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Activation of the type 1 angiotensin II receptor (AT₁) triggers proinflammatory signaling through pathways independent of classical Gq signaling that regulate vascular homeostasis. Here, we report that the AT₁ receptor preformed a heteromeric complex with the receptor for advanced glycation endproducts (RAGE). Activation of the AT₁ receptor by angiotensin II (Ang II) triggered transactivation of the cytosolic tail of RAGE and NF-κB–driven proinflammatory gene expression independently of the liberation of RAGE ligands or the ligand-binding ectodomain of RAGE. The importance of this transactivation pathway was demonstrated by our finding that adverse proinflammatory signaling events induced by AT₁ receptor activation were attenuated when RAGE was deleted or transactivation of its cytosolic tail was inhibited. At the same time, classical homeostatic Gq signaling pathways were unaffected by RAGE deletion or inhibition. These data position RAGE transactivation by the AT₁ receptor as a target for vasculoprotective interventions. As proof of concept, we showed that treatment with the mutant RAGE peptide S391A-RAGE₃₆₂–₄₀₄ was able to inhibit transactivation of RAGE and attenuate Ang II–dependent inflammation and atherogenesis. Furthermore, treatment with WT RAGE₃₆₂–₄₀₄ restored Ang II–dependent atherogenesis in Ager/Apoe-KO mice, without restoring ligand-mediated signaling via RAGE, suggesting that the major effector of RAGE activation was its transactivation.

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Activation of the type 1 angiotensin II receptor (AT₁) triggers proinflammatory signaling through pathways independent of classical Gq signaling that regulate vascular homeostasis. Here, we report that the AT₁ receptor preformed a heteromeric complex with the receptor for advanced glycation endproducts (RAGE). Activation of the AT₁ receptor by angiotensin II (Ang II) triggered transactivation of the cytosolic tail of RAGE and NF-κB–driven proinflammatory gene expression independently of the liberation of RAGE ligands or the ligand-binding ectodomain of RAGE. The importance of this transactivation pathway was demonstrated by our finding that adverse proinflammatory signaling events induced by AT₁ receptor activation were attenuated when RAGE was deleted or transactivation of its cytosolic tail was inhibited. At the same time, classical homeostatic Gq signaling pathways were unaffected by RAGE deletion or inhibition. These data position RAGE transactivation by the AT₁ receptor as a target for vasculoprotective interventions. As proof of concept, we showed that treatment with the mutant RAGE peptide S391A-RAGE₃₆₂–₄₀₄ was able to inhibit transactivation of RAGE and attenuate Ang II–dependent inflammation and atherogenesis. Furthermore, treatment with WT RAGE₃₆₂–₄₀₄ restored Ang II–dependent atherogenesis in Ager/Apoe–KO mice, without restoring ligand-mediated signaling via RAGE, suggesting that the major effector of RAGE activation was its transactivation.

Introduction

Blockade of the renin-angiotensin aldosterone system (RAAS) is one of the most prescribed treatments in all of medicine. The pleiotropic benefits of RAAS blockade on inflammation (1–3), atherogenesis (4, 5), and even longevity (6) are additional to and independent of blood pressure lowering (7). Activation of the type 1 angiotensin II receptor (AT₁) receptor by angiotensin II (Ang II) triggers pathogenic pathways that are distinct from classical Gq protein signaling that induces vasoconstriction and sodium retention (2, 3, 8, 9). However, a complete understanding of this fundamental division and, more important, of any means to specifically block the proinflammatory effects of Ang II remains elusive.

The receptor for advanced glycation endproducts (RAGE) is a type I transmembrane glycoprotein. RAGE is basally expressed at high levels in the vascular endothelium and leukocytes, although RAGE expression may be induced in most cell types following injury, inflammation, or hypoxia (10). RAGE is implicated in a range of diseases, including Alzheimer’s disease (11), malignancy (12), diabetic complications (13), and atherosclerosis (14–17). In each case, dysfunction has been ascribed to activation of the RAGE ectodomain by ligands including S100 proteins (also known as calgranulins) and advanced glycation endproducts (AGEs) that trigger NF-κB–driven proinflammatory gene expression (18). However, we provide evidence for a cognate ligand–independent mechanism for RAGE transactivation that occurs following activation of the AT₁ receptor. Using a variety of mouse models, assay systems, and different RAGE domains, we show that the receptors form a plasma membrane–localized heteromeric complex, which, upon activation, induces NF-κB expression and the subsequent canonical and noncanonical signaling that largely determines the proinflammatory effects of RAAS activation.

Results

Atherogenesis following an infusion of Ang II is reduced in Ager/Apoe–double-KO mice. To demonstrate the role of RAGE in Ang II–dependent atherogenesis, we infused atherosclerosis-susceptible Apoe-KO mice with Ang II (1 μg/kg/min) for 4 weeks. This resulted in increased plaque accumulation across the aortic arch in Apoe-KO but not Ager/Apoe–double-KO (Ager/Apoe-DKO) mice (Figure 1A). Exposure to Ang II also increased aortic expression

Authorship note: RJP, CT, and CJR are co-first authors.

Conflict of interest: KDGP is chief scientific adviser of Dimerix Limited, a spin-off company of The University of Western Australia that has been assigned the rights to Receptor-HIT. KDGP is an inventor on patents covering the technology (World Intellectual Property [WIPO] patent no. WO/2008/055313, held by Dimerix Limited) and is a shareholder of Dimerix Limited.

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of proatherogenic mediators (Supplemental Figure 1A; supplemental material available online with this article; https://doi.org/10.1172/JCI99987DS1) in Apoe-KO mice but not Ager/Apoe-DKO mice. However, we found that systolic blood pressure increased to a similar extent in response to Ang II in both the presence and absence of RAGE (Supplemental Table 1), and neither Ang II nor RAGE deficiency modified total cholesterol levels (no significant difference, P > 0.1).

**Atherogenesis following a low-sodium diet is reduced in Ager/Apoe-DKO mice.** In a second model of Ang II–dependent atherogenesis, 6 weeks of exposure to a 0.05% (low) sodium diet also resulted in RAAS-dependent plaque accumulation in Apoe-KO mice (Figure 1B) and increased aortic expression of proatherogenic mediators (Supplemental Figure 1B). More labeled leukocytes were adherent to the endothelium of aortae taken from Apoe-KO mice after 1 week of exposure to a low-sodium diet when compared with aortae from Apoe-KO mice fed a normal chow diet (Figure 1C). In contrast, we observed that the same low-sodium diet fed to Ager/Apoe-DKO mice did not modify plaque accumulation (Figure 1B) or markers of vascular inflammation (Supplemental Figure 1B), despite comparable activation of the systemic RAAS and sodium retention (Supplemental Figure 1, E–G). Additionally, we found that Ager/Apoe-DKO mice fed a low-sodium diet did not have increased endothelial adhesion of labeled leukocytes (Figure 1C). Blood pressure levels were not reduced by a low-sodium diet in either Apoe-KO or Ager/Apoe-DKO mice (Supplemental Table 1), and a low-sodium diet had no effect on total cholesterol levels (no significant difference, P > 0.1).

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**Atherogenesis in Ace2/Apoe-DKO mice is reduced in Ager/Ace2/Apoe–triple-KO mice.** Atherosclerosis was also explored in a third model of Ang II–dependent atherogenesis, this time using Ace2/Apoe-DKO mice. We found that genetic deficiency of Ace2 in Apoe-KO mice was associated with increased circulating levels of Ang II (Supplemental Figure 1H), elevated systolic blood pressure (Supplemental Table 1), and augmented plaque accumulation (Figure 1D), as previously described (19). The aortae of Ace2/Apoe-DKO mice also had increased expression of proatherogenic mediators (Supplemental Figure 1C) when compared with aortae of Apoe-KO mice. In contrast, the induction of proinflammatory markers (Supplemental Figure 1C) and plaque accumulation (Figure 1D) were attenuated in Ager/Ace2/Apoe–triple-KO (Ager/Ace2/Apoe-TKO) mice, despite a comparable elevation in circulating Ang II levels (Supplemental Figure 1H) and blood pressure (Supplemental Table 1), and Ace2 deficiency had no effect on total cholesterol (no significant difference, P > 0.1).
While RAGE deletion attenuated atherogenesis in RAAS-dependent models, it could be argued that this was due to non-specific antiatherogenic actions of RAGE deletion. However, after 6 weeks of a cholate-containing high-fat (Paigen) diet, a lipid-dependent model of atherosclerosis (20), we found that aortic plaque accumulation was increased in both Aopoe-KO and Ager/Aopoe-DKO mice, with no significant difference between them (Figure 1E).

The induction of proinflammatory mediators by Ang II is dependent on RAGE expression. In isolated aortae from Aopoe-KO mice, ex vivo exposure to Ang II (1 μM) also increased the expression of proatherogenic mediators (Supplemental Figure 1D) and increased the number of adherent leukocytes when compared with untreated aorta (Figure 1F). In contrast, ex vivo treatment of aortae taken from Ager/Aopoe-DKO mice with Ang II did not induce these atherogenic mediators (Supplemental Figure 1D) or increase leukocyte adhesion (Figure 1F).

In primary murine aortic endothelial cells (PMAECs) isolated from the aortae of C57BL/6j mice, exposure to Ang II resulted in a rapid increase in hydrogen peroxide (Figure 2A). This was associated with the induction of canonical and noncanonical signalling via NF-κB-dependent pathways (Figure 2B), leading to increased expression of key proinflammatory genes (Figure 2C) and, functionally, the induction of a proatherogenic phenotype associated with increased adhesion of labeled THP-1 cells to a monolayer of Ang II-treated PMAECs (Figure 2D). We did not observe these Ang II-induced changes in PMAECs isolated from Ager-KO mice. Likewise, silencing of Ager or NF-κB-p65 expression in PMAECs using siRNA (Supplemental Figure 2, A–C) also prevented the NF-κB-dependent induction of ICAM-1 by Ang II or the RAGE ligand S100A8/A9 (Figure 2E). The observed attenuation of Ang II effects in PMAECs from Ager-KO mice was not due to a loss of RAAS-signaling capacity in the absence of RAGE, as exposure of RAGE-deficient PMAECs to Ang II led to a similar increase in inositol monophosphate (IP) (Figure 2F) and downstream induction of the early growth response gene Egr1 (Supplemental Figure 3A). In addition, we found that the classical induction of canonical NF-κB signaling by TNF-α was not attenuated in PMAECs from Ager-KO mice (Figure 2B). The role of RAGE in Ang II-induced proinflammatory signaling was not specific to endothelial cells and was also observed in Ang II-induced activation of macrophages or splenocytes (Supplemental Figure 4, A–C).

To specifically explore the signaling interaction between RAGE and the AT receptor, we transfected CHO cells with AGTR1 and AGER, alone or in combination and observed that expression of the AT1 receptor in CHO cells conferred responsiveness to exogenous Ang II, as demonstrated by the induction of inositol phosphate synthesis (Figure 3A). This was not modified by the coexpression of full-length RAGE. In contrast, NF-κB activation and induction of p65 expression following exposure to Ang II in AT1-CHO cells was dependent on RAGE coexpression (Figure 3B). Similar to this, RAGE expression was also required for the proinflammatory response to the RAGE ligand S100A8/A9 in CHO cells (Figure 3C) and PMAECs (Supplemental Figure 5A).

Ligand-independent transactivation of RAGE following activation of the AT1 receptor by Ang II. Activation of the AT1 receptor is known to trigger transactivation of tyrosine kinase receptors, with one proposed mechanism being protease-mediated extracellular “shedding” of membrane-tethered ligands (21). Activation of the RAAS increased circulating concentrations of RAGE ligands in mice, including S100A8/A9 and AGEs and their reactive dicarbonyl precursors (Supplemental Figure 4, D–F), as well as increased expression of Mac1 (also known as CD11b) in the aorta (Supplemental Figure 1, A–C). However, Ang II–induced proinflammatory signaling was not inhibited by pretreatment with a neutralizing antibody targeting the extracellular ligand–binding domain of RAGE (RAGEab), although RAGE ligand–dependent signaling was fully blocked by this strategy (Figure 3D and Supplemental Figure 5B). Likewise, deletion of the ligand-binding extracellular domain of RAGE did not modify the ability of N-truncated RAGE constructs to mediate Ang II–dependent induction of p65 expression or NF-κB activity in AT1-CHO cells, such that full-length RAGE activity in this assay was retained even by RAGEΔ70-404 (Figure 3E). In contrast, the induction of p65 expression in AT1-CHO cells by S100A8/A9 was lost in cells expressing N-truncated RAGE constructs in which the ligand-binding ectodomain was also deleted (Supplemental Figure 3B).

No role for the phosphorylation of RAGE following activation of the AT1 receptor. The human RAGE cytosolic tail contains 4 potential phosphorylation sites, only 1 of which (S391) is partially conserved across other mammals (Supplemental Table 2). We observed that mutation of S399, S400, or T401 to alanine had no effect on signaling (Supplemental Figure 3C) and that signaling was also retained in E392X-RAGE (Figure 3F), despite the lack of 13 C-terminal residues (E392–P404). In contrast, full-length or C-truncated RAGE constructs in which S391 was deleted (S391X-RAGE) (Figure 3F) or changed to alanine (S391A-RAGE) (Figure 4A) had no ability to mediate proinflammatory signaling in response to Ang II in AT1-CHO cells. However, despite lacking a serine at 391, full-length S391Q-RAGE and S391P-RAGE mutants retained the ability to signal in response to Ang II and S100A8/A9 in AT1-CHO cells (Figure 4B). This was not through redundancy offered by other potential phosphorylation sites, as replacement of the 14 C-terminal amino acids in human RAGE (S399–P403) with those found in murine RAGE (S391–S396, T397–T402, and P403) created a chimeric RAGE (cRAGE), with or without additional S391Q mutation (i.e., Q391EMPGAGGP), did not prevent signaling in response to Ang II or S100A8/A9 (Figure 4C). Moreover, mutation of the C-terminal to further remove potentially phosphomimetic residues (glutamate or aspartate) in addition to all serines and threonines (i.e., PKGPQAGQGGAGGP; no STED) also did not prevent Ang II–dependent signaling via RAGE in AT1-CHO cells (Supplemental Figure 3D).

Although phosphorylation of RAGE was not required for signaling, selective inhibition of PKCζ with a siRNA (siPKCζ) (Supplemental Figure 2D) or a PKCζ pseudosubstrate (PKCζi) prevented the proinflammatory signaling induced by S100A8/A9 and Ang II in PMAECs (Figure 4D). We also observed the inhibitory effects of PKCζi expression in AT1-CHO cells expressing S391Q-cRAGE, despite the absence of serine/threonine targets for phosphorylation in this RAGE construct (Figure 4E). Silencing of PKCζ expression (Supplemental Figure 2E) also inhibited signaling induced in human microvascular endothelial cells (HMEC1) (Figure 4F), which was rescued by overexpression of the purely cytosolic RAGEΔ143-404 but not of the membrane-bound RAGEΔ143-404 (Figure 4F).
Role of diaphanous-1 in signaling via RAGE. The formin-type protein diaphanous-1 (Diaph1) is regarded as a key element for proinflammatory signaling induced by RAGE ligands (22). Consistent with these data, we observed that silencing of Diaph1 expression (Supplemental Figure 2F) prevented the induction of key proinflammatory genes in response to the RAGE ligand S100A8/9 in PMAECs as well as RAGE ligand–independent induction of ICAM-1 (Figure 4, F and G) and leukocyte adhesion to an endothelial monolayer (Supplemental Figure 2H) following exposure to Ang II. Diaph1 and RAGE are thought to interact specifically via a charged patch formed by R366 and Q367 (22), and we observed that R366A-Q367A-RAGE constructs were not able to mediate S100-induced signaling in AT1-CHO cells (Figure 4H), as previously reported (22). In contrast, the R366A-Q367A-RAGE mutation did not alter the ability of full-length or N-truncated RAGE constructs to mediate the proinflammatory signaling induced by Ang II in AT1-CHO cells (Figure 4H). Indeed, the transactivation of a construct lacking this domain (RAGE 370–404) following treatment with Ang II in AT1-CHO cells (Figure 3E) suggests that, unlike RAGE ligand–dependent signaling, this domain is not essential for RAGE ligand–independent signaling. Nonetheless, the finding that silencing of Diaph1 expression in HMEC1 inhibited signaling by endogenous RAGE (Figure 4G) and membrane-bound RAGE343–404 (Figure 4F) demonstrates that Diaph1 still plays a critical role. The finding that silencing of Diaph1 expression did not inhibit transactivation of the purely cytosolic construct RAGE362–404 (Figure 4F) suggests that, like PKCζ, a key target of Diaph1 may also be the single transmembrane domain of RAGE.

Bioluminescence resonance energy transfer indicates close proximity of the AT1 receptor and RAGE when coexpressed in live cells. Constr
indicative of their close proximity, which was reduced upon treatment with Ang II (Figure 5A). Ang II treatment in cells coexpressing AT1-Rluc8 and the signaling and regulatory protein β-arrestin2 tagged with Venus induced a robust and stable ligand-induced BRET signal consistent with recruitment of β-arrestin2 to the AT1 receptor.

Figure 3. Functional response to Ang II or S100A8/A9 in the presence or absence of the human AT1 receptor, RAGE constructs, or both. Treatment with Ang II was at a dose of 1 μM for 2 hours, and treatment with S100A8/A9 was at a dose of 2 μg/ml for 2 hours. (A) Induction of inositol phosphate synthesis. (B) NF-κB activation as measured by a chemiluminescent SEAP reporter gene assay, and induction of gene expression of the NF-κB subunit p65 in the presence or absence of full-length human RAGE. (C) Induction of p65 expression following exposure to the RAGE ligand S100A8/A9 (2 μg/ml for 2 h) in the presence or absence of full-length RAGE. (D) Induction of p65 following exposure to S100A8/A9 or Ang II in PMAECs in the presence or absence of soluble RAGE or a neutralizing antibody targeting the extracellular domain of RAGE. (E) NF-κB activation and the induction of p65 expression in the presence or absence of full-length or N-truncated RAGE constructs in AT1-CHO cells. (F) NF-κB activation and the induction of p65 expression by Ang II in the presence or absence of full-length or C-truncated RAGE constructs in AT1-CHO cells. Data are presented as the mean ± SD. *P < 0.05 versus AT1-CHO plus vector (pCIneo or mCherry); ‡P < 0.05 versus AT1-CHO plus full-length RAGE; †P < 0.05 versus AT1-CHO plus full-length RAGE plus Ang II; §P < 0.05 versus AT1-CHO plus RAGE plus S100A8/A9. P values were determined by 2-way ANOVA.
receptor at the cell surface (Figure 5B) (23). Using the receptor-HIT configuration (24, 25), treatment with Ang II also induced a BRET signal in cells coexpressing RAGE-Rluc8 and β-arrestin2–Venus in the presence of the untagged AT1 receptor, again consistent with this interaction occurring at the cell surface. Notably, we observed no BRET following activation of the CCR4 chemokine receptor by CCL22 (Figure 5C). This was despite the fact that CCR4 was still able to activate G protein (Figure 5D) and trigger the recruitment
of β-arrestin2–Venus following its activation by its cognate ligand (Figure 5B). As further evidence of the functional interaction between RAGE and the AT₁ receptor, we found that RAGE-RluC8 cellular trafficking was induced upon activation of the AT₁ receptor (Figure 5E), as shown by its proximity to Venus-tagged markers of specific cellular compartments (26).

An Ang II–induced BRET signal was still observed between N-terminal mCherry-tagged RAGE₃₃₈–₃₆₁ (the transmembrane and juxtamembrane domains only) and an extracellular luciferase–tagged AT₁ receptor (Nluc-AT₁) (Figure 5F), with the intracellular luciferase–tagged AT₁ receptor (AT₁-Nluc) serving as a negative control for this experiment. Moreover, overexpression of unlabelled RAGE₃₆₂–₄₀₄ inhibited the BRET between full-length RAGE and the AT₁ receptor (Figure 5G). By contrast, RAGE₃₆₂–₄₀₄ had no effect on the BRET between RAGE-Venus and AT₁-Nluc (Figure 5H). To elucidate the functional involvement of the transmembrane domain of RAGE in Ang II–mediated signaling, RAGE₃₃₈–₃₆₁ was overexpressed in HMEC1. This resulted in inhibition of the Ang II–induced ICAM-1 expression mediated via endogenous RAGE, which was able to be rescued by RAGE₃₆₂–₄₀₄ (Figure 5I). This suggests that, while the cytosolic portion of RAGE is necessary for mediating downstream Ang II–induced proinflammatory signaling, the transmembrane domain of RAGE may also be involved in a separate interaction with the AT₁ receptor.

Modulation of RAGE transactivation with RAGE oligopeptides. Although transfection with S391A-RAGE₃₆₂–₄₀₄, or RAGE₃₇₀–₃₉₆, constructs did not have the capacity to be transactivated in AT₁-CHO cells, transfection of HMEC1 with these constructs prevented the induction of ICAM-1 and MCP-1 expression by Ang II via endogenous RAGE in HMEC1 (Figure 6A). To validate the therapeutic potential of directly modulating RAGE ligand–independent signaling via transactivation of RAGE, we generated oligopeptides comprising the C-terminal of RAGE tagged with mCherry fluorescent protein and an HIV-TAT motif to facilitate cellular penetration. Treatment of AT₁-CHO cells with mCherry-RAGE₃₆₂–₄₀₄ (0.4 ng/ml) was able to induce de novo proinflammatory responsiveness to Ang II (Figure 6B) in the absence of RAGE gene expression. Similarly, in PMAECs from Ager/KO mice, we observed that treatment with mCherry-RAGE₃₆₂–₄₀₄ was able to restore proinflammatory responsiveness to Ang II (Figure 6C). Treatment with mCherry-RAGE₃₆₂–₄₀₄ was also able to rescue the inhibition of signaling achieved following treatment with siRNA targeting Ager, Diaph1, or PKCζ in HMEC1 (Supplemental Figure 2E and G and Supplemental Figure 5C).

In contrast to the above findings, treatment of RAGE-replete PMAECs with mCherry-S391A-RAGE₃₆₂–₄₀₄ inhibited the Ang II–mediated induction of NF-κB–dependent genes (Figure 6D). Likewise, mCherry-S391A-RAGE₃₆₂–₄₀₄ inhibited the Ang II–mediated expression of ICAM-1 in HMEC1 (Figure 6E) and HAEcs (Supplemental Figure 5D), which was comparable to that achieved with the AT₁ receptor antagonist irbesartan. We found that pretreatment with mCherry-S391A-RAGE₃₆₂–₄₀₄ also inhibited Ang II–induced signaling in AT₁-CHO cells also overexpressing full-length RAGE or RAGE₃₆₂–₄₀₄ but failed to inhibit signaling mediated by shorter RAGE constructs (Supplemental Figure 5E). We observed that mCherry-S391A-RAGE₃₆₂–₄₀₄ also inhibited Ang II–induced signaling in AT₁-CHO cells expressing S391Q–cRAGE, a construct devoid of targets for phosphorylation (Supplemental Figure 5F). Pretreatment of AT₁-CHO cells with mCherry-RAGE₃₆₂–₄₀₄ did not prevent the inhibitory effects of mCherry-S391A-RAGE₃₆₂–₄₀₄ (Supplemental Figure 6A). Moreover, the inhibition of Ang II–dependent signaling achieved by S391A-RAGE₃₆₂–₄₀₄ was not reversed by mCherry-RAGE₃₆₂–₄₀₄ even at 1000-fold excess (Supplemental Figure 6B). mCherry-S391A-RAGE₃₆₂–₄₀₄ also inhibited RAGE ligand–dependent signaling in PMAECs (Supplemental Figure 6C), but had no effect on ICAM-1 expression induced by TNF-α (Supplemental Figure 6D).

Discussion

Activation of the AT₁ receptor by Ang II leads to the induction of NF-κB (2, 3) and subsequent inflammation through pathways distinct from those mediating classical Gq-induced signaling (8, 9). Here, we show that RAGE ligand–independent transactivation of RAGE following activation of the AT₁ receptor was a key determinant of this division. Consequently, many of the proinflammatory changes induced following AT₁ receptor activation were attenuated when RAGE was deleted or when RAGE transactivation was inhibited. Also, the proinflammatory effects of AT₁ receptor activation were enhanced in the presence of RAGE. In contrast, we observed that AT₁ receptor–dependent signaling via classical Gq pathways was unaffected by RAGE expression and that RAGE deletion had no deleterious effect on vascular homeostasis in the setting of salt deficiency or blood pressure responsiveness to Ang II.

Non-specific crosstalk between the AGE/RAGE axis and the RAAS has long been known, and this crosstalk establishes a vicious cycle for injury and dysfunction (27–34). For example, RAGE ligand–dependent activation of RAGE induces molecular pathways in common with RAAS signaling, including tyrosine phosphorylation of JAK2 and activation of PKC, NF-κB, and
Figure 5. The AT₁ receptor and RAGE form a functional heteromeric complex. (A) BRET saturation curves with AT₁-Rluc8 and RAGE-Venus generated 60 minutes after the addition of Ang II or vehicle. (B) Ang II–induced recruitment of β-arrestin2–Venus to AT₁-Rluc8 and CCL22-induced recruitment of β-arrestin2–Venus to CCR4-Rluc8 as a control. (C) Ang II–induced recruitment of β-arrestin2–Venus proximal to RAGE-Rluc8 only in the presence of AT₁, following exposure to Ang II, and not in the presence of CCR4 following exposure to CCL22. (D) CCL22-induced BRET signal was observed when Gαi-Nluc and Gγ2-Venus were coexpressed in the presence of untagged CCR4, serving as an additional control. (E) Ang II–induced trafficking of RAGE-Rluc8 in the presence of AT₁ using markers of specific subcellular compartments. (F) BRET between mCherry-RAGE_{338-361} and Nluc-AT₁ was increased with Ang II. (G) BRET between AT₁-Rluc8 and RAGE-Venus was inhibited by mCherry-RAGE_{338-361}. (H) BRET between AT₁-Nluc and RAGE-Venus was reduced by Ang II but not affected by mCherry-RAGE_{362-404}. (I) Ang II–mediated proinflammatory signaling in HMEC1 inhibited by RAGE_{338-361} was rescued by mCherry-RAGE_{362-404}.

Data are presented as the mean ± SD. n = 3 (B–D and H), n = 5 (E and F), n = 6–8 (I), or data were combined from 3 independent experiments (A and G).

*P < 0.05 versus vehicle or pretreatment or 0 minutes; #P < 0.05 versus other treated conditions. P values were determined by 2-way ANOVA.
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Figure 6. Effects of treatment with mCherry-RAGE362–404 oligopeptides on proinflammatory signaling following exposure to Ang II in vitro, ex vivo, and in vivo. Treatment with Ang II was at a dose of 1 μM. (A) Induction of ICAM-1 and MCP-1 expression in HMEC1 in the presence or absence of plasmids expressing truncated RAGE constructs. (B) Induction of p65 expression in AT1-CHO cells treated with mCherry-RAGE338–361 or mCherry-S391A-RAGE362–404 (0.4 ng/ml). (C) Expression of proinflammatory genes in PMAEcs from Ager-KO mice in the presence or absence of mCherry-RAGE362–404 (0.4 ng/ml). (D) Expression of proinflammatory genes in (RAGE-replete) PMAEcs induced by Ang II in the presence or absence of mCherry-S391A-RAGE362–404 (0.4 ng/ml). (E) Induction of ICAM-1 expression in PMAEcs in the presence or absence of mCherry-S391A-RAGE362–404 (0.4 ng/ml) or ibesartan (1 μM). (F) Induction of ICAM-1 expression in aortae from Apeko-CHO and Ager/Apoe-DKO mice in the presence or absence of mCherry-RAGE362–404 or mCherry-S391A-RAGE362–404 (8 ng/ml). (G) Plaque area in aortae from Apeko-CHO, Ace2/Apoe-DKO, and Ager/Ace2/Apoe-TKO mice in the presence or absence of mCherry-RAGE362–404 or mCherry-S391A-RAGE362–404 (10 μg/kg i.p. every other day). (H) Plaque area in aortae from diabetic Apeko-CHO and Ager/Apoe-DKO mice in the presence or absence of mCherry-RAGE362–404 or mCherry-S391A-RAGE362–404 (10 μg/kg i.p. every other day). Data are presented as the mean ± SD. n = 6–8 groups. *P < 0.05 versus vehicle or cPcS1 plus Ang II; †P < 0.05 versus Ang II; ††P < 0.05 versus mCherry plus Ang II; †¥P < 0.05 versus mCherry-RAGE362–404 plus Ang II; **P < 0.05 versus Ang II; ‡‡P < 0.05 versus Ang II plus mCherry; ††P < 0.05 versus Ang II plus mCherry-RAGE362–404. P values were determined by 2-way ANOVA.

We revealed additional evidence for the functional interaction between the 2 receptors with the BRET trafficking assay, in which coexpression and activation of the AT1 receptor resulted in trafficking of RAGE-Rlu8 through various cellular compartments, including early endosomes (Rab5), recycling endosomes (Rab11), and forward trafficking from the trans-Golgi network to the plasma membrane (Rab8), consistent with the strong expression of RAGE on the plasma membrane proximal to the Kras marker. This interaction between RAGE and the AT1 receptor appears to be dependent on the transmembrane domain of RAGE, such that BRET was demonstrated between Nluc-AT1 and mCherry-RAGE362–404 and unlabeled RAGE358–561 inhibited the BRET between full-length RAGE and the AT1 receptor. In contrast, the extracellular and cytosolic domains of RAGE were unable to modulate the BRET signal between the AT1 receptor and full-length RAGE. Interestingly, we observed that overexpression of transmembrane RAGE358–561 also inhibited Ang II–dependent signaling by endogenous RAGE in HMEC1, although this inhibition was overcome by cytosolic RAGE362–404. This indicates that, when free from the constraints of being membrane bound, the cytosolic domain of RAGE independently retains the capacity to be transactivated by the AT1 receptor and associated proteins. However, while the 2 domains could function separately, they were mutually dependent for signaling via endogenous membrane-bound RAGE.

Previous studies have suggested that PKCζ-mediated phosphorylation of S391 on RAGE is a key step in RAGE activation (35). Among the 4 potential phosphorylation sites in human RAGE, S391 is conserved among humans, mice, horses, rabbits, dogs, and cats (Supplemental Table 2). However, S391Q-RAGE (as found in camels) and S391P-RAGE (as found in cattle, pigs, sheep, and goats) are both able to be activated by Ang II or S100A8/9, even following the additional mutation of the other possible phosphorylation targets in the cytosolic tail, demonstrating that phosphorylation of RAGE is not essential for its activation. PKCζ is known to associate with the cytosolic domain of RAGE and was thought to mediate ligand-mediated phosphorylation of RAGE (35). PKCζ certainly appeared to be critical for transactivation of full-length RAGE by the AT1 receptor, as selective inhibition of PKCζ was able to prevent activation of NF-kB following exposure to Ang II. However, inhibition of PKCζ also prevented activation of S391Q-RAGE by Ang II or S100A8/9, demonstrating that PKCζ was functioning independently of RAGE phosphorylation. In contrast, signaling via RAGE362–404 was not prevented following silencing of PKCζ, suggesting that PKCζ plays no direct role in transactivation of the cytosolic tail of RAGE.

The interaction of RAGE with Diaph1 is also thought to be important for ligand-dependent signaling (22, 36), possibly because it facilitates the recruitment and Ras-dependent activation of PKCζ, with which Diaph1 also binds (37). In our experiments, silencing of Diaph1 expression also inhibited Ang II–induced proinflammatory signaling via full-length RAGE. However, the role of Diaph1 in RAGE ligand–dependent activation also appears to be different from its role in RAGE ligand–independent transactivation. Specifically, mutation of the charged patch (R366A-Q367A-RAGE) prevents signaling by RAGE ligands, as previously described (22, 36), but has no effect on Ang II–induced RAGE transactivation. Indeed, the activity of.

NADPH oxidase (27–29). Blockade of the RAAS has also been shown to attenuate the effects of AGE-induced activation of RAGE (30–33). Likewise, dicarboxyl scavengers, which reduce the formation of RAGE-activating AGEs, attenuate Ang II–dependent injury (34). In certain cell types, Ang II–mediated signaling is known to transactivate growth receptor tyrosine kinases with a proposed mechanism being metalloprotease-mediated shedding of membrane-tethered ligands (21), and RAAS activation certainly increases RAGE ligands. We had initially speculated that this kind of extracellular pathway may exist for RAGE transactivation. However, we found that a neutralizing antibody targeting the ligand-binding domain of RAGE or a soluble decoy receptor had no effect on Ang II–mediated signaling in vitro. Moreover, deletion of the extracellular domain of RAGE did not modify the ability of N-truncated RAGE constructs to transduce the proinflammatory effects of Ang II. Indeed, we observed that treatment with RAGE362–404 fully restored the Ang II–dependent induction of inflammatory mediators in vitro and ex vivo and atherogenesis in vivo in the absence of full-length RAGE. Nonetheless, the (RAGE-dependent) liberation of RAGE ligands following activation of the AT1 receptor may play a latent role in sustaining or enhancing proinflammatory signaling via RAGE, including the autoinduction of RAGE expression (13). This may partly explain why soluble RAGE, comprising only the ligand–binding domain of RAGE, has some vasculoprotective effects in vivo (16), although other actions including the antagonism of ligand–mediated activation of TLRs may also contribute to this protection.

Here, we provide evidence that RAGE and the AT1 receptor form a heteromeric complex at the plasma membrane, as demonstrated by the generation of a strong and saturable BRET signal between AT1–Rlu8 and RAGE-Venus, as well as the receptor-HIT signal with RAGE–Rlu8 observed when β-arrestin2–Venus was recruited to the AT1 receptor following Ang II–induced activation.

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full-length RAGE is retained by RAGE\textsubscript{370-404} despite the absence of the charged patch in this construct. It may be that AT\textsubscript{1} receptor activation is able to independently induce the activation of Diaph1. Diaph1 is a known downstream effector of Rho-GTPases (38), and Rho-A, a key product of AT\textsubscript{1}, receptor activation, is known to trigger the activation of Diaph1, even in the presence of the R366A-Q367A RAGE mutation (39). However, we found that RAGE\textsubscript{362-404} was not inhibited following the silencing of Diaph1 expression, similar to what we observed following the silencing of PKC\(\zeta\) (Figure 4F). This suggests that the role of RAGE and PKC\(\zeta\) in the propagation of proinflammatory signaling may also involve the transmembrane domain, possibly to regulate membrane localization, similar to what was reported for translocation of the RAGE-binding partner Src (40).

Only some cells (e.g., endothelial cells, leukocytes) express RAGE under basal conditions, meaning that the transactivation pathway described in this report is basally limited to these cell types. Most other cells do not express RAGE, including healthy smooth muscle cells (10) and HepG2, a cell type in which Ang II notably fails to induce canonical NF-\(\kappa\)B signaling (41). In addition to the fact that RAGE expression had no effect on the classical Gq signaling induced following AT\textsubscript{1} receptor activation, this limited basal distribution of RAGE may partly explain why RAGE deletion or inhibition did not affect blood pressure, natriuresis, or other homeostatic functions of the systemic RAAS. However, following injury, inflammation, or hypoxia, the de novo or enhanced expression of RAGE may provide a conduit for the proinflammatory effects of local RAAS activation. In such states, the blood pressure–independent benefits of RAAS blockade may be partly attributable to the inhibition of RAGE transactivation and associated RAGE autoinduction. However, the efficacy of systemic RAAS blockade is often constrained by the limits of blood pressure lowering (e.g., following a myocardial infarction or in severe heart failure) and feedback “escape” activation of the RAAS (42). One potential advantage of targeting RAGE transactivation is that these key limitations of systemic RAAS blockade do not apply.

As a proof of principle, we designed cell-penetrating RAGE oligopeptides that differed by only 1 amino acid and had the capacity to independently mediate (RAGE\textsubscript{362-404}) or selectively inhibit (S391A-RAGE\textsubscript{362-404}) Ang II–induced activation of proinflammatory pathways contributing to RAAS-dependent atherosclerosis. As an inhibitory strategy, the blockade achieved was as effective as an AT\textsubscript{1} receptor antagonist in vitro, but without adverse effects on hemoostasis in vivo. Inhibition with S391A-RAGE\textsubscript{362-404} was achieved at low concentrations and not overcome by WT RAGE\textsubscript{362-404} even at 1000-fold excess. Taken together, these data suggest that conformational changes induced following substitution or deletion of S391 make it more adept at (or less constrained in) interacting with key elements in the RAGE transactivation pathway. Additional mutation or deletion of the Diaph1-binding domain may further reduce structural constraints and improve binding efficiency, explaining how RAGE\textsubscript{370-404} could overcome the inhibition conferred by S391A-RAGE\textsubscript{362-404} over full-length RAGE.

Even though ligand binding to the ectodomain has historically been considered essential for the functions of RAGE (43), the RAGE ligand-independent transactivation pathway described in our study appears to be a dominant mechanism inducing its activation and downstream effector signaling (39). Consistent with this hypothesis, we found that treatment with RAGE\textsubscript{362-404} fully restored the susceptibility of Ager-KO mice to atherogenesis, without restoring the signaling pathways mediated by elevated RAGE ligands, such as AGEs and S100 proteins. This suggests that the relative vasculoprotection observed in Ager-KO mice was largely mediated through inhibition of RAGE transactivation, highlighting its pathophysiological importance and providing a clear rationale for its inhibition. We hypothesize that a mechanistic understanding of the AT\textsubscript{1}-RAGE heteromer will lead to the development of novel therapeutics that are able to specifically target the adverse effects of RAGE transactivation, without compromising blood pressure regulation or inducing feedback “escape” that limits conventional RAAS blockade. These inhibitors may be applicable across a broad range of conditions in which RAGE has been implicated, including atherosclerosis (14), neurodegenerative diseases (11, 44), malignancy (12), diabetic complications (13, 14), and other important inflammatory conditions.

Methods

Animal models

To investigate the impact of RAGE in RAAS-dependent pathology in vivo, we used the Apoe-KO mouse, which is a model associated with enhanced atherosclerotic plaque accumulation following RAAS activation and vasculoprotection with RAAS blockade (4, 5, 7, 19). However, the unusual dyslipidemia, the absence of spontaneous plaque rupture, and the independent proinflammatory effects of Ape deletion potentially limit the applicability of this mouse model to human disease. C57BL/6J mice, Ager-KO mice, Apoe-KO mice, Ager/ApoD-KO, Ager/Acet2/ApoD-KO, and Ager/Acet2/ApoD-TKO mice (all backcrossed >10 generations) were sourced and generated in house, as previously described (14, 43, 45). All mice were bred on a C57BL/6J background. For the experimental studies, male mice aged 6–8 weeks and weighing between 20 and 25 g were used, with at least 8 animals per group.

Ang II infusion. Apoe-KO and Ager/ApoD-KO mice were randomized to receive Ang II (1 \(\mu\)g/kg/min; MilliporeSigma) or vehicle via s.c. infusion for 4 weeks using an osmotic minipump (model 2004, ALZET, BioScientifica) placed into the s.c. space along the dorsal midline of each mouse, under general anesthesia.

Low-sodium diet. Exposure to a low-sodium diet (0.05% sodium) is an experimental model of physiological RAAS activation (46). In this study, Apoe-KO and Ager/ApoD-KO mice were randomized to receive an isocaloric diet containing 6% fat and either low sodium (0.05%) or normal sodium (0.3%) amounts (Specialty Feeds) for 1 week or 6 weeks.

After 1 week of study, Apoe-KO and Ager/ApoD-KO animals were humanely euthanized and the aortae were isolated, cleaned of fat, and mounted in a vessel chamber primed with Krebs buffer and maintained at a physiological pH by infusing carbogen gas (95% O\(_2\), 5% CO\(_2\)) through the buffer at 37°C (19). Vessels (\(n = 6/\text{group}\)) were pretreated with Ang II (1 \(\mu\)M) or vehicle for 4 hours at 37°C. Whole blood labeled with DilC18 (1:1000) was then perfused through the aorta at 0.12 ml/minute. Images and videos of vessel wall–cell interactions were acquired using a fluorescence microscope (Zeiss Discovery. V20), coupled to a digital camera (Hamamatsu ORCA-ER) and analyzed with AxioVision software.
A second group of mice was studied for 6 weeks, after which time the mice were placed into individual metabolic cages (Iffa Credo) for 24 hours, and their weight, water and food (sodium) intake, and urine output were documented. The urinary sodium concentration was measured with a COBAS-INTEGRA 400 Autoanalyzer (Roche Diagnostics) using an ion-sensitive electrode, and the result was adjusted for urinary output (μmol/day).

Ace2-KO mice. In these studies, male Ace2/ApoE-DKO and Ager/Ace2/ApoE-TKO mice were followed out to 18 weeks of age, a time point associated with increased plaque accumulation in DKO mice (19). In a second set of experiments, the mice, from 6 to 8 weeks of age, were further randomized to receive the mCherry-S391A-RAGE_{62-404} oligopeptide (10 μg/kg i.p. every other day), the mCherry-RAGE_{62-404} oligopeptide (10 μg/kg i.p. every other day), or mCherry alone (control; 10 μg/kg i.p. every other day).

Streptozotocin diabetes. Apoe-KO mice and Ager/ApoE-DKO were randomly allocated to receive streptozotocin (55 mg/kg, Sigma Chemical Co.) or buffer (sodium citrate buffer, pH 4.5) delivered i.p. in 5 consecutive daily doses, followed by mCherry-S391A-RAGE_{62-404} oligopeptide (10 μg/kg i.p. every other day), the mCherry-RAGE_{62-404} oligopeptide (10 μg/kg i.p. every other day), or mCherry alone (control; 10 μg/kg i.p. every other day) for 10 weeks. All diabetic mice achieved an HbA1c of more than 8% (median 11.2%) after 10 weeks, with no significant differences between treatment groups (data not shown).

Paigen diet. To exclude the nonspecific antiatherogenic actions of RAGE deletion, Apoe-KO and Ager/ApoE-DKO mice were fed a high-fat (Paigen) diet composed of 18.5% fat, 0.9% cholesterol, and 0.5% cholate for 6 weeks (20). This is considered a lipid-dependent model of atherosclerosis.

Quantitative real-time reverse transcription PCR. Gene expression of key proatherogenic mediators was estimated by quantitative real-time reverse transcription PCR (RT-PCR), performed using the TaqMan system (Thermo Fisher Scientific) on the basis of changes in the percentage of area stained red, as previously described (14, 43, 45).

In vitro studies. PMAECs. To assess the impact of RAGE deficiency on NF-κB-dependent proinflammatory signaling via the AT_1 receptor, PMAECs were isolated from the aortae of (WT) C57BL/6j mice and Ager-KO mice, as previously described (19), and cultured in DMEM/F-12 endothelial cell growth supplement (ECGS) media. Cultured PMAECs were then exposed to Ang II (1 μM for 2 hours), the RAGE ligand S100A8/A9 (2 μg/ml for 2 h), or vehicle, after which the cells were placed in TIRzol, the mRNA extracted, and the cDNA synthesized. Changes in gene expression of the RelA p65 NF-κB subunit gene, the Egr1 gene, or NF-κB-activated target genes were estimated by quantitative real-time RT-PCR, as detailed above. To confirm that the AT_1 receptor signaling cascade was functional in PMAECs from Ager-KO mice, the induction of IκBζ, a stable downstream metabolite of IκB, which accumulates in cells following Gq receptor activation, was measured 2 hours after exposure to Ang II or vehicle using the IP-One HTRF (homogeneous time-resolved fluorescence) assay (CisBio Bioassays).

Functional responsiveness to Ang II was further determined in PMAECs using a static adhesion assay. In brief, PMAECs were seeded at 50,000 cells per well into 6-well plates and allowed to grow to 70% confluency before treatment with 1 μM Ang II for 24 hours. Human THP-1 cells were stained using the CellVue Burgundy Fluorescent Cell Labeling Kit (LI-COR), before seeding them onto the endothelial cell monolayers at 3 × 10^4 viable cells per well, and incubated for 20 minutes at 37°C. The wells were washed with PBS before fixation of the cells with 4% formalin in PBS for 30 minutes. The adhesion of cells was quantitated using the ODYSSEY infrared imager (LI-COR).

In separate experiments, PMAECs from C57BL/6j mice were transfected with siRNAs targeting Ager (Ambion, assay ID: s62119; 2 nM), p65 (RelA) (Ambion, assay ID: s72857; 10 nM), PKCζ (Ambion, s71716; 10 nM), Diaph1 (sense 5’-UACAGAGGAAGGCUGUAAUUGACC, antisense 3’-GGCUUUCAUUCAUACGUCUCCUGUA; Invitrogen, Thermo Fisher Scientific; 10 nM), or scrambled control 1 (Ambion; 2 nM or 10 nM) using Lipofectamine RNAiMAX (Invitrogen, Thermo Fisher Scientific) according to the manufacturer’s instructions. The cells were allowed to recover in media containing 10% FBS for 16 hours prior to treatment with Ang II (1 μM for 2 hours), S100A8/A9 (2 μg/ml for 2 h), or vehicle. Further experiments were also conducted in the presence and absence of a 1-hour pretreatment with a neutralizing antibody against RAGE (R&D Systems, monoclonal mouse IgG2B buffer, and maintained at a physiological pH level by infusing carbonic gas (95% O_2, 5% CO_2) through the buffer at 37°C. The vessels (n = 6/group) were then treated with Ang II (1 μM for 4 h), after which whole blood labeled with Vybrant DiI (1:1000, Lonza) for the fluorescence labeling of leukocytes was then perfused through the aorta at 0.12 ml/minute. Images and videos of vessel wall–cell interactions were observed using a fluorescence microscope (Zeiss Discovery. V20), coupled to a digital camera (Hamamatsu ORCA-ER) and analyzed with AxioVison software.

In a second set of experiments, aortae isolated from Apoe-KO and Ager/ApoE-DKO mice were randomized to receive pretreatment with mCherry (control), mCherry-RAGE_{62-404} or mCherry-S391A-RAGE_{62-404} oligopeptide (all 8 μg/ml). Thirty minutes later, the aortae were exposed to Ang II (1 μM for 4 h), after which changes in the expression of key NF-κB-dependent proinflammatory genes were assessed using real-time RT-PCR.
clone 176902, catalog MAB11451; 1 µg/ml), soluble RAGE (1 µg/ml), or a pseudosubstrate for PKCζ (5 µM, Tocris). As a further control, the inhibitory effect of the AT1 receptor antagonist irbesartan (1 µM; 30 min before dosing with Ang II) was also explored.

To quantify hydrogen peroxide production induced by Ang II in PMAECs, a vessel chamber assay was used to measure peroxide production in real time in PMAECs from Ager-KO and C57BL/6j aortae, induced following exposure to Ang II. PMAECs were grown on collagen-coated coverslips and mounted into a sealed glass chamber when they were 80% confluent. The chambers were perfused with Krebs buffer with 2,7-dichlorofluorescein (DCFH) (Molecular Probes Europe BV) for 1 h, with fluorescence readings taken every 10 minutes to establish baseline values. The cells were then perfused with 1 µM Ang II for 60 minutes, and fluorescence readings were taken every 10 minutes afterward. Fluorescence readings were expressed as the change (Δ) in fluorescence AU from baseline.

Primary bone marrow–derived macrophages and splenocytes. To examine the impact of Ager deficiency on proinflammatory pathways implicated in the development of atherosclerosis, macrophages were isolated from bone marrow using the inflammasome method (19), and their proinflammatory responsiveness was then determined by changes in gene expression, as assessed by quantitative real-time RT-PCR.

Splenocytes were isolated from Ager-KO and C57BL/6j mice as previously described (47). Proinflammatory responsiveness following exposure to Ang II (1 µM for 2 h) was determined by changes in gene expression of Ccr2, Ifng, and the activation marker Mac1, as assessed by real-time RT-PCR (19, 47). Functional responsiveness was estimated by the induction of cellular adhesion between the isolated splenocytes and a monolayer of PMAECs.

Immortalized HMECs. To explore the potential to selectively modulate the transactivation of endogenous RAGE via transfection of human cells, HMECs (HMECI) (American Type Culture Collection [ATCC]) cultured in MCDB 131 medium (10% FCS with 10 mM glutamine, EGF, and hydrocortisone) were used. HMEC1 were transfected with each plasmid combination were then treated with a dose of 0.4 ng/ml for 60 minutes, and fluorescence readings were taken every 10 minutes afterward. Fluorescence readings were expressed as the change (Δ) in fluorescence AU from baseline.

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predicted cleavage site [residue 22] 1–25). The PCR product generated was cloned into pCIneo to generate pCIneo-SP-mCherry. For transmembrane RAGE constructs, portions of the AGER gene encoding residues 343–404 or 338–361 were amplified from pCIneo-AGER WT or the S391A mutant template and subcloned into the 3′ end of pCI-neo-SP-mCherry. For cytoplasmic construct expression, the coding sequence of mCherry was amplified from the vector pmCherry-C1 and subcloned into pCI-neo to generate pCI-neo-mCherry. Portions of the WT or S391A AGER gene encoding residues 362–404 and 370–404 were amplified from the vector pCI-neo-AGER and subcloned into pCI-neo-mCherry to generate cytoplasmic RAGE constructs. In each case, all sequences of the inserts were confirmed by DNA sequencing (Micromon, Monash University).

BRET. To explore the potential for a heteromeric AT, receptor-RAGE complex, we used BRET, an established technology for studying GPCR-protein proximity in live cells (48, 49). The AT, receptor, RAGE, and interacting proteins were linked to a bioluminescent donor enzyme, Rluc8 or Nluc, or an acceptor fluorophore, Venus or mCherry. Whether the donor or acceptor is fused to the N- or C-terminus is denoted by it being written before or after the protein of interest, respectively (e.g., Nluc-AT, vs. AT-Nluc). If in close proximity (<10 nm), energy resulting from the rapid oxidation of a cell-permeable coelenterazine or furimazine substrate by the donor can transfer to the acceptor, which in turn fluoresces at a longer characteristic wavelength. AT, and AT,-Rluc cDNA was provided by Walter Thomas (University of Queensland, St. Lucia, Brisbane, Australia). The Rluc coding region was replaced with Rluc8 cDNA from pcDNA3.1-Rluc8 provided by Andreas Loening and Sanjiv Gambhir (Stanford University, Stanford, California, USA) (50). Nluc-AT was provided by Promega, and AT, -Nluc was generated from AT,-Rluc8 by replacing Rluc8 with Nluc that was synthesized by GeneArt (Thermo Fisher Scientific).

For receptor-HIT (receptor-heteromer investigation technology) assays (Receptor-HIT, Dimerix Ltd.), RAGE-Rluc8 and β-arrestin2-Venus were expressed with untagged AT. In this system, ligand-induced BRET is indicative of recruitment of β-arrestin2-Venus proximal to RAGE-Rluc8, thereby indicating close proximity between RAGE and activated AT. The β-arrestin2-Venus cDNA construct was prepared previously from pCS2-Venus, provided by Atsushi Miyawaki (RIKEN Brain Science Institute, Wako City, Japan) (50). RAGE cDNA was amplified by PCR from pCI-neo-Ager to remove the stop codon and ligated into pcDNA3-Venus and pcDNA3-Rluc8. As a control for β-arrestin recruitment, the cDNA clone for CCR4 was obtained from the cDNA Resource Center (Bloomburg, Pennsylvania, USA). CCR4 cDNA was amplified by PCR to remove the stop codon and ligated into pcDNA3-Rluc8. Gu,-Nluc cDNA was generated by GeneArt (Thermo Fisher Scientific, Germany). Gr,-Venus was provided by Martina Kocan (The Florey Institute of Neuroscience and Mental Health, Parkville, Melbourne, Australia).

To explore changes in the subcellular localization of RAGE induced following ligand binding, BRET between RAGE and subcellular marker proteins was also explored. The plasma membrane marker Venus-Kras, as well as the subcellular compartment marker RabGTPases (Rabs) Venus-Rab5a (early endosomes), Venus-Rab7a (late endosomes/lysosomes), and Venus-Rab11a (recycling endosomes), were provided by Nevin Lambert (Georgia Regents University/Augusta University, Augusta, Georgia, USA). Rab1 (ER trafficking to the cis-Golgi), Rab4 (early endosome recycling), Rab6 (cis-Golgi), Rab8 (trans-Golgi net-
ing at 250 rpm. Cells were harvested by centrifugation (3500 g, 4°C, 20 min) and resuspended in ice-cold lysis buffer (50 mM Tris, pH 7.4, 300 mM NaCl, 10 mM imidazole, 5 mM β-mercaptoethanol). The resuspended cells were sonicated, and insoluble material was pelleted by centrifugation (13,000 g, 4°C, 30 min). The supernatant was filtered (0.2 µm) and applied onto a gravity flow column with a 1-ml Ni-NTA agarose resin bed volume (QIAGEN) preequilibrated with lysis buffer. After the supernatant was passed over the resin, the column was washed with 100 ml wash buffer (20 mM imidazole, 50 mM Tris, pH 7.4, 150 mM NaCl, 5 mM β-mercaptoethanol). The eluted fractions containing protein were pooled and quantified by NanoDrop spectrophotometry (Thermo Fisher Scientific) and BCA Protein Assay (Pierce, Thermo Fisher Scientific). For all in vivo applications, the peptides were buffer exchanged into sterile saline solution and the protein concentration recalculated. Cultured cells were treated with oligopeptides at a dose of 0.4 ng/ml, unless otherwise stated. Mice received respective peptides at a dose of 10 µg/kg i.p. every other day for 10 weeks.

Statistics. Continuous data are expressed as the mean ± SD. Differences in the mean among groups were compared using 2-way ANOVA. Pairwise multiple comparisons were made with Student-Newman-Keuls post hoc analysis to detect significant differences between groups, using Bonferroni’s correction for multiple comparisons. A P value of less than 0.05 was considered statistically significant.

Study approval. All experiments were approved by the animal ethics committee of the Alfred Medical Research Precinct and conducted in accordance with the Guide for the Care and Use of Laboratory Animals (NIH publication 85-23; National Academies Press, 1996).

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