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Highly multiplexed spatial analysis identifies tissueresident memory T cells as drivers of ulcerative and immune checkpoint inhibitor colitis



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Colitis after different checkpoint inhibitor regimens come with unique features

Intra-epithelial CD8⁺ T cells are more abundant in anti-PD-1 \pm anti-CTLA-4 colitis

Cytotoxicity is highest in activated, cycling CD8⁺ tissue-resident memory T

Inflammation is spatially focused below the epithelium, in the lamina

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Highly multiplexed spatial analysis identifies tissue-resident memory T cells as drivers of ulcerative and immune checkpoint inhibitor colitis

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SUMMARY

Colitis is a prevalent adverse event associated with immune checkpoint inhibitor (ICI) therapy with similarities to inflammatory bowel disease. Incomplete mechanistic understanding of ICI colitis curtails evidence-based treatment. Given the often-overlooked connection between tissue architecture and mucosal immune cell function, we here applied imaging mass cytometry (IMC) to gain spatial proteomic insight in ICI colitis in comparison to ulcerative colitis (UC). Using a cell segmentation pipeline that simultaneously utilizes high-resolution nuclear imaging and high-multiplexity IMC, we show that intra-epithelial CD8⁺ T cells are significantly more abundant (and numerically dominant) in anti-PD-1 \pm anti-CTLA-4-induced colitis and UC. We identified activated, cycling CD8⁺ tissue-resident memory T_(RM) cells at the lamina propria-epithelial interface as drivers of cytotoxicity in ICI colitis and UC. Moreover, we found that combined ICI-induced colitis featured highest granzyme B levels both in tissue and serum. Together, these data reinforce CD8⁺ T_{RM} cells as potentially targetable drivers of ICI colitis.

INTRODUCTION

Treatment of various advanced malignancies has considerably improved with the introduction of immune checkpoint inhibitors (ICIs). However, a major downside associated with ICI therapy remains its large variety of immune-related adverse events (irAEs).¹ ICI colitis is among the most prevalent irAEs at approximately 16% with combined anti-cytotoxic T lymphocyte-associated protein 4 (α CTLA-4) and anti-programmed death 1 (α PD-1) therapy (cICI) and is potentially lethal if not adequately treated.^{2,3} Depending on severity, ICI colitis requires ICI discontinuation, administration of high-dose systemic steroids, or selective immunosuppression in steroid-refractory cases.^{4,5} Step-up to selective immunosuppression is largely expert-opinion driven and mostly based on experience in the treatment of patients with inflammatory bowel disease (IBD). Infliximab and vedolizumab are therefore frequently used.⁶ Notwithstanding clinical parallels between ICI colitis and IBD, both conditions should be considered separate diseases. This is underscored by lesser chronicity in histopathology, superior biological response rate, and shorter time-to-response in ICI colitis than IBD.^{6–9}

ICI colitis can be subdivided based on the specific ICI regimen, i.e., α CTLA-4 monotherapy, α PD-1 monotherapy, or combined (c)ICI. Clinically, time-to-onset of α PD-1 colitis is usually longer than for α CTLA-4-based regimes, while response to steroids is better for α PD-1 colitis than for colitis after α CTLA-4 +/- α PD-1.^{10,11} Immunohistochemistry studies have characterized α CTLA-4 colitis as CD4⁺ T lymphocytic disease with increased interferon (IFN)- γ and interleukin (IL)-17 responses, intra-epithelial neutrophils, and erosions.^{12,13} In contrast, α PD-1 colitis has been associated with CD8⁺ intra-epithelial lymphocytosis,¹³ although another study that also reported overlapping histopathology between ulcerative, Crohn's, and α CTLA-4 colitis found lower CD8 staining scores for α PD-1 than for α CTLA-4 colitis.⁹ With the advent of single-cell RNA sequencing (scRNA-seq), enabling highly comprehensive phenotyping, the role of CD8⁺ tissue-resident memory (T_{RM}) cells could be explored in more detail.^{14–17} Some reports indicate a relative decrease of colonic CD8⁺ T_{RM} cells in ICI colitis compared to ICI-treated patients

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without colitis,^{14,16} possibly due to either an influx of circulating effector T cells, or transformation of resting T_{RM} into inflammatory effectors with the loss of residency markers. CD8⁺ T_{RM} cells are also considered pro-inflammatory actors in IBD,¹⁸ although, similar to ICI colitis, a relative decrease in CD8⁺ T_{RM} cells in active IBD versus healthy control tissue has been found.^{19,20}

Interestingly, intra-epithelial CD8⁺ T_{RM} cells have been shown to transcriptionally acquire an innate pro-inflammatory profile in inflammatory sites of Crohn's ileitis,²¹ illustrating the relevance of epithelial-lamina propria localization in relation to cell function. Yet, transcriptomic studies on tissue in ICI colitis inherently lack architectural insight. Incomplete pathophysiologic understanding of causes for the clinically relevant differences observed between ulcerative colitis (UC) and ICI colitis after different ICI regimens prompts a direct in-depth comparison among these groups. Using imaging mass cytometry (IMC), we aimed to integrally characterize immune cell, particularly T cell, infiltrates in different types of ICI colitis and UC, both spatially and highly multiplexed at the same time. We adopted an IMC analysis approach to reliably segment and annotate single cells and we directly compared UC to different subtypes of ICI colitis. Our study identifies CD8⁺ T cells as the dominant immune cell population in α PD-1 and cICI colitis and shows that activated, cycling CD8⁺ T_{RM} cells residing close to the epithelial border are drivers of cytotoxicity across different types of ICI colitis and UC.

RESULTS

Study design and population

Colon biopsies from 18 patients with ICI colitis (4 α CTLA-4 monotherapy, 5 α PD-1 monotherapy, 9 cICI) and 5 patients with UC were included for sequential DAPI nuclear imaging and IMC ("IMC cohort"; Figures 1A and 1B; Table S1). Nine of 23 patients (39%) were female, median age was 67 years (p25–p75, 51–72) and most patients with ICI colitis were treated for melanoma (78%). The majority of patients (61%) were steroidnaive upon endoscopy (median 0 days of steroid use; p25–p75, 0–1). Three of five patients with UC and all ICI colitis patients underwent endoscopy for new-onset disease. In addition, we included serum samples from 80 ICI-treated patients, of whom 39 developed clinically relevant irAEs including 14 (36%) with colitis, for multiplexed proteomics. These serum samples were collected at baseline and upon onset of irAEs or ± 6 weeks after treatment initiation for irAE-free patients ("Serum cohort"; Figure 1A; Table S1). Four cICI-treated patients included in the IMC cohort were also part of this Serum cohort. Finally, we reanalyzed previously published colonic CD3⁺ scRNA-seq data (GEO accession number: GSE144469) to validate main findings from the IMC cohort (Figure 1A).¹⁴

First, we explored to what extent histopathologic and endoscopic disease severity indices, commonly used in UC and increasingly in ICI colitis, were correlated. Within the IMC cohort, a modest association was observed between both the Geboes score for "erosion/ulceration" and the Robarts Histopathology Index (RHI) with the Mayo endoscopic score. The Geboes score for "chronic inflammatory infiltrate" showed a moderate correlation with symptom duration (Figures S1A–S1C). All tissue samples histologically featured inflammation in line with the diagnoses of ICI colitis or UC and were therefore included for further analysis.

CD8⁺ T cells are the dominant immune cell population in *α*PD-1 and combined-ICI colitis

We acquired H&E-stained, DAPI, and IMC images for all 23 patients with colitis in the IMC cohort (Figure 2A). Using the MATISSE cell segmentation pipeline that simultaneously takes advantage of high-resolution DAPI nuclear imaging in combination with high-multiplexity IMC,^{22,23} we identified 215,293 single cells across all patient samples. After removal of cells from excluded tissue regions (including artifacts, lymphoid follicles, and submucosa), 184,975 single cells remained for data normalization, scaling, and lineage determination. These cells were used to generate pseudo-color cell type annotated images as shown in Figure 2A. Due to imaging-inherent overlap of membrane signal between neighboring cells, especially in densely populated infiltrates as seen in colitis, we deliberately developed a *supervised* cell type annotation approach. A comparable strategy previously achieved reliable annotation of segmented intestinal immune cells.²⁴ Indeed, a uniform manifold approximation and projection visualization overlaid with supervised annotation labels confirmed that unsupervised clustering alone would probably insufficiently distinguish biologically distinct cell clusters (Figure 2B). We excluded one UC patient (IBD 1_B) after cell type annotation from further analysis because of ill-annotated cells due to low signal-to-noise ratio. H&E-stained images of the remaining samples are shown in Figures S2–S4.

We confirmed that all cell types were present across patients in comparable and biologically plausible frequencies (Figure 2C). The ~25% non-classified non-immune cells most likely represent stromal cells, which is numerically in line with previous IMC data in IBD patients.²⁴ Based on epithelial masks, we additionally assigned each cell to the intra-epithelial or lamina propria compartment. While some variation in the ratio of intra-epithelial to lamina propria cells existed among subjects, average ratios between colitis groups were similar (Figure S5A). As expected, we observed a high relative fraction of epithelial cells in epithelium and complete absence in lamina propria (Figure 2D). Immune cells were relatively more abundant in lamina propria than epithelium, which was most evident for CD4⁺ T cells. CD8⁺ T cells were prevalent in both the lamina propria and intra-epithelial compartment (Figure 2D). Specifically, CD8⁺ T cells numerically dominated the intra-epithelial immune cell composition of α PD-1 and cICI colitis and were significantly higher compared to α CTLA-4 colitis and UC (Figure 2D). The same trend of CD8⁺ T cell dominance in α PD-1 and cICI colitis could be observed in lamina propria. Epithelial macrophages were significantly reduced in cICI versus α CTLA-4 colitis, while relative abundance of CD4⁺ T cells, B cells, Tregs, dendritic cells, and $\gamma\delta$ T cells was similar between groups per compartment. Confounding of cell composition by age or tissue sectioning was unlikely; age was not associated with abundance of distinct cell types (Figure S5B) and with increasing fractions of (intra-)epithelial cells, only relative abundance of intra-epithelial dendritic cells (DCs) significantly decreased (Figure S5C).







Figure 1. Study design and participant characteristics

(A) All analyses were performed with data from (1) combined DAPI-imaging and imaging mass cytometry (IMC) of colitis tissue (n = 23); (2) paired serum samples from baseline and either 6 weeks after α PD-1 monotherapy or combined α CTLA-4 + α PD-1 treatment initiation, or upon irAEs (n = 80); and (3) previously published colonic T cell single-cell RNA-sequencing (scRNA-seq) data from patients with colitis after α CTLA-4 with or without α PD-1 treatment (n = 8) and healthy controls (n = 8).¹⁴ Created with BioRender.com.

(B) Characteristics of patients in the IMC cohort. Characteristics that apply to a given patient are indicated by yellow spheres. See also Table S1 and Figure S1.

Tissue-resident memory CD8⁺ T cells are more abundant in epithelium than lamina propria and show high cytotoxic potential in most colitis types

The clinical course of cICI colitis is often more acute and severe than ICI monotherapy-induced colitis.^{1,10} We explored if specific characteristics of the immune infiltrate could explain the clinical differences observed among colitis subtypes. Increased expression of cytotoxic effector protein granzyme B in CD8⁺ T cells has been reported in a broad range of inflammatory conditions, including Crohn's disease and UC.²⁵ In our study, CD8⁺ T cell cytotoxicity, quantified by CD8⁺ T cell granzyme B level, was significantly higher in cICI- than in ICI monotherapy-induced colitis or UC (Figures 3A and 3B). The extent of CD8⁺ cytotoxicity did not correlate with any clinical or histological measure of colitis severity, including RHI, Common Terminology Criteria for Adverse Events grade, and Mayo endoscopic score (Figures S6A–S6C). We next evaluated the association between granzyme B levels in peripheral blood and the occurrence of irAEs, in particular ICI colitis. Proteomic









Figure 2. Cell type distribution across individual samples after cell segmentation and supervised lineage determination

(A) Representative sample images from patients with ulcerative colitis (top) and combined α CTLA-4 and α PD-1 colitis (bottom). Shown are hematoxylin & eosinstained images (adjacent serial section) used for histopathologic annotations, DAPI nuclear images, pseudo-color composite mass cytometry images, and finally representations of extracted single-cell events after data normalization, scaling, and lineage determination, pseudo-colored by cell type.

(B) Uniform manifold approximation and projection (UMAP) visualization of a random subset comprising 10% of cells across all patients with IMC data of sufficient quality (n = 22), colored by assigned cell type label with color-coding as in (C).

(C) Stacked bar graph representing fractions of cell types across all patient samples. clCl denotes "combined α CTLA-4 and α PD-1", UC "ulcerative colitis". (D) Boxplots showing relative fraction (range 0–1) of each cell type, stratified by tissue compartment, for all patients. Statistical testing for differences across groups separately per tissue compartment by Kruskal-Wallis test followed by Dunn's post-hoc test with Benjamini-Hochberg false discovery rate correction. Abbreviations clCl and UC as in (C). *p < 0.05. See also Figures S2–S5.

measurements in serum showed that irAEs after cICI indeed are associated with significantly higher granzyme A, B, and H levels as compared to α PD-1 treatment alone (Figure 3C). In addition, significantly higher circulating levels of IFN- γ , IL12R β 1, and TNF- α were observed following cICI, compared to α PD-1 treatment in patients with irAEs. Although limited by smaller sample size, the same trend of an enhanced Th1-asso-

ciated response with higher serum granzyme B levels was visible when only considering patients who developed ICI colitis (Figure S6D). Subsequently, we phenotypically characterized CD8⁺ T cells based on the expression of classical tissue residency markers CD69 and CD103.²⁶ We grouped CD69^{lo}CD103^{lo}, CD69^{hi}CD103^{lo}, and CD69^{hi}CD103^{hi} (termed T_{RM}) CD8⁺ T cells.²⁷ To this end, we defined "low expression" as below-median expression of CD69 or CD103 across all CD8⁺ T cells and vice versa for "high expression". As expected, the classical T_{RM} phenotype was more abundant in epithelium than lamina propria in all colitis groups (Figure 3D). Moreover, we found that granzyme B levels were highest in T_{RM} CD8⁺ T cells in all colitis subtypes, except αCTLA-4 colitis (Figure 3E). Thus, our data suggest that the clinical disparities between different types of colitis may be partly due to difference in abundance of CD8⁺ T_{RM} cells and higher granzyme B levels in CD69^{hi}CD103^{hi} T_{RM} than non-T_{RM} CD8⁺ T cells.

Activated, cycling tissue-resident memory CD8⁺ T cells below the epithelial border are key cytotoxic actors in colitis

IMC uniquely offers the opportunity to enrich phenotypical data with spatial information. We first assessed if CD8⁺ T cell tissue localization was associated with cytotoxic potential and therefore calculated distance to the nearest epithelial border for all CD8⁺ T cells, as shown by example in Figure 3F. Then, we developed a mixed-effects model embedding clinical, phenotypical, spatial, and functional (activation and proliferation status) covariates to delineate the contribution of each variable to total CD8⁺ T cell granzyme B production. Based on this, an activated (MHC-II^{hi}), cycling (Ki67^{hi}) CD8⁺ T_{RM} phenotype was highly significantly ($p < 10^{-50}$ for individual covariates) associated with higher granzyme B production (Figure 3G). The presence of this phenotype was visually confirmed in mass cytometry images of different colitis subtypes (Figure 4). In adjusted analysis, cells adjacent to the epithelium produced more granzyme B than cells distant from epithelium and intra-epithelial cells (Figure 3G), suggesting that inflammation was most pronounced at the lamina propria-epithelial interface. To evaluate how robust these findings were in relation to colitis subtype, we repeated the analysis stratified by colitis group. We found that the significant association between an activated, cycling CD8⁺ T_{RM} phenotype below the epithelial border and high granzyme B production was present across colitis subtypes, although the effect of mucosal localization was not significant in α PD-1 colitis (Figure S7A). In addition, we found that age was not independently associated with CD8⁺ T cell granzyme B expression (Figure S7B). However, CD8⁺ T cell characteristics alone could not completely explain highest granzyme B production in cICI colitis, as indicated by the effect through cICI treatment, adjusted for other covariates (Figure 3G).

Based on increased serum levels of Th1-associated cytokines in clCl relative to α PD-1 irAEs (Figure 3C), we hypothesized that enhanced CD4⁺ T cell help, T cell priming, or activation might additionally underlie high clCl colitis cytotoxicity. Therefore, we also investigated if colitis subtypes differed with respect to cell-cell interactions using neighborhood analysis.²⁸ We assessed interactions between all cell types and visualized interactions between epithelial cells, CD4⁺ and CD8⁺ T cells, CD4⁺ Tregs, DCs, B cells, and $\gamma\delta$ T cells. General tissue organization in epithelium and lamina propria compartments was, as expected, reflected by significant avoidance between CD8⁺ T and epithelial cells, and CD4⁺ T and epithelial cells in both directions (Figure S8). Based on agglomerative clustering, patients could be divided in two groups. Significant interactions among adaptive immune cells, and DCs in addition influencing the adaptive response, presented as an important difference separating the group containing all UC patients from the other group (p = 0.03 by Fisher's exact test, Figure S8). However, none of the ICI colitis subtypes were significantly enriched in one of both adaptive immune cell interaction-based subgroups. In summary, neighborhood analysis did not yield evidence in support of cICI-specific enhanced interactions between CD8⁺ T cells and other (adaptive) immune cells to additionally explain high cICI cytotoxicity.

Activated CD8⁺ $T_{(RM)}$ cells are confirmed as drivers of ICI colitis at the transcriptomic level

To validate CD8⁺ T_{RM} cells as important mediators of inflammation in ICI colitis, we reanalyzed colonic CD3⁺ scRNA-seq data from eight patients with ICI colitis after α CTLA-4 with or without α PD-1 therapy and eight healthy controls.¹⁴ We selected *CD8A*-expressing cells from the CD3⁺ T cell pool, clustered the CD8⁺ T cells, and annotated clusters based on the top 6 cluster-defining genes along with other known markers (Figures 5A, 5B, S9A, and S9B). Clusters 4 and 11 represented (activated) CD103⁺ T_{RM} clusters (*ITGAE*, *KLRB1*, *CD38*, *HLA-DRA*), while cluster 0 was identified as a CD103⁻ T_{RM} subset (*ITGB2*, *KLRG1*). Other subsets included (a mixture of) intra-epithelial cytotoxic and $\gamma\delta$ T cells (clusters 2, 8, and 10; *ITGAE*, *KIR2DL4*, *CD160*, *AREG*, *TRDC*), mucosal-associated invariant T cells (cluster 12; *SLC4A10*, *TRAV1-2*), central memory/naive CD8⁺ T cells (cluster 6; *CCR7*, *SELL*, *CXCR4*), a cycling subset (cluster 13; *MKl67*), and activated cytotoxic effector T cells



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GZMB

GZMA TNF

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IFN_gamma

Adjusted P value <0.05

a ns

160

120

80

40

Figure 3. Phenotypical, functional, and spatial CD8⁺ T cell characteristics related to cytotoxicity

(A) Density plot displaying granzyme B expression levels in CD8⁺ T cells, stratified by colitis subtype. Solid lines within each graph indicate the colitis subtypespecific median expression level. The dashed line displays the overall median expression level of all CD8⁺ T cells together (n = 2,227–13,430 cells per colitis subtype).

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Figure 3. Continued

(B) Boxplots showing fraction of CD8⁺ T cells with higher-than-median cytosolic granzyme B levels across all CD8⁺ T cells, stratified by tissue compartment. Difference between groups (based on averaged lamina propria and epithelium data points) tested by one-way ANOVA, followed by Tukey's post hoc test. clCl denotes "combined α CTLA-4 and α PD-1", UC "ulcerative colitis", *p < 0.05.

(C) Volcano plot showing differential serum protein expression of 92 analytes in irAEs after combined α CTLA-4 and α PD-1 relative to α PD-1 monotherapy (only samples obtained upon irAE onset, n = 39). Differential expression analyzed by Wilcoxon test with Benjamini-Hochberg false discovery rate correction.

(D) Boxplots showing fraction with TRM phenotype of all CD8⁺ T cells, stratified by tissue compartment. No formal statistical comparisons were done.

(E) Boxplots showing fraction of CD8⁺ T cells with higher-than-median cytosolic granzyme B levels across all CD8⁺ T cells, stratified by expression levels of CD69 and CD103. Pairwise comparisons were performed by paired Student's t tests with Benjamini-Hochberg false-discovery rate correction within each colitis subtype. clCl denotes "combined α CTLA-4 and α PD-1", UC "ulcerative colitis", *p < 0.05, **p < 0.01.

(F) Representative example (ICI 7_A) of epithelial cells (gray) and CD8⁺ T cells in lamina propria, color-coded by distance to nearest epithelium (based on epithelial mask).

(G) Visualization of coefficients and $-\log_{10}$ (p values) of fixed effects. A mixed-effects model with fixed effects for all listed covariates and random intercepts for individual patients was developed to explain CD8⁺ T cell granzyme B production, using all 27,548 CD8⁺ T cells across n = 22 patients. Coefficients >0.0 indicate that CD8⁺ T cell granzyme B levels are higher in presence of (or for higher values of) the variable. *p < 0.05. See also Figures S6–S8.

with cytolytic activity (clusters 5 and 7; *HLA-DRA, GZMB, PRF1*), of which cluster 5 was terminally differentiated (*KLRG1*⁺, *IL7R*⁻). *GZMB* expression was significantly higher in clusters 5, 7, 8, and 11 compared to all other clusters (Figure 5C), importantly, including a cluster representing activated CD103⁺ T_{RM} cells (cluster 11). Within cluster 11, *GZMB* expression was only significantly increased in patients with colitis, but not in healthy controls, and the majority of cells in clusters 4 and 11 derived from patients with colitis (Figures S9C and S9D). Consistent with IMC data, the CD103⁻ T_{RM} subset (cluster 0) featured low *GZMB* expression (Figure 5C). Our sub-clustering results fit the possible differentiation path from T_{RM} to cytotoxic effector cells as discovered by Luoma et al.¹⁴ An important role for activated CD103⁺ CD8⁺ T_{RM} cells in granzyme B-mediated cytotoxicity is thus confirmed at the transcriptomic level. Moreover, these data show that enhanced granzyme B expression is specifically associated with inflammation, rather than a general T_{RM} feature, as confirmed on the protein level in Figure 4 (bottom row).

DISCUSSION

ICI colitis and UC show several clinical and histological similarities, but it is unknown to what extent they share a common pathophysiology. Here, we described our approach combining DAPI imaging and IMC to acquire high-resolution, highly multiplexed, single-cell images of inflamed colon from patients with α CTLA-4-, α PD-1-, cICI colitis, or UC. We explored on the protein level what cell types are involved in those subtypes of colitis and assessed phenotypical, spatial, and functional T cell characteristics.

In line with other studies, we found that CD8⁺ T cells are relatively increased in α PD-1 and cICI colitis compared to α CTLA-4 colitis and UC.^{13,15,29} We confirmed that activated cycling CD8⁺ T_{RM} cells are important drivers of inflammation in colitis and additionally our data suggest that their cytotoxic potential was highest below the epithelial border. cICI colitis exhibited higher tissue CD8⁺ T cell granzyme B levels compared to all other colitis types, which was corroborated in serum of cICI- and αPD-1-treated patients who developed irAEs including ICI colitis. The fact that clinical and histological colitis severity indices and CD8⁺ T cell cytotoxicity were not correlated, may suggest that CD8⁺ T cell granzyme B levels possibly reflect the induction of inflammation, while factors beyond CD8⁺ T cell biology may collectively determine clinical severity and thus the need for escalated immunosuppressive therapy. Besides, neither CD8⁺ T cell characteristics nor cell-cell interactions could completely account for the colitis subtype-related differences in cytotoxicity. Nevertheless, others have similarly found enhanced CD8⁺ T cell GZMB or GZMK expression in cICI colitis relative to αPD-1 colitis.^{15,16} We previously showed that, in contrast to αPD-1, irAEs after cICI are strongly associated with peripheral blood effector memory CD4⁺ T cell proliferation amid a mainly Th1-associated response.³⁰ In the present study, we confirmed that cICI irAEs come with stronger peripheral Th1-skewing than αPD-1 irAEs. This indicates that enhanced cytotoxicity observed in cICI colitis might be the result of reinvigorated CD4⁺ T cell help, potentially lowering the threshold for CD8⁺T cell activation. Although our study revealed no clues to support this hypothesis, higher CD28 and TNFRSF4 (encoding CD134) expression in CD8⁺ T cells of clCl colitis than α PD-1 colitis shown previously, suggests enhanced T cell receptor signaling in clCl colitis.¹⁶ Of note, whether CD8⁺ T_{RM} cells may have been activated elsewhere before they entered the tissue, or CD8⁺ T_{RM} cells are directly targeted via PD-1 by ICI therapy, or both, cannot be concluded from these studies.

We detected higher CD8⁺ T cell cytotoxicity at the lamina propria-epithelial interface. Since intraepithelial cells have been shown to migrate dynamically between epithelium and lamina propria,³¹ our observation is compatible with the finding that especially intraepithelial CD103⁺ CD8⁺ T cells adopt a cytotoxic profile in Crohn's ileitis,²¹ and that the highest expression of UC-associated loci is found in intraepithelial CD8⁺ T cells of UC patients.³² Although data are conflicting, others have shown substantial clonal overlap and transcriptional similarity between lamina propria and intra-epithelial lymphocytes in the mucosa of healthy controls.^{33,34} Distinguishing two "canonical" T_{RM} populations classified as CD103⁻ (*KLRG1*, *ITGB2*) and CD103⁺ (*ITGAE*, *KLRB1*) T_{RM} subsets, in healthy tissue, the former were associated with highest cytotoxic potential, while the latter produced multiple cytokines like TNF- α and IFN- γ simultaneously.³³ Interestingly, we showed increased granzyme B production in CD8⁺ T cells with *higher* CD103 expression. Our findings are corroborated by others who reported greatest increase in expression of *GZMB* and *IFNG*, or *IL17A* and *IL26* in *ITGAE*-expressing subsets in ICI colitis.^{15,16} This may present as a dissimilarity with Crohn's disease, in which data are conflicting as to whether CD103⁺ or CD103⁻ CD8⁺ T_{RM} cells are most cytotoxic.^{21,35,36} Although some studies reported decreased abundance of intestinal CD8⁺ T_{RM} cells in patients with ICI colitis compared to ICI-treated patients without co-litis,^{14,16} cytotoxic lymphocytes newly emerging in tissue were clonally related to *bona fide* T_{RM} cells.¹⁴ Compared to active inflammation in





Figure 4. Spatial expression patterns of key phenotypical and functional markers associated with CD8⁺ T cell cytotoxicity Representative sample images from subjects with, in rows from top to bottom, αCTLA-4 colitis, αPD-1 colitis, combined αCTLA-4 and αPD-1 (cICI) colitis, ulcerative colitis (UC) and healthy colon tissue. Shown are pseudo-color composite mass cytometry images of the entire tissue slide with insets as indicated and separate pseudo-colored single-marker channels for insets. Examples of activated, proliferating CD8⁺ TRM cells producing granzyme B are indicated

UC, CD8⁺ $T_{(RM)}$ cells are more abundant and more activated in ICI colitis.^{15,29} In summary, an accumulating body of evidence points toward CD8⁺ T_{RM} cells as drivers of inflammation, potentially playing an even more prominent role in ICI colitis than UC.

with white arrows. Within healthy tissue (bottom row), green and orange arrows respectively indicate a CD103+ CD8+ TRM cell without granzyme B

Thus, intestinal CD103⁺ CD8⁺ T_{RM} cells might serve as a therapeutic target in UC and ICI colitis. Blocking the integrin pair $\alpha 4\beta 7$ with vedolizumab has proven effective in treating both UC and ICI colitis, ^{6,37} but does not target αE^+ (CD103⁺) CD8⁺ T_{RM} cells. Etrolizumab is a monoclonal antibody against the $\beta 7$ integrin subunit, with activity against $\alpha 4\beta 7^+$ and $\alpha E\beta 7^+$ cells. Based on cell type signature scores applied to bulk RNA-seq data from IBD patients, etrolizumab reduced the number of both intestinal CD103⁻ and CD103⁺ CD8⁺ T cells.³⁶ However, to date, clinical results with etrolizumab in Crohn's and ulcerative colitis are disappointing and it is currently unclear if the drug will be clinically taken any further.^{38,39} A possible explanation for the lack of efficacy might be that with the depletion of intestinal CD103⁺ T_{RM} cells, not only pro-inflammatory but also homeostasis-promoting T_{RM} subsets are lost. Apart from the limited efficacy of $\beta 7$ integrin blockade in IBD, this strategy may carry a risk in itself when used in patients with cancer. The presence of CD103⁺ tumor-infiltrating lymphocytes (TILs) has been associated with better outcomes in most cancer types, especially of epithelial origin.⁴⁰ TILs associated with tumors of other origin, including melanoma, also express *ITGAE* and *ITGB7* which can together form dimerized CD103.^{41,42} Directly targeting $\alpha E\beta 7$ could negatively impact intra-tumoral retention of TILs and thus compromise antitumor immunity. In order to avoid this kind of adverse effects of T_{RM}-directed treatment, future research should focus on phenotypical differences between irAE-tissue and tumor-associated T_{RM} cells.

As ICI colitis appears dominantly in Th1/Tc1-mediated disease,^{14,15,17,43} Jak inhibition could be another treatment strategy in ICI colitis.⁴⁴ Its safety from a tumor response perspective is presently not well established, however. Finally, upregulation of *IL*17 and *IL*26 differentiated cICI from αPD-1 colitis.¹⁶ Involvement of the IL-23/IL-17 axis has been demonstrated in various irAEs and is more pronounced after cICI.^{30,45,46} Paradoxically, direct IL-17 blocking, e.g., with secukinumab, has been shown to induce intestinal inflammation.⁴⁷ Pro-homeostatic IL-17

expression and a CD103-CD8⁺ T cell highly expressing granzyme B.







Figure 5. Reanalysis of CD8⁺ T cell clusters in colonic CD3⁺ single-cell RNA-sequencing data from ICI colitis patients and healthy controls (A) Uniform manifold approximation and projection (UMAP) visualization showing sub-clustering of CD8A-expressing cells.

(B) Stacked violin plot indicating expression of key subset-defining genes within CD8A-expressing clusters.

(C) Violin plot indicating GZMB expression in CD8⁺ cell clusters; clusters with statistically significantly higher GZMB expression compared to all other CD8⁺ cell clusters together are indicated as follows (by Wilcoxon test with Benjamini-Hochberg false-discovery rate correction): *P_{adj} = 0.0 #P_{adj} = 5.3 × 10⁻¹⁵²; ¹P_{adj} = 3.1 × 10⁻⁶⁰; ¹P_{adj} = 1.7 × 10⁻¹⁸⁶. All plots (A–C) are based on cells from N = 16 ICI colitis patients and healthy controls. See also Figure S9.

production is independent of IL-23,⁴⁸ and therefore more upstream interventions directed against IL-23, such as ustekinumab, or the IL-6 receptor inhibitor tocilizumab might hold promise.

In conclusion, our study importantly contributes to our understanding of the pathophysiology underlying ICI colitis and UC, including disparities among colitis subtypes, through integrated analysis of phenotypical, functional, and spatial aspects. In this way, we underpin the role of CD8⁺ T_{RM} cells as potentially targetable drivers of ICI colitis.

Limitations of the study

Our study has three main limitations. First, this study did not include healthy controls. However, we confirmed antibody binding specificity both with immunofluorescence on control tissue and after isotope labeling with IMC using a tissue microarray, including uninflamed colon tissue that was included on every slide with study participant tissue. This uninflamed colon tissue sample was also used to confirm scRNA-seq reanalysis findings in healthy controls on the protein level. Moreover, studies comparing colon mucosa from healthy controls with mucosa from ICl colitis cases found CD8⁺ T_(RM) cell numbers to be comparable or higher in colitis, and *GZMB* expression in different CD8⁺ subsets is considerably higher in colitis.^{14,15,29} Second, our IMC antibody panel lacked stromal markers. For this reason, cell types such as fibroblasts could not be classified. However, the primary goal of this study was to compare different types of ICl colitis and UC, delineate the abundance of different immune cells for those colitis subtypes, and relate immune cell phenotypes to their cytotoxic potential *within* patients. Moreover, our lineage definition approach accounted for lacking markers of non-immune cells and, therefore, mis-classification of key immune cells of interest such as CD8⁺ T cells could be minimized. Thirdly, the IMC cohort lacked the power to adjust for possible confounders, such as sex and tumor type. However, we found no significant effect of age on immune cell abundance and CD8⁺ T cell cytotoxicity.

STAR***METHODS**

Detailed methods are provided in the online version of this paper and include the following:

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SUPPLEMENTAL INFORMATION

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AUTHOR CONTRIBUTIONS

Conceptualization, M.E., E.B., J.L., B.O., Y.V., K.S., and F.W.; methodology, M.E., M.B., and J.D.; formal analysis and visualization, M.E.; biobank patient inclusions and sampling, R.V., M.E.; performing experiments, J.L., M.A., and E.D.; histopathologic scoring/annotation, M.L.; writing original draft, M.E.; reviewing & editing, all authors.; approval of final manuscript, all authors.; supervision, K.S. and F.W.; funding acquisition, E.B., B.O., Y.V., and F.W.

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STAR*METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
E-cadherin (clone 24E10)	Cell Signaling Technology	Cat# CST3195BF; RRID: AB_2728822
CD68 (clone KP1)	Thermo Fisher Scientific	Cat# 14-0688-82; RRID: AB_11151139
CD14 (clone EPR3653)	Abcam	Cat# ab214438; RRID: AB_2889158
pS6 (clone D57.2.2E)	Cell Signaling Technology	Cat# CST4858BF; RRID: AB_2721245
CD20 (clone H1)	BD Biosciences	Cat# 555677; RRID: AB_396030
CD45RA (clone HI100)	Biolegend	Cat# 304102; RRID: AB_314406
HLA-DR+DP+DQ (clone CR3/43)	Abcam	Cat# ab7856; RRID: AB_306142
CTLA-4 (clone UMAB249)	OriGene Technologies	Cat# UM800141CF; RRID: N/A
RORγt (clone RORC/2941)	Abcam	Cat# ab268233; RRID: N/A
IFNγ (clone D3H2)	Cell Signaling Technology	Cat# CST8455BF; RRID: AB_2797644
CD45RO (clone UCHL1)	Cell Signaling Technology	Cat# CST55618BF; RRID: AB_2799491
CD103 (clone EPR4166(2))	Abcam	Cat# ab221210; RRID: AB_11142856
TNFα (clone 7B8A11)	Proteintech	Cat# 60291-1-lg; RRID: AB_2833255
FoxP3 (clone 236A/E7)	Abcam	Cat# ab96048; RRID: AB_10861686
CD4 (clone EPR6855)	Abcam	Cat# ab181724; RRID: AB_2864377
T-bet (clone D6N8B)	Cell Signaling Technology	Cat# CST13232BF; RRID: AB_2616022
CD45 (clone D9M8I)	Cell Signaling Technology	Cat# CST13917BF; RRID: AB_2750898
IL-10 (polyclonal)	R&D Systems	Cat# AF-217-NA; RRID: AB_354401
TIGIT (polyclonal)	Abcam	Cat# ab233404; RRID: AB_2827380
CD8α (clone C8/144B)	Thermo Fisher Scientific	Cat# 14-0085-82; RRID: AB_11150240
ICOS (clone D1K2T)	Cell Signaling Technology	Cat# CST89601BF; RRID: AB_2800142
PD-1 (clone EPR4877(2))	Abcam	Cat# ab186928; RRID: AB_2894896
CD56 (clone NCAM1/784)	Abcam	Cat# ab216010; RRID: AB_2909431
IL-17A (polyclonal)	R&D Systems	Cat# AF-317-NA; RRID: AB_354463
Ki67 (clone B56)	BD Biosciences	Cat# 556003; RRID: AB_396287
Granzyme B (clone D6E9W)	Cell Signaling Technology	Cat# CST46890BF; RRID: AB_2799313
CD3 (polyclonal)	Dako	Cat# A045229-2; RRID: N/A
CD11c (clone EP1347Y)	Abcam	Cat# ab216655; RRID: AB_2864379
IL12Rβ2 (polyclonal)	Thermo Fisher Scientific	Cat# PA5-24181; RRID: AB_2541681
CD69 (polyclonal)	Proteintech	Cat# 10803-1-AP; RRID: AB_2074965
TCR-δ (clone H-41)	Santa Cruz Biotechnology	Cat# sc-100289X; RRID: N/A
Histone H3 (clone D1H2)	Cell Signaling Technology	Cat# CST4499BF; RRID: AB_10544537
Human TruStain FcX (Fc receptor blocking solution)	BioLegend	Cat# 422302; RRID: AB_2818986
Biological samples		
Formalin-fixed colon biopsies from patients with immune checkpoint inhibitor or ulcerative colitis	This study	N/A
Serum from immune checkpoint inhibitor treated patients	This study	N/A

(Continued on next page)

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Continued		
REAGENT or RESOURCE	SOURCE	IDENTIFIER
Chemicals, peptides, and recombinant proteins		
DNA intercalator	Fluidigm	Cat# 201192B; RRID: N/A
4′,6-diamidino-2-phenylindole (DAPI)	Sigma-Aldrich	Cat# D9542; RRID: N/A
Toluidine blue	Sigma-Aldrich	Cat# 89640-25G; RRID: N/A
Xylene	Klinipath	Cat# 4055-9005; RRID: N/A
Tween 20	Sigma-Aldrich	Cat# P9416; RRID: N/A
2-amino-2-methyl-1,3-propanediol (Tris base)	Sigma-Aldrich	Cat# 10708976001; RRID: N/A
Ethylenediaminetetraacetic acid (EDTA)	Sigma-Aldrich	Cat# 27285; RRID: N/A
Bovine serum albumin (BSA)	Sigma-Aldrich	Cat# A9647; RRID: N/A
Critical commercial assays		
MaxPar X8 multimetal antibody labeling kit	Fluidigm	Cat# 201300; RRID: N/A
Olink Target 96 Immuno-Oncology panel	Olink	N/A
Deposited data		
Cell-segmented raw imaging mass cytometry data	This study	DataverseNL: https://doi.org/10.34894/N84F12
Serum multiplex proteomic data (Olink)	This study	DataverseNL: https://doi.org/10.34894/N84F12
Colonic CD3 ⁺ single-cell RNA-sequencing data	Luoma et al. ¹⁴	GEO: GSE144469
Software and algorithms		
Imctools (version 2.1.7)	Bodenmiller lab	https://github.com/BodenmillerGroup/imctools; RRID: SCR_017132
Modified Extended Depth of Field (EDF), Fiji plugin	Forster et al. ⁴⁹	https://bigwww.epfl.ch/demo/edf/#soft; RRID: N/A
Microscope Image Stitching Tool (MIST), Fiji plugin	Chalfoun et al. ⁵⁰	https://github.com/usnistgov/MIST; RRID: N/A
llastik (version1.3.3)	Berg et al. ⁵¹	https://github.com/ilastik/ilastik; RRID: SCR_015246
CellProfiler (version 3.1.9)	Carpenter et al. ⁵²	https://github.com/CellProfiler/CellProfiler; RRID: SCR_007358
Fiji (ImageJ) (version 2.9.0)	Schindelin et al. ⁵³	https://github.com/fiji/fiji; RRID: SCR_002285
R (version 4.2.0)	The R Foundation	https://www.r-project.org; RRID: SCR_001905
Rstudio (version 2022.12.0+353)	Posit, PBC	https://posit.co/products/open-source/rstudio/; RRID: SCR_000432
sf (version 1.0.8, R package)	Pebesma ⁵⁴	https://github.com/r-spatial/sf/; RRID: SCR_023393
umap (version 0.2.9.0, R package)	McInnes et al. ⁵⁵	https://github.com/tkonopka/umap; RRID: N/A
ggplot2 (version 3.4.0, R package)	Wickham ⁵⁶	https://github.com/tidyverse/ggplot2; https://ggplot2.tidyverse.org; RRID: SCR_014601
ggpubr (version 0.4.0, R package)	Kassambra ⁵⁷	https://rpkgs.datanovia.com/ggpubr/; RRID: SCR_021139
pheatmap (version 1.0.12, R package)	Kolde ⁵⁸	https://github.com/raivokolde/pheatmap; RRID: SCR_016418
nlme (version 3.1.160, R package)	Pinheiro ⁵⁹	https://github.com/cran/nlme; RRID: SCR_015655
neighbouRhood (version 0.4, R package)	Schapiro et al. ²⁸	https://github.com/BodenmillerGroup/ neighbouRhood; RRID: N/A
Seurat (version 4.2.0, R package)	Hao et al. ⁶⁰	https://github.com/satijalab/seurat; RRID: SCR_016341
glmGamPoi (version 1.8.0, R package)	Ahlmann-Eltze & Huber ⁶¹	https://github.com/const-ae/glmGamPoi; RRID: N/A
FSA (version 0.9.4, R package)	Ogle ⁶²	https://github.com/fishR-Core-Team/FSA; RRID: N/A
MATISSE pipeline	Krijgsman et al. ²³	https://github.com/VercoulenLab/ MATISSE-Pipeline; RRID: N/A
Normalization, scaling, cell type annotation pipeline	This study	Zenodo: https://doi.org/10.5281/zenodo.7858216





RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Femke van Wijk (f.vanwijk@umcutrecht.nl).

Materials availability

This study did not generate new unique reagents.

Data and code availability

- Single-cell segmented raw IMC data derived from participants with colitis and serum multiplexed proteomic data have been deposited online in DataVerseNL. In compliance with national legislation, this repository is accessible with restrictions via https://doi.org/10.34894/N84F12. Serum data from 3 patients who did not give permission for data sharing outside the European Union have been discarded from this data set.
- An R markdown file with code used for IMC data processing and analysis has been deposited on Zenodo, publicly accessible via https://doi.org/10.5281/zenodo.7858216.
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND STUDY PARTICIPANT DETAILS

Study participants

Colon biopsies were obtained from patients treated with aCTLA-4 monotherapy, aPD-1 monotherapy or combined aCTLA-4 and aPD-1 (cICI) who developed ICI-colitis. All biopsies were obtained during routine clinical procedures. Colon biopsies from UC patients and corresponding clinical data were obtained through the University Medical Center Utrecht Research Data Platform. In suspected ICI-colitis patients, biopsies were taken only from inflamed mucosal sites (or at-random in case of endoscopically uninflamed mucosa) in the left-sided colon. Mayo endoscopic scores were retrieved from endoscopy reports or retrospectively assessed from endoscopic images.⁶³ Tissue samples from ICI-colitis patients were only used if histological assessment confirmed active inflammation and no concurrent gastro-intestinal infection was suspected. UC patients from whom samples were included had to be steroid- and biological-free in the six months prior to endoscopy. Serum was obtained from patients participating in the UNICIT biobank study.³⁰ Serum samples collected at baseline and ±6 weeks into treatment (for irAEfree patients) or upon development of Common Terminology Criteria for Adverse Events (CTCAE) v5⁶⁴ grade ≥2 irAEs demanding hospitalization or ≥0.5 mg/kg steroids were selected for multiplex immunoassay. Detailed patient characteristics for both the IMC and Serum cohort are provided in Table S1. Median age was 67 years (p25–p75, 51–72) and 65 years (p25–p75, 55–73), with 9/23 (39.1%) and 23/77 (29.9%) females, in the IMC cohort and Serum cohort, respectively. No data on participants' race, ethnicity or ancestry were collected as part of this study and neither were these data available from routine care. All patients from whom serum is used participated in the UNICIT biobank study and provided written informed consent in accordance with the Declaration of Helsinki. Ethical approval was received from the UMC Utrecht Biobank Review Committee (Toetsingscommissie Biobanken [TC-bio] 18-123) and permission to use human specimens form this biobank was granted (TC-bio 19-704). Use of anonymous or coded leftover material for scientific purposes is part of the standard treatment contract on an opt-out basis with patients in our hospital. Approval for use of clinical metadata from the pseudonymized UC patients was obtained through TC-bio 18-676.

METHOD DETAILS

Sample collection and histological assessment

For each participant, two adjacent 4 µm-thick tissue sections were prepared from colon biopsies fixed in 10% formalin and paraffinembedded (FFPE). One slide was stained with hematoxylin & eosin (H&E) while the other was used for nuclear imaging and IMC. Serum samples were isolated and stored at -80°C within 4 hours after blood collection. H&E-stained slides of all included samples were assessed by an experienced gastro-intestinal pathologist (M.L), who annotated lymphoid follicles in the regions used for IMC and assessed disease activity employing the Geboes scores,⁶⁵ validated for UC but not for ICI-colitis.⁶⁶ These scores were then used to calculate the Robarts Histopathology Index (RHI).⁶⁷ Histological scoring was based on the highest inflammation score found within the entire tissue section available. Hence, IMC regions did not necessarily feature all histologic characteristics that prompted a certain Geboes score in the same patient.

Slide preparation for microscopy and IMC

After confirming antibody binding specificity with immunofluorescence on tonsil and colon FFPE sections, antibodies were conjugated to lanthanide isotopes with the MaxPar antibody labeling kit (Fluidigm, San Francisco, CA, USA) according to the manufacturer's protocols. Tissue microarrays comprising different types of tissue, including non-inflamed colon, ovarian cancer and tonsil, were present on the FFPE slides used for IMC to validate antibody binding and specificity. FFPE tissue slides were baked (1.5 hours, 60°C), deparaffinized with xylene (20 min.) and rehydrated in a gradient of ethanol (100%; 10 min., followed by 95%, 80%, 70%; 5 min. each). Slides were then washed in MilliQ water



(3 min.) and phosphate-buffered saline (PBS) containing 0.1% Tween (PBST; 10 min.). Heat-induced epitope retrieval in 10mM Tris with 1mM EDTA (pH 9.5, 30 min., in a 95°C water bath), cooling to 70°C and washing in PBST (10 min.) were followed by blocking with 3% bovine serum albumin (BSA) and 1:100 Human TruStain FcX in PBST (1 hour, room temperature [RT]). After removal of the blocking buffer, slides were incubated overnight with the antibody cocktail (Table S2) in PBST with 0.5% BSA (4°C). Slides were then washed three times in PBST. Next, slides were stained with 1:400 DNA-intercalator (Ir-191/Ir-193) and 1:1000 4',6-diamidino-2-phenylindole (DAPI, 60 min., RT) in PBS, followed by two washes with MilliQ water, and air-dried.

Fluorescent microscopy imaging

Slides were imaged on a Zeiss Z1 imager using a 20x dry objective (Zeiss, EC Plan-NEOFLUAR 0.5 NA, 420350-9900) with mercury lamp as light source, a 49 filter set and an Axiocam 503 mono camera system. Using ZEN software (2.6), images were acquired in a tiled Z-stack format (9 Z-slices) with 10% overlap between tiles and exported to individual 16-bit tiff tiles. Single in-focus images were created with the Extended Depth of Field plugin in Fiji and tile images were stitched using the MIST algorithm in Fiji.^{49,50,53}

Imaging mass cytometry and cell segmentation

After acquisition of DAPI images, slides were rinsed in MilliQ water and counterstained with 0.1% toluidine blue (5 min., RT), washed with MilliQ water and air-dried. Mass cytometry of ~1 mm² regions per patient was performed on a Helios (Fluidigm) mass cytometer connected to a Hyperion (Fluidigm) laser ablation module (ablation frequency 200 Hz). Imctools was used to convert data to 32-bit tiff files. For two control tissues included within the tissue microarray, single channel expression data are shown in Figures S10 and S11. Single cell segmentation was performed using the MATISSE segmentation pipeline, combining IMC and DAPI images, as described elsewhere.²³ This combined approach achieves superior cell segmentation compared to IMC-based segmentation alone.²² Briefly, high-resolution DAPI images were registered to IMC images. Training data for IMC images (with annotations for epithelial and non-epithelial cell membranes, epithelial and non-epithelial nuclei and background) and DAPI images (only nuclear annotations) were created by two researchers (M.E. and J.L.). IMC channels used for annotations were E-cadherin, CD68, CD14, CD20, CD45, CD45RA, CD45RO, CD4, CD8α, IL-17A, CD3, CD69 and Ir-193. Probability maps for nuclear and cellular objects were created in Illastik,⁵¹ and cell segmentation was performed with CellProfiler using these probability maps.⁵² Next, based on all pixels within a segmented cell, mean IMC signal expression for all channels was extracted for all segmented cells with R.

IMC data clean-up and normalization

A step-by-step description of our approach for data clean-up, normalization, scaling and annotation is provided via the key resources table. First, artefacts (e.g., antibody aggregates), lymphoid follicles as indicated by the gastro-intestinal pathologist and submucosa were manually annotated in Fiji by two researchers (M.E. and J.L.). Coordinates for boundaries of epithelial regions were extracted from Fiji using the E-cadherin channel, smoothened with Gaussian blur ($\sigma = 2$ pixels [µm]). Next steps were exclusively performed in R. Data were natural-log transformed. Single cells within artefact, lymphoid follicle or submucosa tissue regions were excluded and remaining events were labeled as either "intra-epithelial" or "lamina propria" based on their intersection with epithelial masks using the sf package.⁵⁴

IMC data scaling and cell lineage annotation

For data intensity scaling, we calculated scaling factors, derived from modal marker intensities, for each channel and each patient separately, since we assumed that channel intensity variability resulted from both patient- and channel-specific sources. Then, all data were linearly scaled, effectively aligning channel-specific single-cell expression distributions among patients. Normalization and scaling results were visually checked by histogram representations of single-cell data for all patients, as shown in Figure S12A. We confirmed that scaling factors represented a normal distribution, both over channels and over patients (Figure S12B). We also verified that markers for which expression correlated within cells were also biologically related (Figure S12C).

Subsequently, channel positivity thresholds were defined for markers that were selected for lineage determination. For markers with bimodal expression, a threshold separating the marker-positive and -negative populations was applied. For unimodally distributed markers in which "true positive signal" is contained in the heavy right tail, a normal distribution was derived from the left half of the histogram using the "full width at half maximum" method. Then, a threshold for marker positivity was conservatively set at +1.5 standard deviation (SD). Next, candidate cell type annotations were generated using the Boolean rules in the Table below. Due to imaging resolution, single-cell signal will inevitably contain signal of neighboring/overlapping cells, complicating unsupervised cell type annotation based mostly on membrane markers, especially in dense areas. Therefore, we developed a supervised approach in which each cell could be assigned multiple different candidate cell types. Next, expression levels of key markers (indicated in bold in the Table below) were ranked across all cells from all patients pooled together. For each individual cell, ranks of key markers were then compared between the candidate cell types. The key marker ranked highest eventually determined the assigned cell type label for each single cell. Cells that did not fit the criteria for any of the cell types were labeled "non-classified", or "other immune cell" if they expressed any of CD45 isoforms.



Boolean rules for candidate cell type annotation

Candidate cell type	Marker expression (key marker used for ranking in bold)
CD4 ⁺ T cell	CD45 ⁺ CD4 ⁺ FoxP3 ⁻
CD8 ⁺ T cell	CD45 ⁺ CD3 ⁺ CD8 <i>a</i> ⁺
B cell	CD45 ⁺ CD3 ⁻ CD20 ⁺
Regulatory T cell (Treg)	CD45 ⁺ CD3 ⁺ CD4 ⁺ FoxP3 ⁺
γδ T cell	$CD45^+ TCR\delta^+$
Macrophage	CD14 ⁺ CD68 ⁺ CD3 ⁻ CD20 ⁻
Dendritic cell	CD11c ⁺ CD68 ⁻ CD3 ⁻ CD20 ⁻
Epithelial cell	CD45 ⁻ E-cadherin ⁺
Other immune cell	CD45 ⁺ E-cadherin ⁻ CD20 ⁻ CD3 ⁻ CD4 ⁻ CD8a ⁻
CD45 included CD45 and its isoforms CD45RA and CD45	RO. Markers in bold represent the key markers used for lineage determination.

After cell type annotation, we excluded one UC patient (IBD 1_B) from further analysis because of low signal-to-noise ratio and large withinimage intensity variation across channels, which yielded ill-annotated cells (Figure S13).

IMC data dimensional reduction

Uniform Manifold Approximation and Projection (UMAP) visualizations were created with *umap()* and 15 neighbors for a randomly drawn subset containing 10% of all cells.⁵⁵ Mean single-cell intensities of all channels, except RORyt was used, because data normalization was imperfect for this marker, artificially leading to patient-based separation.

IMC neighborhood analysis

Cell-cell interactions for all cell types (including non-classified cells) were investigated by neighborhood analysis as described in Schapiro et al. using the *neighbouRhood* package with n=10,000 permutations.²⁸ An agglomerative clustering heatmap was built based on interactions of prespecified relevant cell types. Hierarchical clustering was performed using Ward's method based on Manhattan distance.

Multiplexed proteomics

Normalized protein expression (NPX) levels of 92 cytokines and chemokines was measured in serum by proximity extension immunoassay using the Olink® Target 96 Immuno-Oncology panel.⁶⁸ This technique is based on the ligation of two complementary DNA strands that are coupled to panel-specific antibodies. After ligation, double-stranded DNA is polymerase chain reaction (PCR) amplified and the detected DNA tags are following corrections for extension- and interpolation controls translated to NPX values on an (inherently) relative log₂-scale. Thus, 1 NPX difference between two samples represents a doubling of protein expression between samples. To correct for batch effects across plates, data were normalized using 12 bridging controls that were included on all measured Olink plates.

scRNA-seq data reanalysis

Previously published (GSE144469)¹⁴ CD3⁺ T cell scRNA-seq data from 8 α CTLA-4 monotherapy or combined/sequentially ICI-treated patients who developed ICI-colitis and 8 healthy controls was reanalyzed using *Seurat* v4.⁶⁰ Cells with 200-3,000 features and <10% mitochondrial transcripts were kept for further analysis. TCR genes were silenced before normalization, scaling and unsupervised clustering because expression of these genes can affect unsupervised T cell clustering in a biologically unmeaningful way.⁶⁹ SCTransform() was used on each sample individually with vars.to.regress='percent.mt' and method='glmGamPoi' to correct for technical variation including batch effect.⁶¹ Then all objects were integrated using SelectIntegrationFeatures() with 2,000 variable features and clustering was performed with FindNeighbors() to construct a K-nearest neighbor (KNN) graph using the first 30 principal components, followed by FindClusters() with esolution=0.8. CD8⁺ T cell clusters were selected based on mean-normalized *CD4* expression <1 and *CD8A* expression >1, with expression >1 indicating above-average expression. Then we performed clustering within the CD8⁺ cell pool using the same parameters. Cell clusters were annotated based on the top 6 markers defining each cluster, using FindAllMarkers() with min.pct=0.1 and logfcthreshold=0.01, along with known population-defining markers. Finally, normalized and scaled expression data of TCR genes was added back to the clustered data set to visualize the expression of TCR genes specifically associated with mucosal-associated invariant T (MAIT) cells and $\gamma\delta$ T cells.

QUANTIFICATION AND STATISTICAL ANALYSIS

All analyses were performed in R version 4.2.0. Continuous variables were compared between multiple groups with a one-way ANOVA followed by Tukey's post-hoc test (for normally distributed data) or the Kruskal-Wallis test followed by Dunn's post-hoc test with



Benjamini-Hochberg false-discovery rate (FDR) correction (for non-normally distributed data). Spearman's rank coefficient was used for correlations between two continuous variables and associations between categorical variables were tested with Fisher's exact test. Differences in normalized protein expression (NPX) obtained in the serum proteomics measurements was compared by Wilcoxon tests with Benjamini-Hochberg FDR correction. The *nlme* package (v3.1-158) was used to analyze CD8⁺ T cell granzyme B production by a linear mixed-effects model with random intercept for patients and fixed effects for all covariates, fit by restricted maximum likelihood. Two-sided *P*<0.05 was considered statistically significant. Statistical details of individual analyses are reported in the figure legends. Unless otherwise specified, n represents the number of participants.