Immunophenotypic characterization and depletion of pulmonary intravascular macrophages of horses

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Abstract – Pulmonary intravascular macrophages (PIMs) are present in horses and are believed to increase their sensitivity to endotoxin-induced cardio-pulmonary shock. However, owing to a lack of a marker for PIMs and the inability to isolate them, their precise contributions in the horse remain unknown. We designed this study to identify an immuno-phenotypic marker for PIMs and to develop a protocol for their transient depletion with gadolinium chloride (GC). GC is a lanthanide that has been used to deplete liver and lung macrophages. The horses (N = 15) were divided into control (n = 5) and GC-treated (n = 10) groups and the lung samples were examined by routine and immunocytochemical light and electron microscopy. GC-treated horses were euthanized at 48 h (n = 6) and 72 h (n = 4) post-treatment. The PIMs reacted with MAC-387 but not with ED-1 and CD-68 anti-macrophage antibodies. GC reduced the number of PIMs in horses at 48 and 72 h compared with the control (p < 0.05). There were increased intravascular TUNEL-positive cells in GC-treated horses and electron microscopy showed apoptotic PIMs in these horses. These data show that MAC-387 is a reliable marker for PIMs and GC is a safe tool to reduce the number of PIMs.

PIMs / gadolinium / MAC-387 / apoptosis / immunohistology / immuno-electron microscopy / TUNEL

1. INTRODUCTION

Macrophages are widely distributed in the body and display structural and functional heterogeneity [13]. In the lungs, macrophages occur in the airways, alveolar spaces, interstitium, and microvessels. Pulmonary intravascular macrophages (PIMs) were recently identified as resident mononuclear phagocytes in cattle, sheep, goats, pigs and horses [4, 5, 7, 29, 35]. PIMs differentiate from monocytes in post-natal life and form firm adhesions with the capillary endothelium [15, 36–38]. These cells phagocytize blood-born tracer particles, bacteria and endotoxins [3, 18, 27, 29, 36]. PIM-containing species show pulmonary hypertension following the administration of tracer particles and endotoxins, and the removal of PIMs suppresses these responses
Neonatal lambs that do not have PIMs show diminished pulmonary vascular response to tracers compared with four-week old lambs that have PIMs [15]. Blecha et al. showed that porcine PIMs secrete thromboxane and are cytolytic [7, 8]. They also reported that the immunological characteristics of PIMs are similar to those of alveolar macrophages [8]. There is a putative association between the presence of PIMs and the susceptibility to acute lung disease [32, 35].

Equine PIMs show structural and functional similarities to their counterparts in other species [5, 29]. Halothane exposure activates equine PIMs as indicated by hyperplastic Golgi complexes and the development of lamellipods [2]. Horses show increases in respiration rate, heart rate and pulmonary arterial pressures following infusions of tracer particles or endotoxins [29]. Endotoxin-provoked increases in mean pulmonary arterial pressure are abolished by the pre-treatment of horses with a detergent called Tyloxapol [16, 18]. Although it is believed that the detergent blocks the PIMs, the mechanisms of these effects remain unknown [17].

Acute lung inflammation is a critical factor in colic-associated equine deaths. These deaths cause serious suffering and substantial financial loss to the equine industry [20]. There is a suggestion that PIMs may play a critical role in acute lung inflammation in horses [17]. However, the functions of PIMs remain unknown largely for technical reasons. First, most attempts to isolate PIMs for in vitro investigations have failed because of their intimate attachment to the lung microvascular endothelium [26]. Second, there is a lack of a phenotypic marker to identify PIMs in situ. Third, it is difficult to attribute particular functions to an individual immune cell in vivo because many cells may contribute to an observed response. Now, we report data on the in situ phenotypic characterization of equine PIMs and a method to reduce their numbers in vivo.

2. MATERIALS AND METHODS

2.1. Animals

The experimental protocols for this study were approved by the Animal Care Committee of the University of Saskatchewan and the work was carried out according to the guidelines of the Canadian Council on Animal Care. Horses (N = 15) of 2–4 years of age, purchased from the local market, were acclimatized to the changed housing for over a week and were regularly physically examined. The horses were randomly divided into two groups: The control (n = 5) and the gadolinium chloride (GC, Sigma Co., USA) treated group (n = 10). GC is a heavy metal lanthanide that has previously been used to remove liver and lung macrophages including the PIMs [25, 28, 39]. In vitro studies show that GC induces apoptosis in macrophages [19].

2.2. Pressure measurements

The day before study, horses were sedated with xylazine hydrochloride (Rompun, Bayer Inc., Canada, 0.5 mg/kg body weight, IV) and under local anesthesia an 8 Fr catheter introducer (Sheath Dilator, Arrow International Inc., USA) was aseptically positioned and secured within the right jugular vein. Next a 7 Fr, 110 cm thermistor tipped, triple lumen pulmonary artery balloon catheter (Edwards Swan-Ganz Catheter, Baxter Corp., Canada) was positioned so that its tip was in the pulmonary artery. The catheter location was confirmed by observation of characteristic blood pressure waveforms. The systolic, mean and diastolic pulmonary arterial blood pressures were measured using a silicon chip strain gauge transducer (Truwave Disposable Pressure Transducer, Baxter Corp., USA), and displayed on a physiological monitor (Propaq 104, Protocol Systems Inc., Canada). The transducer was zeroed to a point approximately level with the
right atrium of the horse and was previously calibrated using a mercury manometer. Blood samples were collected for differential white cell counts.

2.3. Treatments

Control horses (n = 5) were administered 500 mL of physiological saline intravenously for 30 min. Treated horses were given GC (Sigma Co., USA) intravenously at the dose of 10 mg/kg body weight in 500 mL of physiological saline solution over a 30 min period. Pulmonary arterial pressures were recorded three times: immediately before the infusions, after 50% and 100% of the infusions. Horses treated with GC were euthanized with pentobarbital sodium (Euthenol®, Bimeda-MTS Animal Health Inc., Canada, 10 mL/50 kg body weight intravenously) at 48 h (n = 6) and 72 h (n = 4) after treatment while the control horses were euthanized at 48 h.

2.4. Tissue fixation

Following euthanasia, the horses were hoisted off the floor with chains in the necropsy room and the tracheae were exteriorized and opened. Five liters of fixative (2% paraformaldehyde and 0.1% glutaraldehyde in 0.1 M sodium cacodylate buffer; pH 7.2) were poured into the lungs via the tracheae and left in place for 30 min. Then, the lungs were removed en bloc and divided into pre-determined imaginary slices to collect seven pieces (2 cm³) from each lung.

2.5. Tissue processing

2.5.1. Light microscopy

The tissues were fixed in 4% paraformaldehyde for 24 h. These tissues were dehydrated in ascending concentrations of ethanol and xylene and embedded in paraffin.

2.5.2. Routine electron microscopy

Lung tissues were processed for EM as described previously [5]. Briefly, the tissues were fixed in 2% paraformaldehyde and 2.5% glutaraldehyde in 0.1 M sodium cacodylate buffer (pH 7.2) for 3 h and post-fixed for 90 min in 1.5% osmium tetroxide. The tissues were stained en bloc with 0.5% tannic acid in 0.1 M HCl-sodium cacodylate for 30 min, dehydrated and embedded in Jembed 812 resin (J.B. EM Services Inc., USA).

2.5.3. Immuno-electron microscopy

The lung pieces were fixed in 2% paraformaldehyde and 0.1% glutaraldehyde in 0.1 M sodium cacodylate buffer (pH 7.2) for 3 h at 4 °C but were not treated with osmic acid. After dehydration, the tissues were infiltrated with LRWhite (J.B. EM Services Inc., USA) and polymerized in ultraviolet light at –8 °C for 48 h.

2.6. Tissue analyses

2.6.1. Histopathology

5–7 µm sections were cut from all the blocks (7/lung) and stained with hematoxylin and eosin.

2.6.2. Immunohistology with ED-1, MAC-387 and CD68 antibodies

5–7 µm thick sections were prepared from the paraffin-embedded lung tissues and placed on silane-coated glass slides. After removal of wax with xylene and rehydration in an ethanol series, the sections were treated with pepsin (2 mg/mL 0.01 N HCl) for 90 min for antigen retrieval. The sections were incubated in 5% hydrogen peroxide for 20 min to quench endogenous peroxidase and with 1% bovine serum albumin for 30 min to block non-specific binding. We tested three anti-macrophage antibodies. ED-1 (Serotec Ltd., UK) is raised in the mouse and reacts with a 90–100 kDa protein expressed on lysosomal membranes of rat macrophages and has been used to
recognize them in flow cytometry and in routinely fixed tissues [24]. Mouse anti-human monocyte/macrophage antibody (MAC-387, Serotec Ltd., UK) recognizes the L1 cytoplasmic molecule and has been used in immunohistochemistry [11, 12]. CD68 (DAKO A/S, Denmark) is a mouse anti-human macrophage antibody that reacts with a 110 kDa glycoprotein in tissue macrophages [1]. The sections were incubated with anti-human (MAC-387 and CD68, 1:75 each) and anti-rat macrophage (ED-1, 1:75) antibodies for 2 h followed by anti-mouse peroxidase-conjugated secondary antibodies (1:100, DAKO A/S Denmark) for 60 min. The controls included staining the lung sections with only secondary antibody or with rabbit anti-human von-Willebrand Factor antibody (vWF; DAKO A/S, Denmark) which reacts with vWF stored in Weibel-Palade bodies in endothelial cells. This antibody is routinely used to delineate vasculature and as a control in immunostaining protocols [30]. The color was developed with a commercial peroxidase histochemical kit (Vector Laboratories, Canada). Positive cells were counted in 20 oil immersion (×100) randomly chosen fields in one section from each of the blocks from each horse.

2.6.3. Routine electron microscopy

One-micrometer toluidene blue-stained sections were prepared to select the area of interest for ultra-thin microtomy. The sections (80–100 nm) were stained with uranyl acetate and lead citrate, and examined in an electron microscope at 80 kV.

2.6.4. Immuno-electron microscopy

The sections (100 nm) were blocked with 1% normal goat serum followed by incubation with MAC-387 antibody (1:50) and 10 nm-gold conjugated goat antimouse secondary antibody (1:100, British Biocell International, United Kingdom). Control sections were stained without the primary antibody and with rabbit anti-vWF antibody. The sections were stained with uranyl acetate and lead citrate and examined in an electron microscope at 80 kV.

2.6.5. Staining for apoptosis

Lung sections from all the horses were stained using a terminal deoxytransferase-mediated dUTP nick-end labeling (TUNEL) kit (Roche Inc., USA) to detect apoptotic cells. This method is based on the specific binding of terminal deoxynucleotidyl transferase (TdT) to the 3-OH ends of DNA. Sections were nick-end labeled with biotinylated dUTP and TdT, and stained with avidin-conjugated peroxidase. The control sections were processed without the inclusion of the TdT buffer.

2.7. Statistical analyses

Differences in cell counts between the groups were examined by ranking the variables and then performing a one-way analysis of variance (ANOVA) on the ranks. When significant differences were detected, the mean ranks were compared using the Tukey test. Each of the pressure readings (diastolic, mean and systolic) were summed over the three measurement times and the differences in the sums of the pressures were examined using the Wilcoxin Rank Sum test. Probabilities of less than 5% (p < 0.05) were considered significant. The statistical analyses were carried out using a commercial software program (Statistix for Windows, version 7, Analytical Software, Box 12185, Tallahassee, FL 323317-2185, USA).

3. RESULTS

3.1. Hematology

There were no differences in differential blood cell counts between the groups (data not shown).
3.2. Physiological response to GC injection

There were no differences in systolic, diastolic and mean pulmonary arterial pressures between the control and GC-treated horses (data not shown).

3.3. Histopathology

Examination of the sections stained with hematoxylin and eosin revealed no signs of lung inflammation in the control as well as GC-treated horses (Fig. 1).

3.4. Immunohistology

Serial sections were prepared from the same block for immuno-staining. Lung sections from the control or GC-treated horses did not yield staining when incubated with only secondary antibody (Fig. 2A). However, anti-vWF antibody used as a control delineated blood vessels and some blood cells (Fig. 2B). Although we used three anti-macrophage antibodies (ED-1, MAC-387 and CD68, Figs. 2C-2E), only MAC-387 antibody stained large septal cells resembling macrophages in an intense and specific manner.

3.5. Immuno-electron microscopy

Lung sections stained with the MAC-387 antibody and gold-conjugated secondary antibodies showed specific labeling in PIMs but not in microvascular endothelial cells or other vascular cells (Fig. 3A). However, the vWF antibody used as a control
stained Weibel-Palade bodies in endothelial cells (Fig. 3B).

3.6. Effect of GC on equine PIMs

The ability of MAC-387 antibody to recognize PIMs provided us with a tool to precisely quantify the effects of GC on these cells. Horses given a single intravenous injection of GC had lower MAC-387 cell counts at 48 and 72 h post-treatment (Figs. 4A and 4B). Horses sampled 72 h after treatment had a median cell count of 2

Figure 2. This figure shows light micrographs taken from consecutive lung sections from a control horse. Note the lack of staining following exposure to only secondary antibody (2A) or antibodies against ED-1 (2C) or CD68 (2D). The positive reaction (single arrows) is observed with vWF (2B) and MAC-387 (2E) antibodies. Double arrows: bronchiolar epithelium. ×200.
that is significantly less than the median of 9.4 cells observed in the control animals ($p = 0.02$) (Fig. 4C).

Since GC activates apoptosis in macrophages in vitro, TUNEL was used to investigate if it induces a similar phenomenon in vivo. There was a trend suggesting an increase in TUNEL-reactive cells at 48 h and 72 h post-GC treatment compared to the controls (Figs. 5A and 5B). When

**Figure 3.** An electron micrograph (3A) from a lung section stained with MAC-387 antibody shows labeling (single arrows) in a PIM while vWF antibody (3B) reacts with Weibel-Palade bodies (single arrows) in an endothelial cell (E). AS: alveolar space. $\times$37,500.
compared with the controls, the median number of reactive cells in the group sampled 72 h after treatment appeared to be increased but this difference just failed to reach statistic significance \( p = 0.07 \) (Fig. 5C). EM confirmed the presence of many apoptotic PIMs in GC-treated horses (Fig. 6). Unlike normal PIMs, the apoptotic PIMs contained apoptotic bodies and condensed nuclear material as typical ultrastructural features of apoptosis (Fig. 6B).

4. DISCUSSION

Macrophages differentiate from monocytes upon migration out of capillaries and, therefore, generally reside in the
Figure 5. A light micrograph from a lung section from a control horse (5A) shows an occasional TUNEL-positive septal cell (single arrow). Many TUNEL-positive cells (single arrows) in alveolar septae are present in a light micrograph (5B) taken from a section from a GC-treated horse. 5C shows median TUNEL cell counts in 5 control horses (0 h post-treatment), 6 horses (48 h post-treatment) and 4 horses (72 h post-treatment). Boxes enclose the 25th to 75th percentile for each group and are bisected at the value for the median. The whiskers indicate the range of “typical” data values and 0 indicates a probable outlier. Figures A-B: ×200.
Figure 6. An electron micrograph from a lung section of a control horse (6A) shows a normal PIM with nucleus (N), lysosomes (L), mitochondria (m) and a centriole (arrowhead). The micrograph (6B) taken from a lung section from a GC-treated horse shows an apoptotic PIM with many membrane-bound (single arrows) apoptotic bodies (A) and a remnant of a nucleus (N). AS: alveolar space; E: endothelial cell. ×20000.
Equine PIMs

extravascular compartment [13]. There are examples, however, of intravascular macrophages such as those in the liver and the lung [32]. Although liver intravascular macrophages are present in all the known mammalian species, the PIMs are restricted to only a few of the species such as cattle, horses, pigs, sheep and goats [5, 32]. Because the presence or absence of PIMs influences the response of the lung to disease, it is important to study them. Many factors, including a lack of an immuno-phenotypic marker and an inability to isolate equine PIMs for in vitro studies, have hampered the investigation of these cells. Here we report that MAC-387 is a reliable marker for the in situ identification of PIMs and that GC significantly reduces the numbers of PIMs.

Three monoclonal antibodies that identify mononuclear phagocytes in rats and humans have been tested for their reactivity with equine PIMs. Of these, ED-1 and CD68 recognize a 90–110 kDa protein in the lysosomes of rat and human macrophages, respectively, and have been used for immunohistochemistry in the respective species [1, 24]. MAC-387 identifies another cytoplasmic, most probably lysosomal, protein called L1 in the lung and liver macrophages in humans [11, 12]. These antibodies also show some reactivity against monocytes but very weak reaction with peripheral blood granulocytes. Bovine macrophages including PIMs are stained with CD68 antibody [1, 22]. However, equine PIMs and alveolar macrophages did not stain with either CD68 or ED-1. It is possible that further technical refinements may reveal the usefulness of ED-1 and CD68 to identify lung macrophages including PIMs in horses. However, in our hands, only MAC-387 reacted with equine PIMs, which compared to ED-1, is believed to recognize mature tissue macrophages such as PIMs. Immuno-EM confirmed the localization of MAC-387 in the cytoplasm of equine PIMs. Because we have not tested MAC-387 in other organs such as the liver, our data can only be used for the in situ identification of PIMs.

Many other investigators have also used GC to remove alveolar and liver macrophages [6, 23, 28]. It was reported previously that sheep and calves treated with GC show losses of PIMs [22, 28]. Now GC infusion in horses caused a decline in PIM numbers at 48 and 72 h. Even though the differences between 48 h and 72 h post-GC treatment were not significant, the data suggest a steady decline over this period of time. This response was slightly different from that observed in cattle and sheep where an increase was noticed in PIM numbers between 48 h and 72 h after GC treatment [22, 28]. These inter-species differences in the kinetics of PIM depletion notwithstanding, GC treatment offers us a large possibility for the investigation of lung pathophysiology in the absence of PIMs and the comparison with normal horses.

Gadolinium chloride affects macrophages because its phagocytosis is a prerequisite for its cellular actions [19]. For example, GC administered intratracheally or intravenously reduces the number of alveolar macrophages and Kupffer cells, respectively, but has no effect on endothelial or epithelial cells in rats [6, 21, 25, 34]. Single intravenous treatment with the GC solution primarily affects PIMs in sheep and cattle while sparing Kupffer cells and lung microvascular endothelial cells [28]. This targeted depletion of PIMs may be due to the preferential uptake of GC during its “first pass” through the lung microvasculature by these cells. Such a possibility is supported by data that more than 90% of tracers and endotoxins following jugular infusion are phagocytosed by the PIMs [9]. Although we have not examined the effect of GC on liver macrophages in horses, we speculate that these may be unaffected as in sheep [28]. Therefore, GC treatment depletes PIMs and can then be used to investigate lung responses in their absence.

There was a trend indicating an increase, though not significant, in TUNEL-positive
cells in lung alveolar septae of GC-treated horses. Ultrastructural examination strengthening the TUNEL observations and confirmed that the apoptotic cells were PIMs. The PIMs in GC-treated horses showed classical ultrastructural features of apoptosis including apoptotic bodies and condensation of nuclei while other lung cells were unaffected [14]. These in vivo data are similar to in vitro data on GC-mediated apoptosis in alveolar macrophages reported by Mizgerd et al. [19]. Apoptosis is generally believed to be a non-inflammatory mode of cell death since membrane-bound apoptotic bodies prevent the release of hydrolytic enzymes into extra-cellular space and inflammatory damage to neighboring cells [10]. There was neither a pulmonary hypertensive response during GC infusion nor apparent lung damage in GC-treated horses. Therefore, GC appears to be a safe chemical for the removal of PIMs. Previously published data tantalizingly implicate PIMs as a major component in acute lung inflammation in the host species [9, 17, 18, 33, 35]. These methods will facilitate investigations of acute lung inflammation in horses with depleted PIMs to establish their contributions in clinical conditions such as heaves and endotoxemia.

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