Iron addiction: A novel therapeutic target in ovarian cancer

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Abstract

Ovarian cancer is a lethal malignancy that has not seen a major therapeutic advance in over 30 years. We demonstrate that ovarian cancer exhibits a targetable alteration in iron metabolism. Ferroportin (FPN), the iron efflux pump, is decreased, and transferrin receptor (TFR1), the iron importer, is increased in tumor tissue from patients with high grade but not low grade serous
ovarian cancer. A similar profile of decreased FPN and increased TFR1 is observed in a genetic model of ovarian cancer tumor initiating cells (TICs). The net result of these changes is an accumulation of excess intracellular iron and an augmented dependence on iron for proliferation. A forced reduction in intracellular iron reduces the proliferation of ovarian cancer TICs in vitro, and inhibits both tumor growth and intraperitoneal dissemination of tumor cells in vivo. Mechanistic studies demonstrate that iron increases metastatic spread by facilitating invasion through expression of matrix metalloproteases and synthesis of IL6. We show that the iron dependence of ovarian cancer tumor initiating cells renders them exquisitely sensitive in vivo to agents that induce iron-dependent cell death (ferroptosis) as well as iron chelators, and thus creates a metabolic vulnerability that can be exploited therapeutically.

Keywords
iron; ferroptosis; ovarian cancer; tumor initiating cells

INTRODUCTION

Ovarian cancer is the fifth leading cause of cancer-related deaths among women in the US(1), and the most lethal gynecologic malignancy (1)(4). Ovarian cancer is a heterogeneous disease composed of multiple subtypes defined both histologically (2) and molecularly (3, 4). High grade serous ovarian carcinoma (HGSOC) is the most common subtype, accounting for over 50% of ovarian malignancies. The 5-year survival-rate of HGSOC patients is a dismal 9–34%(2). Despite an initial response to therapy, most patients develop recurrent, chemoresistant and ultimately terminal disease(5). One strategy to address this problem has been to develop therapies that effectively target tumor initiating cells (TICs), since this subpopulation of cells in the tumor have been linked to the development of recurrence and resistance to therapy in many types of cancer, including ovarian cancer(6, 7).

Although the origin of ovarian cancer TICs remains an area of debate (8), a growing consensus suggests that stem cells derived from fimbria of the fallopian tubules are the precursors of at least some ovarian cancers, particularly HGSOC (9). By isolating stem cells from normal fimbria and transducing them with defined oncogenes, we recently created a model of serous ovarian cancer tumor-initiating cells (TICs) (10). These cells are capable of multipotent differentiation, and tumors derived from these cells share histological and molecular features of HGSOC.

Alterations in iron metabolism, in particular the acquisition and retention of excess iron, contribute to both tumor initiation and tumor growth (11, 12). This observation has spawned diverse approaches to target intracellular iron in anti-cancer therapy including the use of iron chelators and cytotoxic drug conjugates directed to transferrin receptor 1 (TFR1), a protein that mediates iron import(11). Treating cancer cells with agents that induce iron-dependent cell death (ferroptosis (13)) represents a new strategy for turning the excess iron present in cancer cells to therapeutic advantage. In this manuscript we explore whether HGSOC and HGSOC TICs exhibit targetable alterations in iron metabolism.
RESULTS

HGSOC is characterized by consistent alterations in expression of iron-related genes and proteins

We began by testing whether iron metabolism was altered in HGSOC by assessing levels of transferrin receptor 1 (TFR1) and ferroportin (FPN), proteins that play central roles in iron metabolism by regulating the uptake and efflux of iron, respectively. Immunohistochemistry of normal ovarian epithelium versus tissue from HGSOC patients revealed that TFR1, the iron import protein, was more strongly expressed in malignant than normal ovarian tissue \( (p = 2 \times 10^{-7}) \) and that FPN, the iron efflux pump, was substantially decreased in HGSOC when compared to normal ovarian epithelia \( (p = 2 \times 10^{-11}) \) (Fig. 1). Because of growing evidence that serous tubal intra-epithelial carcinoma (STIC) is a precursor lesion of HGSOC\(^\text{14–16}\)), we next compared normal ovary to STIC. It revealed a similar trend: TFR1 was elevated \( (p = 3 \times 10^{-7}) \) and FPN was decreased \( (p = 5 \times 10^{-6}) \) in STIC relative to the normal ovary. Since HGSOC is suspected to originate in fimbria, we also compared normal fimbria to HGSOC and again observed increased TFR1 \( (p = 0.0001) \) and decreased FPN \( (p = 7 \times 10^{-7}) \) in cancer (Fig. 1). These results identify a phenotype of increased iron retention in ovarian cancer cells that develops early in the evolution of this disease.

Iron regulatory protein 2 (IRP2) regulates TFR1 and FPN levels post-transcriptionally and is overexpressed in some cancers\(^\text{17}\)). Since overexpression of IRP2 would lead to an increase in TFR1 and decrease in FPN, we also assessed levels of IRP2 in our samples of normal and malignant ovary (Fig. 1). Although IRP2 was substantially increased in STIC \( (p = 3 \times 10^{-8}) \) and was modestly elevated in HGSOC \( (p = 0.0008) \) when compared to normal fimbria, it was not statistically different in HGSOC versus normal ovary (Fig. 1), suggesting that differences in post-transcriptional regulation of these proteins, and in particular, differences in levels of IRP2 do not account for all of the observed changes in TFR1 and FPN.

We next tested whether a profile of iron retention was also characteristic of low grade serous ovarian cancer (LGSOC), a less aggressive form of ovarian cancer with a more favorable prognosis that is believed to have a different pathogenesis than HGSOC \(\text{18}\). As shown in Fig. 2, IHC staining for TFR1, FPN and IRP2 revealed that malignant tissue from patients with LGSOC exhibited a staining profile that was quite distinct from HGSOC and more reminiscent of normal tissue, with decreased TFR1 \( (p = 0.001) \), increased FPN \( (p = 0.0001) \) and decreased IRP2 \( (p = 5.9 \times 10^{-6}) \) relative to HGSOC. We also performed IHC staining for ferritin, an iron storage protein composed of H and L subunits that is translationally induced by iron and frequently used as a marker of elevated intracellular iron \(\text{19}\). Consistent with a picture of excess iron acquisition and retention in HGSOC, staining for either ferritin H or ferritin L demonstrated an increase in HG vs LG ovarian cancer \( (p = 1.6 \times 10^{-8}) \) (Fig. 2), as well as in HGSOC vs normal ovary or fallopian tube (Suppl. Fig. 1). Thus, the profile of iron retention that characterizes HGSOC, an ovarian cancer subtype with an especially poor prognosis, does not extend to less aggressive low grade serous ovarian cancer.
Iron metabolism is altered in a model of HGSOC tumor initiating cells

An important clinical problem in ovarian cancer is the development of resistance to chemotherapy, a phenomenon that has been attributed to tumor initiating cells (“cancer stem cells”) resident in the tumor(6, 7, 20, 21). We therefore sought to determine whether alterations in iron metabolism could be detected and potentially targeted in TICs. We used a model of ovarian TIC with limited and defined genetic alterations to help identify specific pathways that drive alterations in iron metabolism (10). This model was created by cloning normal stem cells from human fallopian fimbriae (FTstem cells)(10, 22). We then introduced SV40TAg and hTERT to FTstem cells to create immortalized but non-tumorigenic FT1 cells. Finally FT1 cells were transduced with c-myc to create fully transformed and tumorigenic FTt cells; these give rise to tumors exhibiting the major hallmarks of HGSOC(10). We tested whether changes in iron metabolism occurred during the transition of these normal FTstem cells to malignant FTt cells. We observed a decrease in FPN, an increase in TFR1, and an increase in metabolically available iron (the labile iron pool (LIP)) in FT1 cells expressing hTERT and SV40T; more extensive changes in FPN, TFR1 and the LIP were observed in FTt cells following introduction of c-myc (Fig. 3). Consistent with immunohistochemical staining of ovarian cancer precursor lesions (Fig. 1), these data indicate that perturbations in iron metabolism occur early in the genesis of HGSOC from TICs. They also demonstrate that key elements of the changes in iron metabolism are recapitulated by manipulating p53 and c-myc, two highly prevalent genetic alterations in HGSOC(23, 24).

Ovarian cancer tumor initiating cells exhibit enhanced iron dependence

We tested whether the profile of iron acquisition and retention seen in ovarian cancer TICs was a result of an enhanced dependence on iron (“iron addiction”) (25). We treated TICs with desferroxamine (DFO), an iron chelator, and measured effects on viability. FTt cells were remarkably more sensitive to iron chelation than normal stem cells; immortalized cells exhibited an intermediate sensitivity (Fig. 4a). To confirm the sensitivity of TICs to iron depletion using a genetic rather than pharmacologic approach, we knocked down IRP2 in FTt cells. As expected, knockdown of IRP2 decreased TFR1 and increased ferritin and ferroportin, depleting intracellular iron (Fig. 4b,d). Proliferation FTt cells was dramatically inhibited following this genetically-induced reduction in intracellular iron (Fig. 4c), consistent with the sensitivity of these cells to iron depletion by DFO.

An increase in iron efflux decreases the proliferation of ovarian cancer TICs and HGSOC cells

Our next goal was to determine whether changes in iron metabolism are important drivers or merely incidental passengers in growth and dissemination of ovarian cancer. We therefore selected FPN, a protein that was consistently reduced in ovarian cancer tissues (Fig. 1) and TICs (Fig. 3), and assessed the effect of increasing its expression on the proliferation and tumor-forming capability of ovarian TICs using a conditional doxycycline-driven promoter (FPN-tet-on). Overexpression of FPN reduces intracellular iron by enhancing iron efflux, thus enabling us to test the consequences of altered intracellular iron on the proliferation and tumor-forming ability of ovarian cancer tumor-initiating cells. Following induction of FPN with doxycycline, cells exhibited the predicted increase in FPN and decrease in intracellular
iron (Suppl Fig. 2). This was accompanied by a significant reduction in cell number (Fig. 5a), indicating that a change in the level of FPN is in itself sufficient to modulate TIC proliferation in vitro.

We tested whether cultured cells derived from HGSOC exhibited a similar dependence on iron. We used COV362 cells, an ovarian cancer cell line with mutated p53 and amplified c-myc that is among the top-ranked ovarian cancer cell lines for its similarity in molecular profile to primary HGSOC tumors (26). Like TIC cells, COV362 cells exhibited decreased expression of FPN and increased expression of TFR when compared to normal human ovarian surface epithelial (HOSE) cells (Fig. 6a,b). Conditional overexpression of FPN (Suppl Fig 3) similarly inhibited proliferation and colony-forming ability of COV362 cells (Fig. 6 c,d). Colony formation was not affected in COV362 cells expressing a dysfunctional mutant of ferroportin (FPN A77D) that exhibits attenuated iron efflux activity(27, 28) (Suppl Fig 4).

An increase in iron efflux decreases tumor burden and metastatic spread of ovarian cancer TICs in vivo

To assess whether high intracellular iron contributes to tumor growth in vivo, we injected NSG mice intraperitoneally with FT<sup>t</sup> cells containing FPN-tet-on or empty vector. Mice were treated with vehicle or with doxycycline, and tumors were allowed to grow for 4 weeks before sacrificing the mice. IHC analysis confirmed the induction of FPN in tumors of mice treated with doxycycline (Suppl. Fig. 5). Necropsy revealed widely disseminated tumors in control mice, reminiscent of tumors observed in ovarian cancer patients with advanced disease (Fig. 5b). FPN-overexpressing cells also gave rise to tumors, but the tumor burden and area of metastasis were markedly reduced relative to controls (Fig. 5b–d). Mice injected with tumor cells that did not overexpress FPN also exhibited large blood vessels supplying small and multiple metastases; these were not evident in mice with FPN-overexpressing tumors (Fig. 5b). Quantification revealed a significant decrease in tumor number (p=0.003) as well as a significant decrease in tumor mass (p=4.6x10<sup>−7</sup>) in mice whose tumors over-expressed FPN (Fig. 5c–d).

Iron flux through ferroportin affects invasion of ovarian TICs

In ovarian cancer, the dissemination of tumor within the peritoneal cavity is frequently the only site of metastasis and a major source of morbidity and mortality. To disseminate, tumor cells must seed at multiple peritoneal sites, a process that requires tumor cell invasion of the peritoneum. Given that overexpression of FPN and attendant reduction of intracellular iron markedly reduced the number of tumors (Fig. 5c), we asked whether iron levels in TICs might directly affect tumor invasion by measuring the invasion of FT<sup>t</sup> cells following conditional induction of FPN. Because FPN affects cell proliferation (Fig. 5a), we corrected for cell number by comparing the number of cells that migrated through a layer of basement membrane protein to those that migrated through an uncoated porous membrane. Viable cells expressing FPN showed substantially reduced invasion when compared to control (Fig. 7a). Further, expression of MMP2, MMP9, MMP14 and uPA, enzymes that degrade extracellular matrix, were all reduced in TICs following FPN induction (Suppl. Fig. 6). Thus, iron augments not only proliferation, but also invasion of TICs.

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To understand the mechanism linking iron to invasion, we examined IL6, a cytokine that is elevated in ovarian cancer by a STAT3-mediated pathway, promotes invasion, and contributes to poor patient outcome(29, 30). We found that induction of FPN decreased STAT-3 signaling in FT^T^ cells (Fig 7b) and concomitantly decreased IL-6 mRNA and protein (Fig. 7c,d). Consistent with these results, treatment of cells with the iron chelator DFO also reduced IL6 mRNA (Suppl. Fig. 7). These data suggest that iron accumulation promotes invasion of ovarian TICs at least in part through induction of IL-6.

**Targeting the iron addiction of ovarian TICs through ferroptosis**

Ferroptosis is a recently recognized morphologically and mechanistically distinct form of cell death distinguished by its dependence on iron (13, 31). Ferroptosis causes cell death through iron-mediated accumulation of lipid ROS (31, 32). Ferroptosis can be induced by small molecules such as erastin (13), and inhibited by specifically designed molecules such as Fer-1 (ferrostatin)(31). We reasoned that the excess iron retained by ovarian cancer TICs might render them susceptible to agents that induce ferroptosis. To test this hypothesis, we measured the effect of treating ovarian TICs and non-cancer stem cells with erastin. To confirm that cell death was proceeding via ferroptosis, we treated cells with two different ferroptosis inhibitors, Fer-1 or DFO. Erastin decreased TIC viability (Fig. 8a), and its effects were rescued by both Fer-1 and DFO (Fig. 8b). TICs were significantly more susceptible to erastin treatment than non-cancer stem cells (Fig. 8a).

To test whether erastin was similarly effective *in vivo*, we inoculated NSG mice with FT^T^ cells and treated them with 20 mg/kg erastin daily for 18 days. Erastin treatment produced a marked reduction in both tumor number and mass, indicating that ferroptosis-inducing agents have the potential to successfully target ovarian tumors (Fig. 8c,d).

**DISCUSSION**

Results presented here demonstrate that iron metabolism is dramatically perturbed in HGSOC tumor tissue. Specifically, tumor tissue is characterized by enhanced iron uptake and retention as evidenced by an increase in the iron importer TFR1, a decrease in the iron efflux pump FPN, and an increase in the iron storage protein ferritin relative to non-tumor tissue (Fig. 1,2; Suppl. Fig. 1). The transition to an “iron-seeking” phenotype appears to be an early event in the development of HGSOC, as evidenced by similar perturbations in serous tubal intra-epithelial carcinoma (STIC) and genetically transformed fallopian stem cells (Fig. 1, 3). Further, our results suggest that tumor cells develop an enhanced dependence on iron, or “iron addiction” that can potentially be targeted both by agents that induce iron depletion (Fig. 4) and by agents that depend on iron for their activity, such as the ferroptosis inducer erastin (Fig. 8).

The changes in iron metabolism we observe in ovarian cancer are not merely incidental events that accompany the transition to malignancy: they make important contributions to the growth and metastasis of the cancer. Thus, reduction of iron in TICs through upregulation of ferroportin reduced tumor burden and spread in mice (Fig. 5).
Our studies of a genetic model of ovarian cancer TICs revealed that key elements of iron addiction in ovarian cancer TICs can be recapitulated by introduction of hTERT, SV40T and c-myc into normal fallopian tube stem cells. Although SV40T is not a human oncogene, its validity as an experimental tool in the study of ovarian cancer has been shown by studies demonstrating that its effects can be mimicked by disabling three of its key targets: p53, pRb and protein phosphatase(33). Mutations in TP53 and c-myc are two of the most prevalent genetic alterations in HGSOC(23, 24). We suggest that both p53 inactivation and c-myc expression contribute to altering iron metabolism in TICs, since changes in iron metabolism (decrease in FPN, increase in TFR1 and increase in LIP) were observed in SV40T-transduced FTi cells and further augmented in FTi cells expressing c-myc.

An iron acquisition phenotype has now been reported in a number of cancers: breast(34, 35), glioblastoma(36), prostate(37), and others(11). Iron is required for proliferation and cell cycle progression, and upregulation of these processes underpins at least some of the enhanced dependence of cancer cells on iron. We observed coordinated expression of IL6, MMPs and invasion in response to changes in iron (Fig. 7, Supp. Fig. 6, 7), suggesting a previously undescribed link between iron acquisition, iron-mediated induction of IL6, attendant metalloprotease induction, and tumor cell invasion. IL6 is a key cytokine in the initiation and progression of ovarian cancer(30, 38), affecting tumor cell growth, proliferation, and angiogenesis, as well as contributing to tumor invasion by upregulation of matrix metalloproteases(39). Iron-mediated upregulation of IL6 is consistent with previous work demonstrating that iron chelators inhibit synthesis of IL6 in pancreatic cancer cells (40), and that exogenous iron induces IL6 in hepatocytes (41). Although details of the pathway linking iron to IL6 are yet to be elucidated, STAT3 may play an important role, since a decrease in STAT3 phosphorylation occurred subsequent to iron depletion by ferroportin induction (Fig. 7), and was similarly decreased by iron chelation(40).

Ferroptosis is a recently discovered iron-dependent mode of cell death(13). Agents that trigger ferroptosis potentially represent a new class of anti-cancer agents, and are currently being explored in both preclinical and clinical studies(42). Iron is required to generate the oxidative stress that characterizes ferroptosis, although the precise role played by iron remains speculative(43, 44). The findings we report here suggest that the enhanced levels of iron required for the growth and invasion of HGSOC (and other cancers) may represent an Achilles heel that can be exploited in cancer therapy.

Our evidence of the vulnerability of TICs to ferroptosis may provide not only a new class of compounds that are effective in ovarian cancer, but also an important new target at which ferroptosis therapy can be directed: tumor initiating cells (“cancer stem cells”). We observed that ovarian cancer TICs exhibited decreased FPN, increased TFR1, and decreased LIP relative to non-cancer stem cells (Fig. 3). This was accompanied by an enhanced sensitivity to erastin, a ferroptosis-inducing agent (Fig. 8). Erastin was also able to profoundly inhibit tumor growth in vivo (Fig. 8). TICs are believed to represent a small pool of treatment-refractory cells that contribute to ovarian cancer drug resistance and recurrence (45). Thus the use of ferroptosis-inducing agents in the treatment of ovarian cancer, either alone or in combination with conventional therapies, has the potential to address this important clinical problem.
MATERIALS AND METHODS

Isolation, immortalization, transformation and growth of fallopian stem cells

Adult fallopian stem cells (FTstem) were isolated from fimbrial tissue obtained from discarded surgical specimens of women undergoing benign procedures and cultured on δ-irradiated 3T3 fibroblast feeder cells as described (10). Immortalized and transformed stem cells (FTi, FTt) were cultured in DMEM (GIBCO) containing 10% FBS (BenchMark). All cells were maintained at 37 °C in a humidified incubator at 7.5% CO2. The FTstem cells thus isolated exhibited long term renewability in vitro and the ability to differentiate into secretory and ciliated type cells. The cells were PAX8 positive (marker for fallopian tube epithelium stem cells), E-cadherin positive (marker for epithelial origin), and Ki67 positive (marker for proliferation). Fallopian stem cells (FTstem) were immortalized by infecting with retrovirus expressing hTERT, and SV40 large T antigen as previously described (10). Immortalized cells were further transformed by c-Myc (10). Immortalized cells are referred to here as FTi cells and transformed cells as FTt. Consistent with the expected properties of tumor-initiating cells, these cells can self-renew, form tumorspheres and as few as two thousand FTi cells were sufficient to form palpable tumors in immunodeficient mice in two weeks, and tumor xenografts demonstrated major hallmarks of HGSOC, such as loss of PAX2 and gain of p53, EZH2 and MUC4 expression, and gene expression profiles similar to HGSOC from human patients (10).

Cell culture

COV362 cells were purchased from Sigma and cultured in DMEM (GIBCO) containing 10% FBS (BenchMark). Cells were STR authenticated and tested for mycoplasma by the vendor through the European Collection of Authenticated Cell Cultures (ECACC). Cells were used within three months of receipt from the vendor and were passaged less than five times before use. Human Ovarian Surface Epithelial (HOSE) cells were purchased from ScienCell Research Laboratories and cultured in Ovarian Epithelial Cell Medium (ScienCell Research Laboratories). All cells were maintained at 37 °C in a humidified incubator at 5% CO2.

Infection and preparation of ferroportin-expressing Ft and COV362 cells

Human FPN cDNA was amplified using Open Biosystem clone 4823308 (primers shown in Suppl. Table 1) and introduced into the lentiviral tetracycline (tet) inducible vector pLVX-TetOne-Puro (Takara-Clontech, Mountain View, CA). Lentivirus particles were produced by transient cotransfection of the FPN tet-on expression vector and packaging vectors (VSVG, pMDLG, and RSV-REV) into 293T cells (46). Viral particles containing control empty vector were prepared similarly. COV362 cells were transduced with viral particles in the presence of 4 μg/ml polybrene (Millipore, Billerica, MA, USA). Infected cells were selected with puromycin (2 μg/ml). Mutant FPN (A77D) (27) was generated by site-directed mutagenesis (46) of wild-type FPN and cloned into the pLVX-TetOne-Puro vector. The mutation was authenticated by DNA sequencing.
**Knock down of IRB2 in Ft^1 cells**

IRP2 knockdown was performed using lentiviral shRNAs as described (17).

**Real-time RTPCR**

Briefly, 200 – 400 ng of RNA was reverse transcribed in a total volume of 50 μl with a reverse transcription reagents kit (Applied Biosystems). To make a standard curve, serial dilutions of RNA from one sample were added to the RT reaction. Aliquots (2 μl) of cDNA were added to a 18 μl reaction mixture containing 10 μl of 2× SYBR® Green PCR Master Mix (BioRad) and 400 nm primers. Absence of DNA contamination was confirmed by performing PCR from cDNA without reverse transcriptase. Primers used for each gene are provided in Supplemental Table 1.

**Western blot**

Cells were lysed with NP-40 lysis buffer (25 mM tris (pH 7.4), 1% Triton X-100, 1% SDS, 1% sodium deoxy-cholate, 150 mM NaCl, aprotinin (2 μg/ml), 1 mM phenylmethylsulfonyl fluoride) containing complete protease inhibitor cocktail (Roche Diagnostics). Samples were separated by SDS–polyacrylamide gel electrophoresis and transferred to nitrocellulose membranes before being incubated with primary and secondary antibodies. Bands were detected using chemiluminescence (Thermo Scientific). Membranes were probed with antibodies to β-actin (Sigma, Cat# A3854 ), ferroportin (Novus Biologicals, cat# NBP121502 ), transferrin receptor1 (Invitrogen, cat# 13-6890), IRP2 (Santa Cruz, cat# sc-33682), ferritin H (17), p-STAT3 (cell Signaling, cat# 9131S) and STAT3 (cell Signaling, cat#4904S ). Western blots were quantified using ImageJ software.

**IL6 ELISA**

Cells were seeded in a six well plated in growth media for 24 hours. Conditioned media was collected and viable cells counted. IL6 secretion was measured using a Quantikine IL6 ELISA kit from R & D systems. ELISA results were normalized to number of viable cells at the time of collection of media.

**Cell Invasion assay**

Cell invasion was conducted by Cultrex® BME Cell Invasion Assay kit by Trevigen in a 96 well plate with or without basement membrane extract (BME) coating on the inserts as per manufacturer’s instructions. Because FPN affects cell proliferation, we corrected for cell number by comparing the number of cells that invaded through basement membrane protein to those that migrated through an uncoated 8 micron porous membrane.

**Labile iron pool (LIP Assay)**

Calcein acetoxyxymethyl ester (CA-AM) was obtained from Molecular Probes. The iron chelator, isonicotinoyl salicylaldehyde hydrazone (SIH) (a gift from Dr. P. Ponka, Lady Davis Institute for Medical Research, Montreal, Canada) was prepared as a 50 mM stock solution in dimethyl sulfoxide (DMSO). Briefly, 25,000 – 50,000 cells were cultured in 96-well plates (Black with Clear Bottom purchased from Coring) overnight. Cells were loaded with 2 μM CA-AM for 15 to 30 minutes at 37°C, and then washed with PBS. 100 μM
starch-conjugated desferrioxamine (DFO; a generous gift of Biomedical Frontiers, Inc., Minneapolis, MN) was added to cells to remove extracellular iron. Fluorescence was measured at 485 nm excitation and 535 nm emission with a fluorescence plate reader (BioTek Synergy 2). After the fluorescence signal was stabilized, SIH was added at a final concentration of 10 μM to remove iron from calcein, causing dequenching. The change in fluorescence following the addition of SIH (ΔF) was used as an indirect measure of the labile iron pool.

**Cell viability**

3 × 10^3 cells were seeded in 96 well plates and treated with either deferoxamine mesylate (Sigma), erastin (Selleckchem), or ferrostatin-1 (Selleckchem). Cell viability was assessed at 24–72 hours post-treatment using Cell Titer 96 Aqueous (Promega, Madison, WI, USA). For colony forming assays COV362 cells stably transfected with empty vector (VEC), ferroportin (FPN), or mutant ferroportin (Mutant FPN) were treated with and without doxycycline (0.5 μg/ml) and colony formation was analyzed by crystal violet staining. Colonies from three replicate wells were counted and quantified.

**Immunohistochemistry**

Paraffin-embedded formalin-fixed (PEFF) slides of de-identified human tissues obtained from HGSOC (10 subjects), low grade serous ovarian cancer (10 subjects), STIC (6 subjects), normal oviduct (10 subjects), and normal ovary (10 subjects) were collected from the biorepository of UCHC (IRB IE-08-310-1). Tissues were immunostained with antibodies to human transferrin receptor1 (Invitrogen, cat# 13-6890), ferroportin (47), IRP2 (LSBio, LS-B675), ferritin H (17), and ferritin L (Abcam, cat# ab69090), cytokeratin 7 (abcam, cat# ab9021) and Pax 8 (proteintech, cat# 10336-1-AP). Relevant IgGs were used as negative controls. Images were acquired using a Zeiss Axio Scan Z1. Images of three to four random fields per slide were taken with 40X objectives using Zeiss Axio Imager A.2 and quantified using the DAB application in the open source Fiji software (ImageJ) was used. Reciprocal intensity was measured by subtracting the mean intensity of the stained area of interest from the maximum intensity (48). For each protein, overall differences in staining intensity between groups were calculated using one way ANOVA and individual comparisons were performed using one tailed t-tests. Variance of individual datapoints is shown in the graphs.

**Immunofluorescent staining of cells**

Cells were plated in 8-chamber slides (BD Falcon) and incubated with anti-human ferroportin antibody (47) or anti-human transferrin receptor antibody (Invitrogen, cat# 13-6890). Rodamine-Red (Jackson ImmunoResearch, cat# 109-297-003), Alexa fluor 488 (Invitrogen, cat# A11029), and Alexa Fluor 555-conjugated (Invitrogen, cat# A21429) secondary antibodies were used. Slides were mounted with ProLong Gold anti-fade reagent (Invitrogen). Images were acquired using inverted microscopy (Zeiss Axio Vert.A1).
**Animal Ethics Statement**

All animal studies were conducted in accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the Association for the Assessment and Accreditation of Laboratory Animal Care International (AAALAC). The experiment protocol was approved by the Institutional Animal Care and Use Committee (IACUC) at the University of Connecticut Health Center (protocol # 100881).

**Animal Experiments**

Female NOD.Cg-Prkdcscid Il2rgtm1Wjl/SzJ mice (NSG; ~ 6 weeks of age) were obtained from Jackson Laboratory. \(10^5\) FT cells containing either the FPN tet-on vector or empty vector were injected intraperitoneally (i.p) (n=10/group). FPN was induced in xenografts by adding 2 mg/ml doxycycline to the drinking water and mice were euthanized after 4 weeks. Group size was based on power calculations and was designed to provide 80% power to detect an effect size of 0.5 gm difference in tumor mass using two-sided t-tests. One control mouse died for unknown reasons during the course of the experiment, thus final group sizes were n=10 for FPN-tet-on and n=9 for empty vector. No surviving mice were excluded from the analysis. For erastin treatment, mice were treated with 20 mg/kg Erastin in 2% DMSO/PBS or vehicle alone beginning one day after tumor cell injection. Treatment was continued at five doses per week for 18 days before mice were sacrificed. Tumors were counted and the weight of all tumors found within the abdomen of each mouse was measured. Planned group size for this experiment was n=10; however, a significant difference in tumor size was observed in a pilot experiment using n=6/group; thus, the planned group size of 10 was not used. A control mouse also died of unknown causes in this experiment; thus final group sizes were n=6 for erastin treatment and n=5 for vehicle control. No surviving mice were excluded from the analysis. Mice were not randomized before allocation into control and treatment groups. Outcome assessments were performed by two unblinded investigators. Differences were calculated using two-tailed unpaired Student t-tests on data that were normally distributed and of equal variance.

**Statistical analysis of cell culture experiments**

Statistical analyses were performed using Excel or Prism 6 (Graphpad software). All experiments were performed at least three times using a minimum of three replicates/condition in each experiment. Results of representative experiments are shown in the figures. Comparison tests were performed between two groups and statistical significance assessed using two-tailed unpaired Student’s t tests. Statistics are reported as the mean ± standard deviation. Error bars represent standard deviation.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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Fig. 1. Proteins that control intracellular iron are altered high grade serous ovarian cancer (HGSOC)
Representative images of immunohistochemical staining of normal fimbria, normal ovary surface epithelia, serous tubal intra-epithelial carcinoma (STIC) and HGSOC stained with antibodies against transferrin receptor 1 (TFR1), ferroportin (FPN) and iron regulatory protein 2 (IRP2). Dot plots represent quantification of staining of tissues collected from 8 patients with HGSOC and 5 with STIC compared to 8 subjects with normal fimbria and 6 individuals with normal ovarian surface epithelium. Images of three to four random fields per slide were quantified. Differences in TRFC (p<0.0001), FPN (p<0.0001 and IRP2 (p<0.01) were statistically significant (one way ANOVA). *p<0.001, **p<5x10^{-5}, ***p<5x10^{-6}, ****p<5x10^{-7}, one tailed t test for individual comparisons. Scale bar 1 mm; inset scale bar 10 μm.
Fig. 2. Proteins that control intracellular iron differ in high grade and low grade serous ovarian cancer
Representative images of immunohistochemical staining of tumor tissues from patients with low grade serous ovarian cancer (LGSOC) and high grade serous ovarian cancer (HGSOC). Proteins stained are ferroportin (FPN), transferrin receptor (TFR1), Iron Regulatory Protein 2 (IRP2), ferritin heavy chain (FTH) and ferritin light chain (FTL). Dot plots show quantification of staining of tissues from 5 patients with LGSOC and 8 patients with HGSOC (3 to 4 random fields from each patient tissue slide). *p<0.002. Scale bar 1 mm; inset scale bar 10 μm.
Fig. 3. Iron metabolism is modified in a genetic model of ovarian tumor initiating cells

(a,d) Immunofluorescent staining of ferroportin (FPN) and transferrin receptor (TFR1) in normal fallopian tube stem cells (FTstem), immortalized fallopian tube stem cells (FTi) and transformed fallopian tube stem cells (FTt). FPN and TFR1 in red; nuclei in blue. Scale bar 20 μm. (b,e) Cropped images of western blots and quantification of FPN and TFR1 with β-actin as a loading control. (c,f) qRT-PCR of FPN mRNA and TFR1 mRNA in FTstem, FTi and FTt cells. (n) Labile iron pool (LIP) per 1000 cells. *p<0.02, **p<0.0007, ***p<5x10⁻⁸; ****p<5x10⁻¹¹.
Fig. 4. Tumor-initiating cells exhibit increased iron dependence
(a) Cells were treated for 72 hrs with the indicated concentrations of deferoxamine (DFO) and viability was assessed using an MTS assay. (b) Cropped images of western blots of IRP2, transferrin receptor, (TFR1) ferritin heavy chain (FTH), and ferroportin in FT<sup>t</sup> cells with knockdown of IRP2 (IRP2 KD1, IRP2KD2) or control shRNA (shCtr). (c) Cell proliferation as assessed by trypan-blue exclusion in FT<sup>t</sup> cells treated with control shRNA or IRP2 knockdown vectors. d) Labile iron pool in cells with knockdown of IRP2 or control shRNA
Fig. 5. Increased iron efflux decreases tumorigenicity and invasion of ovarian cancer tumor-initiating cells

(a) Cell viability of FT cells as assayed by MTS assay at indicated time points with and without ferroportin overexpression. Blue line represents control (empty vector with doxycycline), and red line represents FPN-tet-on FT cells (ferroportin tet on with doxycycline). *p<0.0002, one tail t test. (b) Representative images of mice inoculated IP with FPN-tet-on FT cells and left untreated (left) or treated with doxycycline (right) for four weeks to induce expression of ferroportin. Untreated mice have a greater tumor burden and wider area of metastasis (circled in white). (c) Quantification of tumor number/mouse and (d) tumor mass/mouse following implantation of tumor cells containing empty vector (n=9) or FPN tet-on (n=10) following 4 weeks of doxycycline treatment.
Fig. 6. Increased iron efflux reduces proliferation of COV362 ovarian cancer cells
(a) q-RTPCR of FPN (normalized to β-actin) and immunofluorescence staining of FPN in COV362 and HOSE cells: FPN in red; nuclei in blue. Scale bar 20 μm. (b) q-RTPCR of TFR1/β-actin in COV362 ovarian cancer cells and HOSE cells; (c) FPN was induced at time 0 by the addition of doxycycline and cell viability assessed at the indicated timepoints by MTS assay; (e) Colony formation of COV362 cells with and without ferroportin overexpression was analyzed by crystal violet staining. Colonies from three replicate wells were counted and quantified.
Fig. 7. Ferroportin modulates invasion of FT\textsuperscript{4} cells through STAT3 and IL6

(a) In vitro invasion was measured in a simplified Boyden chamber consisting of two chambers separated by an 8 micron porous filter coated with or without basement membrane extract (BME). Invasion was normalized to migration of the cells in the inserts without the BME coating. *p=0.002; **p=5\times10^{-5}. (b) Western blot analysis of p-STAT3 and total STAT-3 in FT\textsuperscript{4} cells transduced with empty vector (EV) or FPN-Tet-on following 48 hrs exposure to doxycycline. Band intensities were quantified using ImageJ. (c) IL6 mRNA was assessed using qRT-PCR; (d) IL6 was measured in culture supernates following 48 hrs’ exposure to doxycycline. *p \leq 0.0002.
Fig. 8. Erastin induces ferroptosis in tumor-initiating cells and reduces tumor number and mass in vivo
(a) 3000 FTrstem and FT cells were treated with indicated doses of erastin and cell viability assessed 48 hours later. (b) FT cells were treated for 48 hrs with 2 μM of erastin alone, the combination of 2μM erastin plus 2 μM ferrostatin-1, the combination of 2μM erastin plus 50 μM DFO, ferrostatin alone, or DFO alone and cell viability assessed. (c,d) Mice were injected intraperitoneally with FT cells and treated for 18 days with either vehicle or 20 mg/kg erastin. The number of tumors/mouse and total tumor mass/mouse were measured in the vehicle-treated group (n=5) and in the group treated with erastin (n=6).