Activated anti-tumor immune infiltrates are associated with improved survival in p53 abnormal endometrial carcinoma

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The authors declare no potential conflicts of interest.

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## Translational Relevance

*TP53* mutated/abnormal (p53abn) endometrial carcinoma is the molecular subtype responsible for most deaths from endometrial carcinoma. Recent clinical trials have demonstrated that a subset of p53abn cancers respond to immune checkpoint inhibitors given alone or in combination with anti-angiogenic agents. We profiled the immune microenvironment of a large well-characterized cohort of p53abn endometrial cancers from across Canada, showing that approximately 50% of tumors are highly infiltrated by immune cells. Immune rich cases associated with longer survival and did not correlate with other therapeutically targetable biomarkers (homologous recombination deficiency and HER2 status). Our findings provide rationale for treatment decisions in p53abn endometrial carcinoma by identifying criteria indicative of an active anti-tumor immune response which may be augmented by immunotherapy. This study demonstrates the potential for rationalized personalized therapy for this deadly disease.

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## Abstract

**Purpose:** *TP53* abnormal (p53abn) endometrial carcinoma (EC) is the deadliest subtype of EC, with 50% of patients dying within 5 years. Most patients show limited response to standard-of-care chemotherapy or newer targeted therapies. Recent clinical trials suggest that some patients with p53abn EC respond to immune checkpoint inhibition. However, the immune microenvironment and its relationship to other therapeutic targets in p53abn EC remains poorly understood.

**Experimental design:** We accrued 256 treatment-naïve p53abn EC samples and systemically profiled T cell, B cell, myeloid and tumor cell populations with multiplex immunofluorescence and guided artificial intelligence analysis to assess the tissue localization and functional status of immune cells. Patterns of immune infiltration were associated with survival outcomes and shallow whole genome sequencing-derived mutational signatures.

**Results:** Mixture modeling divided p53abn EC into tumor infiltrating lymphocyte (TIL)-high and TIL-low subsets. The TIL-high subset was associated with longer overall and disease-specific survival in multivariate analysis, which was particularly pronounced in advanced stage disease. Furthermore, TIL-high cases were associated with increased PD-L1+ macrophages and PD-1+ CD8 T cells. TIL-high p53abn EC cases did not associate with homologous recombination deficient mutational signatures or HER2 amplification.

**Conclusions:** TIL-high p53abn EC correlated with improved survival and did not correlate with homologous recombination deficiency or HER2 amplification. These findings may help inform combination therapies with immune checkpoint inhibition, PARP inhibitors, and anti-HER2 agents.

## Introduction

Endometrial cancer is the second most common type of gynecologic malignancy worldwide and the most common gynecologic malignancy in North America (1). Molecular classification stratifies endometrial cancers into 4 prognostically distinct subtypes: polymerase epsilon-mutated (*POLE*mut), mismatch repair-deficient (MMRd), p53 abnormal (p53abn) and no specific molecular profile (NSMP) (2–5). p53abn cancers have the worst prognosis, comprising only 15% of all endometrial cancers but accounting for 50-70% of mortalities (3,6–8), with extrauterine involvement in over 50% of cases (5,9). Most p53abn cancers recur within five years on standard-of-care carboplatin-paclitaxel chemotherapy with or without adjuvant radiotherapy, highlighting the need for alternative therapeutic options (6,7,10). While several studies have investigated associations between immune infiltration and survival in endometrial carcinomas (11–14), large scale studies investigating the immune microenvironment of p53abn tumors are lacking.

The interplay between different immune cell types within the tumor microenvironment determines the effectiveness of anti-tumor immunity. CD8+ cytotoxic T lymphocytes (CTL) are the main cells that recognize and kill tumor cells. Dendritic cells, macrophages and B cells are professional antigen presenting cells that activate CTL and CD4+ helper T cells (TH), which in turn secrete cytokines that potentiate the activity of CTL and anti-tumor macrophages. CD68+ macrophages can either help activate CTL, by presenting antigens and co-stimulatory molecules, or inhibit CTL, by presenting antigens along with inhibitory ligands (15,16). CD4+FOXP3+ regulatory T cells (Tregs) inhibit anti-tumor immunity by secreting cytokines that block CTL maturation and activity and induce macrophages to express immune inhibiting molecules (17,18). Finally, CD20+ B cells and CD79a+ plasma cells may potentiate anti-tumor immunity via multiple mechanisms (19,20). Key functional molecules that inhibit anti-tumor immunity include indoleamine 2,3-dioxygenase 1 (IDO1) and programmed death-ligand 1 (PD-L1). Macrophages and tumor cells may express IDO1, which limits tryptophan availability, thereby inducing CTL death and proliferation of Tregs (21). Tumors and macrophages may also express PD-L1, blocking CTL mediated killing by ligating programmed cell death-1 (PD-1) on CTL and TH (22). Cytokines such as IFN-gamma are released by CTL upon tumor cell recognition and in turn cause upregulation of both IDO1 and PD-L1 (22); thus, expression of PD-L1 has been used as a marker for an active anti-tumor immune response in multiple different cancer types (23–25).

Immune checkpoint inhibitors disrupt the PD-1 – PD-L1 pathway, reactivating exhausted T cells to attack tumor cells. These treatments are particularly effective in tumors with elevated numbers of mutations that generate neoantigens (26). In endometrial cancer, *POLE*mut and MMRd tumors have over 10 times as many mutations as p53abn and NSMP tumors (2,13) and correspondingly higher TIL densities (14). While systemic therapy is often unnecessary in *POLE*mut cancers due to exceptionally favorable outcomes with hysterectomy alone, anti-PD-1 immune-checkpoint inhibitors have demonstrated remarkable efficacy in advanced, recurrent and persistent MMRd endometrial cancers, even after multiple lines of therapy (27–30). More recently, Mirza et al. (31) and Eskander et al. (32) showed benefit of adding PD-1 inhibitors to chemotherapy in both MMRd and in MMR-proficient (MMRp) endometrial carcinomas. Subgroup analysis showed that the benefit in MMRp was driven by p53abn cases (33), and thus pembrolizumab and dostarlimab received FDA approval for treatment of endometrial carcinoma, regardless of subtype. However, the factors underlying response to immune checkpoint inhibitors in p53abn endometrial cancer remain poorly understood.

Additional classes of targeted therapies under investigation in p53abn endometrial cancer include PARP inhibitors and HER2-directed antibodies (34). PARP inhibitors have become standard-of-care in *BRCA1*/*BRCA2*-mutated or homologous recombination deficient (HRD) cancers in several cancer types (35). In high-grade serous ovarian cancer (HGSOC), HRD tumors have higher immunogenicity than non-HRD tumors, and markers of adaptive immunity are associated with longer overall survival in HRD but not non-HRD tumors (36). In p53abn endometrial cancer, approximately 25% of cases show evidence of HRD and fewer than 5% have *BRCA1/2* mutations (37,38). Combination immune checkpoint and PARP inhibition has shown benefit in homologous recombination-deficient endometrial cancers (39); however, the relative immunogenicity of these cases has yet to be explored. p53abn endometrial cancers show a higher incidence of HER2 amplification than other molecular subtypes, with up to 25% of high-risk cases HER2 amplified (40). Several phase I and II clinical trials assessing HER2-targeted therapies are in progress (41–45), but to date, studies investigating the relative immunogenicity of HER2-amplified p53abn endometrial carcinoma is lacking. Thus, an improved understanding of the relationship between HRD, HER2 status and the immune microenvironment in p53abn endometrial cancers may help inform combination PARP inhibitor, HER2 blockade and immunotherapy in clinical trials (39).

To understand the clinical relevance of the immune response to p53abn endometrial cancer, we systematically profiled the immune cell composition of 256 clinically annotated p53abn endometrial cancers with multiplex immunofluorescence to detect CTL, TH, Tregs, B cells, plasma cells and macrophages. Further, we evaluated the expression patterns of PD1, PD-L1 and IDO1, three pharmacologically actionable immunosuppressive molecules with translational relevance to current clinical trials in endometrial cancer. Finally, we investigated the relationship between immune composition, HRD and HER2 expression/amplification in p53abn endometrial cancer.

## Methods

### Data acquisition

#### Sample acquisition and TMA construction

Ethics approval was obtained from the University of British Columbia (UBC) Research Ethics Board (approval number H18-01652) and the institutional review boards from each center that supplied tissue. The cohort consisted of 256 treatment-naive p53-abnormal endometrial carcinomas collected between 1993 and 2017 in Vancouver and in 10 tertiary and 19 community centers from across Canada (3,4,46). Clinicopathologic and outcome data were collected by chart review. All cancers were classified as p53abn according to the ProMisE algorithm (3) by immunohistochemistry (IHC) for p53 and MMR proteins and next-generation sequencing for *POLE* hotspot mutations. Representative samples of p53abn endometrial carcinomas were cored at 0.6 mm in diameter, in duplicate, and arrayed as described previously (3).

#### Multiplex immunofluorescence

Summary: Full methods are described in **Supplemental Methods 1**. TMAs were cut at 4 um for immunofluorescence. Slides were first deparaffinized with xylene and graded alcohols (Fisher), then incubated for 20 minutes in 10% neutral buffered formalin (Sigma) followed by a wash in deionized water. Diva Decloaker solution (Biocare) was used for antigen retrieval, and slides were stained in an Intellipath FLX Autostainer (Biocare) in six rounds. Endogenous peroxidases were blocked with peroxidazed-1 (Biocare), and non-specific background staining was reduced using Background Sniper (Biocare). Antibodies were diluted in Da Vinci Green Diluent (Biocare) and incubated for the times and concentrations noted in supplementary methods. Either Mach2 Rabbit-HRP polymer (Biocare) or Mach2 Mouse-HRP polymer (Biocare) was added and the specific flour for each round was added as noted in supplementary methods. After each round of staining, the reagent proteins were denatured by microwaving in AR6 (Akoya) in preparation of the subsequent round. Nuclei were counterstained with Spectral DAPI (Akoya) and a coverslip was added using Prolong Diamond coverslipping media (Fisher). Both panels (B/T panel: CD8, CD3, FOXP3, CD20, CD79a, panCK; adaptive response panel: CD68, CD8, IDO1, PD-L1, PD-1, panCK) were imaged using the Vectra Polaris multispectral imaging system (Akoya). They were scanned using the “motif” settings to create a whole slide multispectral QPTIFF file.

### Computational analysis

#### Cell and region counting

Analysis was performed using HALO (Version 3.6.4134.95). Briefly, three tissue segmentation and cell segmentation/phenotyping algorithms were trained using 10 – 30 representative images for each algorithm. Regions were classified as epithelial, stromal, glass, or other (including necrosis) for tissue segmentation. The mean and standard deviation of cell count and region area across all algorithms was calculated for each core and reviewed by a technologist. Any core with a standard deviation of >5 among the three algorithms, or with abnormal features as identified by the technologists, were flagged for review and manual annotation by a pathology resident and/or subspecialist gynecological pathologist. Sarcomatoid areas in carcinosarcomas were considered epithelial. The intraepithelial area that was negative for CD68 and CD8 was designated tumor cell area. All immune cells touching tumor cells were considered intraepithelial. TIL counts and areas for all duplicate cores from each sample were added together. TIL densities were computed by taking the quotient of TIL counts divided by areas (epithelial or stromal), and log-transformed TIL densities were computed as log(TIL density + 1).

#### Mixture modeling of cell counts

The TIL count for a given core , cell type , and region (tumor/stroma) was described as follows:

where the mean follows:

where is the inferred cluster of core , is the area of region in core , and is the mean density (count divided by area) value across all cores . The Gamma distribution was parameterized in terms of a mean and scale parameter, with the scale parameter set to 100 to allow for a fairly uninformed prior.

To reduce the dimensionality of the parameter space, the dispersion parameter was formulated semi-parametrically in terms of a set of Gaussian radial-basis kernels with parameters and , for a specified number of centers uniformly distributed between 0 and the maximum number of counts (47,48), with the default number of centers set equal to 20:

where and follow relatively agnostic lognormal priors as above.

Inference was performed in pymc v5.9.1 with the number of tuning and sampling iterations set to 1000 each. Continuous parameters were sampled with the No-U-Turn Sampler (NUTS) and categorical parameters were sampled with the Metropolis sampler. Convergence was assessed by examining trace plots. To determine the optimal number of clusters , a Dirichlet process prior was first fitted to a version of the model marginalized over cluster membership . Two clusters with proportions of at least 0.05 (5%) were inferred (first cluster with mean proportion 0.588, 95% CI 0.512-0.656; second cluster with mean proportion 0.347, 95% CI 0.289-0.407). The lower bound of the 95% CI for the third most prevalent cluster was 0.004 (i.e. 1 out of the 256 input samples), supporting the presence of only 2 main clusters. Thus, the model was fitted again with the number of clusters fixed at 2 () to determine cluster membership for each input sample.

#### Shallow whole-genome copy number analysis

Copy number (CN) signatures were generated as described in (38) using the process outlined by (49), as implemented by the Utanos R package (<https://github.com/Huntsmanlab/utanos>). In short, CN features such as DNA segment size, CN change points, segment CN, breakpoints per 10Mb, length of segments with oscillating CN, and breakpoints per chromosome arm were extracted using the ExtractCopyNumberFeatures function in Utanos. The optimum number of signatures was determined by the ChooseNumberSignatures function, subsequently, the GenerateSignatures function was applied to the selected number of signatures. Finally, the CN signatures were generated according to the sample by signature exposures. Five CN signatures were identified in our p53abn cohort, one of them (VS5) associated with BRCA1/2 CN Loss. VS5 samples also showed an elevated *BRCA1/BRCA2*-associated HRD CN signature (BS3), first described by (49) in HGSOC samples. HER2 (*ERBB2*) gain was defined as an absolute copy number (ACN) of 2.5-4.5 copies, HER2 amplification was defined by an ACN of 4.5-8.5 copies, and HER2 high-level amplification was defined by an ACN of greater than 8.5 copies. ACN was determined from relative copy number profiles using Rascal (50). These results were corroborated by immunohistochemical staining for HER2.

#### Statistical analysis

All statistical analysis was performed in R (v4.3.2). The Mann-Whitney U test was used to evaluate significance in two-way comparisons. Comparisons for categorical count data were evaluated for significance with Fisher’s exact test. Multiple testing correction was performed with the Holm method. Results with adjusted *P* < 0.05 were considered statistically significant.

*P* values for Kaplan-Meier analyses were computed with log-rank tests. Cox proportional hazards analysis was performed with the survival package (v3.5-7) in R. Samples with missing values were excluded from analysis. Proportional hazards assumptions were evaluated with weighted Schoenfeld residuals (51). Hazard ratios and *P* values from the survival package were validated for consistency with a Bayesian implementation of the Cox proportional hazards model.

Hierarchical clustering was performed with Euclidean pairwise distance and Ward’s method.

#### Code availability

Code associated with this project will be made publicly available at <https://github.com/Irrationone/tfri_halo>.

*Data availability statement*

The data generated in this study are available upon request from the corresponding authors.

## Results

### Cohort

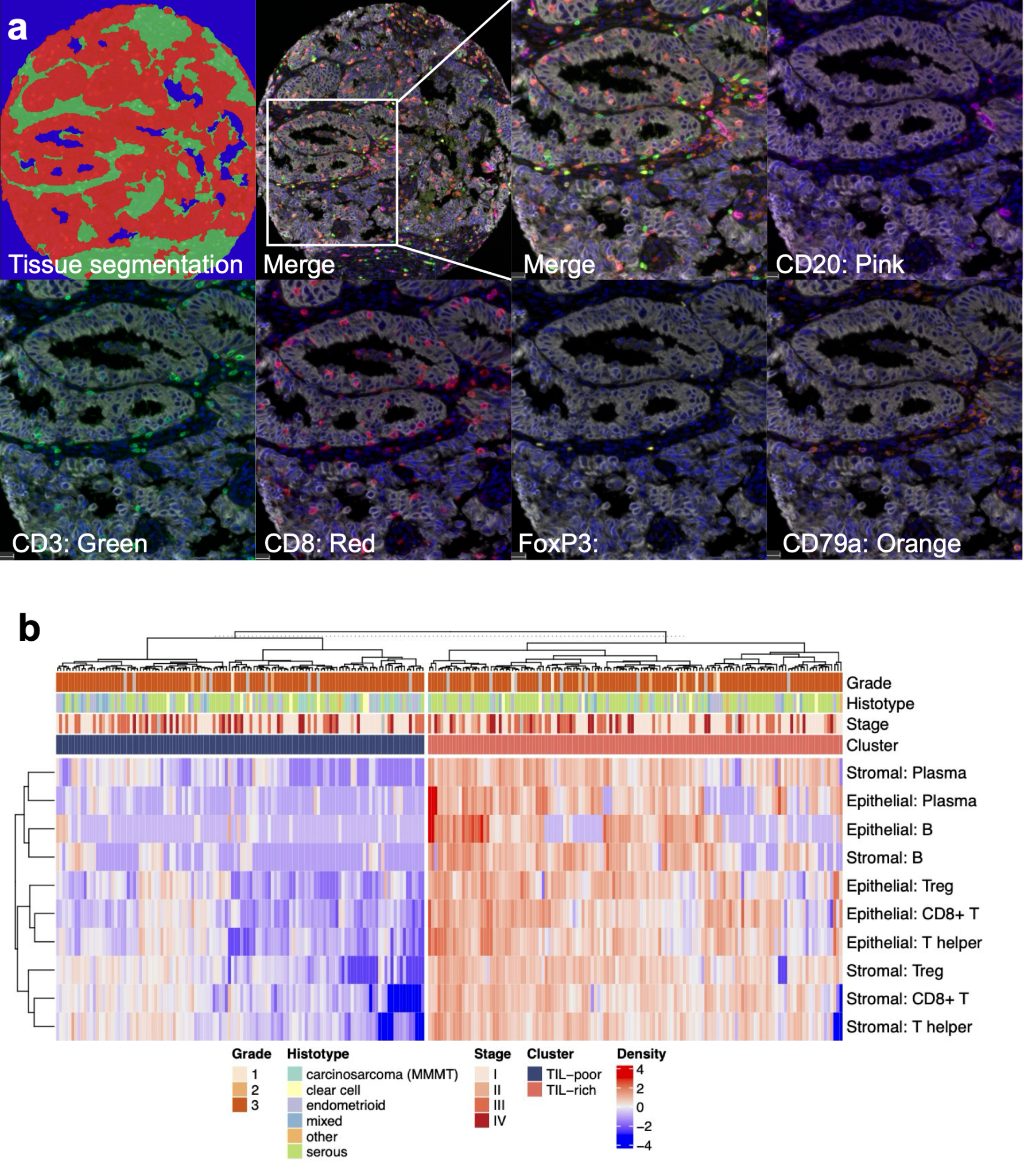
We assembled a cohort of 256 treatment-naive p53abn endometrial carcinomas diagnosed between 1993 and 2017 from institutional cohorts as well as a pan-Canadian initiative (**Figure 1, Methods**) (14). Histotypes of p53abn tumors included serous (n=136), endometrioid (n=52), carcinosarcoma (n=31), clear cell (n=15), mixed (n=17) and other (n=5). All patients received treatment in accordance with standard-of-care at the time of diagnosis, with most patients receiving adjuvant chemotherapy and a smaller proportion receiving adjuvant brachytherapy or radiotherapy (**Figure 1**). No patients received immunotherapy or neoadjuvant chemotherapy.

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| *Figure 1:* Graphical overview of clinicopathologic parameters and data types for each sample, showing the relationships between different parameters. Missing values are colored white. (Myo = Depth of myometrial invasion, LVI = Lymphovascular invasion) |

### Clustering based on immune composition

We performed multiplex immunofluorescence and automated image analysis to segment tumors into epithelial, stromal and ‘other’ regions, followed by quantification of lymphocyte subsets, including CTL (CD3+CD8+), TH (CD3+CD8-FOXP3-), Treg (CD3+CD8-FOXP3+), B cells (CD20+CD79a+) and plasma cells (CD20-CD79a+) (**Figure 2a**). Cells within epithelial and stromal regions were counted separately, and automated counts were manually verified. We found that all TIL densities were highly correlated. Higher correlations between TIL types were seen within specific regions, and the highest correlations were seen between TIL types from the same cell lineage within regions (**Figure S1**).

Next, we clustered tumors based on epithelial and stromal counts normalized by region area with a negative binomial mixture model. As the number of immunologically distinct clusters was unknown *a priori*, the model was formulated to automatically determine the optimal number of clusters using a Dirichlet process (**Methods**). Two distinct clusters were found: TIL-rich and TIL-poor. T and B cell subsets, including CTL, TH, Treg, B cells, and plasma cells infiltrated both tumor stroma and epithelium in TIL-rich tumors (**Figure 2b**). We found no evidence of a subgroup with stroma-restricted TIL, contrary to our findings of an immunologically and genomically distinct subgroup of tumors with stroma-restricted TIL in high-grade serous ovarian carcinoma (36).



*Figure 2: Derivation of TIL-rich and TIL-poor groups.* (a) Representative multiplex immunofluorescence image of the B/T cell panel. Representative segmentation of the tumor (red), stromal (green), and glass/necrosis (blue) are shown in the top left panel. Cytokeratin (white) and DAPI (blue) are shown in each image along with a single immune marker as annotated. (b) Heatmap of log-transformed epithelial and stromal TIL densities for each sample. Results of hierarchical clustering by sample (top dendrogram) are split by TIL cluster.

### TIL-rich tumors are associated with longer survival

We next assessed the relationships between TIL cluster, survival, and other clinicopathologic parameters in our cohort of p53abn endometrial carcinomas. In multivariate Cox proportional hazards analysis, significant associations were identified between overall survival and adjuvant chemotherapy (HR 0.58, *P* = 0.041) and FIGO stage (HR 3.39 and 11.0 for stage III and IV compared to stage I, respectively, both *P* < 0.001), while older age trended towards shorter overall survival (HR 1.02, *P* = 0.14). Importantly, TIL-rich tumors were significantly associated with prolonged overall (HR 0.63, *P* = 0.031) and disease-specific survival (HR 0.58, *P* = 0.037) in multivariate Cox proportional-hazards analysis accounting for age at diagnosis, FIGO stage (52), and adjuvant treatment (**Table 1**), and trended towards longer progression-free survival (HR 0.74, *P* = 0.15). When TIL subtypes were assessed individually, each subtype was associated with mean hazard ratios between 0.77 and 0.98, with statistical significance only in stromal CD8+ T cells, T helper cells and T regs. (**Table S1**). In contrast, univariate Kaplan-Meier analyses failed to identify a significant association between TIL-rich cases or TIL densities and overall, progression-free, or disease-specific survival (all *P >* 0.175) (**Figure 3a**).

To better understand the discordance in the effect of TIL between univariate and multivariate analyses, we assessed the TIL effect stratified by stage and adjuvant therapy. Univariate Kaplan-Meier analysis stratified by stage highlighted that the association between TIL cluster and survival was most pronounced in patients with stage III disease (**Figure 3b-e**). The median 5-year overall survival in stage III disease for TIL-poor cancers was 24.9% (95% CI, 9.6%-64.7%) versus 55.2% (95% CI, 38.3%-79.6%) for TIL-rich stage III cancers (n = 71). In contrast, the median 5-year overall survival in stage I disease for TIL-poor cancers was 74.9% (95% CI, 61.9%-90.6%) versus 79.1% (95% CI, 67%-93.4%) for TIL-rich stage I cancers (n = 118). Interpretability within stage II and IV tumors was limited by smaller sample sizes (**Figure 3c,e**). Thus, TILs were associated with longer survival in p53abn endometrial carcinoma, particularly in advanced stage disease. When we separated the cohort based on adjuvant chemotherapy use, we found that despite higher rates of chemotherapy use in stage III and IV disease in our cohort (*P* < 0.001), TIL-rich cases were associated with significantly longer disease-specific survival (*P* = 0.03) and trended towards longer progression-free survival and overall survival (*P* = 0.067 and 0.09, respectively) only in patients who did not receive chemotherapy. (**Figure 3f,g**). This finding was only significant when all stages were assessed as there were only trends within each stage due to lack of power.

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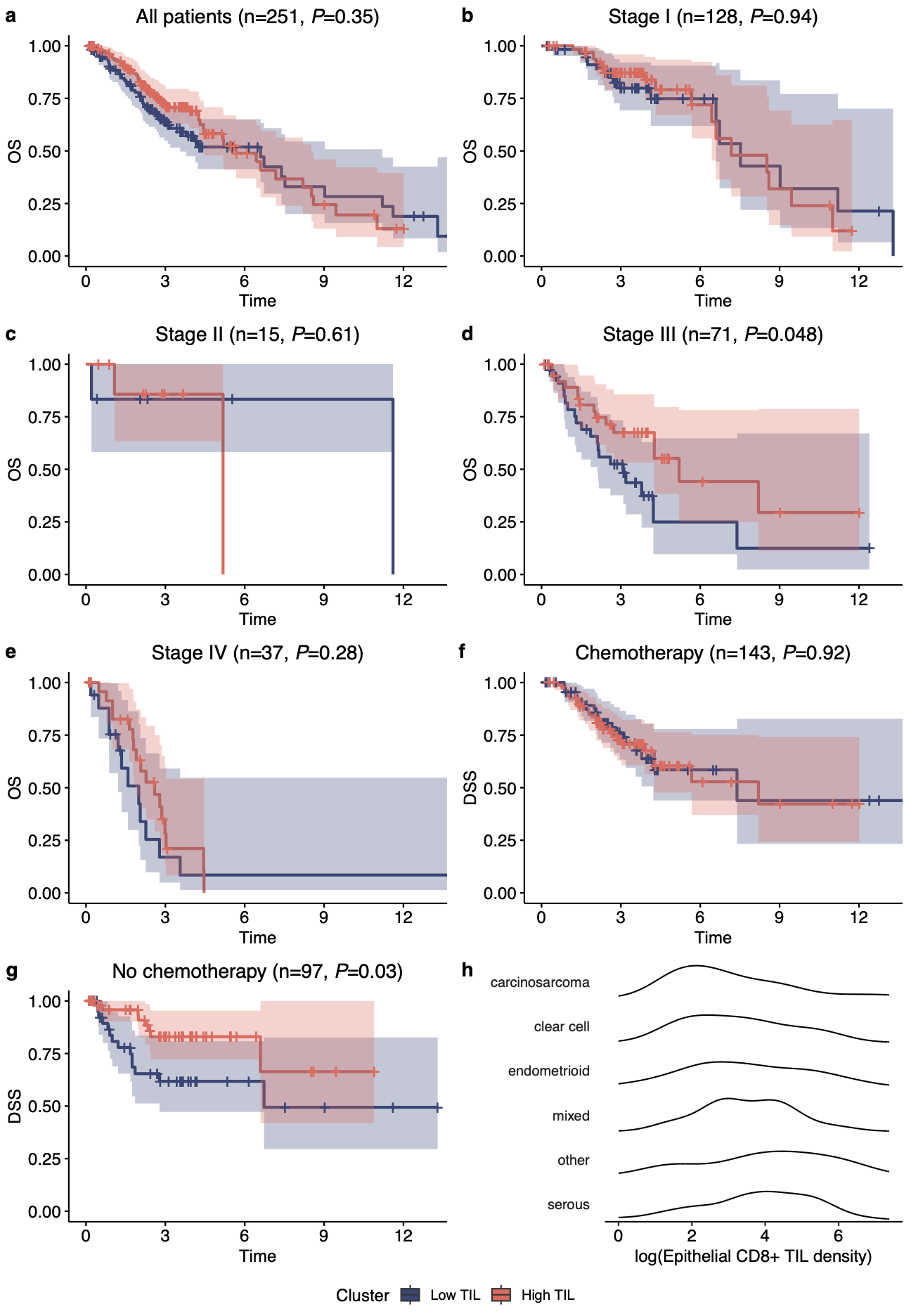
*Table 1:* Hazard ratios, 95% confidence intervals and significance values for TIL cluster and each clinicopathologic variable included in multivariate survival analysis (overall survival, progression-free survival and disease-specific survival).

### Association between TIL cluster and histologic subtypes of p53abn endometrial carcinoma

p53abn endometrial carcinomas comprise a mixture of serous and other histotypes including carcinosarcoma, endometrioid and clear cell. Histotype was significantly associated with TIL cluster (*P* = 6.51e-03), with carcinosarcomas the most TIL-poor histotype and serous carcinomas the most TIL-rich (**Figure 3h**). 74% (23/31) of carcinosarcomas were TIL-poor, compared to only 43% 97/224 of non-carcinosarcoma histotypes (adjusted *P* = 0.011) (**Figure 2b**). Apart from carcinosarcomas, there were no significant pairwise differences in TIL cluster distribution between any of the other non-serous histotypes and serous carcinomas (the most common histotype). In multivariate survival analysis accounting for carcinosarcoma histotype and TIL cluster, carcinosarcomas trended towards shorter overall, progression-free and disease-specific survival, consistent with prior findings (53–55), but this was not statistically significant (all *P* > 0.067). The association between the TIL-rich cluster and overall survival remained significant when accounting for the carcinosarcoma histotype (HR 0.65, 95% CI 0.426-0.989, *P* = 0.044), suggesting that TIL cluster is prognostic independently of histotype.

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| *Figure 3*: Univariate Kaplan-Meier survival curves and log-rank *P* values of overall survival (OS) and TIL cluster, for (a) all tumors in the cohort with known stage and chemotherapy status; (b)-(e) tumors broken down by stage. Univariate Kaplan-Meier survival curves and log-rank *P* values of disease-specific survival (DSS) and TIL cluster (f-g) broken down by adjuvant chemotherapy status. (h) Distribution of epithelial CD8+ TIL densities for each histotype. |

### Immune composition and activity is altered in TIL-rich samples

To further explore anti-tumor immunity and the tumor response to immunity, we assessed CD8+ T cell activation, immune subset composition, and macrophage and tumor expression of immune inhibitory molecules (**Figure S2**). Consistent with our previous findings (14), TIL-rich tumors contained more of both CD8+PD1+ (activated CTL) and CD8+PD1- (naïve T cells), which was expected since CD8+ T cells helped define the TIL-rich group (**Figure S3**). However, activated CTL made up a greater percentage of total CD8 T cells in TIL-rich tumors (*P* = 0.011, **Figure 4a**). Furthermore, the CTL to Treg ratio (CTL:Treg) was significantly elevated within epithelium but not stroma of TIL-rich tumors, suggestive of increased anti-tumor intra-epithelial CTL activity (**Figure 4b**).

Tumor cells and macrophages upregulate PD-L1 and IDO1 in response to CTL and TH expressed cytokines, thereby inhibiting anti-tumor immune attack (21,22). While there was no significant difference in the density of PD-L1-negative macrophages between TIL-rich and TIL-poor tumors, PD-L1+ and PD-L1+IDO1+ macrophages were significantly enriched in TIL-rich tumors (both *P* < 0.001) (**Figure 4c**), as were PD-L1+ and IDO1+ tumor cells (all *P* < 0.001) (**Figure 4d**). Together, these results demonstrate that TIL-rich tumors were not only enriched for immune cells, but also suggest that the immune cells actively participated in anti-tumor immunity that the tumor attempted to resist.

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| *Figure 4*: Expression of immune checkpoint molecules in TIL-rich vs TIL-poor p53abn endometrial cancer. (a) Relative proportions of CD8+ TIL that express PD-1 for each TIL cluster. (b) Relative abundances of CD8+ T cell versus T regulatory cells (CD8/Treg) in TIL-rich and TIL-poor tumors. (c) Densities of CD68+ macrophages that express PD-L1, IDO1, both or neither in TIL-rich vs TIL-poor cases. (d) Proportions of tumor cells that express PD-L1, IDO1 or both in TIL-rich vs TIL-poor cases. *P* values (Mann-Whitney *U* test) corrected for multiple comparisons within each heading are shown. |

### Immune infiltration varies independently of targetable genomic alterations

Finally, we evaluated the relationship between TIL subgroups and therapeutically targetable genomic properties. We leveraged shallow whole genome sequencing data-derived copy number signatures described in endometrial cancer (38) to identify homologous recombination deficient (HRD) tumors (targetable with PARP inhibition) and HER2-amplified tumors (targetable with anti-HER2 antibodies) in a subset of 126 tumors of our cohort. The HRD signature showed no significant association with TIL cluster (**Figure 5a,b**). Furthermore, TIL cluster failed to correlate with HER2 IHC status (*P* = 0.95) or HER2 amplification by shallow whole-genome sequencing (*P* = 0.11) (**Figure 5c,d**). In addition, densities of individual TIL types in epithelial and stromal regions did not significantly correlate with any mutational signature. Thus, the TIL-rich signature is distinct from the HRD and HER2 signatures, possibly expanding the patient cohort amenable to targeted therapies.

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| *Figure 5:* (a,b) Shallow whole genome sequencing derived copy number signature vs TIL-cluster. For each sample (n = 126), the contribution of each of five copy number mutation signatures to each sample is shown in a heat map (a) with comparisons between TIL-poor and TIL-rich cases highlighted in (b). No mutational signature was significantly enriched for TIL-rich cases. (c-d) Number of TIL-rich samples (red) in comparison to TIL-poor samples (blue) by (c) HER2 IHC score and (d) HER2 copy number status (HLAmp = high-level amplification). |

## Discussion

Historically considered one of the less immunogenic subtypes of endometrial cancer, p53abn tumors had, until recently, received minimal attention in immunotherapy research. However recent phase III clinical trials of immune checkpoint inhibitors in addition to standard-of-care chemotherapy demonstrated benefit in both MMRd and MMRp cancers (31,32), with p53abn cases responding substantially better than NSMP cases (mean hazard ratios of 0.41 and 0.87 for overall survival, respectively) (33). These results motivated us to systemically profile the immune microenvironment in a large, multi-institutional cohort of p53abn endometrial cancers. Analysis revealed two immunologically distinct subgroups defined by extensive (TIL-rich) and limited (TIL-poor) infiltration of T cells, B cells and myeloid cells. Over half of cases were highly infiltrated by TIL, challenging the belief that p53abn cancers are immune depleted (56). Moreover, these cases had increased markers of an active immune response, including higher CTL:Treg ratios, higher percentages of CTL expressing PD-1 (indicating active CTL), and increased PD-L1 expression by tumor cells and macrophages, suggesting tumors were responding to active immune attack. In multivariate analysis, we found that TIL-rich tumors were associated with longer overall and disease-specific survival, in contrast to prior findings in smaller cohorts (13,14). Notably, the TIL-rich group was strongly associated with survival in patients with advanced stage III disease and those who did not receive adjuvant chemotherapy. Finally, we found that TIL-rich tumors failed to associate with either homologous recombination deficient (HRD) signature or HER2-overexpressing tumors. Our findings may help to inform personalized treatment of p53abn endometrial carcinoma by helping to identify patients most likely to respond to immunotherapy, PARP inhibition, HER2-directed therapies, and chemotherapy.

In analyzing the immune composition of p53abn endometrial cancer, we found multiple differences from HGSOC and triple-negative breast cancer (TNBC), which morphologically and genomically resemble p53abn endometrial cancer (2). In HGSOC and TNBC, we (36) and others (57) identified a stroma-restricted pattern of TIL, which was associated with inferior prognosis compared to tumors with intraepithelial TIL. In the current study, the stroma-restricted TIL subtype was absent in p53abn endometrial carcinomas. Moreover, whereas immune infiltration correlated with HRD in HGSOC (36,58,59), immune infiltration failed to correlate with HRD or any other known mutational processes in p53abn endometrial carcinoma. The differences we observed in immune infiltration patterns between HGSOC and p53abn EC have also been reflected in immunotherapy clinical trial results: MMRp endometrial carcinomas show striking benefit from immunotherapy compared to HGSOC (31,33,60). Thus, p53abn endometrial carcinomas and HGSOC and TNBC have therapeutically relevant differences in immune microenvironment dynamics and anti-tumor immunity.

Among all p53abn tumors in our cohort, TIL-rich tumors were associated with overall and disease-specific survival in multivariate analysis; however, when stratified by stage and subsequent chemotherapy exposure, we found that the most significant associations were among stage III patients and those who did not receive adjuvant chemotherapy. Stage III and IV endometrial carcinoma encompasses tumors with extension beyond the uterus, either into the uterine serosa, adjacent organs or into lymph nodes. While we failed to identify associations with survival in stage IV disease, likely due to lack of sample size (n = 37), our results suggest that pre-existing anti-tumor immunity plays a greater role in patients with high stage disease compared with tumors confined to the uterus. Our finding that TIL-rich tumors were associated with improved survival in patients that did not subsequently receive chemotherapy highlights several theoretical and practical considerations when choosing adjuvant therapies. Chemotherapy can have conflicting effects on anti-tumor immune responses. It can cause immunogenic tumor cell death (61–63) and release neoantigens into an inflammatory milieu to activate anti-tumor immunity. Conversely, it can inhibit the anti-tumor immune response by eliminating intratumoral anti-tumor T cells and preventing proliferation of tumor-specific immune cells, particularly when delivered at or near the maximum tolerable doses (62,64). Given that the association between TIL-rich tumors and survival was most pronounced in patients who did not subsequently receive chemotherapy, we hypothesize that the immunosuppressive effects of chemotherapy trumped the immunogenic effects in our cohort (which notably did not receive immunotherapy). Conventional chemotherapy alone is strongly associated with survival in p53abn EC and should remain a mainstay of treatment (**Table 1**) (6,65); however, careful titration and selection of chemotherapy agents to optimize immunogenic cell death may help to enable synergy with immunotherapy (62). An alternative hypothesis is that TIL-poor tumors may have converted to TIL-rich tumors by the addition of chemotherapy, obviating the differences between TIL-poor and TIL-rich tumors assessed at the time of surgery (66). Further work including multiplex examination of recurrent disease is required to determine the effects of chemotherapy on anti-tumor immunity in p53abn endometrial carcinoma.

The presence of a TIL-rich group of p53abn endometrial cancers associated with increased survival could help justify improved targeting of immunotherapies beyond molecular subtype-based strategies (10,67). While TIL-rich tumors are more frequent in *POLE*mut and MMRd (13,14) than other molecular subtypes, the variance within molecular subtype is greater than the variance between subtypes. Our data suggest immune profiling may help identify p53abn EC patients with intrinsic anti-tumor immune responses that potentially may be augmented by immunotherapy. Histotype also fails to identify TIL-rich tumors, with carcinosarcoma being the only histotype significantly associated with TIL. Whether carcinosarcomas will show poor response to immune checkpoint inhibition has yet to be determined, but our study indicates that TIL-cluster, not molecular subtype or histotype, may be more informative for stratifying patients with p53abn endometrial cancer for immunotherapy.

Our data examined for the first time the relationship between immune response and mutational processes in p53abn endometrial cancer. Despite genomic similarities between HGSOC and p53abn endometrial cancer (2,36), TIL-rich tumors were not correlated with mutational signatures, including HRD, in p53abn endometrial cancer. In contrast to HGSOC, p53abn endometrial cancer may elicit TIL responses through mechanisms independent of HRD. Other mutational processes generating widespread genomic instability in endometrial cancer (38) may elicit TIL responses through common cGAS-STING pathway activation (68,69). Additionally, we found that TIL groups failed to correlate with HER2 status by immunohistochemistry or whole-genome sequencing. Thus, immunotherapies, anti-HER2 therapies, and PARP inhibitors targeting HRD tumors may represent orthogonal approaches effective in different groups of p53abn endometrial cancers, with some tumors susceptible to multiple agents.

As the use of immunotherapies extends to MMRp endometrial cancers, the immune microenvironment must be considered in addition to molecular subtype as a relevant factor. Our findings highlight properties of the immune microenvironment that may portend susceptibility to immune checkpoint susceptibility in p53abn endometrial cancers and demonstrate the potential for rational personalized therapy for this deadly disease.

## Acknowledgments

The authors would like to acknowledge the expert opinion and guiding mentorship of Dr. Naveena Singh. Her enthusiasm for gynecological pathology and glowing personality will be sorely missed. Our hearts go out to her family and loved ones. We also thank Samuel Leung and the Cross Canada Consortium for data access and help with clinical data acquisition (46).

## Funding statement

This work was funded by the Canadian Cancer Society Uterine Carcinosarcoma and Aggressive Uterine Cancer Research Grant (#707034) the Health Research BC, and the Terry Fox Research Institute (#1116). This team has also been supported through funding from the Canada Research Chairs Program (JMc), the BC Cancer Foundation (Clinician Scientist Award (JMc)), the Chew Wei Chair in Gynecologic Oncology (JMc), the Vancouver Coastal Health Research Institute (Mentored Clinician Scientist Investigator Award (AJ)), and the Miller Mindell Fellowship (AJ) through the VGH & UBC Hospital Foundation.

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