Case report

Comprehensive genomic profiling (CGP) of ovarian clear cell carcinomas (OCCC) identifies clinically relevant genomic alterations (CRGA) and targeted therapy options

Julia A. Elvin, Justin Churab, Laurie M. Gay, Maurie Markman

1. Introduction

Ovarian clear cell carcinoma (OCCC) accounts for 5–13% of ovarian carcinomas, and is associated with higher prevalence in women of Asian descent, poorer prognosis particularly at advanced stages, and relative platinum insensitivity compared to serous carcinomas. Loss of the SWI/SNF chromatin remodeling gene, ARID1A, and frequent PI3K pathway activation has been previously observed in OCCCs. The mechanistic target of rapamycin (mTOR) is a serine/threonine kinase complex that regulates cellular growth, metabolism, and cell cycle progression. mTOR is directly involved in many cell signaling pathways such as the phosphatidylinositol-3-kinase (PI3k)/AKT pathway, PIK3CA mutation or amplification, AKT overexpression, and loss of PTEN [phosphatase and tensin homolog] lead to an increased mTOR activation, which is regarded as an attractive candidate for therapeutic interventions, and inhibitors targeting different components are under various stages of clinical development (Mabuchi et al., 2015).

There are four main categories of inhibitors targeting the mTOR pathway: mTOR inhibitors, PI3K inhibitors, dual mTOR/PI3K inhibitors, and AKT inhibitors. The rapamycin analog everolimus is an oral mTOR inhibitor currently approved for treatment-refractory advanced renal cell carcinoma, advanced pancreatic neuroendocrine tumors, and the tuberous sclerosis complex-associated tumors renal angiomylipoma and subependymal giant-cell astrocytomas (Ray-Coquard et al., 2013). For both ovarian and endometrial cancers, everolimus is currently being evaluated in clinical trials either as monotherapy or in combination with cytotoxic and hormonal agents (Mabuchi et al., 2015; Ray-Coquard et al., 2013; Wheler et al., 2014; Myers, 2013). Everolimus for a patient whose recurrent, metastatic OCCC harbored three alterations affecting PIK3CA and PTEN, detected by comprehensive genomic profiling (CGP) using next-generation sequencing (NGS) techniques. Additionally, CGP results of 125 advanced/recur rent stage OCCCs, including the case presented here, are included to highlight the frequency of clinically relevant GA (CRGA) that predict subsets of patients who may benefit from therapies targeting the PI3K/AKT/mTOR pathway.

2. Methods

Comprehensive genomic profiling (FoundationOne®) was performed on 125 consecutive clinically advanced/recurrent ovarian clear cell carcinomas (Table 1), as described previously (Frampton et al., 2013). In brief, DNA was extracted from 40 µm of formalin-fixed, paraffin-embedded sections, and CGP was performed on hybridization captured libraries of 236 (n = 73) or 315 (n = 52) genes, plus select introns frequently rearranged in cancer, were sequenced to high, uniform coverage (average 609×). Sequence data were analyzed for clinically relevant classes of genomic alterations, including base pair substitutions, insertions/deletions, copy number alterations, and rearrangements. Clinically relevant genomic alterations (CRGA) were defined as genomic alterations associated with response to therapies currently available or in target-driven clinical trials.

One sample analyzed was the liver metastasectomy sample of a 36-year-old woman with recurrent disease after progression following multiple lines of treatment received over the course of 5 years, including surgery, radiation therapy, hormonal therapy (anastrozole) and conventional chemotherapy (taxol/carboplatin, gemcitabine/carboplatin). After genomic analysis of her tumor metastasis, treatment with everolimus for a patient whose recurrent, metastatic OCCC harbored three alterations affecting PIK3CA and PTEN, detected by comprehensive genomic profiling (CGP) using next-generation sequencing (NGS) techniques. Additionally, CGP results of 125 advanced/recur rent stage OCCCs, including the case presented here, are included to highlight the frequency of clinically relevant GA (CRGA) that predict subsets of patients who may benefit from therapies targeting the PI3K/AKT/mTOR pathway.

http://dx.doi.org/10.1016/j.gore.2017.02.007
In May of 2012, 7 months after completing pelvic radiation, the patient had recurrent disease in the liver. A new 2.3 cm FDG avid focus in the right hepatic lobe was identified on a PET-CT scan. Patient then underwent surgical resection of a subcapsular lesion and segment 7 of the liver. Pathology confirmed a metastatic high-grade carcinoma consistent with ovarian primary. Subsequent additional foci suspicious for recurrence in adjacent to the distal left ureter and left posterior urinary bladder prompted a trial of anastrozole 1 mg daily as the tumor was ER positive. In April 2013, the patient experienced another recurrence in the liver. A 6 × 5.3 cm mass in the caudate lobe of the liver was resected and pathologically confirmed as metastatic high-grade ovarian carcinoma. This specimen was subsequently utilized for comprehensive genomic profiling. Disease progression continued with multiple new liver masses and marked left-sided hydronephrosis documented on imaging. The patient started salvage gemcitabine and bevacizumab in October 2013 with a positive treatment response consistent of an interval decrease in the extent of hypermetabolic malignancy in and adjacent to the liver on PET-CT in May 2014. The patient then underwent radiofrequency ablation and radiation therapy to the liver in June 2014 but was in need of additional options for systemic therapy, as the response was considered likely to be short-lived based on the previous disease course.

Genomic profiling of the 2013 liver metastatectomy specimen identified three CRGA affecting PIK3CA and PTEN, which are associated with sensitivity to mTOR inhibitors. CRGA in any of the other 315 genes included in the assay were not detected. Based on the CGP results, the patient began genomically-matched targeted treatment with everolimus 10 mg oral daily in July 2014. A follow-up PET-CT scan in October 2014 demonstrated resolution of previous liver lesions but a worsening of left hydronephrosis (Fig. 2). Everolimus therapy was continued despite a questionable new mild nonspecific elevation in the lateral right second rib on a February 2015 PET-CT scan. Follow-up scans in June 2015 and October 2015 were negative for metabolically active malignancy. As of September 2016, the patient has received 27 months of treatment with everolimus without significant toxicity (Fig. 2) and CA-125 levels remain low at 10.9. The plan is to continue everolimus as maintenance therapy until untoward side effects occur or disease progression.

### Discussion

More than three quarters of OCCC harbor CRGA, most commonly affecting the PI3K/Akt/mTOR pathway, but also other targetable genes (Supplemental Table 1). A significant fraction of OCCC (11.2%) harbor CRGA only in the mTOR pathway, and we present a case of a robust and durable response to treatment with single agent everolimus for a patient with this genomic profile in late stage, heavily pretreated, multiply recurrent disease. It has been shown that patients with PIK3CA mutations are more responsive to PI3K/AKT/mTOR inhibitors than patients without these mutations (Janku et al., 2012, 2011). As a result, screening for these mutations early in the course of the disease could be crucial and can have important implications for selecting the appropriate targeted treatment consistent with the underlying genomic alteration.

The patient described in this study has experienced a durable response to single agent everolimus exceeding 27 months. The costs associated with both comprehensive genomic profiling and everolimus treatment were covered for the patient by their insurance provider. Coverage of the costs associated with everolimus treatment were approved after a peer review of the mechanistic rationale underlying treatment. The authors acknowledge that financials considerations are a component when considering the use of CGP and developing a viable treatment plan. The list prices of commercially available comprehensive genomic profiling assays are commonly several thousand dollars, although the actual cost to a patient or insurance provider may be less, depending on the individual circumstances. The cost of everolimus...
Fig. 1. Distribution of mutations affecting the PI3K/AKT/mTOR pathway in ovarian clear cell carcinoma. A. More than 69% of samples harbor at least one alteration in the MTOR pathway, and 11.2% of samples harbor CRGA only in the MTOR pathway. B. Distribution of mutations across genes that comprise the MTOR pathway. The number of alterations in each gene is listed. The percentage of cases affected is shown in parentheses. Some cases harbor multiple alterations in the same gene. See Supplemental Table 1 for more details on the types of alterations observed.
purchased through a pharmacy can range from $4000 to $13,000 for a 28-day supply. There are several processes that can be explored when CGP indicates treatment with a therapy that is not FDA approved or NCCN indicted, and which may not be immediately covered by most insurance providers. In addition to seeking insurance coverage, when the rationale for treatment is favorably considered, appeals for compassionate use can be made directly to the manufacturers. In our experience advocating on behalf of patients who have received comprehensive genomic profiling, access to therapy has been granted in approximately 60% of cases. Patient advocacy organizations are an additional resource for support and guidance in crafting these appeals. As evidence supporting the use of genomically matched targeted treatments continues to accumulate, an increasing number of clinical basket trials are underway and may be an appropriate venue for treatment. Our analysis indicates that nearly 70% of patients with OCCC may be positive for a biomarker indicating a potential benefit from treatment with mTOR inhibitors, while demonstrating a high level of co-occurrence of additional alterations that could impact the efficacy of single agent everolimus therapy. For example, 5% of OCC in this dataset show co-occurring RAS and MTOR pathway activation, and in a previous report a patient

Fig. 2. Timeline of clinical history and response to radioablation followed by everolimus. A. Clinical history over time. B. Hypermetabolic malignancy in and adjacent to the liver, as observed by PET-CT in May 2014. The patient underwent radiofrequency ablation and radiation therapy to the liver in June 2014. Genomically-matched targeted treatment with everolimus (10 mg oral daily) begun in July 2014. Follow-up PET-CT scan in October 2014 demonstrated resolution of previous liver lesions but a worsening of left hydronephrosis.
responded to combination therapy targeting the PI3K and RAS pathways (Castro et al., 2015). This report supports the use of comprehensive genomic profiling to identify those patients who are positive for an oncogenic event in any one of 15 components of the PI3K/AKT/mTOR pathway (Fig. 1b), and to assess for the presence or absence of additional drivers that may attenuate a response or underlie resistance to treatment with therapies such as everolimus. In this study, a robust response was observed for a patient whose tumor harbored only alterations in the mTOR pathway; for the additional 60% of patients whose OCCC have alterations in the mTOR pathway, combinations of therapies targeting multiple pathways may be required to achieve similar clinical benefit. Clinical trials of rare ovarian carcinoma subtypes, such as clear cell carcinoma, including broad molecular profiling to investigate the efficacy of biomarker matched targeted therapy, are needed.

Conflict disclosure

JAE and LMG are employees of and own equity in Foundation Medicine, Inc. JC and MM have no conflicts to disclose.

Appendix A. Supplementary data

Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.gore.2017.02.007.

References

Husseinzadeh, N., Husseinzadeh, H.D., 2014. mTOR inhibitors and their clinical application in cervical, endometrial and ovarian cancers: a critical review. Gynecol. Oncol. 133:375–381. http://dx.doi.org/10.1016/j.ygyno.2014.02.017.

Malbochi, S., Kuroda, H., Takahashi, R., Sasano, T., 2015. The PI3K/AKT/mTOR pathway as a therapeutic target in ovarian cancer. Gynecol. Oncol. 137:173–179. http://dx.doi.org/10.1016/j.ygyno.2015.02.003.

Ray-Coquard, I., Favier, L., Weber, B., Roemer-Becuwe, C., Bougnoux, P., Fabbro, M., Floquet, A., Joly, P., Plantade, A., Paraíso, D., Pujade-Lauraine, E., 2013. Everolimus as second- or third-line treatment of advanced endometrial cancer: ENDORAD, a phase II trial of GINECO. Br. J. Cancer 108:1771–1777. http://dx.doi.org/10.1038/bjc.2013.183.

Wheler, J.J., Moulder, S.L., Naing, A., Janku, F., Piha-Paul, S.A., Falchouk, G.S., Zinner, R., Tsimberidou, A.M., Fu, S., Hong, D.S., Atkins, J.T., Yelensky, R., Stephens, P.J., Kurzrock, R., 2014. Anastrozole and everolimus in advanced gynecologic and breast malignancies: activity and molecular alterations in the PI3K/AKT/mTOR pathway. Oncotarget 5:3029–3038. http://dx.doi.org/10.18632/oncotarget.1799.

Myers, A.P., 2013. New strategies in endometrial cancer: targeting the PI3K/mTOR pathway—the devil is in the details. Clin. Cancer Res. 19:5264–5274. http://dx.doi.org/10.1158/1078-0432.CCR-13-0615.

Frampton, G.M., Fichtenholtz, A., Otto, G.A., Wang, K., Downing, S.R., He, J., Schnall-Levin, M., White, J., Sanford, E.M., An, P., Sun, J., Juha, F., Brennan, K., Iwanik, K., Maillet, A., Buell, J., White, E., Zhao, M., Balasubramanian, S., Terzic, S., Richards, T., Banning, V., Garcia, L., Mathney, K., Zirzico, Z., Donahue, A., Beltran, H., Mosquera, J.M., Rubin, M.A., Dogan, S., Heddavat, C.V., Berger, M.F., Pusztai, L., Lechner, M., Boshoff, C., Josting, M., Viertz, C., Parker, A., Miller, V.A., Ross, J.S., Curran, J., Cronin, M.T., Stephens, P.J., Lipson, D., Yelensky, R., 2013. Development and validation of a clinical cancer genomic profiling test based on massively parallel DNA sequencing. Nat. Biotechnol. 31:1023–1031. http://dx.doi.org/10.1038/nbt.2696.

Janku, F., Wheler, J.J., Westin, S.N., Moulder, S.L., Naing, A., Tsimberidou, A.M., Fu, S., Falchouk, G.S., Hong, D.S., Garrido-Laguna, I., Luthra, R., Lee, J.J., Lu, K.H., Kurzrock, R., 2012. PI3K/AKT/mTOR inhibitors in patients with breast and gynecologic malignancies harboring PIK3CA mutations. J. Clin. Oncol. Off. J. Am. Soc. Clin. Oncol. 30:777–782. http://dx.doi.org/10.1200/JCO.2011.36.1196.

Janku, F., Tsimberidou, A.M., Garrido-Laguna, I., Wang, X., Luthra, R., Hong, D.S., Naing, A., Falchouk, G.S., Moroney, J.W., Piha-Paul, S.A., Wheler, J.J., Moulder, S.L., Fu, S., Kurzrock, R., 2011. PIK3CA mutations in patients with advanced cancers treated with PI3K/AKT/mTOR axis inhibitors. Mol. Cancer Ther. 10:558–565. http://dx.doi.org/10.1158/1535-7163.MCT-10-0994.

Castro, M.P., Whitcomb, B.P., Zachowksi, D.A., Coleman, R.L., 2015. Successful use of next generation genomic sequencing (NGS)-directed therapy of clear cell carcinoma of the ovary (CCCO) with trametinib and merfinom in a patient with chemotherapy-refractory disease. Gynecol. Oncol. Res. Pract. 2. http://dx.doi.org/10.1186/s40661-015-0013-2.