INTRODUCTION

Placenta is the communication organ that enables the exchange of various materials between mother and fetus. In addition to useful ones, transport of harmful and toxic substances may also take place. These can lead to premature birth or malformations of the fetus [1]. Advancement in biomedicine is increasingly attracting more interest towards the mammalian development, in which one of the key roles is played by the placenta. Understanding the fundamental processes of placentation in different experimental animals, as well as the distinction of morphologies of their placentas, is undoubtedly needed for understanding the processes of human development [2].

The blastocyst, the structure common to all the mammals, is composed of the embryoblast and the trophoblast [3]. Trophoblast is crucial for the maintenance of pregnancy because it establishes the initial contact with the mucous endometrium of the uterus, assisted by expression of a series of glycoproteins on its membrane, e.g. cadherins and integrins [4]. The placentation in the rat occurs on the antimesometrial side of the uterus, whereas placenta is being formed on the mesometrial side of the uterus. On the 8th day of rat gestation, the trophoblast proliferates and gives rise to the ectoplacental cone and on the 9th day maternal lacunae without endothelium are being developed within this ectoplacental cone [5]. At the same time, an offshoot of the allantois is created. The mesenchyme of this offshoot on the 11th day of gestation establishes contact with the ectoplacental cone, i.e., with its basal part, which is called the chorion [6]. Action of allantoic mesenchyme results in angiogenesis, and thus the fetal capillaries are being developed amongst the maternal lacunae. This process leads to the formation of the functionally most important part of the rat placenta - the labyrinth, where nutrients and gasses are being exchanged between mother and the embryo [7]. It is believed that defects in development of the labyrinth can lead to various disorders of placentation, and consequently to growth arrest of the fetus.
Variety of genes exists whose timely expression is essential for proper development of the labyrinth [8]. As the placentation in rats is of hemochorial type, i.e., the trophoblast is in direct contact with maternal bloodstream, the trophoblast invasion is necessary. The invasion is mediated by the trophoblast giant cells (TGC) and glycogen-rich trophoblast cells [9]. TCG are believed to be responsible for the endovascular invasion, whereas the trophoblast glycogen-rich cells are responsible for interstitial invasion in the uterine endometrium [10].

The rat placenta has three main parts, namely the basal plate or trophospongiosis, the labyrinth and the decidual part [11]. The basal part is located close to the decidua and is composed of several cell types: trophoblast stem cells, trophoblast giant cells, and glycogen cells. The labyrinth accommodates maternal lacunae and fetal blood vessels. They form separate bloodstreams segregated by placental membrane [12]. This membrane makes the rat placenta of hemochorial type, because it has three layers of trophoblast [13].

The aim of this research was to determine the dynamics of changes in absolute volumes of placentas between the 16th and the 19th day of the gestation, by applying stereology. Furthermore, changes in absolute volumes of individual components of the placenta: the labyrinth, the basal layer, and the decidua were evaluated. Finally, because of their central role in trophoblast invasion into the uterine endometrium, which is essential for proper hemochorial placentation, the volume density and the absolute volumes of TGC and glycogen cells were determined.

MATERIALS AND METHODS

Placental samples

Adult Fischer female rats (three months old) were mated overnight with males of the same age. Finding of the sperm in the vaginal plug the next morning designated the day 1 of pregnancy. Pregnant animals were euthanized using ether on the day 16 or the day 19 of gestation. Abdomen was opened, and uterine horns were held out with forceps. Uterus was excised and washed in Dulbecco’s phosphate buffered saline (PBS). Uterine horns were dissected under the binocular dissecting microscope, and placentas were isolated. Placentas were weighed and fixed in 10% formaldehyde, embedded in paraffin, processed for routine histology and 10 mm thick sections stained by hematoxylin-eosin (HE).

This study was approved by the Ethical Committee of the Medical School of the University of Zagreb.

Stereological analysis

Stereological analysis was carried out on three randomly selected experimental placentas from the day 16 of gestation and three placentas from the day 19 of gestation. For both days, the stereological analyzes included the absolute volume of total placenta and the absolute volumes of the basal layer, the labyrinth and the decidua. The volume density of the gly- cogen-rich cells and the giant cells of the trophoblast, as well as the absolute volume of these two types of cells in the basal layer of placenta, were also determined.

The absolute volume of placenta

For the determination of absolute volumes of analyzed placentas, we have constructed a test system for stereology in the Ellipse 3D computer program.

Our test system consisted of 494 testing points, superimposed on the vertically oriented, previously photographed, paraffin sections. Analyzed sections were photographed on Olympus BX41 microscope (dissecting microscope Olympus SZX7, camera Olympus DP12) in the Department for Biology and Pathology of Fish and Bees, School of Veterinary Medicine of the Zagreb University. Hits (test points falling on the placental tissue) were counted according to the principle of the Cavalieri-method for the determination of the absolute volume according to formula:

\[ V_P = t \times a_p \times \Sigma \, P \]

Where \( V_P \) is the absolute volume of placenta, \( t \) is thickness of serial section, \( a_p \) is test area belonging to a particular test point, and \( \Sigma \, P \) is total number of hits on all the sections of a single placenta [14]. Thickness of serial sections was determined by preliminary measurements. Determination of the corrective shrinkage factor was carried out by measuring diameters of 200 randomly selected erythrocytes, and compared with the same number of diameters of fresh erythrocytes obtained from the sacrificed pregnant rat. The absolute volume of the placenta was subsequently corrected using this shrinkage factor [14].

Absolute volumes of placental components

Absolute volumes of the labyrinth, the basal layer and the decidua were determined by the same method as used for determination of the absolute volume of the whole placenta. Three testing systems out of 494 test points in the Ellipse 3D stereological program, i.e. one for each placental component (labyrinth, basal layer or decidua) were constructed. Total of volumes of all placental components (phases) was supposed to give the value of the absolute volume of the whole placenta obtained by preliminary measurement, which was used as control of the measurements performed.

The volume density and the absolute volume of TGC and glycogen-rich cells

For determination of the volume density, a testing linear system in Ellipse 3D computer program was constructed.
where the length of the testing line \((L/p)\) was equal to 0.29 mm and the testing area \((A_p)\) was 0.18 mm². This testing system was superimposed on every analyzed section, i.e. it covered the whole cross section of the placenta, so that the trophoblast giant cells and glycogen-rich cells were counted in the entire section, rather than a single visual field, as is the case on the microscope.

As it is most important to define the referent space immediately before the start of measurement, the basal layer of the placenta was defined in this case and hits for this part were counted. Subsequently, hits for glycogenic and giant cell of the trophoblast were counted on the same field.

The volume density \((V_v)\) is the relative stereological variable, which indicates how large is the share occupied by the studied phase in the total space or what is the percentage of the studied phase in unit volume.

The absolute volume of the studied phase was calculated by multiplication of the volume density of the studied phase with the absolute volume of the basal part of placenta, obtained from the previous measurements.

**Statistical analysis**

Mean, standard deviation and standard error were determined using descriptive statistics. For statistical analysis of the mass differences between the two analyzed groups of placentas Student’s \(t\)-test, was used.

**RESULTS**

Placental masses on the day 19 of gestation \((0.353±0.032g)\) were significantly higher than on the day 16 of gestation \((0.299±0.039g)\) \((p<0.05, \text{ Student’s } t\)-test\) (Figure 1).

Analysis of the absolute placental volume on the days 16 and 19 of gestation

Stereological analysis was performed on three randomly selected placentas from each experimental group and analyzed in the total of 660 histological sections. To determine the absolute volume, a “Point grid system” was designed, which consisted of 4 frames (Figure 2). Each of these frames corresponded to one phase – a component of the placenta analyzed.

Mean absolute volume of the placenta on the day 16 of gestation was 0.0638 cm³, out of which the labyrinth occupied 0.0274 cm³ (43%), the basal layer 0.0271 cm³ (42%) and the decidua only 0.0093 cm³ (15%).

On the day 19 of gestation, the mean absolute volume of the placenta has increased to 0.1627 cm³, out of which the labyrinth occupied 0.0922 cm³ (56.5%), the basal layer 0.0596 cm³ (36.5%) and the decidua only 0.0109 cm³ (7%) (Table 1; Figure 3).

The volume density of trophoblast giant cells and glycogenic cells on the day 16 and 19 of gestation

For the analysis of the volume density of trophoblast giant cells and glycogenic cells a “Line System” of equal length of the test...
line and the test areas were constructed (Figure 4). On the day 16 of gestation, the volume density of the trophoblast giant cells was 0.174 cm³, and the volume density of glycogenic cells was 0.379 cm³.

On the day 19 of gestation, as expected, the decline in both volume densities was observed, to 0.107 cm³ for trophoblast giant cells and 0.236 cm³ for glycogenic cells (Table 2).

On the basis of the obtained results on the volume densities, we then calculated the absolute volumes of these cells in the basal layer of the placenta (Table 3). Thus, on the day 19 of gestation, as a consequence of the increase in the absolute volume of the basal layer of the placenta, an increase in the absolute volumes of the trophoblast giant cells was observed as well (Table 3).

**DISCUSSION**

We obtained the accurate data on absolute volumes of individual components of the placental structure and changes in their values during the last third of gestation. These data provide an insight into the adaptation of placenta, particularly its labyrinth part, to the greater exchange of matter between mother and fetus as the gestation nears its end. We discovered the reduction of volume densities of trophoblast giant cells and trophoblast glycogen cells, which coincides with the reduction of their functions at the end of gestation. At that time, the trophoblast invasion is at its end, and the definitive placenta is formed.

This research also provides results on absolute volumes of the whole placenta and its components during normal placental development in the rat, which might be even a better animal model to study placental function than the mouse [15]. Like humans, rodent placenta is of hemochorial type, i.e. the trophoblast cells originating from the chorion are in direct contact with the erythrocytes of the maternal blood. In hemochorial placenta, trophoblast invasion takes place in the uterine endometrium.

Similar stereological studies were carried out on the mouse placenta by Coan et al. and they obtained results analogous to our results for rat, with respect to the differences in gestational age between the two studies. An increase in the absolute volume of the entire placenta but not in its components [14] was found in the mouse. While the absolute volume of the labyrinth increased, the volumes of other placental components (the basal layer and the decidua) exhibited a significant reduction at the end of gestation.

We found that the labyrinth of the rat placenta occupies 43% of the placenta on the day 16 of gestation and 57% of the placental volume on the day 19 of gestation, which is similar to values in the mouse [14]. Comparing our results with those of Coan et al. [14] on the absolute volumes of the placental basal layer and the decidua, there is a similarity in the reduction of their volumes at the end of gestation. Volume share of the basal layer in mouse placenta is about 29% on the day 17 and 24% on the day 19 of gestation, while in our analysis of the rat placenta, the basal layer occupies 43% of the placenta on the day 16 and only 36% on the day 19 of pregnancy. The volume share of the decidua in mouse placenta decreased to 23% on the day 17 of gestation and to 18% on the day 19, while in our case of the rat placenta, the reduction in the volume share of the decidua falls to 15% on the day 16 of gestation and to only 7% of placental mass on the day 19 of pregnancy.

The results of our work are important in view of stereological studies of placenta in normal and in complicated pregnancies, where stereology was successfully applied for interpreting the morphology of human and animal placenta [16], in describing the villous trophoblast growth, trophoblast differentiation, vascular morphogenesis and diffusion transport [17,18].

Our results can be of help in future studies if we assume that the basic stereological variables of individual components of the placenta, and of the whole placenta vary depending
on different harmful environmental influences, or in cases of maternal illness.

The data presented regarding the volume density of TGC and glycogen cells can be very useful in further research of endovascular trophoblast invasion, considering the existing research on human and rat placentas on the effects of pre-eclampsia and other factors that can bring about changes in these processes during pregnancy [19]. Trophoblast invasion in spiral arteries of the uterus is the essential prerequisite for successful development of placental circulation [20], and it is important to understand the dynamics of this process. Crucial roles in this process play trophoblast glycogen cells and trophoblast giant cells, for both of which we have obtained data on their volume densities and absolute volumes. Their volume densities were higher on the day 16 of gestation, when the trophoblast invasion was very intensive, and they diminished by the day 19 of gestation, when the trophoblast invasion ended.

Taking into account the diversity of application possibilities of stereology and the relatively low cost of this technique, stereology should certainly be considered as one of the methods for obtaining quantitative data in both the clinical and basic medical research [21].

CONCLUSION

The results of our study may be used for further research on rodent placentation. The absolute volumes of the labyrinth, decidua and the basal layer, as well as the volume densities and the absolute volumes of the trophoblast giant cells and trophoblast glycogen cells may be used for characterization of effects of various toxic substances and harmful environmental conditions on the components of placenta.

DECLARATION OF INTERESTS

The authors declare no conflict of interest.

REFERENCES