D-Arg\(^0\)-Bradykinin-Arg-Arg, a Latent Vasoactive Bradykinin B\(_2\) Receptor Agonist Metabolically Activated by Carboxypeptidases

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We previously reported hypotensive and vasodilator effects from C-terminally extended bradykinin (BK) sequences that behave as B\(_2\) receptor (B\(_2\)R) agonists activated by vascular or plasma peptidases. D-Arg\(^0\)-BK-Arg-Arg (r-BK-RR) is a novel prodrug peptide hypothetically activated by two catalytic cycles of Arg-carboxypeptidases (CPs) to release the direct agonist D-Arg\(^0\)-BK. N-terminally extending the BK sequence with D-Arg\(^0\) in the latter peptide was meant to block the second kinin inactivation pathway in importance, aminopeptidase P. The affinity of r-BK and r-BK-RR for recombinant B\(_2\)R was assessed using a \[^3\]H]BK binding displacement assay. Their pharmacology was evaluated in human isolated umbilical vein, a contractile bioassay for the B\(_2\)R, in a morphological assay involving the endocytosis of B\(_2\)R-green fusion protein (GFP) and in anesthetized rats instrumented to record hemodynamic responses to bolus intravenous injection of both peptides. r-BK exhibited an affinity equal to that of BK for the rat B\(_2\)R, while r-BK-RR was 61-fold less potent. In the vein and the B\(_2\)R-GFP internalization assay, r-BK was a direct agonist unaffected by the blockade of angiotensin converting enzyme (ACE) with enalaprilat, or Arg-CPs with Plummer’s inhibitor. However, the in vitro effects of r-BK-RR were reduced by these inhibitors, more so by enalaprilat. In anesthetized rats, r-BK and r-BK-RR were equipotent hypotensive agents and their effects were inhibited by icatibant (a B\(_2\)R antagonist). The hypotensive effects of r-BK were potentiated by enalaprilat, but not influenced by the Arg-CPs inhibitor, which is consistent with a minor role of Arg-CPs in the metabolism of r-BK. However, in rats pretreated with both enalaprilat and Plummer’s inhibitor, the hypotensive responses and the duration of the hypotensive episode to r-BK were significantly potentiated. The hypotensive responses to r-BK-RR were not affected by enalaprilat, but were reduced by pre-treatment with the Arg-CPs inhibitor alone or combined with enalaprilat. Therefore, in vivo, Arg-CPs activity is dominant over ACE to regenerate the B\(_2\)R agonist r-BK from r-BK-RR, a prodrug activator of the B\(_2\)R. A B\(_2\)R agonist activated only at the level of the microcirculation by resident peptidases could be developed as an intravenously infused drug for ischemic diseases.

Keywords: bradykinin, angiotensin converting enzyme, arginine carboxypeptidases, B\(_2\) receptors, hypotension, D-Arg\(^0\)-BK-Arg-Arg
INTRODUCTION

Bradykinin (BK), a short-lived peptide, is the prototype of a family of peptides, the kinins, formed by the action of kallikreins on blood kininogens; the multi-molecular kallikrein-kinin system includes 2 G-protein coupled receptors termed B$_1$ and B$_2$ receptors [B$_1$R, B$_2$R; (Leeb-Lundberg et al., 2005)]. While the B$_1$R appears to have limited distribution and is generally absent in normal tissues, but is strongly induced under conditions of inflammation and tissue damage (Bouthillier et al., 1987; Marceau et al., 1988; Marceau and Regoli, 2004), the B$_2$R is constitutively expressed in a variety of tissues, and account for most of the vascular and metabolic actions of BK (Regoli and Barabe, 1980; Heitsch, 2003; Veeravalli and Akula, 2004; Marketou et al., 2010; Potier et al., 2013). BK exerts a variety of actions implicated in several physiological and pathological process, such as inflammatory reactions, due to its ability to cause vasodilation, hyperemia, vascular leakage, and pain sensation (Leeb-Lundberg et al., 2005; Moreau et al., 2005). However, apart from being a pro-inflammatory mediator, BK is also recognized as a regulator of blood pressure and several vascular and renal functions, mainly triggered by the synthesis and release of the vasorelaxant, anti-hypertrophic and anti-atherosclerotic endothelial mediators nitric oxide, prostaglandins, and tissue-type plasminogen activator (Brown et al., 2000; Pretorius et al., 2003; Griol-Charhbili et al., 2005; Leeb-Lundberg et al., 2005; Moreau et al., 2005; Kakoki et al., 2007). These cardioprotective effects of BK during hypertension and other clinical and experimental conditions, such as cardiac failure, ischemia, myocardial infarction, and pulmonary hypertension are believed to be B$_2$R-mediated (Heitsch, 2003; Veeravalli and Akula, 2004; Marketou et al., 2010; Sharma and Al-Banoon, 2012; Potier et al., 2013). Consequently, B$_2$R agonists may have important clinical value in the treatment and prevention of various cardiovascular disorders by mimicking the beneficial effects of BK.

Inspired by a “prodrug” strategy where a therapeutic B$_2$R agonist would be activated only at the level of the microcirculation by resident peptidases, we have previously explored the design of potential peptide drugs that are latent B$_2$R agonists activated by peptidases in isolated vascular systems (Gera et al., 2011; Charest-Morin et al., 2014) and in vivo (Jean et al., 2016). One of the most interesting, BK-Arg, massively lost affinity for recombinant B$_2$Rs but could regenerate active BK after reaction with arginine-carboxypeptidases (Arg-CPs) present in vascular tissue and blood plasma (Charest-Morin et al., 2014; Jean et al., 2016). The model was supported by the inhibition of BK-Arg biological activities by Plummer’s inhibitor, a high affinity blocker of Arg-CPs that is an arginine analog (Plummer and Ryan, 1981). Other BK sequences C-terminally extended with 2 residues were tested as angiotensin converting enzyme (ACE) substrates (Charest-Morin et al., 2014), but the cleavage rule(s) that lead to BK regeneration were not clear in vivo, possibly involving multiple peptidases (Jean et al., 2016). On the other hand, B$_2$R agonists that are resistant to multiple inactivation pathways, such as the peptide B-9972 (Jean et al., 2016) or the amphibian BK homolog maximakinin (Charest-Morin et al., 2017), may not be desirable vasodilators due to their propensity to activate extravascular B$_2$Rs on afferent nerve terminals, epithelia and other tissues, and are not remarkably more longer acting than BK in vivo when administered as intravenous boluses. BK, itself, is highly susceptible to intravascular inactivation mainly by ACE (Cyr et al., 2001; Fryer et al., 2008).

We report here a second round of the development of BK prodrug/soft drug design based on a peptide that is also C-terminally extended. The basic assumption is that prolonged BK sequences massively lose affinity for the B$_2$R, and also regenerate the C-terminal sequence of BK upon cleavage. Novel aspects include the block of the second kinin inactivation pathway in importance, aminopeptidase P (Cyr et al., 2001; Fryer et al., 2008), by N-terminally extending the BK sequence with D-Arg$^5$-BK (r-BK) by 2 cycles of hydrolysis by Arg-CPs from D-Arg$^5$-BK-Arg-Arg (r-BK-RR) (Figure 1). Circulating carboxypeptidase N and membrane-bound carboxypeptidase M are Arg-CPs strategically located to limit the regeneration of r-BK in the vasculature.

Although the protective role of kinins in the circulation is increasingly recognized, there have been very few attempts to use BK or a derivative in cardiovascular therapeutics. Therefore, development of a new drug class, designed to exploit vascular and blood plasma peptidases to stimulate the most desirable effects of endothelial B$_2$Rs and where circulatory benefits are generated, might find application in intensive care situations where an intravenous line is available (unstable angina, myocardial infarction, perhaps decompensated congestive heart failure) and possibly, in more chronic ailments (e.g., pulmonary hypertension).

MATERIALS AND METHODS

Drugs

Bradykinin was purchased from Bachem (Torrance, CA, United States), the B$_2$R antagonist icatibant, from Phoenix Pharmaceuticals (Burlingame, CA, United States), enalaprilat dihydrate, from Kemprotect Ltd. (Maltby, Middlebrough, United Kingdom) and Plummer’s inhibitor (mercaptopemethyl-3-guanidinoethylthiopropanoic acid) from Calbiochem (La Jolla, CA, United States; sequence of BK-related peptides in Figure 1). B-9972 {D-Arg$^5$-[Hyp$^3$, Igl$^5$, Oic$^7$, Igl$^8$]}-BK, (Bawolak et al., 2007; Jean et al., 2016) was a gift from Dr. Lajos Gera (University of Colorado Denver).

Design of Novel Prodrugs

D-Arg$^5$-BK (r-BK) and D-Arg$^5$-BK-Arg-Arg (r-BK-RR) were custom synthesized by CanPeptide, Inc. (Pointe-Claire, QC, Canada) via standard solid-phase methodology and provided as ≥98.3% pure reagents (mass spectroscopy and HPLC analyses). The first peptide, r-BK, putatively is an N-terminally protected direct agonist of the BK B$_2$R by virtue of its intact C-terminal BK sequence (Figure 1). The other C-terminally prolonged peptide theoretically retains little affinity for the B$_2$R, which has been experimentally verified. r-BK-RR was designed as a potential
FIGURE 1 | The C-terminally extended r-BK-RR sequence as potential “prodrug” agonist of the B\(_2\)R activated by peptidases. r-BK is itself potentially degraded by several peptidases that terminate its signaling at B\(_2\)Rs, but not by the major kininase aminopeptidase P. ACE, angiotensin converting enzyme; APP, aminopeptidase P; Arg-CPs, arginine carboxypeptidases. Marker 1: two cycles of Arg-CPs may regenerate r-BK from r-BK-RR. Marker 2: alternatively, ACE acting as a carboxydipeptidase may directly activate r-BK-RR. Marker 3: in turn, Arg-CPs and ACE will limit the half-life of r-BK by attacking his unprotected C-terminus. Marker 4: r-BK is postulated to be the only efficient high affinity B\(_2\)R agonist in the system. Marker 5: effects mediated at the B\(_2\)R are abated by the BK-sequence related B\(_2\)R antagonist icatibant. Modified from Jean et al. (2016).

prodrug needing 2 cycles of reaction with Arg-CPs to release r-BK.

\([\text{\textsuperscript{3}}\text{H}]\text{BK}\) Binding Competition Assays

Affinity for the B\(_2\)R was evaluated using a radioligand binding competition assay performed at 0°C in the presence of peptidase inhibitors that included captopril and PMSF (Charest-Morin et al., 2014). Briefly, the binding of 3 nM \([\text{\textsuperscript{3}}\text{H}]\text{bradykinin}\) (Perkin Elmer Life Sciences; 90 Ci/mmol) to adherent intact Human Embryonic Kidney (HEK 293) cells stably expressing the myc-tagged rat B\(_2\)R construction (Charest-Morin et al., 2017) was applied to construct binding competition curves for a series of unlabeled peptides.

Competition of an ACE Substrate by Synthetic Kinins

We applied a previously described enzymatic assay based on the internally quenched fluorogenic substrate Abz-Phe-Arg-Lys(Dnp)-Pro-OH, obtained from Bachem (Torrance, CA, United States), knowing that it has an approximately equal low micromolar affinity for the two separate catalytic sites of ACE (Araujo et al., 2000), to determine the affinity of the novel extended BK sequences for this peptidase. The source of enzyme was whole HEK 293a cells grown in 24-well plates and transfected as described with a vector coding for the human ACE-mCherry fusion protein, an active enzyme (Charest-Morin and Marceau, 2016). The culture medium was removed from cell wells, which were rinsed twice with PBS. The fluorogenic substrate (final concentration 20 \(\mu\)M), and, optionally, a competitor (BK, r-BK, or r-BK-RR in variable concentrations, or the ACE inhibitor enalaprilat 1 \(\mu\)M) were added in 250 \(\mu\)l PBS in each well; the plates were then incubated for 15 min at 37°C. At this point, the well supernatants were transferred in Eppendorf tubes, centrifuged 30 s at 12500 rpm to remove debris and 100 \(\mu\)l of this fluid transferred in wells of 96-well plate (black background) to read the fluorescence (excitation 320 nm, emission 420 nm) using a TECAN Infinite \(\textsuperscript{\text{\textregistered}}\) 200 PRO microplate reader. Results were expressed as the percent of the fluorescence in wells containing ACE-mCherry and the substrate without competitors. Controls included mock-transfected cells and enalaprilat-treated cells.

Human Umbilical Vein Contractility Assay

The institutional research ethics board (CHU de Québec) approved the anonymous use of human umbilical cord segments obtained after elective cesarean section deliveries (file number: 2012-323). Informed written consent was obtained from mothers. Umbilical vein rings, used as a contractile bioassay for the BK B\(_2\)R, were prepared and suspended in organ baths and submitted to equilibration in Krebs’ solution as described (Marceau et al., 1994; Gera et al., 2011). The vascular preparation was used to assess the effect of the peptidase inhibitors (introduced 30 min before the agonist) on the apparent potency of r-BK and r-BK-RR. The full cumulative concentration-effect curves were recorded for each peptide; a large concentration of BK (9.4 \(\mu\)M) was added to record the maximal contractile effect mediated by the B\(_2\)Rs for low potency agonists. Tissues were used only once and discarded; controls curves were obtained from other vascular rings from the same vein.
Microscopy of B₂R-GFP

Epifluorescence of GFP-tagged rabbit B₂R (B₂R-GFP) was observed in HEK 293 cells that stably express this construction, in order to detect the effect of r-BK and of r-BK-RR on receptor endocytosis and recycling, a morphological response to receptor stimulation. This system was previously shown to exhibit BK-induced endosomal internalization of the fluorescent receptor, maximal 30 min after stimulation but with gradual recycling to the cell surface in 1–3 h (Bachvarov et al., 2001; Charest-Morin et al., 2013). This system, based on a serum-containing culture medium, contains ACE and Arg-carboxypeptidase activities (Bachvarov et al., 2001; Charest-Morin et al., 2014). r-BK or r-BK-RR were added to 35-mm Petri dishes containing the cells with optional pre-treatment with peptide inhibitors and the cells were observed as described after 30 min of incubation at 37°C (Charest-Morin et al., 2014). The cells were observed in epifluorescence microscopy at 1000× magnification and photographed using an Olympus BX51 microscope coupled to a CoolSnap HQ digital camera (filters for GFP and fluorescein: excitation 460–500 nm, emission 510–560 nm, objective lens 100× oil UPlanApo, Olympus).

In Vivo Hemodynamics in Anesthetized Rats

All surgical and experimental procedures were reviewed and approved by the Animal Care and Handling Committee of Laval University, in accordance with the Canadian Council on Animal Care. Experiments were performed on male Sprague-Dawley rats (300–375 g) purchased from Charles River Laboratories (St-Constant, QC, Canada). The rats were housed in a light-controlled (12:12-h light-dark cycle (lights on at 0600)) and temperature-regulated room (22 ± 1°C). Animals had free access to normal chow diet and tap water. They were allowed to acclimate to their environmental conditions for 1 week prior to being studied. At the end of the acclimation period, the rats were anesthetised with sodium pentobarbital (50 mg kg⁻¹, i.p., supplemented as required) and had one catheter implanted into the right jugular vein [for intravenous (i.v.) injection] and one into the left femoral artery [for direct and continuous measurement of blood pressure and heart rate as previously described (Jean et al., 2016)]. Experiments started at least 20 min following the end of surgery in anesthetised rats.

Baseline measurements of heart rate and phasic and mean arterial blood pressure were recorded over a period of 15 min in anesthetised rats. A dose response curve was then obtained by recording changes in blood pressure and heart rate elicited by i.v. injection of peptide vehicle followed by increasing doses (0.025–12.8 µg/kg) of r-BK or r-BK-RR. Peptides were dissolved in isotonic saline (0.9% NaCl) containing 0.1% BSA to prevent the adsorption of peptide to the glassware and plastic surfaces. All i.v. injections were given as 100 µl boluses which were washed in with a further 100 µl of saline (the dead space of the catheter). Only one peptide was tested per group of rats and each injection started with saline-BSA 0.1% followed by the lowest dose of peptide. The next dose was administered once all recorded cardiovascular parameters had returned to baseline after the previous injection (usually 2–10 min). At the end of the experiments each animal was euthanized with an overdose of sodium pentobarbital (240 mg/kg, i.v.).

The mechanism subserving the cardiovascular responses to random i.v. injections of r-BK and r-BK-RR at the dose of 1.6 µg/kg, was evaluated in rats pretreated with specific antagonists. In these experiments, the rats were separated in four groups depending on the antagonist tested, and the cardiovascular responses to the peptides agonists were compared with those elicited in the untreated control group of rats. In the first group of treated rats, the long acting and selective B₂R antagonist, icatibant (Hoe 140) (D-Arg-[Hyp³,Thi⁵,D-Tic⁷], Thi⁵, D-Tic⁷, D-Tic⁷,

TABLE 1 | Nanomolar IC₅₀ values and their 95% confidence limits (between parentheses) derived from the competition assay for [³H]BK binding to B₂R-Rs (Figure 2A).

| Peptide | [³H]BK binding competition, rat myc-B₂R |
|---------|---------------------------------------|
| BK      | 17.2 (13.6–21.6)                      |
| r-BK    | 13.3 (10.3–17.3)                      |
| r-BK-RR | 1048 (448–2455)                       |
bradykinin (Hock et al., 1991; Wirth et al., 1991; Rhaleb et al., 1992; Marceau et al., 1994) was intravenously administered as bolus (75 µg/kg, 0.1 ml) following a 10 min period of baseline measurements of heart rate and blood pressure. Fifteen minutes later, recording changes in blood pressure and heart rate elicited by i.v. injection of peptide vehicle (saline-BSA 0.1%) followed by random injections of r-BK and r-BK-RR, were made as described above. The next peptide was administered once all recorded cardiovascular parameters had returned to baseline after the previous injection. Further experiments were made in a second group of rats pretreated with the ACE inhibitor, enalaprilat. In these experiments, enalaprilat was intravenously administered as bolus (0.1 m/kg, 0.1 ml) 15 min before the i.v. injections of peptide vehicle and both customized peptides, as above. Further experiments were also carried out in rats pretreated with the Plummer’s inhibitor (mercaptomethyl-3-guanidinioethylthiopropanoic acid) a high affinity inhibitor of arginine carboxypeptidases that is an arginine analog (Plummer and Ryan, 1981). The inhibitor was intravenously administered as bolus (0.75 mg/kg, 0.1 ml) followed 15 min later by the i.v. injections of saline-BSA 0.1%, and then r-BK and r-BK-RR (1.6 µg/kg) in random order. A fourth group of rats was pretreated with a combined i.v. injection of enalaprilat (0.1 mg/kg) and the Plummer’s inhibitor (0.75 mg/kg) 15 min before the randomized i.v. injections of both customized peptides, as above. The doses of different inhibitors were based on preliminary experiments and from previous studies performed by us and others (Ishida et al., 1989; Wirth et al., 1991; Muto et al., 2003; Jean et al., 2016). At the end of the experiments the rats were euthanized with an overdose of sodium pentobarbital (240 mg/kg, i.v.).

Data Analysis
Results are presented as means ± SEM. Radioligand binding data were fitted by non-linear regression to a one-site competition equation using a least-square method (Prism 5.0, GraphPad Software Inc., San Diego, CA, United States) and IC$_{50}$ values calculated from this procedure. The same computer program was used to draw concentration-effect curves generated with the contractility assay (least square fitting of sigmoidal dose-response equation with variable slope) and to derive contractile EC$_{50}$ values. Data describing the dose-response relationship of novel peptides vs. hypotension and heart rate changes, the baseline values of heart rate and mean arterial blood pressure and the effect of drugs on the hemodynamic responses to r-B K and r-BK-RR were assessed by using one-way analysis of variance (ANOVA) followed by the Dunnett’s test (repeated comparison with a common control; Prism 5.0 software).

RESULTS

Affinity of Peptides of the r-BK Series for the Rat B$_2$R
A [³H]BK binding competition assay performed at water-ice temperature in the presence of protease/peptidase inhibitors was exploited to determine the true receptor affinity of r-BK and r-BK-RR. Thus, r-BK was found to exhibit an affinity practically equal to that of BK for the rat myc-B$_2$R construction (Figure 2A; IC$_{50}$ values and their 95% confidence limits in Table 1). As expected, the C-terminal prolongation of r-BK decreased the receptor affinity, 61-fold for r-BK-RR (Figure 2A and Table 1). Thus, any significant activity of the C-terminally prolonged analog in

| Inhibitor Peptide        | r-BK EC$_{50}$ (µM) | r-BK-RR EC$_{50}$ (µM) |
|--------------------------|---------------------|------------------------|
| Control                  | 62.4 (49.7–78.4)    | 431 (348–534)          |
| Plummer’s inhibitor 1 µM| 68.4 (52.0–90.1)    | 850 (683–1057)         |
| Enalaprilat 1 µM         | 61.6 (41.9–90.5)    | 1506 (1129–2009)       |
| Enalaprilat 1 µM + Plummer’s inhibitor 1 µM | 45.1 (32.8–62.0) | 1760 (1377–2250) |

TABLE 2 | Nanomolar EC$_{50}$ values and their 95% confidence limits (between parentheses) derived from the human umbilical vein contractility assays (Figure 3).
a B2R-mediated bioassay of BK must be dependent of the regeneration of the direct agonist r-BK by precise cleavage rules.

**Affinity of Peptides of the r-BK Series for the Recombinant ACE**

If r-BK or r-BK-RR were ACE substrates, it would be predicted that they would compete for the hydrolysis of another ACE substrate. Thus, we used the fluorogenic ACE substrate Abz-Phe-Arg-Lys(Dnp)-Pro-OH and recombinant ACE-mCherry to determine, by competition, the affinity of r-BK and r-BK-RR over that of BK for ACE. As shown in Figure 2B, BK and r-BK competed with a similar affinity for the recombinant enzyme in a concentration-dependent manner, exhibiting nearly complete inhibition at a concentration equal to that of the substrate (20 µM). Unexpectedly, r-BK-RR was also active in this assay, only slightly less potent than the two other peptides, indicating...

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**FIGURE 4**

Epifluorescence microscopy studies of live HEK 293 cells stably expressing B2R-GFP. (A) Cells were stimulated for 30 min with r-BK or r-BK-RR (100 nM) with optional pretreatment with a peptidase inhibitor, as indicated. Control cells generally exhibit sharply defined plasma membrane-associated green fluorescence. Two different fields are shown for r-BK-RR. (B) Lack of persistence of the translocation of fluorescent receptor from the plasma membrane to endosomal structures in cells stimulated with BK or r-BK for 30 min or 3 h, as indicated. B-9972 is an inactivation-resistant agonist with a persistent morphologic response. Original magnification 1000×.
that r-BK-RR may be an ACE substrate. Controls made with cells without ACE (mock-transfected with mCherry) catalyzed the reaction to a small extent that may represent a contamination of cells by ACE from the serum-containing culture medium. ACE-expressing cells treated with enalaprilat did not catalyze the reaction.

**Pharmacological Profile of Peptides of the r-BK Series in an Isolated Vascular Preparation**

The human isolated umbilical vein is a contractile bioassay for the endogenous B₂R, and was exploited to further study the pharmacology of r-BK and R-BK-RR. Cumulative concentration-effect curves were constructed for r-BK and its homolog r-BK-RR based on the contractility of this preparation (Figures 3A,B; IC₅₀ values and their 95% confidence limits reported in Table 2). Specific peptidase inhibitors were used in separate tissues, as indicated. r-BK was a potent contractile agent essentially unaffected by the blockade of ACE with enalaprilat, and/or Arg-CPs with Plummer’s inhibitors. On a molar basis, r-BK-RR was about sevenfold less potent than r-BK. This peptide was designed to regenerate r-BK after 2 cycles of reaction with Arg-CPs (Figure 1), but the corresponding inhibitor, Plummer’s inhibitor, only slightly decreased the apparent potency of r-BK-RR (Figure 3B and Table 2). Unexpectedly, the ACE inhibitor enalaprilat was more effective to shift the concentration-effect curve of r-BK-RR to the right by a factor 3.5; the likely explanation is the removal of the Arg-Arg extension in a single step by this carboxydipeptidase (Figure 1). The combination Plummer’s inhibitor + enalaprilat was not significantly more effective than enalaprilat alone (Figure 3), and the residual effect of r-BK-RR in the presence of ACE blockade may approach the direct micromolar affinity of the peptide for B₂R, as suggested by its potency in the [³H]BK binding competition assay (Figure 2A). Thus, these results support a metabolic activation by ACE of the latent B₂R agonist, r-BK-RR, in this preparation.

**Effects of r-BK and r-BK-RR on B₂R-GFP Cycling**

The stable transfecant HEK 293 cell line expressing B₂R-GFP at the level of the plasma membrane was applied to detect the effect of r-BK and r-BK-RR on receptor endocytosis and recycling, a morphological response to receptor stimulation. Thus, after 30-min stimulation period with r-BK (100 nM; Figure 4A) in the serum-containing culture medium at 37°C, the fluorescent receptor was internalized. The cell morphology in green epifluorescence was that of disrupted plasma membrane continuity with abundant cytosolic inclusions close to the plasma membrane and discrete intracellular structures. This assay was also applied to r-BK-RR, which exerts, at 100 nM, little competition on the binding of [³H]BK to B₂R: (Figure 2A). After 30 min of treatment with r-BK-RR, the translocation of B₂R-GFP-associated fluorescence from the plasma membrane to endosomes was similar to that seen with r-BK, observed in the vast majority of cells (Figure 4A).

While HEK 293 cells do not express ACE unless transfected with the corresponding expression vector (Morissette et al., 2008), evidence was provided that their serum-supplemented culture medium contains soluble ACE (Bachvarov et al., 2001), as well as soluble Arg-CPs activity (Charest-Morin et al., 2014). The specific inhibitors of these peptidases, enalaprilat and Plummer’s inhibitor, respectively, were applied separately or together, 15-min before an additional 30-min treatment with either peptide in additional experiments reported in Figure 4. While none of these inhibitors influenced the >90% proportion of cells that exhibited r-BK-induced internalization of B₂R-GFP, treatment with enalaprilat alone or combined with Plummer’s inhibitor was found to suppress the effect of r-BK-RR (Figure 4A), consistent the metabolic activation of r-BK-RR by ACE. Treatment with Plummer’s inhibitor alone had no effect on r-BK-induced internalization of the fluorescent receptor, but had only a partial effect on r-BK-RR-induced translocation of plasma membrane fluorescence to endosomal structures (Figure 4A).
Bradykinin induces the rapid endocytosis of the B₂R-GFP construction toward the endosomal pathway in HEK 293 cells, but this is followed by extensive recycling to the plasma membrane following 3 h of incubation (Bachvarov et al., 2001). This is replicated in experiments reported in Figure 4B, where BK was compared to r-BK and the inactivation-resistant B₂R agonist B-9972. The effect of r-BK was reversible as a function of incubation time, as that of BK, while B-9972 elicits a prolonged agonist B-9972. The effect of r-BK was reversible as a function of incubation time, as that of BK, while B-9972 elicits a prolonged agonist B-9972. The effect of r-BK was reversible as a function of incubation time, as that of BK, while B-9972 elicits a prolonged agonist B-9972. The effect of r-BK was reversible as a function of incubation time, as that of BK, while B-9972 elicits a prolonged agonist B-9972. The effect of r-BK was reversible as a function of incubation time, as that of BK, while B-9972 elicits a prolonged agonist B-9972. The effect of r-BK was reversible as a function of incubation time, as that of BK, while B-9972 elicits a prolonged agonist B-9972. The effect of r-BK was reversible as a function of incubation time, as that of BK, while B-9972 elicits a prolonged agonist B-9972. The effect of r-BK was reversible as a function of incubation time, as that of BK, while B-9972 elicits a prolonged agonist B-9972. 

**In Vivo Hemodynamic Responses to r-BK and Its C-Terminally Extended Homolog, r-BK-RR**

We previously described the brief hypotensive responses associated with tachycardia in response to the i.v. injection of increasing doses of BK in anesthetized rats and the strong potentiation of the responses following pharmacologic ACE blockade (Jean et al., 2016). Using the same methods, we found that intravenous injection of increasing doses of r-BK, as bolus in anesthetized rats, produces dose-dependent hypotensive and tachycardiac responses; comparison with the effects of BK assayed under the same experimental conditions shows that r-BK is 2.5–3-fold more potent than BK as an hypotensive agent in anesthetized rats (Jean et al., 2016). In the present experiments, r-BK and r-BK-RR are virtually equipotent hypotensive agents (typical blood pressure tracings, Figure 5; dose-response curves, Figure 6A). These hypotensive responses are significantly greater than that of the BSA-saline vehicle for peptide doses greater or equal to 400 ng/kg. Further, the two peptides exhibit similar rapid action and cessation of action (half-time for the recovery of the hypotensive episodes; Figure 6C). After a brief bradycardiac episode that follows by the bolus injection (an injection artifact), the major effect of r-BK and r-BK-RR on the heart rate usually was a brief tachycardic response, statistically significant only at the highest tested doses (Figures 5, 6B). This suggests a particularly effective conversion of r-BK-RR into the active peptide r-BK in vivo.

**Effects of Icatibant and Peptidase Inhibitors on Cardiovascular Responses to r-BK and r-BK-RR**

Baseline values for mean arterial blood pressure and heart rate measured in the untreated control group or 15 min after i.v. pretreatment with icatibant or the peptidase inhibitors are shown in Table 3. While no significant changes in basal values of MAP were noted between the treated groups and the untreated control group, slight but significant increases in basal heart rate were noted between the treated groups and the control group.

As shown in Figure 7 and consistent with its direct agonist action on B₂R, the hypotensive responses to r-BK (1.6 µg/kg) were significantly inhibited by pretreatment with icatibant (P < 0.001, Dunnett’s test), while the heart rate response and the duration of the residual hypotensive episode were comparable to those seen in the untreated group. In rats pretreated with enalaprilat, the hypotensive effect of r-BK was significantly potentiated (P < 0.05, Dunnett’s test) as was the tachycardic

| Pre-treatment            | Mean arterial blood pressure (mmHg) | Heart rate (bpm) | n  |
|-------------------------|------------------------------------|------------------|----|
| Control                 | 93.2 ± 3.3                         | 352 ± 10         | 13 |
| Icatibant               | 93.7 ± 3.3                         | 396 ± 10*        | 10 |
| Enalaprilat             | 89.3 ± 4.1                         | 412 ± 11**       | 9  |
| Plummer’s inhibitor     | 90.3 ± 3.6                         | 407 ± 10**       | 8  |
| Enalaprilat + Plummer’s inhibitor | 87.8 ± 3.9 | 406 ± 13**       | 10 |
| ANOVA                   | N.S.                               | P < 0.001        |    |

*P < 0.05, **P < 0.01, Dunnett’s test vs. control.
response ($P < 0.05$, Dunnett’s test). However, the duration of the hypotensive episode was not different from that seen in untreated rats (Figure 7). Pretreatment with the Plummer’s inhibitor alone had no effect on the hypotensive and tachycardic responses to r-BK, which is consistent with a minor role of Arg-CPs in the metabolism of BK (Cyr et al., 2001; Fryer et al., 2008). In rats pretreated with both enalaprilat and the Plummer’s inhibitor, the hypotensive effect of r-BK, as well as the duration of the hypotensive episode were both significantly potentiated ($P < 0.01$, Dunnett’s test), while the heart rate response was not different from that seen in untreated rats (Figures 7, 8A).

Therefore, in the presence of enalaprilat, which contributes to amplify the hypotensive response to r-BK, the Plummer’s inhibitor reveals the involvement of Arg-CPs in cessation of the hypotensive episode.

The C-terminally extended analog r-BK-RR is predicted to be an indirect activator of the B$_2$R, via its conversion to r-BK (Figure 1). Consistently, pretreatment with icatibant was found to strongly reduce the hypotensive response to intravenously administered (1.6 $\mu$g/kg) r-BK-RR ($P < 0.01$, Dunnett’s test), while the heart response or the duration of the residual hypotensive episode were not different from those of controls (Figure 7). r-BK regeneration from r-BK-RR should involve Arg-CPs, but perhaps also the carboxypeptidase ACE, based on in vitro results (Figures 1–4). Interestingly, pretreatment with enalaprilat had no significant effect on the hypotensive response to r-BK-RR, but it was found to significantly increase the duration of the hypotensive episode ($P < 0.001$, Dunnett’s test) (Figures 7, 8B). Pretreatment with the Plummer’s inhibitor alone or combined with enalaprilat was found to significantly reduce the hypotensive response to r-BK-RR, when compared to the untreated group ($P < 0.05$, Dunnett’s test) (Figure 7), while the heart rate response and the duration of the residual hypotensive episode were not different from those seen in untreated rats. Altogether, in vivo results indicate that Arg-CPs activity is dominant over ACE to regenerate a B$_2$R agonist from r-BK-RR, but ACE inhibition potentiates the reaction product r-BK.

**DISCUSSION**

In the present study, we evaluated a “prodrug” peptide extended around the BK sequence, as a potential B$_2$R agonist activated by peptidases present in the microcirculation. We explored the possibility of a controlled release of the direct agonist r-BK by two cycles of hydrolysis by Arg-CPs from r-BK-RR. Briefly, r-BK exhibited an affinity practically equal to that of BK for the rat B$_2$R construction, while r-BK-RR was 61-fold less potent. In the vein and the B$_2$R-GFP internalization assay, r-BK behaved as a direct agonist unaffected by the blockade of ACE with enalaprilat, or Arg-CPs with Plummer’s inhibitor. However, the in vitro effects of r-BK-RR were reduced by these inhibitors, more so by enalaprilat. In anesthetized rats, r-BK and r-BK-RR caused equipotent hypotensive effects that were inhibited by icatibant (a B$_2$R antagonist). The hypotensive effects of r-BK were potentiated by enalaprilat, but not influenced by the Arg-CPs inhibitor, which is consistent with a minor role of Arg-CPs in the metabolism of r-BK. However, in rats pretreated with both enalaprilat and Plummer’s inhibitor, the hypotensive responses and the duration of the hypotensive episode to r-BK were significantly potentiated. The hypotensive responses to r-BK-RR were not affected by enalaprilat, but were reduced by pre-treatment with the Arg-CP inhibitor alone or combined with enalaprilat, indicating that, in vivo, Arg-CPs activity is likely dominant over ACE to regenerate the B$_2$R agonist, r-BK, from r-BK-RR, a prodrug activator of the B$_2$R.
The development of protease-activated “soft pro-drugs” might be an original and highly useful approach to deliver drugs in areas where protease expression is higher than in normal. This might contribute to reduce off-target side effects and to prolong beneficial effects by controlled and progressive regeneration of active drug. BK, a high-affinity, direct B₂R agonist that is also an effective ACE substrate, plays an important role in the regulation of blood pressure, renal and cardiac functions, through its effects on nitric oxide, prostaglandins and tissue plasminogen (t-PA), which effects mainly arise from endothelial cell B₂Rs activation (Brown et al., 2000; Veeravalli and Akula, 2004; Kakoki et al., 2007; Marketou et al., 2010; Potier et al., 2013). Consequently, B₂R agonists may have important clinical value in the treatment and prevention of various cardiovascular disorders such as hypertension, ischaemic heart disease and other. Thus, using a “prodrug” strategy design and exploiting the distribution of ectopeptidases expressed in the vasculature and blood plasma, we recently provided pharmacological evidence, that interesting vascular effects can be extracted from C-terminal extended BK sequences that behave as latent peptidase-activated B₂R agonists (Charest-Morin et al., 2014; Jean et al., 2016). In continuity with these studies, we have designed and evaluated a new “prodrug” peptide extended around the BK sequence, as a potential peptidase-activated B₂R agonist, in vitro and in anesthetized rats. Novel aspects of the present study include the block of the second kinin inactivation pathway in importance, aminopeptidase P (Cyr et al., 2001; Fryer et al., 2008), by N-terminally extending the BK sequence with D-Arg⁸. This was done to explore the possibility of a controlled and progressive release of the direct agonist r-BK by two cycles of hydrolysis by Arg-CPs from r-BK-RR.

**Pharmacological Profile of BK Extended Peptides**

We found that r-BK shares similar affinity than BK for the rat B₂R, while the C-terminal prolongation of r-BK results in a severe decrease of receptor affinity, 61-fold for r-BK-RR. Therefore, any significant vasoactive effect of the latter BK analog must derive from its cleavage, leading to a more conventional receptor agonist. r-BK-RR, was designed to regenerate active r-BK after two cycles of reaction with Arg-CPs present in vascular tissue and blood plasma. Interestingly, r-BK-RR was found to be a contractile agonist of the human umbilical vein less potent than r-BK but more potent than anticipated from the radioligand binding competition assay. Moreover, the contractile potency of r-BK-RR was only slightly decreased by Plummer’s inhibitor, whereas the ACE inhibitor was more effective to shift the concentration-effect curve of r-BK-RR to the right, suggesting the removal of the C-terminal dipeptide in a single step by this carboxypeptidase. This finding is in line with the enzymatic assay applied to recombinant ACE indicating that r-BK-RR may be an ACE substrate. It is also consistent with the demonstration that r-BK-RR-induced internalization of B₂R-GFP is selectively suppressed by enalaprilat in HEK 293 cells, a system where ACE is supplied by serum-containing culture medium (Bachvarov et al., 2001). The residual contractile effect of r-BK-RR in the presence of ACE blockade may largely depend on the direct micromolar affinity of the peptide for B₂R, as
suggested by its potency in the [3H]BK binding competition assay.

Although ACE presence in the umbilical vein is functionally revealed by the metabolic activation of prodrug peptides that regenerate BK, enalaprilat failed to potentiate r-BK in the human umbilical vein contractility (Figure 3A). The lack of effect of ACE inhibitors on the apparent potency of BK has been previously reported in this preparation (Marceau et al., 1994; Gobell et al., 1996; Bawolak et al., 2007). Previous immunohistochemistry and immunofluorescence studies showed that the umbilical vein possesses a relatively thick media (>20 layers of smooth muscle cells), while immunoreactive ACE is limited to the single endothelial cell layer of the luminal surface of the vein (Koumbadinga et al., 2010; Gera et al., 2013). Therefore, considering the low endothelium/muscle ratio found in human umbilical vein preparation, it is suggested that ACE is not abundant enough in the tissue to impair the equilibrium between BK concentrations in the bathing fluid and those at the vicinity of B2R that mediate the contraction of venous muscle cells (Marceau et al., 2010).

The B2R-GFP internalization assay also suggests that ACE is dominant over Arg-CPs to activate the pro-drug r-BK-RR, based on the effect of peptidase inhibitors (Figure 4A). Both in this cell culture setting and in the isolated umbilical vein model, the absolute and relative abundance of ACE and Arg-CPs may not fully represent their in vivo role at the level of resistance arterioles. Further, the translocation of the fluorescent receptor from the plasma membrane to endosomal structures is not more persistent (3 h of stimulation as compared to 30 min) in cells stimulated with r-BK than in those exposed to BK (Figure 4B). r-BK, unlike B-9972, appears to be a "soft drug" perhaps less prone to exert extravascular effects if infused in the circulation.

**Cardiovascular Effects of BK Extended Peptides in Rats**

The most salient finding we made in anesthetized rats was the demonstration that r-BK and r-BK-RR are equipotent in causing rapid and transient and dose-related hypotensive responses and tachycardia following their systemic administration. While BK is a negative inotropic stimulus in the isolated rat heart (Rastaldo et al., 2001), massive vasodilation determines a reflex tachycardic response to BK injection in vivo (Jean et al., 2016). As expected, the hypotensive response to systemic administration of r-BK was greatly enhanced by pretreatment with enalaprilat (as was the tachycardia), extensively inhibited in the presence of a B2R antagonist and remained unchanged in the presence of the Plummer’s inhibitor (Figure 7), which is consistent with a minor role of Arg-CPs in the metabolism of r-BK. These results underscore the important role played by ACE, as the main r-BK-inactivating peptidase in the extracellular space. However, in rats pretreated with both enalaprilat and Plummer’s inhibitor, the hypotensive responses and the duration of the hypotensive episode to r-BK were significantly potentiated, revealing the involvement of Arg-CPs in cessation of the hypotensive episode when the metabolism of r-BK by ACE is inhibited.

Pharmacological evidence of B2R-mediated hypotensive response to r-BK-RR (shown to have very little direct affinity for B2R) was obtained as icatibant significantly inhibit the hypotensive effect of the peptide. However, in contrast to what we previously found in the human umbilical vein contractile assay, and despite the enzymatic assay indicating that r-BK-RR may be an ACE substrate, the cleavage rule leading to r-BK regeneration in vivo does not appear to follow a single catalytic step mediated by ACE. Indeed, in anesthetized rats, pretreatment with enalaprilat had no effect on the hypotensive response to r-BK-RR, but it was found to increase the duration of the hypotensive episode. Pretreatment with the Arg-CPs inhibitor alone or combined with enalaprilat was found to strongly reduce the hypotensive response to r-BK-RR. Altogether, the in vivo results indicate that Arg-CPs activity is dominant over ACE to regenerate a B2R agonist from r-BK-RR, but ACE inhibition potentiates the reaction product r-BK. Thus, the present findings indicate that the C-terminally extended analog r-BK-RR is an indirect activator of the B2R, via its conversion to r-BK, and support the basic postulated cleavage rule leading to r-BK regeneration following two catalytic step mediated by Arg-CPs (Figure 1).

r-BK-RR is presumably a pro-drug B2R agonist peptide activated by Arg-CPs expressed in the vicinity of vascular endothelial cells and a soft drug, because the active reaction product r-BK is itself cleared by ACE and in the endosomes. The development of such a drug, selective for the vascular system and stimulating the most desirable endothelial B2R effects, might prove to be very useful in specific intensive care situations where the stimulation of vascular B2Rs has been proposed to have therapeutic value, such as myocardial infarction and ischemic stroke.

**AUTHOR CONTRIBUTIONS**

HB and FM conceived and designed the experiments, analyzed the data, wrote the paper, prepared the figures and/or tables, and reviewed the drafts of the paper. XC-M performed and designed some of the experiments and reviewed the drafts of the paper. All authors approved the version to be published.

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**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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