus AngII + amlodipine 5.8 ± 2.1 % versus $0.3 \pm 0.1\%$; P=0.05).

Conclusions: Amlodipine inhibited AngII-induced aortic aneurysms in both the abdominal and ascending regions, and atherosclerosis in hypercholesterolemic mice.

Citation: Chen X, Rateri DL, Howatt DA, Balakrishnan A, Moorleghen JJ, et al. (2013) Amlodipine Reduces AnglI-Induced Aortic Aneurysms and Atherosclerosis in Hypercholesterolemic Mice. PLoS ONE 8(11): e81743. doi:10.1371/journal.pone.0081743

Editor: Elena Aikawa, Brigham and Women's Hospital, Harvard Medical School, United States of America

Received June 28, 2013; Accepted October 24, 2013; Published November 14, 2013

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Funding: This study was supported by a funding from the National Institutes of Health (R01 HL107319 to AD) and a McKusick Fellowship grant from the National Marfan Foundation to XC. Mass Spectrometry equipment used for plasma amlodipine measurement was provided by funds from the National Center for Research Resources (P20 RR021954) and the National Institute of General Medical Sciences (P20 GM103527). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing interests: The authors have declared that no competing interests exist.

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Introduction

Aortic aneurysmal and atherosclerotic diseases may occur coincidently in humans [1-5]. Excessive stimulation of the renin angiotensin system (RAS) has been implicated in many human cardiovascular diseases, including aortic aneurysms and atherosclerosis [6,7]. There are substantial experimental data that excessive RAS activation also promotes these diseases in animal models. AnglI infusion into LDL receptor -/-, apoE -/- or normolipidemic mice induces development of aneurysms in the suprarenal region of the abdominal aorta [8-11]. More recently,

it has been noted that AngII infusion also promotes pronounced dilation of the thoracic aorta, which is restricted to the ascending region [12,13]. In addition to promoting formation of aortic aneurysms, many studies have reported that chronic subcutaneous infusion of AngII into mice augments development of atherosclerosis in both apoE -/- and LDL receptor -/- mice [8,9,14].

Since AngII infusion into hypercholesterolemic mice promotes diverse aortic pathologies, this model permits simultaneous determination of effects of an intervention on abdominal and thoracic aortic aneurysms as well as atherosclerosis. Several studies have demonstrated that interventions can have differential effects on abdominal aortic aneurysms and atherosclerosis. These include pharmacological interventions with doxycycline, bosentan, and pioglitazone [15-17], and surgical interventions such as orchidectomy [18]. Genetic manipulations have also provided divergent effects including whole body genetic deletion of interferon-gamma, CXCL10, receptor-associated protein, or Rag-1 [19-21], and cell-specific deletion of PPAR gamma or AT1a receptors [17,22]. In contrast, several interventions have similar effects on these pathologies including pharmacological inhibition or whole body genetic deficiency of AT1a receptor [23,24], an AT2 receptor antagonist [23] and whole body deficiency of CCR2 [12]. Whole body deficiency of AT1a receptor or CCR2 also decreased AnglI-induced thoracic aortic aneurysms [12,22]. Findings from these studies implicate that the aortic pathologies in aneurysms and atherosclerosis may be attributed to a combination of common and distinctive mechanisms.

Anti-hypertensive drugs, including calcium channel blockers, are commonly prescribed for hypertensive patients who frequently also have aortic aneurysmal or atherosclerotic diseases. Although one study indicates that this class of drugs may have detrimental effects on abdominal aortic aneurysms in humans [25], calcium channel blockade has been shown to decrease atherosclerosis in humans [25-30]. However, this has not been determined in prospective studies. The purpose of this study was to determine effects of a commonly used calcium channel blocker, amlodipine, on aortic pathologies in a mouse model that develops aortic aneurysms and atherosclerosis simultaneously.

Materials and Methods

Ethics Statement

Recommendations from the Guide for the Care and Use of Laboratory Animals (National Institutes of Health) were followed. All procedures were approved by the University of Kentucky Institutional Animal Care and Use Committee (Protocol # 2006-0009). Mice were examined daily for signs of distress, dehydration, lack of weight gain, or bilateral hind leg paralysis. No adverse effects were noted during this study. Mice were terminated by overdose of a ketamine/xylazine mixture.

Mice

Male low-density lipoprotein (LDL) receptor -/- mice (stock # 002207; > N10 background into the C57BL/6 background) were purchased from the Jackson Laboratory (Bar Harbor, ME, USA). Mice were housed in individually ventilated cages with negative air pressure (Allentown Inc; Allentown, NJ, USA). Filtered drinking water by reverse osmosis system was provided ad libitum. Normal mouse diet (Global 18% protein rodent diet; Diet # 2918; Harlan Teklad; Madison, WI, USA) was fed. Light and dark cycle of the room was 14 hours of light and 10 hours of dark. Ambient temperature ranged from 20 to 23°C and humidity was 50-60%. Necropsy was performed on mice that died during AngII infusions within 12 hours of

expiration. Systolic blood pressure was measured by a tail cuff based technique using Coda 8 machines (Kent Scientific, Torrington, CT, USA) as described previously [31].

Diet and Drug Administration

To induce hypercholesterolemia, mice were fed a diet enriched in saturated fat (milk fat 21% wt/wt) and 0.2% wt/wt cholesterol (Diet # TD.88137; Harlan Teklad) 1 week prior to osmotic mini-pump implantation and throughout the 4 weeks of infusions.

Four groups (n=10 per group) of mice were infused with: 1. saline + vehicle [50% dimethyl sulfoxide (DMSO)]; 2. saline + amlodipine (5 mg/kg/day); 3. Angll (1,000 ng/kg/min) + vehicle (50% DMSO); and 4. Angll + amlodipine. For osmotic pump preparation (Durect Corporation; Cupertino, CA, USA), AngII (1,000 ng/kg/min; Cat # H-1705; Bachem; Torrance, CA, USA) was dissolved in saline. Angll and saline were infused using Alzet model 1004. Amlodipine besylate (5 mg/kg/day; Cat # A5605; Sigma-Aldrich; St. Louis, MO, USA) was dissolved in 50% DMSO. Amlodipine and vehicle (50% DMSO) were infused using Alzet model 2004. After anesthesia with ketamine/xylazine (90 and 10 mg/kg body weight, respectively), a Model 1004 pump and a Model 2004 pump were implanted subcutaneously into the flanks of each mouse [9]. A topical analgesic, LMX4 (Ferndale Laboratories; Ferndale, MI, USA), was used to provide relief from pain associated with surgery.

Plasma Measurements

Plasma cholesterol concentrations were measured using a commercially available enzymatic kit (Cat # Cholesterol E 439-17501; Wako Chemicals; Richmond, VA, USA) as described previously [32]. Plasma lipoprotein fractions were resolved by size exclusion chromatography and cholesterol concentrations were measured using the cholesterol enzymatic kit as described previously [32].

Plasma amlodipine concentrations were measured by HPLC electrospray ionization tandem mass spectrometry using adaptation of a previously reported method [33]. Briefly, individual plasma samples (50 µl) were extracted with ethyl ether: dichloromethane (70:30). After centrifugation, the supernatant was evaporated to dryness. Dried samples were reconstituted in a mixture of ammonium formate/ methanol/ acetonitrile (30:50:20; 50 µl). Samples (10 µl) were analyzed using a Shimadzu automated HPLC system with an Agilent Eclipse XDB C18 column (Agilent; Santa Clara, CA, USA) coupled with an AB Sciex 4000-Qtrap hybrid linear ion trap triple quadrupole mass spectrometer (AB Sciex; Framingham, MA. USA) that was operated in multiple reaction monitoring mode. Tizanidine (Cat # T6950; Sigma Aldrich) was used as an internal standard. Quantitation was accomplished by reference to an offline calibration generated using amlodipine that was independently guantitated by accurate mass measurements.

Plasma renin concentrations were measured as described previously [34]. Briefly, individual plasma samples (8 μ l) harvested with EDTA (1.8 mg/ml) were incubated with an excess of rat angiotensinogen at 37 °C for 30 minutes. The generated Angl was quantified by radioimmunoassay using a



Figure 1. Plasma concentrations of amlodipine. Plasma concentrations of amlodipine were measured at the end of study. Circles and triangles represent individual mice, diamonds are means of groups, and error bars are SEMs. * denotes P<0.0001 comparing amlodipine groups with saline + vehicle and AngII + vehicle groups by two way ANOVA. doi: 10.1371/journal.pone.0081743.g001

commercially available kit (Cat # 1553; DiaSorin; Stillwater, MN, USA).

Quantification of Aortic Aneurysms and Atherosclerosis

Aortas were dissected from the root to iliac bifurcation and placed in 10% neutrally buffered formalin overnight. After removal of adventitia, aortas were pinned and photographed. Maximal ex vivo diameters of suprarenal aortas were measured with ImagePro Plus software (Media Cybernetics; Rockville, MD, USA) to quantitate AAAs [11,35]. Subsequently, aortas were cut open longitudinally, secured with pins, and photographed. To determine formation of thoracic aortic aneurysms, dilation of ascending aortas was quantified by measurement of intimal area of the ascending aortic region [12]. Atherosclerosis was quantified on the intimal surface of the aorta by an en face technique as described previously [36,37].

Histological and Immunohistochemical Staining

Aortic tissues were fixed with 4% paraformaldehyde, and then sectioned on a cryostat at a thickness of 10 µm. Sections were histologically stained using Verhoeff's hematoxylin to visualize the integrity of elastin. Immunohistochemical staining was performed as described previously [38]. Smooth muscle cells were detected using smooth muscle α -actin antibody (2 µg/ml; Abcam, cat# ab5694) and macrophages using an CD68 antibody (5 µg/ml; Serotec, cat# MCA1957GA). Positively reactive areas were visualized via application of an ABC kit (Vector) and subsequent detection with AEC chromogen (Vector). Images were captured using a Nikon Eclipse E600 scope and a Nikon DXM1200F digital camera.

Statistical Analyses

Data for continuous variables are summarized as means \pm standard error of means (SEM). Version 9.2 of SAS (SAS Institute, Cary NC) was used for data analysis (SAS Institute Inc.; Cary, NC, USA). Effects of AngII and amlodipine were assessed by two-way analysis of variance (ANOVA) if normality and equal variance assumptions were satisfied (or if violations of these assumptions could be addressed by transforming the data or reweighting the observations) and were assessed posthoc by nonparametric Kruskal-Wallis test otherwise. P≤0.05 was considered significantly different.

Results

Characteristics of Study Mice

Amlodipine was infused subcutaneously through osmotic pumps to maintain a constant plasma concentration throughout the study. Plasma concentrations of amlodipine were not significantly different between the groups that were co-infused with either saline or AnglI (32 \pm 2 ng/ml and 27 \pm 2 ng/ml, respectively), while the drug was not detectable in saline + vehicle or AngII + vehicle infused mice (Figure 1). Amlodipine administration had no effect on body weight, plasma cholesterol concentrations, and plasma lipoprotein-cholesterol distributions in mice infused with either saline or AngII (Table 1). Amlodipine also had no significant effect on systolic blood pressure in mice infused with AnglI (150 ± 3 versus 148 ± 4 mmHg for AngII + vehicle versus AngII + amlodipine-infused mice, respectively). Angll infusion significantly decreased plasma renin concentration (P=0.005); however, addition of amlodipine significantly increased plasma renin concentration (P<0.0001; Table 1).



Figure 2. Survival curve for study mice. The survival curve for the four groups of mice during the infusion interval. Mice infused with AngII + vehicle had 3 deaths (occurred on days 3, 4, and 8) due to aortic rupture, while no deaths occurred in the other groups. doi: 10.1371/journal.pone.0081743.g002

Table 1. Characteristics of study mice.

Infusions	Saline		Angli	
	Vehicle	Amlodipine	Vehicle	Amlodipine
n	10	10	7*	10
Body weight (g)	28.0 ± 0.5	27.6 ± 0.5	26.9 ± 0.6	26.9 ± 0.5
Plasma cholesterol (mg/dl)	1038 ± 41	949 ± 41	1168 ± 49	1119 ± 41
Plasma renin (ng/ml/30 min)	3.0 ± 0.3	8.5 ± 1.3 [†]	1.1 ± 0.1 [#]	5.2 ± 1.1 ^{†#}
Body weight, plasma cholesterol and renin concentrations were measured after 28				
days of infusions. Values are represented as mean \pm SEM. * Ten mice were				
initiated in the study, but 3 succumbed to aortic rupture, so data were derived from				
7 surviving mice. [†] P<0.0001, compared to vehicle within saline or AngII groups. [#]				

P=0.005, compared to saline infusion within vehicle or amlodipine groups.

doi: 10.1371/journal.pone.0081743.t001

Amlodipine Prevented Development of Angll-induced Aneurysms in Both Abdominal and Ascending Aortic Regions

Three of 10 mice infused with AngII died of aortic rupture within 10 days after initiating AngII infusion, while no aortic ruptures occurred in the other 3 groups as shown in Figure 2.

To determine effects of amlodipine on development of AAAs, maximal diameters of suprarenal aortas were measured ex vivo. Amlodipine had no effect on suprarenal aortic width in saline-infused mice. As described previously, infusion of AngII led to profound increases of maximal aortic width (saline + vehicle versus AngII + vehicle: 0.86 ± 0.02 versus 1.72 ± 0.26 mm; P=0.0006). Amlodipine administration significantly attenuated AngII-induced abdominal aortic dilation (1.02 ± 0.14 mm; P=0.003; Figure 3). Aortas from AngII-infused mice administered with amlodipine were grossly normal as demonstrated by histological analysis (Figure 4). To determine

whether any pathological changes were discernible, aortic tissues were sectioned and immunohistochemically and histologically stained. As the example shown in Figure 4, suprarenal aortas from amlodipine infused mice were grossly normal with consistent smooth muscle α -actin immunostaining, intact elastin fibers in the media, and no accumulation of macrophages.

En face measurement of intimal surface area of the ascending aorta was used as an index to determine ascending aortic dilation [12]. Administration of amlodipine had no effect on ascending aortic area in saline-infused mice. Angll infusion significantly increased mean intimal area of ascending aortas (saline + vehicle versus Angll + vehicle: 8.5 ± 0.3 versus $12.5 \pm 1.1 \text{ mm}^2$; P=0.001). Co-infusion of Angll with amlodipine ablated Angll-induced ascending aortic dilation ($8.6 \pm 0.2 \text{ mm}^2$; P=0.03; Figure 5). As with the suprarenal aorta, infusion of amlodipine led to preservation of a normal histological appearance of the ascending aorta (Figure 4).

Amlodipine Reduced Atherosclerosis in AnglI-infused Hypercholesterolemic Mice

Atherosclerotic lesion size was only minor in mice infused with saline + vehicle and saline + amlodipine after 5 weeks of feeding the saturated fat-enriched diet. Consistent with previous studies [9,39], mice infused with AngII had significantly increased lesion area in the thoracic aorta (saline + vehicle versus AngII + vehicle: 0.22 ± 0.18 % versus 5.76 ± 2.29 %; P=0.02; Figure 6). Co-infusion of amlodipine reduced atherosclerosis in AngII-infused mice (0.27 ± 0.14 %; P=0.05).

Discussion

Angll infusion simultaneously induces aneurysms in the suprarenal and ascending regions of the aorta and augments



Figure 3. Amlodipine ablated AnglI-induced abdominal aortic aneurysms. *Ex vivo* suprarenal aortic diameter was measured after termination. Circles and triangles are individual measurements, diamonds means of the groups, and error bars SEMs. Representative images of aortas are shown below the graph. Statistical analyses were performed using nonparametric Kruskal-Wallis test. * denotes P=0.0006 for comparison between AngII + vehicle and saline + vehicle, and # denotes P=0.003 for comparison between AngII + amlodipine and AngII + vehicle groups. doi: 10.1371/journal.pone.0081743.g003

atherosclerosis in hypercholesterolemic mice, as demonstrated in the present and previous studies [9,12,14,40]. Despite markedly different vascular pathologies promoted by chronic Angll infusion, the present study provides evidence that continuous infusion of a dihydropyridine calcium channel blocker, amlodipine, has pronounced effects on inhibition of these diverse pathologies that developed in mice rendered hypercholesterolemia by deletion of LDL receptor and feeding a diet that has a high saturated fat content (42% by calories), compared to normal laboratory diet (18% fat by calories).

Amlodipine has commonly been administrated by gavage, mixed with diet, or dissolved in drinking water [41-46]. In the present study, amlodipine was given using osmotic minipumps, as used in a previous study [47]. This approach was used to provide stable calcium channel blockade throughout the 28 days of AngII infusion. In addition to providing a mode of constant drug delivery, we also measured plasma concentrations of this drug in mouse plasma. The detected concentrations (mean concentration of saline + amlodipine and AngII + amlodipine: 29.3 ± 1.7 ng/ml) are above the reported EC50 of amlodipine [48,49]. By comparison to humans, a single dose of 10 mg amlodipine to healthy male volunteers resulted in plasma concentrations of 5.9 ± 1.2 ng/ml. Repeated daily dosing led to mean plasma concentrations of 14.5 ± 5.8

ng/ml with peaks and troughs of 18.1 ± 7.1 and 11.8 ± 5.3 ng/ml, respectively [50,51]. Therefore, plasma concentrations of amlodipine achieved in this mouse study were in a similar range to those achieved in humans.

Previous studies have demonstrated that increased systolic blood pressure observed during AngII infusion is not the determinant of aortic aneurysmal formation or atherosclerosis augmentation [14,40,52,53]. In agreement with these previous reports, the present study also found that amlodipine reduced all vascular pathologies without resulting in any measurable change in systolic blood pressure. This result extends the findings by Kanematsu et al. [54], who reported that amlodipine reduced both systolic blood pressure and aortic aneurysmal formation in normocholesterolemic mice.

We also found that amlodipine administration increased plasma renin concentrations. Previous studies suggest that amlodipine has biphasic effects on plasma renin activity in rodents, with reductions at low doses and stimulation of renin at higher doses [55]. The basis for amlodipine exerting these effects on plasma renin concentrations has not been defined. In our experience, plasma renin concentrations in mice are inversely related to plasma concentrations of AngII [24,56,57], as also demonstrated in the present study by pronounced reductions in plasma renin concentrations in AngII-infused



Figure 4. Amlodipine preserved normal aortic structure in the suprarenal and ascending regions. Tissue sections from AngII-infused mice also infused with amlodipine were immunostained with alpha actin and CD68 to detect smooth muscle cells and macrophages, respectively. Positive reactivity is show by intense red color. Verhoeff's hematoxylin staining was performed to facilitate visualization of elastin.

doi: 10.1371/journal.pone.0081743.g004

mice. Therefore, the observed elevations in plasma renin concentration by amlodipine suggest a reduction in endogenous production of AngII. This may have contributed to the ability of amlodipine to reduce vascular pathologies.

Although both abdominal and ascending aortic aneurysms are defined by pronounced aortic luminal expansion, pathologies in these two locations are distinct [12,22,58-60]. In AnglI infused mice, abdominal aortic aneurysms appear to be initiated by rapid medial macrophage accumulation co-localized with focal elastin fragmentation, and these changes are followed by rapid luminal expansion and marked leukocyte infiltration in the adventitia [61,62]. In contrast, during the initiation of AnglI-induced ascending aortic aneurysms, one of the earliest pathological changes is hemorrhage restricted to the outer medial layers, and subsequent concentric medial changes and luminal dilation, while leukocyte infiltration is not as pronounced as in abdominal aortic aneurysms [60]. In spite of significant differences of these two aortic aneurysms, the present study demonstrated that amlodipine reduced aortic expansion in both abdominal and ascending regions. In agreement with our findings, a recent study reported that amlodipine profoundly reduced incidence of abdominal and thoracic aortic aneurysms in normocholesterolemic mice coadministered either Angll or deoxycorticosterone acetate-salt with a lysyl oxidase inhibitor, beta-aminopropionitrile [54]. This

model creates profound dilation of a translucent aorta from the middle of the ascending region to aortic orifice of the subclavian arterial branch. Therefore, this pathology differs from that formed during AngII infusion in which dilation is restricted to the full length of the ascending aorta and the media is thickened and opaque [12]. Despite differences in gross appearance, amlodipine markedly decreased thoracic aortic pathologies in both animal models. Currently, information from humans with ascending aortic aneurysms is limited to a small study inferring benefits in young patients with Marfan's syndrome who were administered a non-dihydropyridine calcium channel blocker, verapamil [63].

Calcium channel blockade has also been studied in a rat model of abdominal aortic aneurysms induced by intra-aortic infusion of elastase. Two different members of the dihydropyridine class of calcium channel blockers, azelnidipine and nifedipine, reduced AAAs formed in this model [64,65]. In contrast to these reports and what was found in the present study, a retrospective investigation reported that calcium channel blockade was associated with the presence of abdominal aortic aneurysms. However, administration of calcium channel blockers did not affect aortic dilation rate. The authors acknowledged potential compounding factors in this retrospective analysis and noted the need for prospective studies [25]. Currently, two prospective studies are evaluating



Figure 5. Amlodipine attenuated AnglI-induced ascending aortic aneurysms. Ascending aortic areas were measured using an en face method. Circles and triangles are individual measurements, diamonds are means of groups, and error bars are SEMs. Representative images of aortas are shown below the graph. Statistical analyses were performed using nonparametric Kruskal-Wallis test. * denotes P=0.001 for comparison between AnglI + vehicle and saline + vehicle groups, and # denotes P=0.03 for comparison between AngII + amlodipine and AngII + vehicle groups. doi: 10.1371/journal.pone.0081743.g005

effects of amlodipine on growth of AAAs in humans (NCT01425242 and NC01118520). Completion of these studies will provide insight into the validity of translating results from AngII-induced AAAs in mice to humans.

In contrast to the limited information on aortic aneurysms, there have been many experimental and human studies to determine effects of multiple calcium channel blockers on atherosclerosis [66]. Several of these studies have investigated effects of amlodipine in mouse atherosclerosis models with either apoE deletion or mutation. These reports provided a range of results with amlodipine reducing atherosclerotic lesion size [41,42,46,67-69] or having no effect [43-45,70]. In the present study, we used saturated fat-fed LDL receptor -/- mice infused with AngII, and found that amlodipine decreased atherosclerosis. The conflicting findings in the literature may be partially explained by different doses of amlodipine used and mode of drug delivery; however, potential mechanisms of the conflicting findings remain to be unraveled.

The overt differences in the appearance of AnglI-induced abdominal aortic aneurysms, ascending aortic aneurysms, and

atherosclerosis imply that the octapeptide is stimulating several mechanisms to produce these diverse pathologies. Therefore, the profound effects of amlodipine on all these pathologies also imply that the drug is potentially interfering with many mechanisms. Amlodipine has been demonstrated to influence many processes that have been invoked in AngII-induced vascular pathologies include reductions in oxidant stress [41], matrix metalloproteinases [71], nitric oxide synthesis [72], and leukocyte migration [47]. However, the contribution of each of these mechanisms to be determined.

In conclusion, administration of amlodipine was studied in a mouse model that simultaneous developments aortic aneurysms and atherosclerosis. Although several studies have observed disparities in response to different AngII-induced vascular pathologies, this study demonstrated that continuous infusion of amlodipine, at a dose that achieved therapeutically relevant plasma concentrations, markedly reduced all these pathologies.



Figure 6. Amlodipine reduced AnglI-induced atherosclerosis. Percent atherosclerotic lesion area of thoracic aortas were measured by an en face method. Circles and triangles are individual measurements, diamonds are means of the groups, and error bars are SEMs. Statistical analyses were performed using nonparametric Kruskal-Wallis test. * denotes P=0.02 for comparison between AngII + vehicle and saline + vehicle groups, and # denotes P=0.05 for comparison between AngII + amlodipine and AngII + vehicle groups.

doi: 10.1371/journal.pone.0081743.g006

Acknowledgements

We acknowledge Sunkara Manjula for mass spectrometry measurements of plasma amlodipine, and Vicki English for plasma renin measurements. We thank Dr. Hong Lu for assistance in editing the manuscript.

Author Contributions

Conceived and designed the experiments: XC AD. Performed the experiments: XC DR DAH AB JM. Analyzed the data: XC

References

- Bengtsson H, Sonesson B, Bergqvist D (1996) Incidence and prevalence of abdominal aortic aneurysms, estimated by necropsy studies and population screening by ultrasound. Ann NY Acad Sci 800: 1-24. doi:10.1111/j.1749-6632.1996.tb33294.x. PubMed: 8958978.
- Homme JL, Aubry MC, Edwards WD, Bagniewski SM, Shane Pankratz V et al. (2006) Surgical pathology of the ascending aorta: a clinicopathologic study of 513 cases. Am J Surg Pathol 30: 1159-1168. PubMed: 16931961.
- Duncan JL, Harrild KA, Iversen L, Lee AJ, Godden DJ (2012) Long term outcomes in men screened for abdominal aortic aneurysm: prospective cohort study. BMJ 344: e2958. doi:10.1136/bmj.e2958. PubMed: 22563092.
- Golledge J, Muller J, Daugherty A, Norman P (2006) Abdominal aortic aneurysm. Pathogenesis and implications for management. Arterioscler Thromb Vasc Biol 26: 2605-2613. doi:10.1161/01.ATV. 0000245819.32762.cb. PubMed: 16973970.
- Alcorn HG, Wolfson SK Jr, Sutton-Tyrrell K, Kuller LH, O'Leary D (1996) Risk factors for abdominal aortic aneurysms in older adults enrolled in The Cardiovascular Health Study. Arterioscler Thromb Vasc Biol 16: 963-970. doi:10.1161/01.ATV.16.8.963. PubMed: 8696960.
- Daugherty A, Lu H, Rateri DL, Cassis LA (2008) Augmentation of the renin-angiotensin system by hypercholesterolemia promotes vascular diseases. Future Lipidol 3: 625-636. doi:10.2217/17460875.3.6.625. PubMed: 19802341.

DR DAH AB JM AJM RC LAC AD. Contributed reagents/ materials/analysis tools: AJM AD. Wrote the manuscript: XC DR DAH AB JM AJM RC LAC AD.

- Rader DJ, Daugherty A (2008) Translating molecular discoveries into new therapies for atherosclerosis. Nature 451: 904-913. doi:10.1038/ nature06796. PubMed: 18288179.
- Daugherty A, Cassis L (1999) Chronic angiotensin II infusion promotes atherogenesis in low density lipoprotein receptor -/- mice. Ann NY Acad Sci 892: 108-118. doi:10.1111/j.1749-6632.1999.tb07789.x. PubMed: 10842656.
- Daugherty A, Manning MW, Cassis LA (2000) Angiotensin II promotes atherosclerotic lesions and aneurysms in apolipoprotein E-deficient mice. J Clin Invest 105: 1605-1612. doi:10.1172/JCI7818. PubMed: 10841519.
- Deng GG, Martin-McNulty B, Sukovich DA, Freay A, Halks-Miller M et al. (2003) Urokinase-type plasminogen activator plays a critical role in angiotensin II-induced abdominal aortic aneurysm. Circ Res 92: 510-517. doi:10.1161/01.RES.0000061571.49375.E1. PubMed: 12600880.
- Uchida HA, Poduri A, Subramanian V, Cassis LA, Daugherty A (2011) Urokinase-type plasminogen activator deficiency in bone marrowderived cells augments rupture of angiotensin II-induced abdominal aortic aneurysms. Arterioscler Thromb Vasc Biol 31: 2845-2852. doi: 10.1161/ATVBAHA.111.234997. PubMed: 21868698.
- 12. Daugherty A, Rateri DL, Charo IF, Owens AP, Howatt DA et al. (2010) Angiotensin II infusion promotes ascending aortic aneurysms:

attenuation by CCR2 deficiency in apoE-/- mice. Clin Sci (Lond) 118: 681-689. doi:10.1042/CS20090372.

- Rateri DL, Moorleghen JJ, Knight V, Balakrishnan A, Howatt DA et al. (2012) Depletion of endothelial or smooth muscle cell-specific angiotensin II type 1a receptors does not influence aortic aneurysms or atherosclerosis in LDL receptor deficient mice. PLOS ONE 7: e51483. doi:10.1371/journal.pone.0051483. PubMed: 23236507.
- Weiss D, Kools JJ, Taylor WR (2001) Angiotensin II-induced hypertension accelerates the development of atherosclerosis in apoEdeficient mice. Circulation 103: 448-454. doi:10.1161/01.CIR. 103.3.448. PubMed: 11157699.
- Manning MW, Cassis LA, Daugherty A (2003) Differential effects of doxycycline, a broad-spectrum matrix metalloproteinase inhibitor, on angiotensin II-induced atherosclerosis and abdominal aortic aneurysms. Arterioscler Thromb Vasc Biol 23: 483-488. doi: 10.1161/01.ATV.0000058404.92759.32. PubMed: 12615694.
- Suen RS, Rampersad SN, Stewart DJ, Courtman DW (2011) Differential Roles of Endothelin-1 in Angiotensin II-Induced Atherosclerosis and Aortic Aneurysms in Apolipoprotein E-Null Mice. Am J Pathol 179: 1549-1559. doi:10.1016/j.ajpath.2011.05.014. PubMed: 21718678.
- Subramanian V, Golledge J, Ijaz T, Bruemmer D, Daugherty A (2010) Pioglitazone-induced reductions in atherosclerosis occur via smooth muscle cell-specific interaction with PPAR{gamma}. Circ Res 107: 953-958. doi:10.1161/CIRCRESAHA.110.219089. PubMed: 20798360.
- Henriques TA, Huang J, D'Souza SS, Daugherty A, Cassis LA (2004) Orchidectomy, but not ovariectomy, regulates angiotensin II-induced vascular diseases in apolipoprotein E-deficient mice. Endocrinology 145: 3866-3872. doi:10.1210/en.2003-1615. PubMed: 15105380.
- King VL, Lin AY, Kristo F, Anderson TJ, Ahluwalia N et al. (2009) Interferon-gamma and the interferon-inducible chemokine CXCL10 protect against aneurysm formation and rupture. Circulation 119: 426-435. doi:10.1161/CIRCULATIONAHA.108.785949. PubMed: 19139386.
- Wang S, Subramanian V, Lu H, Howatt DA, Moorleghen JJ et al. (2012) Deficiency of receptor-associated protein attenuates angiotensin II-induced atherosclerosis in hypercholesterolemic mice without influencing abdominal aortic aneurysms. Atherosclerosis 220: 375-380. doi:10.1016/j.atherosclerosis.2011.11.013. PubMed: 22153700.
- Uchida HA, Kristo F, Rateri DL, Lu H, Charnigo R et al. (2010) Total lymphocyte deficiency attenuates AnglI-induced atherosclerosis in males but not abdominal aortic aneurysms in apoE deficient mice. Atherosclerosis 211: 399-403. doi:10.1016/j.atherosclerosis. 2010.02.034. PubMed: 20362292.
- Rateri DL, Moorleghen JJ, Balakrishnan A, Owens AP 3rd, Howatt DA et al. (2011) Endothelial cell-specific deficiency of Ang II type 1a receptors attenuates Ang II-induced ascending aortic aneurysms in LDL receptor-/- mice. Circ Res 108: 574-581. doi:10.1161/ CIRCRESAHA.110.222844. PubMed: 21252156.
- Daugherty A, Manning MW, Cassis LA (2001) Antagonism of AT2 receptors augments angiotensin II-induced abdominal aortic aneurysms and atherosclerosis. Br J Pharmacol 134: 865-870. doi:10.1038/sj.bjp. 0704331. PubMed: 11606327.
- 24. Cassis LA, Rateri DL, Lu H, Daugherty A (2007) Bone marrow transplantation reveals that recipient AT1a receptors are required to initiate angiotensin II-induced atherosclerosis and aneurysms. Arterioscler Thromb Vasc Biol 27: 380-386. PubMed: 17158350.
- Wilmink AB, Vardulaki KA, Hubbard CS, Day NE, Ashton HA et al. (2002) Are antihypertensive drugs associated with abdominal aortic aneurysms? J Vasc Surg 36: 751-757. doi:10.1016/ S0741-5214(02)00129-5. PubMed: 12368736.
- 26. Lichtlen PR, Hugenholtz PG, Raffenbeul W, Hecker H, Jost S et al. (1990) Retardation of angiographic progression of coronary artery disease by nifedipine: Results of International nifedipine Trial on Antiatherosclerotic therapy (INTACT). Lancet 1: 1109-1113.
- Waters D, Lespérance J, Frncetish M, Causey D, Théroux P et al. (1990) A controlled clinical trial to assess the effect of a calcium channel blocker on the progression of coronary atherosclerosis. Circulation 82: 1940-1953. doi:10.1161/01.CIR.82.6.1940. PubMed: 2242520.
- Zanchetti A, Bond MG, Hennig M, Neiss A, Mancia G et al. (2002) Calcium antagonist lacidipine slows down progression of asymptomatic carotid atherosclerosis: principal results of the European Lacidipine Study on Atherosclerosis (ELSA), a randomized, double-blind, longterm trial. Circulation 106: 2422-2427. doi:10.1161/01.CIR. 0000039288.86470.DD. PubMed: 12417537.
- Pitt B, Byington RP, Furberg CD, Hunninghake DB, Mancini GBJ et al. (2000) Effect of amlodipine on the progression of atherosclerosis and

the occurrence of clinical events. Circulation 102: 1503-1510. doi: 10.1161/01.CIR.102.13.1503. PubMed: 11004140.

- Nissen SE, Tuzcu EM, Libby P, Thompson PD, Ghali M et al. (2004) Effect of antihypertensive agents on cardiovascular events in patients with coronary disease and normal blood pressure: the CAMELOT study: a randomized controlled trial. JAMA 292: 2217-2225. doi: 10.1001/jama.292.18.2217. PubMed: 15536108.
- Daugherty A, Rateri D, Lu H, Balakrishnan A (2009) Measuring blood pressure in mice using volume pressure recording, a tail-cuff method. J Vis Exp 1291: ([MedlinePgn:]) PubMed: 19488026.
- Daugherty A, Rateri DL (1994) Presence of LDL receptor-related protein/alpha 2-macroglobulin receptors in macrophages of atherosclerotic lesions from cholesterol- fed New Zealand and heterozygous Watanabe heritable hyperlipidemic rabbits. Arterioscler Thromb 14: 2017-2024. doi:10.1161/01.ATV.14.12.2017. PubMed: 7526898.
- 33. Bhatt J, Shah B, Kambli S, Subbaiah G, Singh S et al. (2007) Rapid and sensitive method for the determination of sibutramine active metabolites in human plasma by reversed-phase liquid chromatography-tandem mass spectroscopy. J Chromatogr Sci 45: 91-96. doi:10.1093/chromsci/45.2.91. PubMed: 17425138.
- Cassis LA (1992) Downregulation of the renin-angiotensin system in streptozotocin-diabetic rats. Am J Physiol, 262: E105-E109. PubMed: 1733240.
- 35. Owens AP 3rd, Rateri DL, Howatt DA, Moore KJ, Tobias PS et al. (2011) MyD88 deficiency attenuates angiotensin II-induced abdominal aortic aneurysm formation independent of signaling through toll-like receptors 2 and 4. Arterioscler Thromb Vasc Biol 31: 2813-2819. doi: 10.1161/ATVBAHA.111.238642. PubMed: 21960563.
- Daugherty A, Whitman SC (2003) Quantification of atherosclerosis in mice. Methods Mol Biol 209: 293-309. PubMed: 12357958.
- Daugherty A, Rateri DL (2005) Development of experimental designs for atherosclerosis studies in mice. Methods 36: 129-138. doi:10.1016/ j.ymeth.2004.11.008. PubMed: 15893934.
- Lu H, Rateri DL, Daugherty A (2007) Immunostaining of mouse atherosclerosis lesions. Methods Mol Med 139: 77-94. doi: 10.1007/978-1-59745-571-8_4. PubMed: 18287665.
- Findeisen HM, Gizard F, Zhao Y, Cohn D, Heywood EB et al. (2011) Telomerase deficiency in bone marrow-derived cells attenuates angiotensin II-induced abdominal aortic aneurysm formation. Arterioscler Thromb Vasc Biol 31: 253-260. doi:10.1161/ATVBAHA. 110.218545. PubMed: 21088250.
- Rateri DL, Howatt DA, Moorleghen JJ, Charnigo R, Cassis LA et al. (2011) Prolonged infusion of angiotensin II in apoE(-/-) mice promotes macrophage recruitment with continued expansion of abdominal aortic aneurysm. Am J Pathol 179: 1542-1548. doi:10.1016/j.ajpath. 2011.05.049. PubMed: 21763672.
- Yoshii T, Iwai M, Li Z, Chen R, Ide A et al. (2006) Regression of atherosclerosis by amlodipine via anti-inflammatory and anti-oxidative stress actions. Hypertens Res 29: 457-466. doi:10.1291/hypres.29.457. PubMed: 16940709.
- Trion A, de Maat M, Jukema W, Maas A, Offerman E et al. (2006) Antiatherosclerotic effect of amlodipine, alone and in combination with atorvastatin, in APOE*3-Leiden/hCRP transgenic mice. J Cardiovasc Pharmacol 47: 89-95. doi:10.1097/01.fjc.0000195603.65858.27. PubMed: 16424791.
- Candido R, Allen TJ, Lassila M, Cao Z, Thallas V et al. (2004) Irbesartan but not amlodipine suppresses diabetes-associated atherosclerosis. Circulation 109: 1536-1542. doi:10.1161/01.CIR. 0000124061.78478.94. PubMed: 15023892.
- 44. Delsing DJM, Jukema JW, van de Wiel MA, Emeis JJ, van der Laarse A et al. (2003) Differential effects of amlodipine and atorvastatin treatment and their combination on atherosclerosis in ApoE*3-Leiden transgenic mice. J Cardiovasc Pharmacol 42: 63-70. doi: 10.1097/00005344-200307000-00010. PubMed: 12827028.
- 45. Doran DE, Weiss D, Zhang Y, Griendling KK, Taylor WR (2007) Differential effects of AT(1) receptor and Ca(2+) channel blockade on atherosclerosis, inflammatory gene expression, and production of reactive oxygen species. Atherosclerosis 195: 39-47. doi:10.1016/ j.atherosclerosis.2007.02.018. PubMed: 17224157.
- Nussberger J, Aubert JF, Bouzourene K, Pellegrin M, Hayoz D et al. (2008) Renin inhibition by aliskiren prevents atherosclerosis progression: comparison with irbesartan, atenolol, and amlodipine. Hypertension 51: 1306-1311. doi:10.1161/HYPERTENSIONAHA. 108.110932. PubMed: 18391092.
- Das R, Burke T, Van Wagoner DR, Plow EF (2009) L-type calcium channel blockers exert an antiinflammatory effect by suppressing expression of plasminogen receptors on macrophages. Circ Res 105: 167-175. doi:10.1161/CIRCRESAHA.109.200311. PubMed: 19520970.

- Jeppesen P, Bruun J, Nielsen-Kudsk F (1998) Amlodipine dynamic effects and myocardial pharmacokinetics in the isolated and perfused guinea-pig heart. Pharmacol Toxicol 82: 250-256. doi:10.1111/j. 1600-0773.1998.tb01433.x. PubMed: 9646331.
- Noguchi K , Takahashi K, Higuchi S (1998) In-vitro negative chronotropic and inotropic effects of a novel dihydropyridine derivative, CD-832, in the guinea-pig: comparison with calcium-channel antagonists. J Pharm Pharmacol 50: 329-334. doi:10.1111/j. 2042-7158.1998.tb06869.x. PubMed: 9600727.
- Faulkner JK, McGibney D, Chasseaud LF, Perry JL, Taylor IW (1986) The pharmacokinetics of amlodipine in healthy volunteers after single intravenous and after 14 repeated oral doses given once daily. Br J Clin Pharmacol 22: 21-25. doi:10.1111/j.1365-2125.1986.tb02874.x. PubMed: 2943308.
- Bhatt J, Singh S, Subbaiah G, Shah B, Kambli S et al. (2007) A rapid and sensitive liquid chromatography-tandem mass spectrometry (LC-MS/MS) method for the estimation of amlodipine in human plasma 21. Chromatogr: Biomed. pp. 169-175.
- 52. Cassis LA, Gupte M, Thayer S, Zhang X, Charnigo R et al. (2009) Angiotensin II infusion promotes abdominal aortic aneurysms independent of increased blood pressure in hypercholesterolemic mice. Am J Physiol Heart Circ Physiol 296: H1660-H1665. doi:10.1152/ ajpheart.00028.2009. PubMed: 19252100.
- Lu H, Cassis LA, Daugherty A (2007) Atherosclerosis and arterial blood pressure in mice. Curr Drug Targets 8: 1181-1189. doi: 10.2174/138945007782403829. PubMed: 18045096.
- Kanematsu Y, Kanematsu M, Kurihara C, Tsou TL, Nuki Y et al. (2010) Pharmacologically induced thoracic and abdominal aortic aneurysms in mice. Hypertension 55: 1267-1274. doi:10.1161/HYPERTENSIONAHA. 109.140558. PubMed: 20212272.
- Schricker K, Hamann M, Macher A, Krämer BK, Kaissling B et al. (1996) Effect of amlodipine on renin secretion and renin gene expression in rats. Br J Pharmacol 119: 744-750. doi:10.1111/j. 1476-5381.1996.tb15735.x. PubMed: 8904650.
- 56. Lu H, Balakrishnan A, Howatt DA, Wu C, Charnigo R et al. (2012) Comparative effects of different modes of renin angiotensin system inhibition on hypercholesterolaemia-induced atherosclerosis. Br J Pharmacol 165: 2000-2008. doi:10.1111/j.1476-5381.2011.01712.x. PubMed: 22014125.
- Lu H, Rateri DL, Feldman DL, Charnigo RJ Jr, Fukamizu A et al. (2008) Renin inhibition reduces hypercholesterolemia-induced atherosclerosis in mice. J Clin Invest 118: 984-993. PubMed: 18274671.
- Daugherty A, Cassis LA, Lu H (2011) Complex pathologies of angiotensin II-induced abdominal aortic aneurysms. J Zhejiang Univ Sci B 12: 624-628. doi:10.1631/jzus.B1101002. PubMed: 21796801.
- Bruemmer D, Daugherty A, Lu H, Rateri DL (2011) Relevance of angiotensin II-induced aortic pathologies in mice to human aortic aneurysms. Ann N Y Acad Sci 1245: 7-10. doi:10.1111/j. 1749-6632.2011.06332.x. PubMed: 22211965.
- Lu H, Rateri DL, Bruemmer D, Cassis LA, Daugherty A (2012) Involvement of the renin-angiotensin system in abdominal and thoracic aortic aneurysms. Clin Sci 123: 531-543. doi:10.1042/CS20120097. PubMed: 22788237.
- Saraff K, Babamusta F, Cassis LA, Daugherty A (2003) Aortic dissection precedes formation of aneurysms and atherosclerosis in

angiotensin II-infused, apolipoprotein E-deficient mice. Arterioscler Thromb Vasc Biol 23: 1621-1626. doi:10.1161/01.ATV. 0000085631.76095.64. PubMed: 12855482.

- Barisione C, Charnigo R, Howatt DA, Moorleghen JJ, Rateri DL et al. (2006) Rapid dilation of the abdominal aorta during infusion of angiotensin II detected by noninvasive high-frequency ultrasonography. J Vasc Surg 44: 372-376. doi:10.1016/j.jvs.2006.04.047. PubMed: 16890871.
- Rossi-Foulkes R, Roman MJ, Rosen SE, Kramer-Fox R, Ehlers KH et al. (1999) Phenotypic features and impact of beta blocker or calcium antagonist therapy on aortic lumen size in the Marfan syndrome. Am J Cardiol 83: 1364-1368. doi:10.1016/S0002-9149(99)00101-0. PubMed: 10235096.
- Yokokura H, Hiromatsu S, Akashi H, Kato S, Aoyagi S (2007) Effects of calcium channel blocker azelnidipine on experimental abdominal aortic aneurysms. Surg Today 37: 468-473. doi:10.1007/ s00595-006-3367-6. PubMed: 17522763.
- Tomita N, Yamasaki K, Izawa K, Kunugiza Y, Osako MK et al. (2008) Inhibition of experimental abdominal aortic aneurysm progression by nifedipine. Int J Mol Med 21: 239-244. PubMed: 18204791.
- Mancini GB (2002) Antiatherosclerotic effects of calcium channel blockers. Prog Cardiovasc Dis 45: 1-20. doi:10.1016/ S0033-0620(02)70007-8. PubMed: 12138411.
- Weiss D, Taylor WR (2008) Deoxycorticosterone acetate salt hypertension in apolipoprotein E-/- mice results in accelerated atherosclerosis: the role of angiotensin II. Hypertension 51: 218-224. doi:10.1161/HYPERTENSIONAHA.107.095885. PubMed: 18158357.
- van de Poll SW, Delsing DJ, Jukema JW, Princen HM, Havekes LM et al. (2002) Raman spectroscopic investigation of atorvastatin, amlodipine, and both on atherosclerotic plaque development in APOE*3 Leiden transgenic mice. Atherosclerosis 164: 65-71. doi: 10.1016/S0021-9150(02)00055-2. PubMed: 12119194.
- van de Poll SW, Delsing DJ, Wouter Jukema J, Princen HM, Havekes LM et al. (2003) Effects of amlodipine, atorvastatin and combination of both on advanced atherosclerotic plaque in APOE*3-Leiden transgenic mice. J Mol Cell Cardiol 35: 109-118. doi:10.1016/ S0022-2828(02)00284-5. PubMed: 12623305.
- Nakandakare ER, Charf AM, Santos FC, Nunes VS, Ortega K et al. (2008) Dietary salt restriction increases plasma lipoprotein and inflammatory marker concentrations in hypertensive patients. Atherosclerosis 200: 410-416. doi:10.1016/j.atherosclerosis. 2007.12.034. PubMed: 18262533.
- Yue H, Uzui H, Shimizu H, Nakano A, Mitsuke Y et al. (2004) Different effects of calcium channel blockers on matrix metalloproteinase-2 expression in cultured rat cardiac fibroblasts. J Cardiovasc Pharmacol 44: 223-230. doi:10.1097/00005344-200408000-00012. PubMed: 15243304.
- 72. Kataoka C, Egashira K, Ishibashi M, Inoue S, Ni W et al. (2004) Novel anti-inflammatory actions of amlodipine in a rat model of arteriosclerosis induced by long-term inhibition of nitric oxide synthesis. Am J Physiol Heart Circ Physiol 286: H768-H774. doi:10.1152/ajpheart. 00937.2002. PubMed: 14592942.