PulmoBind, an Adrenomedullin-Based Molecular Lung Imaging Tool

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Previous studies showed that adrenomedullin (AM) could be a promising agent for molecular imaging of the pulmonary circulation, with abundant specific binding sites at the pulmonary vascular endothelium. The purpose of this work was to design an AM-based compound that encompasses the desired imaging properties without posing safety issues for clinical applications. Methods: AM analogs were synthesized through solid-phase peptide synthesis. They were evaluated for 99mTc labeling efficiency and in vivo lung uptake. Biodistribution and hemodynamic characteristics of the lead compound were determined in anesthetized dogs as well as by a dosimetric analysis. Lung perfusion was evaluated in the monocrotaline model of pulmonary arterial hypertension in rats. Results: A cyclic AM (residues 22–52) analog encompassing a polyethylene glycol spacer and a tetrapeptide chelating moiety was found to possess the desired characteristics, with 90.7% ± 0.3% (mean ± SD) labeling efficiency, 40% lung uptake at 10 min after injection, and a favorable safety profile. Lung uptake of the 99mTc-labeled compound was markedly reduced in rats with pulmonary arterial hypertension. Conclusion: This lead compound could be a suitable clinical imaging agent for the molecular diagnosis of disorders of the pulmonary circulation.

Key Words: lung; molecular imaging; adrenomedullin; nuclear medicine; pulmonary hypertension

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Pulmonary embolism is a potentially lethal lung perfusion defect that commonly develops in patients with deep vein thrombosis (1). Pulmonary hypertension, a disorder with various etiologies, leads—in its most severe forms—to right ventricular failure and death (2). Diagnosis of these pathologies is challenging because patients often have a nonspecific symptom, shortness of breath. At present, nuclear imaging of the pulmonary circulation can be achieved through the use of 99mTc-albumin macroaggregates, particles that are trapped in the microvasculature because of their size (10–150 μm) in proportion to the regional blood flow (3). However, this technique lacks the ability to detect functional perfusion defects, and its ability to detect small-blood-vessel blockade is limited. Moreover, this tracer blocks pulmonary blood vessels in subjects with an already compromised pulmonary circulation and increases the potential risk for infections because it is prepared from human albumin.

We recently showed that adrenomedullin (AM) is a promising molecular lung imaging tool for the diagnosis of pathologies such as pulmonary embolism and pulmonary hypertension (4,5). AM, a peptide from the calcitonin family, allows specific lung imaging through binding with its specific receptor, AM1 (CRLR/RAMP2), which is expressed at a high density in pulmonary vessels (6,7). Enhanced imaging efficiency can be associated with cell internalization after receptor activation by its agonist (8,9), but this process is also correlated with inherent biologic activity. Conversely, antagonists do not evoke cellular activation in receptor binding but are able to bind with uniform affinity to the total pool of receptors (10,11).

AM, a 52-amino-acid multifunctional regulatory peptide, is widely distributed and acts mainly in the cardiovascular system, for which it is classically described as a vasodilator (12). Previous structure–activity relationship studies demonstrated that truncating the N-terminal stretch while maintaining the cyclic structure formed by a disulfide bridge between residues 16 and 21 generated agonists, such as AM with residues 13–52 [AM(13–52)] and AM(16–52), with affinities similar to that of the full-length peptide (13). Further truncation at the N terminus to remove the ring structure yielded the only fully characterized human AM antagonist, that is, AM (22–52); in contrast, removal of the C-terminal amino acid greatly diminished binding affinity and peptide activity (13,14). Interestingly, analogs composed only of the ring structure, that is, AM (15–22) and AM (16–21), increased systemic arterial blood pressure (15), but AM(1–25) and AM(1–40) were unable to displace 125I-AM in a dog lung homogenate binding experiment (6). Moreover, linearization of the full-length peptide resulted in a loss of affinity (6,13). Finally, although AM is a long peptide, the incorporation of a radionuclide can affect its binding ability; therefore, AM-derived nuclear imaging tools should favor radiolabeling at the N terminus of the peptide (16).

AM(22–52) is an antagonist that has 10 times less affinity for its receptors than AM(13–52) in competition displacement experiments but that has higher selectivity for AM1 than for AM2 (CRLR/ RAMP3) (6,14). Thus, this peptide appeared to be a good starting point for generating new AM antagonists suitable for lung nuclear imaging. Such analogs would enable pulmonary microcirculation imaging without any side effects on blood pressure. Furthermore, and as pointed out for AM, such molecules not only would be...
valuable for detecting perfusion defects but also could allow the detection of early endothelium-related deficiencies, such as pulmonary hypertension.

In this article, we describe the design, synthesis, and characterization of PulmoBind, an AM-based analog for pulmonary circulation imaging. This new molecular lung imaging tool has good binding affinity, high $^{99m}$Tc labeling efficiency, and significant lung uptake even after 1 h without any effect on mean arterial pressure (MAP).

This molecule could make it possible to detect not only large-pulmonary-vessel blockade, such as a pulmonary embolism, but also anatomic and, potentially, functional pulmonary microcircular disorders caused by various forms of pulmonary hypertension. Thus, PulmoBind could provide an efficient, safe, and noninvasive method for probing the status of the pulmonary circulation.

**MATERIALS AND METHODS**

All animal experimental procedures were performed in accordance with regulations and ethical guidelines of the Canadian Council for the Care of Laboratory Animals and received approval from the Animal Ethics and Research Committee of the Montreal Heart Institute.

**AM Derivative Synthesis and Purification**

AM-derived peptides were synthesized and purified as previously described (5). The Fmoc-$\alpha$-(polyethylene glycol)$_4$-COOH acid used as a spacer in the peptide synthesis was purchased from Quanta Bidesign Ltd. Disulfide bond formation to yield cyclic derivatives was performed by overnight air oxidation after dissolution of the linear peptide in an aqueous solution (pH 8.8) at a concentration of 0.1 mg/mL. Peptides were purified by reversed-phase high-performance liquid chromatography (HPLC). The amino acid sequences of the synthesized peptides are shown in Table 1, along with the purity of the final products and the molecular masses, as evaluated by matrix-assisted laser desorption ionization–time-of-flight mass spectrometry (Voyager DE; Applied Biosystems).

**$^{99m}$Tc Labeling**

Linear AM was radiolabeled through its cysteine residues as previously described (5). For AM analogs containing a chelating moiety, 2.29 nmol of lyophilized peptide were suspended in 20 mM acetate buffer (pH 5.5), to which 200 μL of 1 M NaH$_2$PO$_4$ (pH 4.5) was finally added. Radiochemical purity and labeling efficiency were assessed by instant thin-layer chromatography (ITLC) on silica gel–impregnated glass fiber filter paper with acetone for the dosage of free $^{99m}$Tc and 37.5% (v/v) butanol–7.5% (v/v) acetic acid–30% (v/v) pyridine in water to discriminate $^{99m}$Tc-colloids (bottom) from radiolabeled peptides (top).

For purifying $^{99m}$Tc-labeled AM derivatives, the radiolabeling reaction mixture was injected into a 1-mL (100-mg) C$_{18}$ SepSep Waters (Eke) cartridge. The cartridge was then washed with 3 mL of 1 mM hydrochloric acid and eluted with 3 mL of a 50% (v/v) solution of ethanol and water. Fractions of 0.5 mL were collected into sterile propylene tubes. Fractions with the highest counts were pooled, and 200 μL of 10× concentrated phosphate-buffered saline (pH 7.4) were added for stabilization.

**In Vivo Biodistribution and Lung Uptake**

Mongrel dogs (20–30 kg) were anesthetized by intravenous injection of sodium pentobarbital (50 mg/kg), intubated, and mechanically ventilated. Purified $^{99m}$Tc-labeled AM derivatives (185 MBq) were injected intravenously via the jugular vein through a 3-way stopcock 18-French catheter for lung uptake evaluation. In vivo whole-body biodistribution of radiolabeled peptides was evaluated by use of a dual-head γ-camera (Siemens) equipped with a low-energy parallel-hole collimator. Dynamic acquisitions were recorded for a 30-min period, and static whole-body scans were obtained at 30, 60, and 120 min after $^{99m}$Tc-labeled peptide injection at a scan speed of 10 cm/min. Dynamic and static acquisitions were evaluated with MATLAB version 7.01 image analysis tool software. Data correction was applied for radioactive decay, table adjustment (dorsal images only), geometric mean, and organ attenuation based on the transmission factor. Results were expressed as a percentage of the total radioactivity injected.

**Binding Assays**

The binding ability of PulmoBind was evaluated with dog lung homogenates prepared as previously described (6). In brief, lungs were diced, homogenized, and centrifuged at 49,000g for 20 min at 4°C. Pellets were washed twice, and nonhomogenized tissue was removed by gentle centrifugation (500g; 5 min). Aliquots of the supernatant...

#### Table 1

Human AM (hAM) and Peptide Analogs Synthesized

<table>
<thead>
<tr>
<th>Compound</th>
<th>Sequence</th>
<th>Purity (%)</th>
<th>Theoretic molecular mass</th>
<th>Observed molecular mass</th>
</tr>
</thead>
<tbody>
<tr>
<td>hAM(22–52)</td>
<td>Thr-Val-Gln-Lys-Leu-Ala-His-Gln-Ile-Tyr-Gln-Phe-Thr-Asp-Lys-Asp-Asp-Asn-Val-Ala-Pro-Arg-Ser-Lys-Ile-Ser-Pro-Gln-Gly-Tyr</td>
<td>95</td>
<td>3,576.03</td>
<td>3,578.58</td>
</tr>
<tr>
<td>1</td>
<td>Gly-Gly-$\alpha$-Ala-$\alpha$-PEG$_4$-$\alpha$hAM(22–52)</td>
<td>92</td>
<td>4,064.16</td>
<td>4,065.55</td>
</tr>
<tr>
<td>2 (PulmoBind)</td>
<td>Gly-Gly-$\alpha$-Ala-$\alpha$-Cys-$\alpha$-PEG$_4$-$\alpha$-Cys-$\alpha$hAM(22–52)</td>
<td>97</td>
<td>4,270.82</td>
<td>4,271.71</td>
</tr>
<tr>
<td>3</td>
<td>Gly-Gly-$\alpha$-Ala-$\alpha$-Cys-(Amc)$\alpha$-PEG$_4$-$\alpha$-Cys-(Amc)$\alpha$hAM(22–52)</td>
<td>96</td>
<td>4,412.82</td>
<td>4,410.34</td>
</tr>
</tbody>
</table>

hAM = human AM; PEG = polyethylene glycol; Amc = acetalmodimethyl.
ments were based on previously described protocols \(6,16\), and the specific activity was 81 MBq/mmol.

Competitive displacement experiments were carried out by incubating dog lung homogenates (100 μL; 0.24 mg) for 90 min at room temperature in binding buffer (50 mM Tris, 100 mM NaCl, 4 mM MgCl₂, and 0.1% [w/v] bovine serum albumin; pH 7.4) with various concentrations \(10^{-12} - 10^{-5}\) M of either unlabeled PulmoBind or AM in the presence of 1 nM ⁹⁹mTc-labeled PulmoBind. For saturation binding experiments, dog lung homogenates were incubated as described for competitive displacement experiments with increasing concentrations \((0.01-5\) nM) of ⁹⁹mTc-labeled PulmoBind in the absence (total binding) or presence of 1 μM unlabeled ligand. Incubations were stopped by rapid filtration through glass fiber filter papers presoaked in 0.3% polyethyleneimine with a Millipore 1225 sampling vacuum manifold. After 3 washings, the activity in the papers was counted with an LKB Wallac 1272 automatic γ-counter.

Hemodynamic Evaluation

Mongrel dogs (male and female) were used to study cardiovascular effects induced by cumulative intravenous injections of human AM (1–52), the agonist human AM (13–52), the antagonist human AM (22–52), and PulmoBind. The heart rate of anesthetized and ventilated animals was monitored with cutaneous electrocardiographic leads, and a catheter installed in the right femoral artery was used to monitor blood pressure. Pulmonary artery pressure was evaluated with a Swan–Ganz catheter. In brief, the catheter was introduced through the jugular vein. From this entry site, it was threaded by fluoroscopy through the right atrium of the heart, through the right ventricle, and subsequently into the pulmonary artery until a wedge pressure tracing was noted. Next, the Swan–Ganz catheter balloon was deflated; deflation was confirmed by the reappearance of pulmonary artery pressure contours.

Dosimetric Evaluation

The PulmoBind tracer was injected into 7 dogs to estimate the mean biodistribution on whole-body scintigraphic images. Before peptide injection, a whole-body transmission scan was obtained with the γ-camera and a ⁵⁷Co flood source. The injection syringe was measured before injection under the camera. Immediately after intravenous injection, dynamic acquisitions were recorded for a 30-min period, and static whole-body scans were obtained 35, 60, and 120 min later at a scan speed of 10 cm/min. The empty syringe was measured again to determine residual activity.

The biodistribution into organs with significant uptake was evaluated with MATLAB version 7.01 image analysis tool software. Dynamic and whole-body images were first corrected for radioactive decay. Regions of interest were drawn on each organ in anterior and posterior views and on the transmission map. The geometric mean of the activity in each organ was determined and then corrected for average attenuation over the organ. Syringe image data were used to convert the results to percentages of the total radioactivity injected.

These animal studies with dogs were performed to obtain a first approximation of the expected radiation dose absorbed in human subjects. Despite differences between animal and human pharmacokinetic behaviors, an extrapolation process was used to estimate human dosimetry from the calculated dog biodistribution. To compensate for the physiologic time difference between species, allometric scaling was applied to the biodistribution curves, assuming human physiologic time to be 1.19 times slower than canine physiologic time (fourth root of the mass ratio) \((17)\).

RESULTS

Lung Perfusion in Pulmonary Arterial Hypertension

Male Sprague–Dawley rats weighting 200–225 g received a 0.5-mL intraperitoneal injection of either 0.9% saline \((n = 4)\) or monocrotaline at 60 mg/kg \((n = 4)\). Five weeks later, rats were anesthetized for hemodynamic measurements and nuclear medicine experiments with PulmoBind as previously described \((4)\).

Statistics

Statistical analyses were assessed with Prism 4.0 software (GraphPad) and an unpaired Student \(t\) test or a 1-way ANOVA.
linear AM had been directly labeled through its free cysteine residues and purified with a SepPak cartridge to obtain a final product with a high radiochemical purity (≥95%) (5). However, the initial yield of 99mTc-labeled AM, that is, before the purification step, was only about 65% (Fig. 1). Therefore, we added a chelating moiety to trap the radioisotope. A simple tetrapeptide having 99mTc chelating properties, Gly-Gly-Δ-Ala-Gly (19), was used. Thus, compound 1 was prepared by complexing the antagonist AM(22–52) with this tetrapeptide, 99mTc chelating moiety through a molecular linker, the heterobifunctional polyethylene glycol derivative Fmoc-Δ-(polyethylene glycol)1-COOH. Two cysteine residues were introduced into the AM analog construction to generate cyclic compound 2. Finally, compound 3 was designed with Cys acetamidomethyl (Acm) residues, which cannot form a disulfide bridge because their sulfhydryl moieties are blocked by an Acm group, to verify the impact of cyclization on lung uptake and the influence of cysteine residues in the labeling process.

All peptide syntheses yielded only 1 major product that was isolated by reversed-phase HPLC and confirmed to be the desired AM analog by matrix-assisted laser desorption ionization–time-of-flight mass spectrometry. Formation of the disulfide bond was monitored by analytic reversed-phase HPLC, the cyclized peptide having a shorter retention time than its linear counterpart (results not shown).

99mTc Labeling

As expected, the introduction of a chelating moiety greatly improved the 99mTc labeling yield of AM analogs over that of linear AM (Fig. 1). In fact, the labeling of AM through its free cysteine residues resulted, on average, in 65% 99mTc-labeled AM, whereas the labeling yield was higher than 80% when the tetrapeptide was used as the 99mTc chelator. Moreover, the labeling yield obtained with compounds 1, 2, and 3 was not markedly affected by the nature of the peptide, although yields varied slightly with compound 2, which had the greatest ability to retain 99mTc (Fig. 1).

In Vivo Biodistribution and Lung Uptake

Because satisfactory 99mTc labeling yields were obtained with the 3 AM analogs, their lung uptake ability was evaluated in anesthetized dogs. Purified labeled compound 1, 2, or 3 (radiochemical purity of ≥95%, as evaluated by ITLC) was injected intravenously into dogs, and then static evaluation was performed. All 3 compounds showed reduced lung uptake compared with that of AM (Fig. 2). However, compound 2 had a lung first pass comparable to that observed with AM (~75% of the injected dose), rapidly followed in the first minute by a 40% uptake plateau maintained for up to 1 h (Fig. 2). For compounds 1 and 3, initial lung uptake was considerably reduced (~55% of the injected dose), and the compounds were continuously cleared from the pulmonary circulation (Fig. 2).

Interestingly, at 30 min after injection, heart uptake of compound 2 was low (3.1% ± 0.3% [mean ± SD])—even lower than that of AM (4.9% ± 1.0%)—although not significantly different, an important asset for obtaining good lung imaging (Fig. 3). Moreover, liver uptake of compound 2 was comparable to that of AM. Thus, with liver uptake of less than 10% (8.2% ± 0.8%) of the injected dose, lung uptake at least 5 times as high as heart uptake, and lung retention lasting up to 1 h, compound 2 appeared to be a good candidate for lung imaging (Fig. 4). Further analyses were performed to characterize this compound, renamed PulmoBind.

Binding Assays

For further characterization of this new lung imaging agent, binding assays were performed with dog lung homogenates and 99mTc-PulmoBind as the tracer. Saturation experiments demonstrated that PulmoBind bound to specific binding sites in the dog lung at a density of 2.317 ± 0.320 fmol/mg (maximum binding potential [Bmax]), with an affinity in the nanomolar range (dissociation constant [Kd], 2.6 ± 0.8 nM) (Figs. 5A and 5B). These values indicated that PulmoBind could occupy more binding sites than AM in the dog lung but at a lower affinity. Indeed, a similar set of experiments was performed with AM and the same preparation (16), thus allowing a comparison. It was found that AM bound specific binding sites at a density of 1.222 ± 0.148 fmol/mg, with a Kd of 0.17 ± 0.07 nM.

For verification that PulmoBind had a binding site different from that of AM in the dog lung, competitive displacement experiments were performed. With either unlabeled AM or PulmoBind, displacement curves were statistically indistinguishable, suggesting that AM and PulmoBind competed for the same binding site in the dog lung (Fig. 5C).

Hemodynamic Evaluation

AM is a known vasodilator. When injected intravenously into anesthetized dogs, increasing doses of AM produced a drop in mean arterial blood pressure (MAP) that was accompanied by an elevation in heart rate (Fig. 6A). Similarly, the truncated AM agonist, human AM(13–52), generated decreases in MAP proportional to the increasing doses of peptide injected. It was noteworthy that the accompanying...
increase in heart rate was less pronounced than that observed with AM, although the decreases in MAP were similar (Fig. 6B). When cumulative doses of human AM(22–52), a specific AM antagonist, were injected, no changes in MAP or heart rate were observed, even at a dose 50 times the initial injection (Fig. 6C). In a similar manner, cumulative intravenous injections of PulmoBind, which is derived from the latter antagonist, did not produce significant variations in MAP and heart rate, suggesting that this new lung imaging agent also acts as an antagonist (Fig. 6D). Finally, mean pulmonary artery pressure did not vary significantly after injections of human AM(22–52) and PulmoBind and was within expected values.

Dosimetric Analysis

A dosimetric analysis of PulmoBind is shown in Table 2. This analysis was extrapolated to humans from the biodistribution obtained in dogs. The analysis revealed a favorable profile, with rapid elimination of the tracer into urine and the digestive tract, with the kidneys receiving the highest radiation dose (0.034 mGy/MBq) (Table 2). The total effective doses were determined (under allometric scaling) to be 0.0075 and 0.0094 mSv/MBq for men and women, respectively, after 2 h (Table 3).

Pulmonary Arterial Hypertension Model

As shown in Figure 7, the in vivo biodistribution revealed markedly reduced lung uptake of PulmoBind, from 12 ± 2 percentage injected dose in control rats to 4 ± 1 percentage injected dose in rats with pulmonary arterial hypertension (P < 0.001).

DISCUSSION

Nuclear medicine offers clinicians novel avenues for the diagnosis and therapy of various pathologic conditions with noninvasive and rapid techniques. In fact, the success of nuclear medicine relies on the effectiveness of radiopharmaceutical compounds. These compounds, consisting of a target-specific moiety and a radionuclide, must be well designed and finely tuned to attain the desired results for a particular purpose. Because of their inherent specificity, antibodies are biomolecules of interest for generating pharmaceutical compounds. However, their poor pharmacokinetics and their tendency to evoke an immunogenic response (20) limit their effectiveness. On the other hand, peptides, which are also highly specific biomolecules, proved to be useful targeting moieties for generating therapeutic or imaging agents (21,22) with the advantages of being practically devoid of immunogenicity and showing favorable pharmacokinetics (rapid clearance from blood). Moreover, they are quite flexible in terms of chemical modification, allowing radiolabeling (23).

We previously showed that the linear AM peptide, once labeled with 99mTc, was an attractive lung imaging agent for the diagnosis of pulmonary embolism (5). Lung perfusion scintigraphy with 99mTc-labeled macroaggregates of albumin (99mTc-MAA) is generally con-
logic effect, the safety of clinical procedures is a crucial issue. Imaging agents are generally used at a dose devoid of any biogenicity, a less desirable outcome. However, AM is recognized as a cardioreactive peptide. As a vasodilator, it could be perceived as beneficial, especially if used for the diagnosis of pulmonary hypertension. In fact, AM is a biologically active peptide. As a vasodilator, it could be perceived as beneficial, especially if used for the diagnosis of pulmonary hypertension. In fact, AM is recognized as a cardioreactive peptide. As a vasodilator, it could be perceived as beneficial, especially if used for the diagnosis of pulmonary hypertension.

FIGURE 6. Hemodynamic evaluation in dogs of AM (A), agonist AM(13–52) (B), antagonist AM(22–52) (C), and PulmoBind (D). Anesthetized and ventilated mongrel dogs were injected intravenously with cumulative doses of each peptide, and heart rate (green line), mean arterial pressure (black line), and mean pulmonary artery pressure (red line) were monitored. Initial dose (lung scan dose) was maximal dose anticipated for humans (555 MBq; 18.5 μg) in phase 1 clinical study. Animal dose equivalence was computed for body surface area to monitor hemodynamic effects. Each graph depicts typical trace obtained from 2–7 evaluations. bpm = beats per minute.

We demonstrated the relevance of this approach using a monocrotaline-induced pulmonary hypertension rat model (4). However, AM is a biologically active peptide. As a vasodilator, it could be perceived as beneficial, especially if used for the diagnosis of pulmonary hypertension. In fact, AM is recognized as a cardioreactive modulator (25). However, AM also stimulates angio genesis, a less desirable outcome (26). Although nuclear medicine imaging agents are generally used at a dose devoid of any biologic effect, the safety of clinical procedures is a crucial issue that prompted us to develop a lung imaging agent with antagonist pharmacology.

Many chelators trap 99mTc; these include diethylenetriaminepentaacetic acid, mercaptoacetyltriglycine, and 6-hydrazinopyridine-3-carboxylic acid. However, a simple tetrapeptide having 99mTc chelating properties appeared to be more attractive because it was introduced to the peptide chain during synthesis, in accordance with the procedure used for all other amino acids. Incorporation of this peptide chelating moiety greatly improved the labeling yield without any significant difference among the 3 compounds synthesized (Fig. 1). A difference appeared when lung uptake was evaluated. As a matter of fact, the presence of a cyclic structure proved to be crucial for efficient lung uptake because at just 10 min after injection, about 20% of the linear AM analogs (compounds 1 and 3) remained in the lungs, whereas more than 40% of AM and the cyclic AM analog (compound 2) were retained in the organ; this activity was maintained even after 30 min (Fig. 2). These results are in accordance with the results of structure–activity relationship studies of peptides belonging to the calcitonin family. Circular dichroism analyses of AM demonstrated that the peptide structure is composed of 28% α-helix and 18% β-sheet. Interestingly, the antagonist AM(22–52) shares these structural features (14). However, the ring structure composed of 6 residues linked by a disulfide bridge is a characteristic common to all calcitonin peptide family members, despite their low sequence homology, and it has been shown to be important for proper binding and subsequent signaling (13,26). Moreover, amino acid substitution within the ring structure does not seem to markedly alter the biologic activity of the peptide (26). Therefore, combining AM(22–52) with a flexible ring structure composed of 2 cysteine residues linked by a polyethylene glycol spacer as a substitute for the 6 residues generated a molecule encompassing all of the AM structural features important for ensuring proper receptor binding while avoiding a biologic response. In fact, the use of the polyethylene glycol spacer allowed appropriate spacing without introducing reactive or sensitive chemical groups or

TABLE 2

<table>
<thead>
<tr>
<th>Organs</th>
<th>Radiation dose (mGy/MBq)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lungs</td>
<td>0.005</td>
</tr>
<tr>
<td>Liver</td>
<td>0.012</td>
</tr>
<tr>
<td>Kidneys</td>
<td>0.034</td>
</tr>
<tr>
<td>Spleen</td>
<td>0.006</td>
</tr>
<tr>
<td>Bladder</td>
<td>0.010</td>
</tr>
<tr>
<td>Testes</td>
<td>0.002</td>
</tr>
<tr>
<td>Ovaries</td>
<td>0.005</td>
</tr>
<tr>
<td>Total body</td>
<td>0.003</td>
</tr>
</tbody>
</table>
increasing too much the hydrophobicity of the molecule. Furthermore, the chelating moiety was added at the N terminus of the peptide because it was demonstrated that labeling of the peptide at any other position could affect receptor interaction (16).

The results of our binding study correlated with AM structure–activity relationships. Indeed, AM had a \( K_d \) of 0.17 ± 0.07 nM for dog lung homogenates (16), and PulmoBind (compound 2) had a \( K_d \) of 2.6 ± 0.8 nM for the same preparations (Fig. 5). Given that the same conditions were used to evaluate binding, AM(22–52) had a binding affinity about 100 times lower than that of AM, and even linear AM (reduced disulfide bridge) had a markedly reduced affinity (6). Thus, adding a cyclic structure to AM(22–52) to generate PulmoBind enhanced binding significantly. Moreover, the number of accessible binding sites appeared to be higher for PulmoBind than for AM because \( B_{\text{max}} \) values were 2,317 ± 320 fmol/mg (Fig. 5) and 1,222 ± 148 fmol/mg (16), respectively. This observation can be related to studies of somatostatin, corticotrophin-releasing factor, and bombesin demonstrating that radioactive ligand antagonists label more receptor binding sites than corresponding agonists (11,27,28). Therefore, it appears that the slight loss of binding affinity of PulmoBind is compensated for by a larger number of accessible binding sites, allowing pulmonary circulation imaging similar to that observed with the full-length AM peptide labeled with \(^{99m}\text{Tc}\).

As previously mentioned, the AM disulfide ring structure is implicated in proper receptor interaction and the receptor signaling process. Accordingly, the PulmoBind cyclic moiety has enhanced receptor binding, but this analog appears to still act as an antagonist, like its parent molecule, as observed in an in vivo hemodynamic evaluation (Fig. 6). In fact, AM(22–52) did not evoke any changes in MAP, heart rate, or mean pulmonary artery pressure even at 50 times the imaging dose, and similar observations were obtained for PulmoBind; these observations suggest that an AM-based imaging agent such as PulmoBind should not have any acute toxicity effects on the pulmonary vascular system during the course of the examination. In contrast, the truncated AM analog comprising the ring structure, that is, AM(13–52), elicited a vasodilation response at only 2.5 times the imaging dose, just like AM. Interestingly, the increase in heart rate was less important with the N-terminally truncated analog, suggesting that this portion of the peptide may be responsible, at least in part, for this specific biologic response.

In the design of targeted imaging compounds, the goal is to obtain high tissue specificity with a high signal-to-noise ratio, appropriate pharmacodynamics, and good pharmacokinetics. Injection of PulmoBind into anesthetized dogs generated a good lung image with low heart and liver uptake, an essential feature for pulmonary vasculature imaging, even at 60 min after injection (Fig. 4). The lung kinetic profile of PulmoBind was characterized by an uptake plateau, suggesting irreversible binding to its pulmonary receptor. As determined in our in vivo biodistribution study, PulmoBind was retained not only in the pulmonary circulation but also in the kidneys (Fig. 3). As a matter of fact, labeling of the kidneys and bladder was higher than that of other tissues at later time points, with a concomitant decrease in lung, heart, and liver uptake, indicating that the molecule was eliminated mainly through renal excretion. This clearance route was also observed with \(^{99m}\text{Tc}\)-labeled AM (5). Thus, PulmoBind appears to offer good pulmonary circulation imaging properties, without adverse hemodynamic effects and with urinary elimination, qualities that make it a promising compound for clinical use. However, this elimination route was associated with our finding that the kidneys were the critical organ in terms of radiation dose. Moreover, the close proximity of the urinary tract to radiation-sensitive tissues, such as the reproductive organs, could be a safety concern. However, our dosimetric analysis revealed that the testes and ovaries received the least

### Table 3

<table>
<thead>
<tr>
<th>Radiopharmaceutical</th>
<th>Administered activity (MBq)</th>
<th>Critical organs</th>
<th>Total dose (mSv)</th>
<th>Effective dose (mSv/MBq)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(^{99m}\text{Tc})-PulmoBind</td>
<td>185–555</td>
<td>Kidneys</td>
<td>1.4–4.2</td>
<td>0.0075</td>
</tr>
<tr>
<td><strong>Men</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Women</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(^{99m}\text{Tc})-MAA</td>
<td>185</td>
<td>Lungs</td>
<td>2.0</td>
<td>0.011</td>
</tr>
<tr>
<td>(^{99m}\text{Tc})-DTDA</td>
<td>370</td>
<td>Bladder</td>
<td>1.8</td>
<td>0.0061</td>
</tr>
<tr>
<td>(^{99m}\text{Tc})-DMSA</td>
<td>370</td>
<td>Kidneys</td>
<td>3.3</td>
<td>0.0088</td>
</tr>
</tbody>
</table>

DTPA = diethylenetriaminepentaacetic acid; DMSA = dimercaptosuccinic acid.
radiation among the evaluated organs (Table 2), and even the kidneys received lower doses than other critical organs identified with clinically used 99mTc-labeled radiopharmaceuticals (Table 3). Finally, with the proposed injected activity (185–555 MBq), the total effective radiation dose (1.7–5.2 mSv for a woman) would be comparable to those used in actual nuclear medicine procedures because doses of 0.2–14 mSv are typically used in clinical radiographic examinations (29).

Finally, we demonstrated that radiolabeled PulmoBind could detect a lung microcirculatory perfusion defect associated with pulmonary arterial hypertension using the monocrotaline model. This model causes progressive obliteration of pulmonary vessels, and we demonstrated that it was associated with reduced lung expression of the specific heterodimeric component of the AM receptor, RAMP2 (4). Evidence for the abundant distribution of AM receptors in the pulmonary microcirculation was also obtained previously. A study evaluating the distribution of the calcitonin receptorlike receptor, the other heterodimeric component of the specific AM receptor (CRLR/RAMP2), demonstrated intense staining in the lung capillaries (30). Another study demonstrated colocalization of the endothelial cell marker and the calcitonin receptorlike receptor in pulmonary capillaries and lung endothelial cell expression of the calcitonin receptorlike receptor and RAMP2 (31).

CONCLUSION

Because nuclear medicine definitely benefits from precise targeting, the development of specific radiotherapeutic drugs and medical imaging agents has attracted much interest, with peptides and proteins forming a large part of the starting material. Many peptides have been used to create new specific radiopharmaceutical compounds, but our work supports studies of somatostatin indicating the usefulness of peptide antagonists as nuclear medicine agents (27,32). PulmoBind, an AM analog, binds AM receptors found at a high density in the pulmonary microvascular endothelium with a high affinity and without causing any adverse hemodynamic response after systemic injection. Once labeled with 99mTc, it provides good pulmonary circulation imaging with a nuclear medicine camera for at least 1 h, and the compound is efficiently eliminated through renal excretion. Thus, PulmoBind labeled with 99mTc is a radiopharmaceutical with the potential to provide new molecular modalities for pathologies such as pulmonary hypertension and pulmonary embolism.

DISCLOSURE

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