

Involvement of chymase-mediated angiotensin II generation in blood pressure regulation

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Angiotensin I-converting enzyme (ACE) inhibitors are thought to lower blood pressure in hypertensive patients, mainly by decreasing angiotensin II (Ang II) formation. Chymase, a human mast cell protease, has recently been proposed to play a role in blood pressure regulation because of its Ang II-forming activity. Here we show that the predominant chymase mRNA species in the mouse aorta are those for types 4 and 5 isoforms, and that both are efficient Ang II-forming enzymes. Evaluation of ACE-dependent and ACE-independent Ang II-forming pathways in mast cell-deficient (Kit^w/Kit^{w-v}) mice and their mast cell-sufficient littermate (MC^{+/+}) controls revealed that, in contrast to the latter, Kit^w/Kit^{w-v} mice fail to express chymase mRNAs in the vasculature and have almost no ACE-independent Ang II-forming activity in either isolated blood vessels or homogenates. Moreover, in MC^{+/+} but not in Kit^w/Kit^{w-v} mice, a contribution of ACE-independent Ang II generation to blood pressure regulation was evident by a 1.6-fold greater maximal reduction in mean arterial pressure with acute ACE inhibition plus AT₁ receptor blockade than with ACE inhibition alone. Thus, mast cells are the source of the vascular ACE-independent pathway, and the antihypertensive benefit of combining ACE inhibitor therapy with AT₁ receptor antagonist therapy is most likely due to negation of chymase-catalyzed Ang II generation.

Introduction

An angiotensin I-converting enzyme-independent (ACE-independent) pathway for the conversion of angiotensin I (Ang I) to the vasoconstrictor hormone angiotensin II (Ang II) was demonstrated in hamster cheek pouch blood vessels more than two decades ago (1). This pathway, which is alternate to the ACE pathway, has been demonstrated in the heart and blood vessels of several species, including humans (2-4). More recently, a mast cell chymase was identified as the major Ang II-forming pathway in the human heart (5). The significance of these studies is twofold. First, the presence of an ACE-independent pathway of Ang II formation in human cardiovascular tissues suggests that the clinical effectiveness of ACE inhibitors in lowering blood pressure may be limited, because these agents do not block chymase-mediated Ang II formation. Second, these studies suggest that AT1 receptor antagonist therapy could allow a more complete interruption of the renin-angiotensin system by blocking both ACE-dependent and -independent Ang II formation. Nevertheless, given that ACE inhibitors also affect bradykinin degradation, as well as other hormonal pathways, it remains unclear whether the ability of AT1 receptor blockers to inhibit not only ACEdependent but also ACE-independent Ang II generation is mechanistically important in terms of their antihypertensive effects.

Although several isolated organ bath studies have shown that the Ang I vasoconstrictor response is incompletely inhibited by

Nonstandard abbreviations used: angiotensin I (Ang I); Ang I-converting enzyme (ACE); angiotensin II (Ang II); mouse mast cell protease (MMCP); physiological salt solution (PSS); suprarenal aortic constriction (SAC); turnover number (k_{cat}).

Conflict of interest: The authors have declared that no conflict of interest exists. Citation for this article: *J. Clin. Invest.* **114**:112–120 (2004). doi:10.1172/JCI200420805.

ACE inhibitors, in vivo studies with Ang I, where the prohormone is administered intravenously, demonstrate an abrogation of the Ang I response after ACE inhibition (6, 7). Insights into this apparent paradox have come from electron microscopyimmunohistochemistry and microdialysis studies (8-10). These indicate that chymase is localized to the interstitial compartment of the heart and blood vessels, whereas ACE is chiefly in the luminal compartment, and that circulating Ang I does not readily penetrate into the interstitial tissue compartment. In vivo evidence for chymase functionality has also come from studies with the Ang I analog [Pro¹¹,DAla¹²]Ang I – a substrate for Ang II generation by chymase, but not by ACE (11, 12). Moreover, the demonstration that transgenic overexpression of an Ang II-forming rat chymase in mouse VSMCs causes hypertension, suggests that chymase regulates vascular tone (13, 14). This transgenic overexpression of a vascular chymase in mouse VSMCs, which establishes in the mouse the high ACE-independent Ang II-forming activity seen in VSMCs of spontaneously hypertensive rats, also produces vasculopathy. These pharmacological and transgenic studies show chymase functionality in vivo but do not address the question of the relative importance of the ACE-dependent versus ACE-independent pathways of Ang II formation in regulating blood pressure.

Chymase is the major Ang II-forming enzyme in human heart and blood vessels (10), which suggests that chymase deficiency should cause a loss of Ang II generated by ACE-independent pathways. Here, we show that in the vasculature of mice with genetic mast cell deficiency (Kit^w/Kit^{w-v}), expression of various chymase isoforms and ACE-independent Ang II-forming activity are markedly attenuated relative to those in their mast cell-sufficient littermate (MC^{+/+}) controls. These findings indicate that mast cells are the main source of the vascular ACE-independent Ang II-form-





ing pathway. Using these mice as model systems, we explore the in vivo importance of ACE-dependent and -independent pathways of Ang II formation in blood pressure regulation.

Results

Chymase-subtype expression in the aortae of $MC^{*/*}$ and Kit^{w}/Kit^{w-v} mice. Figure 1A shows the expression of chymase isoforms in the aortae of $MC^{*/*}$ mice. Transcripts for the 2, 4, and 5 isoforms were readily detected in the aortae of $MC^{*/*}$ mice. However, chymase 1 mRNA, which was readily detected in the stomach (data not shown), was not readily detected in the aorta (Figure 1A). When normalized for GAPDH expression, chymase mRNA levels in $MC^{*/*}$ mouse aortae were highest for isoforms 4 and 5 and two- tot threefold lower for isoform 2. However, mRNAs for these isoforms were not detected in the aortae of Kit^w/Kit^{w-v} mice (Figure 1A).

Ang II-forming activity of recombinant mouse chymase isoenzymes. Because of species and subtype differences in chymase-isoenzyme kinetic properties, we investigated the Ang II-forming activity of mouse chymases detected in the aorta. Recombinant forms of mouse chymases 1, 2, 4, and 5 tagged at the C-terminus with a c-myc epitope (referred to hereafter as *r-c-myc*-tagged chymases 1, 2, 4, and 5) were generated and partially purified. *r-c-myc*tagged chymases 1, 2, 4, and 5 were detected by Western blotting using a *c-myc* antibody (data not shown). *r-c-myc*-tagged chymases 1, 4, and 5 hydrolyzed the Phe⁸-

Figure 1

GAPDH-normalized chymase 1, chymase 2, chymase 4, and chymase 5 mRNA levels in aortae from vehicle- or captopril-treated MC^{+/+} and Kit^w/Kit^{w-v} mice. Three-week-old MC^{+/+} and Kit^w/Kit^{w-v} mice were given vehicle (controls) (**A**) or captopril (50 mg/kg/d) (**B**) in drinking water for 4 weeks. Thus, all measurements are from aortae of 7-week-old mice. Values are mean \pm SEM, and n = 7 in each group. Comparisons between MC^{+/+} and Kit^w/Kit^{w-v} mouse groups were made using the unpaired Student's *t* test; **P* < 0.01, ***P* < 0.001. In all cases, differences in transcript ratios between vehicle- and captopril-treated groups were not significant.

His⁹ and Tyr⁴-Ile⁵ bonds in Ang I (DRVY⊻IHPF±HL; cleavage sites indicated by arrows), thus simultaneously forming and then degrading the generated Ang II (DRVYIHPF) (data not shown).

Because Ang I is degraded at two sites, it became difficult to define the net Ang II-forming activity of these recombinant mouse chymases using Ang I alone as a substrate. We addressed this issue by studying the kinetics of Ang II formation and of degradation separately, using the Ang I fragments Ang I-(5-10) (IHPF±HL) and Ang II (DRVY \pm IHPF), respectively, as substrates. *r-c-myc*-tagged mouse chymases cleave these substrates at a single site (indicated by the arrow). Conversion of Ang I-(5-10) to Ang II-(5-8) is expected to reflect the potential of the chymases to convert Ang I to Ang II, because the P_4 to P_1 and P_1' to P_2' residues of the substrate are known to be crucial in determining catalytic specificity (15, 16). In Ang I-(5-10) and Ang I, the substrate P4 to P1 residues (i.e., IHPF) and P_1' to P_2' residues (i.e., HL) are identical. Table 1 summarizes the kinetic constants for Ang I-(5-10) and Ang II hydrolysis. The specificity constant (turnover number $[k_{cat}]/K_m$) ratios for Ang II formation/Ang II degradation for r-c-myc-tagged mouse chymases 1, 4, and 5 are 17:1, 5:1, and 13:1, respectively; thus, these mouse chymase subtypes are net Ang II-forming enzymes. The specificity constants for the hydrolysis of the Phe⁸-His⁹ bond in Ang I-(5-10) by r-c-myc-tagged chymases 4 and 5 were more than 460-fold higher than that for *r*-*c*-*myc*-tagged chymase 1.

Vascular ACE-dependent and -independent pathways of Ang II formation in vitro in MC^{+/+} and Kit^w/Kit^{w-v} mice. To study the effect of mast cell deficiency on vascular ACE-dependent and -independent Ang II-forming pathways, we examined the effect of ACE inhibition on the Ang II-forming activity of aortic homogenates from MC^{+/+} and

Table 1

Kinetic constants for the formation of Ang II-(5–8) from Ang I-(5–10) or Ang II by purified recombinant *c-myc*–tagged mouse chymases (*r-c-myc*chymases) 1, 2, 4, and 5

Enzyme	Substrate	<i>K</i> տ (μM)	k _{cat} (s ⁻¹)	<i>k</i> _{cat} / <i>K</i> _m (s ⁻¹ μM ⁻¹)
<i>r-c-myc</i> -chymase 1	Ang I-(5–10)	12 ± 1.1	0.075 ± 0.006	0.0063
<i>r-c-myc</i> -chymase 2	Ang I-(5–10)		ND	
<i>r-c-myc</i> -chymase 4	Ang I-(5–10)	76 ± 6.4	220 ± 18	2.9
<i>r-c-myc</i> -chymase 5	Ang I-(5–10)	35 ± 6.8	220 ± 31	6.3
<i>r-c-myc</i> -chymase 1	Ang II	5,500 ± 1,800	2.0 ± 0.67	0.00036
<i>r-c-myc</i> -chymase 2	Ang II		ND	
<i>r-c-myc</i> -chymase 4	Ang II	67 ± 3.2	37 ± 0.7	0.55
<i>r-c-myc</i> -chymase 5	Ang II	28 ± 1.9	13 ± 0.2	0.46

Ang II-(5–8), IHPF; Ang I-(5–10), IHPFHL; Ang II, DRVYIHPF; ND, activity not detectable. Values are mean ± SEM of three independent determinations.

Table 2

Effect of chronic captopril treatment on aortic ACE-dependent and ACE-independent Ang II–forming activities in MC^{+/+} and Kitw/Kitw-v mice

Ang II–forming activity (nmol Ang II formed/min/mg protein) ^A								
	ACE-deper	ndent	ACE-inde	ACE-independent				
Genotype	Veh	Cap	Veh	Cap				
MC+/+	1.3 ± 0.56	ND	0.87 ± 0.023	2.1 ± 0.49^{B}				
Kit ^w /Kit ^{w-v}	0.98 ± 0.3	ND	0.11 ± 0.09 ^c	0 ± 0^{D}				

Values are mean \pm SEM. In each group, n = 6. All measurements were made in 7-week-old mice. Because the ACE inhibitor–tissues ACE complexes dissociate in in vitro assays, the measurement of ACE activity in ACE inhibitor–treated animals is considered not to reflect true residual tissue ACE activity in vivo. Hence, we do not report these values here. ^Ang II–forming activity: Total, that determined in the absence of any inhibitors; ACE-dependent, that level of the total that is inhibited by the ACE inhibitor lisinopril (10 μ M); and ACE-independent, the total minus the ACE-dependent activity. Only ACE-dependent and -independent activities are shown. Intra-genotype comparisons were made between vehicle and captopril groups; ^BP < 0.05. Inter-genotype comparisons: ^CP < 0.05, ^DP < 0.001. Cap, captopril (~50 mg/kg/d) given orally in the drinking water to 3-week-old mice for 4 weeks. Veh, control drinking water without captopril. ND, not determined.

Kit^w/Kit^{w-v} mice. The ACE-dependent Ang II–forming activity was 1.5-fold higher than the ACE-independent activity in MC^{+/+} aortic homogenates (Table 2). In contrast, in the Kit^w/Kit^{w-v} mouse, the aortic ACE-dependent Ang II–forming activity was about 8.9-fold higher than the ACE-independent activity (Table 2). Total Ang II–forming activity (ACE-dependent plus ACE-independent) was 1.9-fold higher in MC^{+/+} mouse aortic homogenates than in Kit^w/Kit^{w-v} mouse aortic homogenates (P < 0.05), and this difference was almost entirely due to an approximately eightfold higher ACE-independent Ang II–forming activity in MC^{+/+} mouse aortae than in Kit^w/Kit^{w-v} mouse aortae (Table 2).

To determine whether the greater dependence of mast cell-deficient mice on ACE for their vascular Ang II generation in vitro influences the enzymatic pathway used for the functional conversion of Ang I to Ang II in intact blood vessels, we examined the vasoconstrictor responses of Ang I in isolated mouse carotid arteries in the presence or absence of the ACE inhibitor lisinopril. As shown in Figure 2, carotid artery contraction produced by 1 nM Ang I was approximately 50% lower (P < 0.001) in those vessels from MC^{+/+} mice pretreated with the ACE inhibitor lisinopril (10 μ M) than in those vessels from MC^{+/+} mice that were naive with respect to ACE inhibition, indicating the presence of functional ACE-dependent and -independent pathways of Ang II formation in the vasculature of mast cell-sufficient mice. In contrast, in Kit^{w/} Kit^{w-v} mice, 1 nM Ang I-dependent constriction of isolated carot-

Figure 2

Reactivity of untreated (black bars) or lisinopril-treated (Lis; 10 μ M; white bars) isolated carotid arteries from 7-week-old MC^{+/+} (left panel) and Kit^{w/}Kit^{w-v} mice (right panel) to 1 nM Ang I. Each carotid artery preparation was subjected to Ang I only once to preclude the effects of tachyphylaxis. The results are expressed as a percentage of the vasoconstrictor effect induced by 1 nM Ang I when given in the absence of ACE inhibitor pretreatment. Values are mean ± SEM, and *n* = 6 in each group. Comparisons between Ang I and Lis + Ang I groups were made using the unpaired Student's *t* test; **P* < 0.001. †*P* < 0.001 for inter-genotype comparisons.



id arteries pretreated with 10 μ M lisinopril was approximately 1% (P < 0.001) of that observed with 1 nM Ang I in other carotid artery preparations that were naive with respect to ACE inhibition (P < 0.001), indicating the near absence of a functional vascular ACE-independent pathway in mast cell-deficient mice.

We further examined whether mast cell deficiency influences the vascular ACE-independent Ang II-forming pathway in isolated mesenteric arteries (resistance vessels). The difficulty of achieving complete ACE inhibition across the different vascular compartments (luminal and abluminal) in this closed system necessitated a strategy different from that used in the studies of isolated carotid artery rings (described above), in which the ACE inhibitor easily accessed both vascular compartments. We used [Pro11,DAla12]Ang I (Asp-Arg-Val-Tyr-Ile-His-Pro-Phe-His-Leu-Pro-DAla; the Ang II sequence is italicized), a chymase-selective Ang I analog that we have previously described (11). [Pro¹¹,DAla¹²]Ang I takes advantage of substrate-specificity differences between ACE and chymase. In [Pro¹¹,DAla¹²]Ang I, the penultimate proline makes it resistant to ACE, and the C-terminal DAla makes it resistant to carboxypeptidases. Chymase, an endopeptidase, cleaves [Pro¹¹,DAla¹²]Ang I at the Phe-His bond to generate Ang II. Without this conversion, [Pro¹¹, DAla¹²]Ang I is unable to bind or activate the AT₁ receptor.

As shown in Figure 3, Ang II (1 μ M, a concentration that produces maximal vasoconstriction) produced similar vasoconstrictor responses in mesenteric arteries isolated from either MC^{+/+} or Kit^w/Kit^{w-v} mice. In MC^{+/+} mouse arteries, [Pro¹¹,DAla¹²]Ang I (1 μ M) produced a degree of vasoconstriction that was similar to that produced by Ang II. This was reduced by more than 90% by pretreatment with the AT₁ receptor antagonist losartan (10 μ M), indicating that vasoconstriction by [Pro¹¹,DAla¹²]Ang I was indeed due to its conversion to Ang II and, thus, that an ACE-independent pathway for Ang II generation is active in MC^{+/+} vasculature. In contrast to these responses in MC^{+/+} vessels, and consistent with the virtual abolition of an ACE-independent Ang II-forming pathway in Kit^w/Kit^{w-v} vasculature, the vasoconstrictor effect of [Pro¹¹,DAla¹²]Ang I (1 μ M) in Kit^w/Kit^{w-v} mesenteric arteries was 86% less than that of Ang II (1 μ M).

Comparison of the acute blood pressure–lowering effect of ACE inhibitor alone versus a combination of ACE inhibitor and AT_1 receptor antagonist in vivo in $MC^{+/+}$ and $Kit^{w/}Kit^{w-v}$ mice. To quantify the level of resting blood pressure in anesthetized mice that is due to steady-state conversion of endogenous Ang I to Ang II by the ACE-independent pathway, we determined the difference between the maximal blood pressure–lowering effect of the ACE inhibitor captopril and that produced by a combination of captopril and the AT_1 receptor antagonist losartan. The rationale for these experiments was as follows. The vasodilator effect of ACE inhibition is the result of both suppression of the effects of ACE-mediated Ang II formation, and alterations in other non-angiotensinergic mechanisms, such as





Figure 3

Vasoconstrictor effects of Ang II (1 μ M) or [Pro¹¹, DAla¹²]Ang I (1 μ M) in mesenteric artery preparations from 7-week-old MC^{+/+} (left panel) and Kit^w/Kit^{w-v} mice (right panel). Results are expressed as the peak change in vessel diameter (μ m), where a decrease in vessel diameter represents vasoconstriction. Each mesenteric artery preparation was subjected to Ang II or [Pro¹¹, DAla¹²]Ang I only once to preclude the effects of tachyphylaxis. Values are mean ± SEM, and *n* = 5–6 in each group. Comparison between the [Pro¹¹, DAla¹²]Ang I responses in mesenteric artery preparations from MC^{+/+} mice in the presence or absence of 10 μ M losartan was made using the unpaired Student's *t* test; **P* < 0.001. †*P* < 0.001 for inter-genotype comparisons.

inhibition of bradykinin degradation. Combined ACE inhibition and AT₁ receptor blockade will additionally inhibit the actions of Ang II formed by the ACE-independent pathway. Thus, the difference in the vasodilator response between combined blockade and ACE inhibition alone is that which is attributable to Ang II generated by the ACE-independent pathway.

As shown in Table 3, base-line aortic blood pressure, heart rate, and cardiac contractility (dP/dt_{max}) were similar in anesthetized MC^{+/+} and Kit^w/Kit^{w-v} mice (Table 3). In the MC^{+/+} mice, acute administration of captopril plus losartan (2.5 mg/kg and 80 mg/kg, respectively, doses shown in initial studies to be twofold greater than those required by each agent, when given alone, to produce maximal falls in blood pressure; data not shown) resulted in a significantly greater reduction (1.6-fold) in mean aortic blood pressure than administration of captopril alone (2.5 mg/kg) (P < 0.05) (Figure 4). In contrast, in Kit^w/Kit^{w-v} mice, the difference in the blood pressure-lowering effect of combined therapy versus captopril monotherapy was not statistically significant (Figure 4).

In both MC+/+ and Kitw/Kitw-v mice, acute administration

of captopril (2.5 mg/kg) plus losartan (80 mg/kg) produced a greater (P < 0.05) fall in blood pressure than losartan alone (80 mg/kg), and the additional decrements in pressure observed with the combined agents were genotype-independent (1.9fold in MC^{+/+} vs. 2.1-fold in Kit^w/Kit^{w-v} mice, P not significant) (Figure 4). These findings indicate that, at least acutely, Ang II-independent vasodilator activity of captopril is similar in both MC^{+/+} and Kit^w/Kit^{w-v} mice. Effect of chronic ACE inhibition on blood pressure and aortic ACEindependent Ang II–forming activities in $MC^{+/+}$ and Kit^{w}/Kit^{w-v} mice. As shown in Figure 5, chronic ACE inhibition with captopril (50 mg/kg/d for 4 weeks) lowered mean proximal aortic blood pressures in both anesthetized Kit^w/Kit^{w-v} mice (53% decrease, P < 0.001) and $MC^{+/+}$ mice (38% decrease, P < 0.001), although the decrements in pressure were greater in the former (1.4-fold, P < 0.05). The less marked reductions in blood pressure in the $MC^{+/+}$ animals were associated with a twofold increase in ACE-independent aortic Ang II–forming activity (P < 0.05) (Table 2), a response that was not due to enhanced chymase gene expression, since steady-state mRNA levels for the 1, 2, 4, and 5 chymase isoforms remained unchanged with captopril treatment (Figure 1).

Effect of chronic ACE inhibition on blood pressure in MC^{+/+} and Kit^w/ Kit^{w-v} mice with suprarenal aortic constriction. To evaluate the antihypertensive effects of chronic ACE inhibition in the setting of renin-angiotensin system activation (17), MC^{+/+} and Kit^w/Kit^{w-v} mice were subjected to suprarenal aortic constriction (SAC). This resulted in similar increases in mean proximal aortic blood pressure (MC^{+/+}, +33%, P < 0.001; Kit^w/Kit^{w-v}, +27%, P < 0.01, respectively), as compared with those in the respective sham-operated controls (Figure 6). In a parallel short-term study, whereas captopril (50 mg/kg/d, orally, for 3 days) administered from the time of SAC attenuated the rise in blood pressure in MC^{+/+} mice by only about 50% (P < 0.05), in the Kit^w/Kit^{w-v} animals it completely prevented it (Figure 6).

Discussion

Here we show that the vascular ACE-independent pathway of Ang II formation is deficient in mice with genetic mast cell deficiency, but not in their mast cell-sufficient littermate controls. Furthermore, in the former, but not the latter, the blood pressure-lowering effect of ACE inhibitor monotherapy is equivalent to that of combined ACE inhibitor plus AT₁ antagonist therapy. These findings suggest an important role for the ACE-independent pathway in the physiological regulation of blood pressure.

In support of this contention, vascular ACE-independent Ang II-forming activity in aortic homogenates was about eightfold lower in Kit^w/Kit^{w-v} mice compared with their MC^{+/+} littermate controls. Despite electron microscopic-immunohistochemical evidence for chymase localization in the interstitial compartment of human cardiac left ventricles (8), it is possible that assays for Ang II-forming activity in tissue homogenates, while indicating overall enzymatic potential in a tissue, do not provide information on the extracellular component of the pathway. For example, in intact tissues, if mast cell Ang II-forming proteases are mainly in intracellular granules, they may not be available for the physiological conversion of Ang I to Ang II in the extracellular milieu. To specifically address this issue, we examined total and

Hemodynamic measurements in anesthetized 7-week-old MC^{+/+} and Kit^w/Kit^{w-v} mice

Genotype	п	MABP (mmHg)	SABP (mmHg)	DABP (mmHg)	HR (beats/min)	dP/dt _{max} (mmHg/s)	dP/dt _{min} (mmHg/s)
MC ^{+/+}	13	89 ± 4	116 ± 3	75 ± 4	353 ± 20	8,084 ± 555	-9,444 ± 420
Kit ^w /Kit ^{w-v}	13	87 ± 4	116 ± 4	72 ± 5	332 ± 21	7,446 ± 526	-9,359 ± 573

Values are mean \pm SEM. MABP, SABP, and DABP; mean, systolic, and diastolic proximal aortic blood pressure, respectively. HR, heart rate.

Table 3



Figure 4

Acute mean proximal aortic blood pressure–lowering (BP-lowering) effects of infusion of 2.5 mg/kg captopril (Cap), 80 mg/kg losartan (Los), or 2.5 mg/kg captopril combined with 80 mg/kg losartan (Los + cap) in anesthetized 7-week-old MC^{+/+} (left panel) and Kit^{w/}Kit^{w-v} mice (right panel). The data are maximal changes in mean blood pressure. Values are mean \pm SEM, and n = 6-10 in each group. Intra-genotype comparisons shown in the figure were made using an unpaired Student's *t* test. Inter-genotype comparisons, using an unpaired Student's *t* test, were found not to be significantly different.

ACE-independent vasoconstrictor actions of Ang I in isolated carotid arteries. An ACE-independent vasoconstrictor effect of Ang I was not observed in Kit^w/Kit^{w-v} mice but was about 50% of the total response in MC^{+/+} mice. In addition, we showed that [Pro¹¹,DAla¹²]Ang I, an analog of Ang I that is converted to Ang II by chymase but not ACE (11), produces a vasoconstrictor response in isolated mesenteric artery preparations from MC^{+/+} mice. Thus, the homogenate enzyme activity assays and the functional isolated organ bath studies both indicate that mast cell deficiency causes almost complete loss of vascular ACE-independent Ang II formation and, further, that resident mast cells are a constitutive source of interstitial chymase.

In a recent study, Rabinovitch and colleagues (14) showed that VSMC chymase mRNA levels are about tenfold higher in aortae of spontaneously hypertensive rats than in control Sprague-Dawley rats. Moreover, through conditional transgenic studies, they provided evidence that this vascular expression contributes to the hypertensive phenotype (13). Does a homolog of the rat VSMC chymase participate in the vascular ACE-independent pathway in MC^{+/+} mice? The most compelling evidence that it does not is the finding that in Kit^w/Kit^{w-v} mice, aortic ACE-independent Ang II-forming activity is, at best, only 12% of that in MC^{+/+} mice. It is unlikely that this minor aortic ACE-independent Ang II-forming activity is due to a homolog of rat VSMC chymase, since (a) PCR primers, specific for rat vascular smooth muscle chymase, do not amplify a homolog using MC^{+/+} mouse genomic DNA as a template (data not shown), and (b) the two mouse genes most closely related to rat vascular smooth muscle chymase - mouse chymases 1 and 4 (18) - are not expressed in aortae of Kit^w/Kit^{w-v} mice. Although it is possible that the minor non-ACE Ang II-forming activity that occurs in Kitw/Kitw-v mouse aorta homogenates is due to non-ACE Ang II-forming enzymes other than chymases (for example, cathepsin G and tonin), the virtual absence of an ACE-independent Ang I response in blood vessels from these animals suggests that this minor activity is not functionally important.

Kit^w/Kit^{w-v} mice have mutations in *c-Kit* that cause a defect in the tyrosine kinase activity of this receptor (19). This results not only in mast cell deficiency, but also in a mild anemia, an absence of skin melanocytes, and infertility (19). It is unlikely, however, that loss of ACE-independent Ang II formation in the mast cell-deficient mice is due to these extra-mast cell effects, as they are not related to blood vessel functioning.

Mast cells synthesize and secrete three main groups of enzymes: chymases, tryptases, and carboxypeptidases. Tryptases and carboxypeptidases are not Ang II-forming (18, 20). In the mouse, chymase-subtype expression in mast cells depends on their extant hormonal or cytokine environment and is thought to be a product of the state of mast cell differentiation. In the mouse aorta, mRNAs for chymase isoforms 4 and 5 predominate. In Kitw/Kitw-v mice, no chymase mRNA species are detectable, as has been reported previously (21). Chymases 1 and 4 have been demonstrated to be Ang II-forming (18), but chymase 2 lacks activity (22). Based on theoretical considerations regarding the enzyme S₁ subsite, chymase 5 has been reported to be an elastase that lacks the crucial chymotrypsin-like activity necessary for the conversion of Ang I to Ang II (23). As demonstrated here, recombinant mouse chymase 2 lacks Ang II-forming activity. The 1, 4, and 5 chymase isoforms, on the other hand, do have Ang II-forming activity in in vitro assays but, unlike human chymase, also possess Ang II-degrading activity. Nevertheless, since with all three isoforms the former activity predominates, in vivo they are expected to be net Ang II-forming enzymes. Given that the specificity constant (k_{cat}/K_m) for the Ang II-forming activity of mouse chymase 1 is more than 400-fold lower than that for chymase isoforms 4 and 5, and that chymase 2 lacks activity, the 4 and 5 isoforms are likely to be the key enzymatic determinants of ACE-independent Ang II generation in mouse blood vessels. However, definitive proof of their relative roles in converting Ang I to Ang II in vivo must await the development of isoform-specific chymase inhibitors.

At the outset, the finding that blood pressure levels in Kit^w/Kit^{w-v} mice are similar to those in $MC^{*/*}$ mice appeared not to support a



Figure 5

Proximal aortic blood pressure (BP) in anesthetized MC^{+/+} and Kit^{w/}Kit^{w-v} mice after 4 weeks of vehicle control (veh; **A**) or captopril administration (cap; 50 mg/kg/d; **B**) in drinking water. Captopril or vehicle administration was initiated in MC^{+/+} and Kit^{w/}Kit^{w-v} mice at 3 weeks of age. Thus, all measurements are from 7-week-old mice. Values are mean \pm SEM, and n = 6 in each group.



Figure 6

Effect of chronic captopril treatment (cap; 50 mg/kg/d orally; started at the time of the operation) on proximal aortic blood pressures in anesthetized 7-week-old MC^{+/+} (**A**) and Kit^{w/}Kit^{w-v} mice (**B**) 3 days after SAC (black bars) compared with sham controls (white bars). Values are mean \pm SEM, and n = 6 in each group.

role for the ACE-independent pathway in blood pressure regulation. However, a conclusion based solely on this finding is limited, since compensatory mechanisms, both angiotensinergic and nonangiotensinergic, could maintain blood pressure in the absence of ACE-independent Ang II generation.

To dissect out the influence of ACE-dependent from the influence of ACE-independent Ang II formation on blood pressure, we compared the acute effects of the ACE inhibitor captopril given alone, with those of this agent given in combination with the AT₁ receptor blocker losartan. In both cases, we used doses that were maximally hypotensive when used alone in either Kit^w/Kit^{w-v} or MC^{+/+} mice. Ang II-independent blood pressure effects of ACE inhibition (e.g., on bradykinin degradation) are expected to be identical in the single-agent and combination treatments. Thus, the difference in the vasodilator effect between these two treatments in mice of a given genotype is likely to be due to ACE-independent Ang II formation.

In Kit^w/Kit^{w-v} mice, we found that the reduction in aortic blood pressure by combined ACE inhibition and AT₁ blockade, though slightly greater than that caused by ACE inhibition alone, was not significantly greater. This result is expected, since biochemical and isolated organ bath studies show a near absence of a vascular ACEindependent pathway in these mast cell-deficient mice. In MC+/+ mice, the mean aortic blood pressure-lowering effect of combination therapy was 1.6-fold greater than that of ACE inhibition alone (P < 0.05). This result, combined with the demonstration of a vascular ACE-independent pathway of Ang II formation in MC+/+ mice, indicates that an Ang II-dependent but ACE-independent mechanism contributes to blood pressure regulation in MC^{+/+} mice. It is also of note that in human cardiovascular tissues, chymase levels are much higher than in the mouse (4, 24); this suggests an even more important role for the ACE-independent pathway in blood pressure regulation in humans than that demonstrated here in the mouse.

In these studies we also demonstrate that a chronic loss of vascular chymase is not associated with a compensatory increase in vascular ACE activity. Hence, the total Ang II-forming activity of the vessel is not normalized in mice with chronic mast cell deficiency. It is, therefore, not surprising that the acute hypotensive effects of ACE inhibition are equivalent in normotensive Kit^w/ Kit^{w-v} and MC^{+/+} mice. In contrast to these findings, we found that chronic ACE inhibition lowered blood pressure to a greater extent in normotensive Kit^w/Kit^{w-v} mice than in MC^{+/+} mice. Moreover, whereas chronic ACE inhibition normalized blood pressure in Kit^w/Kit^{w-v} mice with renovascular hypertension, it was only about 50% as effective in MC^{+/+} mice with an equivalent degree of hypertension. The ACE-independent pathway of Ang II formation, therefore, appears to limit the vasodilator potential of chronic ACE inhibitor therapy.

Why should the vasodilator effect of chronic ACE inhibition be different from that of acute ACE inhibition in MC^{+/+} and Kit^w/ Kit^{w-v} mice? In MC^{+/+} mice we found that chronic ACE inhibition causes an approximately 2.4-fold increase in vascular ACE-independent Ang II-forming activity. The mechanism of this increase appears not to be due to an increase in chymase mRNA expression levels but could be due to an increase in chymase release, an increase in tissue retention of the released chymase, and/or a decrease in degradation. In this regard, tissue retention of released chymase through ionic interactions with ECM proteins has been proposed to play an important role in determining chymase levels in the human heart (8). Chronic ACE inhibition has also been shown to increase circulating Ang I levels (25). Increases in chymase activity or Ang I levels are expected to increase vascular Ang II levels, because physiological levels of Ang I are well below the K_m of chymase (5, 25). This line of reasoning suggests that limitations to the therapeutic efficacy of ACE inhibitors resulting from ACEindependent Ang II generation are not static. Rather, responses to chronic ACE inhibition are likely to be influenced by progressive compensatory increases in vascular chymase-like activity, as well as by increased production of Ang I. Thus, the contribution of the ACE-independent pathway to vascular tone could become exaggerated with chronic ACE inhibitor therapy.

Reduction in Ang II formation by ACE inhibitors, and inhibition of the cardiovascular effects of Ang II by AT₁ receptor antagonists, are mechanisms exploited clinically to lower blood pressure in hypertensive patients. The demonstration that combined ACE inhibitor plus AT₁ receptor antagonist therapy lowers blood pressure more effectively than does ACE inhibitor monotherapy suggests that ACE inhibitor–dependent suppression of the reninangiotensin system is incomplete (26, 27). Based on the findings of this study, we propose that this incomplete suppression is due to the chymase pathway of Ang II formation.

Methods

Animals. Both Kitw-v (background, C57BL/6) and Kitw (background, WB/ReJ) and their wild-type littermates were purchased from The Jackson Laboratory (Bar Harbor, Maine, USA), and two lines were maintained as the sources of the Kit^{w-v} and Kit^w heterozygotes. Mating Kit^{w-v} heterozygotes with Kit^w heterozygotes resulted in WBB6F1/J-Kit^w/Kit^{w-v} double heterozygotes (identified as having black eyes and white coat), wild-type littermates (black coat), Kit^{w-v/+} heterozygotes (gray with light belly and white spot, light tail), and Kit^w/⁺ heterozygotes (black with white belly; occasional occurrence of a white spot on the head with a white streak radiating from the spot; tail with white tip). Only mast cell-deficient WBB6F1/J-Kit^w/Kit^{w-v} mice (black eyes and white coat) and their mast cell-sufficient wild-type littermate controls (black coat) were used in this study. Their phenotypes have been reviewed previously (19). Multiple (>30), synchronous breedings were initiated, according to protocols provided by The Jackson Laboratory, to generate the required numbers of male double heterozygotes and their wildtype littermates, the latter being used as controls. All procedures

were performed in accordance with institutional guidelines of animal care and treatment.

Determination of chymase mRNA expression in the mouse aorta. For the measurement of mRNA transcripts, aortae of 8-week-old MC+/+ or Kit^w/Kit^{w-v} mice were freshly dissected. Total RNA was extracted from aortae with TRIzol reagent (Invitrogen Corp., Carlsbad, California, USA). mRNA was reverse-transcribed using Transcriptor reverse transcriptase (Roche Diagnostics Corp., Indianapolis, Indiana, USA) according to the manufacturer's protocols. Realtime PCR reactions were carried out using iCycler and iQ SYBR Green Supermix (Bio-Rad Laboratories, Hercules, California, USA). The primer sets for chymases 1, 2, 4, and 5 and GAPDH, and their PCR product sizes (in parentheses), were as follows (28): for mouse chymase 1, 5' primer was 5'-GGAAAACTGGAG-AGAAAGAACCTAC and 3' primer was 5'-GACAGCTGGGGA-CAGAATGGGG (460 bp); for mouse chymase 2, 5' primer was 5'-ATTTCATTGCCTAGTTCCTCTGAC and 3' primer was 5'-CAGGATGAGAACAGGCTGGGAT (525 bp); for mouse chymase 4, 5' primer was 5'-GTAATTCCTCTGCCTCGTCCTTC and 3' primer was 5'-GGACAGGATGGACACATGCTTT (454 bp); for mouse chymase 5, 5' primer was 5'-GGCAGAACAAACGTGAAT-GAGCC and 3' primer was 5'-AAGAACCTTCTGGAAGCTCAGGG (418 bp); and for GAPDH, 5' primer was 5'-ATGGTGAAGGTCG-GTGTG and 3' primer was 5'-ACCAGTGGATGCAGGGAT (633 bp). In the mouse, these chymases were originally referred to as mouse mast cell proteases (MMCPs; i.e., MMCP1, MMCP2, etc.), but not all MMCPs are chymases. Here, we have retained their number designation to describe the subtype, but we refer to them as chymases (29). Real-time PCR conditions for these primers were optimized as 3 mM Mg²⁺, 5 pmol primer, and 50 ng cDNA in a total reaction volume of 25 µl. The amplifying conditions were determined empirically and were 95°C for 30 seconds, 62°C for 30 seconds (or 59°C for 30 seconds for GAPDH), and 72°C for 60 seconds for 45 cycles. All PCR products appeared as a single band of the expected molecular size by 1.5% agarose gel fractionation; these PCR products were cloned and sequenced to identify target cDNAs. In addition to confirming that all PCR products appeared as a single band of the expected molecular size, we also verified that each PCR product consisted of only a single species, as determined by melting-curve analysis. To determine copy number, standard curves for each target transcript were established using purified PCR products. Standard curves were linear, and the correlation coefficients were greater then 0.99.

Construction of the mouse chymase expression vectors. cDNAs were obtained by RT-PCR from MC+/+ mouse stomach total RNA. The primers were designed to amplify the entire coding sequence and to contain restriction sites for insertion of the cDNA into the expression vector. The 5' and 3' primers used for amplifying mouse chymase cDNAs were 5'-ATCTCGAG-AAAAGAATTATTGGTGGTGTTGAGGCTAG and 5'-ATGCG-GCCGCCTACTTGCCATTTATGACTG for mouse chymase 1; 5'-ATCTCGAGAAAAGAATTATTGGTGGTGTTGAGGCTAAAC and 5'-ATGCGGCCGCCTAGGGGCAGACTTGATGCAG for mouse chymase 2; 5'-ATCTCGAGAAAAGAATTATTGGTGGT-GTTGAGTCTAG and 5'-ATGCGGCCGCTTTTCACTTCT-GCCCTTTATAA for mouse chymase 4; and 5'-ATCTC-GAGAAAAGAATCATTGGAGGCACGGAGTGC and 5'-ATGCGGCCGCTTAATTCTCCCCTCAAGATCTTATTG for mouse chymase 5. The cDNAs were synthesized using Transcriptor Reverse Transcriptase (Roche Diagnostics Corp.) according to manufacturer's protocol. For all primer sets, RT-PCR was carried out at 95°C for 30 seconds, 58°C for 60 seconds, and 72°C for 60 seconds for 30 cycles. PCR products were separated on 1.5% agarose gels and then purified using QIAEX II Gel Extraction Kit (QIAGEN Inc.). These purified PCR products were ligated into pGEM-T-Easy Vector (TA vector) (Promega Corp.) and subsequently expressed in *E. coli*. The expressed plasmid DNAs were sequenced to confirm identity.

Pichia pastoris expression vector pPICZα A (Invitrogen Corp.) was used to express mouse chymases. First, all the chymase genes were transferred into pPICZα A vector from TA vector using XhoI and NotI restriction cutting sites. The native N-terminus (IIGG-) of MMCP was introduced flush with the Kex2 cleavage site of the vector in order to allow endogenous activation of chymase during protein expression. To identify recombinant chymase expression in vitro, c-myc-tagged versions of the vectors were created using sitedirected mutagenesis. In these constructions, coding sequences for c-myc epitope tags were introduced immediately C-terminal to the mature chymase sequence. All mutants were verified by sequencing to check for appropriate positioning of the epitope tag.

Expression of recombinant mouse chymases in P. pastoris. Chymase cDNA inserted in pPICZ α A vector DNA (5–10 µg) was linearized by digestion with BstXI. Linearized DNA was transformed into P. pastoris strain GS115 by electroporation (charging voltage 1,500 V, capacitance 25 microfarads, and resistance 400 ohms). The transformants were cultured onto YPDS plates (yeast extract peptone dextrose medium with 1 M sorbitol) containing 100, 500, or 2,000 µg/ml zeocin to select multicopy recombinants in vivo. To verify that the chymase DNA construct was integrated into P. pastoris genomic DNA, the multicopy recombinant strain, the vector-alone transformant, and the parent strain were cultured in minimal dextrose medium at 30°C for 16 hours. Genomic DNA of the cultures was isolated, and total DNA was used as a template for PCR with individual chymase-specific primers (not shown).

To determine appropriate expression conditions for the recombinants, we identified the Mut phenotypes by growing the multicopy recombinants onto MDH (minimal dextrose with histidine) and MMH (minimal methanol with histidine) plates. The Mut phenotype-determined recombinant *P. pastoris* strains were grown in buffered methanol-complex medium at 30°C in a shaking incubator. Methanol was added to a final concentration of 0.5% every 24 hours to maintain induction of expression. Ang I-degrading activity in the culture media was also checked every day for a week to determine the expression level of the recombinant mouse chymases.

Purification of recombinant mouse chymases. Recombinant mouse chymase 1, 4, or 5 in the culture media was purified by ion-exchange chromatography on Bioscale Q2 HPLC columns (Bio-Rad Laboratories) using linear KCl gradients as described previously (30). Recombinant mouse chymase 2 could be detected in the culture media by Western blot analysis, but because it lacked activity it was not studied further. Mouse chymases 1, 4, and 5 each eluted in broad peaks from the ion-exchange column coincident with KCl concentrations between 0.15 and 0.4 M. Peak-activity fractions were pooled. The presence of chymase in the activity-enriched fractions was confirmed using a c-myc antibody (Santa Cruz Biotechnology Inc., Santa Cruz, California, USA) in a West-ern blotting protocol that detects the c-myc tag on the recombinant chymase (data not shown).

Determination of kinetic constants. Twelve concentrations of Ang II or Ang I-(5–10) (Bachem California Inc., Torrance, California,

USA) ranging from 5 to 2,000 µM were incubated with the purified recombinant mouse chymases in 20 mM Tris-HCl buffer, pH 7.5, containing 0.5 M KCl, in a total volume of 50 µl at 37 °C for 30 minutes (30). Chymase concentrations were adjusted so that less than 15% of the substrate (at the lowest substrate concentration) was consumed during the assay to ensure that product formation in the assay remained linear with respect to time. The reactions were terminated by addition of 50 µl ice-cold 0.25% trifluoroacetic acid. The resulting reactions (100 μ l) were applied to a C₁₈ reversephase HPLC column (XTerra RP18 3.5 µm, 4.6 × 50 mm column; Waters Corp., Milford, Massachusetts, USA), which was developed using 8-minute linear gradients with acetonitrile containing 0.1% trifluoroacetic acid. The flow rate was 1 ml/min. The elution position of Ang I, Ang II, Ang I-(5-10), and Ang II-(5-8) was determined using pure synthetic angiotensin peptides as standards. The corresponding product peak areas were used to determine the amount of product formed. Km (Michaelis-Menten constant) and Vmax (maximal velocity) were calculated with nonlinear regression using the equation $v = V_{\text{max}} \times [S]/(K_{\text{m}} + [S])$, where v is the initial rate and [S] is the substrate concentration. Correlation coefficients were mostly greater than 0.99, but never less than 0.97. The k_{cat} (turnover number) was calculated by the formula $k_{cat} = V_{max}/[E_0]$, where $[E_0]$ is the total active-chymase concentration in the assay. The concentration of active-chymase molecules, as opposed to improperly folded and hence inactive-chymase molecules, was determined using 3H-diisopropylfluorophosphate (3H-DFP) labeling (30).

Measurement of ACE and ACE-independent Ang II-forming activities in the mouse aorta. Aortic tissue was homogenized in 2 ml of 50 mM NaH₂PO₄ buffer, pH 7.4, using a Polytron homogenizer (Kinematica AG, Lucerne, Switzerland) at 9,000 rpm for 15 seconds at 4°C and centrifuged at 30,000 g for 20 minutes at 4°C, and the pellet was retained. This procedure was repeated twice. Resulting supernatant fractions, S-1 and S-2, were kept. The pellet was resuspended in 0.5 ml 50 mM NaH₂PO₄ buffer, pH 7.4, containing 100 mM NaCl and 10 mM MgCl₂, and 5 µl of this was added to 35 µl of assay buffer (20 mM Tris-HCl, pH 8.0, containing 0.5 M KCl and 0.01% Triton X-100) containing either no further additive or 10 µM lisinopril and preincubated for 30 minutes at 0°C. Ten microliters of 1 mM Ang I was added to 40 µl of the preincubated aortic extract, and incubations were for 40 minutes at 37°C. Reactions were terminated by the addition of 300 µl ethanol. Precipitated proteins were removed by centrifugation, and the supernatant containing angiotensins was dried. The residue, resuspended in 125 µl of distilled water, was applied to a C18 HPLC column (Grace Vydac, Hesperia, California, USA); angiotensins were separated as described previously (30). Total and ACE-independent Ang II-forming activities were determined in duplicate from assays containing either buffer or lisinopril, respectively. ACE-dependent Ang II-forming activity was taken as total minus ACE-independent activity. Assays were optimized to ensure that Ang II generation was linear with time. With crude membrane preparations, positively charged chymases are quantitatively retained in the pellet, as is ACE, which is membrane bound. Ang II-forming activity in S-1 and S-2 was negligible.

Measurement of Ang I responses in isolated carotid artery preparations. Carotid arteries (560–580 µm internal diameter) were dissected from 7-week-old MC^{+/+} or Kit^w/Kit^{w-v} mice and mounted as ring preparations (2 mm long) in the chamber of an isometric myograph (J.P. Trading, Aarhus, Denmark) as described previously (31). In this apparatus, the entire vessel ring is immersed in the bath, such that any agents added to the bath access both the luminal and abluminal surfaces. The vessel was set up at 37°C in a 15-ml bath containing physiological salt solution (PSS; composition in mM: NaCl 119, KCl 4.7, KH₂PO₄ 1.18, MgSO₄ 1.17, NaHCO₃ 25, CaCl₂ 2.5, EDTA 0.026, glucose 5.5) saturated with 95% O₂/5% CO₂. Then it was stretched to an internal circumference equivalent to 90% of the internal diameter of the vessel if it had been relaxed and under a transmural pressure of 100 mmHg. Vessels were allowed to equilibrate for 1 hour and were then maximally activated with high-potassium PSS (PSS with an equimolar exchange of KCl for NaCl; termed KPSS) and the response recorded. A new baseline was established when the high-potassium solution was washed out. The vessel was then subjected to either vehicle or 10 µM lisinopril for 15 minutes. Following this treatment, the vessels were subjected to 1 nM Ang I and the peak response recorded. We have previously shown using an ACE-selective substrate, [Pro¹⁰]Ang I, that this concentration of the ACE inhibitor is sufficient to fully inhibit ACE activity in isolated tissue preparations (32). Arteries were exposed to Ang I for 4 minutes or until a peak response had been reached, whichever was sooner, because tachyphylaxis developed even after a single dose of Ang I. In preliminary experiments (not shown), we found that 1 nM Ang I produced a submaximal vasoconstrictor response. To avoid the effects of tachyphylaxis, each artery preparation was subjected to Ang I only once. Thus, responses to 1 nM Ang I in the presence of 10 µM lisinopril (relative to the maximal KCl effect in that blood vessel) were determined in one set of isolated carotid artery preparations from MC^{+/+} mice, and responses to 1 nM Ang I in the absence of ACE inhibitor pretreatment (again, relative to the maximal KCl effect in that blood vessel) were obtained in a second set of isolated carotid artery preparations from MC^{+/+} mice. The results are expressed as a percentage of that effect caused by 1 nM Ang I in the absence of ACE inhibitor pretreatment. A parallel approach was used to study the effect of 10 µM lisinopril pretreatment on the vasoconstrictor response to 1 nM Ang I in isolated carotid artery preparations from Kit^w/Kit^{w-v} mice.

Measurement of [Pro¹¹,DAla¹²]Ang I and Ang II responses in isolated mesenteric artery preparations. Using procedures described previously (33), mesenteric artery segments (150-200 µm in internal diameter, 2-3 mm in length) were dissected from excised guts of 7-week-old MC^{+/+} or Kit^w/Kit^{w-v} mice anesthetized with a mixture of 100 mg/kg ketamine and 20 mg/kg xylazine. Mesenteric arteries were mounted in a pressure myograph (Living Systems Instrumentation, Burlington, Vermont, USA), and their diameters were measured by video microscopy (33). Arteries were maintained in a PSS at 37°C, pH 7.4. The pO₂ was 160 mmHg and the pCO₂ 37 mmHg. The intraluminal pressure was set at 60 mmHg. Arterial diameters were recorded continuously. Vessels were allowed to stabilize for at least 30 minutes. Thereafter, each isolated mesenteric artery segment from an MC+/+ or Kit^w/Kit^{w-v} mouse was exposed to either 1 µM [Pro¹¹, DAla¹²]Ang I or 1 µM Ang II. Each vessel was exposed only once to an angiotensin. In initial studies, we determined that 1-µM concentrations of these peptides produced maximal responses. Results are expressed as maximal reductions in resting diameter. In MC^{+/+} mouse isolated mesenteric arteries, in which responses to 1 µM [Pro¹¹,DAla¹²]Ang I were significantly greater than base line, the mechanism of action of the [Pro¹¹,DAla¹²]Ang I effect was further explored by determination of the effect of AT_1 receptor antagonism by 10 μ M losartan on the [Pro¹¹,DAla¹²]Ang I response. [Pro¹¹,DAla¹²]Ang I was purchased from Bachem California Inc.

Induction of pressure overload in MC^{+/+} or Kit^w/Kit^{w-v} mice. Pressure overload was produced by constriction of the abdominal aorta (SAC). Mice were anesthetized by intraperitoneal injection of a mixture of 100 mg/kg ketamine and 20 mg/kg xylazine. The abdominal aorta was banded just above the renal vessels with a 7-0 silk suture tied around both the aorta and a blunted 29-gauge needle; the needle was then immediately withdrawn. All animals were given drinking water and food ad libitum. Eight-week-old male mice were divided into the following groups: (a) MC^{+/+} or Kit^w/ Kit^{w-v} mice allowed to recover for 3 days after a sham operation; (b) MC^{+/+} or Kit^w/Kit^{w-v} mice allowed to recover for 3 days after SAC; (c) MC^{+/+} or Kit^w/Kit^{w-v} mice allowed to recover for 3 days after SAC but given approximately 50 mg/kg/d captopril from the time of SAC. Captopril was given in the drinking water, and the concentration in the drinking water was adjusted to about 50 mg/kg/d captopril based on the volume of fluid consumed.

Blood pressure measurements. Proximal aortic blood pressure and dP/ dt measurements were performed as described by us previously (34), in mice anesthetized intraperitoneally with 100 mg/kg ketamine and 20 mg/kg xylazine. The right femoral vein was cannulated for infusions (0.1 μ l/g body weight, in <5 minutes) of captopril (2.5 mg/

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kg), losartan (80 mg/kg), or a combination of losartan (80 mg/kg) and captopril (2.5 mg/kg). Maximal responses were recorded.

Statistics. Data are presented as mean ± SEM. Statistical analysis was performed using the unpaired Student's *t* test or Scheffe's test after ANOVA indicated significant differences. *P* values less than 0.05 were considered significant.

Acknowledgments

This work was supported in part by National Health and Medical Research Council of Australia grant 109001.

Received for publication December 12, 2003, and accepted in revised form May 4, 2004.

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