RELATIONSHIPS OF MEAN CORPUSCULAR VOLUME WITH DIAMETER AND SURFACE AREA OF CANINE ERYTHROCYTES

Indira Silva1, K. Thananjayan1, DRA Dissanayake1, WCR Fernando2 and M. Murugananthan3

1 Veterinary Teaching Hospital, University of Peradeniya, Sri Lanka 
2 Department of Animal Production and Health, Sri Lanka 
3 Department of Parasitology, Faculty of Medicine, University of Jaffna, Sri Lanka

SUMMARY: Assessment of erythrocyte morphology is an important aid in diagnostic haematology. Anisocytosis which is changes in RBC size, correspond to changes in diameter and surface area (SA) of the cell, and are not always reflected in the Meand Corpuscular volume (MCV). This paper discusses the relationship of MCV with surface area and diameters (vertical and horizontal) in 2D views of erythrocytes of clinically healthy dogs using confocal microscopy. This information would be valuable for early detection of cellular changes in dogs in subclinical or clinical diseases.

The average diameters, average SA, and MCV values of RBCs studied were in normal distribution and the mean values of 7.169 ± 0.648 µm for horizontal diameter (D1), 7.1245 ± 0.6646 µm for vertical diameter (D2), and 41.061 µm2 ± 6.866 for SA did not reveal a significant correlation or a strength of association with the MCV values, indicating that the MCV value has limitations as an objective measurement of detecting anisocytosis.

INTRODUCTION

Examination of blood is virtually the universal first step in the evaluation of health and disease in human and animal patients as physiological and pathological changes in most tissues are reflected directly or indirectly in blood. Erythrocyte morphology is routinely assessed microscopically and is an important aid in diagnostic haematology (Thrall, 2006; Jain, 1993; Proctor et al., 1976). The morphology of red cells is categorized according to colour, size, shape, intra or epicellular structures, and their arrangement on blood films, which are unique for each animal species (Ford, 2013; Thrall, 2006).

The typical 3D biconcave disk shape of RBCs in animal species show very little variability in size and shape between cells (Aditi et al., 2017). Anisocytosis is defined as size variations resulting from microcytes, macrocytes or both (Proctor et al., 1976). Parameters of red cell volumes, such as, Mean Corpuscular Volume (MCV), Red Cell Distribution Width (RDW) and/or RDW-SD are tools generally used to quantify anisocytosis. Changes in RBC size correspond to changes in cell diameter and not necessarily to changes in RBC volume, and changes in cell volumes are not always reflected in the MCV. Furthermore, the surface area and volume changes in disease do not occur equally in all RBCs in a sample (Jay and Rowlands, 1975). Hypochromic microcytic erythrocytes in iron-deficiency anemia show decreased MCV when determined electronically, but show normal diameter (Thrall, 2006).

Changes in diameter may not be reflected in the MCV obtained as a direct measurement from automated hematology analyzers, which is the average volume of RBCs in a mixture of cells (Weiss and Wardrop, 2010), and does not indicate individual changes in RBC sizes. Therefore, it is important and timely to study morphometry of RBCs, such as, the relationship between the diameter, the surface area (SA) and the MCV of individual RBCs. Several techniques have been proposed for morphological studies of biological specimens, such as optical coherent tomography and confocal microscopy. The objective of this study was to analyze the relationship of MCV with SA and diameters in 2D views of RBCs of clinically healthy dogs using...
confocal microscopy. The information presented would be valuable for early detection of cellular changes in dogs with subclinical or clinical diseases.

**MATERIALS AND METHODS**

Dogs presented to Veterinary Teaching Hospital (VTH), Peradeniya with normal Full Blood Counts (FBC) analyzed using a fully automated veterinary hematology analyzer (MS9-5V, MELET Schloesing laboratories, France) were selected for this study. A volume of 1ml blood was collected to K<sub>3</sub>EDTA coated tubes with the consent of the owners. The anticoagulant K<sub>3</sub>EDTA is recommended by ICSH (International Council for Standardization in Hematology) and CLSI (Clinical and Laboratory Standards Institute) as the choice of anticoagulant for blood cell counting.

Blood smears were prepared from each sample, stained with Leishman stain and transported to the Parasitology laboratory, Faculty of Medicine, University of Jaffna to study morphometry of RBC in 2D view using a Laser scanning confocal microscope (Carl Zeiss LSM 700, Germany). Up to 500 RBCs in the middle one third of the monolayers of each slide were randomly examined by cross sectional method under oil immersion (×100) using the above microscope. A laser beam (405 nm - 639 nm wavelength) was passed through the confocal optical path, and images were obtained with Zeiss Axio Imager Z2 (high resolving and high sensitive cameras) on the monitor at 2560 × 1920 resolution to generate the 2-dimensional object information of the RBCs (Cortada, 2013; Bernhard, 2012 in LSM 700 Operating Manual Black edition 2012). The digital images were acquired at 2560 × 1920 resolution using a Laser scanning confocal microscope (Carl Zeiss LSM 700, Germany) to measure perpendicular diameters (horizontal, D1 and vertical, D2) and surface area (SA) of each RBC in micrometer using the ZEN lite (blue edition) software from Carl Zeiss. The entire system was controlled through a standard high-end Personnel Computer (PC) with Windows operating system. The approximate perpendicular diameters (horizontal, D1 and vertical, D2) and surface area (SA) of each RBC was measured in micrometers up to 3 decimals using the ZEN lite (blue edition) software from Carl Zeiss. These values were analyzed with the MCV values in the FBC report.

**Data analysis**

Anderson Darling normality test (ADNT) was performed using 'MINITAB' software to analyze whether the data obtained for MCV, diameters and SA of RBCs follows a normal distribution (Ghasemi and Zahediasl, 2012). Mean diameter, standard deviation and variance, minimum, maximum data for D1, D2 and SA of 500 RBC per sample were calculated using the 'MINITAB' software. The Pearson Correlation Coefficient test was performed using the same software to assess the strength and direction of association of MCV values with diameters (D1, D2, average diameters) and SA individually. The Linear regression test was performed to predict the value of the dependent variable (MCV) based on the value of the independent variable (SA, D1 and D2). Box plots were created separately to visualize the pattern of association between the MCV, D1, D2 and SA using the Box plot R software.

**RESULTS AND DISCUSSION**

Morphometry provides higher reliability and reproducibility of cytological and histopathological diagnoses in human medicine (Albertini et al., 2003). This paper compares the relationship of MCV with surface area (SA) of erythrocytes and diameters in 2D views of RBCs. The Full Blood Count (FBC) on blood collected from 14 apparently healthy dogs (9 females, 5 males) of different breeds (German shepherd, Lion shepherd, and Rottweiler, Doberman and Cross breeds), with varying ages (4 months to 13 years) were measured within 4 hours of collection.

The perpendicular diameters (horizontal, D1 and vertical, D2) and surface area (SA) of each RBC in micrometer up to 3 decimals using the ZEN lite (blue edition) software from Carl Zeiss are shown in Figure 1. The above software allowed accurate morphometric measurements, similar to other software, such as OPTIKATM Vision Pro software (Adili et al, 2017), than the conventional method of measuring the diameters of erythrocytes in a blood film using a scale on an oil immersion lens in a light microscope. The use of software reduces errors due to the human factor that is involved in studies with ocular micrometer.

The reference values of MCV for dogs in Sri Lanka range between 52-76 fl and 60-77 fl (Silva and Mallawa, 2010). The Anderson Darling normality test revealed that the average diameter, average SA, and MCV values of RBCs were in normal distribution (P=0.075, 0.198, and 0.135, respectively). The mean diameters of 7.169 (± 0.648 µm) for D1, 7.1245 (± 0.6646 µm) for D2, and 41.061 (± 6.866) for SA calculated were similar to the ZEN lite (blue edition) software from Carl Zeiss. The reference values of MCV for dogs in Sri Lanka range between 52-76 fl and 60-77 fl (Silva and Mallawa, 2010). The Anderson Darling normality test revealed that the average diameter, average SA, and MCV values of RBCs were in normal distribution (P=0.075, 0.198, and 0.135, respectively). The mean diameters of 7.169 (± 0.648 µm) for D1, 7.1245 (± 0.6646 µm) for D2, and 41.061 (± 6.866) for SA calculated were similar to values obtained by Adili et al., (2017) for diameter (6.29-7.77 µm) and surface area (25.55-47.05 µm) of RBCs of Sloughi and German Shepherd dogs.

The Pearson Correlation coefficient did not reveal significant relationships between MCV and SA (r= -0.023, p=0.140), MCV and D1 (r=0.011, p=0.140), or between MCV and D2 (r=0.05, p=0.001). Linear regression analysis between MCV (dependent variable) and SA did not reveal a significant correlation (r=0.011, p=0.140), or a strength of association (P=0.474, R-sq. -0.1%). A significant linear relationship or significant strength of association was not evident between MCV and D1(r=0.474, R-sq. -0.0%) or between MCV and D2 (p=0.001, R-sq.0.1%). The box plots also did not reveal a pattern of association between the MCV and SA (Figure 2) or between MCV and D1 (Figure 3) or D2 (Figure 4). Adili et al, 2017 also have not seen significant differences between the diameter and the SA of RBCs of dogs.

Anisocytosis in a blood sample, depicted by parameters of red cell volumes, such as, MCV and RDW do not complement the changes in RBC size in relation to diameter or surface area (Proctor et al, 1976). Our results
show that diameters and surface areas of RBCs do not correspond with the MCV values calculated by the automated analyzer, indicating that the MCV value has limitations as an objective measurement of detecting anisocytosis. Practitioners may interpret changes in test results as significant, when in reality, they are close to reference range, and they also should be able to recognize changes in a patient's test results that are due to random error before the test results are reported (Houwen, 1989). Electronic hematology analyzers are
stable and accurate, however, random errors can be caused by preanalytical and postanalytical problems, rather than by instrument or reagent problems, resulting in changes in one or more parameters tested which cannot be adequately explained by changes in the patient’s clinical condition (Houwen, 1989). Preanalytical errors include specimen collection, improper specimen handling and labeling, or misidentification of the patient, while postanalytical errors include failure to correct test values for dilution, transcription errors, and misinterpretation of test results (Houwen, 1989). Transcription errors in automated hemoanalyzers can occur at a rate of approximately 1 per 500 samples or more, while the frequency of random errors in most laboratories exceeds 1 per 200 to 500 samples (Houwen, 1989). Therefore, a blood smear must be examined if and when an automated instrument produces highly improbable results which may be factitious (Bain, 2005). Future morphometric studies are warranted for further investigations on the relationship between these parameters.

CONCLUSION

Morphometry using a Laser scanning confocal microscope and ZEN lite (blue edition) software from Carl Zeiss showed that diameters and surface area of canine erythrocytes did not correspond with the MCV values calculated by the automated analyzer, indicating that the MCV value has limitations as an objective measurement of detecting anisocytosis.

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Figure 4. Variation of vertical diameter of red blood cells with MCV