

# Multinucleated giant extravillous trophoblast cells in the decidua of placentas from pregnancies complicated with preeclampsia

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**UNIVERSITY OF SPLIT  
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**MULTINUCLEATED GIANT EXTRAVILLOUS TROPHOBLAST  
CELLS IN THE DECIDUA OF PLACENTAS COMPLICATED  
WITH PREECLAMPSIA**

**Diploma thesis**

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## **1. Introduction**

## 1.1. Preeclampsia

### 1.1.1. Definition

Preeclampsia (PE) is a pregnancy related disorder, characterized by raised blood pressure and signs of kidney damage. The onset of this condition is usually after 20 weeks of gestation.

According to the American College of Obstetricians and Gynecologists (ACOG), the criteria for the diagnosis of PE include onset of a new episode of hypertension during pregnancy, characterized with persistent hypertension (systolic blood pressure  $\geq 140$  mmHg or diastolic blood pressure  $\geq 90$  mm Hg) and substantial proteinuria ( $>300\text{mg/L/24 hours}$ ) (1). In case of absence of proteinuria or proteinuria below the diagnostic threshold, other diagnostic criteria are required. They include impaired liver function, new-onset thrombocytopenia, pulmonary edema, visual or cerebral disturbances or renal insufficiency (1).

PE can progress to severe PE, which includes liver dysfunction, edema, pain, low platelet count as well as vascular and cerebral disturbances. Less than 1% of women with PE proceed to the most severe form, named eclampsia, which is characterized by eclamptic seizures (2).

### 1.1.2. Epidemiology

Hypertensive disorders of pregnancy (HDP) complicate up to 10% of all pregnancies. They include gestational hypertension, chronic hypertension, chronic hypertension superimposed with PE and PE. The later one affects about 3% of all pregnancies, making up a substantial part of HDP (3).

PE/eclampsia is a major cause of maternal and fetal mortality and morbidity. The maternal death rate of eclampsia and preeclampsia is 0-1.8% in developed countries and as high as 14% in developing countries. Fetal mortality is estimated to be 13-30% (4).

Women with a previous history of PE are at 7 times higher risk of developing the condition in following pregnancies (5). Other risk factors include: age over 40 years, first pregnancy (primipara), maternal obesity, maternal diabetes (type 1 or type 2), previous hypertension, coagulation disorders, systemic diseases, like systemic lupus erythematosus (SLE), kidney diseases and multiple pregnancy (6).

### 1.1.3. Etiology

The etiology of PE remains unknown. One of the most accepted theories characterized PE as a multisystem disorder due to abnormal response of the maternal vasculature to placentation.

In normal pregnancies, structural modifications of the spiral arteries by invasion of the cytotrophoblasts (CTB) causes them to be highly dilated and thus to become less resistant. This allows them to be less sensitive to vasoconstrictive substances. The described physiological change plays a vital role for the supply of nutrients to the placenta and eventually the foetus (7).

In placentas from pregnancies complicated with PE, this remodelling process has failed and the remaining increased uterine arterial resistance is preserved or even increased. In addition, the preserved sensitivity to vasoconstrictive substances induces a condition of chronic placental ischemia and oxidative stress (8).

Fetal complications develop as sequel to placental ischemia whereas maternal complications are caused by oxidative stress triggering the release of certain substances into the maternal circulation. These substances include free radicals, oxidized lipids, cytokines, and serum soluble vascular endothelial growth factor 1 (9).

They are responsible for the vascular endothelial dysfunction seen in PE, including vascular hyperpermeability, thrombophilia and hypertension (8).

Another mechanism contributing to the development of PE is a disturbed trophoblast life cycle. Changes in apoptosis, meaning programmed cell death of the trophoblast, is thought to be one of the core factors for evoking an inflammatory response from the mother, as seen in PE. The normal trophoblast lifecycle, which is approximately 25 days long (10), includes differentiation and proliferation of mononucleated CTB with eventual fusion into multinucleated syncytiotrophoblasts, followed by the formation of syncytial knots. Those syncytial knots represent dying remnants of the trophoblast and they are finally shed from the surface of villi into the maternal circulation, where maternal pulmonary macrophages eliminate them. The described process is also known as trophoblast turnover (11).

In PE however, several studies revealed an imbalance between proapoptotic and antiapoptotic factors and therefore an increased apoptosis, which is believed to be crucial in the development of PE (11,12). In this context, aponecrosis, meaning incomplete apoptosis and eventual necrosis with formation of impaired syncytial knots, is another important parameter. Compared to

apoptosis with normal formation of syncytial knots, those produced from aponecrosis are still capable of evoking an inflammatory response from the mother (13).

PE is a disorder with a high risk of progression to more severe conditions like eclampsia, which includes convulsions. The risk of developing cardiovascular disorders and chronic kidney disease due to metabolic stresses and vascular injury induced by PE is increased. In case of severe PE, the clinical findings despite hypertension and proteinuria also include signs of microangiopathy, systemic endothelial dysfunction and the HELLP syndrome (haemolysis, low platelet count and elevated liver function tests) (14).

Other symptoms include swelling, shortness of breath and visual disturbances. Preterm delivery, placental abruption, caesarean delivery and even stillbirth show an increased occurrence in pregnancies complicated with PE. The need for preterm delivery, as only known cure for PE, poses a risk for the neonatal wellbeing. Late preterm neonates appear to have a significantly higher mortality rate than term infants (15-19).

The decreased uteroplacental blood flow during pregnancy and the resulting ischemia increases the risk for the foetus to develop intrauterine growth retardation. Neonatal thrombocytopenia, increased risk for neutropenia, bronchopulmonary dysplasia and variable neurodevelopmental outcomes are further complications for the neonate (20).

## 1.2. Placenta

### 1.2.1. Development of placenta and its role in normal pregnancy

Six days after fertilization, the blastocyst attaches to the endometrium. The trophoblast, which is the outer layer of the blastocyst, proliferates and differentiates into the outer syncytiotrophoblast (STB) and the inner CTB to enable the implantation of the embryo. The CTB give rise to amnioblasts, which eventually form the amnion. The STB on the other hand, invade the endometrium and form lacunae, which become filled with maternal blood. Primitive intervillous spaces form by fusion of several lacunae. Primitive villi develop from CTB proliferation. The chorion is formed by extraembryonic mesoderm and the primitive villi. Chorionic villi, which are in contact with the maternal blood in the lacunae develop from the villous trophoblast (VTB). The chorionic plate is build by rapidly proliferating villi along the decidua basalis. Extravillous trophoblast (EVTB) that is formed by CTB in the tips of villi, penetrate the maternal vessels and



help to anchor the placenta. Besides, EVTB is further classified into interstitial and the endovascular trophoblast. The later is responsible for the remodeling of the spiral arteries shortly after implantation. This process is crucial in the development of the placenta. It ensures the high flow, low resistance uteroplacental vasculature in the normal placenta. The remodelling of spiral arteries by endovascular trophoblast has two peaks in time. The first one is around 8 to 10 weeks of gestation and involves the invasion into decidual segments of spiral arteries. At 16 to 18 weeks of gestation the second peak occurs with invasion into myometrial segments. This mechanism ensures gradual loss of spiral artery musculature and replacement with fibrinoid material to enable a successful pregnancy (21).

Umbilical vessels formed by the mesoderm in the core of the villi, the allantois and remnants of the yolk sac build up the umbilical cord. In the end there is a fusion of amnion and chorion to form the amniochorionic membrane, which eventually fuses with the decidua capsularis and the decidua parietalis as the embryo grows (22,24).

The placenta is fully functional by the end of the third month of gestation. Its role is to provide nutrients and oxygen for the foetus as well as to eliminate waste products. Furthermore it has an immunologic role, as IgG antibodies may pass to provide passive immunity for the foetus. It also serves as endocrine organ, secreting human chorionic gonadotropin to ensure that the pregnancy may take place as well as a set of different other hormones. Since the placenta carries much more blood than actually needed by the foetus, it also has a compensatory function in case of diminished blood flow (25).

When fully developed, on gross examination, the placenta is composed of the placental disk, extraplacental membranes and the umbilical cord (Figure 1).

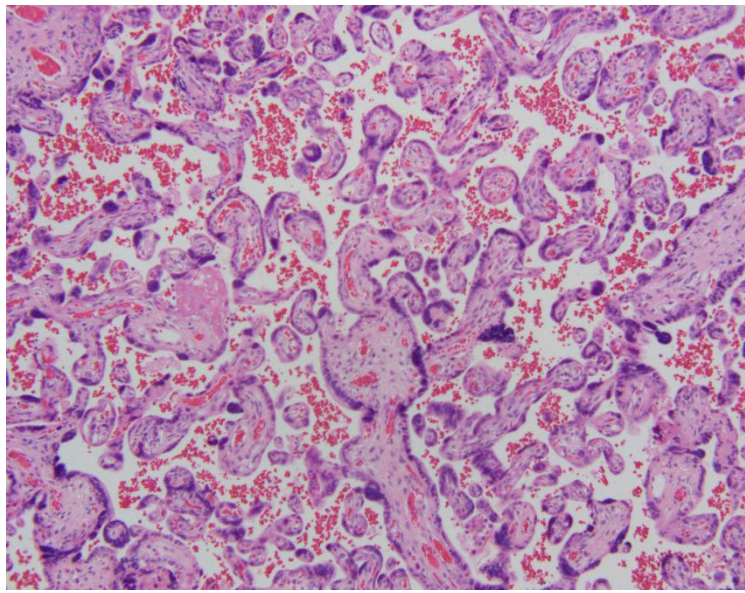


**Figure 1.** Gross image of term placenta. A. Maternal side of the term placenta. B. Foetal side of the placenta with the umbilical cord. (*Images taken from the personal archive of the mentor with mentor's permission*).

Cotyledons and the basal plate can be found on the maternal side of the placental disk, whereas foetal side, which is covered with the amnion, is called chorionic plate. The latter is the place for umbilical cord insertion. Extraplacental membranes are usually attached to placental disk margins (23).

Microscopically, the basal plate includes decidua basalis, EVTB and maternal blood vessels. The chorionic plate is composed of subchorionic fibrin, umbilical blood vessel branches, chorion and amnion. In between basal and chorionic plate, stem, intermediate and terminal villi as well as intervillous spaces are seen, building the placental parenchyma. Septa, made out of EVTB are subdividing the basal plate into cotyledons. The membranes consist of amniotic epithelium, connective tissue and chorion, which is composed of EVTB and involuted villi, attached decidua parietalis and capsularis as well as maternal blood vessels and yolk sac remnants (Figure 2).

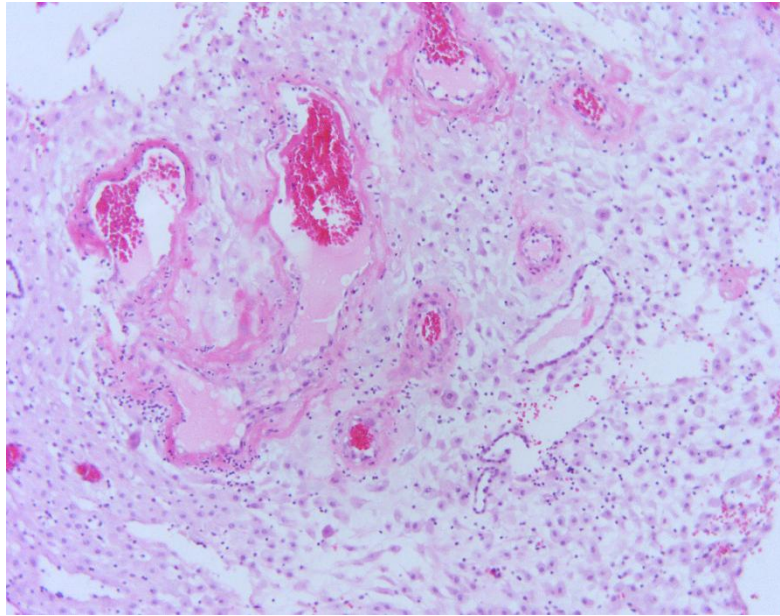
Finally, the umbilical cord is made of two arteries, one vein, Wharton's jelly and surface epithelium (23).



**Figure 2.** Histology of term placenta with placental villi, syncytial knots and intervillous spaces with maternal blood (Hematoxylin&Eosin, magnification 100x, Olympus BX41 microscope; *image taken by candidate at the Pathology Department*).

#### 1.2.2. Placenta in preeclampsia

Abnormalities in the placenta are believed to be the major components in the PE development. Diminished blood flow through the placenta due to defects in the maternal vasculature to adapt to normal pregnancy is the core pathological feature of PE. At about 10 weeks of gestation, the intervillous space becomes vascularized and CTB undergoes changes to respond to the previous low oxygen environment. The differentiation into the invasive trophoblast phenotypes allows the spiral arteries to be modified into flaccid, funnel shaped large diameter vessels, which are insensitive to humoral or neural vasoconstrictive substances. In PE however, these vessels remain contractile and cause a large amount of oxidative stress. Acute atherosclerosis, the occlusion of vessels by fibrinoid surrounded by foam cells, is another change seen microscopically in the unmodified spiral arteries, contributing to the state of underperfusion (Figure 3) (26,27).



**Figure 3.** Acute atherosclerosis of blood vessels in the decidua basalis from placenta with preeclampsia (Hematoxylin&Eosin, magnification 100x, Olympus BX41 microscope; *image taken from the personal archive of the mentor with the mentor's permission*)

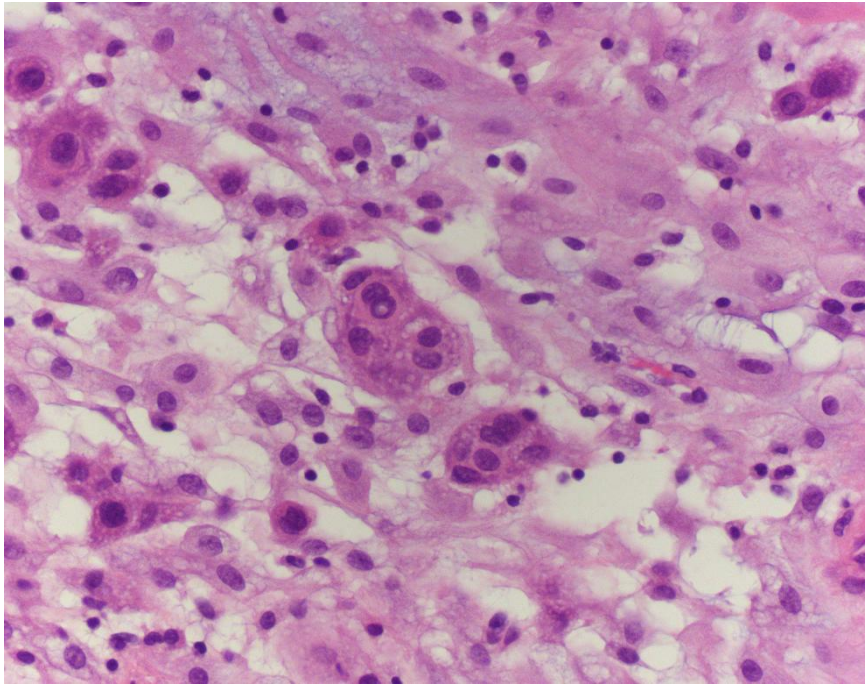
Histologically, numerous large syncytial knots, small sclerotic villi and accelerated villous branching are seen due to hypoxia and resulting reactive oxygen species. Especially in severe PE, also gross morphological changes like infarctions and abnormal placental shape might be noted. Despite some characteristic changes in placentas of pregnancies complicated with PE, the exact linkage between abnormal trophoblast invasion and the maternal disease remains unknown (26,27).

### 1.2.3. Multinucleated Giant Cells

Multinucleated Giant Cells (MGC) constitute a subgroup of EVTB near the site of implantation, which have lost their invasive properties (28,29). Their trophoblastic origin was confirmed by immunohistochemical labelling and observation with a light microscope by Pijnenborg *et al.* as well as Wells and Bulmer (30,31).

MGCs appear to be larger than the surrounding trophoblast and contain two or more nuclei, which is probably attributed to cell fusion (Figure 4).





**Figure 4.** Multinucleated giant cells (MGC) in the decidua basalis from placenta with preeclampsia (Hematoxylin&Eosin, magnification 400x, Olympus BX41 microscope; *image taken by candidate at the Pathology Department*).

MGC mostly appear in mid pregnancy but they can also be seen later (28).

Usually MGCs can be found as a normal component of the implantation site but they appear to be increased in placentas of pregnancies complicated by PE and decreased in placenta creta (32-35). Hence, they are not merely a characteristic of maternal vascular underperfusion in PE but also linked to other types of pregnancy related disorders associated with placental hypoxia. Their significance lies in being very specific and therefore MGC might be used to retrospectively clarify perinatal mortality and morbidity, meaning when seen in excess number in placentas from full term pregnancies, they could be indicative of pre-existing hypoxia (29).

## **2. Hypothesis**

Placentas from pregnancies complicated with PE show an increased number of MGC in the decidua basalis compared to placentas from normal pregnancies.

### **3. Material and Methods**



The study was conducted upon approval of the local Ethics Committee and after all participants provided informed consent.

Overall this study included 30 placentas.

The studied group was represented by 15 placentas from singleton term pregnancies complicated with PE, which was diagnosed based on following criteria: systolic blood pressure >140 mmHg and diastolic blood pressure >90 mmHg; proteinuria >300 mg/L per 24 hours (1).

The remaining 15 placentas were collected from singleton term pregnancies without clinical signs of PE and posed the control group.

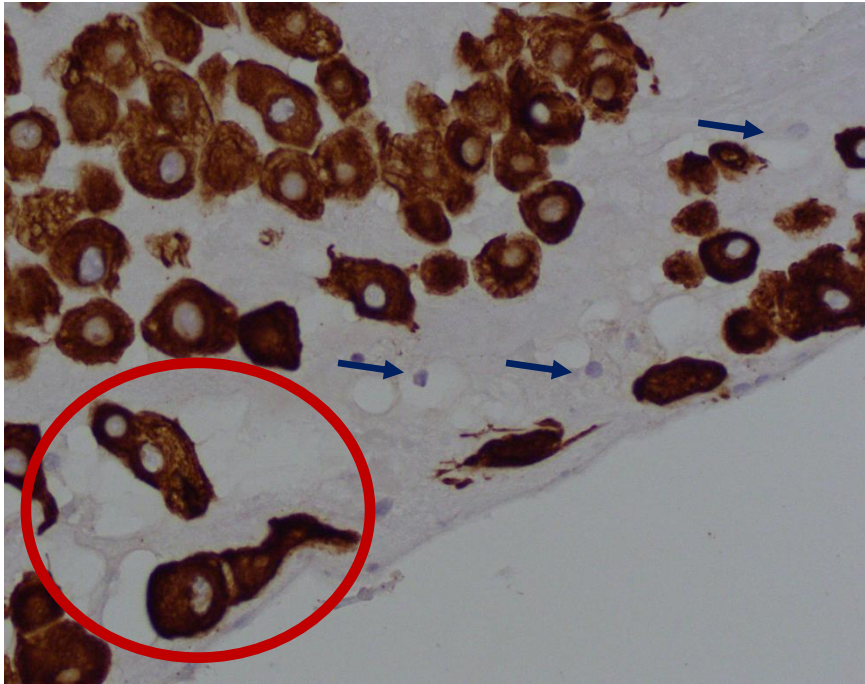
General exclusion criteria for both groups were: diagnosis of HELLP syndrome (platelets <100x10<sup>9</sup>/L, AST and ALT >70 U/L, and LDH >600 U/L) (14), renal diseases, vascular diseases like hypertension, or any other maternal condition existing prior to pregnancy.

Within 20 minutes after delivery, placentas were collected from the delivery room of the University Hospital of Split, fixed in 4% buffered formalin and sent to the Pathology Department for further evaluation.

Placental samples, all containing decidua basalis, were taken from the macroscopically normal part of the placental disk, near to the umbilical cord insertion site. Sections of 4-μm thickness were cut from previously paraffin embedded placental samples. Afterward they were mounted on silanized slides and dried at 37°C. After hematoxylin and eosin staining, histologic examination was performed using an Olympus BX 41 microscope (Olympus, Tokyo, Japan).

Immunohistochemistry was done after the paraffin slices were mounted on super frost slides (Thermoscientific, Darmstadt, Germany) and further processed in the automatic stainer (Ventana Bench Mark Ultra autostainer, Tucson, USA). CK7 antibody (rabbit monoclonal antibody, clone SP52, Ventana, Tucson, USA) was used to distinguish EVTB from decidual cells (DC) in the decidua basalis with Ultra view Universal DAB detection kit (Ventana, Tucson, USA). DC were CK7 negative whereas EVTB cells were CK7 positive.

Since MGC are considered to be EVTB, they were also CK7 positive (Figure 5).



**Figure 5.** Immunohistochemical staining with CK7, brown membranous and cytoplasmic staining of extravillous trophoblast (EVTB) and multinucleated giant cells (MGC, marked with red circle). Decidual cells (DC) are negative (blue arrows). (Magnification 400x, Olympus BX41 microscope; *image taken by candidate at the Pathology Department*).

For each sample, the number of MGC, EVTB and DC in the decidua basalis was determined by the use of Olympus BX 41 microscope and Olympus cell D1 image analyser.

Statistical analysis was done by the MedCalc software (MedCalc softare, Mariakerke, Belgium) using the chi-square-test, t-test for independent samples and Mann-Whitney test, depending on the result distribution. Statistical significance was set at  $p < 0.05$ . Results were expressed as mean with standard deviation or median with minimum and maximum value, depending on the result distribution determined by Kolmogorov-Smirnov test.

## **4. Results**

Fifteen placentas from PE were included in the studied group and the same number of placentas from normal pregnancies in the control group. There was no statistically significant difference regarding maternal age ( $p=0.390$ ;  $t=0.872$ ;  $DF=27$ ), gestational age ( $p=0.644$ ;  $t=0.468$ ;  $DF=28$ ), birth weight ( $p=0.657$ ;  $t=0.449$ ;  $DF=28$ ) and neonatal gender ( $p=0.695$ ;  $\chi^2 = 0.153$ ;  $DF=1$ ) between investigated groups (Table 1).

There was no statistically significant difference in the number of DC and EVTB between both groups ( $p=0.112$ ;  $t=1.643$ ;  $DF=28$ ); ( $p=0.115$ ;  $Z=3.846$ ) as well as in MGC/EVTB ratio ( $p=0.113$ ;  $Z=1.587$ ) (Table 2).

The number of MGC was significantly higher in the PE group compared to control group ( $p<0.001$ ;  $Z=3.765$ ). Likewise MGC/DC ratio was significantly higher in the PE group compared to control ( $p<0.001$ ;  $Z=1.587$ ) (Table 2).

Table 1. Age of participants, gestational age at birth as well as birth weight and gender of the neonate

	PE group (N=15)	Control group (N=15)	p
Age of the mother (years)	$34 \pm 7.14$	$31 \pm 7.86$	0.390*
Gestational age (weeks)	$38 \pm 1.36$	$38 \pm 0.94$	0.644*
Birth weight (grams)	$3304 \pm 971.88$	$3429 \pm 462.34$	0.657*
Gender (male sex)	11	10	0.695†

\*Student t-test

†chi-square tests

The values are presented as arithmetic mean  $\pm$  standard deviation or general number.

Table 2. The number of multinucleated giant cells (MGC), decidual cells (DC) and extravillous trophoblast (EVTB) as well as the ratio between MGC and DC and MGC and EVTB

	PE group (N=15)	Control group (N=15)	p
MGC	3 (0-15)	0 (0-4)	<0.001*
DC	205.46 ± 227.64	325.46± 167.95	0.112†
EVTB	880 (153-2290)	420 (61-2438)	0.115*
MGC/DC	0.03 (0-14)	0 (0-0.01)	<0.001*
MGC/EVTB	0 (0-0.04)	0 (0-0.02)	0.113*

\*Mann-Whitney U test

† Student t-test

The values are presented as median with minimum and maximum values or as arithmetic mean ± standard deviation.

## **5. Discussion**

PE is a very heterogeneous condition with many influencing factors and a still unknown etiology. Many of those factors are being investigated to show their interference with normal placental development. This study aimed to show the number of placental MGC in the decidua basalis from PE placentas and to compare the results with MGC number in placentas from normal pregnancies.

Results of this study have shown that there is a significant difference in the number of MGC in PE placentas compared to the control group, MGC being higher in the studied group.

To the best of my knowledge, it is the first study to investigate the number of MGC in the decidua basalis from PE pregnancies solely and to compare them with the results from normal placentas.

MGC can also be found in normal pregnancies, especially in mid pregnancy (28). Their function is largely unknown. It is thought that they are created from EVTB by cell fusion (29-31).

A significant increase of MGC in the studied group suggests that MGC formation is being triggered by a hypoxic injury pattern. There are many different placental responses to hypoxia. Not only the duration of hypoxia plays a role, but also whether it is a preuterine, uterine or postuterine pattern of hypoxia as Stanek pointed out (36). He described that MGC in the decidua basalis, as well as excessive EVTB numbers are less commonly seen in preuterine pattern of chronic placental hypoxia (e.g. maternal anemia). Also, he emphasized that this pattern has a better prognosis than other types of placental hypoxia and concluded that it is most probably due to hypoxic preconditioning. This implies that the specific hypoxic injury pattern plays a major role in the production of MGC. Hence, they are not increased in all hypoxic patterns, which makes them relatively specific for certain types of hypoxia, as for chronic uterine hypoxic pattern seen in PE (36). Still it is not clear, if there are other factors being involved, which might trigger the development of MGC in conditions like PE.

Cowden Dahl et al. discuss hypoxia-induced changes on molecular level that might be involved in MGC production (37). Especially hypoxia-inducible factors (HIFs) seem to be included in giant cell formation and differentiation (37). This means, that although hypoxia might be the main mechanism to trigger MGC production, many different pathways on molecular level seem to be of importance for the formation of MGC.

Stanek et al. reported a significantly increased number of MGC, not only in PE, but also in other high-risk pregnancies associated with this chronic uterine hypoxic injury pattern. These findings suggested the high value of MGC in explaining perinatal mortality and morbidity (29), because it might function as indicator for hypoxia.

The number of MGC was likewise found to be significantly higher in placenta previas by Biswas et al. (38), which might also be explained by hypoxia due to bleeding, inadequate attachment or changes in spiral arterioles, and therefore placental malperfusion. This would support the theory that placental hypoxia is the main trigger for MGC increase. Van Beekhuizen et al. on the other hand, reported a decreased number of MGC in retained placentas (placenta adhesiva) and concluded that the missing fusion of trophoblasts and hence the decreased number of MGC might be a factor involved in the pathology of placenta adhesiva (39). This finding implies, that MGC play a role in the regulation of placental adhesion and are needed in order to prevent the event of retained placenta whereas their abundance is a sign of placental hypoxia.

As Stanek et al. pointed out, MGC measurement could be used retrospectively to explain adverse pregnancy outcomes (29). Since the number of MGCs is also elevated in other pregnancy related complications than PE, it is not specific for this condition (38). Also it is not clear whether MGCs contribute to the development of PE or are they just an indicator of placental hypoxia.

Moreover this study showed a significant increase in the MGC/DC ratio in the studied group compared to control, but there was no statistically significant difference in the number of DC and EVTB between both groups, meaning that the number of DC and EVTB did not have any influence on the increased number of MGC.

This study was limited by a relatively small sample size (N=15 in each group). Also, it was difficult to distinguish between EVTB and DC on the basis of standard hemalaun and eosin (H&E) slides, so CK7 antibody staining was introduced to avoid possible errors.



## **6. Conclusion**

Concluding, this study suggests that there is an increased number of MGC in the decidua basalis in placentas from pregnancies with PE compare to placentas from normal pregnancies. This finding might be a useful marker in the workup of pregnancies that have been complicated by PE. The exact etiology of the formation of MGCs and possible contribution to the course of PE is yet to be determined.

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## **8. Summary (English)**

## **MULTINUCLEATED GIANT EXTRAVILLOUS TROPHOBLAST CELLS IN THE DECIDUA OF PLACENTAS COMPLICATED WITH PREECLAMPSIA**

**Objectives:** To compare the number of Multinucleated Giant Cells (MGC) in the decidua basalis of placentas complicated with preeclampsia (PE) and placentas from normal pregnancies.

**Subject and Method:** Fifteen placental samples from pregnancies complicated with PE and the same number of placental samples from normal pregnancies were taken. For each placental sample the number of MGC, extravillous trophoblast (EVTB) and decidual cells (DC) was counted in the decidua basalis. Immunohistochemistry with primary antibody CK7 was used to distinguish EVTB and MGC from DC. The MGC/DC ratio and MGC/EVTB ratio was also determined.

**Results:** MGC number was significantly higher in placentas complicated with PE. Likewise MGC/DC ratio was found to be significantly higher in the studied group. There was no statistically significant difference in EVTB and DC number as well as MGC/EVTB ratio.

**Conclusion:** Increased MGC number in the decidua basalis could be a morphological sign of PE and hypoxic conditions in the placenta.



## **9. Summary (Croatian)**

## **MULTINUKLEARNE ORIJASKE STANICE EKSTRAVILOZNOG TROFOBLASTA U DECIDUI POSTELJICA IZ TRUDNOĆA S PREEKLAMPSIJOM**

**Ciljevi:** Usporediti broj multinuklearnih orijaških stanica (MOS) u decidui basalis posteljica iz trudnoća s preeklampsijom (PE) i posteljica iz normalnih trudnoća.

**Materijali i metode:** U ispitivanu skupinu je uključeno petnaest uzoraka posteljica iz trudnoća s PE. Isti broj uzoraka posteljica iz normalnih trudnoća je činio kontrolnu skupinu. U svakom uzorku je određen broj MOSa, stanica ekstraviloznog trofoblasta (EVTB) i decidualnih stanica (DS) u decidui basalis. Za razlikovanje stanica EVTB i MOSa od DSa korišteno je imunohistokemijsko bojanje s primarnim protutijelom CK7. Određen je i odnos MOS/DS i MOS/EVTB.

**Rezultati:** Broj MOS je bio značajno veći u ispitivanoj skupini posteljica u odnosu na kontrolnu skupinu. Odnos MOS/DS je također bio značajno veći u ispitivanoj skupini posteljica. Nije bilo statistički značajne razlike u broju stanica EVTB, DS kao i u odnosu MOS/EVTB između dvije skupine.

**Zaključak:** Povišeni broj MOSa u decidui basalis jedan je od morfoloških znakova koji mogu ukazivati da se radi o posteljici trudnoće s PE, to jest o postojanju hipoksičnih uvjeta.

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