# 1 DEFICIENT RESIDENT MEMORY T-CELL AND CD8 T-CELL RESPONSE TO

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3 Short title: Deficient resident memory T-cells in IBD

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COMMENSALS IN INFLAMMATORY BOWEL DISEASE

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#### 30 Author Contributions

A.N. and S.C.K. designed the studies and wrote the manuscript. A.N. performed the experimental work and analyzed the data. L.D., P.H., D.R., M.J.O. and T.P. assisted with the experimental work and data interpretation. L.H. and A.L.M. isolated the bacterial strains and provided microbiota expertise. R.M., J.S., S.P.C., P.H., S.B., D.L., P.D., A.W., N.A., A.A. and A.L.H. recruited patients and provided clinical samples. N.A., A.A. and A.L.H. provided clinical insights into the studies. S.R.C. contributed to study design, data interpretation and manuscript preparation. All authors provided input on the manuscript.

Abbreviations: IBD, inflammatory bowel disease; Trm, resident memory T-cells; CD, Crohn's
 disease; UC, ulcerative colitis; HC, healthy control; Treg, regulatory T-cell; IEL, intraepithelial
 lymphocytes; LPL lamina propria lymphocytes; IEM, intraepithelial microbes; ATRA, all-trans
 retinoic acid; FICZ, 5,11-Dihydroindolo[3,2-b]carbazole-6-carboxaldehyde; TNF, tumour
 necrosis factor; IFN, interferon; TGF, transforming growth factor; pDC, plasmacytoid
 dendritic cell; mDC, myeloid dendritic cell; Tfh, follicular helper T cell.

#### 44 Abstract

Background & Aims: The intestinal microbiota is closely associated with resident memory
lymphocytes in mucosal tissue. We sought to understand how acquired cellular and humoral
immunity to the microbiota differ in health versus inflammatory bowel disease (IBD).

Methods: Resident memory T-cells (Trm) in colonic biopsies and local antibody responses to
intraepithelial microbes were analyzed. Systemic antigen-specific immune T- and B-cell
memory to a panel of commensal microbes was assessed.

Results: Systemically, healthy blood showed CD4 and occasional CD8 memory T-cell 51 52 responses to selected intestinal bacteria but few memory B-cell responses. In IBD, CD8 memory T-cell responses decreased although B-cell responses and circulating plasmablasts 53 increased. Possibly secondary to loss of systemic CD8 T-cell responses in IBD, dramatically 54 reduced numbers of mucosal CD8<sup>+</sup> Trm and  $\gamma\delta$  T-cells were observed. IgA responses to 55 intraepithelial bacteria were increased. Colonic Trm expressed CD39 and CD73 56 ectonucleotidases, characteristic of regulatory T-cells. Cytokines/factors required for Trm 57 differentiation were identified, and in vitro-generated Trm expressed regulatory T-cell 58 function via CD39. Cognate interaction between T-cells and dendritic cells induced T-bet 59 60 expression in dendritic cells, a key mechanism in regulating cell-mediated mucosal 61 responses.

62 Conclusions: A previously unrecognized imbalance exists between cellular and humoral 63 immunity to the microbiota in IBD, with loss of mucosal T-cell-mediated barrier immunity 64 and uncontrolled antibody responses. Regulatory function of Trm may explain their 65 association with intestinal health. Promoting Trm and their interaction with dendritic cells

66	rather than immunosuppression may reinforce tissue immunity, improve barrier function
67	and prevent B-cell dysfunction in microbiota-associated disease and IBD etiology.
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#### 75 Introduction

76 Large numbers of lymphocytes reside in the intestinal mucosa and play a key role in barrier 77 function and immune surveillance. Immunity against infection is provided by long-lived memory T-cells reactive to foreign antigens as well as antibody<sup>1</sup>. Memory T-cells can be 78 broadly categorized into circulating (central memory and effector memory) and tissue-79 resident, non-recirculating cells called resident memory T-cells (Trm)<sup>2</sup>. Trm provide potent 80 81 barrier immunity in mucosal tissues due to their high motility, rapid re-activation of effector 82 function and ability to recruit further immune responses via cytokine (e.g. IFN-y) secretion. However, the role of Trm in human disease is unclear, and how they interact with resident 83 84 microbes that make up the intestinal microbiota is not understood.

Inflammatory bowel disease (IBD) is thought to be perpetuated by intestinal microbial 85 dysbiosis leading to episodic colitis (ulcerative colitis, UC) or localized inflammation 86 anywhere along the gastrointestinal (GI) tract (Crohn's disease, CD), mediated by Th17 or 87 88 other subsets of CD4 T-cells<sup>3,4</sup>. Disease etiology involves interaction of multiple genetic 89 susceptibilities with environmental factors including diet and lifestyle factors that can affect 90 the microbiota. However, most studies to date have focussed on sequence-based profiling 91 of microbiomes in disease; how different microbial species interact with the immune system is not well understood. Mouse studies indicate the colonic microbiota is essential for 92 recruiting sufficient CD4 Foxp3-expressing regulatory T-cells (Treg) to the colon to prevent 93 inflammation<sup>5,6</sup>. This suggests that IBD results from a failure of Treg-mediated tolerance to 94 commensals in the GI tract. 95

96	Here we studied memory T-cell responses to a panel of intestinal commensal bacteria in IBD
97	patients and healthy controls, and analyzed Trm populations in the epithelium and lamina
98	propria colonic tissue where they are in close proximity to mucosa-associated microbes. Our
99	data show that underlying disease in human IBD is related to a lack of associated with
100	reduced CD8 T-cell responses to commensal bacteria leading to Trm deficiency in the colon,
101	and chronic B-cell activation and excess IgA secretion associated with a loss of barrier
102	immunity. We also show that human Trm express Treg function and propose specific
103	mechanisms to explain how loss of Trm:dendritic cell interaction could contribute to the
104	development of inflammatory disease.

#### 107 Methods

#### 108 Study Design

The study aimed to determine the role of resident memory T cells in IBD. Donors (age 16-80) 109 were recruited to the study from outpatient clinics of St Mark's Hospital and included those 110 111 with a diagnosis of CD or UC, and healthy donors undergoing investigative endoscopy. None 112 of the CD patients had a history of obstruction, perianal disease or ileitis alone. Patients were recruited over a fixed period determined by ethical permission, no data were excluded 113 114 at the end of the study. Additional healthy blood donors were recruited from hospital staff 115 and visitors. Ethics approval was obtained from the Health Research Authority UK and London Brent Research Ethics Committee. Written informed consent was received from 116 participants prior to inclusion in the study. 117

# 118 Colonic intraepithelial lymphocytes (IEL), lamina propria lymphocytes (LPL) and 119 intraepithelial microbe (IEM) isolation

5 left colon and 5 right colon biopsies (10mg tissue each) were obtained from uninflamed 120 tissue at routine colonoscopy as described<sup>7</sup>. IEL and IEM were released from biopsies using 121 DTT/EDTA and harvested by centrifugation at 300g (5min). IEM were obtained by 122 centrifugation of resulting supernatants at 4500g (20min). LPL were obtained by collagenase 123 digestion of remaining tissue; all cells were phenotyped and counted by flow cytometry. 124 Cells were washed in PBS and stained for viability using LIVE/DEAD Fixable-near-IR stain 125 126 (ThermoFisher) before addition of surface-staining antibodies in fetal calf serum. In some 127 cases cells were then fixed/permeabilized for intranuclear staining using the Foxp3 buffer 128 set (ThermoFisher, as instructions). Antibodies used are listed in supplementary file 1. All

samples were analyzed on a BD Biosciences FACS Canto II and data analyzed by FlowJo
software (Tree Star), with volumetric sampling determined using Perfect-Count
microspheres<sup>™</sup> (Cytognos, S.L).

#### 132 In vitro differentiation of Trm-like cells from human PBMC

133 Naïve CD8 T-cells were purified by magnetic selection from healthy donor PBMC using the 134 naïve CD8 T-cell isolation kit (Miltenyi Biotec) and were >98% CD8<sup>+</sup> and >98% CD45RA<sup>+</sup>. Naïve CD8 T-cells were stimulated with plate-bound anti-CD3 (1µg/ml), soluble anti-CD28 135  $(1\mu g/ml)$  and IL-2 (5ng/ml, Peprotech). Further additions of TGF- $\beta$  (3 ng/ml, R&D Systems), 136 137 IFN-β (10 ng/ml, R&D), all-trans retinoic acid (ATRA, 10nM, Sigma), FICZ (AhR agonist, 100nM, Tocris Bioscience) were made at the start of the 7 day culture. Cultured cells were 138 washed in PBS, stained for viability and surface or intracellular markers as above. Tc1/Trm-139 like cells were analysed for cytokine production by re-stimulation with PMA 140 (20ng/ml)+ionomycin (400ng/ml)+monensin (3µM) for 4h before staining using Foxp3 141 142 staining buffer set.

#### 143 Commensal-specific T and B-cell memory proliferative responses

Commensal species were isolated from the cecum of healthy donors with the exception of *Collinsella aerofaciens,* which was from feces<sup>8,9,10</sup>. Strains were grown anaerobically in Hungate tubes containing Wilkins-Chalgren broth (37°C for 24h). Aliquots (1ml) were centrifuged (13,000rpm for 10 min), supernatants removed and cell pellets snap-frozen with dry ice before storage at -80°C. PBMC were obtained over Ficoll gradients and labelled with CellTrace Violet<sup>TM</sup> (1 $\mu$ M, Life Technologies) according to manufacturer's instructions, then cultured at 4x10<sup>6</sup>/ml in XVIVO15 serum-free medium (Lonza, +50 $\mu$ g/ml gentamycin (Sigma)

and penicillin/streptomycin (Life Technologies, 1/100)). 2x10<sup>5</sup> killed bacteria from 19 species
(as in Fig 5) were added to 0.2ml cultures and microbe-specific CD4<sup>+</sup>/CD8<sup>+</sup> T-cell and B-cell
responses were determined after 7 days culture. Cultured cells were analyzed by staining
with LIVE/DEAD stain, CD4/CD8/CD19/integrin-β7/CLA/CD39.

#### 155 Suppression assays

156 A fraction of healthy donor PBMC were cryopreserved before isolation of naïve CD8 T-cells and differentiation into Tc1- or Trm-like cells as described above. Cells were cultured at 157 0.5x10<sup>6</sup>/ml in 0.4ml cultures; Tc1 cells were generated with anti-CD3/28+IL-2 only and Trm-158 159 like cells with addition of TGF- $\beta$ , IFN- $\beta$ , ATRA and FICZ. After 7 days cells were washed and autologous PBMC thawed before labelling with CellTrace Violet<sup>™</sup>. Labelled target cells were 160 cultured in U-bottom wells (0.2ml XVIVO-15) at  $10^6$ /ml with or without addition of 161 unlabelled Tc1/Trm cells and CD39 inhibitor ARL67156 (200µM, Tocris Bioscience). Cells 162 were stimulated by addition of SEB (0.1µg/ml, Sigma) and stained after 4 days with 163 164 LIVE/DEAD stain, CD3/CD4/CD8/ CD25. Cells were gated for CellTrace Violet<sup>+</sup> CD8<sup>+</sup> T-cells 165 and fractions of cells which had divided and upregulated CD25 assessed.

#### 166 Induction of transcription factors and cytokines in DC

PBMC from healthy donors were cultured at  $10^6$ /ml in RPMI-1640 medium (Sigma) supplemented with 10% newborn calf serum (Sigma) and antibiotics as above. LPS (1µg/ml), SEB (10ng/ml, both Sigma), anti-IFN-γ (50µg/ml) or isotype control IgG1 were added. After overnight culture all cells were stimulated with LPS+poly I:C (1µg/ml, Sigma) + monensin (3µM) for a further 4h and stained for lineage markers, HLA-DR/CD123/CD11c/T-bet/TNF-α /IFN-α and LIVE/DEAD stain using Foxp3 buffer set. Gating for singlet mDC and pDC was

performed as shown in supplementary file 1. <u>Strict gating for CD11c-negative cells was used</u>
 to exclude mDC precursors from the pDC gate – this was confirmed by lack of staining for
 <u>CD33, CX3CR1 and Axl.</u>

#### 176 Measurement of antibody responses

177 IEM were labelled with SYBR Green DNA stain (Life Technologies, 1/100,000), anti-IgA-178 APC/anti-IgG-APC/Cy7 and analyzed by flow cytometry to determine proportion (%) of bacteria coated with antibodies in the gut. Circulating antibodies to commensal species 179 were determined by incubation of plasma (1/10 in 0.1% BSA, 0.5ml) with  $1x10^5$  bacteria 180 181 (30min), followed by centrifugation (12,000g, 10 min) and staining as for IEM (or isotype controls for each sample). Intact microbes were gated according to SYBR Green, and ratio of 182 geometric mean fluorescence intensity of staining for test sample vs isotype control was 183 used as measure of antibody titre. Plasma IgG antibodies to viruses were measured using 184 ELISA kits from Abcam according to instructions. 185

186 Statistical analysis

GraphPad Prism 7 software (GraphPad, San Diego, CA) was used to plot and analyze the data. Clinical data were analyzed by two-tailed unpaired t testsone-way ANOVA or where populations were skewed, two-tailed Mann-WhitneyKruskal-Wallis tests. For in vitro experiments, data were analyzed using two-tailed paired t tests or one-way ANOVA for multiple experimental conditions. P values less than 0.05 were considered significant and indicated by: \*:p<0.05; \*\*:p<0.01; \*\*\*:p<0.001.

195 Human colonic Trm are identified by CD103 and Runx3 and express Treg markers CD39 and

196 *CD73* 

To evaluate the role of Trm in IBD, we first identified Trm in intraepithelial lymphocytes (IEL) 197 198 and lamina propria lymphocytes (LPL) from healthy control (HC), CD and UC colonic biopsies 199 (non-inflamed tissue; clinical and demographic patient characteristics are shown in Table 1). In healthy IEL (Fig 1a) all T-cells including CD8<sup>+</sup> or  $\gamma\delta$  T-cells, expressed the CD69 putative 200 201 Trm marker. However, CD103 distinguished Trm from effector memory T-cells, which was 202 confirmed by their intranuclear expression of the Runx3 transcription factor, which controls 203 the Trm transcriptional program in mice<sup>11</sup>. A fraction of Trm expressed T-bet, which controls 204 the Th1 program of differentiation<sup>12</sup>. In LPL (Fig 1b), main populations were CD4<sup>+</sup> and CD8<sup>+</sup> T-cells and the latter not only contained a much larger proportion of CD103<sup>+</sup> Trm-like cells 205 206 but also had higher levels of Runx3 and T-bet than their CD4 counterparts. IEL lacked CD4 cells and no  $\gamma\delta$  T-cells were found in LPL. Trm and  $\gamma\delta$  T-cells expressed high levels of the 207 Treg markers CD39 and CD73, suggesting immunosuppressive function (Figs 1c/1d). CD8 208 Trm were nearly all conventional T-cells expressing CD8αβ heterodimer (Fig 1c) and no 209 Foxp3 transcription factor (Fig 1d). The vast majority of cells expressing CD39 and CD73 210 211 were also Foxp3<sup>-</sup> (barring a small fraction of the CD4 Trm) suggesting that the Trm 212 themselves contribute to maintaining tissue homeostasis.

#### 213 Trm are deficient in Crohn's disease and ulcerative colitis

Using this analysis we next compared numbers and phenotype of CD8<sup>+</sup> and CD4<sup>+</sup> Trm and  $\gamma\delta$ T-cells obtained from right colon of HC, CD and UC donors (Fig 2). We found a dramatic decrease in both CD8 Trm and  $\gamma\delta$  T-cells in IEL in IBD patients (Fig 2a; 84% and 61% for CD8

217 Trm; 90% and 87% for γδ T-cells in CD and UC, respectively). Both total numbers and 218 percentages of Trm relative to total live cells were reduced in IBD donors. The phenotype of 219 CD8 Trm was unchanged, but significantly decreased vo T-cell expression of CD39 and CD73 220 was seen in UC and CD, respectively, suggesting impaired regulatory function. In LPL, there 221 were also fewer Trm, although the deficiency was less dramatic than in IEL (61% and 44% for CD8; 28% and 68% for CD4 in CD and UC respectively) and did not reach statistical 222 223 significance for CD4 Trm in CD or when expressed in percentage terms (Fig 2b). There was no change in phenotype of LPL Trm in disease. Total yields of viable cells in IEL and LPL were 224 225 unchanged in IBD (Fig 2c); thus deficiencies in Trm were selective and could not be explained by loss of epithelium. We also found a strong correlation between numbers of 226 CD8 Trm in IEL and γδ T-cells (Fig 2d), suggesting co-dependence of these populations. In left 227 colon biopsies we found significantly fewer Trm (supplementary file 2, Fig S1) and no 228 229 significant changes in IBD.

Human Trm development in vitro is controlled by TGF-8, IFN-8, retinoic acid and AhR
receptor agonists

232 To determine possible mechanisms contributing to the deficiency of Trm in IBD we studied 233 micro-environmental factors. We developed an in vitro model to induce human Trm-like cells from naïve CD8 T-cells purified from healthy donor PBMC (Fig 3). Seven days 234 differentiation with anti-CD3/CD28 and IL-2 yielded effector cells with few markers of Trm 235 with the exception of the CD73 Treg-associated molecule, and were designated Tc1 type. 236 We tested addition of TGF- $\beta$ , a mucosal cytokine known to promote mouse Trm 237 development<sup>2</sup>; IFN- $\beta$ , since type 1 interferon in the gut can control colitis<sup>13</sup>; all-trans retinoic 238 239 acid (ATRA), known to induce CD10314; and FICZ (5,11-Dihydroindolo[3,2-b]carbazole-6-

240 carboxaldehyde), an aryl hydrocarbon receptor (AhR) agonist known to promote development of IEL<sup>15</sup>. Different Trm markers were induced differentially by each factor or 241 combinations thereof. CD103 expression was dependent on TGF-B alone; CD39 was induced 242 243 by IFN- $\beta$  and FICZ, as was the Trm transcription factor Runx3. Integrin- $\beta$ 7, not a Trm marker 244 but indicative of gut-homing potential, was induced by a combination of TGF- $\beta$  and ATRA, as 245 was CD69, a Trm marker expressed on all intestinal T-cells. Cells expressing all Trm-246 associated markers simultaneously (Fig 3b) were maximal using A a combination of all four 247 factors, which were-was-therefore used to induce Trm-like cells in further functional 248 experiments. IL-15, although involved in mouse Trm development, had no effect in this 249 model.

#### 250 B-cells are dysregulated in quiescent IBD patients

251 To determine if reduced CD8 T-cells in colonic tissue of IBD patients was 252 indicative of imbalance in cell-mediated immunity versus humoral immunity towards the 253 microbiota, we examined proportions of follicular helper (Tfh-like) cells, key inducers of 254 antibody production through interaction with B-cells in germinal centres, alongside the gut-255 homing function of T-cells in PBMC (Fig 4a and Fig S3). Tfh-like cells expressing CD4 and 256 CXCR5 were unchanged in IBD, as were proportions of integrin- $\beta$ 7<sup>+</sup>, gut-homing cells. CD8 Tcells did not express CXCR5 but showed high levels of integrin-\$7 indicating their gut-257 homing capacity was not impaired in IBD. Analysis of B-cell subsets in PBMC (Fig 4b, Table 2) 258 showed significantly increased proportions of plasmablasts (CD38<sup>hi</sup> CD27<sup>+</sup> B-cells) in both CD 259 and UC; these are a highly activated subset destined to become plasma cells in tissues<sup>16</sup>. 260 Other B-cell subsets, including those switched to IgA or IgG production were unchanged. 261 262 Consistent with increased B-cell activity, IEM released from colonic biopsies showed

significantly higher levels of IgA coating in both CD and UC than in HC (Figs 4c/4d), although
numbers of microbes obtained was unchanged.

T- and B-cell memory responses to commensal bacteria indicate skewing to humoral
 immunity in IBD

267 We then analyzed whether antigen-specific T- and B-cell responses to specific commensal 268 bacteria were imbalanced in IBD (Fig 5). We selected 19 commensal strains mainly isolated 269 from healthy human cecum, covering as many genera as possible. Killed bacteria were added to PBMC for 7 days to identify specific memory CD4/CD8 T-cell or B-cell proliferative 270 271 responses. Results showed responses were highly specific to individual species (Fig 5a/5b) and showed high degrees of variability both between individual donors and between HC and 272 IBD patients (Fig 5c). Variability within responses of individual donors was noted after 273 around one year (Fig S2), indicating such memory is dynamic and not long-term. As 274 275 expected CD4 T-cell responses were the predominant memory response in all groups; 276 however, total numbers of positive responses for each donor were unchanged in health vs 277 IBD (Fig 5d). By contrast, numbers of the less frequent CD8 T-cell responses were 278 significantly reduced in CD compared with HC, with the same trend apparent in UC (Fig 5d). 279 B-cell memory responses to bacteria were rare in health but significantly increased in both CD and UC (Fig 5d). T-cells proliferating in response to microbes expressed integrin- $\beta$ 7 (gut-280 homing marker), CLA (skin-homing) and the CD39 Treg marker, while B-cells only expressed 281 282 CLA in response to microbes (Fig 5a/Fig S3).

To investigate a possible link between circulating CD8 memory to commensals and recruitment of CD8 Trm to mucosa, we correlated numbers of CD8 responses to the 19

285 bacteria with CD8 Trm (IEL) numbers in donors where both blood and biopsies were 286 obtained; this indicated a significant positive correlation (Fig 5e). We then examined the magnitude of individual responses, as reflected by the proportion of divided cells, which is 287 related to antigen-specific precursor frequency. Results for the most immunogenic species 288 289 from each phylum (Fig 5f) show the high level of variability between donors, with significant differences in CD8 response in health vs IBD revealed for Staphylococcus epidermidis and 290 Escherichia coli only, the two most immunogenic species. B-cell responses however were 291 significantly increased for all species in CD and UC. We also categorized numbers of 292 293 proliferative responses against the four phyla of bacterial species (Fig S4). The same trends 294 were observed in all phyla, with the most significant differences in CD8 and B-cell responses seen in Actinobacteria and Firmicutes. We also performed assays for commensal-specific 295 antibodies in plasma, a more conventional readout for B-cell immunity. Circulating IgG 296 297 specific for the most immunogenic species in the 19-strain panel, as shown in Fig 6a, showed that antibody was increased in CD but not UC- indeed levels in UC were the same as 298 those in HC and significantly lower than in CD. Antibodies to less immunogenic species were 299 detectable but not significantly different between health and IBD. The divergent findings 300 301 with B-cell proliferative response vs circulating IgG were reflected in poor correlations 302 between the levels of each in individual donors; the only statistically significant correlation 303 was found with E. coli (Fig 6b). Circulating IgA levels were much lower than IgG (Fig S5) or 304 undetectable, and did not show significant differences between health and disease.

We tested whether immune deviation seen in IBD was specific to microbiota or reflected a systemic bias affecting responses to other antigens. We chose to assess responses to classic CD8 T-cell-inducing viral antigens. Plasma were assayed for IgG to three non-enteric viruses

encountered in childhood – varicella-zoster (VZV), measles and respiratory syncytial virus (RSV) (Fig 6c). Antibody to VZV showed the same pattern as commensals, with significantly increased levels in CD but not UC. Antibodies to measles were detectable in all patient groups but did not differ significantly, while few positive titres of RSV IgG were detected.

#### 312 Mechanisms of immune deviation in IBD

313 The above data clearly indicated a pattern of immune deviation between cellular/cytotoxic 314 and humoral immunity to members of the intestinal microbiota as an underlying feature of IBD. To examine potential pathogenic mechanisms, we first pursued the hypothesis that 315 Trm express regulatory T-cell function. We performed suppression assays using 316 conventional Tc1-type CD8 effector cells and Trm-like cells, generated using our in vitro 317 model system (Fig 3). The targets used in the assays were autologous PBMC stored in liquid 318 nitrogen. Effector cells were added at a 1:4 ratio and an inhibitor of CD39 ectonucleotidase 319 320 activity (ARL67156) used to determine whether suppressive activity was CD39-dependent. 321 Target CD8 T-cell proliferation assayed after 4 days revealed suppressive activity in Trm but 322 not Tc1 cells, which was partially reversed in the presence of the CD39 inhibitor (Fig 7a). The 323 cultured Tc1 and Trm cells were also tested for cytokine production (Fig 7b), which revealed 324 that Trm cells had similar capacity for production of pro-inflammatory cytokines IFN-y, TNF-325  $\alpha$  and IL-17 compared to Tc1; however IL-10, the key immunoregulatory cytokine in the GItract<sup>17</sup>, was significantly increased in Trm cells. Since dendritic cells (DCs) are critical for 326 327 controlling immune deviation and tolerogenic responses and are a target of Treg, we 328 analyzed the interaction between T-cells and DCs, again using a model system with healthy PBMC. To stimulate DC: T-cell cognate interactions we used SEB superantigen, compared to 329 330 TLR-mediated DC stimulation using LPS (Fig 7c). We also added LPS + poly I:C + monensin to

331	cultures for the final 4h in order to assess DC cytokine production. SEB but not LPS strongly
332	induced T-bet transcription factor expression in both myeloid (mDC) and plasmacytoid DC
333	(pDC), and those DC expressing T-bet produced less TNF- $\alpha$ and more IFN- $\alpha$ , cytokines with
334	opposing roles in colitis^{13,18-20}. In pDC, overall TNF- $\alpha$ production was suppressed by SEB,
335	whilst in mDC the reduced levels in T-bet^ DC were counterbalanced by increased TNF- $\alpha$ in
336	T-bet $ cells. The effect of T-cell:DC interaction on T-bet was partially dependent on IFN-\gamma, as$
337	shown by a neutralizing IFN- $\!\gamma$ antibody, but effects on cytokine production appeared IFN- $\!\gamma$ -
338	independent.

#### 341 Discussion

342 IBD is characterized by acute inflammatory episodes and pathology, and current treatments 343 aim to suppress symptoms using a plethora of immunosuppressive strategies. Our studies here, focussing mainly on patients with little or no active inflammation, reveal 344 345 that underlying disease is characterized by reduced CD8 T-cell immunity to commensal microbes associated with a paucity of Trm, potentially explaining the loss of 346 barrier immunity which characterizes IBD and drives pathology. Reduced CD8 347 348 response can also explain the skewing of immunity towards B-cell-mediated antibody production, and loss of immunoregulation in the local mucosa due to the reduced numbers 349 of cells expressing key Treg molecules CD39 and CD73. The mutually antagonistic 350 relationship between cell-mediated and humoral immunity was first noted in the 1970s<sup>21</sup> 351 and was subsequently attributed to the Th1/Th2 axis<sup>22,23</sup>, as was the hygiene hypothesis in 352 immune-mediated disease<sup>24</sup>. CD8 T-cell responses skew immunity away from humoral and 353 towards cellular immunity - our study is the first to examine such responses to the intestinal 354 microbiota in humans and points towards novel strategies in IBD treatment. Studies of anti-355 inflammatory commensal-induced pathways in the gut have focussed on CD4 Foxp3 Treg, 356 357 which form a small fraction of the LPL and are absent from IEL in human colon. By contrast, 358 our data show Trm could provide a gatekeeper function, controlling access of mucosal antigens to germinal centres in lymphoid tissue, and thus Tfh:B-cell interaction, whilst 359 simultaneously controlling inflammation through breakdown of extracellular ATP<sup>25,26</sup>. Runx3 360 361 has recently been defined as a master transcription factor for development of murine Trm<sup>11</sup>. 362 Our data show that human gut Trm preferentially express Runx3, and further co-express CD39 and CD73, key functional molecules on Treg cells<sup>27</sup>. CD39 is essential for in vitro 363

suppressive activity of Foxp3<sup>+</sup> Treg cells due to its ability to degrade extracellular ATP<sup>28</sup>, which activates DC<sup>29</sup>, and CD73 assists further nucleotide breakdown to adenosine, an immunosuppressive molecule<sup>30</sup>. ATP is released in mucosal tissue by injury but is also secreted by bacteria<sup>31</sup>, explaining the necessity for high expression of these molecules by Trm, especially IEL, in comparison to circulating T-cells. IEL and in vitro-derived Trm-like cells expressed lower levels of CD73, suggesting that further breakdown of ADP towards adenosine occurs further into the mucosa.

371 Foxp3<sup>+</sup> Treg are critical in systemic tolerance and in establishing tolerance to self-antigens in early life<sup>32</sup>. Foxp3 was not expressed in CD8 T-cells in the colon, which outnumber CD4 T-372 373 cells. Foxp3<sup>+</sup> Treg were present at a modest percentage in the CD4 LPL population (around 374 5%) and were vastly outnumbered by Foxp3-negative CD4 and CD8 T-cells, mostly Trm, expressing high levels of CD39 and CD73. Arguably the low number of Foxp3<sup>+</sup> Treg in human 375 376 colon is insufficient to maintain tolerance in the presence of such large antigenic loads from the microbiota, necessitating accumulation of Trm populations with regulatory capacity. 377 378 Since Trm do not differentiate until they reach the tissue<sup>33</sup>, this would explain why we found tolerance to commensal bacteria was not systemic, but localized to the gut. Circulating T-379 380 cells reactive to commensals would not express regulatory function until resident in the 381 tissue and would require local tissue factors such as type I IFN and AhR agonists to maintain their function. This picture contrasts with that emerging from mouse models, most likely 382 383 due to far greater antigenic experience and maturity of the adult human immune system 384 compared to laboratory mice. We found higher proportions of conventional CD8 $\alpha\beta$  T-cells in 385 tissue than reported in mice, which may rely more on innate mechanisms and thymus-386 derived Foxp3<sup>+</sup> Treg due to their short lifespan. We also found strong correlation between

 $\alpha\beta^+$  and  $\gamma\delta^+$  T-cells in IEL, implying  $\alpha\beta^+$  may support  $\gamma\delta^+$  cell populations in a fashion 387 analogous to that demonstrated in the thymus<sup>34</sup>, or co-dependence on tissue-specific 388 environmental factors. A further correlation was shown between numbers of memory CD8 389 responses to commensal bacteria and colonic CD8 Trm, suggesting that such responses are 390 391 required to recruit and maintain healthy Trm populations. CD8 Trm have recently been 392 shown to be recruited to skin in response to skin resident microbes in a non-classical MHCrestricted fashion<sup>35</sup>. This mouse study demonstrated that such Trm exhibited an unusual 393 phenotype with expression of immunoregulatory genes and wound-healing activity, thus 394 395 improving barrier function without inflammation. Our data suggest a similar phenomenon 396 occurs in the colon but is dependent on classical responses to a wide range of bacterial 397 antigens.

One recent study demonstrated a pro-inflammatory role for Trm cells in active IBD<sup>36</sup>, thus 398 suggesting that Trm can exhibit both pro- and anti-inflammatory activities dependent on the 399 context. CD4 and CD8 Trm were increased in the lamina propria of inflamed IBD tissue in 400 this study, and T-cell transfer colitis experiments in mice confirmed that T-cells adopt a Trm 401 402 phenotype soon after recruitment to lamina propria in active disease<sup>36</sup>. The pathologic role 403 of Trm was dependent on their pro-inflammatory cytokine production regulated by 404 Hobit/Blimp-1 transcription factors expressed in Trm. However, deletion of Hobit/Blimp-1 in mouse CD4 T-cells had no effect on their regulatory function or development into Trm<sup>36</sup>. A 405 further study compared proportions of CD103<sup>+</sup> cells within total gut T cell populations in 406 inflamed vs uninflamed biopsies from CD and UC patients . This study showed decreased 407 408 proportions of CD103<sup>+</sup> cells in inflamed tissue; however this could have been due to influx of

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409 <u>effector memory-type cells.</u> Further work in murine systems targeting regulatory function in

Trm and effects on disease susceptibility are therefore warranted. We propose that dual functionality of Trm cells in homeostatic versus inflammatory conditions would allow balanced immunity to occur across large areas of tissue exposed to high antigenic loads.

Murine studies have not clearly described the biology of CD4<sup>+</sup> Trm and unlike CD8 cells, 413 Runx3 expression in human CD103<sup>+</sup> CD4 cells in LPL was low, so we focussed on CD8<sup>+</sup> Trm 414 activities. Dietary factors retinoic acid and AhR agonists could play a role in expression of 415 the Trm phenotype within tissue or in Trm survival, in addition to the mucosal cytokine TGF-416  $\beta$  and type 1 interferon, both cytokines associated with suppression of colitis<sup>13,37-39</sup>. Gut type 417 1 interferon production could be influenced by the enteric virome, which is also altered in 418 419 IBD<sup>40</sup>. ATRA is derived from vitamin A by antigen-presenting cells<sup>37</sup> while AhR agonists are dietary factors contained in cruciferous vegetables and key to gut health<sup>15</sup>. Combining these 420 factors in vitro allowed us to develop the first in vitro differentiation model for human Trm, 421 but also revealed distinct regulation of individual Trm-associated markers. Interestingly 422 CD39 was co-regulated with Runx3 while CD103 and CD73 were independently regulated. 423

424 Circulating IgG antibody to certain commensals was increased in CD but not UC, despite B-425 cell dysfunction in UC with increased circulating plasmablasts and IgA secretion. Since inflammatory lesions penetrate deeper into intestinal tissue in CD than UC<sup>41</sup>, and there is 426 more involvement of mesenteric lymph nodes in CD than UC<sup>42,43</sup>, it is possible that longer-427 lived, higher affinity antibody responses are generated in CD as antigens could access 428 germinal centres in lymph nodes driving affinity maturation. Circulating B-cell proliferative 429 430 responses may reflect shorter term responses with more broadly reactive antibody synthesis focussed on mucosa. Indeed, repeat assays on individual healthy donors showed 431 432 memory responses to commensal bacteria could change within a year, and long-term

433 memory is not required for non-pathogenic organisms. The excessive IgA response to 434 mucosa-associated microbes was apparent in both CD and UC but was not accompanied by 435 increased circulating Tfh-like cells. Future work could examine microbe-specific Tfh cells but 436 these might be sequestered in lymphoid tissue.

Mechanisms through which CD8 T-cells might control immune deviation are not entirely 437 clear, but they are known to regulate CD4 T-cell development. Here CD4 responses were not 438 altered in IBD, but we did not examine their cytokine profiles, which are skewed towards a 439 440 Th17 profile in IBD<sup>4</sup>. Th17 development is strongly inhibited by IFN- $\gamma^{44}$ , the major product of CD8 T-cells. Our data show an additional mechanism could be via induction of T-bet in DCs, 441 either via interaction with tissue-resident T-cells reactive to microbial antigens, or in 442 443 draining lymphoid tissues. T-bet expression in DC is critical in preventing colitis in mice, since it represses production of TNF- $\alpha^{18,45}$ . Consistent with this concept, we found 444 interaction with T-cells suppressed TNF-a in pDC whilst enhancing IFN-a production in DC, 445 although these effects were less specific to T-cells and IFN-y. This novel pathway may 446 contribute to immune deviation and allow acquired immune memory to reinforce DC 447 activity in tissues. Dialogue between Trm and DC in tissue may inform appropriate type of 448 449 memory response as well as directing tissue migration of effector cells. Current dogma 450 states that DC direct T-cell responses after integrating signals from innate immunity and tissue damage. However, additional dialogue between tissue DC and Trm would allow for 451 452 more intelligent decision-making based on host immunological experiences, thus allowing 453 the gut immune system to learn which bacteria are pathogenic over time.

IBD is a clear example of a "western" disease associated with dysbiosis and disrupted
 immunoregulation<sup>46</sup>. Our data establish IBD as a disease of B-cell dysfunction and point

456 towards a lack of deficient CD8 T-cell priming to the microbiota as key to its etiology. IBD 457 often exhibits extra-intestinal manifestations<sup>47</sup>, and several other diseases are associated with intestinal dysbiosis. It is therefore possible that lack of reduced Trm-priming is a general 458 mechanism underlying the hygiene hypothesis in immune-mediated disease, and 459 460 associations of microbiota with tumour development and cancer therapy<sup>48,49</sup>. Memory CD8 461 T-cells migrate to multiple tissues and escape homeostatic control mechanisms that limit their numbers in the circulation<sup>50</sup>, so numbers of Trm can accumulate throughout life in 462 response to immunological experiences. Notably, CD8 Trm accumulate throughout 463 childhood in humans<sup>51</sup>, when IBD is often first diagnosed. Since CD8 responses are typically 464 465 utilized for dealing with highly pathogenic organisms, a lack of exposure to enteric pathogens in early life could result in weakened tissue immunity and thus an altered 466 microbiota. Evidence for this in IBD was provided by the increased antibody response to VZV 467 468 seen in CD. VZV is latent and requires constant immune surveillance by cytotoxic T-cells; thus increased antibody may reflect weaker cytotoxic control of virus, although this was not 469 the case for measles and RSV. 470

471 Manipulating immunity to intestinal microbiota through vaccination may address the 472 underlying disease process, unlike current immunosuppressive strategies. It may prove of 473 greater therapeutic benefit than changing the microbiota itself in a range of diseases associated with dysbiosis, since every patient will respond differently to any particular 474 microbe/cohort due to MHC differences. Mice which lack T-bet expression in their innate 475 immune system develop altered microbiota which is colitogenic<sup>18</sup>, indicating dysbiosis is 476 477 secondary to immune changes. Vaccination would need specifically to target CD8 T-cell 478 responses; inducing cytotoxic activity against target microbes may eliminate them from the

479	microbiota, thus preventing pathology. The concept that immunization to induce CD8 T-cell
480	responses can suppress inflammatory pathology may be counterintuitive, but proof of
481	principle for this was demonstrated in mouse models of airway disease <sup>52,53</sup> . Such
482	vaccination could provide long-lasting effects on the highly plastic DCs that direct immune
483	responses into pathways associated with health or disease.

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#### 627 Figure Legends

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Figure 1. Human colonic Trm are identified by CD103 and express Runx3, T-bet and 629 regulatory T-cell markers but not Foxp3. A: CD8 T-cell and  $\gamma\delta$  T-cell populations were 630 identified in IEL fractions and stained for CD69/CD103 surface Trm markers; gated CD103<sup>+</sup> 631 cells were stained for intranuclear Runx3 and T-bet. B: CD4 and CD8 T-cell populations were 632 identified in LPL fractions and stained for CD69/CD103; gated CD103<sup>+</sup> cells were stained for 633 634 Runx3 and T-bet. C: IEL CD8<sup>+</sup> and γδ T-cell populations were stained for CD39/CD73 Treg-635 associated ectonucleotidases and CD8 $\alpha\beta$  to distinguish conventional vs innate-type 636 lymphocytes. CD8aß and CD8aa subsets were separately gated and T-bet and Runx3 637 expression shown, including isotype control staining for transcription factors. D: LPL CD4+ and CD8<sup>+</sup> Trm-like populations were stained for surface CD39/CD73 and intranuclear Foxp3. 638 639 Right panels show isotype control for Foxp3 stain. Staining is from right colon biopsies of 640 healthy donors and is representative of at least 5 individual donors. Similar data were 641 obtained in left colon.

Figure 2. Quiescent IBD is associated with reduced numbers of Trm and  $\gamma\delta$  T-cells in colonic 642 643 tissue. A: Numbers and percentages of CD8<sup>+</sup> γδ TCR<sup>-</sup> CD103<sup>+</sup> Trm recovered from IEL (upper 644 graphs) and  $\gamma\delta$  CD103<sup>+</sup> T-cells in IEL (lower graphs), alongside CD39 and CD73 expression on 645 these populations. B: Numbers and percentages of CD4<sup>+</sup> CD103<sup>+</sup> Trm (upper graphs) and 646 CD8<sup>+</sup> CD103<sup>+</sup> Trm (lower graphs) recovered from LPL, alongside CD39 and CD73 expression. 647 Full gating strategies are shown in supplementary methods. C: Total live cell numbers in 648 IEL and LPL fractions (including epithelial cells). D: Correlation of CD8 Trm and  $\gamma\delta$ 649 T-cell numbers in IEL populations from right colon biopsies of all donors. HC: healthy

controls, n=25; CD: Crohn's disease, n=12; UC: ulcerative colitis, n=20. Median values ± 95%
 confidence intervals are shown; statistically significant differences between groups (Mann
 WhitneyKruskal-Wallis test) are indicated. Spearman correlation coefficient was calculated
 in D.

Figure 3. Human CD8 Trm development in vitro is regulated by cytokines, vitamins and 654 dietary factors. A: Effects of combinations of TGF- $\beta$ , IFN- $\beta$ , all-trans retinoic acid (ATRA) and 655 an AhR agonist (FICZ) on Trm, Treg and homing markers in CD8 effector cells derived from 656 657 CD8 naïve T-cells differentiated with anti-CD3/CD28+IL-2 for 7 days. Graphs show mean ± 658 SEM from 5 independent experiments; groups compared using 1-way ANOVA with Dunn's 659 test for multiple comparisons applied. B: Cells expressing all Trm-associated markers simultaneously were analyzed as in A. C: Staining profiles as in A, showing example of cells 660 cultured in anti-CD3/28 + IL-2 only (Tc1 cells) or with addition of TGF- $\beta$ , IFN- $\beta$ , ATRA and 661 FICZ (Trm-like cells). 662

663 Figure 4. Immunopathology of quiescent IBD reflects B-cell dysregulation. A: Proportions of 664 Tfh-like (CXCR5<sup>+</sup>) CD4/CD8 T-cells and gut homing (integrin- $\beta$ 7<sup>+</sup>) T-cells in PBMC of healthy control and IBD donors. B: Circulating plasmablasts (CD38<sup>hi</sup> CD27<sup>+</sup> CD19<sup>+</sup>) and B-cell subsets 665 666 in healthy and IBD donors. Mann WhitneyKruskal-Wallis tests were used to compare groups 667 (n=23 HC; n=18 CD; n=17 UC). C: IgA coating of IEM obtained from right colon biopsies of example HC, CD and UC donors, after gating on SYBR Green<sup>+</sup> events. D: Pooled data showing 668 proportions of IgA<sup>+</sup> IEM in donor groups. Unpaired t-tests were1-way ANOVA was used to 669 compare groups (n=25 HC; n=9 CD; n=19 UC). 670

Figure 5. T- and B-cell memory responses to commensal bacteria show skewing from cellmediated to humoral immunity in IBD. A: CD4 and CD8 T-cell memory responses to selected

673 commensals in healthy PBMC, showing examples of CellTrace Violet dilution in CD4/CD8-674 gated populations in cultures showing positive and negative responses alongside SEB positive control; integrin- $\beta$ 7 staining indicates gut-homing potential of expanded antigen-675 specific cells. B: CD19<sup>+</sup> B-cell responses to selected commensals in example CD and UC 676 677 patient PBMC; as in A but gated on CD19<sup>+</sup> events. C: Representative proliferation data in 678 PBMC from a HC, CD and UC donor, showing responses to a panel of 19 bacteria after 7 days stimulation and gating for CD4<sup>+</sup> CD8<sup>+</sup> and CD19<sup>+</sup> cells. D: Pooled data as in C, showing 679 680 numbers of positive responses within panel of 19 commensals. Mann-WhitneyKruskal-Wallis 681 tests were used to compare groups; n=18 HC, n=16 CD&UC. E: Correlation of CD8 proliferative responses in PBMC with CD8 Trm in IEL from autologous biopsies (n=15). Upper 682 panel shows pooled data with lower panels showing individual correlations for HC and IBD 683 684 samples. F: Magnitude of proliferative responses to 4 individual species, one from each 685 phylum. Pearson correlation coefficient was calculated in E.

686 Figure 6. Circulating specific IgG antibodies to immunogenic commensal species are raised in 687 CD but not UC donors. A: Plasma was assayed for IgG antibodies using a coating assay. Ratios of median fluorescence intensity of anti-IgG-stained vs isotype control for each 688 689 sample are shown (median and 95% CIs). Mann-WhitneyKruskal-Wallis tests were used to 690 compare groups; n=30 HC, n=18 CD&UC. B: Correlation of B-cell proliferative responses to E. 691 coli against circulating IgG in 40 matched donors, including Pearson correlation coefficient. Correlations with other species were not significant. C: Antibodies against non-enteric 692 viruses in health vs IBD. Plasma were assayed by ELISA for IgG to viral antigens and results 693 694 expressed in arbitrary units. Grey lines represent cut-off points below which titres are 695 considered negative. Unpaired t tests were 1-way ANOVA was used to compare groups.

696 Figure 7. Mechanisms of immune deviation in IBD. A: Trm-like cells have Treg function partially dependent on CD39 nucleotidase activity - Tc1 or Trm-like cells were added to 697 autologous fluorescent labelled PBMC and suppression of CD8 target cell activation was 698 699 determined in the presence or absence of the CD39 inhibitor ARL67156. Example staining 700 and pooled data showing % suppression of proliferation from 3 independent experiments mean±SEM, paired t-tests used to compare groups. B: Trm-like cells have increased capacity 701 for IL-10 secretion. Tc1 and Trm-like cells were restimulated with PMA/ionomycin/monensin 702 and stained for intracellular cytokines. Example staining and pooled data showing % staining 703 704 from 4 independent experiments - mean±SEM, paired t-tests used. C: Induction of T-bet expression in DC by cognate interaction with T-cells mediated by superantigen is associated 705 706 with altered cytokine synthesis - PBMC were cultured overnight with LPS or SEB plus control 707 antibody/anti-IFN-y, followed by 4h with LPS+poly I:C+monensin. mDC/pDC populations were gated according to CD11c/CD123 expression after gating on singlet, viable DC using 708 lineage vs HLA-DR plots. Example staining and pooled data from 4-5 independent 709 experiments is shown; paired t-tests were used to compare groups. 710

# Table 1. Clinical characteristics of St Mark's Hospital colonoscopy patientsdonating colonic biopsies

Characteristic	HC	CD	UC
n	23	11	18
Male/female	13/10	4/7	12/6
Median age (95% CI) at sampling	51.5 (41-57)	43 (28-57)	53.5 (47-60)
Median age (95% CI) at diagnosis		26 (18-47)	35 (29-42)
Inflammation scores CD:			
Normal		5	
Erythema		1	
Active ulceration		4	
Severe ulceration/chronic inflammatory change		1	
Ileocolonic disease		<u>3</u>	
Inflammation scores UC:			
Mayo 0			8
Mayo 1			3
Mayo 2			6
Mayo 3			1
IBD Medications at sampling:			
Aminosalicylates		4	15
Azathioprine/6-mercaptopurine		4	1
Buscopan		1	0
Adalimumab		1	0
None		4	3
Non-IBD medications at sampling:			
Metformin/gliclazide/statin		0	2
Ondansetron		1	0
Certirazine		1	0
None		9	16

Demographic and clinical data analyzed in Figs 2, 4d, S1

# Table 2. Clinical characteristics of St Mark's Hospital blood donors andhealthy volunteers

Characteristic	НС	CD	UC
	40	47	
n Na kula	18	1/	14
Madian and (05% CI) at someling	13/5	8/9	///
Median age (95% CI) at sampling	41.5 (30-54)	43.5 (33-01)	54.5 (35-03) 20 5 (25 47)
Median age (95% CI) at diagnosis		25 (20-44)	38.5 (25-47)
Symptoms at sampling:			
Diarrhoea/loose stools		1	2
Occasional loose motion/watery stool		2	1
Abdominal pain		2	1
Peri-anal pain/itch/disease		2	1
Lethargy		2	0
Proctitis		0	1
None		9	8
IBD Medications at samplina:			
Aminosalicylates		8	8
Azathioprine/6-mercaptopurine		8	2
Corticosteroids		1	0
Vedolizumab		1	0
Antibiotics		1	0
Methotrexate		1	0
None		1	6
Non-IBD medications at samplina:			
Vitamins D/D3/B12/multi		4	3
Metformin/gliclazide		0	2
Statins		1	2
Hydroxychloroguine		0	1
Proton pump inhibitor		0	1
Alendronate		1	0
Finasteride		1	0
Loperamide		1	1
None		12	9

Demographic and clinical data analyzed in Figs 4, 5, 6, S3, S4