Chronic inflammation in endometriosis and endometriosis-associated ovarian cancer: New roles for the “old” complement pathway

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Immune escape is consequential for cancer development. Identifying abnormalities of the immune microenvironment during early carcinogenesis can provide insight into disease pathogenesis and unravel new preventive or therapeutic targets. We recently conducted a comprehensive immune gene expression analysis in endometriosis and endometriosis-associated ovarian cancer and explored new mechanistic roles for the complement pathway.

Introduction

Despite concerted efforts in the last decade, the rate of ovarian cancer mortality has only improved marginally. The majority of patients continue to present with late-stage disease, typically associated with low 5-year survival. The lack of effective screening procedures for early detection is partly due to the highly diverse pathology of ovarian epithelial tumors. For several types of cancer (cervical, colon, pancreas, breast, and prostate), the premalignant lesions are well defined (human papillomavirus-associated cervicitis, colon polyposis, pancreatitis, breast intraductal neoplasia, and prostate intraepithelial neoplasia, respectively) and prevention and/or early detection of these conditions can be designed and implemented.1 Consequently, it is estimated that the screening and treatment of the above precursors prevent a significant number of cancers each year, although some approaches (e.g., in cervical cancer) are more effective, whereas others (such as in prostate cancer) still struggle to prove efficacy. Notably absent from the list is a precursor for ovarian cancer, the most lethal gynecologic malignancy.

Several studies in the past decade suggest that endometriosis, a commonly diagnosed, chronic benign inflammatory disease, may predispose to endometrioid, clear-cell, and low-grade serous ovarian cancers, also termed endometriosis-associated ovarian cancer (EAOC).2 Endometriosis consists of endometrial-like tissue (glandular epithelia and stromal cells) ectopically implanted outside the uterine cavity and is usually treated through a combination of hormone therapy and surgical excision of lesions. Endometriosis implants develop in a highly inflammatory milieu and are often recurrent, resistant to treatment, and debilitating. Furthermore, these lesions have attributes that resemble ovarian cancer, including multifocality, genetic instability, increased proliferation, and capacity to invade surrounding tissues. In advanced stages, epithelial cells in endometriotic lesions display signs of “atypia,” defined by histological criteria such as cellular crowding and large nuclei with moderate to marked pleomorphism. Often found adjacent to a tumor, atypical endometriosis (AE) may represent the transitioning entity from benign lesions to malignant variants.3

In response to the invasive and sometimes recurrent endometriotic lesions, the host’s immune system often makes ineffective attempts to “repair and renew” that are believed to contribute to the transition from endometriosis to EAOC. Thus, a better understanding of the immune microenvironment in precursor lesions may provide mechanistic insight to EAOC pathogenesis. To date, cellular and molecular characterization of the endometriosis-associated inflammatory milieu has mostly focused on circulating chemokines and peritoneal immune cells (macrophages, natural killer [NK] cells).4 We recently reported results from the first comprehensive immune gene expression analysis of pelvic inflammation in benign endometriosis, AE, and EAOC.5 Our 120-case cohort revealed gene expression changes associated with each disease state. Benign endometriosis cases displayed a mixed immune gene profile, with one third of the patients showing immune responses more similar to cancer. However, when analyzing the AE cases, we observed that the majority (85%) of patients had a cancer-like immune environment further supporting AE as a tumor precursor lesion (Fig. 1).

Using differentially expressed genes in all disease categories, we identified complement as the most significantly

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Using this preclinical model, we explore this mechanism, we employed tri-immune stoma interaction. To further a potential mechanism of epithelium–dent, complement-mediated immunity as immunity, suggesting antibody-dependent alterations in serous ovarian cancer, suggesting that complement involvement may be specific to endometriosis and EAOC. We also identified activation of humoral immunity, suggesting antibody-dependent, complement-mediated immunity as a potential mechanism of epithelium–immune stroma interaction. To further explore this mechanism, we employed triple transgenic mice that progress to EAOC closely mirroring the human disease. Using this preclinical model, we demonstrated that complement upregulation in epithelial cells occurs early in carcinogenesis. However, exposure of complement-expressing ovarian tumor epithelial cells to in vivo-generated antibodies that can recognize and bind to the cell-surface mucin 1 (MUC1) tumor antigen does not trigger antibody-induced, complement-mediated cell death, demonstrating that, at least in this system, complement engagement is a rather ineffective tumor lytic mechanism. Furthermore, inhibition of 1 of the effector complement genes (C7) triggered suppression rather than stimulation of tumor cell proliferation, generating the hypothesis that complement activation, despite antibody presence, may bolster tumor growth (Fig. 1). This hypothesis has been supported by a series of recent studies that demonstrate a tumor-supporting role for complement, via several mechanisms that coexist in the tumor environment: directly, by stimulating tumor cell proliferation; or indirectly, by stimulating immune suppression and neovascularization.

Currently, there are no standardized screening methods for endometriosis or early ovarian carcinogenesis. Serum CA125 continues to be the mostly widely used protein biomarker, despite its limited specificity and sensitivity. Large studies on transvaginal sonography have failed to provide support for this approach. As a concerted effort to identify molecular abnormalities during the transition from endometriosis to EAOC, we have profiled micro-RNA (miRNA) expression in tissue (manuscript in preparation) and in patients’ plasma, and determined that plasma miRNA expression signatures could be used as biomarkers for endometriosis-to-EAOC progression. The immune gene expression profile further demonstrates that cancer-like signatures may develop early in patients with precursor lesions. Given that immune effectors can distinctly sense early molecular changes of cells undergoing transformation, it is conceivable that profiling the immune response in tissue may provide molecular clues for patients with EAOC risk and new targets for preventive or therapeutic vaccines. We hope our hypothesis will stimulate further studies, in larger cohorts, that can validate (or disprove) the clinical applicability of immune profiles as biomarkers and of the complement pathway as a potential therapeutic target.

Disclosure of Potential Conflicts of Interest
No potential conflicts of interest were declared.

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