ABSTRACT:
Endometriosis is a gynecological condition recognized by the existence of ectopic endometrial tissue outside the uterus. It is predominantly present in females of reproductive age group and is one of the main causes of infertility. Even with a predictable prevalence of 11% in females and considerable historical explanations adopted from the seventeenth century, the diagnosis of endometriosis still remains doubtful. The conventional concepts on histological basis of endometriosis are explained by a number of theories. Medical signs of endometriosis contain prolonged pelvic ache, dyspareunia, repeated menstrual discomfort and chronic pelvic pain which can severely affect the excellence of life and health of the patient. In this review we will discuss the prevalent theories for the diagnosis of endometriosis and suggestions to identify the condition well in time for better control and management.

Key words: Endometriosis, Endometriotic lesion, angiogenesis, vascularization, vasculogenesis, endothelial progenitor cells

INTRODUCTION:
Endometriosis is well-defined as the existence of endometrium in uncommon or ectopic position. Histopathologically, there is the existence of tissue or glands of the endometrial type external to the uterine cavity. This is a gynecological condition that depends on the hormones that are observed more frequently in child bearing age women. Frequency of endometrium is among 5% and 10% in premenopausal females and be able to extent up to 35% in females suffering of subfertility, as this might be main reason for infertility. The hazardous cause of endometriosis is menarche starting at the age of 11 years as well as prolonged and heavy periods. These two causes may raise the extra uterine environment for menstrual flow and the threat to endometriosis. The common locations for pelvic endometriosis are fallopian tubes ovaries, Douglas-fir pouch and uterine ligaments (broadly broad and uterosacral ligaments). Endometriotic embeds are also present outside the pelvis, i.e. diaphragm, lungs, gastrointestinal tract, pericardium and abdomen.

There are 3 main types of endometriosis present in pelvisperitoneal, ovarian, and infiltrating endometriotic disease. Structurally, there are 3 forms of endometriotic lesions: red lesions, white lesions and black lesions. Red are characterize by great vascularization, although white are at late stages of red lesions, assuming the development of fibrosis and inflammation. The black lesions have tissue breakdown and scarring with successive development of scar tissue. The histological basis of endometriosis are explained by a number of theories.

METHODOLOGY:
Literature search was carried out regarding the studies conducted on Theories of Endometriosis with the key words endometriosis, endometriotic lesions, angiogenesis, vascularization, vasculogenesis, endothelial progenitor cells. Through Literature searches were performed in PubMed, Medline and Google scholar for English articles. The searches included both animal and human studies (Figure 1)

Literature Review: The various theories were discussed in 43 articles for the diagnosis of endometriosis.

Molecular and Cellular Theories on Pathophysiology of Endometriosis
To date, the pathologic process of endometriosis is quiet debatable even with many years of investigations. Numerous concepts of pathology have been suggested in current years: i) implantation theory; ii) metaplasia theory; iii) induction theory; iv) Epigenetic theory; v) stem cell based theory; vi) Perineural theory. In recent times, it is suggested that an additional means of expressing the development and pain associated with endometriosis on the basis of inflammatory processes and initiation of nerve terminals to menstrual debris resultant retrograde and addition aluterinomenstrual flow of endometriosis.
i) Implantation theory:
The best frequently recognized implantation theory stands on hypothesis that small and early disease is recognized the consequent progression and invasion indicates towards advanced problem. These conserved endometriotic particle shave capability for attachment to the peritoneum, multiply, categorize, and attack the adjoining tissues. Additionally the distribution of endometrotic cells thru lymph nodes impacts over the source of disease at distant places such as cerebellar or thoracic endometrotis.

As per requirement towards sustenance of implantation theory a number of causes take place: i) existence of backward menstruation; ii) occurrence of sustainable endometrotic tissues to reversing refluxed menstrual flow; and iii) binding ability of endometrotial cells on peritoneum together with implantation and propagation.

The peritoneal fluid, (PF) is filled into the peritoneal cavity, variation of fluid resulting eg. macropage exudations, ovarian exudate, retrograde tubal fluid, transudate and reflux endometrotic material through reflux menstruation are therefore a significant component of the peritoneal surrounding. This conversion of fluid into the pelvic cavity may explain in part the anatomical distribution of endometrotic lesions, which corresponds well to the principles of implantation biology and is therefore favorable to the theory of implantation. In contrast, endometrotis is only seen in a subcategory of females, despite of detail that FP comprises endometrotic tissue in equal to 59% of patients, regardless of endometrotic lesion present or menstrual cycle at any stages. However, a persistent as well as increasing menstrual flow noticed in females having endometrotis might because of retrograde fluid into the pelvis compared to healthy females having patent fallopian tubes. Furthermore, the sealed uterine peristalsis might be the reason for the interruption of additional basal endometrium and, hence, growing extent of stem cell-sembling cells existing in retrograde flow of menstrum.

ii) Metaplasia theory:
The theory of metaplasia further more remains redirected in developing rest theory since poorly placed Mullerian/ endometrotic tissues might be stimulated on the way to suffer metaplasia. These data are corroborated by recent evidence that emigrant primary epithelium remains or endometrium-like ectopic glands are able to create beside fetal woman reproductive system sustainingas likely cause for endometrotic disease. Conversely, endometrotial lesions also take place at additional positions outer to the Müller canals.

iii) Induction theory:
The induction theory associates the theories of implantation and metaplasia and hypothesizes that unidentified materials released from the degenerating endometrium will induce the undifferentiated mesenchyme to form a tissue similar to that of the endometrium. In summing-up, the overhead concepts emphasis over the occurrence of endometrotial lesions however remain unsatisfactory to describe the happening of severe endometrotis. The progressive development of short-term disease to initial endometrotial lesions and severe types approaching benign cancers could remain elucidated by cellular alterations initiating after genetic or epigenetic modifications besides is treated into the theory of endometrotic lesion.

iv) Epigenetic theory:
In Support of the epigenetic concept, it is found that cystic ovarian endometrotis is of clonal origin then certain endometrotial particles are disturbing in vitro, related with loss of epithelial -cadherin appearance, a process commonly detected in cancer biology. Furthermore, there is a combined proposal of tendency of germ line to endometrotis. Family reunification of endometrotis in humans and rhesus monkeys moreover to the increased prevalence between blood relatives of females having all diseases, associated to the wide spread population has been described. In addition, the suffered non-twin sisters are similar in ageof onset for symptoms and show onisistancy in monozygotic twins. Furthermore, other risk factors, such as prolonged exposure of digoxins, may to oplay a part in etiology of disease. These comments may lead to the assumption that endometrotis is probably having complicated genetic trait where several genes work together and with the surroundings to produce the phenotype of disease. The endometrium have a tremendously regenerative power and this is not an amazing that endometrium has stem cell properties.

v) Stem cell-based theory:
After confirmation, that endometrotis may possibly be a stem cell-based situationstops after the statement that newly isolated endometrotial stromal and epithelial cells comprise a uncommon population of cells having clonogenic property fictional in colony-forming units. (CFUs)

The CFUs in endometrotial stromal fractions are analogous with mesenchymal stem cells (MSCs) with respect to their differentiation potential across several lineages. Development of endometrotial type MSC cells (eMSC) is likely through expression of PDGFR-Rb and CD146 perivascular cell markers. The epithelial and strontial cells Clonogenicity of the endometrium shows non-substantial predisposition dependent on stage of the menstrual cycle, with greater than before clonogenicity at the growing stage of stromal cells, at secretory stage of epithelial cells. CFUs can similarly be identified in the non-cycling endometrium. MSC inappropriate retrograde in the pelvis hence a serious element in forming an initial endometrotic disease. Further significance, the menstrual blood comprises plasticity particles, i.e. the re-forming cells of the endometrium (ERC). ERCs be similar to MSCs by their presence, growing...
possessions and prospective for differentiation into several cell categories. But, unlike MSCs, they direct matrix metalloproteases (MMP-3 and MMP-10), angiogenic factor ANG-2 and cytokines (GM-CSF, PDGF-BB) exposed by proteome investigations\(^{34}\).

The morphology of menstrual blood-derive MSC (discussed as MMCs or MenSCs) for example unique like as fibroblast and similar to bone marrow-derived MSCs\(^{35}\).

An additional research established the comprehensive plasticity of Men SCs\(^{36}\).

Generally, Men SC shave advanced rate of proliferation, clonogenicity and migration than angiogenic potential in vitro and bone marrow-derived MSCs in vivo studies\(^{37}\). Hida et al. confirmed the power of Men SC stowards there sortation developments in Myocardial Infarction rat model\(^{38}\). Now, Men SCs contributed in reestablishment for diminished cardiac physiology thrudistinguishing into Cardiomyocytes derived from Men SC over transplant position. Therefore, menstrual blood comprises plastic cells that provide a new basis for cell-based additional managements\(^{39}\). These outcomes obviously designate that back ward flow of menses be able to carry stem cell in the pelvic cavity and also there may be additional cell types using putative stem / progenitor cell properties. Investigations on blood and menstrual-derived plastic cells are still in its infancy. This is likewise the objective why numerous researchs describe the appearance of diver seimmuno phenotypic sketches of MenSC\(^{40}\). A consistent attitude to segregate and describe

*Figure 1; Flow Chart of Literature Review*

(figures 2) Interplay of theories of endometriosis adopted from PAB Klemm, A Starzinski - Powitz - Current Women's Health (2018)
stem cells in menstrual blood is important in deciphering their title role in the pathogenesis of endometriosis\textsuperscript{31}.

\textbf{vi) Perineural theory:}

It was suggested the perineural extent of endometriotic lesion with in inferior hypo gastric plexus, established concept of extent of endometriosis into nerve tissues in pelvic cavity\textsuperscript{15}. Meanwhile, further studies are demonstrated participation of nerves arising after lumbosacral plexus, as well as sciatic and obturator nerves\textsuperscript{42}. Newly revealed biological concepts of general importance (e.g. miRNAs, stem cell-based) are also significant to the pathogenesis of endometriosis\textsuperscript{32}. One challenge in endometriosis research will be to evaluate non-steroidal signaling pathways as targets for new therapeutics for the treatment of endometriosis. This may be an opportunity to substitute E2 depletion therapies to reduce undesirable effects.

\textbf{CONCLUSION:}

The molecular features of endometriosis include a hormone-dependent (estrogen-dependent, progesterone-opposition) and inflammatory state with an epi-genetic predisposition that is most likely driven by cells with plasticity.

\textbf{REFERENCE:}


