Blood flow measurements in rats using four color microspheres during blockade of different vasopressor systems

K. De Angelis1,4, V.M. Gama1,2, V.A.M. Farah3 and M.C. Irigoyen1,2

Abstract

The use of colored microspheres to adequately evaluate blood flow changes under different circumstances in the same rat has been validated with a maximum of three different colors due to methodological limitations. The aim of the present study was to validate the use of four different colors measuring four repeated blood flow changes in the same rat to assess the role of vasopressor systems in controlling arterial pressure (AP). Red (150,000), white (200,000), yellow (150,000), and blue (200,000) colored microspheres were infused into the left ventricle of 6 male Wistar rats 1) at rest and 2) after vasopressin (aAVP, 10 µg/kg, iv), 3) renin-angiotensin (losartan, 10 mg/kg, iv), and 4) sympathetic system blockade (hexamethonium, 20 mg/kg, iv) to determine blood flow changes. AP was recorded and processed with a data acquisition system (1-kHz sampling frequency). Blood flow changes were quantified by spectrophotometry absorption peaks for colored microsphere components in the tissues evaluated. Administration of aAVP and losartan slightly reduced the AP (-5.7 ± 0.5 and -7.8 ± 1.2 mmHg, respectively), while hexamethonium induced a 52 ± 3 mmHg fall in AP. The aAVP injection increased blood flow in lungs (78%), liver (117%) and skeletal muscle (>150%), while losartan administration enhanced blood flow in heart (126%), lungs (100%), kidneys (80%), and gastrocnemius (75%) and soleus (94%) muscles. Hexamethonium administration reduced only kidney blood flow (50%). In conclusion, four types of colored microspheres can be used to perform four repeated blood flow measurements in the same rat detecting small alterations such as changes in tissues with low blood flow.

Microsphere techniques provide more detailed information about regional perfusion between and within organs than the use of flow probes. Microspheres are also easier to use and give higher resolution than methods based on molecular tracer washout (1). The use of colored microspheres (CM) represents a simple and safe method, especially...
because it avoids the radiation associated with radioactive spheres (2-4). Frequently, repeated blood flow measurements must be performed in the same animal (same tissue or organ) (5-7). For this purpose, microspheres labeled with different color components are injected, and the components are then measured simultaneously. However, the determination of the amount of the different components may be disturbed by interference between the different absorption spectra. Indeed, CM have been further validated (4,6,8) and employed extensively, although usually with a maximum of three different colors (7). The present study was designed to test the possibility to use four different microsphere colors: white (370 nm), yellow (448 nm), red (530 nm), and blue (672 nm).

To assess the individual contribution and the interaction of three vasopressor systems in blood pressure control and blood flow distribution we used four repeated estimates in the same rat, i.e., a basal one and three more after vasopressin, renin-angiotensin and sympathetic system blocker injection. Thus, we obtained four repeated blood flow estimates using a combination of different numbers and colors by adapting the standardized infusion protocols, dye extraction procedures and microsphere measurements.

Six male Wistar rats from the Animal House of Universidade de São Paulo, São Paulo, SP, Brazil, weighing 290 ± 5 g, received standard laboratory chow ad libitum and were housed in individual cages in a temperature-controlled room (22°C) with a 12-h dark/light cycle. All surgical procedures and protocols used were in accordance with the Guidelines for Ethical Care of Experimental Animals and Use Committee.

The rats were anesthetized with a mixture of ketamine (80 mg/kg) and xylazine (12 mg/kg). Catheters filled with saline were implanted into the femoral artery and vein and into the left ventricle. PE-10 catheters were positioned into the abdominal aorta and the vena cava by way of the femoral artery and vein for direct measurements of arterial pressure (AP) and for drug injections, respectively. A third PE-50 catheter was inserted into the left ventricle via the right carotid artery for infusion of the CM. The position of the catheter was determined by observing the characteristic left ventricular pressure waveform at surgery and confirmed at autopsy. The catheters were anchored with silk sutures and exteriorized on the back of the neck. Rats receiving food and water ad libitum were studied one day after catheter placement; the animals were conscious and allowed to move freely during the experiments.

The femoral artery cannula was connected to a strain-gauge transducer (Narco Bio-Systems Miniature Pressure Transducer RP 1500, Houston, TX, USA) and blood pressure signals were monitored continuously, except during microsphere infusion and withdrawal of a reference blood sample. The AP signals were recorded and processed with a microcomputer equipped with an analog-to-digital converter board (CODAS, 1-kHz sampling frequency; Dataq Instruments, Inc., Akron, OH, USA). The recorded data were analyzed on a beat-to-beat basis to quantify changes in mean AP and heart rate (HR) as described elsewhere (9).

After 20 min of basal AP recording, a vasopressin V1 receptor antagonist (aAVP, 10 µg/kg; Sigma, St. Louis, MO, USA) was administered iv and the AP signals were recorded over a period of 10 min. When AP returned to basal values (≅25 min), losartan, an angiotensin II AT1 receptor antagonist (10 mg/kg; Du Pont Merck, Wilmington, DE, USA), was injected iv and AP was recorded for 10 min. Finally, after a period of recovery (≅25 min) AP returned to baseline values and hexamethonium, a sympathetic ganglion blocking drug (20 mg/kg; Sigma), was administered iv and blood pressure was recorded over a period of 10 min (10). The efficacy of blockade was determined by using vasopressin and angiotensin II after aAVP.
Flow measurements using four color microspheres and losartan administration, respectively. The responses to vasopressor system blockade were analyzed by averaging mean arterial pressure (MAP) values obtained during a 5- to 10-min recording period after the administration of each drug.

Dye-Trak CM (15 µm; Triton Technology, San Diego, CA, USA) were infused at rest (red: 150,000 CM) and 10 min after aAVP (white: 200,000 CM), losartan (yellow: 150,000 CM) and hexamethonium (blue: 200,000 CM) administration for blood flow and cardiac output (CO) determination. Microsphere infusion and processing were performed by the method of Hakkinen et al. (4).

Vials of commercial stocks of microsphere infusates were sonicated for 5 min and inverted several times immediately before dilution to the desired concentration with 0.9% saline containing 0.01% Tween 80. To determine the average dye content of the microspheres, 200 µl of the diluted commercial suspensions was taken after 3 min of suspension sonication. These aliquots were placed in a 15-ml tube and were centrifuged, dried, and extracted with dimethylformamide (DMF; Sigma). The mean absorbance was determined with a spectrophotometer and divided by the average microsphere concentration [absorbance units (spheres ml⁻¹)]⁻¹.

The sphere dilutions were sonicated for 3 min before infusion. A coil of PE-50 tubing (75 cm) was then filled with 180 µl of the infusate of CM and interposed between the left ventricular catheter and a 1-ml syringe containing 0.5 ml of prewarmed (40°C) saline. This syringe was mounted on a variable speed infusion pump (Infusion Pump 22; Harvard Apparatus, South Natick, MA, USA). With a withdrawal pump (Infusion and Withdrawal Pump; Harvard Apparatus), reference blood samples were drawn from the abdominal aorta catheter at the rate of 0.5 ml/min into a pre-heparinized and weighed 1-ml disposable syringe. The withdrawal of a reference blood sample was started 10 s before the beginning of microsphere infusion and was continued for 75 s. The microspheres and saline were infused for 50 s at a rate of 0.36 ml/min. After microsphere infusion the animals were killed with a thiopental overdose and heart, lung, kidneys, liver, and the entire gastrocnemius, soleus and brachial triceps muscles were removed to determine regional blood flow.

The reference blood samples and tissues were processed as described by Hakkinen et al. (4). To extract the dyes from the isolated dried microspheres, 250 µl DMF was added to each tube, which was briefly but vigorously vortexed. The samples were centrifuged at 2000 g for 10 min and the absorbance of the supernatant was determined with a DU 640 spectrophotometer (<1.8-nm slit width; Beckmann Instruments, Inc., Fullerton, CA, USA) using a 200-µl quartz cuvette (Sigma). The absorption spectrum peaks for the white, yellow, red, and blue microspheres were obtained at 370, 448, 530, and 672 nm, respectively. The absorbances were corrected for overlapping using a matrix inversion technique (Dye-Trak® matrix macro-inversion for Excel; Triton Technology, San Diego, CA, USA). The minimum acceptable absorbance was 0.010 absorbance units (AU).

For each infusion, the tissue flow rates were calculated according to (4):

\[ \text{Qt} = \frac{\text{At} \times (\text{Qb} \div \text{Ab})}{\text{Reference blood sample weight} \div 1.05 \text{ g/ml}} \]

where Qt and Qb represent flow in the sample tissue and in the reference blood, respectively; At and Ab represent the peak absorbance (AU) of the tissue sample and of the reference blood, respectively. The Qb, in ml/min, was calculated by:

\[ \text{Qb} = \frac{\text{Reference blood sample weight} \div 1.05 \text{ g/ml}}{\text{Reference blood sample volume} \div 0.5 \text{ ml/min}} \]

where 1.05 g/ml is the specific gravity of blood and 0.5 ml/min the withdrawal rate.

Blood flow rates were divided by tissue
weights to yield ml min\(^{-1}\) g\(^{-1}\). CO was calculated by the following formula (11):

\[
\text{Total number of injected microspheres} \times \frac{\text{reference rate (0.5 ml/min)}}{\text{Number of microspheres in the reference blood sample}}
\]

Data are reported as mean ± SEM, and the paired \(t\)-test and repeated measure ANOVA were used to compare data, followed by the Tukey test. A \(P\) value of 0.05 was considered to be statistically significant.

Figure 1 shows the spectra of the four different colors of microspheres (red, white, yellow, and blue, injected at rest, post-aAVP, post-losartan, and post-hexamethonium, respectively) presented in the dye solution obtained from individual samples of heart, right and left kidneys, soleus and gastrocnemius muscles from one rat. Arrows indicate the wavelength of the absorbance of white (370 nm), yellow (448 nm), red (530 nm), and blue (672 nm) colors for the calculation of tissue blood flow. These results showed individual absorption spectrum peaks with no interference for the white (370 nm), yellow (448 nm), red (530 nm), and blue (672 nm) components of the microspheres in the heart, lungs, kidneys, and the gastrocnemius and soleus muscle tissues, demonstrating the ease of accurately measuring each color used. Similar spectrum peaks were obtained for these tissues and for the liver and triceps brachial muscle in all rats studied.

After blockade of each vasopressor system, AP was recorded (10 min) and the microspheres were infused. Before starting each protocol step, a 25-min interval was allowed to elapse between the use of the different blockers for blood pressure to re-
Flow measurements using four color microspheres

123

Flow measurements using four color microspheres

Braz J Med Biol Res 38(1) 2005

Flow measurements using four color microspheres

Braz J Med Biol Res 38(1) 2005

Flow measurements using four color microspheres

Braz J Med Biol Res 38(1) 2005

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Braz J Med Biol Res 38(1) 2005

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Braz J Med Biol Res 38(1) 2005

Flow measurements using four color microspheres

Braz J Med Biol Res 38(1) 2005
nique, which usually requires a larger number of CM than that required by the radioactive method to produce a minimum detectable signal.

In the present study, infusion and withdrawal rates were changed to allow measurements of a slight reduction in AP such as that observed with aAVP and/or losartan administration. The rates previously described by Hakkinen et al. (4) induced increased AP although it was established that to accurately evaluate small blood flow changes, microsphere infusion should not change hemodynamic parameters. In the present study, we used a spectrophotometer for colorimetric measurements instead of a microplate reader (4) or HPLC (8), which reduced the cost of the necessary equipment, but increased the overall time required for these measurements.

Heymann et al. (11) have provided a comprehensive review of the use and validation of the microsphere method. Recently, Prinzen and Bassingthwaighte (1) discussed the pros and cons of radioactive, colored and fluorescent microspheres. The CM technique allows the measurement of regional blood flow in a manner quite similar to that of radioactive microspheres (4,7,8,13) while avoiding the disadvantage of radioisotope decay, limiting their lifetime, and the handling of radioactive materials. In a study that used fluorescent and CM simultaneously and compared them with radioactive microspheres, no differences were found in the accuracy of blood flow values obtained with the two types of non-radioactive spheres (7). In the present study, the basal CO and regional blood flow values were similar to values obtained by using radiolabeling and dye extraction methods in rats (4,14). Moreover, the similar spectra and blood flows observed in the right and left kidneys during the four color evaluations provided evidence for good mixing and distribution of the microspheres in the circulation, since changes in kidney flow are used to analyze this quality parameter (4).

The changes in MAP observed in the present experiment were similar to those obtained in a previous study (10,15). The V1 blockade did not change basal AP values, probably because the increase in lung, liver and skeletal muscle blood flow was accompanied by a slight increase in CO. Similar findings were observed with losartan administration, with the increase in regional blood flow being associated with no changes in blood pressure. Indeed, Ullman et al. (16) have reported increased regional blood flow changes without concomitant effects on systemic hemodynamics.

The marked decrease in MAP observed after hexamethonium injection agreed with the data reported by Fink and Ploucha (17). However, these investigators reported that sympathetic blockade reduced blood pressure by vasodilation alone since no effect on CO was observed. The data reported in the present study did not show significant changes in regional blood flow after hexamethonium injection, except for the reduction in kidney blood flow. This reduction may be associated with decreased CO. However, the blood pressure decrease induced by sympathetic blockade produced only a non-significant reduction in CO (20%), indicating that other changes in regions in which measurements were not performed should be considered.

Associated with this impaired reduction in blood pressure, hexamethonium administration induced an increase in HR, probably representing a baroreflex-mediated tachycardia induced by the marked decrease in AP. This HR increase is probably due to a vagal withdrawal, since sympathetic activity was blocked. Interestingly, the magnitude of the peak depressor response seems not to be associated with V1/AT1 receptor blockade since Santajuliana et al. (10) have previously demonstrated no changes in the depressor response with and without V1/AT1 blockade.
The results of the present study demonstrate that four repeated blood flow measurements in the same rat with the use of four CM (white, yellow, red, and blue) can be performed accurately to detect small alterations in low blood flow tissues, suggesting that this technique is potentially useful for estimating blood flow in the same rats in at least four different situations. Using this approach, it was possible to describe changes in regional blood flow and CO induced by vasopressor system blockade in conscious rats.

References