

1 **An association between the gut microbiota and immune cell dynamics in humans.**

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21 **ABSTRACT**

22 The gut microbiota influences development and homeostasis of the mammalian immune system¹⁻³,
23 can alter immune cell compositions in mice⁴⁻⁷, and is associated with responses to immunotherapy
24 that rely on the activity of peripheral immune cells⁸⁻¹². Still, our understanding of how the microbiota
25 modulates immune cells dynamics remains limited, particularly in humans where a lack of deliberate
26 manipulations makes inference challenging. Here we study hundreds of hospitalized—and closely
27 monitored—patients receiving hematopoietic cell transplantation as they recover from chemotherapy
28 and stem cell engraftment. This aggressive treatment causes large shifts in both circulatory immune
29 cell and microbiota populations, allowing the relationships between the two to be studied
30 simultaneously. We analyzed daily changes in white blood cells from 2,235 patients, and 10,680
31 longitudinal microbiota samples to identify bacteria associated with those changes. Bayesian
32 inference and validation across patient cohorts revealed consistent associations between gut bacteria
33 and white blood cell dynamics in the context of immunomodulatory medications, clinical metadata
34 and homeostatic feedbacks. We contrasted the potency of fermentatively active, obligate anaerobic
35 bacteria with that of medications with known immunomodulatory mechanism to estimate the potential
36 of the microbiota to influence peripheral immune cell dynamics. Our analysis establishes and
37 quantifies the link between the gut microbiota and the human immune system, with implications for
38 microbiota-driven modulation of immunity.

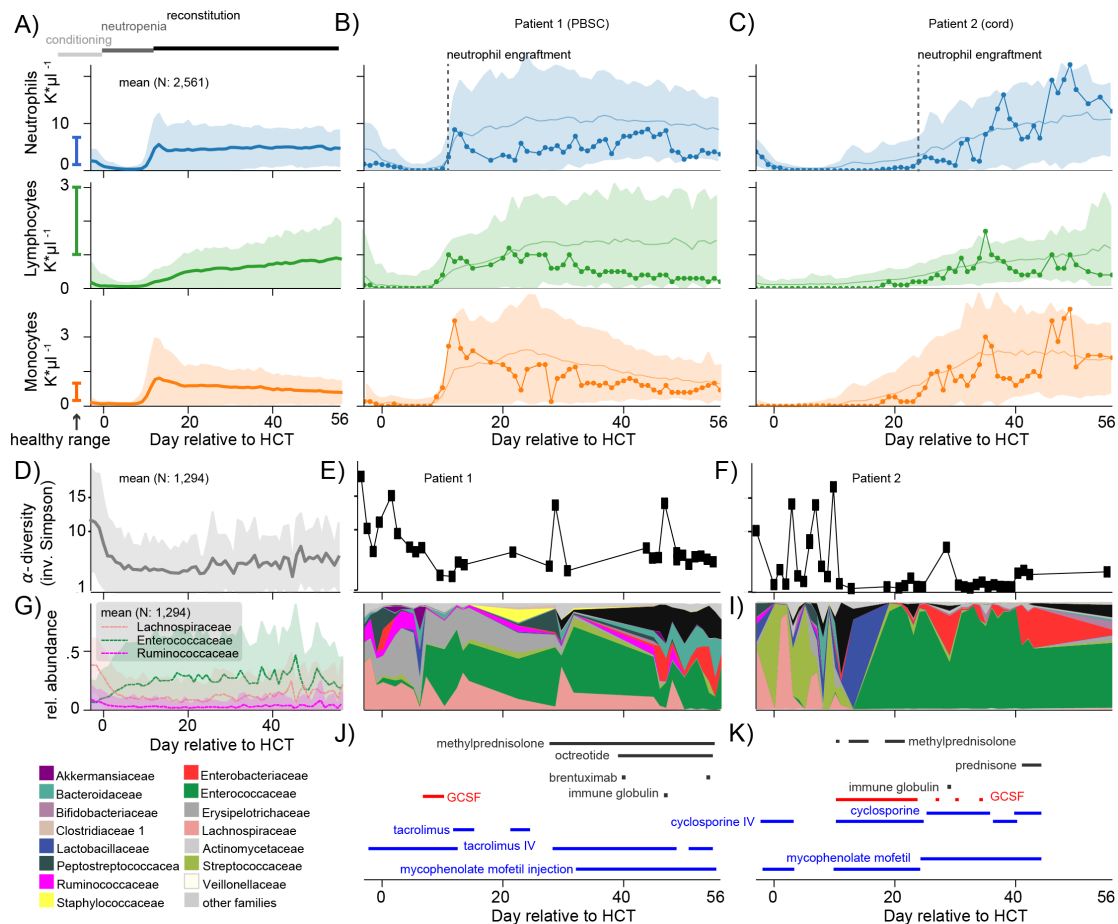
39 **MAIN TEXT**

40 Experiments in mice provide evidence that the mammalian intestinal microbiome influences the
41 development¹⁻³ and homeostasis of its host's immune system^{4-7,13-15}. In humans, inflammatory bowel
42 diseases correlate with functional dysbiosis in the gut microbiota^{16,17}. Children born preterm and at
43 term have different gut microbiome compositions and differ in the development of immune cell
44 populations in their blood¹⁸. The composition of the gut microbiota may also influence the success of
45 immunotherapies⁸⁻¹¹. Immune checkpoint inhibitor therapy relies on activation of circulating T-cells
46 and its success has, independently, been associated with abundances of intestinal anaerobic genera
47 such as *Akkermansia*⁹ and *Faecalibacterium*¹⁰. It is therefore an intriguing prospect to augment
48 treatments such as cancer immunotherapy¹⁹, including the burgeoning field of chimeric antigen
49 receptor (CAR) T-cell therapy²⁰, by leveraging microbiome-driven immune system modulation.

50 Our understanding of how the microbiota influences the dynamics of immune cells in
51 humans, and how this compares to deliberate immunomodulatory interventions nevertheless remains
52 limited. Experiments with animals may not always be sufficient to study mechanisms of
53 microbiome-immune interactions and translate them to human biology as the microbial ecology in the
54 gut of an animal model may be different from humans receiving treatment²¹. On the other hand,
55 studies directly in patients may be criticized when they have small subject numbers, are cross-
56 sectional, lack statistical power, or disregard key confounders such as medications²¹.

57 To overcome these limitations, we conducted a large-scale longitudinal study of the gut
58 microbiota and day-by-day changes in circulatory immune cell counts. We investigated immune
59 reconstitution dynamics after allogeneic hematopoietic cell therapy (HCT) within all 2,926 patients
60 who underwent HCT at Memorial Sloan Kettering for various hematological malignancies, including
61 leukemia, between 2003 and 2019 (**Figure 1A, Table S1**). The conditioning regimen of radiation and
62 chemotherapy administered to HCT patients is the most severe perturbation to the immune system
63 deliberately performed in humans and thus offers a unique opportunity to investigate dynamic links
64 between the gut microbiota and the immune system directly in humans. Conditioning depletes white
65 blood cell counts (**Figure 1A**) and can lead to prolonged periods of neutropenia (<500 neutrophils per
66 μ l blood). Immune reconstitution begins after transplanted stem cells have matured sufficiently to

67 release granulocytes from the bone marrow (neutrophil engraftment is defined as 3 consecutive days
 68 with >500 neutrophils per μl blood, **Figure 1A-C**). The blood of each patient is carefully monitored



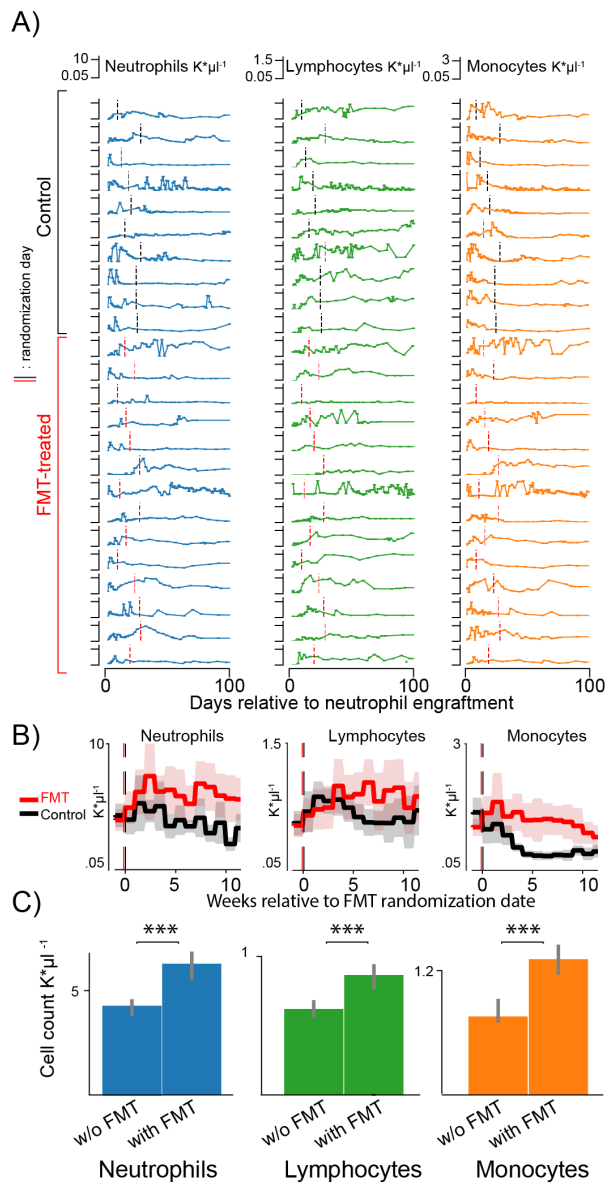
69 throughout this recovery, and medications are administered to modulate the immune cell dynamics,
 70 including granulocyte-colony stimulating factor (GCSF) to increase neutrophil counts, and
 71 immunosuppressants such as mycophenolate mofetil or tacrolimus to prevent complications such as
 72 graft-vs-host disease (**Figure 1 J,K**). To investigate if the composition of the gut microbiota is
 73 associated with the dynamics of circulating white blood cells, we analyzed detailed blood and clinical

74 metadata of our patients between 3 days before HCT and until 100 days post neutrophil engraftment
75 (excluding pediatric patients, and other exclusion criteria: N=2,235, supplementary methods, **Figure**
76 **S1**). During this period patients are monitored carefully, and our analysis included over 140,000 host
77 phenotype measurements in the form of complete blood counts which quantify the most abundant
78 white blood cells—neutrophils, lymphocytes, monocytes, eosinophils—as well as platelet counts
79 (**Figure 1, S1**). We started collecting patients' fecal microbiota data in 2009²², and by now obtained
80 10,680 high frequency, longitudinal microbiota compositions.

81 HCT patients lose gut microbiota biodiversity and commensal microbial families during their
82 treatment (**Figure 1D-I**); this figure generated from N=1,294 HCT patients clarifies the preliminary
83 trends observed in previous studies from smaller datasets^{23,24}. We have shown recently that mice with
84 a depleted intestinal flora had worse recoveries of white blood cells after bone-marrow
85 transplantation²⁵. In our patients, microbial diversity usually recovers slowly during white blood cell
86 reconstitution (**Figure 1D**); however, microbiota recovery as well as immune reconstitution can vary
87 strongly between patients and treatment types (**Figure 1B,C,J,K, S1**). This variation is illustrated by
88 the distinct trajectories of patient 1 who received a graft of peripheral blood stem cells (PBSC),
89 retained high microbiota diversity and engrafted earlier (**Figure 1B,E,H**), and patient 2 who received
90 a graft of umbilical cord blood (cord), lost microbiota diversity and engrafted later (**Figure 1C,F,I**).
91 Low microbiota diversity at the time of neutrophil engraftment has been associated with 5-fold
92 increased transplant-related mortality²⁶, suggesting that that the joint recovery of the microbiota and
93 white blood cells in circulation is critical for clinical outcomes.

94 To detect a directional and causal link between the microbiota and white blood cells, we first
95 used data from a recent prospective randomized trial of autologous fecal microbiota transplantation
96 (auto-FMT), which is a microbiota manipulation experiment done directly in our patients²³
97 (supplementary methods). Twenty-four patients (**Figure 2A, Table S2**) underwent randomization,
98 resulting in 10 untreated control and 14 treated patients, including patient 3 in **Figure S2**. To
99 investigate if auto-FMT affected white blood cell reconstitution, we compared the 24 patients'
100 neutrophil, lymphocyte, monocyte (**Figure 2**) and total white blood cell counts (**Figure S3**) post-
101 engraftment (i.e. when the transplanted hematopoietic cells begin producing new white blood cells,

102 **Figure 2A,S3).** FMT procedures were conducted at variable time points relative to neutrophil
 103 engraftment, but overall, we observed higher counts of each white blood cell type in patients who
 104 received an auto-FMT during the first 100 days post neutrophil engraftment ($p < 0.001$, **Figure 2B,S3-**
 105 **S6).**
 106



107 **Figure 2: Neutrophil, lymphocyte and monocyte counts increased in FMT-treated patients**
 108 **during the weeks following the treatment compared to control patients.** A) Absolute counts of
 109 neutrophils (blue), lymphocytes (green) and monocytes (orange) in 24 patients enrolled in a randomized
 110 controlled trial to receive an autologous fecal microbiota transplant post-neutrophil engraftment (10 control:
 111 black vertical line, 14 FMT treated: red vertical line,). B) Weekly mean cell counts aligned to the date of
 112 randomization into FMT treatment arm (red) or control (black). Line: mean per week, shaded region: 95%-CI.
 113 C) Results from a linear mixed effects model with random effects per patient and per day relative to neutrophil
 114 engraftment confirms that neutrophil, lymphocyte and monocyte counts are higher in patients receiving auto-
 115 FMT after their treatment as compared to control patients after the randomization date; bars and confidence
 116 intervals for the averages of observed white blood cell counts without FMT and post-FMT (***: $p < 0.001$).
 117

118 The increased white blood cells in patients receiving auto-FMT could be due to the
119 reconstitution of a complex microbiota that we saw in these patients²³ and the associated metabolic
120 capabilities^{6,7,25}, or it could be a systemic response to a severe therapy which introduced billions of
121 intestinal organisms at once via an enema (no enema was administered to control patients²³).
122 Moreover, while the mixed-effects model accounted for patient-specific HCT treatments in this
123 randomized patient cohort, chance differences in extrinsic factors such as different
124 immunomodulatory drug exposures may have affected this result due to the small cohort size.
125 Nonetheless, observing that auto-FMT recipients had increased white blood cell counts supports the
126 notion that the microbiota can modulate the peripheral immune system. High counts of lymphocytes
127 during immune reconstitution has been associated with improved clinical outcomes²⁷. Additionally, in
128 our HCT patients, a higher average level of white blood cells measured across a period of 100 days
129 after neutrophil engraftment (supplementary methods) confirms a positive association with 3-year
130 survival (hazard ratio: 0.91, p:0.04). Determining which taxa modulate immune dynamics could open
131 new ways to improve robust immune reconstitution, which is critical for clinical outcomes²⁷⁻²⁹.

132 To address this question, we next investigated the link between the gut microbiota and the
133 dynamics of white blood cell recovery in our large observational cohort of HCT patients. Homeostasis
134 of circulatory white blood cell counts is a complex, dynamic process: neutrophils, lymphocytes and
135 monocytes are formed and released into the blood *de novo* by differentiation of hematopoietic
136 progenitor cells from the bone marrow, and they can be mobilized from thymus and lymph nodes
137 (lymphocytes), spleen, liver and lungs (neutrophils); they can also migrate from the blood to other
138 tissues when needed³⁰. These processes are dynamic sources and sinks of circulatory white blood
139 cells, and they can be modulated by drugs administered to patients receiving HCT. To identify factors
140 associated with these dynamic source- and sink-processes—including the microbiota—we developed
141 a two-stage approach analyzing the changes of white blood cell counts between two days (i.e. the
142 rates of cell count increases and decreases). Stage 1 served as a feature selection stage where we used
143 data of 1,096 patients (after filtering for qualifying samples and applying exclusion criteria, see
144 supplementary methods) *without* available microbiome information to identify associations between
145 clinical metadata, including immunomodulatory medications (methods), and changes of white blood

146 cell counts from one day to the next (**Figure 3A**). Stage 2 was performed on data from an independent
147 cohort of 841 different patients at MSK from whom concurrent microbiome samples were available.
148 Stage 2, our main analysis, sought to reveal associations between the abundance of microbial taxa and
149 the daily changes in blood cell counts in context of immunomodulatory medications, additional
150 clinical metadata and the current state of the blood itself.

151 In stage 1 we calculated the changes in neutrophil, lymphocyte and monocyte counts during
152 patients' recovery from >20,000 pairs of post-engraftment blood samples separated by a single day
153 (**Figure 3B, S7-9**, supplementary methods). Using a cross-validated feature selection approach, we
154 detected medications and HCT treatment parameters that were associated with different rates of
155 change in neutrophil, lymphocyte and monocyte counts, including, as expected, GCSF and the graft
156 stem cell sources (**Figure S7-S9, Table S1**).

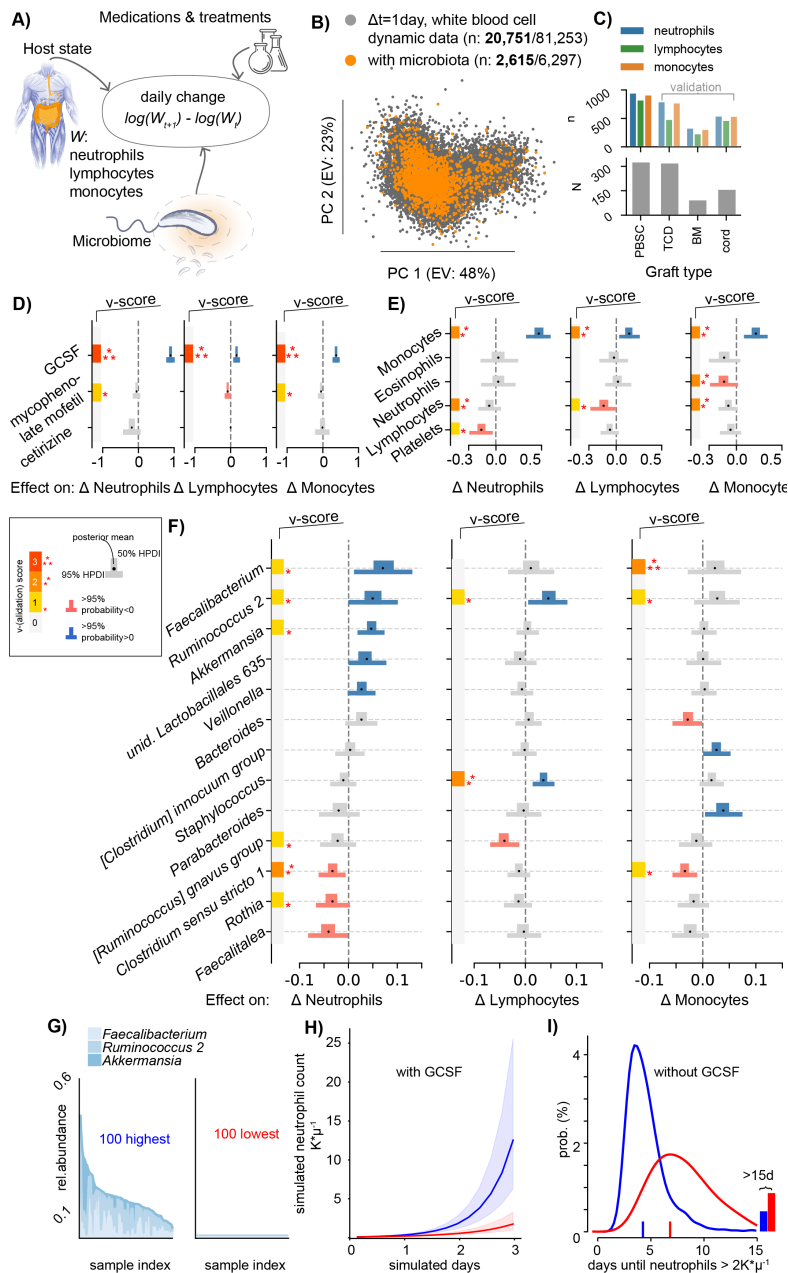
157 During stage 2 we sought to identify associations between bacterial taxa of the gut microbiota
158 and the dynamics of immune cells in circulation. For this, we performed Bayesian inferences using
159 data from different sets of patients *with* available microbiome samples. Stage 1 had identified—as
160 expected—that stem cell graft sources are associated with immune reconstitution kinetics (e.g. cord
161 on average slower compared with PBSC³¹), and we therefore stratified our patients by graft source in
162 stage 2. The model of stage 2 now included microbial genera as predictors of observed changes in
163 white blood cell counts, in addition to the medications selected in stage 1, clinical features
164 (conditioning intensity, age, sex), and the current state of the blood in the form of counts of
165 neutrophils, lymphocytes, monocytes, eosinophils, and platelets. The data comprised 841 patients, but
166 approximately 60% of the stool samples paired with a daily change in white blood cell counts were
167 taken before neutrophil engraftment (**Figure 3B, Table S1**, supplementary methods), i.e. when blood
168 cells counts were zero. In total, we analyzed 2,615 post-engraftment observations of changes in
169 neutrophil counts during immune reconstitution (lymphocytes: 2,006, monocytes: 2,534) with paired
170 stool samples which provided a large sample of observed white blood cell dynamics (**Figure 3B,**
171 **Table S3,S4**). We first focused on the data from the largest (**Figure 3C**) cohort—patients who
172 received a PBSC graft—and withheld the other cohorts (bone marrow, BM; T-cell depleted graft
173 (*ex-vivo*) by CD34+selection, TCD; and cord) to use as independent validation cohorts. For this

174 validation, we analyzed TCD, BM, and cord patients' data in the same way as PBSC patients' data
175 and compared the resulting posterior coefficient distributions (methods). We assigned coefficients
176 obtained from the PBSC cohort a validation score (v-score) between 0 and 3, representing the number
177 of times that the focal coefficient was validated in the other cohorts; but, conservatively, the score was
178 always set to zero if we observed counter-evidence among any of the other data sets, i.e. evidence that
179 coefficients had the opposite sign, ensuring only the most consistent associations were considered as
180 validated. Finally, we analyzed data from another patient cohort consisting of 493 bone marrow
181 transplantation recipients treated at Duke University including 9,603 blood samples and a total of 629
182 microbiota samples from 218 patients, albeit with lower sampling density, and we used the results
183 from this analysis for further validation.

184 Notably, as a verification of our approach, we detected associations between the
185 administration of immunomodulators and increased or decreased rates of immune cell count changes
186 consistent with the known biological mechanism of these medications (**Figure 3C,S10-S13**). The
187 strongest across all predictors is the well-known neutrophil-increasing effect of GCSF³²; GCSF
188 administration—used to accelerate recovery from chemotherapy-induced neutropenia³²—was
189 associated with a +140% increase in the rate of neutrophil changes from one day to the next ([+114%,
190 +170%], 95 percent probability density interval [HPDI95]). This finding was observed in all MSK
191 validation data sets (v-score=3, **Figure 3D**), as well as among Duke University patients (**Figure**
192 **S14,S15**). We furthermore found a GCSF-associated increase of +43% ([+30%, +58%]HPDI95, v-
193 score=3) in monocyte rates, and, although smaller, in lymphocyte rates (+16%, [+5%, +27%]HPDI95,
194 v-score=3). Both neutrophil and lymphocyte rates decreased following the exposure to antihistamine
195 or immunosuppressive medications (cetirizine -18%, [-35%, +5%]HPDI95, mycophenolate
196 mofetil -8% [-15%,+1%]HPDI95, respectively). Finally, less intensive chemotherapeutic conditioning
197 regimens (non-ablative conditioning and reduced intensity) were associated with larger lymphocyte
198 and monocyte count growth rates during immune reconstitution (**Figure S10C**)

199 Beyond associating medications in agreement with their known biological mechanism, our
200 analysis detected associations between the current count of white blood cells and their rate of change:
201 a negative association among lymphocytes, negative associations between counts of neutrophils and

202 lymphocytes with the rates of monocytes, and a negative association between the counts of platelets
 203 and lymphocytes and the rates of neutrophils (**Figure 3E**). Conversely, we found positive associations
 204 between monocytes and the rates of each of the investigated white blood cell subsets. These
 205 associations, derived from daily counts of white blood cells, could reflect a complex network
 206 underlying the regulation of blood immune cell composition³⁰. More importantly, the associations
 207 quantified for medications and potential homeostatic feedbacks provided a benchmark against which
 208 we could compare associations from gut microbial taxa.



209

210 **Figure 3: Bayesian inference conducted in context of the host state (white blood cell and platelet counts)**
 211 **and clinical variables including immunomodulatory medications reveals the microbiota potential to affect**

212 **daily changes in circulatory white blood cell counts.** A) Cartoon of the model: observed changes in white
213 blood cell counts between two consecutive days are associated with the current state of the host in the form of
214 blood cell counts in circulation, the administration of immunomodulatory medications, patient and clinical
215 metadata, and the state of the microbiome. B) Visualization of the dynamic white blood cell data; scatter plot of
216 the principal components (PC) of observed daily changes of neutrophils, lymphocytes and monocytes without
217 (grey, see **Figures S7-9**) and with (orange) concurrent longitudinal microbiome data (bold: post engraftment
218 sample counts). C) PBSC patients (N=312) provided the most blood samples with simultaneous microbiome
219 data (n=995) relative to TCD, BM and cord patients, who were used as validation data sets. D-F) PBSC patient
220 data inference results; bars show the posterior parameter estimate distributions (thin: 95% highest posterior
221 density intervals, HPDI95, thick: HPDI50) for the jointly inferred associations between treatments (D), white
222 blood cells counts (E), and fecal microbiota genus log-relative abundances (F) with the observed daily changes
223 in neutrophils, lymphocytes and monocytes. v-score: number of validation cohort confirming associations,
224 *always* set to zero if invalidated in any of the TCD, BM, or cord cohorts (additional coefficients in **Figure S10**).
225 G) 100 microbiota samples with highest (left) or lowest (right) relative abundances of *Faecalibacterium*,
226 *Ruminococcus 2* and *Akkermansia*. H) Simulation of the neutrophil population over time in presence of GCSF
227 with microbiota compositions sampled either from those high (blue) or low (red) in *Faecalibacterium*,
228 *Ruminococcus 2* and *Akkermansia* relative abundance as shown in G); line: median of 1,000 simulations, shaded
229 regions: quartile range of simulated neutrophil trajectories. I) In absence of GCSF, equivalent simulations to H)
230 predict that the time to reach neutrophil counts $>2K*\mu l^{-1}$ for the first time after HCT when the microbiota is
231 high (red) in *Faecalibacterium*, *Ruminococcus 2* and *Akkermansia* compared with when these genera are low
232 (blue) will decrease from 6.8 (95%-confidence interval, CI: [6.5, 7]) days to 4.4, (CI: [4.3, 4.5]).
233

234 We identified microbial genera that consistently associated with increases or decreases in
235 white blood cell counts by first using data from the PBSC patients and then validating the associations
236 in the other cohorts (**Figure 3F**). Higher abundances of *Faecalibacterium* (+8%, [+1%,
237 +14%]HPDI95 per log₁₀), *Ruminococcus 2* (+5%, [0%, +10%]HPDI95) and *Akkermansia* (+4%,
238 [+1%, +7%]HPDI95) were associated with greater neutrophil increases, whereas increased *Rothia* (-
239 3%, [-7%, 0%]HPDI95), and *Clostridium sensu stricto 1* (-3%, [-6%, 0%]HPDI95) relative
240 abundances associated with reduced neutrophil rates. These results were validated in univariate
241 analyses conducted in the Duke University cohort (**Figure S14, S15**). We also conducted the
242 inference using total genus abundances as predictors instead of relative abundances; this analysis
243 confirmed *Faecalibacterium* as most strongly associated with neutrophil dynamics (**Figure S16**,
244 supplementary methods). *Staphylococcus* was positively associated with lymphocyte rates (+4%,
245 [+1%, +6%]HPDI95) and, again, *Ruminococcus 2* was also associated with faster lymphocyte
246 increases (+5%, [+1%, +9%]HPDI95). Both *Faecalibacterium* as well as *Ruminococcus 2* also
247 associated with increases in monocytes, and while this association was validated in other cohorts (v-
248 score 3 and 1, respectively), there was higher uncertainty of the association estimate (HPDI50>0).
249 Again, *Clostridium sensu stricto 1* (-3% [-5%, -1%]HPDI95) associated consistently with decreased
250 rates of monocytes. The associations we identified—and validated in other cohorts—between

251 microbial taxa in the gut and daily changes in white blood cell counts support the idea that
252 hematopoiesis and mobilization respond to the composition of the gut microbiome, influencing
253 systemic immunity³³.

254 Most of the taxa that strongly associated with white blood cell dynamics were obligate
255 anaerobes. *Rothia*, was a notable exception: this aerobic genus is typically found in the oral cavity³⁴
256 but can become an opportunistic pathogen in immunosuppressed patients and is not known to provide
257 metabolic functions to the host³⁵. Some obligate anaerobes, on the other hand, produce short-chain
258 fatty acids^{36,37} and bacterial cell-wall molecules^{1,38,39} that modulate immune responses and
259 granulopoiesis⁶. Nutritional support from the intestinal microbiota improved hematopoietic
260 reconstitution in a mouse model²⁵. To identify a similar association in our patients, we estimated a
261 microbiota potency by multiplying the log₁₀-relative abundances of microbial genera in a sample with
262 their corresponding posterior coefficients. We analyzed shotgun metagenomics sequences from 124 of
263 the samples and observed that samples with positive microbiota potency were associated with
264 enrichment in cholate degradation and vitamin-B1 synthesis related pathways, as well as butanoate
265 formation (**Figure S17**). Our findings are in line with evolutionary theory⁴⁰ that essential but broadly
266 available microbial traits such as the production of B vitamins, secondary bile acid metabolism, and
267 fermentation to short-chain fatty acids⁴¹ could be co-opted by the host's immune system as part of the
268 homeostatic interplay between immune system and a complex microbiota^{42,43}. For example,
269 *Ruminococcus 2* is a genus that contains *R. bromii*, a keystone species necessary for the efficient
270 release of energy from complex starch in the normal diet⁴⁴. Reassuringly, the genera
271 *Faecalibacterium*, *Ruminococcus 2* and *Akkermansia* that we associated with faster rates of white
272 blood cells (**Figure 3F**) were among those best reconstituted by auto-FMT²³, potentially explaining
273 why we found higher counts of neutrophils, monocytes and lymphocytes in patients who received the
274 auto-FMT (**Figure 2B,C**).

275 The associations we reveal are interpretable as potential effectors on sources and sinks of
276 white blood cell counts in circulation. Intestinal bacteria may affect white blood cell counts in
277 circulation by influencing either their sources in the bone marrow or their cytokine profiles⁴⁵ and
278 proliferation rates in the blood, their sinks in different organs, or both. The human immune system in

279 turn can interact with the microbiota and modulate its composition, for example via
280 immunoglobulin A responses targeting specific bacteria as studied in mice^{43,46,47}. To investigate a
281 reverse effect of the peripheral immune cell system onto bacterial populations, we employed an
282 analogous approach to the stage 1 analysis of white blood cell dynamics. Dynamics of white blood
283 cells can be estimated from changes in absolute cell counts, and to obtain the necessary measurements
284 in absolute bacterial abundances, we measured total bacterial 16S rRNA gene copies per gram of stool
285 for a subset of our samples (3,995 samples from 481 patients). Using absolute abundances of bacteria
286 as predictors in addition to medications, we jointly inferred the association network of dynamics
287 between the gut bacterial ecosystem and the peripheral immune system. All of our patients receive
288 antibiotics on some days during their treatment²⁴ and their strong effects on microbiota dynamics
289 were the dominant effects that survived cross-validated regularized elastic net regression (**Figure**
290 **S18**). Relaxing the strength of the regularization (methods), however, revealed several bi-directional
291 relationships between immune cells in circulation and bacterial dynamics in the gut (**Figure S19**). Of
292 note, we detected a negative association of absolute [*Ruminococcus*] *gnavus* group abundance with
293 lymphocytes dynamics, confirming our main result based on relative bacterial abundances (**Figure 3**).
294 In the reverse direction, we saw a positive association of [*Ruminococcus*] *gnavus* group dynamics
295 with lymphocyte counts. This result agrees with findings that *Ruminococcus gnavus* thrives in and
296 promotes inflammatory conditions such as Crohn's disease and other inflammatory bowel diseases
297 (IBD)⁴⁸; our analysis suggests it may drive high neutrophil to lymphocyte ratios that are broadly
298 characteristic for poor disease outcomes in IBD^{49,50} and beyond^{51,52}.

299 Overall, our analysis identified that the microbiome is associated with immune cell dynamics
300 in addition to medications. The effects should be interpreted as net effects since they do not
301 distinguish, for example, how the microbiota impacts *de novo* hematopoiesis in isolation from its
302 impact on other sources and sinks. Unlike the plausible role of obligate anaerobe fermenters in
303 augmenting hematopoiesis via nutritional support²⁵, the positive association detected between
304 *Staphylococcus* and lymphocyte dynamics could instead result from reduced extravasation of T cells
305 from circulation into the gut epithelium⁵³, especially since high abundances of *Staphylococcus* are

306 associated with low gut microbiota diversity ($p < 0.001$, **Figure S20**), which indicates a depleted
307 microbiota.

308 Nevertheless, our approach allows us to leverage the chronology of events and assess
309 “mathematical causality”⁵⁴. Of course, due to the observational nature of these data there are risks of
310 confounding that could explain some of the associations found, but the close temporal
311 correspondence⁵⁴ between microbiota and blood cell dynamics, and the validation across cohorts
312 reduces the number of plausible confounders. Our results, therefore, quite naturally suggest candidate
313 microbial taxa to manipulate if we seek to steer complex hematopoietic dynamics and utilize the
314 microbiota as an immunomodulatory component of the human body. Intriguingly, members of
315 *Faecalibacterium* and *Ruminococcus* in one study¹⁰, and *Akkermansia* in another⁹, were identified as
316 enriched in patients with better responses to anti-PD-1 immunotherapy, which suggested a
317 disagreement between the two studies⁵⁵. Our results, however, revealed *Faecalibacterium*,
318 *Ruminococcus 2*, and *Akkermansia* as the most strongly associated taxa with increases in white blood
319 cell counts from one day to the next. Therefore, our results agree with the findings of both anti PD-1
320 therapy studies that these taxa are associated with immune modulation in humans. Our results also
321 allow us to compare the potency of manipulating these intestinal commensals to that of
322 immunomodulatory drugs. While these genera are common in the gut microbiota of healthy people¹⁷,
323 the relative abundance of each genus can drop below detection in our patients during the intestinal
324 damage related to HCT²⁴. Therefore, realistic ranges of 3-5 orders of magnitude in bacterial log-
325 relative abundances (**Figure 3G, S21**) can yield effect sizes similar to that of homeostatic feedbacks
326 between white blood cells and several immunomodulatory medications (e.g. a change in
327 *Ruminococcus 2* from below detection to 1% relative abundance associated with a +67% and +63%
328 increases in neutrophil and lymphocyte rates, respectively). Therefore, while the effect sizes of
329 intestinal bacteria at first may appear smaller than those of immunomodulatory drugs, the herein
330 estimated homeostatic effects of gut bacteria may not be that small since their coefficients refer to
331 changes in exponential rates of white blood cells and accumulate each day. To better demonstrate how
332 this accumulation of effects would work, we conducted simulations of the inferred dynamic system of
333 white blood cells using our posterior coefficient distributions (methods). We simulated 1,000 time

334 series for microbiota compositions either chosen from the 100 samples highest or lowest in
335 *Faecalibacterium*, *Ruminococcus 2* and *Akkermansia* (**Figure 3G**), in presence (**Figure 3H**) or
336 absence (**Figure 3I**) of GCSF administration. Simulations predict that a microbiota enriched in these
337 genera could accelerate immune reconstitution, and reduce the time until neutrophils reach $>2K*\mu l^{-1}$
338 in absence of GCSF by 2.4 days, from predicted 6.8 (CI: [6.5, 7]) to 4.4 days (CI: [4.3,4.5]) days. Gut
339 bacteria, in concert and over time, could therefore have significant impact on systemic immunity even
340 in individuals with less severely injured microbiomes.

341 In sum, our work links the gut microbiota to the dynamics of the human immune system via
342 peripheral white blood cell populations. Our analysis uses white blood cells counted directly from
343 patients, which are coarse-grained clinical analyses conducted at large scale but lack details such as
344 lymphocyte and other immune cell subsets. Nonetheless, because it is in humans, this study fills an
345 important gap at a critical time for microbiome research when the clinical relevance of animal models
346 of microbiome-immune interaction has been questioned²¹. By studying a large number of patients
347 over time, we were able to infer and quantify for the first time the association of microbiota
348 components on systemic immune cell dynamics, and our results help to consolidate previous
349 findings^{10,9} that seemed in conflict with each other⁵⁵. Our study demonstrates that the composition of
350 the microbiota does indeed modulate systemic immune cell dynamics, a link that could be used in the
351 future to improve immunotherapy and help identify microbiota treatments for inflammatory
352 diseases^{9,10,56-60}.

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516 convergence of MCMC. *arXiv* (2019).
- 517
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519 **Supplementary Information**

520 Appended and available online.

521

522 **Acknowledgments and Conflicts of Interest**

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540

541 **Author Contributions**

542 J.S. and J.B.X. wrote the manuscript. J.S. and J.B.X. designed the analyses with expert help from
543 R.N., J.U.P. and Y.T. contributed to the clinical data preparation, B.P.T. provided the 16S data
544 processing pipelines, A.D. provided the shotgun processing pipelines. All authors contributed to the
545 writing and interpretation of the results.

546 **SUPPLEMENTARY INFORMATION**

547 **Methods**

548 **Complete blood count collection and characterization**

549 Absolute white blood cells count data were obtained from routine complete blood counts ordered by
550 clinicians during normal clinical practice, used to obtain informative diagnostic and monitoring
551 information. Blood samples received in the clinical hematology laboratory were analyzed using
552 Sysmex XN automated hematology analyzers (Sysmex, Lincolnshire, IL) and, when needed based on
553 specific flags and parameters as per MSKCC standard operating procedures, were validated manually
554 using the Sysmex DI-60 Slide Processing System or CellaVision DM9600 Automated Digital
555 Morphology System (Sysmex, Lincolnshire, IL).

556

557 **16S rRNA gene amplification and multiparallel sequencing**

558 For each sample, duplicate 50- μ l PCRs were performed, each containing 50 ng of purified DNA, 0.2
559 mM deoxynucleotide triphosphates, 1.5 mM MgCl₂, 2.5 U Platinum Taq DNA polymerase, 2.5 μ l of
560 10 \times PCR buffer, and 0.5 μ M of each primer designed to amplify the V4-V5: 563F (5'-nnnnnnnnn-
561 NNNNNNNNNNNN-AYTGGGYDTAAAGNG-3') and 926R (5'-nnnnnnnnn-NNNNNNNNNNNNN-
562 CCGTCAATTYHTTTRAGT-3'). A unique 12-base Golay barcode (Ns) precedes the primers for
563 sample identification⁶¹, and one to eight additional nucleotides were placed in front of the barcode to
564 offset the sequencing of the primers. Cycling conditions were 94°C for 3 min, followed by 27 cycles
565 of 94°C for 50 s, 51°C for 30 s, and 72°C for 1 min. For the final elongation step, 72°C for 5 min was
566 used. Replicate PCRs were pooled, and amplicons were purified using the QIAquick PCR Purification
567 Kit (Qiagen). PCR products were quantified and pooled at equimolar amounts before Illumina
568 barcodes and adaptors were ligated, using the Illumina TruSeq Sample Preparation protocol. The
569 completed library was sequenced on an Illumina MiSeq platform following the Illumina
570 recommended procedures with a paired-end 250 \times 250 bp kit

571

572 **Sequence analysis**

573 The 16S (V4-V5) paired-end reads were merged and demultiplexed. Amplicon sequence variants
574 (ASVs) were identified using the Divisive Amplicon Denoising Algorithm (DADA2) pipeline
575 including filtering and trimming of the reads⁶². Reads were trimmed to the first 180 bp or the first
576 point with a quality score $Q < 2$. Reads were removed if they contained ambiguous nucleotides (N) or
577 if two or more errors were expected based on the quality of the trimmed read. We assigned taxonomy
578 to ASVs using a 8-mer based classifier trained by IDTaxa⁶³ using the SILVA database⁶⁴. We
579 determined the copy number of 16S rRNA genes per gram of stool for 4,158 of our samples as
580 reported previously²⁴, by quantitative PCR on total DNA extracted from fecal samples.

581

582 **Quantification of total microbiota density per gram of stool and estimation of total genus**
583 **abundances.**

584 Quantitative PCR (qPCR) was performed on DNA extracted from the 1g wet weight of a stool sample
585 using DyNAmo SYBR Green qPCR kit (Finnzymes) and 0.2 μ M of the universal bacterial primer 8F
586 (5'-AGAGTTTGATCCTGGCTCAG) and the broad-range bacterial primer 338R
587 (5'-TGCTGCCTCCCGTAGGAGT-3'). Standard curves were prepared by serial dilution of the PCR
588 blunt vector (Invitrogen) containing 1 copy of the 16s rRNA gene. Cycling conditions were 95°C for
589 10 minutes followed by 40 cycles of 95°C for 30 seconds, 52°C for 30 seconds, and 72°C for 1
590 minute. We used the measurements of total 16S rRNA gene counts per gram of stool to multiply the
591 relative abundances of taxa obtained from 16S amplicon sequencing to obtain the estimate of their
592 total abundance per gram of stool (supplementary methods). Importantly, this does not account for
593 16S copy number variation between taxa, but the observed dynamic ranges in total abundances of taxa
594 in our data set span up to 9 orders of magnitude, exceeding the potential inaccuracies due to copy
595 number variation.

596

597 **Diversity calculations**

598 Microbiome alpha-diversity was measured by the inverse Simpson (IS) index of a sample. It was
599 calculated by $IS_i = \frac{1}{\sum_{j=1}^N p_{ij}^2}$, where p is the relative abundance of the j th ASV out of N total ASVs in
600 sample i .

601

602 **Linear mixed-effects model of white blood cell counts**

603 To study the effect of auto-FMT on white blood cells, we investigated the white blood cell counts of
604 24 enrolled patients of this trial from the day of neutrophil engraftment until 100 days after. FMT
605 occurred on different days relative to neutrophil engraftment. Thus, we performed an analogous
606 analysis to that conducted in the original publication that demonstrated how FMT re-established a
607 diverse microbiome in the post-FMT period²³. To answer if white blood cell counts differed post-
608 FMT, we used a linear mixed effects model of white blood cell counts, y , modeled as a function of the
609 FMT treatment as well as patient and timepoint specific random effects. We included random
610 intercept terms for each day i and each patient j , and a fixed effects term for the post-FMT period with
611 associated coefficient “armpost”, using the indicator variable “FMT”, that is 1 when a patient was
612 from the FMT treated arm of the trial and day was greater than or equal to the day of the FMT
613 procedure. We conducted independent analyses for neutrophil, lymphocyte and monocyte counts.
614 This resulted in the following model of a cell count, y , for patient j on day i :

$$615 \quad y_{ij} = \beta_0 + \text{armpost} * \text{FMT}_{ij} + \text{day}_i + \text{patient}_j + \varepsilon_{ij}, \quad i = 0, \dots, D, j = 1, \dots, P$$

616 with prior distributions $\text{day}_i \sim \mathcal{N}(0, \sigma_{\text{day}}^2)$, and $\text{patient}_j \sim \mathcal{N}(0, \sigma_{\text{patient}}^2)$, independent error
617 $\varepsilon_{ij} \sim \mathcal{N}(0, \sigma^2)$ and fixed intercept β_0 , for the D days post neutrophils engraftment and P patients,
618 ($D=100, P=24$). For convenience of those interested in reanalyzing our data, the part of our data
619 concerning the auto-FMT analysis is available in tidy format (supplementary data 11), and the
620 analysis code conducted in the R programming language is available as an exported notebook
621 (fmt_effect_on_wbc.pdf) on github: https://github.com/jsevo/wbcdynamics_microbiome/.⁶⁵ We
622 conducted an additional analysis with “day” as a continuous predictor which did not change our
623 conclusions (supplementary methods).

624

625 **Dynamic systems analyses**

626 We analyzed factors associated with the observed changes of absolute counts of neutrophils,
627 lymphocytes and monocytes between two days. In the following we describe how chronology of
628 events and biological samples were encoded, and the models used to infer a role of medications,
629 clinical parameters and the microbiome on dynamics of white blood cells.

630 To reveal factors that associate with day-to-day changes in white blood cell counts, we started
631 from a first-order differential equation of white blood cell (W) dynamics:

$$632 \quad \frac{d(W)}{dt} = W(gr + \sum_{j=1}^P \beta_j X_j)$$

633 Where gr represents the intercept, i.e. the base line rate of change during immune reconstitution, and
634 β_j are the to-be-estimated coefficients of the P predictors $X_j, j \in P$, of the white blood cell dynamics.

635 This equation was then linearized to

$$636 \quad \frac{d \ln(W)}{dt} = gr + \sum_{j=1}^P \beta_j X_j$$

637 And we parameterized the corresponding discrete difference equation:

$$638 \quad \frac{\Delta \ln(W)}{\Delta t} = gr + \sum_{j=1}^P \beta_j X_j$$

639 where $\Delta \ln(W)$ is the log-difference between single days of neutrophils, lymphocytes or monocytes
640 counts, and $\Delta t=1$ for all intervals. Predictors include the counts of neutrophils, lymphocytes,
641 monocytes, eosinophils and platelets during an interval (homeostatic feedbacks), immunomodulatory
642 medication and clinical observations such as a blood stream infection and the onset of graft versus
643 host disease, HCT parameters such as graft types and conditioning regimens, and, additionally, the
644 microbiota composition in “stage 2” of our analysis (supplementary methods for data exclusion and
645 additional details on interval definitions). Importantly, by parameterizing a dynamic equation and
646 analyzing *rates of change*, our coefficient estimates have an immediate causal interpretation within
647 our modeling framework (i.e. a $\beta_j > 0$ implies that higher levels of the corresponding X_j *increases* the

648 respective white blood cell type, W). To differentiate such results from other associations, analyses of
649 this type have been termed “mathematical causality”⁵⁴.

650

651 *Stage 1 analysis: Feature selection. Identifying medications and clinical observations associated with*
652 *white blood cell dynamics from patients without microbiome data*

653 Stage 1 uses data of patients without any available microbiome samples and the following model of
654 white blood cell changes, y :

$$655 \quad y = gr + \sum_{j=1}^p \beta_j X_j,$$

656 with intercept, gr . The predictors, X , include dummy variables for the HCT graft type, patients’ age on
657 the date of HCT, sex, 13 most frequently observed positive blood cultures with remaining other blood
658 stream infections grouped into a separate category “other infections”, an indicator for the onset of
659 graft versus host disease, administrations of 55 different, most common immunomodulatory
660 medications and platelet transfusion events, and HCT conditioning intensity regimens as well as the
661 log-transformed geometric mean counts of neutrophils, lymphocytes, monocytes, eosinophils and
662 platelets during the respective interval. We used elastic net regression⁶⁶ for feature selection using the
663 sklearn package for the Python programming language⁶⁷. For elastic net regression with 50% L1-
664 penalty, predictors were scaled between zero and 1, and we used 10-fold cross validation (i.e. leaving
665 out 10% of patients at each cross-validation step) to choose the regularization strength, λ , solving for

$$666 \quad \operatorname{argmin}_{gr, \beta} \left\{ \frac{1}{2N} \sum_{i=1}^N (y_i - gr - \sum_{j=1}^p x_{ij} \beta_j)^2 + \frac{1}{2} \lambda \sum_{j=1}^p |\beta_j| + \frac{1}{2} \lambda \sum_{j=1}^p \beta_j^2 \right\}$$

667
668 Stage 1 yielded a sparse coefficient matrix of predictors used to design the model in stage 2.

669

670 *Expanded analysis on patients with microbiome data – stage 2*

671 To identify associations between microbiota and white blood cell dynamics, we conducted an
672 analogous, Bayesian regression using the package PyMC3 for the Python programming language⁶⁸.
673 Stage 1 identified important difference between transplant types, and we therefore stratified our data
674 into 4 cohorts according to their stem cell graft source. Using data independently from each cohort,

675 we applied “no U-turn” sampling⁶⁹ to produce 10,000 posterior samples from 5 independent MCMC
676 chains that parameterized the model:

$$677 \quad y \sim \mathcal{N}(\mu, \sigma^2)$$

$$678 \quad \mu = gr + \sum_{j=1}^{\hat{P}} x_j \beta_j$$

679 with uninformative prior distributions

$$680 \quad gr \sim \mathcal{N}(\text{mean} = 0, \text{standard deviation} = 100)$$

$$681 \quad \beta_j \sim \mathcal{N}(\text{mean} = 0, \text{standard deviation} = 100)$$

$$682 \quad \sigma \sim \text{HalfCauchy}(\text{beta} = 2)$$

683 where y is the observed daily change of a focal white blood cell type as in stage 1 with normal
684 distributed mean, μ , and σ , the model uncertainty with a thick-tailed half Cauchy prior (importantly,
685 our posterior estimates do not depend on this choice as we obtain the same results with an inverse
686 Gamma prior, figure S19). μ was a function of the baseline growth rate, gr , and predictors, \hat{P} :
687 medications with non-zero coefficients in stage 1, the white blood cell counts, patient age and sex,
688 and HCT conditioning intensities; additionally, \hat{P} now included the log-abundances of microbial
689 genera as measured by 16S sequencing from DNA in the stool collected on the second day of a daily
690 interval (see supplementary methods for details). We considered taxa that were among the 100 most
691 abundant, or had reached maximum relative abundances of at least 10%, and selected those who were
692 non-zero in more than 75% of our samples. White blood cell counts and microbiota data present
693 during a daily interval were log-transformed, and zeros were filled with half of the minimum observed
694 non zero counts (i.e. $0.5e3$ and $2e-6$, respectively). We focused on the largest cohort (PBSC) and
695 used the independent inference results from TCD, BM, and cord cohorts for validation.

696

697 *Validation score*

698 Coefficients learnt from the PBSC patient cohort were assigned a “validation score” based on the
699 results obtained from the other three MSK patient cohorts. Our requirements for validation were
700 conservative; we required evidence from our validation data sets as well as absence of counter

701 evidence. For regression results from each of the validation graft type cohorts, i.e. TCD, BM, and
702 cord, we checked if a coefficient had more than 75% probability (50%HPDI) to have the same sign as
703 the mean of the PBSC coefficient posterior for a given predictor. If so, this was considered evidence
704 of validation, and we summed the evidence over the three validation sets (i.e. maximum score of 3, 1
705 from each of TCD, BM, and cord cohorts). Conversely, if we found more than 75% probability
706 among any of the validation data sets that a given predictor had the *opposite* sign as the posterior
707 mean calculated from PBSC data, this was considered counter evidence and the validation score was
708 always set to zero.

709

710 *Analysis of white blood cell dynamics with absolute bacterial abundances as predictors instead of*
711 *relative abundances*

712 We conducted an ordinary least squares regression using the statsmodels package in the Python
713 programming language of the same model as in the main Bayesian analysis using total bacterial
714 abundances as predictors. This was only possible on a subset of 389 neutrophil, 331 lymphocyte and
715 376 monocyte rate observations from PBSC patients.

716

717 *Forwards simulation of predicted immune system reconstitution kinetics*

718 To assess the impact of the estimated microbiota coefficients on immune system dynamics, we
719 conducted 1,000 simulations of the system of 3 differential equations describing the dynamics of
720 neutrophils, lymphocytes and monocytes. We ran 1,000 simulations four times: in presence and
721 absence of GCSF, each with microbiota compositions enriched or depleted in *Faecalibacterium*,
722 *Ruminococcus 2* and *Akkermansia*. To identify these compositions, we ranked the observed
723 microbiota compositions by these taxa, and chose randomly either from the top or bottom 100. The
724 coefficients for white blood cell interactions, interactions with the microbiota and the effect of GCSF
725 were sampled from our posterior coefficient distributions. Using these coefficients sampled at the start
726 of the simulation, and using 50 cells* μl^{-1} of neutrophils, lymphocytes and monocytes as initial values,
727 we simulated these differential equations forwards in time using the odeint function of the scipy
728 package for the Python programming language.

729

730 *Validation on data from Duke University*

731 We analyzed 9,603 blood samples with 25,581 associated administrations of immunomodulatory
 732 medications, and 741 microbiota samples from Duke as an orthogonal data set to validate our
 733 findings. The temporal resolution of this data was much lower, and after filtering for samples from the
 734 relevant post neutrophil engraftment period, and by requiring daily intervals, 83 valid, complete data
 735 points were available. Using these data, we correlated daily blood cell changes individually in
 736 univariate, or jointly in a partial least squares regression, with those predictors that achieved more
 737 than 95% probability density in the positive or negative domain in the PBSC data regression. For each
 738 of these predictors, we present the sign of slopes and Bonferroni corrected p -values from individual
 739 linear regressions.

740

741 *Joint analysis of the effect of antibiotics and white blood cell counts on the microbiota and the*
 742 *microbiota and immunomodulatory medications on white blood cell counts*

743 Analogous to *stage 1*, we performed cross-validated, regularized linear regressions (ElasticNet) using
 744 the scikit-learn package for the Python programming language to jointly estimate the association
 745 network between microbiota and circulatory white blood cells. For this, we constructed a block matrix
 746 \mathbf{X} of predictor matrices \mathbf{X}_i that include the absolute bacterial abundances, drug data (antibiotics for
 747 bacterial dynamics and immune modulators for white blood cell dynamics), as well as the counts of
 748 white blood cells and a separate intercept term per block. Each block \mathbf{X}_{n_l, p_l}^l , with n_l observations and p_l
 749 predictors ($l=0\dots k$), on the diagonal of \mathbf{X} corresponds to the indices of the observed daily log-changes
 750 of one of the 41 bacterial genera considered in our main analysis or the log changes in neutrophil,
 751 lymphocyte and monocyte counts from PBSC patients contained in \mathbf{Y} (in total we calculated 15,833
 752 rates from 256 patients). Our regression problem can thus be written as:

$$753 \quad \underset{\beta}{\operatorname{argmin}} (\mathbf{Y} - \mathbf{X}\beta) \quad \text{where } \mathbf{X} = \begin{matrix} \mathbf{X}_{n_0, p_0}^0 & \cdots & \mathbf{0}_{n_0, p_k} \\ \vdots & \ddots & \vdots \\ \mathbf{0}_{n_k, p_0} & \cdots & \mathbf{X}_{n_k, p_k}^k \end{matrix}$$

754 with $k=44$, i.e. 41 bacterial genera and 3 white blood cell types, the to-be estimated coefficient vector
 755 β and $\mathbf{0}$ the zero matrix. This system is underdetermined and we therefore chose the same approach as

756 in stage 1, elastic net regression, for feature selection. Predictors were scaled between zero and 1, and
757 we used 3-fold cross validation, leaving out 1/3rd of the patients at each iteration to identify a global
758 regularization strength, λ , solving for

$$759 \quad \operatorname{argmin}_{\beta} \left\{ \frac{1}{2\eta} \sum_{i=1}^{\eta} (y_i - \sum_{j=1}^{\rho} x_{ij}\beta_j)^2 + \frac{1}{2}\lambda \sum_{j=1}^{\rho} |\beta_j| + \frac{1}{2}\lambda \sum_{j=1}^{\rho} \beta_j^2 \right\}$$

760
761 where η is the total number of observed daily log changes in genera and white blood cells, and ρ the
762 total number of predictors. This yielded a strongly regularizing λ_s , and thus few predictors. To
763 characterize potential bidirectional relationships between white blood cell counts and the gut
764 microbiota, we iteratively reduced the regularization strength until the strongest interaction between
765 microbiota and white blood cell dynamics, i.e. *Faecalibacterium* with neutrophil dynamics, was
766 detected. We then re-ran the regression with this reduced regularization strength, λ_r .

767

768 *Shotgun sequencing*

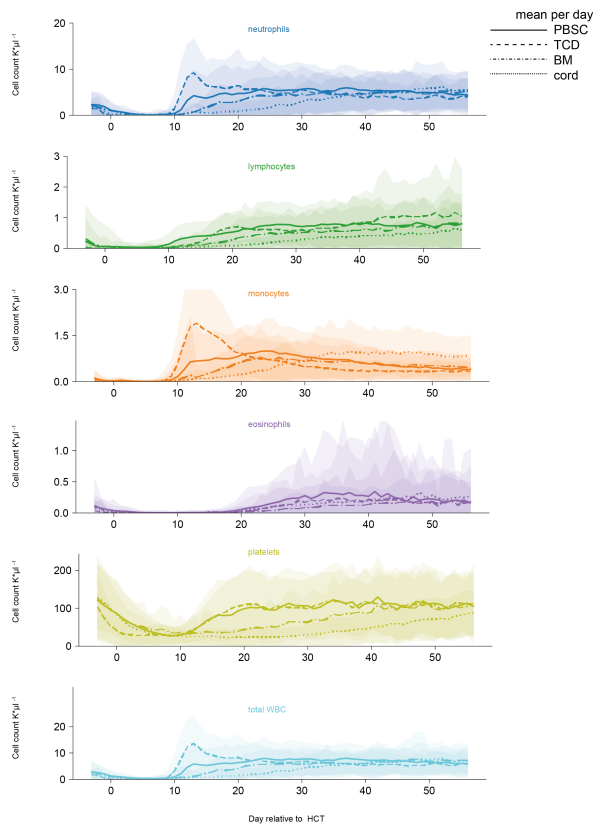
769 Sequencing of 124 post-neutrophil engraftment was conducted on the Illumina HiSeq platform. For
770 details and the processing of the FASTQ files, see supplementary methods. We used the HUMAnN2
771 pipeline⁷⁰ with default settings for functional profiling of our samples, with the UniRef90 data base
772 and ChocoPhlAn for alignment, and we renormalized our samples by library depth to copies per
773 million. We used MetaCyc to obtain stratified and unstratified pathway abundances.

774

775 *Statistical analysis of shotgun data*

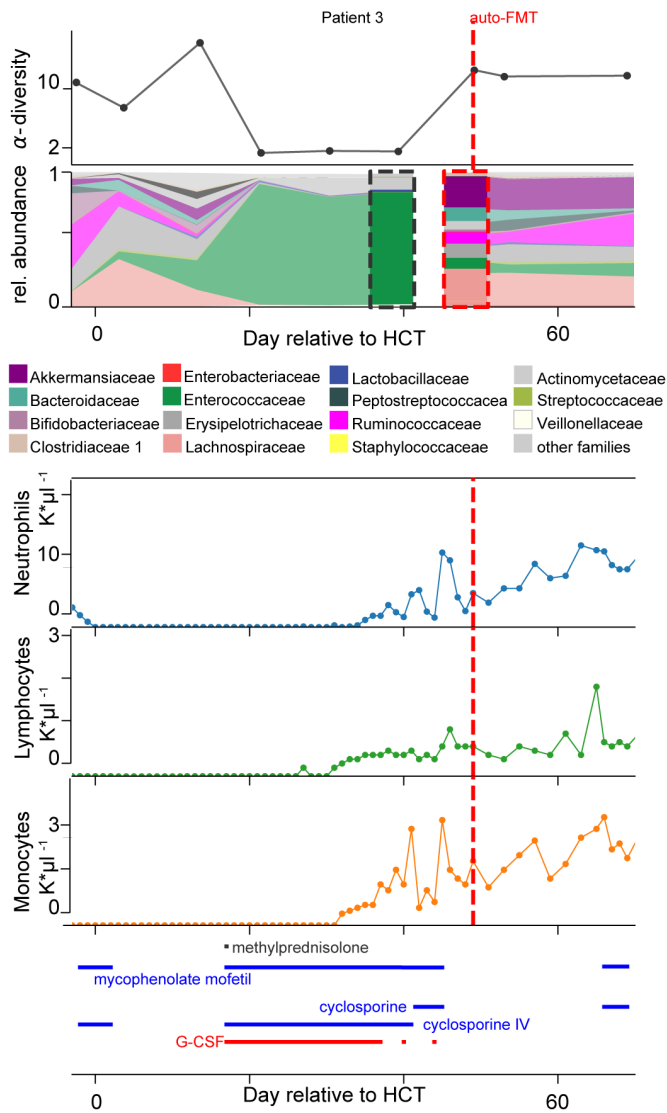
776 We calculated the predicted microbiota potency score for each sample and separately for neutrophils,
777 lymphocytes and monocytes, by multiplying the abundances of taxa in each of the 124 samples with
778 the corresponding posterior coefficients obtained from the PBSC inference. To distinguish the sets of
779 metabolic functions that separate samples with positive and negative predicted potencies, we
780 converted the pathway abundances into presence and absence profiles. We performed a linear
781 discriminant analysis between positive and negative potency samples with a least squares solver and
782 automatic shrinkage using the Ledoit-Wolf lemma using the sklearn package for the Python
783 programming language⁶⁷. To assess differences in the presence or absence of pathways between
784 samples with positive and negative potency, we used Fisher's exact test for each pathway.

785 **Supplementary Figures**



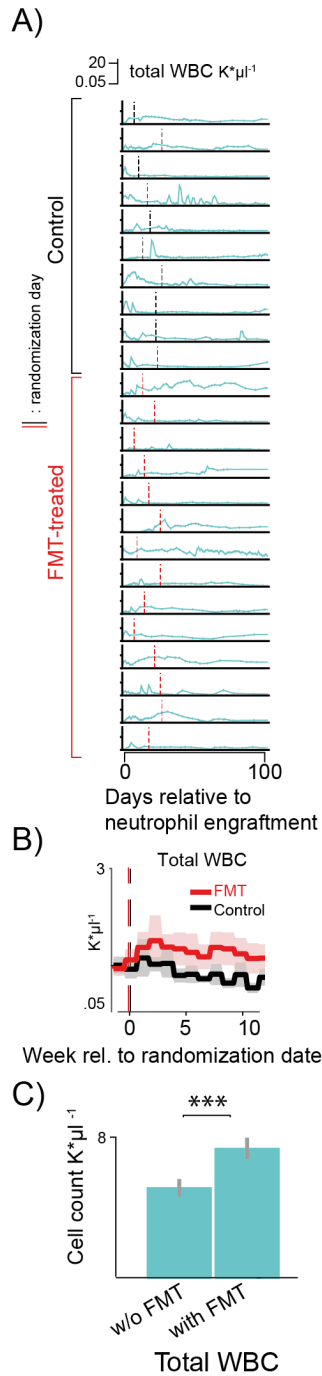
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Figure S1: White blood cell counts and platelet counts per graft source over the first 100 days post HCT per day relative to HCT; lines: mean, shaded: \pm standard deviations).



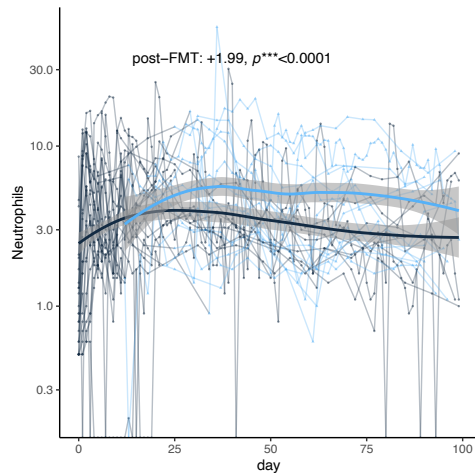
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Figure S2: HCT patient who received an autologous fecal microbiota transplant (auto-FMT, dashed red line) that restored commensal microbial families and ecological diversity in the gut microbiota, with concurrent cell counts of peripheral neutrophils, lymphocytes and monocytes and immunomodulatory drug administrations.



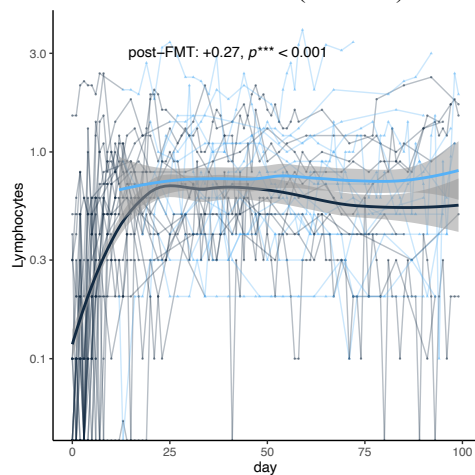
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Figure S3: Total counts of white blood cells increased in FMT-treated patients in the weeks following the treatment compared to control patients. A) Total white blood cell counts in 24 patients enrolled in a randomized controlled trial to receive an autologous fecal microbiota transplant post-neutrophil engraftment (10 control: black vertical line, 14 FMT treated: red vertical line,). B) Weekly mean cell counts aligned to the date of randomization into FMT treatment arm (red) or control (black). Line: weekly mean, shaded region: 95%-CI C) Results from a linear mixed effects model with random effects per patient and per day relative to neutrophil engraftment confirms that the total white blood cell counts is higher in patients receiving auto-FMT after their treatment as compared to control patients after the randomization date, bars and confidence intervals for the averages of observed white blood cell counts without FMT and post-FMT (***: $p < 0.001$).



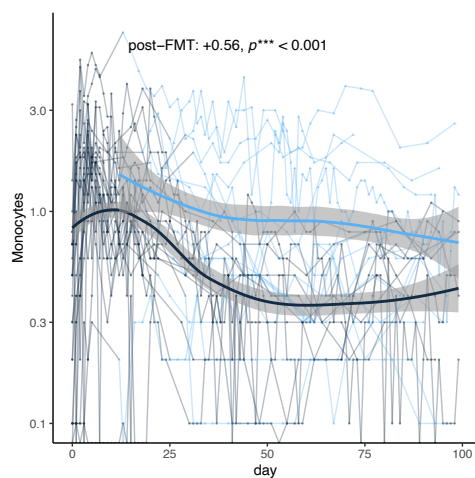
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Figure S4: Neutrophil counts in 24. FMT trial patients. Thin lines: raw data (blue: post-FMT); thick black: mean per day, thick blue: mean+post-FMT coefficient. Means and confidence intervals (shaded region) from linear mixed effects model (methods).



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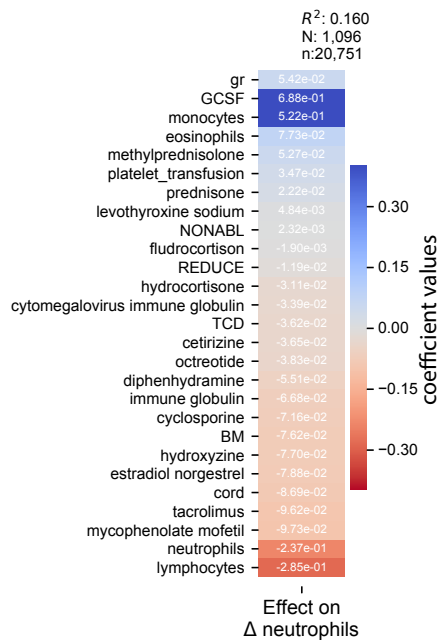
Figure S5: Lymphocyte counts in 24. FMT trial patients. Thin lines: raw data (blue: post-FMT); thick black: mean per day, thick blue: mean+post-FMT coefficient. Means and confidence intervals (shaded region) from linear mixed effects model (methods).



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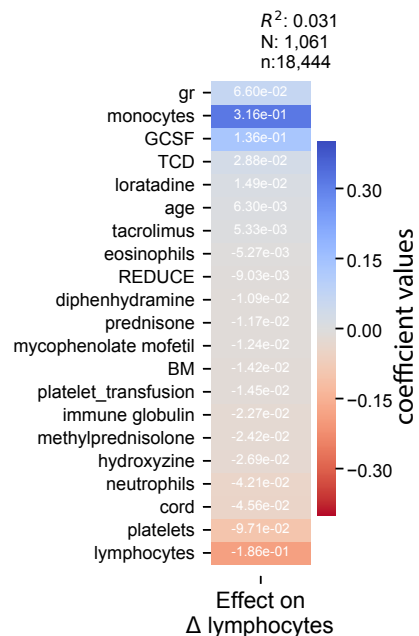
Figure S6: Monocyte counts in 24. FMT trial patients. Thin lines: raw data (blue: post-FMT); thick black: mean per day, thick blue: mean+post-FMT coefficient. Means and confidence intervals (shaded region) from linear mixed effects model (methods).

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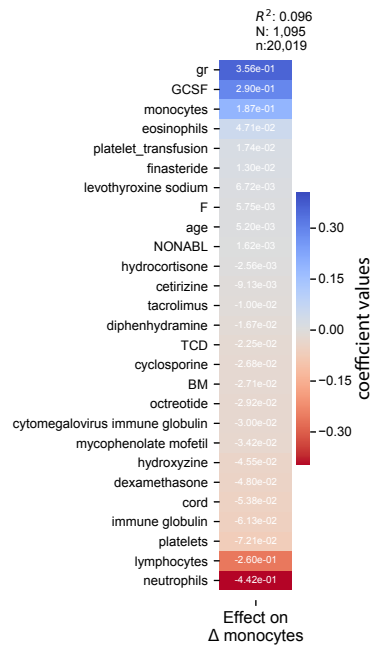
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Figure S7: “Stage 1” regression on neutrophil dynamics on patients without microbiome data. Coefficients from 10-fold cross-validated elastic net regression daily changes in neutrophils. gr: intercept; TCD: T-cell depleted graft (ex-vivo) by CD34+selection; PBSC: peripheral blood stem cells; BM: bone marrow; cord: umbilical cord blood; NONABL: Nonmyeloablative; REDUCE: reduced-intensity conditioning regimen; F: female; N: patients, n: samples (daily changes in neutrophils).



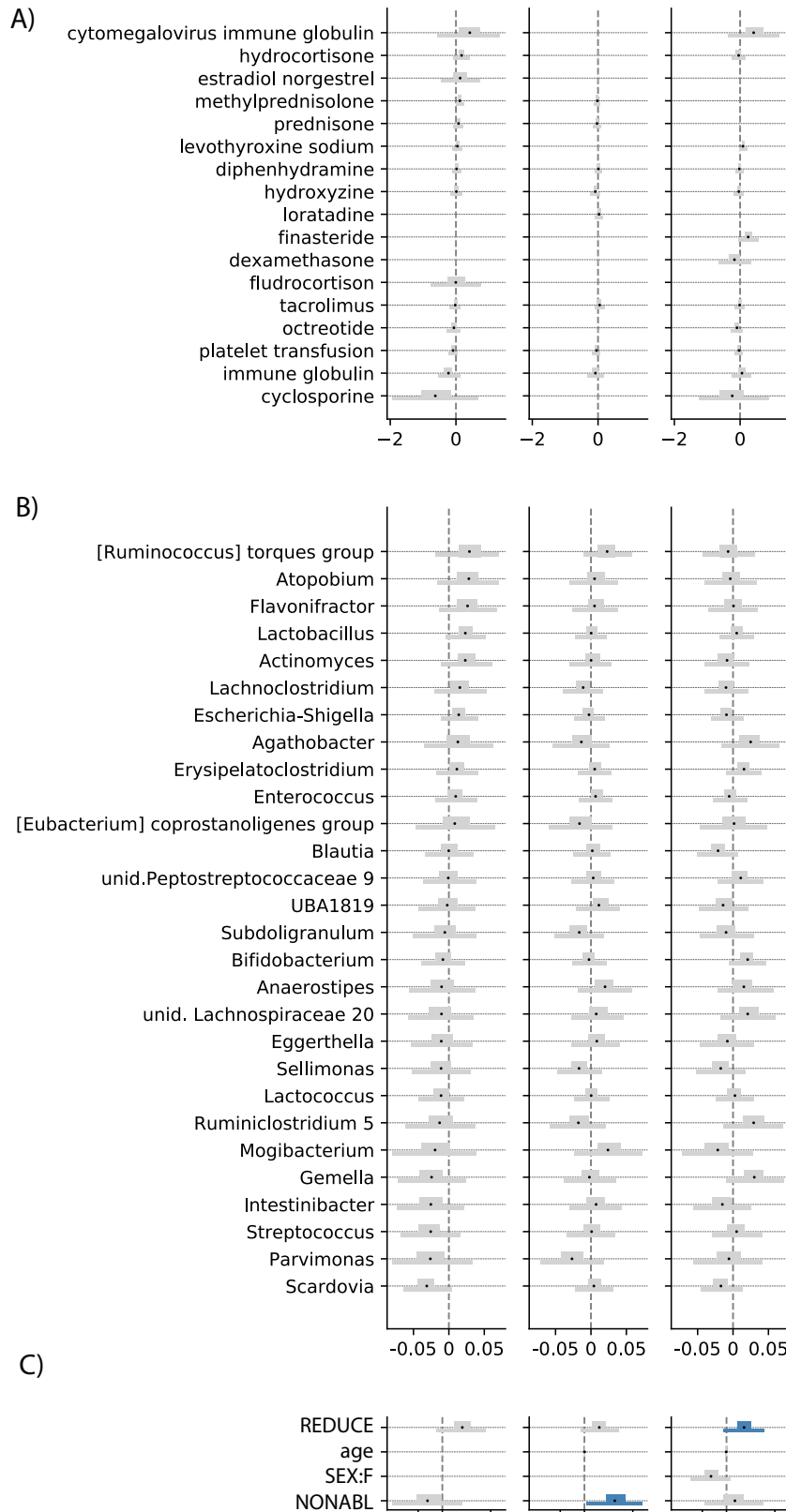
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Figure S8: “Stage 1” regression on lymphocytes dynamics on patients without microbiome data. Coefficients from 10-fold cross-validated elastic net regression daily changes in lymphocytes. gr: intercept; TCD: T-cell depleted graft (ex-vivo) by CD34+selection; PBSC: peripheral blood stem cells; BM: bone marrow; cord: umbilical cord blood REDUCE: reduced-intensity conditioning regimen; F: female. N: patients, n: samples (daily changes in lymphocytes).



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Figure S9: “Stage 1” regression on lymphocytes dynamics on patients without microbiome data. Coefficients from 10-fold cross-validated elastic net regression daily changes in lymphocytes. gr: intercept; TCD: T-cell depleted graft (ex-vivo) by CD34+selection; PBSC: peripheral blood stem cells; BM: bone marrow; cord: umbilical cord blood REDUCE: reduced-intensity conditioning regimen; F: female. N: patients, n: samples (daily changes in lymphocytes).



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Figure S10: Additional coefficient estimates of medications (A), additional genera (B) and metadata (C) from the Bayesian regression, see also Figure 3. REDUCE: reduced-intensity conditioning regimen; NONABL: non-meloablative conditioning regimen. F: female



Figure S11: Posterior sampling converged. Histograms of the ranked posterior draws from the model of neutrophil dynamics in PBSC patients (ranked over all chains), plotted separately for each chain (see supplementary methods), show no substantial differences between chains⁷¹.

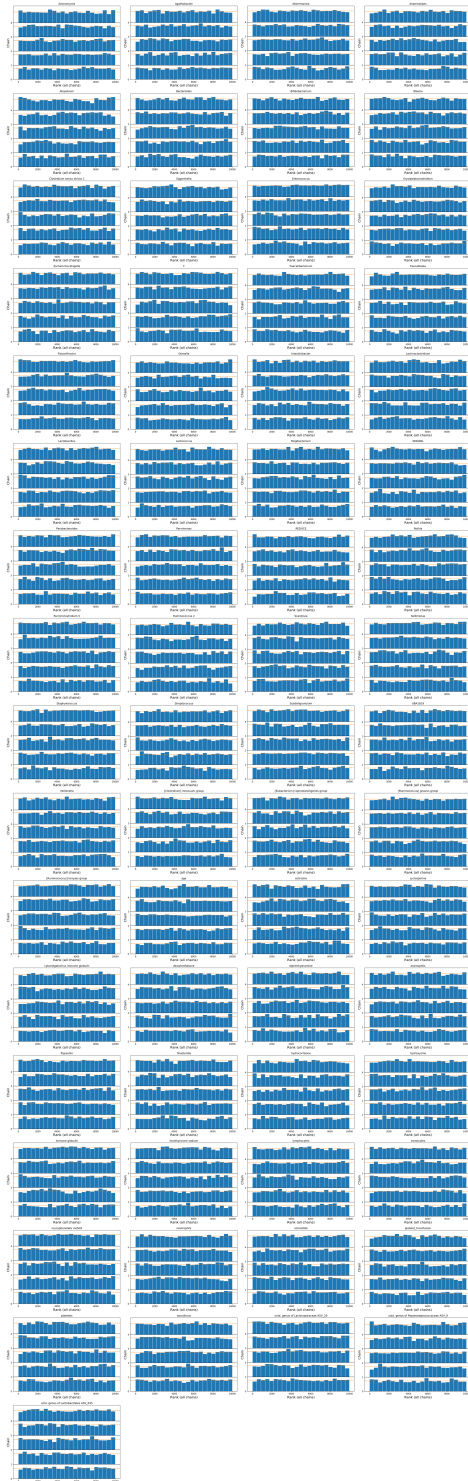
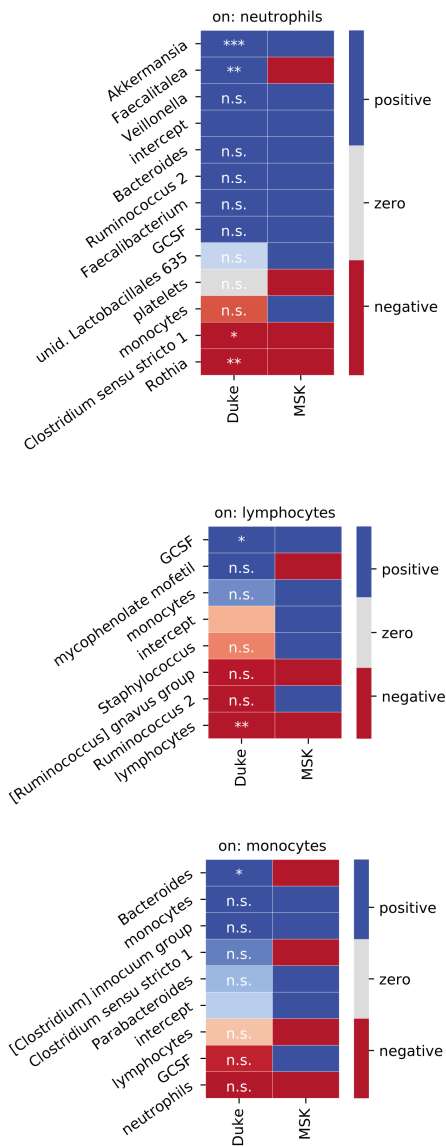


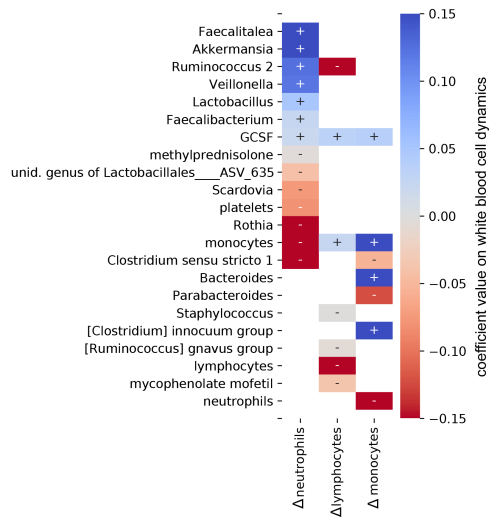
Figure S13: Posterior sampling converged. Histograms of the ranked posterior draws from the model of monocyte dynamics in PBSC patients (ranked over all chains), plotted separately for each chain (see supplementary methods), show no substantial differences between chains⁷¹.

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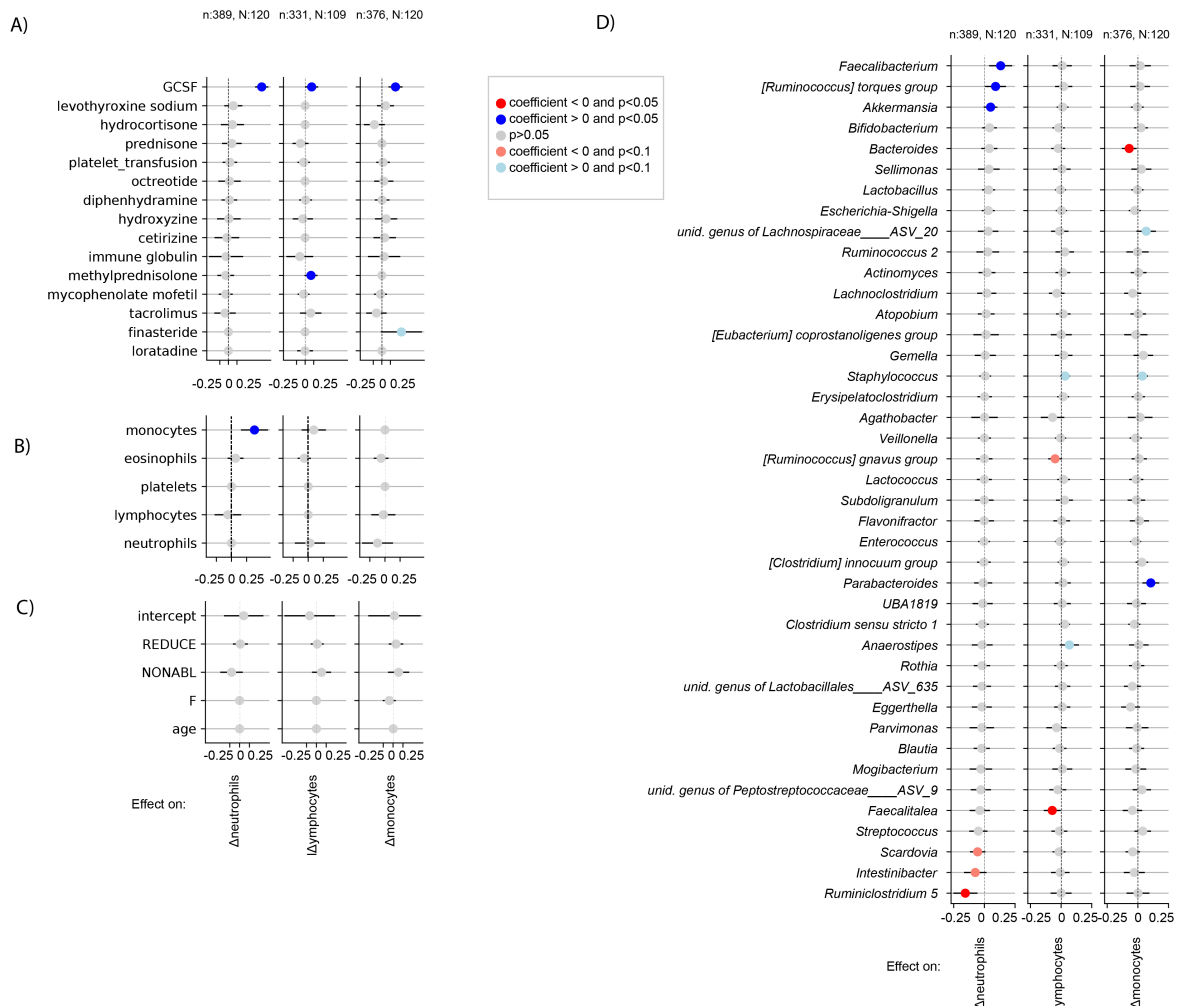
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Figure S14: Validation analysis of predictors on white blood cell dynamics using data from patients treated at DukeHealth. Individual univariate regressions of microbiome and clinical predictors identified in stage 2 of our analysis on daily changes in neutrophils, lymphocytes and monocyte. Bonferroni corrected p-values: ***<0.001, **<0.01, *<0.05; p>0.05: n.s. Sign of coefficients from MSK PBSC patients for comparison.



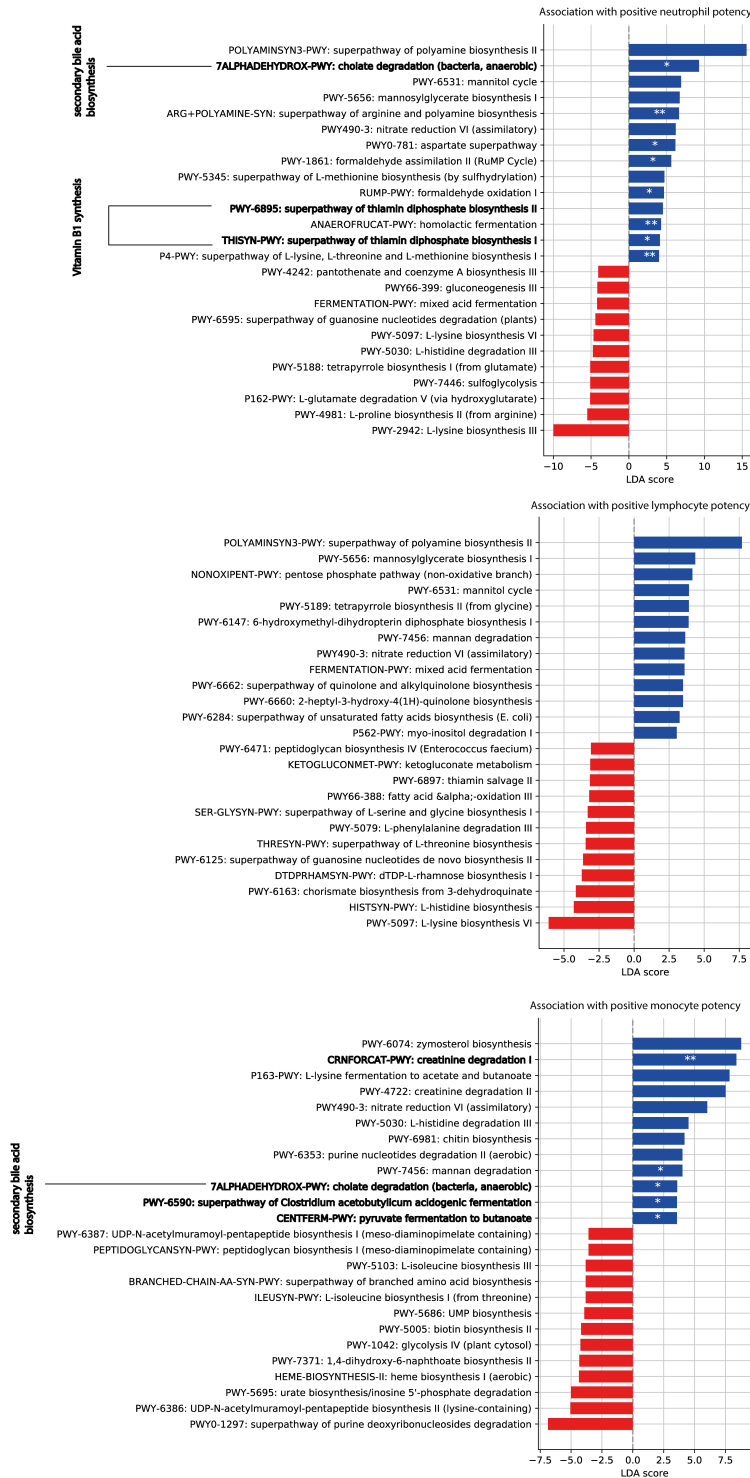
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Figure S15: Validation analysis of predictors on white blood cell dynamics using data from patients treated at DukeHealth. Partial least squares regression of microbiome and clinical predictors identified in stage 2 of our analysis on daily changes in neutrophils, lymphocytes and monocyte.



883

884 **Figure S16:** Validation analysis of the main model using absolute bacterial abundances as predictors instead of
 885 relative abundances in Figure 3. Results show coefficients from a least squares regression for medications (A),
 886 white blood cell feedbacks (B) metadata (C) and total genus abundances (D) of neutrophil, lymphocyte and
 887 monocyte daily log-changes. This was only possible for only a subset of the data for which we obtained absolute
 888 bacterial abundance estimates (methods), n: samples, N: patients.
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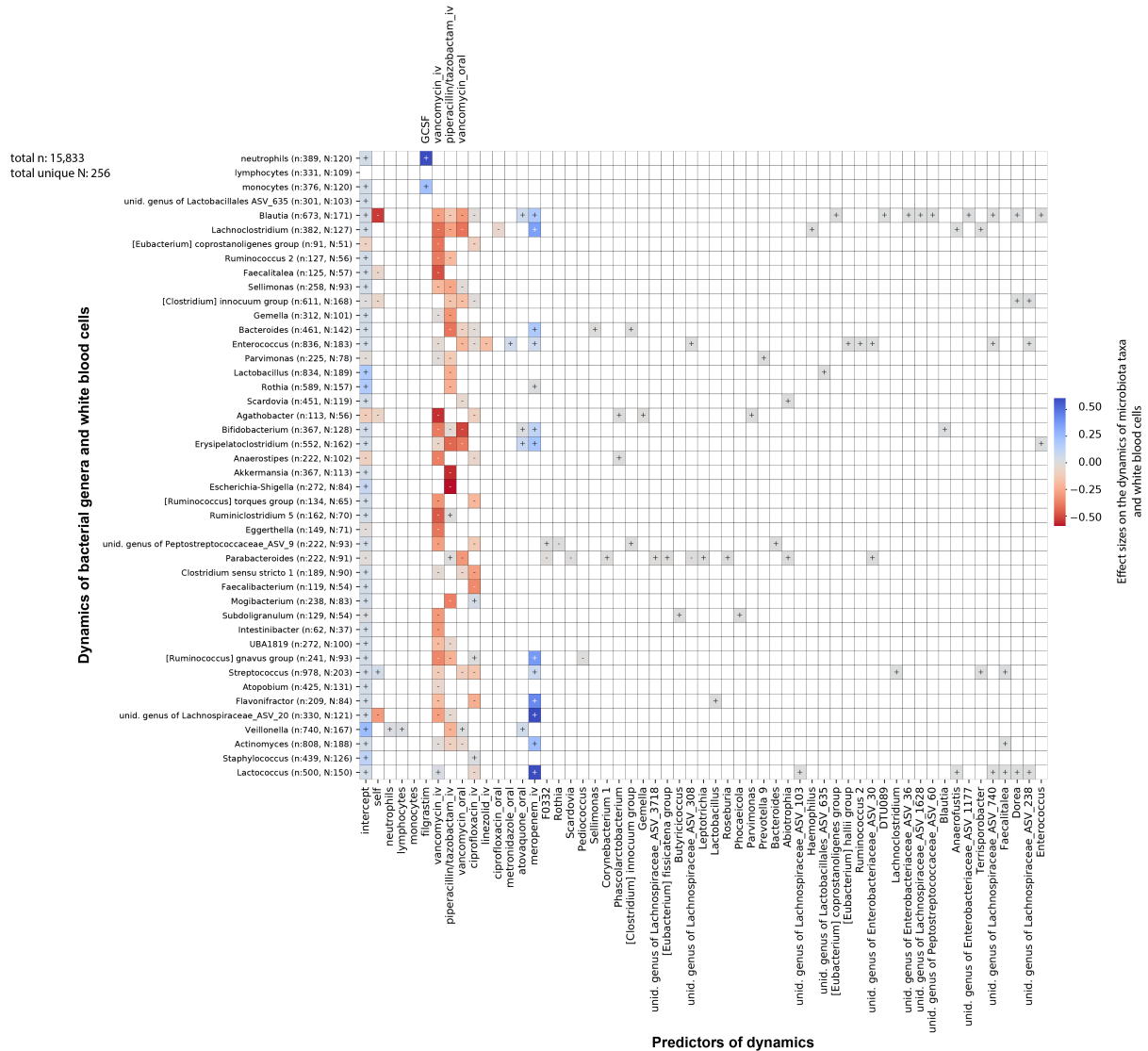


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Figure S17: Functional analysis of microbiota samples. To distinguish samples predicted to increase rates of white blood cells, a microbiota potency score was calculated from posterior coefficients (Figure 3, methods) and the relative abundance of taxa in samples. Bars show linear discriminant analysis (LDA) scores of MetaCyc

896 pathway profiles from 124 shotgun sequenced samples that distinguished positive and negative potency samples
 897 the most (LDA-score magnitude in the 95th percentile). Highlighted pathways are discussed in the main text. For
 898 each pathway, we tested differences between positive and negative potency samples using Fisher's exact test; p-
 899 value <0.001: ***, <0.01: **, <0.05: *.

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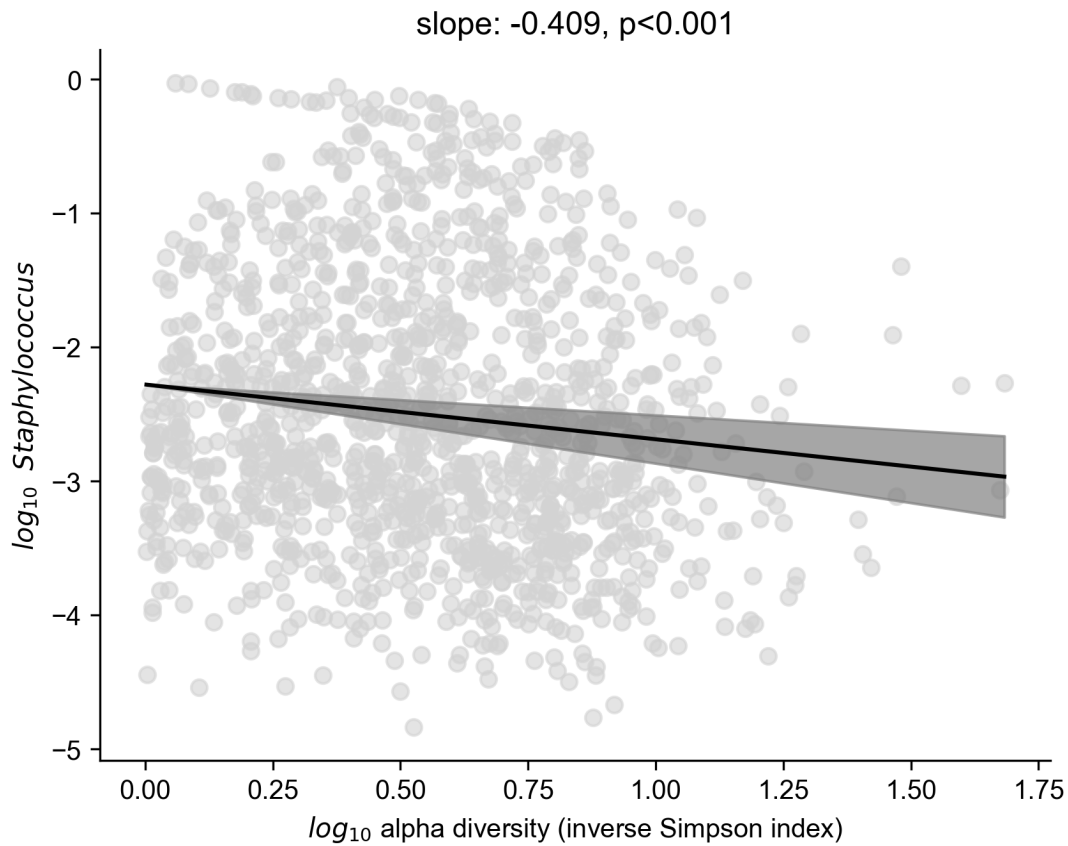
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Figure S18: Jointly inferred association network between white blood cell and bacterial genus dynamics (methods). Strong regularization yields few non-zero coefficients and antibiotics dominate the dynamics.



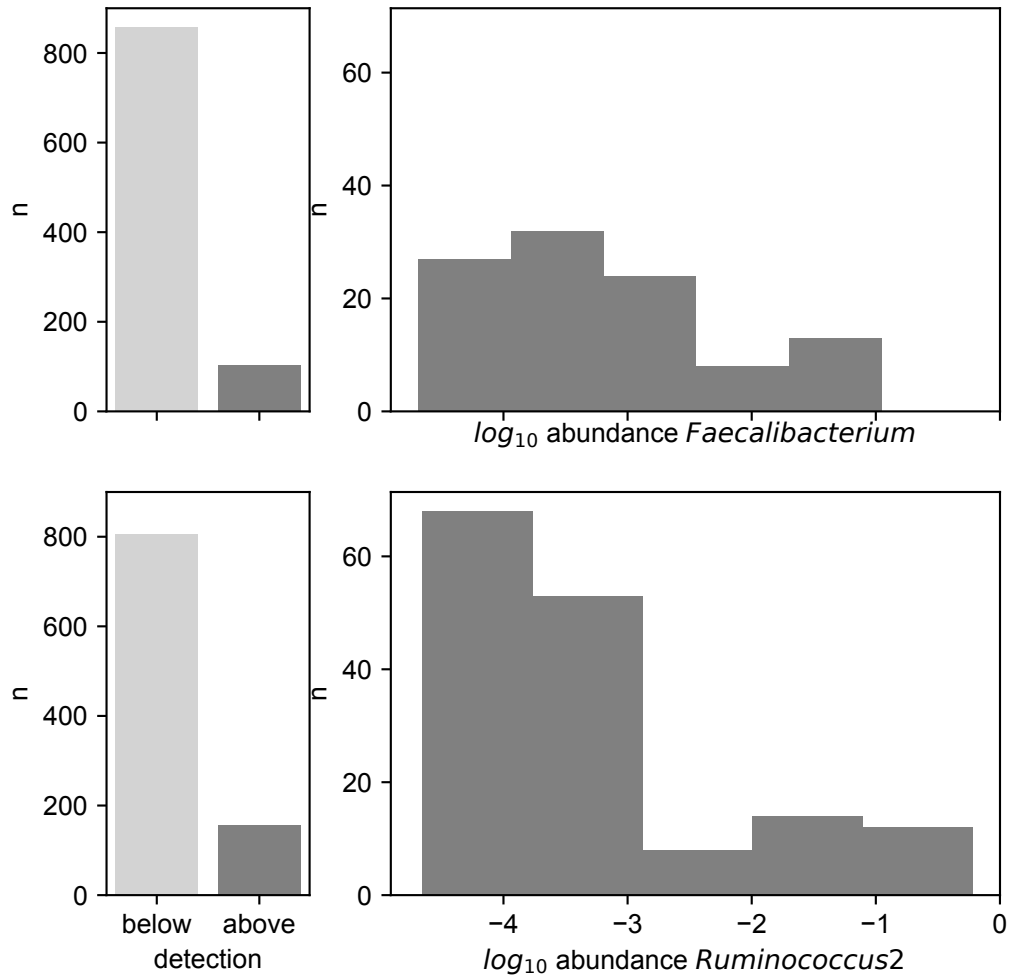
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Figure S19: Jointly inferred association network between white blood cell and bacterial genus dynamics with reduced regularization (methods) indicates potential bidirectional feedbacks, e.g. between lymphocytes and *[Ruminococcus] gnnavus* group (highlighter green boxes, and cartoon).



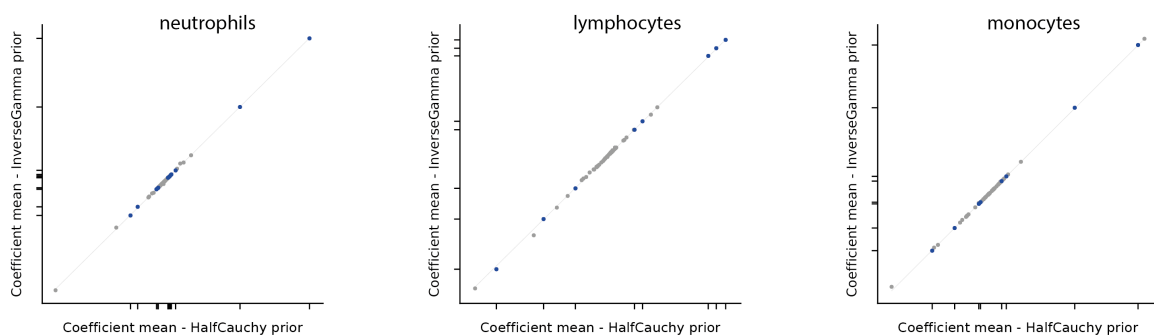
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Figure S20: The relative non-zero abundance of *Staphylococcus* is inversely related to microbiome alpha diversity, shaded: 95% confidence intervals.



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Figure S21: Abundance profiles of the two genera, *Faecalibacterium* and *Ruminococcus 2*, most strongly associated with white blood cell increase; number of times detected (left) and \log_{10} abundance distribution when above detection (right).



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Figure S22: Posterior association coefficients do not depend on the choice of prior for σ in the main Bayesian model. Plotted are the posterior means from our main analysis against the equivalent inference with an inverse Gamma prior ($\alpha=1$, $\beta=1$).

942 Table S1: Data set summary and patient characteristics. HCT-graft types: TCD: T-cell depleted graft (*ex-vivo*)
 943 by CD34+selection; PBSC: peripheral blood stem cells; BM: bone marrow; cord: umbilical cord blood;
 944 Conditioning intensity: Bacigalupo classification, graded categories from most to least intense (ABLATIVE,
 945 REDUCE, NONABL).
 946

patients		2,926
HCT therapies*		3,060
blood samples	total	450,635
	between HCT-day -21 and HCT-day 183	193,396
Disease	Leukemia	1,635
	Non-Hodgkin's Lymphoma	415
	Multiple Myeloma	170
	Hodgkin's disease	88
	other	752
HCT graft type	TCD	1,106
	PBSC unmodified	959
	BM unmodified	617
	cord	378
Conditioning intensity	ABLATIVE	65%
	REDUCE	21%
	NONABL	13%
Gender	M	58%
	F	42%
Age of adults (years)	25%-tile	39
	mean	50
	75%-tile	62
Microbiome samples	total	12,633
	- from patients with blood data	10,680
	- of those, post engraftment	4,179
	- of those with daily change in WBC	2,615
	patients with microbiome sample	1,290

947 *) some patient received several HCTs

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Table S2: Patient and HCT characteristics of 24 patients enrolled in the randomized controlled FMT trial.

	control	FMT treated
N patients	10	14
ABLATIVE	6	7
REDUCE	4	7
BM unmodified	1	3
PBSC unmodified	3	4
TCD	5	3
cord	1	4

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959 Table S3: Patient and HCT characteristics of the subset of patients who donated microbiota samples.

patients		1,294
Disease distribution	Leukemia	51%
	Non-Hodgkin's Lymphoma	15%
	Multiple Myeloma	8%
	Hodgkin's disease	3%
	other	23%
HCT graft type	TCD	37%
	PBSC unmodified	38%
	BM unmodified	9%
	cord	16%
Conditioning intensity	ABLATIVE	55%
	REDUCE	34%
	NONABL	11%
Gender	M	59%
	F	41%
Age (years)	25%-tile	46
	mean	54
	75%-tile	65

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962 Table S4: Patient and HCT characteristics of the subset of patients who did not donate microbiota samples.

patients		1,010
Disease distribution	Leukemia	53%
	Non-Hodgkin's Lymphoma	17%
	Multiple Myeloma	5%
	Hodgkin's disease	5%
	other	20%
HCT graft type	TCD	40%
	PBSC unmodified	31%
	BM unmodified	17%
	cord	12%
Conditioning intensity	ABLATIVE	65%
	REDUCE	14%
	NONABL	21%
Gender	M	58%
	F	42%
Age (years)	25%-tile	36
	mean	47
	75%-tile	59

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966 Table S5: Patient and HCT characteristics of the Duke University patient cohort.

patients		493
Disease distribution	Lymphoma	11%
	Leukemia	50%
	Non-Hodgkin's Lymphoma	4%
	Multiple Myeloma	8%

HCT graft type	Hodgkin's disease	4%
	other	24%
	TCD	0%
	PBSC unmodified	72%
Conditioning intensity	BM unmodified	11%
	cord	16%
	ABLATIVE	92%
Gender	NONABL	7%
	M	65%
Age (years)	F	35%
	25%-tile	41
	mean	49
	75%-tile	57

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Data availability

970 The data used in our study is organized in supplementary tables (data-tables.zip), with corresponding
971 filenames (*italic*):

- 972 1. *cGENUS.csv*: relative taxon abundances in fecal microbiota samples from 12,633 stool
973 samples
- 974 2. *cHCTMETA.csv*: HCT characteristics
- 975 3. *cINFECTIONS.csv*: positive blood culture results
- 976 4. *cMISAMPLES.csv*: NCBI SRA accession number, diversity (inverse Simpson index), total
977 16S (where available), stool consistency for each fecal microbiota sample
- 978 5. *cMED.csv*: medication data
- 979 6. *cPIDMETA.csv*: anonymized patient demographics
- 980 7. *cWBC.csv*: absolute counts of neutrophils, lymphocytes, monocytes, eosinophils, and platelets
981 with indication if included in analyses
- 982 8. *cDUKE_GENUS.csv*: relative taxon abundances in fecal microbiota samples from 12,633
983 stool samples
- 984 9. *cDUKE_WBC.csv*: absolute counts of neutrophils, lymphocytes, monocytes, eosinophils,
985 and platelets with indication if included in analyses
- 986 10. *cDUKE_MED.csv*: medication data
- 987 11. *cFMT_analysis.csv*: convenience table for Figure 2

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Code availability

991 The relevant scripts for stage 1, stage 2, and the model assessing the effect of FMT on white blood
992 cell counts are on github:https://github.com/jsevo/wbcdynamics_microbiome.