FOXP3+ T cells in uterine sarcomas are associated with favorable prognosis, low extracellular matrix expression and reduced YAP activation

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Uterine sarcomas are rare but deadly malignancies without effective treatment. Immunotherapy is a promising new approach to treat these tumors but has shown heterogeneous effects in sarcoma patients. With the goal of identifying key factors for improved patient treatment, we characterized the tumor immune landscape in 58 uterine sarcoma cases with full clinicopathological annotation. Immune cell characterization revealed the overall prevalence of FOXP3+ cells and pro-tumor M2-like macrophages. Hierarchical clustering of patients showed four tumor type-independent immune signatures, where infiltration of FOXP3+ cells and M1-like macrophages associated with favorable prognosis. High CD8+/FOXP3+ ratio in UUS and ESS correlated with poor survival, upregulation of immunosuppressive markers, extracellular matrix (ECM)-related genes and proteins, and YAP activation. This study shows that uterine sarcomas present distinct immune signatures with prognostic value, independent of tumor type, and suggests that targeting the ECM could be beneficial for future treatments.

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INTRODUCTION
Uterine sarcomas are a heterogeneous group of rare neoplasms comprising 3–4% of all uterine malignancies. Despite their rarity, they are responsible for considerable mortality and morbidity by frequently recurring and metastasizing distantly. Uterine sarcomas are classified into various histopathological subtypes, and treatment is subtype-specific. These subtypes include the most frequently diagnosed, uterine leiomyosarcoma (LMS), and rarer subtypes like low-grade endometrial stromal sarcoma (ESS) and high-grade endometrial stromal sarcoma including YWHAE-FAM22 translocation-bearing tumors (YFAM), adenosarcoma, and undifferentiated uterine sarcoma (UUS). Current treatments include surgery, chemotherapy, radiotherapy, and hormone therapy. However, their poor effectiveness highlights the urgent need for new therapies.

Immunotherapies are promising treatments that exploit the immune system to eliminate cancer cells. Immune evasion is essential for cancer cells to survive and is achieved by manipulating immune checkpoint pathways, normally used to maintain self-tolerance. Checkpoint inhibitors have shown promising results in the treatment of various tumor types but the efficiency of these treatments in uterine sarcomas remains low. The success of immunotherapies relies on the ability to manipulate the tumor immune microenvironment (TIME) into an anti-tumorigenic state. Checkpoint inhibitor treatments can accomplish this, but their efficacy depends on the expression of targetable immune regulatory proteins (IRPs) and the cellular composition of the TIME. Among these IRPs, the PD-1/PD-L1 pathway, IDO1, and B7-H4 have sparked special interest.

The immune cell repertoire of the TIME is another determining factor for the efficacy of immunotherapies. Tumors rich in CD8+ cytotoxic T lymphocytes (CTLs), frontline defensive cells for targeting and killing tumor cells, often are associated with good prognosis and immunotherapy response. Moreover, targeting exhausted CTLs can enhance antitumor immunity. Other promising targetable components of the TIME include regulatory T cells (Treg), CD4+ T cells, and macrophages. The immune inhibitory function of Tregs is generally considered to have pro-tumorigenic effects. Similarly, although macrophages are presented as a phenotypic spectrum, anti-inflammatory or M2-like macrophages are generally associated with poor prognosis while pro-inflammatory or M1-like macrophages are a marker of good prognosis in various cancer types.

In this study, we aimed to characterize the TIME in uterine sarcomas in order to identify immune signatures with therapeutic potential.

RESULTS
Patient cohort
Clinicopathologic characteristics are presented in Supplementary Table 1. A total of 58 malignant tumor cases were included, comprising 13 leiomyosarcomas (LMS), 16 endometrial stromal sarcomas (ESS), 26 undifferentiated uterine sarcomas (UUS), and 3 YWHAE-FAM22 endometrial stromal sarcomas (YFAM). In addition, there were 14 benign leiomyomata controls (LM). All the samples were obtained from primary tumor cores of patients that had not received any prior treatment. The mean follow-up time for LMS

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was 4.99 years (range 0.55–16.42), for ESS 10.88 years (range 2.93–20.67), for UUS 5.02 years (range 0.09–20.38), and for YFAM 11.16 years (range 2.61–18.79). For LMS 11/13 (84.6%) patients were deceased at last follow-up, for ESS 2/16 (12.5%), for UUS 22/26 (84.6%), and for YFAM 2/3 (66.7%).

Multiplex immunofluorescence reveals T cell and macrophage heterogeneity across sarcoma subtypes

To characterize the TIME in uterine sarcomas, tumors were analyzed for T cell and macrophage infiltration, and for the expression of immune regulatory molecules. Characterization of T cells included the CTL marker CD8 and the Treg marker FOXP3 (Fig. 1a). Besides cells expressing a single marker, co-expression of CD8+ and FOXP3+ was also detected. Double CD8+FOXP3+ cells have been described as Treg cells with phenotypic resemblance to classical CD4+FOXP3+ Treg cells.15,16 However, for the purposes of this paper, they have been analyzed separately. Infiltration of T cells was heterogeneous within each tumor type, with no specific type-dependent differences (Fig. 1b–e). The exception was CD8+FOXP3+ cells, which were detected in roughly half of ESS and UUS cases (Fig. 1f, g) but were completely absent in all other cases.

Macrophages were characterized using the CD68 and CD163 markers, allowing quantification of M1-like (CD68+CD163−) and M2-like (CD68+CD163+) populations (Fig. 1h). Macrophage infiltration was heterogeneous within each tumor type, with no specific type-dependent differences (Fig. 1i, j). Yet, anti-inflammatory M2-like macrophages were more prevalent than M1-like macrophages in all tumor types, with most tumors presenting over 60% M2-like macrophages (Fig. 1k).

The expression of potentially targetable IRPs, including PD-1, PD-L1, B7-H4, and IDO1, was also characterized by immunofluorescence (Fig. 2a). Expression of PD-1, B7-H4, and IDO1 was heterogeneous in the different tumor types (Fig. 2b–d) with LMS tumors presenting a higher B7-H4 expression than ESS and UUS. Expression of PD-L1 was not detected in any group (Fig. 2e, simultaneously stained control tissue showed expected positive staining, Supplementary Fig. 1a).

Further classification of cells co-expressing PD-1 with CD8 and
FOXP3, indicative of T cell exhaustion or activation, revealed that PD-1 is expressed in a subset of FOXP3+ but not in CD8+ cells (Fig. 2f, simultaneously stained control tissue showed CD8+ PD-1+ expressing cells, see Supplementary Fig. 1b). Moreover, the proportion of PD-1-expressing FOXP3+ cells was generally low (Fig. 2g; 7.0% in CD8-FOXP3+ and 3.9% in CD8-FOXP3-PD-1+). Interestingly, a large proportion of PD-1+ cells did not express either CD8 or FOXP3. As CD8-FOXP3+ and CD8-FOXP3-PD-1+ cells are likely to be CD4+ Treg and exhausted or activated CD4+ T cells, respectively, we assessed the CD4+ cell infiltration in these tumors by immunohistochemistry (Fig. 2h). Infiltration of CD4+ cells was heterogeneous in all tumor types and positively correlated with CD8-FOXP3+ and CD8-FOXP3-PD-1+ cell infiltration (Fig. 2i, j), suggesting that a considerable proportion of these cells are CD4+.

Fig. 2  The expression of immune regulatory proteins (IRPs) is heterogeneous in uterine mesenchymal tumors. a Representative IF staining of an IRPs-expressing tumor (left), its corresponding H&E staining (right), and representative images of cells expressing PD-1, B7-H4, and IDO1 (bottom). Upper scale bars indicate 100 μm and lower panel scale bar indicates 20 μm. b–e Quantification of IRP-expressing cells shows a heterogeneous expression of PD-1+ (b), IDO1+ (c), B7-H4+ (d), whereas PD-L1 is not detected in any tumor group (e). f Percentage types of PD-1 expressing cells. g PD-1 expression in FOXP3+ cells. h Representative examples of IHC staining for CD4+ T cells representing high and low infiltration tumors. Scale bar indicates 100 μm. i Quantification of CD4+ T cell density shows heterogeneity in CD4+ expression in all tumor types. j Scatter plot shows the significant correlation between IHC stained CD4+ and IF stained CD8-FOXP3+ and CD8-FOXP3-PD-1+ T cells (r indicates Pearson correlation coefficient). Plots indicate average ± s.d. Significance is indicated as **(P < 0.01).
Uterine mesenchymal tumors can be divided into four intrinsic immune cell subtypes

To better understand the nature of the immune cell microenviron-
ment, we performed unsupervised clustering of individual
patients based on immune cell infiltration and expression of IRPs.
This analysis resulted in clustering of patients based on the tumor
immune cell infiltrate rather than the specific tumor type (Fig. 3a).
Three distinct immune cell groups were identified. These immune
groups were defined by macrophages, FOXP3+ IRP negative, and
IRP together with CD8+ cells (Fig. 3b). Patient clustering revealed
two major clusters. One cluster included tumors with high
immune cell infiltration (hot tumors; four LMS, three ESS, six
UUS, and three LM). A second major cluster generally presented
low immune cell infiltrates (cold tumors; nine LMS, 13 ESS, 20 UUS,
and three YFAM). Clustering of cold tumors further revealed three
specific subclusters. These included a FOXP3-high (immune
excluded; one LMS, four ESS, one UUS, one YFAM, and two LM),
a group of tumors expressing IRP5 (immune ignored; four LMS,
four ESS, 11 UUS, and eight LM), and a third subgroup expressing
either FOXP3+PD-1+ or some IRP/CD8+ markers (nonspecific;
four LMS, five ESS, eight UUS, two YFAM, and one LM). These
results again indicate that immune cell infiltrates are largely
independent of the current tumor subtype classification.

Finally, we assessed the prognostic value of these patient
groups, excluding the benign cases. Comparing patient overall
survival between hot and cold tumors did not show any major
differences in overall patient survival (Fig. 3c). Similarly, overall
survival (OS) did not show statistical differences when cold tumors
were further divided into the three subgroups (Fig. 3d).

Correlation with clinicopathologic data shows that FOXP3+ T
cell and M1-like macrophage infiltration are associated with
improved overall survival

We next evaluated the relationship between OS and individual
immune cell infiltrates. Considering all cases, improved OS was
significantly associated with the presence of FOXP3+ cells
(Fig. 4a–c) but not with CD8+ or CD4+ cells (Fig. 4d, e and
Supplementary Fig. 2a). Moreover, an abundance of M1-like
macrophages was associated with better OS, while M2-like
macrophages or the M1/M2 ratio showed no association (Fig. 4f
and Supplementary Fig. 2b, c). Dividing the cases by tumor subtype,
the FOXP3+ survival improvement was observed in UUS, and as a
nonsignificant trend in ESS (p = 0.0269 and p = 0.0954,
respectively), but not in LMS (Supplementary Fig. 3). The M1-like
macrophage prevalence was associated with survival improvement,
specifically in UUS patients (p = 0.0296, Supplementary Table 2).
Neither PD-1, IDO1, nor B7-H4 expression showed general or tumor
type-specific prognostic value (Supplementary Fig. 4a–c and
Supplementary Table 3). However, absence of PD-1-expression in
FOXP3+ cells was a marker of good prognosis (Fig. 4g–j). Contrarily,
the infiltration of PD-1+ cells that do not express either CD8 or
FOXP3 was associated with improved OS (Fig. 4k).

In other tumor types, the CD8+/FOXP3+ ratio (CFR) has been
defined to have a stronger association with patient survival than
individual cell types17–19. We investigated the impact of this ratio
on survival in ESS and UUS as these contained both CD8+ and
FOXP3+ cells. Cases with few tumor-infiltrating lymphocytes (Low
TILs) were considered as a third group. Low CFR (<1) (i.e., tumors
with a larger proportion of FOXP3+ than CD8+ cells) was
associated with significantly improved survival, compared to both
CFR[High] and low TILs (Fig. 4l). This association was stronger in ESS
than in UUS (Supplementary Fig. 5a, b). Together, these data show
that infiltration of FOXP3+ cells (either as an absolute density or
CFR), M1-like macrophages, and CD8-FOXP3-PD-1+ cells
(i.e., presumed CD4+PD-1+ cells), are associated with improved
overall survival, suggesting that these cells are involved in
modulating tumor aggressiveness.

High CFR associates with ECM signaling genes

To further understand the biological underpinnings of the
observed survival differences, we first compared the expression
of immune marker and cytokine genes between CFR[High] and
CFR[Low] or Low TILs. CFR[High] tumors presented upregulation of
markers associated with immunosuppression such as the T cell
exhaustion markers HAVCR2 (TIM3) and ENTPD1 (CD39), and
immune evasion markers including the inhibitor of macrophage
phagocytosis function CD47 and CD274 (PD-L1) (Supplementary
Fig. 6a). However, no significant differences were observed in the
expression of cytotoxicity markers, while the T cell activation
marker CD69 was upregulated in CFR[High] tumors. Cytokine gene
expression showed the anti-inflammatory HGF as the most highly
upregulated gene, as well as TGFβ1 and FG7, upregulated in
CFR[High]. Several pro-inflammatory cytokine genes such as IL2 and
TNFSF13B (BAFF) were instead downregulated in these tumors
(Supplementary Fig. 6b). These results suggest that CFR[High] tumors
present generally more anti-inflammatory and immune-escaping
characteristics compared to CFR[Low] tumors.

Subsequently, to capture other differences between these
tumors, we compared the general gene expression of CFR[High] and
all other tumors, which revealed 267 differentially expressed
genes (Supplementary Data 1). Specifically, comparing CFR[High]
and CFR[Low] tumors resulted in 524 differentially expressed genes,
indicating that these tumors differ considerably at the gene
expression level. Contrarily, transcriptomic analysis of tumors
expressing high and low M1-like macrophages showed 18
differentially expressed genes, indicating that these tumor groups
are transcriptionally similar (Supplementary Data 2). Then, we
conducted a pathway analysis to uncover major pathway
alterations between CFR[High] and other tumors. Major alterations
were found in pathways that included extracellular matrix
organization, proteoglycans in cancer, regulation of cell adhesion,
and integrin-mediated signaling (Fig. 5a). Similarly, independently
comparing tumors with high vs low CD8+ cells or tumors with
high vs low FOXP3+ cells, showed ECM-related pathways as the
most significantly altered ones (Supplementary Fig. 7). MCODE
analysis for protein–protein interactions showed two densely
connected neighborhoods (Fig. 5b). Gene ontology enrichment
analysis of these two MCODE neighborhoods linked MCODE1 with
non-integrin membrane–ECM interactions, integrin pathway,
and laminin interactions, and MCODE2 with smooth muscle
contraction, muscle contraction, and actin cytoskeleton organization.
These results indicate that differences in tumor CFR are related to
ECM-cell adhesion and signaling at the transcriptional level.

Previously, we identified an ECM gene signature in a subgroup
of UUS tumors characterized by a trend towards an increased
incidence of lymphovascular invasion and very poor patient
outcome20. Comparing tumor CFR and ECM signature revealed
that five out of eight CFR[High] tumors presented the ECM gene
signature (Fig. 5c). Moreover, 75% (six out of eight) of CFR[High]
cases presented lymphovascular invasion compared to 62% (8 out
of 13) of the other cases. Finally, to further uncover the biological
characteristics of CFR[High] tumors, we identified likely upstream
regulators and drugs targeting the genes upregulated in these
tumors by Ingenuity Pathway Analysis and Connectivity Map,
respectively. These analyses revealed TNF, TGFβ1/2, PDGF-BB, and
TEAD as upstream regulators (Fig. 5d), all of which are involved in
tissue fibrosis21–23. Furthermore, verteporfin and chlorophenesin
were uncoupled as potential gene perturbing drugs (Fig. 5e).
Interestingly, verteporfin is a suppressor of the YAP-TEAD complex
downstream of ECM-integrin signaling, and chlorophenesin is a
muscle relaxant24,25. Thus, to investigate whether the YAP-TEAD
complex was differentially activated in the different CFR groups,
we compared the expression of known YAP-TEAD target genes26.
Out of 22 target genes, eight were significantly upregulated in
CFR[High] compared to CFR[Low] tumors, indicating higher YAP-TEAD
activity (Fig. 5f). Altogether, these results suggest that there is a link between ECM signaling and infiltration of distinct T cell populations in uterine sarcomas.

**CFR**<sup>low</sup> tumors show reduced ECM protein expression and YAP nuclear localization

To validate the ECM gene expression differences, we analyzed the protein expression of collagen I, VI, fibronectin, and MMP14, as these proteins are significantly enhanced in tumors with the ECM gene signature. In UUS tumors, the expression of collagen I, collagen VI, and fibronectin was lower in CFR<sup>low</sup> than in the CFR<sup>high</sup> group, although fibronectin was not significant (p = 0.0699) (Fig. 6a). Similarly, the expression of these proteins was lower in CFR<sup>low</sup> ESS tumors compared to CFR<sup>high</sup>, although MMP14 expression was similar (Fig. 6b). To validate the possible implication of YAP signaling in defining the tumor CFR, we compared the YAP nuclear to the cytoplasmic ratio in the different CFR-based groups. In conjunction with ECM protein expression and YAP-TEAD target gene expression, the proportion of nuclear YAP was higher in CFR<sup>high</sup> compared to CFR<sup>low</sup> and Low TILs, significantly in UUS tumors (Fig. 6c, d). Together, these data...
support the correlation between ECM expression, YAP activation, and high CD8+ /low FOXP3+ in filtration in ESS and UUS, suggesting a mutual regulation of these components in the tumor microenvironment.

DISCUSSION

Uterine sarcomas are aggressive and rare mesenchymal tumors that currently lack effective treatment. The biology of sarcomas and their interaction with their microenvironment is complex and influences the potential utility of immunotherapy. There is a critical need to understand the sarcoma immune microenvironment and the molecular mechanisms behind it, in order to identify new therapies.

In the work reported here, we have used multiplex immunofluorescence to identify TIL and macrophage populations across a variety of uterine sarcomas and to investigate the expression of potentially targetable IRPs. Further, we have used transcriptomic and protein level data to identify potential pathways involved in the observed TIME differences. These experiments have led to several surprising findings.

First, the immune microenvironment is largely independent of tumor type. Previous studies have shown that immune cell accumulation is associated with diverse sarcoma subtypes, where dominant infiltration of TAMs was observed in UUS and undifferentiated pleomorphic sarcomas. In line with these reports, our results indicate that the immune microenvironment depends on factors other than the tumor type. This is seen when considering the infiltration of individual immune cell types and seen again in our identification of intrinsic immune cell subgroups. This finding carries an important translational relevance; namely, that immunotherapy trials should be performed with a well-planned translational component, so that observed responses can be related to the TIME in those tumors. These hypothetical predictive biomarkers may then be "agnostic" to the sarcoma subtype and should be evaluated as such. This could greatly assist with the clinical management of sarcomas, where a large number of histopathologic subtypes can make traditional "subtype-specific"
trials difficult. Instead, trials should focus on the varieties of TIME and intrinsic immune cell subgroup present.

Second, we have demonstrated that the nature of the immune cell infiltrate has an influence on overall survival. The presence of FOXP3+ cells and M1-like macrophages was associated with better prognosis. This association was observed in both tumors enriched with CD8− and CD8+ FOXP3+ cells. Classifying tumors according to their CD8 to FOXP3 ratio (CFR) indicated that low CFR is associated with improved survival. This result is counter-intuitive, given that in most tumors it is the presence of CD8+ cells that is associated with a better prognosis. Our results indicate that CD8+ infiltration plays a much less important role in uterine sarcomas. The current knowledge of FOXP3+ cell function in uterine sarcomas is limited. However, our data indicate that these cells play a central role in uterine sarcoma progression and are associated with better prognosis. Contrarily, a previous study identified FOXP3+ Treg cells to associate with high tumor grade and poor survival in a cohort including various soft tissue sarcoma types, suggesting that our findings are limited to uterine sarcomas or that the detected FOXP3+ cells are not functionally Tregs. The good prognosis associated with FOXP3+ cells opposes the general tumor-promoting role of these cells, but similar observations have been made in colorectal cancer, head and neck carcinoma, and Hodgkin lymphoma, among others. One possible explanation for these effects is that a significant proportion of the FOXP3+ cells detected are not suppressive Treg cells, rather non-suppressive cells with low expression of FOXP3, which secrete...

Fig. 5 Tumors with high CD8+FOXP3+/−FOXP3+ ratio show upregulation of ECM-integrin interaction pathways in our UUS cohort. a Metascape pathway analysis reveals the most significantly altered pathways in the CFRHigh group (CD8+FOXP3−/FOXP3+ >1) and their interactions. b MCODE-identified neighborhoods of densely connected proteins based on upregulated genes in the high CFR group showing their corresponding enriched Gene Ontologies. e Heat map of gene expression of significantly altered genes in the CFR-defined groups with patient clustering indicating their corresponding ECM signature as defined by Binzer-Panchal et al. d Ingenuity pathway analysis (IPA) indicates the potential upstream regulators of the differential gene expression between CFR high and low tumors. e Connectivity Map analysis shows potential gene expression-altering drugs for the upregulated genes in the CFRHigh compared to CFRLow groups. f Expression of 22 YAP-TEAD target genes in CFR groups and Low TILs. Significance is indicated as * (P < 0.05), ** (P < 0.01), *** (P < 0.001).
pro-inflammatory cytokines, and are associated with better prognosis in colorectal cancer\(^31\). This hypothesis is supported by the expression of pro-inflammatory factors observed in high FOXP3-expressing CFR\(^{\text{High}}\) uterine sarcoma tumors. As these non-Treg FOXP3\(^+\) cells lack the expression of CD45RA, including this marker for further characterization of this cell population may shed light on the function of FOXP3\(^+\) cells in uterine sarcomas and other tumors.

Third, the results from our transcriptomic analysis indicate that pathways such as extracellular matrix organization, proteoglycans in cancer, regulation of cell adhesion, and integrin-mediated signaling are associated with the observed differences in the TIME. Predicted drug regulators of these altered genes highlighted verteporfin and chlorphenesin. Verteporfin is an inhibitor of the YAP-TEAD transcription factor complex\(^32\). There is currently a clinical trial planned for the use of liposomal verteporfin in the treatment of recurrent glioblastoma multiforme, and previous reports indicate that verteporfin can modulate the TIME through PD-L1 inhibition\(^33\). Although our findings show that uterine sarcomas present low PD-L1 expression, independent responses of the YAP-TEAD inhibition may lead to the regulation of the TIME in these tumors.

Finally, analysis of protein expression confirms the role of the ECM regulatory proteins in the observed differences in immune cell infiltration. The aggressive nature of CFR\(^{\text{High}}\) tumors may emanate from the enhanced ECM expression and YAP activation observed through their immune regulatory and invasion-promoting functions. Accumulation of CD8\(^+\) T cells occurs in collagen-rich areas\(^34\). This indicates that, although the density of CD8\(^+\) cells may be enhanced in collagen-rich tumors, their motility and function may be impaired. Moreover, the phenotype-modulating function of the ECM has been shown to promote M2 polarization in macrophages and TGF\(\beta\)-dependent Treg differentiation\(^35,36\). Thus, the lower expression of ECM and TGF\(\beta\) in high FOXP3\(^+\) CFR\(^{\text{Low}}\) tumors further supports the notion that these cells may not have a Treg function.

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**Fig. 6** ECM protein and nuclear YAP expression is enhanced in UUS and ESS tumors with high CFR. **a, b** Representative IHC images and quantification of indicated protein expressions show higher expression in the CFR\(^{\text{High}}\) group in UUS (a) and in ESS patients (b). Scale bars indicate 100 \(\mu\)m. **c, d** Representative IHC images of YAP expression and quantification of its nuclear/cytoplasmic ratio shows a higher ratio in the CFR\(^{\text{High}}\) patient group in UUS (c) and in ESS patients (d). Upper panels scale bars indicate 100 \(\mu\)m and lower panels scale bars indicate 20 \(\mu\)m. Plots indicate average ± s.d. Significance test is indicated *\((P < 0.05)\), **\((P < 0.01)\).
The differential YAP activity between tumor groups indicates that YAP signaling may play a role in the distinct behavior of uterine sarcomas. YAP is clearly a central regulator of ECM signaling and is involved in sarcomagenesis. Thus, our data suggests that YAP is involved in uterine sarcoma progression and may have immune regulatory functions. Indeed, tumoral YAP expression prevents the activation of CTLs by inhibiting PRF1 and GZMB expression in pancreatic cancer.

Altogether, these results show that, independently of tumor type, tumors with an ECM signature present a similar immune microenvironment. Thus, characterizing tumors for specific TME signatures including immune cells and ECM might be a better approach to precision therapy than solely using traditional histopathologic subtypes.

These results have numerous translational implications. First, they indicate that clinical trials evaluating immune therapies will need to include a strong translational component to identify what immune cell signature can be related to a response. Furthermore, any immune cell signature that can be related to a response should then be evaluated in a tumor agnostic manner, given the heterogeneity observed. The central role of YAP and ECM pathways in these differences indicates that therapies focusing on modulation of the ECM should be considered.

METHODS
Patient cohort and central review
The retrospective cohort used in this study contains material from two collaborating centers, the Karolinska University Hospital and Skåne University Hospital. Ethical approval was obtained from the relevant authorities. All cases were reviewed centrally, and the cases of endometrial stromal sarcoma and UUS were analyzed with RT-PCR for detection of the YWHAE-FAM22 translocation. Details of the histopathological review have been described previously by Binzer-Panchal et al. Patients gave their written informed consent. This study was approved by Stockholm’s regional ethical review board (Dnr2010/1916-31/1).

Tissue microarray preparation
Formalin-fixed paraffin-embedded (FFPE) primary tumors were punctured at 1 mm diameter in two different sites of the tumor core. Therefore, using two distinct tumor core sites per patient minimized the effect of the inherent heterogeneity of the tumor immune microenvironment. Consecutive cuts of these TMAs were used to conduct different stainings.

Multiplex immunofluorescence
Tissue microarray (TMA) slides were air-dried at room temperature, deparaffinized, and then rehydrated in phosphate-buffered saline (PBS) (P4417 Sigma-Aldrich, St. Louis, MO, USA) for 15 min. Primary antibodies (Supplementary Table 4) were diluted in 0.3% Triton X-100 (X100, Sigma-Aldrich, St. Louis, MO, USA) containing 0.1% NaN3 in PBS pH7.4 and applied to the slides for 16 h at 4°C. Afterward, sections were washed three times in Tris–Tween 20 (0.05%) (TBS–Tween 20, T9039 Sigma-Aldrich, St. Louis, MO, USA) for 15 min each. The sections were then incubated in a Tris–Tween 20, T9039 Sigma-Aldrich, St. Louis, MO, USA) for 15 min. Primary antibodies were then incubated in Tyramide Signal Amplification Kit, or with 0.03% H2O2 for 10 min, 1 min H2O, and 10 min PBS for collagen I, collagen VI, and MMP (LEM) antibody (Supplementary Table 4). The TSA method was used to visualize the immunohistochemistry signals.

Immunohistochemistry
GenChip ST Arrays (GeneChip Human Gene 2.1 ST Array Plate) were hybridized, washed, stained, and finally scanned with the GeneTitan Multichannel (MC) Instrument, according to the GeneTitan Instrument User Guide for Expression Array Plates (PN 702933, Thermo Fisher, Scientific Inc., Life Technologies).

The RNA raw data were normalized and compared using the software Affymetrix Expression Console Software provided by Thermo Fisher. Pathway analysis of differentially expressed genes was conducted by Metascape.

Immunofluorescence quantification
Qupath (v.0.2.0-m2) (https://github.com/petebankhead/qupath) open-source software, was used for single-cell detection on 3 TMAs with IF staining for CD8, FOXP3, PDCD1, CD68, CD163, and DAPI. TMA slides that held 80 of the cases were dehydrated, and then detected.

Unsupervised patient clustering
The average cell densities of two cores from each patient was used to create unsupervised patient clustering. The analysis was conducted in R v.4.0.3 (R Core Team, 2020) with a heatmap2 package.

Gene expression analysis
The RNA expression arrays method has been previously described. Briefly, RNA from each sample was used to generate amplified and biotinylated sense-strand cDNA from the entire expressed genome according to the Sensation Plus FFPE Amplification and WT Labeling Kit (P/N 702088, Rev.4 Thermo Fisher Scientific Inc., Life Technologies). GeneChip ST Arrays (GeneChip Human Gene 2.1 ST Array Plate) were hybridized, washed, stained, and finally scanned with the GeneTitan Multichannel (MC) Instrument, according to the GeneTitan Instrument User Guide for Expression Array Plates (PN 702933, Thermo Fisher, Scientific Inc., Life Technologies).

Statistical analysis
Descriptive statistics were calculated and presented in tables. For the comparison of the density within groups and intragroup, non-parametric ANOVA (Kruskal–Wallis test) and one-way ANOVA testing were used, respectively. The overall survival (OS) probabilities were estimated and presented by Kaplan–Meier survival curves. The correlation between densities was obtained using a two-tailed Pearson correlation coefficient. All statistical tests were two-sided. P values less than 0.05 were considered statistically significant. Prism v8.0 software (GraphPad) and R programming were used for statistical analysis.

Reporting summary
Further information on research design is available in the Nature Research Reporting Summary linked to this article.
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Conception and design and development of methodology: J.W.C. Acquisition of data: O.G., J.G.-M., E.H., G.K., L.M.-G., N.M., J.M., A.I., K.L. and J.W.C. Analysis and interpretation of data: O.G., J.G.-M., E.H., D.S., K.L. and J.W.C. Writing, review, and/or revision of the manuscript: O.G., J.G.-M., E.H., D.S., K.L. and J.W.C. Data organization and database construction: O.G., J.G.-M., E.H., J.M., A.I. and J.W.C. Study supervision: K.L. and J.W.C.

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The authors declare no competing interests.

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