Comparison of the protocols for obtaining platelet-rich plasma in dogs: a cellular study

André William Masseaux Vidal Júnior1* Ricardo Siqueira da Silva2
Ana Paula Lopes Marques2 Heloisa Justen Moreira de Souza2

1Programa de Pós-graduação em Medicina Veterinária, Universidade Federal Rural do Rio de Janeiro (UFRJR), 23890-000, Seropédica, RJ, Brasil. E-mail: andremasseaux@gmail.com. *Corresponding author.
2Departamento de Medicina e Cirurgia Veterinária, Universidade Federal Rural do Rio de Janeiro (UFRJ), Seropédica, RJ, Brasil.

ABSTRACT: This study aimed to evaluate two protocols (PA and PB) that are used to obtain canine platelet-rich plasma (PRP) for cellularity. Twenty healthy dogs were used. Blood samples were collected and placed in two tubes of 3.2% sodium citrate. PA used double centrifugation at 210 x g and 370 x g and PB used double centrifugation with 140 x g and 330 x g. The PRP samples from the protocols were examined in terms of their platelet, erythrocyte, and leukocyte count in the Neubauer chamber, differential leukocyte count and platelet morphological observation in blood smears. Data (mean and standard deviation) were analyzed with the 95% probability t-test (P <0.05) using Pearson’s correlation to test the relationship between platelets and erythrocytes, platelets, and leukocytes, and the leukocyte count versus the erythrocytes. Very weak negative correlation between platelets and leukocytes (p = -0.03), weak negative correlation between platelets and erythrocytes (p = -0.3) and a strong positive correlation between leukocytes and erythrocytes (p = 0.75) were noted. Although, BP did not reach the desired mean of one million platelets (979300 ± 79631 cells / μL), both protocols, A and B (4.42 ± 1.61 and 3.85 ± 1.55 times more platelets than total blood, respectively) (p <0.05) were efficient in concentrating platelets. Platelet activation was present in 26.55 ± 6.72% of the PA platelets and 26.25 ± 7.03% in PB (p> 0.05). PA and PB presented low erythrocyte concentration (p> 0.05), and PA had more leukocytes (p <0.05) than PB, with higher concentrations of basophils that were segmented, and lymphocytes.

Key words: growth factors, platelet-rich plasma, cell therapy, canines.

INTRODUCTION
Platelet-rich plasma (PRP) is generally autologous and consists of a small volume of plasma with a high concentration of platelets, growth factors (GF), and cytokines for epithelial migration, bone and connective tissue formation (MARX, 2004; CROVETTI et al., 2004; HENDERSON et al., 2003). The PRP quality is mainly measured by the ability to concentrate platelets. The PRP should have three to five times more platelets than whole blood (WHITLOW et al., 2008). However, other studies have indicated that a concentration > 1 million platelets/μL is necessary (MARX et al., 1998; LEMOS, 2002).
Dogs can benefit from PRP, associated or not with other biomaterials, in bone grafts and bone tissue regeneration, and integration in cases of experimental fractures (YOU et al., 2007; CASATI et al., 2007). The semi-automatic PRP processing method using a conventional centrifuge in a laboratory or outpatient environment provides low-cost PRP and greater applicability in veterinary medicine (VENDRAMIN et al., 2006).

All protocols should increase the platelet count to a concentration well above that reported in whole blood, preserving the platelet structure and function. If platelets are activated by releasing the GF earlier, the therapeutic action of PRP will be impaired (LOPEZ et al., 2012).

High leukocyte concentrations in PRP may delay tendon and joint healing processes due to an increased expression of inflammatory cytokines (McCARREL et al., 2012; SCHNABEL et al., 2007; CARMONA et al., 2007). However, in contaminated skin or bone lesions, the use of PRP with higher leukocyte concentration is beneficial (MARX & GARG, 1999; BIELECKI et al., 2007; BARBOSA et al., 2008). Red blood cells in PRP may increase gel viscosity after platelet activation (ROSSI, 2001), but pain perception may increase after use due to local irritation (HALPERN et al. 2012).

In this context, the evaluation of two protocols was proposed to obtain autologous canine PRP. These semi-automatic protocols are easy to perform in an outpatient clinic and have good quality levels (platelet concentration capacity, qualitative evaluation of platelet morphology, qualitative and quantitative leukocyte classification, and reduced contamination with red blood cells).

MATERIALS AND METHODS

The study was approved by the committee on animal research and ethics of the Federal Rural University of Rio de Janeiro (CEUA-UFRRJ) under protocol no. 1827171016. All animal owners signed an informed consent form.

This study included twenty dogs (*Canis lupus familiaris*) of both sexes treated at the Veterinary Hospital of the Federal Rural University of Rio de Janeiro (UFRRJ), Brazil. All dogs were aged between 1 and 7 years, with a mean age of 4 years. All presented with no clinical changes (routine consultations or elective castration procedures) and underwent blood collection. After shaving and disinfecting the cervical region with 70% alcohol, the external jugular vein was punctured using a 22G needle and 10 mL syringe to collect 10 ml of blood. Of this, 2 ml of blood was transferred to a tube containing ethylenediamine tetraacetic acid (EDTA) anticoagulant for a complete blood count (CBC) and the remaining 8 ml was divided into two 4 ml tubes containing 3.2% sodium citrate for two PRP protocols.

An impedance hematological analyzer was used for the CBC. An immersion optical microscope with a 100x objective was used for differential leukocyte count using whole blood smear slides that were instantaneously stained (Panótico Rápido®) (JAIN, 1993). Plasma concentrations of total proteins were measured using refractometry (COLES, 1987).

The blood was centrifuged (Daiki Centrifuge 80-2B, Centribio®) at 210 x g for 10 min for protocol A (PA) (ALEIXO et al., 2011, modified). After separation of the supernatant plasma, leukocyte layer, and red blood cell fraction, the supernatant plasma was divided into two fractions using an automatic micropipette. Subsequently, 50% (the upper half) of the supernatant plasma was discarded, and the lower half and the leukocyte layer were stored in a Vacutainer tube without additives for the second centrifugation at 370 x g for 10 min. Thereafter, the supernatant plasma and a red blood cell button were visualized. The automatic micropipette was used to discard 2/3 of the supernatant plasma, and the rest was considered PRP obtained with protocol A (Figure 1). The same procedures were repeated for protocol B (PB) (KIM et al., 2002, adapted) with different centrifugation speeds and plasma fractions. The first centrifugation was at 140 x g for 10 min, and all supernatant plasma was centrifuged a second time at 330 x g for 10 min. The automatic micropipette was used to discard 2/3 of the supernatant plasma, and the rest was considered PRP obtained with protocol B. The centrifugation strength and time of each protocol was based on protocols described by ALEIXO et al. (2011) and KIM et al. (2002) that are considered efficient in concentrating platelets.

PRP samples obtained from each protocol were diluted and homogenized to count platelets, red blood cells, and leukocytes in a Neubauer chamber after resting in a humidified chamber for 5 min. Counts were duplicated, using the mean values obtained for each sample. PRP was diluted at 1/200 (10 μl PRP in 1990 μl Brecher liquid with 1% ammonium oxalate) to determine the number of platelets. The Neubauer chamber was filled with a capillary tube, and after 20 min, an optical microscope with a 40x objective was used for counting the central reticulum (25 central medium squares = 1 mm² area). To determine the number of red blood cells, the PRP was diluted at 1/200 (10 μl PRP in 1990 μl 0.9% saline solution), the
Neubauer chamber was filled with a capillary tube, and after 5 min an optical microscope with a 40x objective was used for counting the central reticulum (25 central medium squares). For leukocytes, the PRP was diluted at 1/20 (20 μl PRP in 380 μl Thoma Liquid), the Neubauer chamber was filled with a capillary tube, and after 5 min an optical microscope with a 40x objective was used for counting the four largest lateral squares (4 mm² area).

The formula used to obtain the total number of cells, platelets, red blood cells, and leukocytes per μl of PRP was the number of cells counted x dilution x 10 (0.1 mm height) x area, as described by COLES (1984).

Differential leukocyte counts were performed using thin, uniform PRP smear slides that were instantaneously stained (Panótico Rápido®) and visualized under an optical microscope in immersion with a 100x objective. The count started from the middle region of the smear and extended towards the tail, zigzagging through the slide. In total, 100 leukocytes were counted, and the result was presented as a percentage (relative leukocyte count). The absolute specific leukocyte count of each PRP was determined by the relationship between the global and relative specific leukocyte counts. The smear slides were used to visualize platelet morphology, with 100 platelets being counted. Their conformation was observed through pseudopod formation when activated, with the result presented as a percentage.

The confidence level was 95%, with a maximum error of 5%, and the standard deviation of the population in the pilot test did not exceed 10 points. Results are presented as a mean, standard deviation, and minimum and maximum values. The data were analyzed using the t-test with a 95% probability (p<0.05). Pearson’s correlation was used to test the relationship between platelet and red blood cell counts, platelets and leukocytes, and leukocytes and red blood cells. Prism version 6.0 software for Windows (GraphPad Software, California, USA) and Microsoft Excel 2010 software were used for the analyses.

RESULTS AND DISCUSSION

Table 1 presents a summary of the data of the variables (platelets, red blood cells, and leukocytes) in whole blood and in each protocol evaluated. Figure 1 shows that in both the PRP protocols the number of platelets differed (p<0.05) in relation to whole blood using a manual method for platelet concentration counting in PRP for its concomitant morphological evaluation and activation analysis.

The protocols were selected for being simple, low-cost, and efficient at concentrating platelets (ALEIXO et al., 2011; KIM et al., 2002). However, leukocytes and red blood cells were not evaluated in previous studies.

Platelet concentration in PRP should be three to five times higher than in whole blood.

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Moreover, both protocols, PA and PB (4.42 ± 1.61 and 3.85 ± 1.55 times more platelets than whole blood, respectively) were efficient in concentrating platelets but differed (P<0.05) from one another (Table 2). However, other studies considered a concentration higher than 1 million platelets/µL (MARX et al., 1998; LEMOS, 2002). According to these authors, PB would not be considered a quality test for presenting a mean platelet concentration of 979,300 ± 79,631 cells/µL (Table 1). Platelet activation did not differ (p>0.05) between protocols and was relatively low (Table 2). The microscopic morphological visualization of platelets using a PRP smear is recommended to evaluate the quality of the blood component, since pseudopods are formed after activation (WEISS; WARDROP, 2010), releasing growth factors in advance (EPPLEY et al., 2004). A previous study showed no morphological changes in PRP platelets obtained by direct microscopy (ALEIXO et al., 2011). However, a study using electronic microscopy to visualize platelet morphology in PRP detected 49% of platelets with uncertain activation, 41% at rest, 9% activated, and 1% damaged (ZANDIN et al., 2012). The present study detected small cytoplasmic prolongations evidencing platelet activation in 26.55 ± 6.72% of the PA platelets, and 26.25 ± 7.03% of PB platelets, with no statistical difference between them. However, the optimal amount of inactive platelets has not been established.

Red blood cells are considered PRP sample contaminants (MARX, 2004). Both protocols were considered of good quality, as demonstrated by reduced red blood cell concentration with no difference (p>0.050) among the evaluated PRPs (Table 2). A lower number of red blood cells was previously correlated with lower gravitational force (PEREIRA, 2012), which corroborates our data since PB showed lower gravitational force (140 x g and 330 x g) than PA (210 x g and 370 x g) and concentrated less red blood cells. According to PEREIRA (2012), lower gravitational force protocols concentrate fewer platelets, which would reduce the quality of the final product. This differs from studies with equine blood, in which higher platelet counts were associated with lower centrifugal forces (VENDRUSCOLO et al., 2012). When evaluating the number of leukocytes in the PRPs, PA showed more white blood cells, significantly differing (p<0.05) from PB (Table 2).

A study on cell proliferation in tendinous lesions using plasma rich in leukocyte platelets (L-PRP) and plasma rich in pure platelets (P-PRP) showed that both treatments induced the differentiation of progenitor tendon cells into active tenocytes. However, L-PRP induced predominantly catabolic and inflammatory changes in differentiated tenocytes, increased the expression of catabolic marker genes, matrix metalloproteinases (MMP), interleukin-1beta (IL-1β), IL-6, tumor necrosis factor-alpha (TNF-α), and prostaglandin E2 (PGE 2) production. Contrastingly, P-PRP mainly induced

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Table 1 - Mean values of the platelets, red blood cells, and leukocytes obtained by the two protocols using whole blood and plasma rich in platelets from 20 dogs.

|                   | WHOLE BLOOD | Protocol A | Protocol B |
|-------------------|-------------|------------|------------|
| **PLATELETS (cells/µl)** |             |            |            |
| Mean              | 285000      | 1135300    | 979300     |
| SD                | 101359      | 158779     | 79631      |
| Minimum           | 118000      | 934000     | 830000     |
| Maximum           | 609000      | 1536000    | 1162000    |
| **RED BLOOD CELLS (cells/µl)** |         |            |            |
| Mean              | 6577000     | 67800      | 59600      |
| SD                | 906480      | 38437      | 48300      |
| Minimum           | 5040000     | 20000      | 10000      |
| Maximum           | 8970000     | 148000     | 188000     |
| **LEUKOCYTES (cells/µl)** |         |            |            |
| Mean              | 12360       | 2740       | 1665       |
| SD                | 6080        | 1056       | 1154       |
| Minimum           | 6100        | 1250       | 550        |
| Maximum           | 33100       | 4850       | 4350       |
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There were improvements in bone and soft tissue healing processes with the use of L-PRP (YUAN et al., 2012). The PRP with a high leukocyte concentration associated with experimental fracture of the tibia in dogs resulted in a faster and more efficient repair when compared to the control group without PRP (BARBOSA et al., 2008). Therefore, PRP obtained by protocol A would be the most appropriate in fracture cases. Some studies cite the protective effect PRP against infections due to leukocytes (MARX & GARG, 1999; BIELECKI et al., 2007).

A previous study showed the presence of white blood cells, particularly concentrated in PRP, predominantly lymphocytes (UBEZIO et al., 2014). They showed that the total biomodulator content, such as GF, in autologous plasma rich in platelets is not influenced by the lymphocyte status (UBEZIO et al., 2014). A longer time in centrifugation slightly increased platelet recovery and decreased leukocyte concentrations in the upper layer. Therefore, time can be a control parameter when low levels of white blood cells, such as lymphocytes, are required in the PRP sample (PEREZ et al., 2014).

**CONCLUSION**

Both protocols concentrated platelets efficiently, with no difference between platelet

### Table 2 - Statistical summary of the percentage of platelets, mean percentage of activated platelets, percentage of red blood cells, and percentage of leukocytes in both protocols (PA and PB) to obtain platelet-rich plasma from 20 dogs.

|                         | Protocol A         | Protocol B         | Protocol A         | Protocol B         | Protocol A         | Protocol B         | Protocol A         | Protocol B         |
|-------------------------|--------------------|--------------------|--------------------|--------------------|--------------------|--------------------|--------------------|--------------------|
| Mean ± SD               | 442.13 ± 161.93    | 385.70 ± 155.05b   | 26.55 ± 6.72c      | 26.25 ± 7.03c      | 1.04 ± 0.60d       | 0.90 ± 0.72d       | 26.41 ± 16.10e     | 15.25 ± 11.84f     |
| Minimum                 | 205.25             | 164.53             | 11                 | 13                 | 0.30               | 0.15               | 10.27              | 5.13               |
| Maximum                 | 918.64             | 893.22             | 39                 | 57                 | 2.42               | 3.07               | 72.13              | 45.90              |
| p-value                 | 0.89               | 0.89               | 0.48               | 0.01               |                    |                    |                    |                    |

Data on the same line, followed by different letters indicate a statistical difference with 95% confidence.

*Mean values found after individually correlating the value in whole blood in relation to protocols A and B for each animal.

Anabolic changes, increased the genetic expression of anabolic genes, smooth muscle actin, and type I and III collagen (ZHOU et al., 2015). Only the use of P-PRP in intra-articular injections was indicated as safe to avoid catabolism and failure in tissue recovery (MATTEO et al., 2016).
activations. However, only PA reached a mean concentration higher than 1 million platelets/μL.

PA resulted in PRP with a higher leukocyte concentration than PB. Both protocols concentrated more lymphocytes in relation to other leukocyte types. PA yielded PRP with higher concentrations of basophils, neutrophils, and lymphocytes than PB.

There was no difference in the red blood cell concentration obtained between the two protocols.

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Table 3 - Mean values of the absolute specific leukocyte count using the two protocols to obtain platelet-rich plasma from 20 dogs.

| SPECIFIC LEUKOCYTE COUNT (cells/μL) | Protocol A | Protocol B |
|-------------------------------------|------------|------------|
|                                     | Mean ± SD  | Mean ± SD  |
| Basophil                            | 8.15 ± 15.04a | 0 ± 0b  |
| Minimum                             | 0          | 0          |
| Maximum                             | 45         | 0          |
| p-value                             | -0.02      | -0.97      |
| Eosinophil                          | 19.12 ± 42.74a | 18.65 ± 42.66a |
| Minimum                             | 0          | 0          |
| Maximum                             | 176        | 168        |
| p-value                             | -0.97      | -0.39      |
| Banded neutrophil                   | 79.45 ± 38.79a | 64.15 ± 69.15a |
| Minimum                             | 42         | 5.5        |
| Maximum                             | 178        | 300        |
| p-value                             | -0.39      | -0.001     |
| Segmented neutrophil                | 410.25 ± 143.44a | 244.25 ± 162,83b |
| Minimum                             | 196        | 83         |
| Maximum                             | 780        | 713        |
| p-value                             | -0.001     | -0.005     |
| Lymphocyte                          | 2214.72 ± 922.92a | 1340,2 ± 949,58b |
| Minimum                             | 988        | 451        |
| Maximum                             | 4074       | 3785       |
| p-value                             | -0.005     |            |
| Monocyte                            | 2.22 ± 9.95a | 0 ± 0a   |
| Minimum                             | 0          | 0          |
| Maximum                             | 45         | 0          |
| p-value                             | -0.32      |            |

Data on the same line, followed by different letters indicate a statistical difference with 95% confidence.

BIOETHICS AND BIOSecurity COMMITTEE APPROVAL

This study was approved by the Committee on Animal Research and Ethics of the Federal Rural University of Rio de Janeiro (CEUA-IV, UFRRJ), under protocol no. 1827171016. All animal owners signed a free and informed consent form.

DECLARATION OF CONFLICT OF INTERESTS

The authors declare no conflict of interest. The founding sponsors had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript; and in the decision to publish the results.
AUTHORS' CONTRIBUTIONS

The authors contributed equally to the manuscript.

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