Histological evaluation of vitamin-D deficiency on the endometrial decidualization in early pregnancy in rat model

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ABSTRACT

Introduction: Vitamin-D is a pleiotropic secosteroid hormone that exerts classical effect in calcium uptake and bone metabolism. In addition, it has non-classical effects on variety of organs. Endometrial decidualization, placentation, and the development of the embryo itself have all pivotal roles for normal pregnancy outcomes after embryo implantation. Early maternal decidua and fetal trophoblast cells illustrate expression of vitamin-D receptor and 1α-hydroxylase enzyme that catalyzes vitamin-D into the active metabolite, calcitriol.

Aim: To record the histological effect of vitamin-D deficiency on endometrial decidualization during early pregnancy on day 8 post-coitum.

Materials and Methods: Thirty female albino rats were divided into 2 groups: control group (GpI) and vitamin-D deficient group (GpII). After establishment of pregnancy, the animals were sacrificed on the 8th gestational day. The bicornuate pregnant uteri were processed for histological analysis using H&E and PAS stains in addition to immunohistochemical staining for osteopontin (OSP), as a marker for decidualization, and for CD34 to assess the decidual angiogenesis. Morphometric and statistical studies were performed.

Results: Substantial decrease in the number of implantation sites in vitamin-D deficient rats was noted by gross examination. Histological examination revealed significant decrease in endometrial thickness at the antimesometrial region with remarkably impaired stromal cells differentiation. PAS staining illustrated fewer PAS positively stained uterine natural killer cells compared to control. Weak cytoplasmic OSP immunostaining was observed in the luminal as well as glandular epithelium, vascular endothelium and few stromal cells in mesometrial region. CD34 immunostaining demonstrated impaired angiogenesis in the lateral decidua as compared to control. These findings were confirmed by morphometric and statistical analysis.

Conclusion: Vitamin-D deficiency impairs the uterine stromal cells differentiation, affects the immunity tolerance and weakens decidual angiogenesis during early pregnancy in rat model.

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Key Words: CD34, decidualization, osteopontin, vitamin-D deficiency.

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INTRODUCTION

Endometrial decidualization, placentation, and the development of the embryo itself are all important for normal pregnancy outcomes after blastocyst implantation. Decidualization is a precise remodelling of the maternal uterine tissue to provide nourishment for the developing blastocyst before placental formation and to protect the embryo from being attacked by maternal immunity[1]. The endometrial decidualization includes terminal differentiation of stromal cells into decidual cells, uterine angiogenesis and abundant recruitment of immune cells especially uterine natural killer cells (uNK) and macrophages[2].

Rats have been effectively used as an appropriate model for studying the remodeling of uterine tissue during pregnancy and the physiological mechanisms controlling it, in contrast to human[3]. In rats, implantation occurs between days 4 and 5 of pregnancy, during which the uterus is receptive to blastocysts and known as window of implantation[4]. Decidual reaction in rat occurs in response to blastocyst implantation or artificial stimuli[5].

Decidualization of the stromal cells in the rat occurs in two stages. It first starts in the antimesometrial region on day 6 of pregnancy and reach its maximum development on day 10 then enters a period of regression until they are entirely absent by day 16. The second stage develops in the mesometrial region from 8th to 12th day post-coitum, and includes transforming the uterine stromal cells into the decidual cells[4]. Decidual angiogenesis is considered the initial fetal-maternal exchange apparatus. Thus, it is a crucial process for embryonic survival and maintenance of pregnancy[6]. Decidualization of each region is dependent on expression of hormone receptors, local factors and
growth cytokines mediated by progesterone\(^7\).

Vitamin-D is a major regulator of calcium and phosphate homeostasis. Calcitriol is the most potent metabolite of vitamin-D. It is also known as 1,25-dihydroxy-vitamin D3 (1,25(OH)\(_2\)D\(_3\)), which has been documented to possess variety of functions not immediately related to ion homeostasis\(^8\). Calcitriol has a key role in immune tolerance by regulating T-cell maturation into effector and regulatory T cells\(^9\). Progesterone substantially enhances the T-cell response to calcitriol. Thus, there is a clear synergistic corporation between vitamin D and calcitriol in regulating T cells and tolerating decidual immunity in pregnancy\(^10\). Noteworthy, early pregnant maternal decidua and fetal trophoblast cells can express both vitamin D receptor and 1α-hydroxylase enzyme that converts 25(OH)D\(_3\) into 1,25(OH)\(_2\)D\(_3\)\(^11\).

**MATERIALS AND METHODS**

**Animals:**

Thirty adult female Sprague Dawley albino rats, 4-weeks old bred in the Animal House of Ophthalmology Research Center were included in the study. All procedures were held under ethical guidelines for the use and care of animals. The animal protocol was approved by the Institutional Animal Care and Use Committee of Cairo University. Rats were divided into 2 equal number groups 15, each as followed:

- Control group (GpI) rats fed ad libitum a standard routine chow diet for 6 weeks before mating and throughout the 7 days of gestation. They were maintained in daylight controlled room with light/dark cycle 12:12 h. Normal serum blood level of vitamin-D (Vit-D) was assured before matting.

- Vitamin D deficient rats (GpII) were kept for 6 weeks before mating and throughout the 7 days of gestation on a Vit-D deficient diet (containing 20% lactose, 2% Calcium and 1.25% Phosphate). In addition, animals were housed under incandescent lighting to avoid Vit-D activation by ultraviolet rays. The model was verified by serum blood low level of vitamin-D (below 15 nmol/L) in all included animals before matting\(^13\).

**Establishment of pregnancy**

The sexual maturity of the experimental rats was attained by monitoring their estrous cycle using vaginal smear. Nulliparous 10-weeks estrous female rats were bred overnight with fertile males of the same strain (female to male ratio = 1:1) to allow matting. Morning examination was done every day for detection of the vaginal semen plug and sperms intermingled with the cornified cells in the vaginal smear that indicates day 1 pregnancy\(^13\). Continuous di-oesstrus vaginal smears confirmed the pregnancy\(^9\). Thirteen out of the fifteen rats get pregnant in GpI, while ten out of fifteen rats get pregnant in GpII. The dams underwent only follow up until the 8th day of pregnancy. Ten pregnant rats from each group were sacrificed and subjected to histological assessment.

**Histological Analysis**

All animals were anesthetized using intraperitoneal injection of thiopental sodium (50 mg/kg)\(^14\) and sacrificed at the 8th day of gestation. The uterus was exposed through laparotomy and the right and left pregnant horns were isolated, cleaned from attached fatty tissue. The pregnant uteri possessed scattering nodes (implantation sites), indicating decidualization. The uteri were transversely cut at the widest area of these nodes and placed in Bouin's solution overnight. Paraffin blocks were processed and 5-7μm thick sections were subjected to:

1- Hematoxylin and eosin (H&E) staining to determine structural changes\(^15\) and PAS staining to visualize the uterine natural killer cells (uNK)\(^16\).
2- Immunohistochemical staining using anti-osteopontin polyclonal antibody (Ab) (Thermo Scientific, RB-9097-R7) as a marker of decidualization\(^16\), and anti-cluster differentiation 34 (CD34) monoclonal Ab (Thermo Scientific, MA1-10202) as a marker for endothelial progenitor cells and endothelial cells of blood vessels to assess the decidual angiogenesis\(^9\). Application of the primary Ab was followed by incubation in humidity chamber for about 60 min at room temperature. Immunostaining was proceeded by the use of Ultravision detection system. Sections were co-stained with Meyer's hematoxylin to visualize the nucleus.

**Morphometry and Statistical Analysis**

The following parameters were accomplished:
1- The number of implantation sites in the uterine horns/rat in both groups was assessed by gross examination.
2- The thickness of the decidualized endometrium at the antimesometrial region in H&E stained sections at low power field (x40) using Olympus-DP27 digital camera (Japan). Ten readings were taken for each rat.
3- The number of uNK cells/ high power field (HPF) in PAS stained sections was counted in 10 non-overlapping randomly chosen fields/rat using Olympus-DP27 digital camera.
4- The area % of positive (+ve) osteopontin immune-staining in 10 non-overlapping randomly chosen HPF/ rat (x400) using "Leica Qwin 500 C" image analyzer (Cambridge, UK).

Quantitative data were summarized as means ±standard deviations (SD) and compared using T-test. The probability value <0.05 was considered statistically
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significant. Calculations were made on Statistical Package of Social Science software (SPSS), Version 19 (Chicago, CA).

RESULTS

Gross Examination

The uterine horns of Vit-D deficient rats (GpII) had significant less number of implantation sites in uterine horns (6± 0.81) compared to control rats (GpI) (11.4±0.97) [Figure 1], indicating impaired pregnancy in vitamin-D deficiency.

Hematoxylin and Eosin Staining:

The histological analysis of the decidualized endometrium demonstrated antimesometrium, mesometrium and lateral decidua [Figure 2]. In GpI, the antimesometrial region exhibited remarkable full differentiation of stromal cells into numerous large and small decidualized cells. In addition, the proliferative capacity of decidual cells was demonstrated by presence of many mitotic figures and dividing cells [Figure 3a and 3b]. While the mesometrial region showed smaller cells with irregular shape and a single nucleus. Large venous sinusoids containing leukocytes were noted in the lateral decidua [Figure 4a and 4b]. Vit-D deficiency rats (GpII) showed remarkably impaired stromal cells differentiation with significant decrease in the mean endometrial thickness (131.6±3.5) at the antimesometrial region compared to control animals (201.5±5.4) [Figure 3c and 3d]. Small irregular shaped stromal cells with single nucleus at the mesometrial region and small blood vessels with narrow lumen at the lateral decidua were observed [Figure 4c and 4d].

PAS Staining

Numerous uNK cells with PAS +ve reaction were detected in GpI. While Gp II exhibited significant decrease in the number of PAS positively stained uNK cells [Figure 5]. The mean number of counted cells/ HPF was 12.8±1.8 in GpI versus 4.3± 1.2 in GpII.

Immunohistochemistry Staining

On day 8 of pregnancy, OSP immunostaining in GpI showed strong +ve cytoplasmic localization in the luminal and glandular epithelium, vascular endothelium and numerous stromal cells in the mesometrial region. In addition, numerous decidual cells in the antimesometrial region with +ve strong cytoplasmic immunostaining were detected [Figure 6a and 6b]. GpII demonstrated weak cytoplasmic OSP immunostaining in the luminal and glandular epithelium, vascular endothelium and few stromal cells in the mesometrial region, in addition to few decidualized cells in the antimesometrial region [Figure 6c and 6d]. The mean area % of OSP immunostaining in GpII was significantly decreased as compared to GpI [Figure 7].

Immunostaining of CD34 showed numerous elongated dilated vascular sinusoids, closely related to decidual cells in the in the lateral decidua (the transition zone between mesometrial and antimesometrial region) of GpI. Impaired formation of decidual angiogenesis was observed in the GpII in the lateral decidua of GpII [Figure 8].

Fig. 1: Gross uterine morphology 8th day of pregnancy showing (a): The uterus of the control group showing eleven implantation sites in both horns, (b): The uterus of vit-D deficiency group illustrating one implantation site in the left horn and five implantation sites in right, in addition they appear invariably smaller.
Fig. 2: Photomicrograph of H&E stained section in the implantation chamber of control rat at 8th day of pregnancy showing different regions of the decidua (x original 40). (AM) Antimesometrial region, (M) Mesometrial region, (LD) Lateral decidua.

Fig. 3: Photomicrograph of H&E stained sections showing the antimesometrial region of the rat uterus at day 8 of pregnancy. GpI illustrating (a) thick endometrial decidualization (from the lumen to the myometrium), (b) obvious decidualization of the stromal cells into numerous large and small rounded decidual cells with abundant eosinophilic cytoplasm, large round vesicular nuclei and prominent 1-4 nucleoli. Dividing decidual cells (B) and mitotic figure (arrow) are observed. GpII exhibited (c) thinner endometrial decidualization with (d) impaired differentiation of most of stromal cells into decidual cells. Few cells are demonstrating vesicular nuclei with prominent nucleoli (a and c x40, b and d x400). Scale bar 200 µm and 20 µm.
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Fig. 4: Photomicrograph of H&E stained sections showing the mesometrial and lateral decidua regions of the rat uteri at day 8 of pregnancy (x400). Gp I: (a) the mesometrial region illustrates small cells with irregular shape and a single nucleus. (b) the lateral decidua demonstrates formation of large venous sinusoids (arrow heads) harboring leukocytes. Gp II: (c) the mesometrial region displays small irregular shaped and spindle shape cells with single nucleus. (d) the lateral decidua exhibited small blood vessels with narrow lumen. Scale bar 20µm.

Fig. 5: Photomicrograph of PAS stained sections of the rat uteri at day 8 of pregnancy (x400). (a) Gp I displaying numerous uNK cells with PAS +ve reaction (arrowhead). (b) Gp II exhibiting few PAS positively stained uNK cells. Scale bar 20µm.
Fig. 6: Photomicrograph of osteopontin immunostained sections in rat uteri at day 8 of pregnancy (x400). Gp I demonstrating (a): strong +ve osteopontin (OSP) immunostaining in the cytoplasm of luminal (curved arrow) and glandular (arrow) epithelium, vascular endothelium (bifid arrow) and numerous stromal cells (S) in the mesometrial region. (b) Numerous decidual cells in the antimesometrial region strongly expressed +ve cytoplasmic OSP immunostaining (arrowhead). Gp II exhibiting (c): weak cytoplasmic OSP immunostaining in the luminal (curved arrow) and glandular (arrow) epithelium, vascular endothelium (bifid arrow) and few stromal cells (S) in the mesometrial region. (d): Weak cytoplasmic OSP immunostaining is observed in few cells in the antimesometrial region (arrowhead). Scale bar 20µm.

Fig. 7: Quantification of the mean area % of OSP +ve immunostaining in the pregnant uterus at day 8 of pregnancy at: (a) the mesometrial and (b) antimesometrial regions. * Significant decrease as compared to control group.
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Fig. 8: Photomicrograph of cluster differentiation 34 (CD34) immunostained sections showing the lateral decidua of the rat uteri at day 8 of pregnancy. Gp I demonstrating: (a) numerous elongated well developed venous sinusoids (arrows) that appear in close contact with the decidual cells with (b) +ve CD34 immunostaining of endothelial cells (arrowhead). Gp II exhibiting: (c) poorly developed vascular sinusoids (arrows) displaying (d) +ve CD34 immunostaining of endothelial cells (arrowhead). (a and c x200, b and d x400). Scale bar 50 µm and 20µm.

DISCUSSION

The present work indicated that vitamin-D deficiency provides direct evidence of impaired pregnancy and decreased receptivity of the endometrium. This study succeeded in modeling vitamin-D deficiency in pregnant rats within 6 weeks before mating and throughout the 7 days of gestation which was verified by low serum blood level of Vit-D. Vitamin-D deficiency is defined by the Endocrine Society Clinical Practice Guideline as 25(OH)D less than 20 ng/ml (50 nmol/L), and vitamin-D insufficiency as 25(OH)D range between 21–29 ng/ml (52.5–72.5 nmol/L)\[17\]. Noteworthy, vit-D deficiency was substantially associated with decrease in achieving pregnancy following IVF trails\[18\]. Besides, vit-D deficiency is strongly documented as a key regulator in women fertility problems\[19\].

Histological analysis of the decidua in early pregnant control rats revealed striking full differentiation of stromal cells into numerous decidualized cells in the antimesometrial region. In addition, dividing cells and mitotic figures were observed. In rodents, decidualization is anatomically polarized process. Once blastocyst invasion, differentiation of the stromal cells of antimesometrial region is settled initially. It begins in the stromal cells from the subepithelial zone, then spread to the surrounding stromal cells. On 8th day of gestation, the decidual cells expand the whole thickness of the antimesometrial region\[4\].

Vit-D deficient rats showed remarkably impaired stromal cells differentiation and proliferation with subsequent significant decrease in the mean endometrial thickness at the antimesometrial region, which could affect the pregnancy outcome and even the embryo growth. Vitamin-D joins its nuclear receptor that forms a complex which controls transcriptions of more than 200 genes implicated in metabolism, anabolism and mineral homeostasis and cell cycle control\[20\]. Moreover, vitamin-D exerts pro-differentiation activity in a variety of cell types achieved by enhancing the expression of growth factors and cytokines. In female reproductive system, it stimulates the production of progesterone which mediates the uterine function decidualization\[21\].

The present work illustrated numerous uterine natural killer (uNK) cells with strong PAS +ve reaction in control rats. The uNK cells are the most prominent immune cells in the decidua during the first trimester of pregnancy, accounting >70% of all leukocytes in the decidua\[22\]. In rodents, 2 types of uNK cells are identified by PAS and Dolichos Biflorus Agglutinin (DBA) reaction. PAS+DBA− cells are involved in spiral arterial remodeling, while PAS+DBA+ cells are involved in spiral arterial remodeling, while PAS+DBA+ cells produce angiogenic factors\[23\]. In
on endothelial colony forming cells. Thus, vitamin-D promotes the formation of capillary-like structures\textsuperscript{26}. Maternal vitamin-D deficiency in mice exhibited decreased in the vascular diameter in the labyrinth zone, which is the highly vascular zone of the rodent placenta responsible for nutrient and waste exchange\textsuperscript{27}. In addition, vit-D deficiency was accused in reduction of angiogenic factors expression, like vascular endothelial growth factor (VEGF), which is normally highly up-regulated during early decidualization for induction of angiogenesis. Reduction of VEGF was observed in maternal vit-D deficient mice with adverse consequences for the progression of pregnancy, placentation and the health of offspring\textsuperscript{28}.

CONCLUSION

Vitamin-D deficiency substantially impairs the decidualization of the endometrium during early pregnancy in rat model by affecting the differentiation of the stromal cells into decidual cells. In addition, it influenced the OSP expression, the immunity tolerance and uNK cells recruitment and weakened the decidual angiogenesis.

CONFLICT OF INTEREST

There are no conflicts of interest.

REFERENCES


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الملخص العربي
التقييم الهستولوجي لنقص فيتامين د على تكوين الغشاء الساقطي لبطانة الرحم في بداية الحمل في نموذج الفأر
سماء سمير قمر
قسم الهستولوجيا الطبية ، كلية الطب ، جامعة القاهرة ، القاهرة ، مصر

الملخص:
فيتامين د هو هرمون سيكوستيرويد متعدد المظاهر الذي يمارس التأثير الكلاسيكي في امتصاص الكالسيوم واستقلاب العظام بالإضافة إلى ذلك ، له تأثيرات غير تقليدية على مجموعة متنوعة من أجهزة الجسم. لكل من تكوين الغشاء الساقطي لبطانة الرحم والمشيمة وتطور الجنين أدوار محورية تؤثر على نتائج الحمل بعد زرع الجنين. ولقد أوضح الغشاء الساقطي الامومي المبكر وخلايا الأرومة الغامضة الجنينية عن مستقبلات فيتامين د وأنزيم 1- ألفا هيدروكسيلاز الذي يحفز تحويل فيتامين D إلى مَئِيْضَ نشط، الكالسيترويل.

الهدف من البحث:
محاولة تسجيل التأثير النسيجي لنقص فيتامين د على الغشاء الساقطي لبطانة الرحم أثناء بداية الحمل في اليوم 8 بعد الجماع.

مواد وطرق البحث:
أدرج ثلاثون من أنثى الفئران البيضاء في الدراسة وقسمت إلى مجموعتين: المجموعة الضابطة (المجموعة الأولى) و مجموعة نقص فيتامين د (المجموعة الثانية). بعد حدوث الحمل ومتابعة الحيوانات في يوم الحمل الثامن، وتمت معالجة الرحم ثنائي القرن الحامل للتحليل النسيجي باستخدام صبغة الهيماتوكسلينإيوسين. وتمت المعالجة النسيجية باستخدام إنزيمات السيتوبلازمية للأستيوبونتين كدلالة على تكوين الغشاء الساقطي، ولسي دي 34 لتحديد تكوين الأوعية الدموية في الغشاء الساقطي.

النتائج:
 lokبظ إنخفاض كبير في عدد مواقع تكوين الغشاء الساقطي في الفئران التي تعاني من نقص فيتامين D من خلال الفحص البصري. وكشف الفحص النسيجي عن انخفاض ملحوظ ذو دلالة إحصائية في سمك بطانة الرحم في المنطقة المقابلة لمِسْراقُ الرَّحِم مع اختلال ملحوظ في تكوين الخلايا النسيجية. كما أوضحت صبغة السيتوبلازمية أن عدد أقل من الخلايا الفائقة الطبيعية الإجابة للبيأبة إس مقارنة بالمجموعة الضابطة. كما لوحظ ضعف الصبغة المناعية السيتوبلازمية للأستيوبونتين في الظهارة المبطنة للتجويف و للغدد والأوعية الدموية و عدد قليل من الخلايا النسيجية في منطقة مِسْراقُ الرَّحِم. أوضحت الصبغة المناعية لسي دي 34 ضعف في تكوين الأوعية الدموية في الغشاء الساقطي الجانبى مقارنة بالمجموعة الضابطة. تم تأكيد هذه النتائج من خلال القياسات المترية الشكلية والتحليل الإحصائي.

الاستنتاج:
أشارت هذه النتائج إلى أن نقص فيتامين D يضعف تمايز الخلايا النسيجية للرحم ، ويؤثر على المناعة ويضعف تكوين الأوعية الدموية أثناء بداية الحمل في نموذج الفأر.