Epithelial ovarian cancer: A feasible plan for adjunctive treatment using simultaneous acyclovir, ambrisentan, captopril, disulfiram, fluvoxamine-augmented ramelteon, icatibant, imiquimod peritoneal lavage, and plerixafor

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Abstract

Background: To improve current treatment options in epithelial ovarian cancer, this paper outlines how nine non-cytotoxic drugs, eight of which are marketed for non-cancer indications, can be expected to augment current cytotoxic chemotherapy. By matching what we know about the pathophysiology of epithelial ovarian cancer with the pharmacologic attributes of currently marketed non-cytotoxic drugs, nine drugs were found to match eight identified important growth facilitating paths. The nine drugs: 1) The old anti-viral drug acyclovir inhibits indolamine 2, 3- dioxygenase, an action expected to lower immunoinhibitory Treg lymphocytes; 2) Ambrisentan, marketed to lower pulmonary hypertension can be expected to lower endothelin-1 mediated growth stimulation; 3) The old anti-hypertension drug captopril inhibits matrix metalloproteinases-2 and -9. Both are used for invasion and growth signaling; 4) The old drug disulfiram used to treat alcoholism can be expected to inhibit stemness; 5 and 6) Ramelteon, a melatonin agonist marketed as a sleep aid also stimulates interleukin-2 synthesis. The antidepressant fluvoxamine increases ramelteon levels thereby amplifying interleukin-2 stimulating effects; 7) Imiquimod is a topical Toll-like receptor-7 agonist marketed to treat basal cell carcinoma and other skin cancers by host immunostimulation. It cannot be given systemically but is tolerated well intraperitoneally and can be expected to behave in a similar way in ovarian cancer, causing regression of topically exposed micrometastases after imiquimod peritoneal lavage; 8) The CXCR4 chemokine signaling receptor that plays an important role in many cancers including epithelial ovarian cancer, is blinded by plerixafor, a drug approved for bone marrow stem cell mobilization; 9) Bradykinin is active in generating ovarian cancer ascites. Also captopril risks generating supraphysiological bradykinin levels. Both aspects necessitate using icatibant, a bradykinin blocker. Such a large number of drugs opens the way to unexpected side effects but there is no immediately apparent interaction of concern with simultaneous use of these. If effective as predicted, this type of drug mix could be elevated to a principle of Comprehensive Growth Factor Inhibition as a new core treatment path in cancer.

Preamble: “Little minds try to defend everything at once, but sensible people look at the main point only; they parry the worst blows and stand a little hurt if thereby they avoid a greater one. If you try to hold everything, you hold nothing.” Frederick the Great, mid-1700’s.
with violent weaponry” [5]. CGFI generally, and the nine drugs specifically discussed in this note for EOC, constitute our non-kinetic operations. Surgery, irradiation, and cytotoxic chemotherapies correspond to our kinetic operations.

The Leaky Bucket problem
With many overlapping growth promoting signaling systems, inhibition of any one or even few of these systems will result in no or little increase in overall survival. This situation can be likened to a bucket with ten holes in the bottom. The effect of plugging one or two holes would not be evident; the remaining open holes would just take a little more flow to compensate. It would only be after most holes are plugged that a longer emptying time (corresponding to longer overall survival) would be noticed.

NINE PROPOSED DRUGS AND THE CORRESPONDING EOC SYSTEMS

1. Inhibit EOC endothelin-1 signaling with ambrisentan:
Expression of both the 21 amino acid peptide signaling molecule endothelin-1 and its receptor ET-A have been extensively documented in EOC cells, [6,7] where expression has been shown to enhance epithelial-to-mesenchymal transition and relative resistance to current cytotoxic chemotherapies [6,7,8]. Ambrisentan is a 378 Da orally available ET-A inhibitor, approved and marketed for treatment of pulmonary hypertension in both the EU and USA/Canada [9]. By blocking endothelin-1 signaling in the course of chemotherapy for EOC we would deprive EOC cells of this growth and angiogenesis ET-A signaling path, thereby enhancing chemotherapy effectiveness [6].

2. Reduce MMP-2 and MMP-9 activity with captopril:
Rodent studies, both in vitro [Matrigel invasion, histochemistry, and similar] and in vivo with EOC cell lines have previously shown a prominent role for two proteases, matrix metalloproteinase-2, MMP-2, also known as 72 kDa gelatinase-A, and MMP-9, also known as 92 kDa gelatinase-B [10-14]. Studies directly looking at MMP-2 and MMP-9 in resected human EOC tissues [15-23] show MMP-2/MMP-9 overexpression, with evidence that degree of MMP overexpression is inversely related to survival duration [lower expression, longer survival] in some studies [15,19,21].

MMP-2 and -9 are thought to have roles in degradation of the extracellular matrix allowing EOC migration as well as in triggering growth-promoting surface receptors via proteolytic clipping of surface-tethered ligands for these receptors [24]. MMP-2 and MMP-9 are synthesized as pro-molecules that require proteolytic cleavage themselves before they gain their enzymatic activity [16,22]. Human EOC biopsy tissue contains the necessary protease to generate active MMP-2 from INACTIVE pro-MMP-2 [16].

Starting with the 1993 in vitro data of Sorbi et al. [25] showing that gelatinase activity of both MMP-2 and MMP-9 were inhibited by captopril, several human studies of this have been carried out, reviewed below. Captopril is an angiotensin converting enzyme, ACE, inhibitor in use since 1979 to treat hypertension and as a renal protective agent [26]. Currently, due to captopril’s short half-life requiring multiple daily doses, more recently introduced ACE inhibitors allowing once daily dosing are usually used clinically.

Captopril inhibited in vitro migration and MMP-2 and MMP-9 activity of two glioblastoma cell lines and short term biopsy tissue culture [27]. Nakagawa et al. showed earlier that captopril inhibited both gelatinase and migration in glioma cell line T98G [28]. A single 75 mg oral captopril dose reduced CSF ACE level by 60% indicating robust captopril blood-brain barrier penetration [29].

Captopril inhibited MMP-2 activity in peritoneal dialysis fluid [30]. Serum MMP-9 increases during acute Kawasaki disease and this MMP-9 activity was inhibited by ex vivo serum exposure to captopril [30]. In vitro plasma from acute myocardial patients showed inhibited ACE and MMP-9 activity [31]. Heart failure patients had increased MMP-9 within the failing myocardium that was inhibited by in vitro exposure to captopril [32]. MMP-2 and MMP-9 activity was reduced to half at captopril concentrations of 30 to 50 nM [25], levels easily clinically achieved in humans.

Although most of the research on MMP’s and ACE inhibitors has been done with captopril, other commercially marketed pharmaceutical ACE inhibitors do have evidence of MMP-2 and -9 inhibition. Cardiac biopsy material showed upregulated MMP-2 and or MMP-9 in various heart failure states that was inhibited in vitro by both captopril and the related ACE inhibitor ramiprilate [33]. The recent demonstration by Efen et al [34] that ramiprilate inhibits Crohn’s fistula MMP-9 activity [34] is strong support for the clinical use of captopril or the related ACE inhibitor ramiprilate to decrease EOC’s ability to use MMP-2 and MMP-9 to grow and thrive.

3. Decrease Tregs with acyclovir:
Tregs are a sub-population of lymphocytes that are thought to inhibit or down-regulate other lymphocytes’ immune responses [35]. Tregs bear CD4+, CD25+ FoxP3+ surface markers. Among several immune system inhibitory factors identified in EOC, prominent among these are elevated Tregs in EOC [36-45] with some researchers finding higher Tissue Tregs correlate with shorter survival [41-45].

Indolamine 2-3-dioxygenase, IDO, is the enzyme that converts tryptophan to N-formylkynurenine, which is the initial step in tryptophan catabolism. IDO helps generate immunosuppressive Tregs [46-48] leading to obvious pharmaceutical attempts to increase immune responses in cancer treatment by inhibiting IDO [fewer Tregs] and alternatively damping immune responses [more Tregs] by increasing IDO as for example a treatment path in multiple sclerosis or systemic lupus erythematosus [46,49].

The old drug acyclovir, one of the first anti-viral drugs, was shown to inhibit IDO [50]. If the pre-clinical data holds, such IDO inhibition will decrease Tregs, a potentially useful intervention in EOC given the documented Treg upregulation in this disease. Particularly intriguing and important in this
context are recent reports of increased IDO in EOC [51,52]. This could interestingly enough account for the findings of elevated circulating blood Tregs in EOC compared to controls, and most tellingly the fall in Tregs after surgical debulking of EOC [53].

Others are thinking along similar lines, seeking to reduce Tregs during EOC [54,55]. One approach is to use low-dose cyclophosphamide to reduce Tregs prior to and during experimental vaccination in EOC patients in the effort [56]. Another approach is to use a toxin linked to the natural ligand for CD25 [57], when binding would then be suicidal for Tregs and any other CD25 bearing cell. If the preclinical data holds in the actual clinical situation then acyclovir might be a safer, more effective substitute for either of these.

4. Defeating stemness with disulfiram:
Disulfiram is a small molecule inhibitor of aldehyde dehydrogenase, ALDH. Ethanol is catabolized to acetaldehyde that in turn is broken down to acetic acid by ALDH [58]. By inhibiting this last step, disulfiram, commonly given as a single 250 mg tablet once daily, results in large amounts of acetaldehyde building up if ethanol is ingested. This in turn gives intense nausea, vomiting, malaise, flushing. In the absence of ethanol ingestion disulfiram usually is un-noticed by the patient and is side effect free. Disulfiram has been in continuous use worldwide to treat alcoholism for over sixty years [58,59].

High ALDH activity is quite commonly associated with other attributes of stemness in both cancer and normal stem cell sub-populations [60,61]. Stem cells with a higher expression of ALDH show a higher proliferation rate, higher clonal plating efficiency, and increased resistance to cyclophosphamide and doxorubicin [61]. These authors documented good ALDH inhibition by disulfiram and a corresponding reduction in stemness attributes [61], thereby confirming earlier predictions [59] that disulfiram can defeat stemness via ALDH inhibition.

Human EOC stem cells over-express ALDH and this ALDH over-expressing sub-population shows enhanced proliferation or growth advantage [over the lower or non-ALDH expressing sub-population], better plating efficiency, and greater chemotherapy resistance [62-66].

5. Interleukin-2 increase with fluvoxamine-enhanced ramelteon:
Melatonin is a serotonin metabolite that is synthesized by many cells of the body, including lymphocytes [67]. A large research database supports an immune enhancing role for melatonin by unknown mechanisms [67,68] leading to suggestions that melatonin might have a role in immune enhancement in cancer or during cancer vaccine trials [67,69].

A melatonin analogue marketed as a sleep aid, ramelteon, stimulates and binds to melatonin receptors. With several times greater affinity than the natural ligand melatonin [70]. Since melatonin receptors are present on a sub-population of lymphocytes and melatonin and agonists at melatonin receptors stimulate interleukin-2, IL-2, synthesis and release [71-73] this is thought to be a prominent immunostimulatory path of melatonin. Fluvoxamine increases ramelteon levels by a factor of x16 thereby potentiating ramelteon's stimulation of the melatonin receptor [71].

A further benefit could derive from the observation that we have evidence that IL-2 converts Treg to T effector cells and lose their immunosuppressive capacity [44,74]. Note that Tregs bear CD25+, the outer cell membrane receptor for IL-2. Thus Treg decreases and increased IL-2 could both be achieved by fluvoxamine-augmented ramelteon. “Ovarian cancer is recognized as a paradigm for tumor-associated immune suppression” [75] and the GFI points 3. and 5. above and 6. below are paths that may well reverse this aspect of tumor promotion, providing both “immune-activating strategies [and] elimination of immune-suppressing mechanisms” for better immunotherapy of ovarian cancer [76].

6. Treat micrometastasis-seeded peritoneum surface with imiquimod lavage:
Primary EOC surgery strives to remove as much tumor as possible but the risk/benefit for best extent of blind radicality of resection is still uncertain [77]. Often unseen [not visible to the naked eye] micrometastases are left, seeded throughout the peritoneum. These go on to kill. To combat these micrometastases the current practice of open perioperative hyperthermic peritoneal lavage with cytotoxic chemotherapy agents at the time of primary surgery has somewhat improved the prognosis of EOC and is generally well-tolerated [78,79].

Imiquimod is a Toll-like receptor-7, TLR-7, agonist approved in USA/Canada and in the EU as a topical cream for treatment of basal cell carcinoma and other cutaneous malignancies [80,81]. It cannot be given systemically due to provocation of severe systemic inflammation. But it can be given topically without severe systemic effects, and this includes intraperitoneally in lavage fluid as a form of topical administration [82]. Since mice tolerated imiquimod peritoneal lavage without apparent discomfort [82] and topical imiquimod is effective in a wide variety of cancers resident in skin [81], it was suggested that open peritoneal lavage with an imiquimod solution be undertaken as treatment adjunct at the time of primary EOC surgery [83].

The data of Hackl et al. in 2011 [84] provides an interesting intersection with points 3. and 5. above. They showed that treatment with TLR-7 agonists like imiquimod strongly suppressed Treg number and function [84].

7. Block signaling at CXCR4/CXCL12 with plerixafor:
As in many other cancers [85,86], CXCR4 receptor signaling by 8 to 12 kDa peptide chemokine CXCL12 [also termed stromal cell-derived factor-1, SDF-1] is important to EOC growth, migration and angiogenesis [87-89]. “Abundant" CXCL12 protein was found in most EOC tissue samples but was absent in normal ovary tissue [87]. siRNA-mediated CXCL12 knockdown in an EOC cell line constitutively expressing CXCL12 and CXCR4 reduced proliferation in vitro, and tumor growth in vivo [88]. We can expect similar inhibition of human EOC by treatment with a low
molecular weight, orally available CXCR4 antagonist, plerixafor. Plerixafor triggered necrosis and apoptosis in EOC cells [88]. "SDF-1-CXCR4 interaction confers on EOC cells a remarkable potential to activate MMPs for subsequently invading the peritoneal cavity" [19], findings supported by immunohistochemical studies of others on the importance of CXCR4 in EOC [90]. This provides a happy intersection of captopril-based inhibition of MMP-2 and MMP-9 with plerixafor-based blocking of CXCR4 mediated stimulation of MMP-2 and MMP-9.

Since plerixafor is now approved and marketed for bone marrow stem cell mobilization prior to apheresis harvest for bone marrow transplantation, plerixafor was recently suggested as treatment adjunct in glioblastoma [86] where CXCR4 signaling also plays an important role. It is therefore now straightforward to suggest plerixafor in EOC based on the data above showing activity in a CXCL12 sensitive EOC rodent model and the ample evidence that the CXXCR4/CXCL12 axis plays an important role in growth stimulation in human EOC [87-89].

8. Blocking bradykinin signaling with icatibant:

Bradykinin is a ~1 kDa, nine amino acid, inflammation-related signaling peptide, generated by proteolysis from a high molecular weight precursor kininogen [91]. Bradykinin is rendered inactive by ACE leading to excess bradykinin during ACE inhibitor treatment. That causes the characteristic cough proclivity side effect of ACE inhibitors [91].

Ascites is common in EOC. Evidence points to the bradykinin system as at least contributory to this [92,93]. In humans, ascites development as a consequence of iatrogenic (non-malignant) ovarian hyperstimulation occurs as a consequence of a peritoneal hyperpermeable state largely caused by excess ovary-produced bradykinin [94]. In experimental models, ovarian hyperstimulation ascites is, as expected, made worse by captopril [94]. Active bradykinin and inactive bradykinin catabolites are found in abundance in EOC ascites [95].

Icatibant is a bradykinin receptor blocker recently approved to treat acute hereditary angioedema bouts [96, 97]. In addition icatibant is reportedly effective when used off-label to treat ACE inhibitor-related angioedema [98]. For both reasons identified above, a) the evidence that bradykinin plays at least a contributory role in development of EOC ascites, and b) the clear risk that we increase active bradykinin by the use of captopril (point 2. above), the use of a commercially marketed bradykinin inhibitor, icatibant, will be advisable.

Conclusion

To improve the prognosis in EOC an effort to comprehensively block growth factors that have been identified as active in human EOC as suggested in this paper may prove fruitful. Past research indicates that several already-approved and marketed drugs might do this. The nine drugs have no clearly discernable interaction with each other and none would be expected to interfere with concomitant current cytotoxic chemotherapy regimens although such cannot be excluded. Given the safety of the nine drugs individually, and the poor prognosis of an EOC seeded peritoneum, the risk of unexpected side effects or interaction I believe is worth taking.

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Abbreviations

ACE: angiotensin converting enzyme;
CGFI: Comprehensive Growth Factor Inhibition
EOC: epithelial ovarian cancer; interleukin-2, IL-2; TLR-7, Toll-like receptor-7.

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