Comparison of cryoprotective methods for histological examination of rat and porcine lung tissue

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Abstract

Proper histological evaluation of lung tissue and accurate antigen detection by immunodetection techniques require histological tissue processing to preserve tissue reactivity and open alveolar spaces. In this study, we focused on testing and comparing different procedures of tissue cryopreservation. Sucrose or Tissue Tek were used with several methods of freezing samples by supercooled liquids and liquid nitrogen. Changes in tissue caused during the freezing of samples and the effect of cryoprotectants on the tissue were recorded. Rat and porcine pulmonary tissues were used in this experiment. This study aimed to optimize the process of lung cryopreservation with emphasis on enabling proper anatomical evaluation and preserving a high tissue immunoreactivity. The best results were obtained by inflating pulmonary parenchyma with a 1 : 1 mixture of O.C.T. with phosphate buffered saline (PBS) frozen in supercooled n-heptane placed on dry ice. Pulmonary tissue prepared in this way enables to perform proper histological evaluation and to detect target molecules by immunohistochemical analysis.

Alveolar microstructure, cryopreservation, frozen sections, immunohistochemistry, pig

Pulmonary tissue shows a very fragile and unique structure, formed by open alveolar spaces, where air exchanges between the organism and its environment. Under physiological conditions, lungs are filled with air that maintains a typical lung morphology. Lungs removed at necropsy normally collapse due to the loss of negative pleural pressure leading to a quite unnatural appearance of both gross and histological specimens (Hausmann et al. 2004). A proper histological evaluation of lung tissue is easier to perform if the alveolar spaces of the lungs are open. The deflated pulmonary parenchyma has a wider alveolar tissue structure resulting from inflammatory infiltrate. This is because the alveoli tend to collapse, which may appear as the presence of fibrosis or inflammation, thus providing a distorted picture for the pathological examination of the lungs (Churg 1983).

The method of formalin-fixed paraffin embedded tissue (FFPE) histology provides a good preservation of lung microstructure. Perfusion of lungs with buffered formalin at necropsy is an accepted technique to improve the histology of the pulmonary architecture (Hausman et al. 2004). On the other hand, the use of formalin reduces the immunoreactivity of many antigenic structures (Gatter et al. 1984). The process of processing paraffin slides also harms the quality of nucleic acid isolation, where formalin crosslinks between proteins and nucleic acids occur (Werner et al. 2000). Long-term storage of paraffin preparations causes considerable degradation of DNA, RNA, and many antigens (Von Ahlfen et al. 2007). A gentler and faster method of histological preparation is the frozen sections method (FS), which allows to leave out denaturation fixatives and keeps most of the biologically active substances in the native form, even in the long-term storage of samples. The FS provides...
ideal conditions for both immunological and molecular analyses, such as immunodetection or gene expression measurements (Gianoulis et al. 1988).

Preparing the tissue correctly is a basic step in the detection of antigenic molecules by using the immunohistochemical analysis (IHC), which serves to accurately locate, for example, inflammatory markers of immune cells during the study of respiratory diseases of pigs. At this time, porcine respiratory diseases, such as porcine respiratory and reproductive syndrome (PPRS) or pneumonia caused by actinobacillus pleuropneumoniae (APP), are currently a topic of animal research. Both of these diseases often cause great economic and breeding losses in swine breeding (Zimmerman et al. 2002). Identification of inflammatory processes in the lungs of infected animals, as well as an understanding of the interaction of the immune system with pathogens, can help to elucidate the pathogenesis and to improve protection of pigs against these diseases.

There are several established methods for the cryopreservative processing of lung tissue, and the selection of a suitable method depends on the target evaluation of preparation. After tissue sampling, specimens were portioned, and each portion handled and processed differently, to stabilize or preserve different biomolecules. Each specialized method preserves one class of biomolecules at the expense of others (Baatz et al. 2014).

Cryopreservation of the tissue is an important point in its treatment, as to avoid the formation of ice crystals arising when water, contained in the tissues, expands during solidification. Ice crystals disrupt cell membranes and cause so-called Swiss Cheese artifacts (Peters 2010). One way to prevent these problems is to treat the tissue with a suitable cryoprotectant which interferes with interactions between the polar bonds of the water molecule. Most commonly used cryoprotectants are sucrose, glycerol, and polyethylene glycol (Pegg 2007). Sucrose was used for tissue cryopreservation by many authors (Vacciana et al. 1994) in combination with formaldehyde (Hosoda et al. 2001) or 4% paraformaldehyde (Ling et al. 2009).

For proper evaluation of lung tissue it is necessary to maintain open alveolar spaces using the instillation of cryoprotective media into inner spaces of the lung. There are many different preservative options for filling lung tissue with various cryoprotectants (Baijnath et al. 2016). A former study (Rabinovitch et al. 1981) reports the filling of the pulmonary parenchyma with glutaraldehyde. In another study (Halbower et al. 1994), preservation of alveolar structure was performed by filling lungs with agarose. In a different study (Braber et al. 2010), formalin together with agarose was used for fixation of the lung microstructure in mice.

Pulmonary parenchyma cryopreservation by using sucrose and commercially produced cryoprotective polymer Tissue Tek “optimal cutting temperature” compound (O.C.T. Compound; Sakura-Finetek, Tokyo, Japan) was tested. Moreover, some methods of tissue freezing in supercooled liquids (isopentane, n-heptane), in liquid nitrogen or freezing in a cryostat were sampled and compared. The aim of this study was the preservation of alveolar microstructure of lungs and maintaining high tissue immunoreactivity. Anticipated results were to minimize the compromise between selection of molecular or protein analysis and anatomical evaluation, as well as to provide a simple and reproducible method of tissue preparation.

**Materials and Methods**

**Experimental material**

The experiment was performed in compliance with the law of the Czech Republic (Act No. 501/2020 Coll.) for the protection of animals against cruelty and with the agreement of the Branch Commission for Animal Welfare of the Ministry of Agriculture of the Czech Republic (approval no. 31674/2018-MZE-17214).

Eight lungs of laboratory rats (*Rattus norvegicus*) were used for this experiment. Rats were anaesthetized with ether and sacrificed by cranio-cervical dislocation. After dissection, isolation of whole lungs was performed, the pulmonary tissue was portioned and processed by several different cryopreservation procedures. A standard
fixation of the samples by 10% formalin was omitted in order to preserve lipids and proteins in their native form.

Additionally, the lungs of two eight-week-old pigs (*Sus scrofa f. domestica*) without anomaly or vaccination were used. The pigs were kept in the accredited barrier-type animal facilities of the Veterinary Research Institute (VRI). Before being sacrificed, the pigs were allowed to acclimatize in animal facilities for one week. Experimental material was collected by trained staff from an authorized and registered slaughterhouse at VRI. Following the isolation of the whole lungs, pulmonary tissue treatment was performed immediately using optimized cryopreservative methods which were found in the rat models.

**Experimental design**

**Sampling of rat pulmonary tissues**

Cryopreservation by a 10% sucrose (S) (Ha & Ka CZ s.r.o., Mnichovo Hradiště, Czech Republic) solution were tested. Parts of the lung samples were immersed in a 10% S solution at 4 °C for 24 h overnight (O/N). Other lung samples were inserted into O.C.T and frozen. For enabling the preservation of open alveolar spaces, some lung samples were filled with 10% S solution, others by O.C.T. The cryoprotective materials were injected into airways by cannulation of the bronchi. Lungs samples were cut and tissue blocks were trimmed to the size of 5 × 5 × 5 mm and then frozen.

The freezing of samples was performed in four different ways. Samples were frozen in liquid nitrogen (LN), in supercooled isopentane (n-pentane) (n-P) (PENTA s.r.o., Praha, Czech Republic), and n-heptane (n-H) (PENTA s.r.o.) placed on dry ice or frozen in the cryomicrotome (CM) at –20 °C. Metal containers with a capacity of 250 ml were used for freezing in supercooled liquids. One of the containers was filled with 200 ml n-P, the second with 200 ml n-H. These metal containers were wholly embedded into dry ice and liquids were pre-cooled for 30 min. Samples were embedded into the vessel of an aluminum foil and some of them contained O.C.T. All samples were frozen. During freezing, samples were held beneath the surface of the liquid.

Two tissue blocks were used from each cryopreservation procedure. Samples frozen in the CM were stored at –20 °C. Samples frozen in LN and super-cooled liquids (n-P and n-H) were stored at –80 °C. Ten various combinations of tissue cryopreservation were tested in this part of the experiment.

**Variants of rat pulmonary tissue processing**

Ra1 samples were frozen in supercooled n-H without any cryoprotective treatment.

Ra2 samples were immersed in a 10% S solution O/N and frozen in supercooled n-H.

Ra3 samples were immersed in a 10% S solution O/N and frozen in the CM.

Ra4 lung inflation and immersion by a 10% S solution O/N and frozen in supercooled n-H.

Ra5 lung inflation and immersion by a 10% S solution O/N and frozen in the CM.

Ra6 samples were embedded in O.C.T. and frozen in LN.

Ra7 samples were embedded in O.C.T. and frozen in supercooled n-H.

Ra8 lung inflation with O.C.T., embedded in O.C.T. and frozen in LN.

Ra9 lung inflation with O.C.T., embedded in O.C.T. and frozen in supercooled n-P.

Ra10 lung inflation with O.C.T., embedded in O.C.T. and frozen in supercooled n-H.

**Sampling of porcine pulmonary tissues**

Porcine lung tissue was processed by only those procedures that showed the best results in rats. For cryopreservation, only O.C.T was used; the freezing of the samples was done in LN and n-H. Inflation of lungs with O.C.T. was optimized by diluting O.C.T. in phosphate buffered saline (PBS) (Bio Whittaker, Lonza, Basel, Switzerland) at a ratio of 1:1. The process of freezing and storage of porcine samples was performed in the same way as mentioned above.

**Processing of porcine lungs**

After lung isolation, caudal right and left lung lobes were selected. The first step was the trimming of the lung lobe into large blocks of approximately 5–10 × 5–10 × 5–10 cm, then cryoprotective medium was instilled into bronchi, and the tissue was cut into blocks and trimmed to the size of 5–7 × 5–7 × 5–7 mm before being frozen. Three tissue blocks from each procedure were used in this part of the experiment.

**Variants of porcine pulmonary tissues processing**

Su1 samples were embedded in O.C.T. and frozen in LN.

Su2 samples were embedded in O.C.T. and frozen in supercooled n-H.

Su3 lung inflation with pure O.C.T., embedded in O.C.T. and frozen in supercooled n-H.

Su4 lung inflation by diluted O.C.T., embedded in O.C.T. and frozen in supercooled n-H.

Su5 lung inflation by diluted O.C.T, embedded in O.C.T. and frozen in LN.

**Cryohistology**

The tissue samples were cut to a thickness of 5–10 μm on the cryostat (Leica Microsystems, CM 1900, GmbH, Wetzlar, Germany) at a temperature of –20 °C. The cuts of tissue were placed on slides, the sections allowed to dry at room temperature, fixed in pre-cooled acetone (PENTA s.r.o.) at –18 °C for 5 min and stained with Mayer...
haematoxylin (MH) (PENTA s.r.o.). A total of 140 histological slides of 4 sections each were evaluated using the light microscope Olympus BH-2 (Olympus, Tokyo, Japan) at × 100 and × 400 magnifications. Images of sections were taken with a VH-Z500R high resolution zoom lens (Keyence, Itasca, U.S.A.) mounted on the Keyence VHX-5000 digital microscope (Keyence). Two magnifications, × 500 and × 1000 were used.

Immunohistochemistry

Immunohistochemical analysis (IHC) confirms detection of antigens in the cryopreserved tissue. The endogenous peroxidase was blocked with Dual Endogenous Enzyme-Blocking Reagent (DAKO, Glostrup, Denmark) for 10 min. The slides were washed and the Protein Block (DAKO) applied for 5 min. Then, anti-CD163 (Serotec, Oxford, UK), the antibody was applied and the slides were incubated for 60 min at 37 °C in a humid chamber. The slides were washed and EnVision reagent (HRP, Mouse, DAKO) was added. The slides were incubated for 30 min at 37 °C in a humid chamber. Then they were washed and stained with DAB+ (Liquid, DAKO) for approximately 30 s. The next step was washing the slides with distilled water, lightly counterstained with MH and mounted in glycerol-gelatine. As a negative control, immunohistochemistry was performed using secondary antibodies without the primary antibody. Preservation of the lung microstructure was the primary task. Subsequent IHC staining was performed only in samples treated by the procedure with the best cryopreservative properties and for a confirmation of tissue immunoreactivity.

Results

Photomicrographic assessment of rat lung parenchyma

In the Ra1 samples (frozen in supercooled n-H without any cryoprotective treatment), the alveolar structure was not preserved. Alveoli were totally collapsed, showing atelectasis (Plate XIX, Fig. 1A, detail 1a).

In Ra2 samples (immersed in a 10% S solution O/N and frozen in supercooled n-H), a total destruction of the pulmonary parenchyma occurred. Haematoxylin precipitate was observed, complicating the histological evaluation of the sample (Fig. 1B, detail 1b).

In Ra3 samples (immersed in a 10% S solution O/N and frozen in the CM at −20 °C), the alveolar structure was not preserved. Collapsed alveoli were observed, cells and cell nuclei were not distinguishable. Swiss cheese artifacts were found (Fig. 1C, detail 1c).

In Ra4 samples (lung inflation and immersion in a 10% S solution O/N and frozen in supercooled n-H), the alveolar structure was not preserved. Collapsed alveoli were observed, cells and cell nuclei were not distinguishable. Swiss cheese artifact and air bubbles in cryoprotective medium were found (Fig. 1D, detail 1d).

In Ra5 samples (lung inflation and immersion in a 10% S solution O/N and frozen in the CM at −20 °C), a little partial contour of the alveolar structure was shown. Due to a very blurred microstructure, it was not possible to distinguish alveoli, intraalveolar septa or cell nuclei. Haematoxylin precipitate and air bubbles in cryoprotective medium were found (Fig. 1E, detail 1e).

In Ra6 samples (embedded in O.C.T. and frozen in LN), the alveolar structure was preserved. The microstructure was very blurred. Dense cellularity was observed in some areas. Some connective tissue was separated. Cell nuclei could not be observed in poorly visible cells (Plate XX, Fig. 2F, detail 2f).

In Ra7 samples (embedded in O.C.T. and frozen in supercooled n-H), poor quality of alveolar structure with blurred microstructure was observed. Alveolar walls and intraalveolar septa were wider, some alveoli were collapsed. Densely cellular areas were found. Pneumocytes were difficult to distinguish, macrophages were without visible nuclei. Haematoxylin precipitate was present (Fig. 2G, detail 2g).

In Ra8 samples (lung inflation with O.C.T., embedded in O.C.T. and frozen in LN), the alveolar structure was preserved. Unnaturally enlarged intraalveolar septa were observed. The connective tissue was separated from alveolar walls. Cells were difficult to distinguish. Contamination with unfamiliar particles was found (Fig. 2H, detail 2h).

In Ra9 samples (lung inflation with O.C.T., embedded in O.C.T. and frozen in supercooled n-P), the alveolar structure was preserved. Collapsed alveoli were observed, cells and cell nuclei could not be distinguished properly. The connective tissue was separated from
alveolar walls. Massive contamination with separated connective tissue and unfamiliar particles was found (Fig. 2I, detail 2i).

In Ra10 samples (lung inflation with O.C.T., embedded in O.C.T. and frozen by supercooled n-H) the alveolar structure was preserved. Alveoli and intraalveolar septa had a natural appearance. Some connective tissue was separated from alveolar walls. Cells in alveolar walls could be distinguished (Fig. 2J, detail 2j).

Photomicrographic assessment of porcine lung parenchyma

In Su1 samples (embedded in O.C.T. and frozen in LN), the alveolar structure was not preserved. Alveoli were collapsed, showing partial atelectasis. Massive dense cellularity was observed (Plate XXI, Fig. 3A, detail 3a).

In Su2 samples (embedded in O.C.T. and frozen in supercooled n-H) preserved alveolar structure was found. Some alveoli were collapsed. Cells and cell nuclei were visible and could be distinguished (Fig. 3B, detail 3b).

In Su3 samples (lung inflation with O.C.T., embedded in O.C.T. and frozen in supercooled n-H), preserved alveolar structure was found. Some collapsed alveoli and torn areas were observed. Visible cell and cell nuclei could be distinguished (Fig. 3C, detail 3c).

In Su4 samples (lung inflation by diluted O.C.T., embedded in O.C.T. and frozen in supercooled n-H), preserved alveolar structure was found. A natural appearance of alveoli and an optimal thickness of alveolar walls were observed. Visible cell and cell nuclei could be distinguished (Fig. 3D, detail 3d).

In Su5 samples (lung inflation by diluted O.C.T, embedded in O.C.T. and frozen in LN), the alveolar structure was preserved. Unnaturally enlarged intraalveolar septa were observed. Cells and cell nuclei were visible and could be distinguished (Fig. 3E, detail 3e).

The difference between the inflated part and collapsed part of the interalveolar septa, thickening by extensive inflammatory cell infiltration in rat and in porcine lungs is shown in Fig. 4A and B (Plate XXII).

Immunohistochemistry

IHC analysis was performed only in the best cryopreserved pulmonary tissue by the method of lung inflation by diluting O.C.T., embedding in O.C.T., and freezing in supercooled n-H. Open and well-visible alveolar spaces enabled accurate detection and space localization of the CD163 molecule in porcine lungs (Fig. 4C, D).

Discussion

This study aimed to compare methods of lung cryopreservation and furthermore, to find a simple and applicable procedure for tissue cryopreservation, and to create useful histological specimen that could be properly evaluated. Pulmonary rat and porcine tissues were used in this experiment. First, cryopreservation of the rat lung tissue was tested. The rat was selected for experimental purposes due to low cost and quick and easy acquisition of the biological material. The purpose of this experiment was to establish an adequate preservation method of the porcine lung parenchyma and subsequently, to determine the space localization of immunological markers.

The sections showed that the method of freezing samples without any cryoprotective treatment led to the atelectasis, collapsed airways, and also alveoli (Fig. 1A). This method was included for identifying and assessing changes that occurred during the freezing of untreated tissue, and for a subsequent comparison with cryopreserved tissues. According to the Bearer and Orci (1986), the tissue can be frozen without cryoprotection, but freezing must be done as quickly as possible with a vapour of liquid nitrogen.
Cryopreservation by a 10% sucrose solution was tested. By using sucrose, we assumed that the tissue would be protected from the formation of ice crystals and the tissue tension would be maintained. The effect of 10% sucrose solution was totally destructive for the lung parenchyma (Fig. 1B-E) and did not prevent the formation of ice crystals (Fig. 1C, D). Inflation with a 10% sucrose solution did not preserve the alveolar microstructure (Fig. 1D, E). In contrast to Hosoda et al. (2001) who used 20% sucrose in combination with formalin in their study, we did not achieve satisfactory results. In our experiment, formalin fixation was omitted to preserve lipids and proteins in their native form for subsequent IHC analysis.

Lung samples cryopreserved only by insertion into O.C.T. and freezing showed a poor quality of the alveolar microstructure. The alveolar walls appeared to be much wider than in their physiological state (Fig. 2F, G, B) or showed totally collapsed alveoli (Fig. 3A). We chose the method of inflation of bronchoalveolar spaces with pure embedding media O.C.T. according to Gianoulis et al. (1988) for the preservation of lung morphology. The alveolar microstructure was partially preserved during this procedure. The connective tissue was separated from the alveolar walls (Fig. 2 H–J), some parts of the microstructure were torn (Fig. 3C) probably due to the high density of the media. The density of O.C.T. was not suitable for inflation, especially in the rat lungs which are smaller and less accessible than the porcine lungs. Similarly to Myung et al. (2008), we tested the dilution of O.C.T. with PBS at a ratio of 1:1. In comparison with Prince and Porter (1975) who used a 1:2 mixture of O.C.T. and PBS injected intratracheally into fresh lung tissue, our histological sections showed that a 1:1 dilution of O.C.T. with PBS was sufficient. The specimens showed correctly preserved alveolar microstructure. Clear and sharp alveolar walls had optimal thickness, well-visible nuclei in cells, and a minimum of freezing artifacts (Fig. 3D, E). In this case, the smallest negative impact on the lung parenchyma microstructure was found. This embedding medium inflation technique may be applied to basic research. Morphology preserved in the frozen tissue may be helpful in pertinent interpretation of molecular data.

Several methods of freezing samples were tested in this experiment. Sections showed that freezing in the cryomicrotome at –20 °C produced artifacts in the form of white holes, the so called Swiss Cheese artifacts (Fig. 1C). Similarly Peters (2003) reported that the slow freezing of tissue samples in the cryostat caused numerous artifacts. Swiss Cheese artifacts appeared also in a specimen frozen in n-heptane (Fig. 1D), probably due to improperly pre-cooled freezing media and by the slow freezing process.

Better results were obtained by freezing samples in supercooled liquids isopentane and n-heptane placed on dry ice. Both procedures provided uniform freezing, adequate freezing rate, and no cracks in frozen samples. Due to numerous contaminations of the samples frozen in n-pentane (Fig. 2I), we selected only n-heptane for subsequent freezing. Similar results were obtained by Peters (2003). Faster freezing of the sample in liquid nitrogen or super-cooled liquids limits the creation of frosty artifacts but also limits the precision of isolation. Our results showed that freezing in liquid nitrogen was an aggressive method for the pulmonary tissue. The samples often cracked during the freezing and the evaluated sections had a widespread alveolar structure probably due to the boiling of the liquid nitrogen (Fig. 2H, 3E).

Some sections revealed the interalveolar septum inflated by diluted O.C.T. with a preserved alveolar microstructure in contrast to the adjacent collapsed part of the septum, where collapsed alveoli resulting in dense cellularization (Fig. 4A, B).

Pulmonary tissue processed by the inflation method with diluted O.C.T. and frozen by supercooled n-heptane showed the most suitable parameters for IHC analysis (Fig. 4C, D). In this part of the experiment, antigen CD163 was chosen as a membrane sign of myeloid cells such as monocytes/macrophages that naturally occurred in lungs. Antigen CD163 was
used to detect alveolar mononuclear phagocytes and confirmed the preservation of tissue immunoreactivity.

In summary, we have described a simple technique for lung tissue inflation by embedding medium O.C.T. diluted with PBS at a ratio of 1:1 and freezing in supercooled n-heptane placed on dry ice. This method is simple and quick to perform in comparison with other methods, providing perceptible morphological details of lung microstructure, and enabling alveoli to be held open, thus making a thorough pathological evaluation possible, with a quality comparable to paraffin sections. Tissue prepared in this way allows the detection and localization of the antigen by using the immunohistochemistry method. Optimizing cryopreservation of the pulmonary tissue will lead to a better detection of pathological processes in lungs when studying respiratory diseases of pigs.

Conflict of interest
The authors declare no conflict of interest.

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Fig. 1. Mayer haematoxylin stained frozen section of rat lungs: A - frozen by supercooled n-heptane (n-H) without any treatment by cryoprotective agent, detail (a); B - lungs immersed in a solution of 10% sucrose (S) O/N and frozen by supercooled n-H, detail (b); C - immersed in a 10% S at O/N and frozen in cryomicrotome (CM) at −20 °C, detail (c); D - instillation of a 10% S, immersed O/N and frozen by supercooled n-H, detail (d); E - instillation of 10% S, immersed O/N and frozen in the CM at −20 °C, detail (e).
Fig. 2. Mayer haematoxylin stained frozen section of rat lungs: F - embedded in optimal cutting temperature compound (O.C.T.) and frozen in liquid nitrogen (LN), detail (f); G - embedded in O.C.T. and frozen by supercooled n-heptane (n-H), detail (g); H - instillation of O.C.T. and frozen in LN, detail (h); I - instillation of O.C.T. and frozen by supercooled n-pentane (n-P), detail (i); J - instillation of O.C.T. and frozen by supercooled n-H, detail (j).
Fig. 3. Mayer haematoxylin stained frozen section of porcine lungs: A - embedded in optimal cutting temperature compound (O.C.T.) and frozen in liquid nitrogen (LN), detail (a); B - embedded in O.C.T. and frozen by supercooled n-heptane (n-H), detail (b); C - instillation of O.C.T., embedded in O.C.T. and frozen by supercooled n-H, detail (c); D - instillation of diluted O.C.T., embedded in O.C.T. and frozen in supercooled n-H, detail (d); E - instillation of diluted O.C.T., embedded in O.C.T. and frozen LN, detail (e).
Fig. 4. A - inflated interalveolar septum shows preserved alveolar microstructure in contrast to adjacent collapsed part of septum, with collapsed alveoli resulting in dense cellularization in rat lung. (B) in porcine lung. Magnification × 100. Immunohistochemical localization of the CD163 molecule on frozen section of porcine lungs inflated with a mixture of optimal cutting temperature compound (O.C.T.) and phosphate buffered saline (PBS) (1:1). C - interalveolar septum and connective tissue. D - alveoli. CD163 positive cells are stained by brown colour. Magnification × 100.