Development and maturation of the lung in fetuses of *Galea spixii* and expression of markers

Desenvolvimento e maturação do pulmão em fetos de *Galea spixii* e expressão de marcadores

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**ABSTRACT**

The aim of this research was to study the development of the lung in *Galea spixii*, by gross anatomy, histological analysis and immunohistochemical techniques. Totally, 8 fetuses were used and allocated into three groups of age: Group I (33-35 days), Group II (38-40 days), and Group III (43-45 days) with given crown-rump - CR lengths. According to the gross morphology, there were no differences in relation to the lung morphology among groups. In relation to lung maturation, the Group I showed early formation of the bronchi and bronchioles, which were richly surrounded by mesenchyme and small blood vessels, typical features of the pseudoglandular stage. Individuals from Group II showed higher amounts of tubular formations in the lung parenchyma and reduced mesenchyme, reaching the canalicular stage. The lung from individuals of the Group III was completely formed, reaching the alveolar phase. In the immunohistochemical analysis, the lung of individuals from both Groups I and II were positive for Pcna, Oct-4, and VEGF. In contrast, there was not labeling on samples from Group III. Thus, the *G. spixii* lung anatomy resembles the agouti (*Dasyprocta* sp.) as regards the number of lobes and fissures. In regard to the lung development, *G. spixii* showed more similarity with the human fetal lung, since both are born in the stage of the alveolar phase, different to other rodent species.

**Key words**: respiratory system, alveoli, morphology, rodent, experimental model.

**INTRODUCTION**

The lower respiratory tract starts its development from an endoderm epithelial bud, from the primitive gut, as a medial laryngotracheal groove in the early larynx floor, which deepens to form the respiratory diverticulum (MOORE & PERSAUD, 2008).

Lung maturation is divided into four periods: (a) pseudoglandular, (b) canalicular (c) terminal sac, and (d) alveolar phases. Alveolar
epithelial type II cells are responsible for the synthesis of pulmonary surfactant, a lipoprotein complex that contains about 30% protein and 70% lipids. The surfactant develops several important functions, but the most obvious is to reduce the surface tension of the alveoli, without which they would tend to collapse during exhalation (JUNQUEIRA & CARNEIRO, 2006). Thus, the alveolar epithelium can be considered the active metabolic unit of the lung (REECE, 2006).

The need of experimental models that may reproduce human lung diseases has led researchers to study the development of the respiratory system in different species as in murines (METZGER et al., 2008), suine (BALAZS et al., 1994), rabbit (BARONE, 1997), and non-human primates (MCCULLOUGH et al., 1978). Especially rodents have contributed significantly as experimental models for understanding various diseases (CABRAL, 2013).

The cavy (Galea spixii) shows interesting characteristics related to reproduction and embryonic development, such as a long gestation period of around 48 days (OLIVEIRA et al., 2012) and reduced time of growth (ROBERTS & PERRY, 1974) when compared to other rodents. In this way, it was studied the development and maturation of the lung in the cavy in order to describe the macroscopic and histological characteristics, including cell proliferation and the presence of pluripotent cells during fetal life.

MATERIALS AND METHODS

Eight cavy fetuses at different gestational ages obtained from Cemas – Center of Multiplication of Wild Animals, at the Universidade Federal Rural do Semi-Árido (UFERSA), Mossoró-RN, Brazil were used. Fetuses were already fixed in formalin 10% or buffered glutaraldehyde 2.5% and were used in other previous studies related to placentation (OLIVEIRA et al., 2012). The research was registered with the IBAMA (Protocol 1478912) and was approved by the Ethics Committee of the UFERSA (Protocol 23091.001975/10-24).

Biometric data and gross morphology

The specimens were measured using a caliper in order to obtain the “crown-rump - CR” (EVANS & SACK 1973). With the use of a digital scale (0.001 grams – MARTE) it the weight (g) was determined before fixation of the animals. Fetuses were allocated into 3 Groups (Group I, II, and III), according to their CR and external features. The nomenclature used was based on International Committee on Veterinary Gross Anatomical Nomenclature, International Committee on Veterinary Histological and Embryologica Nomenclatura, 2012.

Histology

Samples of lungs were dehydrated in increasing concentrations of ethanol (70 to 100%), for 1 hour in each, followed by diaphanization in xylene, for 2 hours and then, samples were embedded in paraffin (Histosecâ - MERCK, lote K91225309). Sections of 5µm were obtained in an automatic microtome (Leica, RM2165) and stained with hematoxylin and eosin - HE and Masson trichrome. The material was analyzed using a light microscope (Olympus BX40, Zeiss KS400).

Immunohistochemistry

Primary antibody were applied for PCNA (PC10, sc-56, dilution 1:100, Santa Cruz Biotechnology, Santa Cruz, CA, USA), in order to verify cellular proliferation; Oct-4 (sc-4420, dilution 1:100, Santa Cruz Biotechnology, Inc, Europe), to identify pluripotent cells, and VEGF (ab1316, diluição 1:100, Cambridge, UK) to proliferation of blood vessels. For that, it was followed a formal protocol (OLIVEIRA et al., 2012). Negative controls was performed using IgG (Goat anti-Mouse IgG - AP 308F, Chemical International).

Scanning electron microscopy (SEM)

One sample from each Group fixed in 2.5% glutaraldehyde in 0.1M buffer, pH 7.4, and postfixed in osmium tetroxide 1% (SEM®, Hatfield, Pennsylvania, PA) were dehydrated in increasing concentrations of etanol (70-100%) and dried in a critical point dryer (Balzers Union® CPD 020, Liechtenstein, Germany). Then, the material was placed in stubs and covered with gold in a sputter coater (Emitech® K550, Ashford, Kent, England), and were analyzed using a SEM (ME Leo 435 VP-Zeiss, England).

RESULTS

General characteristics

The main external features of Group I (days 33-35) were: pigmented retina, forelimb and hind limb enlarged with separated digits, rounded cephalic and occipital regions, short external ear, and short tail. Group II (days 38-40) showed: pronounced external ear and nasal region, and increased size of the whole body regions. Finally, Group III (days 43-45) showed as main characteristics the upper and lower eyelids, keratinized claws, pigmented skin, sensorial...
hair in the nasal region, and complete developed external ear (Figure 1A).

Gross morphology and scanning electron microscopy (SEM)  
There were no significant differences on lung morphology among the 3 groups. It was noted only a simple difference in the size of the whole organ size when compared the fetuses of different groups. Externally, the lungs were divided by pronounced fissures on both sides, right and left, corresponding to the divisions of the lobes. Regarding to the lobation, it was identified on the right side the following lobes: right cranial lobe, middle lobe, right caudal lobe, and accessory lobe. On the left side, the left cranial lobe was divided into cranial and caudal segments through the cranial intralobar fissure, and caudal lobe as observed in the gross morphology analysis and SEM (Figures 1B-1D).

The trachea was composed of hyaline cartilage rings and connective tissue (Figures 1D and 1E). It was observed that the primary right and left bronchi arise from the bifurcation of the trachea, immediately dorsal to the base of the heart. Then, the bronchi give rise to the lobar bronchi, which continue to branch to form the bronchial tree.

Histology and immunohistochemistry  
In the Group I, the initial development of the bronchi and bronchioles was observed in the lung parenchyma, which were surrounded by abundant fetal mesenchyme and blood vessels (Figures 2A-2C), typical characteristics of the pseudoglandular stage. The developing bronchioles had a cubic simple epithelium, with globular cells with rounded nuclei located in the basal region of the cytoplasm. These epithelial cells were supported by a layer of flat cells which maintained contact with the fetal mesenchyme and blood vessels (Figure 2C).

In Group II was observed an increased amount of the number and caliber of the bronchi and bronchioles (Figures 2D and 2E), decreased mesenchyme and the initial opening of the internal lumen of the bronchioles (Figure 2F). In Group II, the number of bronchi and bronchioles and the lung reached the canalicular stage gradually increased (Figure 2D).

Lungs of Group III were completely developed. Inside the lung parenchyma, bronchi were composed by prismatic epithelium surrounded by cartilage hyaline (Figures 2G and 2H). During the maturation of the lung, the parenchyma in this phase was composed of alveolar cells, thereby reaching the alveolar phase. With the rapid cell differentiation, a larger bronchial lumen was observed (Figure 2I), with early development of alveoli.

Figure 1 - Gross morphology of the fetuses and lung of cavy (Galea spixii). In [A] comparative morphology of the three Groups: I (33-35 days), II (38-40 days), and III (43-45 days). Note on figure A: short external ear (ee), occipital region (oc), cephalic region (ce), pigmented retina in the optic vesicle (ov), forelimb (fl), hind limb (hl), short tail (st), nasal region (na), upper eyelid (ue), lower eyelid (le), keratinized claw (cl), sensorial hair (sh). In [B] and [C] ventral and dorsal view of the lung, respectively; [D] and [E] scanning electron microscopy of the lung and trachea. Note on the right side: right cranial lobe (RCrL), medium lobe (ML), accessory lobe (AL), and right caudal lobe (RCaL), and on the left side: left cranial lobe which was divided into cranial (CrLCrL) and caudal (CaLCrL) segments, and the left caudal lobe (CLL). The trachea (T) was composed by hyaline cartilage rings (C) and connective tissue (TC).
Both, Group I and II showed similarities in regarding to the expression of PCNA, VEGF, and Oct-4 markers. In contrast, the samples from Group III were negative for all markers. Thus, throughout early pregnancy it was observed an intense proliferative activity on epithelial cells lining the bronchi and bronchioles, hialine cartilage, and mesenchyme on Groups I and II, as demonstrated on figure 3A for the Group I. In the same regions there was no labbelering for Group III (Figure 3B). It was observed the expression of VEGF in the epithelial region of the bronchi, bronchioles, and mesenchyme in the Groups I and II (Figure 3C), and its absence of expression on Group III (Figure 3D). In relation to the presence of pluripotent cells, it was verified the co-location of these Oct-4+ cells with those who were on proliferative phase, especially in the epithelial bronchi and bronchioles, and also on alveolar cells of the Groups I and II (Figure 3E). In contrast, it was not identified Oct-4+ cells in the Group III (Figure 3F).

**DISCUSSION**

Currently, the most common experimental models used for the study of acute and chronic lung diseases are represented by small rodents and mini pigs, due to the easy and practical management of these animals in animal facilities and low costs (CABRAL, 2013). However, there is a constant search for new animal species that may present, at

Ciência Rural, v.46, n.9, set, 2016.
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Ciência Rural, v.46, n.9, set, 2016.

Figure 3 - Immunohistochemical analysis for proliferation (PCNA) [A and B], angiogenesis (VEGF) [C and D], and pluripotency (Oct-4) [E and F] in the lung of fetuses of cavy (*Galea spixii*). Group I (33-35 days) and Group III (43-45 days). Positive labeling was observed in epithelial cells of the bronchioles (brl), hyaline cartilage (hc), mesenchyme cells (ms), alveoles (al), and epithelial cells (ep) of individuals from Groups I and II. In contrast, the Group III was negative for all markers [B, D, F].

With regard to the lung development pattern, this is described in several species of mammals, although times of development of each phase are significantly different among the species, especially due to the period of gestation and the relative degree of lung maturation at birth (PINKERTON & JOAD, 2000). According to PINKERTON & JOAD (2000), the pseudoglandular phase in humans is the most critical stage for the development of the conductive portions. In the cavy, the pseudoglandular phase was associated with the development of the bronchi and bronchioles with a simple cuboid epithelium, as described for mice (WARBURTON et al., 2010). This is different to the morphological descriptions of the bronchi and bronchioles epithelium in humans, for which a columnar epithelium is characteristic (PINKERTON & JOAD, 2000). According to ELHAFEZ (2012), the canalicular stage is an important step in the development of the rabbit lung, since the terminal bronchioles branches into 2-4 acinar channels, from which some cells of the epithelium line these channels and differentiate into type I pneumocytes. In the descriptions for Group II, the most remarkable features were: increased number of bronchi and bronchioles, division of the lung in lobes, and mesenchymal differentiation. Moreover, results showed progressive decreased expression of markers involved in cell proliferation (PCNA), angiogenesis (VEGF) and pluripotency.
(Oct-4) during pregnancy. Recent results showed the importance of these markers for differentiation processes, morphogenesis and lung growth (PINKERTON & JOAD, 2000). The decreased expression of these markers during gestation can be explained by the decrease of proliferative activity of the cells directly before birth. It is known that the vascular endothelium growth factor (VEGF) is highly expressed in epithelial cells located near to microvascularization tissue (BERSE et al., 1992). Indeed, it was observed an intense staining for VEGF in the lung parenchyma, which is richly vascularized. In regard to the Oct-4, which was expressed in the lung of the cavy in Groups I and II, it represents a marker expressed by pluripotent cells and its expression in cell and tissue cultures is related to cell differentiation (PALMIERI et al., 1994). In addition, it plays an important role in self-renewal of somatic stem cells and maintenance of tissue homeostasis (WU & SCHOLER, 2014). This indicates that the early stages of development of the cavy lung shows undifferentiated cells with high capacity of differentiation.

In most mammals with short gestation periods, such as rodents and lagomorphs, the birth occurs before the formation and maturation of the alveoli in the lungs, so, in these species the alveolarization can be considered late and/or an postnatal event (PINKERTON & JOAD, 2000). However, the cavy development pattern differs from other rodents, since the alveolar period occurs earlier when compared to other rodents and lagomorphs.

Finally, an important feature of the development of the cavy lung is that they are born in the alveolar stage, as well as in humans, whereas rats and mink are born in the saccular stage (SCHITTNY, 2007), allowing infer that the cavy can be used as an experimental model for understaing the development of early respiratory diseases in humans.

CONCLUSION

Among the variety of mammalian species for which lung embryology is known, the cavy (G. spixii) showed more similarity with human fetuses than other rodent species even from the same systematic group, especially because they born in the alveolar stage. It was demonstrated that there is a high intensity markers expression involved in cell proliferation, angiogenesis and pluripotency in the earliest stages of development, showing that these cells have a high capacity for proliferation and pluripotency, suggesting its use in future studies, mainly related to the establishment of cell cultures for regenerative medicine.

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