Endometrial Curettage When and for Who

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Abstract: In this study, the histopathological diagnoses pertaining to curettage material and the clinical necessity and results of endometrial curettage in age groups of patients in our region who had curettage due to abnormal uterine bleeding were studied. The histopathological diagnoses of patients with the above-mentioned criteria were scanned, excluding those related to pregnancy. 409 cases were re-examined and evaluated by a group consisting of three pathologists. Diagnoses were classified into 10 different groups. According to this classification system, out of the total number of cases, 100 were defined as endometrial polyp (24.4%), 72 cases as secretory endometrium (17.6%), 59 cases as proliferative endometrium (14.4%), 51 cases as stromal/glandular degeneration (12.5%), 32 cases as irregular proliferation (7.8%), 26 cases as insufficient (6.4%), 22 cases as chronic endometritis (5.6%), 18 cases as decidualization (4.4%), 17 cases as endometrial hyperplasia (complex atypical)/adenocarcinoma (4.2%), and 12 cases as others (inactive, atrophic, basal and menstrual endometrium) (2.9%). The average age of the 409 cases was 44.8±7.8. Upon evaluation of curettage due to abnormal uterine bleeding in the approximate 3 year period in our region, and taking into consideration the age distribution, it was observed that the clinical pre-diagnoses of patients especially with endometrial polyps and precancerous lesions were coherent with the corresponding histopathological diagnoses.

Keywords: Endometrium, Curettage, Histopathological Diagnosis, Age

1. Introduction

Abnormal uterine bleeding (AUB) is the most common reason for admissions to gynecology clinics after vaginal infections. Chronic endometritis, endometrial polyp, endometrial hyperplasia and adenocarcinoma are usually considered to be among the reasons underlying AUB (3).

Malignant or premalignant endometrial lesions are most commonly determined by AUB (3). Endometrial sampling is important for accurate histopathological diagnosis. Starting from delivery to pathology laboratory, the endometrial sample passes through a series of diagnostic processes. Both macroscopic and microscopic descriptions are valuable for histopathological diagnosis (1-13). The descriptions are as follows:

Insufficient term: Occasionally the material may only consist of blood and mucus or just fragments of endometrium surface epithelium. Curettage of menopausal or post-menopausal endometrium may be void of any tissues.

Decidualization term: Indicates transition of stromal cells from proliferative stage to cells with nuclear cytoplasmic boundaries, large oval and round vesicle nucleus, with eosinophilic or clear cytoplasm.

When this transition is less evident, it is referred to as predecidual stage and is an indication of the late secretory period of normal menstruation cycle.

Secretory endometrium: It has been stated that the most appropriate time for verification of ovulation is on the 22nd day of cycle or afterwards. The first reliable histological change specific to ovulation is observed at the 17th day of the cycle.

These changes are edema, folding of the spiral arterioles...
and stroma predecidualization.

Maximal stromal edema occurs on 22nd day.

Endothelial proliferation causes folding in the endometrium arterial system. In histological sections vascular groups are visible on the upper functional layer.

Stromal predecidualization begins on days 22-23. On the functional layer, it is on the spiral arteriole and capillary surroundings. Predecidual transformation of stromal cells below superficial epithelium occurs on 25th day. The joint decline of estrogen and progesterone on day 25 of the cycle fails to maintain the integrity of the lysosomal enzymes. As a result, lysosomal enzymes are released inside the cells and in the cavity between them.

Proliferative endometrium: Vascular endothelial cells, together with glandular and stromal cells are characterized by proliferation. This proliferation results in increased mucosa of the uterus. It consists of numerous mitoses. The normal proliferative glands of endometrium consist of oval-round-shaped glands that exhibit changes in diameter and fold according to early, middle or late stage and include nuclear pseudo-stratification and mitosis. As glands in early stage of proliferation advance to the middle and late stages, they acquire amoeboid and voluminous shape compared with straight tubules. In addition, the epithelial cells of surface epithelium and especially gland lumens gain cilia and microvilli. Lymphoid aggregates reminiscent of follicles may be seen in endometrial stroma, especially during the proliferative stage.

Endometritis: Endometrial inflammation is a pathology generally belonging to the fertility period, which can appear in infectious or non-infectious cases. Hence, it is actually a histopathological entity. Clinical correlation is not always possible. It is possible to classify endometrium inflammations as specific or non-specific. Non-specific endometritis are more commonly seen. Since some inflammation elements may be present in the endometrium stroma under normal conditions, the inflammatory cells of the endometrium must always be evaluated as a component of endometritis. Especially the granulocites of secretion phase and polymorphonuclear leukocytes of the menstrual phase must not be accepted as symptom of inflammation. In acute inflammation there is an infiltration consisting of polymorphonuclear leukocytes, which collect in gland lumens and form micro-abcesses and are also present in surface epithelium. The presence of PNL alone is not an evidence for inflammation as they are also present in the menstrual phase. In chronic endometritis there is an inflammation with a predominancy of plasma cells, lymphocytes and macrophages. Small amounts of polymorphonuclear leukocytes and eosinophils may be present.

Endometrial polyps: Since these polyps are seen as fragments in curettage material, it is sometimes difficult to distinguish them from normal endometrium and endometrial hyperplasia. Polyps appear as larger particles compared with normal endometrium, and are round-shaped particles surrounded by surface epithelium. The stroma is usually dense and includes thick-walled vessel sections. The glands do not have the uniform orientation of normal endometrium and have variable shapes and sizes. At times they exhibit the normal typical secretory or proliferative activity, in which case they are named as “functional polyps”.

In pathology reports, polyps are classified as
- adenomatous
- adenofibromatous
- adenofibromyomatous
- adenomyomatous

according to the appearance and ratio of glandular and stromal elements.

The polyps located at the lower segment and proximal endocervix sometimes have both endometrial and endocervical type epithelium on the surface or glands. These are called “endocervical-endometrial polyps”.

Endometrial Hyperplasia: The uterus is usually large. The curettage material fills 3 or more cassettes. Hyperplasia is usually a diffuse abnormality; it covers most of the endometrium and is often polypoid-occurring. Gland and stroma together may exhibit increase. In some cases, especially atypical hyperplasia, it can be a localized lesion. The atypical diagnosis is primarily based on nuclear features; most of them change from oval to round shape, become enlarged, irregularly-shaped, and have non-uniform membrane with 2-4 lines true stratification. There is irregular stratification and loss of polarity. There is variable mitotic activity. Chromatin forms clusters along the nuclear membrane and lead to the vesicular appearance characteristic of endometrial atypia. The nucleolus may be apparent. Since the atypia in hyperplasia is usually focal, the cytoplasmic cosinophilia may be the first symptom to warn the pathologist.

Endometrioid adenocarcinoma: Often occurs on the basis of endometrial hyperplasia. They are defined as endometrial carcinoma due to their resemblance to the glandular structure of normal endometrium.

Below items must be taken into account in differentiating endometrioid adenocarcinoma diagnosis from atypical hyperplasia
- Complex and cribriform pattern
- Epithelial bridge
- Bizzare-shaped glands
- Mitosis (sometimes atypical)
- Nucleus abnormalities: nucleomegaly, chromatin transparency, evident nucleolus
- Presence of luminal neutrophile

Moreover, evident desmoplastic stroma separating glands and gland masses can be observed in endometrioid adenocarcinoma.

Irregular proliferation: Developes as a result of prolonged, unfulfilled estrogen effect (similar to chronic anovulatory cycles). It is not a real premalign lesion.

Stromal/glandular degeneration: A non-specific symptom. It can also be observed in bleedings accompanying menstrual endometrium and different organic diseases. Endometrium areas that have remained healthy must be carefully examined.

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for specific pathological findings (such as polyps, hyperplasia and malignancy. Nuclear debris in stroma (apoptotic), fibrin thrombi, stromal collapse, stromal cell aggregates, papillary syncytial changes in glands, glandular impingement, nuclear debris in basal cytoplasm of glandular cells (apoptotic), storage of hemosiderin pigment, fibrosis and hyalinization occur.

Menstrual endometrium: Occurs on day 28 of cycle following glandular and stromal degeneration. Apoptotic bodies become evident in stroma and glands. Clustered stromal cell groups (stromal blue balls) gather together and are seen in the degenerated stroma.

Following degeneration, glandular cells take a form similar to micropapillary structure, this is defined as a regenerative change and called papillary syncytial metaplasia.

Inactive endometrium: An endometrium with sufficient thickness but failing to exhibit morphological characteristics toward hormonal stimuli, such as active proliferative or secretory activity. These morphological changes can be found in most menopausal and post-menopausal women with decreased ovarian hormones, which are not sufficient enough to start endometrial proliferation. It recalls proliferative endometrium, however, glands are usually parallel to surface epithelium, in contrast to perpendicular orientation in proliferative endometrium. They have pseudostratified nucleus without mitosis, and ciliated cells are rarely observable. The stroma has dense characteristic regardless of basal and functional distinction.

Atrophic endometrium: Consists of small atrophic tubules and cystic dilated glands scattered in fibrous stroma in wide spaces, usually exhibiting no proliferative activity in post-menopausal stage (13).

2. Findings

In this study, retroactive analysis of histopathological diagnoses of curettage materials obtained in 2011-2013 as a result of AUB was performed. The histopathological diagnoses were scanned, excluding those related to pregnancy, and 409 cases were obtained. The cases were re-examined and evaluated by 3 pathologists. These were then gathered under 10 different diagnosis headings in histopathological terms. According to this classification system, out of the total number of cases, 100 were defined as endometrial polyp (24.4%), 72 cases as secretory endometrium (17.6%), 59 cases as proliferative endometrium (14.4%), 51 cases as stromal/glandular degeneration (12.5%), 32 cases as irregular proliferation (7.8%), 26 cases as insufficient (6.4%), 22 cases as chronic endometritis (5.6%), 18 cases as decidualization (4.4%), 17 cases as endometrial hyperplasia (complex atypical)/adenocarcinoma (4.2%), and 12 cases as others (inactive, atrophic, basal and menstrual endometrium) (2.9%), (Graph 1). The ages of 409 cases were analyzed, with a minimum value of 20 and maximum value of 70, and average age of 44.8±7.8. The average age of each diagnostic group was calculated separately (Table 1). The difference between diagnostic groups and average age was found to be significant. Single direction variance analysis was used as $P=0.01$ test. The difference originated from the averages of groups 2, 6 and 9.

![Graph 1. Distribution of patients according to their diagnosis.](image-url)
The importance of its application for endometrial bleeding cases, and the unnecessity of its routine application in menometroragia cases, yet with the possibility of pregnancy complications.

Most of the endometrial samples are obtained from women in pre-, peri- or post-menopausal stage (3, 6). The traditional method is cervical dilatation and curettage. It is believed that this method is therapeutic and has a good sample forming property for histopathological evaluation (13).

In our study, the histopathological diagnoses were scanned, excluding those pertaining to pregnancy.

In the same study, chronic endometritis was detected in some patients, and in one patient epithelioid granuloma supporting tuberculosis was detected (1).

In conclusion, in the retrospective analysis of 409 cases carried out by Lasmar et al. (2) on hysterectomy patients, 14.8% of endometrial polyps with diameters larger than 15 mm hyperplasia were found to have hyperplasia, and therefore concluded that the increase in polyp diameter increases the possibility of hyperplasia development.

All 409 cases in our study were endometrial curettage material and 100 cases were endometrial polyp.

In a research carried out by Sweet et al. (6) on patients defined as AUB for various reasons, they have concluded that the most effective treatment of pre-menopausal women is hysterectomy.

Oğuz et al. (12) have detected that in 12 out of 58 hyperplasia patients, atypical endometrial hyperplasia has developed into adenocarcinoma.

In compliance with literature, since the possibility of complex atypical hyperplasia developing into adenocarcinoma is high, we have formed one of the diagnostic groups as hyperplasia/adenocarcinoma.

In conclusion, in the retrospective analysis of 409 cases and evaluation in light of age range, the highest age (70) was detected in the insufficient category, the second highest age (66) in stromal/glandular degeneration category, and the third highest age (63) in endometrial hyperplasia/adenocarcinoma and endometrial polyp categories. According to these findings, due to the average age being 44.8±7.8, it has been found that the clinical/histopathological diagnoses of endometrial curettage carried out on patients in reproductive age are in agreement with one another.

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