Background Outcomes for high grade serous ovarian cancer (HGSOC) patients have remained dismal due to the inevitable development of chemotherapy resistance with recurrent disease. In order to better tailor treatment approaches and uncover opportunities for novel treatments, we need to better understand factors contributing to chemotherapy resistance. Recent studies have shown that immune-related gene expression profiles may serve as prognostic indicators of response to chemotherapy and clinical outcomes in solid tumors, including ovarian cancer. Moreover, immunologic factors have been shown to mediate chemotherapy resistance. Reports in the literature show that common ovarian cancer therapeutic approaches, including chemotherapy, PARP inhibitors, and bevacizumab, modulate tumor cell expressed PD-L1 levels through immunologic signaling pathways. However, very little research has addressed the effect of these treatments on other immune ligands or the differences in immunologic responses between platinum-sensitive and platinum-resistant HGSOC cell lines.

Methods The HGSOC cell lines OVCAR4 (naturally platinum-resistant), PEO1 and PEO4 (matched platinum-sensitive and -resistant lines from the same patient), were treated with common ovarian cancer therapeutics (carboplatin/paclitaxel, olaparib, and bevacizumab), in the presence or absence of peripheral blood mononuclear cells. Western blot was employed to identify levels of immune ligands of interest and a proteome profiler was used to detect broad immunologic changes in response to standard of care therapeutics.

Results Olaparib and bevacizumab treatment strikingly upregulated levels of tumor cell expressed immune ligands ICOSL and PVRL2. Platinum status or presence of an immune component had no bearing on the effect. Moreover, blockade of PVRL2 using siRNA or monoclonal antibodies suppressed STAT3 signaling. When examining the effect of these therapeutics on cytokine levels in HGSOC cell lines treated in immune cell co-culture, OVCAR4 cells displayed marked changes in cytokine levels, particularly CXCL10, CXCL12, SERPINE1, IL1A, and IL1RA. While PEO1 and PEO4 cells displayed more subtle cytokine changes compared to OVCAR4 cells, differences in basal levels and treatment responses were observed between the platinum-sensitive and -resistant lines, respectively. High levels of SERPINE1 and CCL5/RANTES in PEO4 cells, and a robust increase in IL8 levels in response to chemotherapy in only PEO1 cells and not PEO4.

Conclusions In conclusion, common ovarian cancer chemotherapeutics and targeted agents induce tumor cell intrinsic immunologic effects that could potentially be exploited as combinatorial therapeutic targets. Differences in immunologic responses may help define platinum-sensitive and -resistant disease. These results will require further exploration in immune-competent mouse models and human HGSOC tissue.

REFERENCES