MINI-REVIEW

Molecular Therapy as a Future Strategy in Endometrial Cancer

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Abstract

Of all gynecologic cancers, endometrial cancer is the most common cancer in the US and Europe. In addition, it is presently the second most common gynecologic cancer in the world. As a result of increasing menopausal, obese and tamoxifen use women, the incidence of the cancer seems to be on the increase. Surgery is the major treatment, whereas postoperative radiation therapy in high-intermediate risk patients many prevent locoregional recurrence. Adjuvant chemotherapy can improve progression free survival in advanced or recurrent cancers. Molecular targeted therapies are now a focus of attention including anti-vascular endothelial growth factor (VEGF), mammalian target of rapamycin (mTOR) inhibitor and tyrosine kinase inhibitor (TKI). They may provide useful future strategies for control of endometrial malignancies in developing countries and across the world.

Keywords: Endometrial cancer - molecular therapy - VEGF - mTOR - TKI

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Introduction

Incidence of endometrial cancer has been growth since the world menopausal women populations were increasing when compared to that in the year 2000 (Parkin et al., 2001). In addition, there are risks of the cancer following obeities and tamoxifen users (Bergman et al., 2000) which are expanding. The treatment strategies for endometrial cancer have to be more focuses on effectively eradicating of the cancer and minimizing adverse effects. The latest world report shows that there were more than 280 thousands new cases of endometrial cancer in 2008 which was the second most common gynecologic cancer (Ferlay et al., 2010). The age specific incidence rate was 8.2 per 100,000 women-year in 2008 (Ferlay et al., 2010). It was found more than 73 thousands deaths from the cancer in that year. In US, 47,130 estimated new endometrial cancer cases, based on 1995-2008 incidence rates, are expected to be diagnosed with 8,010 deaths in 2012. In addition to the first rank among gynecologic cancer in US, it was the third rank followed cervical cancer and ovarian cancer in developing countries as well as in Thailand (Thanapprapasr, 2010). Most cases were diagnosed in early stages and good prognoses. However, almost thirty percent of cases were regional or distant diseases at the time of diagnosis (Howlader et al., 2012), and some resisted to the contemporary therapy. In addition of a high five-year survival rate in all stages in the US report (81.8%) (Howlader et al., 2012), it was a low rate in developing countries (67%) (Parkin et al., 2005).

Current technique of an accurate staging system and surgical technique approach could be primary managing strategies. Benefits from evidences of adjuvant radiation approach and chemotherapeutic strategies in selected patients give the better outcomes. For the best goal, therapeutic advanced in molecular biology and targeted therapy are evidence in preclinical studies, both phase I and some of phase II trials. There are going to be evidence in phase III trials. Accumulating evidences and discussions are herein. To move forward passing through the tragic endometrial cancer together are the aims.

Endometrial Cancer Treatment Situations

Particularly, seventy percent of patients had localized diseases within the uterus, and twenty percent of them had regional diseases (Howlader et al., 2012). Surgery is the mainstay of treatment. Surgical staging is performed in all early diseases, while cytoreductive surgery was operated in the patients with advanced diseases. Radical hysterectomy was suitably performed in patients with cervical involvement. New FIGO (International Federation of Gynecology and Obstetrics) staging 2009 (Pecorelli, 2009) is well included in managing strategies in Thailand and others.

Since one-fifth of patients were premenopause, conservation of hormone production from their normal appearance ovaries in patients with early stages and low grade tumor were reported. There were reported no worse overall survival in ovarian preservative patients than the patients who had bilateral salpingo-oophorectomy during their surgical staging (Richter et al., 2009; Wright et al., 2009). In contrast, two reported cases, who were preserved their ovaries, had subsequently diagnosed

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ovarian carcinoma (Zivanovic et al., 2009). In conclusion, ovarian preservation is a non-standard strategy, it is still controversy. Post-operative radiation therapy decreased the loco-regional recurrence of the patients with high-intermediate risk (Creutzberg et al., 2000; Keys et al., 2004). Adjuvant chemotherapy improved progression free survival in advanced or recurrent cancer (Johnson et al., 2011). Nevertheless, the systematic review shows unchanged overall survivals (Johnson et al., 2011). Platinum based regimens give effective alternatives or added to radiation therapy for patients with high-risk, advanced or recurrent diseases. Concurrent chemoradiation therapies are ongoing evidence of Post-Operative Radiation Therapy in Endometrial Cancer (PORTEC) 3 trial and Gynecologic Oncology Group (GOG) 258 trial. Future and trends of molecular targeted therapies are values and of the most interest strategies especially in patients with persisted, advanced or recurrent diseases.

**Molecular Therapy**

Therapeutic advances in women’s cancer are molecular therapies. These could be directly cytotoxic cancer cells, omitting or at least adverse side effects. Each patient has different molecular signature of the cancer cells. In addition, molecular therapy may play role as individualized therapy for each of them.

Over ten percent of gynecologic cancer related patients died annually, in despite of the most effective cytotoxic, surgical and radiation strategies currently. Large numbers of toxicities were also from these. Drugs against molecular pathways directly to cancer cells’ survival should be targeted. Understanding biology of cancer cells could lead to developing novel effective treatment strategic approaches against them (Thanapprapasr et al., 2012). Preclinical studies show different genetic materials and pericellular proteins between normal tissues and cancerous tissues (Shahzad et al., 2011). There were revealed some proteins which effectively blocked the cancerous cascades. These lead to pharmaceutical designs and developing novel drugs. Currently, phase I, II trials have progressed on several studies.

The two types of endometrial cancer are classified on different characteristics (Bokhman, 1983). The type I clinicopathological characteristics are endometrioid carcinoma histology and tumor grading 1 or 2. Nonendometrioid carcinomas (serous or clear cell carcinoma) are type II endometrial cancer. Most common endometrial cancer is type I, which almost have Phosphate and tensin homolog on chromosome ten (PTEN) mutations. PTEN is a tumor suppressor gene which controls Phosphotidylinositol-3-kinase-Serine/threonine-specific protein kinase (PI3K-AKT) pathway in normal cells. Mammalian target of rapamycin (mTOR) is a protein in the cytoplasm of cancer cell which acts as an antiapoptotic factor by inhibiting G1 arrest action responsibly following AKT action (Figure 1). The inhibitor (temsirolimus) was developed and studied (Oza et al., 2011). The inhibitor was given as a single agent to the patients with recurrent or metastatic chemotherapy-naive or chemotherapy-resisted endometrial cancer, 25 mg intravenously weekly in 4-week cycles. It had results in some responses including 4% partial responses and forty-eight percent stable diseases in the patients with chemotherapy-resisted endometrial cancer (Table 1). The adverse events in temsirolimus used patients were not severe including fatigue, rash, mucositis, and asymptomatic pneumonitis (42%). Hematologic adverse events were generally mild, and the most common was lymphopenia.

Everolimus, an oral mTOR inhibitor, was studied and reported in phase II trial in patients with measurable recurrent, endometrioid histology, endometrial cancer who had failed at least 1 prior chemotherapeutic regimens (Slomovitz et al., 2010). A dose of 10 mg daily for 28-day cycles was administered until disease progression or toxicity. Twenty-eight patients were evaluated. Twelve (43%) patients had not developed disease progression at the time of the first objective evaluation (8 weeks). All these patients had stable disease (median, 4.5 cycles; range, 2-10 cycles). They discontinued treatment because

![Figure 1. Pathways of Molecular Targeted Drugs in Endometrial Cancer Cells and Cancer Related Endothelial Cells](image-url)

**VEGF-A=Vascular endothelial growth factor A. VEGFR=Vascular endothelial growth factor receptor. mTOR=Mammalian target of rapamycin. EGFR=Epidermal growth factor receptor. HER-2=Human epidermal growth factor receptor 2. TK=Tyrosine kinase. FAK=Focal adhesion kinase. PI3K=Phosphotidylinositol-3-kinase. PIP2=Phosphotidylinositolphosphate-2. PIP3=Phosphotidylinositolphosphate-3. AKT=Serine/threonine-specific protein kinase. PTEN=Phosphate and tensin homolog on chromosome ten. EZH2=Enhancer of zeste homolog 2. Dll4=Delta-like ligand 4**
Table 1. Actions and Responses of Molecular Targeted Drugs in Phase II Trials of Endometrial Cancer

<table>
<thead>
<tr>
<th>Molecular targeted drugs</th>
<th>Number of patients</th>
<th>Actions</th>
<th>Responses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bevacizumab (Aghajanian, Sill et al. 2011)</td>
<td>52</td>
<td>Anti VEGF-A</td>
<td>1CR, 6PR</td>
</tr>
<tr>
<td>Temsirolimus (Oza, Elit et al. 2011)</td>
<td>25</td>
<td>mTOR inhibitor</td>
<td>1(4%)PR, 12(48%)SD</td>
</tr>
<tr>
<td>Everolimus (Slomovitz, Lu et al. 2010)</td>
<td>28</td>
<td>mTOR inhibitor</td>
<td>12(43%)SD</td>
</tr>
<tr>
<td>Erlotinib (Oza, Eisenhauer et al. 2008)</td>
<td>32</td>
<td>EGFR inhibitor</td>
<td>4(12.5%)PR, 15(47%)SD</td>
</tr>
<tr>
<td>Trastuzumab (Fleming, Sill et al. 2010)</td>
<td>33</td>
<td>Monoclonal anti HER-2 antibody</td>
<td>12(36%)SD</td>
</tr>
<tr>
<td>Sorafenib (Nimeiri, Oza et al. 2010)</td>
<td>40</td>
<td>RTK inhibitor</td>
<td>2(5%)PR, 17(42.5%)SD</td>
</tr>
</tbody>
</table>

*VEGF=A-Vascular endothelial growth factor. mTOR=Mammalian target of rapamycin. EGFR=Epidermal growth factor receptor. HER-2=Human epidermal receptor
2. RTK=Tyrosine kinase receptor. CR=Complete response. PR=Partial response. SD=Stable disease

Antiangiogenesis Therapy

Since before 1971, it has been recognized that one is almost forced to the conclusion that there is, associated with the viable growing tumor, some blood vessel growth stimulating factor (Folkman, 1971; 1990; Folkman et al. 1971). Vascular endothelial growth factor (VEGF) is the major angiogenic agent for endothelial cell proliferation in endometrial carcinoma (Sivridis, 2001). Bevacizumab, a recombinant humanized monoclonal antibody anti-VEGF-A, is approved by the U.S. FDA (Food and Drug Administration) for metastatic colorectal, non-small cell lung, renal cell, and breast cancers. In gynecologic cancers, it is considered in the patients with ovarian cancer who have progressed on chemotherapy. The most common drug-related side effect is hypertension. Bevacizumab is well tolerated and active based on progression free survival (PFS) at 6 months in recurrent or persistent endometrial cancer and warrants further investigations. Phase II, GOG 229 trial recruited 52 patients with persistent or recurrent endometrial cancer previously receiving one or two prior cytotoxic regimens (Aghajanian et al., 2011). Bevacizumab at a dose of 15 mg/kg intravenously every three weeks were given until progression or prohibitive toxicity. No gastrointestinal perforations or fistulae were reported. Seven patients (13.5%) experienced clinical responses (one complete response and six partial responses). Median response duration was six months. Twenty-one patients (40.4%) survived with progression free of the tumor for at least 6 months. Median progression free survival (PFS) and overall survival (OS) times were 4.2 and 10.5 months respectively. Actions and responses of these molecular targeted drugs in phase II trials of endometrial cancer were summarized in Table 1.
of VEGFR reacted to VEGF-A ligand, may have activity as in ovarian cancer (Coleman et al., 2011). More targets other than VEGF (vascular endothelial growth factor) are in attentive studies. Focal adhesion kinase (FAK) (J. N. Bottsford-Miller; Thanapprapasr), Enhancer of zeste homolog 2 (EZH2), Delta-like ligand 4 (DLL4) are active molecules for angiogenesis and cancer growth (Thanapprapasr et al., 2012) (Figure 1). Agents against these targets may be challenged and worthwhile. A number of preclinical studies would be further conducted.

**Future and Trends**

Advanced in the future, it should be further evaluate studies on possibility of molecular therapeutic advantages. Antiangiogenesis is one of the pivotal strategies. Addition to current chemotherapy regimens (Korets et al., 2011), these molecular targeted drugs may further improve tumor responses. Either neoadjuvant molecular therapy, concurrent with radiation therapy or sequential adjuvant postoperative therapy in advanced and recurrent cancer would be reveal in further studies. Synergistic effects from combined antiangiogenesis and other molecular targeted therapies are revealed in preclinical studies (Li et al., 2012) and ongoing trials. All these targeted drugs have been review(Thanapprapasr, 2013). Thus, these would also be possible in further clinical trials.

**Conclusion**

Staging surgery is the major treatment, whereas postoperative radiation therapy in high-intermediate risk patients decreased their locoregional recurrences. Adjuvant traditional chemotherapy improved progression free survival in advanced or recurrent cancer. Molecular targeted therapies are in focus including anti-vascular endothelial growth factor (VEGF), mammalian target of rapamycin (mTOR) inhibitor and tyrosine kinase inhibitor (TKI). They may be beneficially in future strategies in patients with endometrial cancer.

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