The Impact of Parity on the Number of Ovarian Cortical Inclusion Cysts

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Abstract

Objectives: Ovarian cancer is the most lethal gynecologic cancer. Understanding the origin and pathogenesis of epithelial ovarian cancers is one of the most challenging issues of gynecology. According to one of the theories, ovarian cortical inclusion cysts can cause epithelial ovarian cancer, but there are a lot of criticisms against the theory. In this study, the effect of oral contraceptive pills (OCP) on the number and type of inclusion cysts were investigated. Parity is a strong factor in the prevention of ovarian epithelial cancer. The base of this study is that if parity reduced the total number of cortical inclusion cysts (CIC) and tubal metaplasia, the theory would be strengthened.

Materials and Methods: One hundred twenty-five patients under the total abdominal hysterectomy and bilateral salpingo-oophorectomy (TAH & BSO) due to causes other than conflict ovaries were considered for study. The patients were categorized in three groups based on their parity. Slides related to normal ovaries were considered for study by pathologist and the results analyzed.

Results: The one-way analysis of variance (ANOVA) of total number of cysts revealed a statistically significant main effect, Welch’s F (2, 55.09) = 3.51, P < 0.05, indicating that not all groups had the same number of total number of cysts. Also, the results indicated that patients with 0-2 parity had a significantly higher total number of cysts than patients with 6 and over (P < 0.05). Moreover, the results of Games-Howell post hoc procedure indicated that patients with 0-2 parity had a significantly higher number of inclusion cysts than patients with 6 and over (P < 0.05).

Conclusion: Based on the results of this study, parity has significant effect on reducing total number of inclusions cysts, but number of tubal type inclusion cyst was not different.

Keywords: Cysts, Epithelium, Ovary, Parity

Introduction

Ovarian cancer is the fifth most common cause of death from cancer in women. The highest proportion of fatality per one case is seen among all genital cancers. A woman with probability of 1%-1.5% faces the risk of ovarian cancer at some point in her lives. The risk of dying from ovarian cancer per woman is about 0.5% (1).

The peak incidence of invasive epithelial ovarian cancer is age 56-60 years. This cancer is relatively uncommon in women under 45 years (2).

Unfortunately, so far no progress has occurred in screening and early diagnosis of the ovarian cancer and understanding its pathogenesis. Investigator believes that determination of the source of the malignancy can help in screening and early diagnosis. (2)

For many years the hypothesis that epithelial ovarian cancer originates from ovarian cortical inclusion cysts (CIC) is lived. According to this hypothesis, cells—from ovarian surface epithelium (OSE)—that line these cysts multiply and mutate in response to bio-active factors and oncogenes (3).

Another interesting theory in recent decades is that serous ovarian cancer originates from tubal intraepithelial neoplasia (serous tubal intraepithelial cancer, STIC). Despite the overwhelming evidence in favor of the second theory, investigators failed to put the OSE theory aside (4). In this connection, there is much evidence; for example, metaplastic changes in epithelial inclusion cysts, from one type of epithelium to another, which can predispose to neoplastic changes (5).

Ovarian cancer is associated with infertility and low parity. The most important factors are the duration of fertility and reproductive history (6). Every pregnancy reduces the risk of ovarian cancer by about 10%.

Protection against ovarian cancer is one of the main advantages of oral contraceptive pills (OCP). The risk for all histologic type of ovarian cancer is reduced by 40% in OCP users compared with women who do not use this method. Many of these risk-reducing factors such as OCP and parity verify frequent ovulation hypothesis as etiology of ovarian cancer. According to this hypothesis, the main cause of ovarian cancer is dysfunction in the process of repairing the surface epithelium (7).

 Accordingly, the effect of parity on the number of inclusion cysts and tubal metaplasia of the cyst are investigated.
It is believed that if cortical inclusion cyst is the source of epithelial ovarian tumors, high number of pregnancies would reduce the number of cysts and reduce tubal metaplasia. Three theories have been proposed in regard to the origin of ovarian cancer:

1) The theory of OSE: According to this theory, OSE is origin of most ovarian stromal and epithelial cancers and can be differentiated to tissues with Mullerian types and create ovarian epithelial cancer (8). When culturing purified human OSE, it is converted to aggressive neoplasms similar to high-grade and low-undifferentiated grade serous carcinomas (9).

2) Theory of STIC: In this theory, the origin of high-grade ovarian serous tumors is malignant lesion in the fimbrial epithelium (10).

3) The theory of junction location of tubal fimbria epithelium with peritoneum (tubo-peritoneal junction [TPJ]): This location is the hot spot of the female genital tract for the generation of tumors (11).

The purpose of this study is to evaluate the effect of the number of pregnancies as one of the reducing decisive factors for ovarian cancer on the number of inclusion cysts in superficial epithelium as the source of ovarian tumors.

Materials and Methods
The study is descriptive analytic. Women who underwent hysterectomy and oophorectomy in Zahra hospital during 2011-2013 were included. For this purpose, the hospital records of patients who underwent hysterectomy and oophorectomy from any cause had been studied and those with normal ovaries were selected. The questionnaire was filled for each patient according to the data. In the case of incomplete records they were called. Patients with accurate information about the age, number of pregnancies, number of delivery and type of contraceptive method and were divided into three main groups based on the type of contraceptive methods:

1. Patients who had 2 pregnancies over 20 weeks.
2. Patients who had between 3 and 5 pregnancies over 20 weeks.
3. Patients who had at least six pregnancies over 20 weeks.

Patients who had normal ovaries in the final pathology report and their ovarian size were normal were selected. The paraffin blocks of patients’ ovarian tissue were obtained from the pathology archive of the hospital; from the paraffin blocks, sections with a thickness of 5 micrometers were prepared. Then, after consolidating the tissue in neutral buffered formalin and staining with hematoxylin and eosin, routine histological examination was carried out.

Slides obtained by a woman pathologist were reviewed to determine the presence of ovarian surface epithelial in obtained slices, the number of ovarian cortical cysts and existing tubal cell types in cyst epithelium and OSE. The pathologist, during examination of the slides, was not aware of the group of each patient.

For analysis of morphological results in the experimental

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The Effect of Parity on the Number of Tubal Cysts

Due to the nature of the data that it had over distribution, the assumption of homogeneity of variance was violated (P=0.001), the Welch's F test was used. The results of one-way ANOVA revealed a statistically significant differences between three groups in terms of number of inclusion cysts, Welch's F (2, 55.89) = 3.48, P<0.05. As depicted in Figure 3, the results of Games-Howell post hoc procedure indicated that patients with 0-2 parity had a significantly higher number of inclusion cysts than patients with 6 and over (P<0.05).

Discussion

Ovarian cancer is the fifth most common cause of death due to cancer in women. Ovarian cancer is called “silent killer,” as it has no symptoms and is detected at advanced stages (1).

The discussion about the pathogenesis and origin of epithelial ovarian cancer is still one of the hottest and most challenging topics in pathological obstetrics. Fortunately, in recent years, there have been major advances in this matter which can improve our understanding of the pathogenesis of ovarian cancer and it can certainly lead to advances in prevention, screening and early detection of this disease. The question is that whether ovarian cancer and in particular High Grade Serous Carcinoma (HGSC) originate from OSE, CIC and/or fallopian tubal epithelium (especially fimbria).

Until recently, the prevailing view was that these tumors originate from OSE, malignant transformation in OSE or CIC. In fact, despite the fact that these tumors are mainly epithelial, recently, histological studies on HGSC present compelling evidence that these tumors originate from the fimbrial mucosa of fallopian tubes. Briefly based on STIC theory, the high grade ovarian serous cancer HGSC are not primary ovarian neoplasms, but metastases from fallopian tube to the ovary. Although there is much evidence in favor of this theory, but STIC is associated with only 60% of these neoplasms and the question arises that where is the origin of remaining 40% (12)?

We actually planned to answer this question. In fact, we believe that even though there are arguments in favor of origin of HGSC from STIC, but the OSE and ovarian CIC as a source of additional and perhaps important neoplasms cannot be ignored. If CIC are the origin of ovarian epithelial cancer, proved risk factors for ovarian cancer should increase the number of cysts and increase the tubal metaplasia in cysts and in OSE. This attitude can be subject for extensive research to find the origin of this malignancy. It is not so far-fetched that age increases the numbers of ovarian inclusion cysts, as well as an increase in tubal type cysts.

Banet and Kurman in a study in 2014 analyzed the relationship between aging and the number of CIC in the ovaries. They used PAX8 as an indicator of Mullerian (tubal type cells) and calretinin as mesothelial indicator (Flat cells). They did not find CIC with tubal type epithelium in pre-menarche girls. They found a significant correlation between tubal type CIC that increases with aging. These findings mean that the existence of two types of CIC (one flat epithelium and the other one with columnar and tubal view epithelium) and the absence of CIC with tubal epithelial in the ovaries of the girls before menarche give ground for following finding: ciliary pattern of CIC is formed over time with the process of ovulation and im-
planted fimbrial cells in ruptured area of OSE during ovulation (13). If the findings of Banet and Kurman were correct, then, parity must decrease the number of inclusion cysts and especially, the cysts with tubal epithelial lining. In our study the total number of inclusion cysts in group with the parity number more than 6 inclusion cysts was significantly lower. But number of tubal type inclusion cysts was same. Auersperg et al showed that OSE normal cells, express markers of stem cells. The OSE in response to changes in the microenvironment and different growth factors can convert into different types of cells (14). The process of metaplasia from flat or cubic cells of CIC and OSE to cylindrical cells similar to fimbria epithelium is approved by IHC not only morphologically but also functionally (15). The culturing of OSE cells in the culture medium with endometrial stroma and estrogen cause endometrioid metaplasia in OSE (16).

**Conclusion**

Based on the results of this study, ovarian CIC were significantly lower in the group with more than 6 parity. Types of tubal inclusion cysts were not significantly different between the 3 groups. Ovarian surface epithelium in all groups was same. Normal and tubal type OSE was not significantly different. Our study was based on morphological features only. For a better survey we should assess tissue specific markers such as calretinin and PAX8. In the current study we included patients who use OCP for 3 years or less than 3 years. Due to the low number of samples, the number of cysts in both groups showed no significant differences, probably because we had samples (patients) that had used OCP for less than 5 years and this distorted the study finding. Another study should be arranged over samples (patients) who have not used OCP in order to look closely at the impact of parity on the OSE.

**Ethical Issues**

Since the research and laboratory works were performed over pathology samples after surgery, there was no need to get informed consent. No additional cost was imposed to the patients. All information about patient records and phone calls was kept confidential.

**Conflict of Interests**

The authors declare no conflict of interests.

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**References**


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