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Article

Chronic lymphocytic leukemia B-cell-derived TNF& impairs bone marrow myelopoiesis

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Summary

TNF α is implicated in chronic lymphocytic leukemia (CLL) immunosuppression and disease progression. TNF α is constitutively produced by CLL B cells and is a negative regulator of bone marrow (BM) myelopoiesis. Here, we show that co-culture of CLL B cells with purified normal human hematopoietic stem and progenitor cells (HSPCs) directly altered protein levels of the myeloid and erythroid cell fate determinants PU.1 and GATA-2 at the single-cell level within transitional HSPC subsets, mimicking ex vivo expression patterns. Physical separation of CLL cells from control HSPCs or neutralizing TNF α abrogated upregulation of PU.1, yet restoration of GATA-2 required TNF α neutralization, suggesting both cell contact and soluble-factor-mediated regulation. We further show that CLL patient BM myeloid progenitors are diminished in frequency and function, an effect recapitulated by chronic exposure of control HSPCs to low-dose TNF α . These findings implicate CLL B-cell-derived TNF α in impaired BM myelopoiesis.

Introduction

Chronic lymphocytic leukemia (CLL), a chronic lymphoproliferative neoplasm characterized by accumulation of CD19⁺CD5⁺ B cells, is associated with global immunosuppression that clinically manifests as an inability to respond appropriately to immunologic challenges. As CLL B cells are widespread in the host, the immunosuppression may be directly induced by organ infiltrative accumulation of clonal leukemic cells or indirectly via leukemic cell-induced remodeling of tissue microenvironments (Forconi and Moss, 2015; Goldman, 2000; Kipps et al., 2017). In addition to accumulation in blood and secondary lymphoid organs, CLL B cells infiltrate the bone marrow (BM), the primary site of adult hematopoiesis. This latter feature is thus a threat to the normal and critical function of BM.

Steady-state BM hematopoiesis sustains blood cell genesis throughout life (immune cells, erythrocytes, and platelets) (Doulatov et al., 2012) and is a tightly regulated process (Novershtern et al., 2011). Importantly, BM hematopoiesis becomes myeloid-skewed with aging (Pang et al., 2017), a critical consideration as CLL is largely a disease of the aged. Aberrant hematopoiesis and BM dysfunction in CLL is understudied and historically suggested to most likely be an infiltrative remodeling that disrupts normal hematopoietic niches (Lagneaux et al., 1993). Reduced numbers of BM hematopoietic stem cells (HSCs) and their progeny may manifest due to competition with CLL B cells for hematopoietic niches, local increases in CLL-derived factors, or CLL-induced alterations in resident immune cells or components of the BM microenvironment (Fecteau and Kipps, 2012; Sala et al., 1998).

Given the chronic residence of CLL cells in BM, their likely competition with hematopoietic progenitors for essential niches, and the increased potential for leukemic and HSPC interactions, we hypothesized that BM hematopoietic capacity may be compromised in CLL patients. Indeed, our previous report confirmed significantly reduced frequencies of BM HSCs and their progeny in CLL patients compared with age-matched controls (Manso et al., 2019). In addition, we demonstrated hematopoietic functional impairment as evidenced by significantly reduced colony forming unit (CFU) capacity, revealing diminished production of myelo-erythroid progenitors. In this initial work we also showed that select BM hematopoietic progenitor subsets displayed aberrant transcription factor (TF) profiles at the protein level. Importantly, the collective features of impaired BM hematopoiesis in CLL patients were independent of the extent of BM infiltration by

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(A-C) Flow cytometry was performed on freshly isolated BM from controls and CLL patients to determine MFI (representative of protein levels) of (A) PU.1, (B) GATA-2, and (C) GATA-1 among HSPC progenitor subsets. Control n = 11 for PU.1, n = 11 for GATA-2, and n = 8 for GATA-1. CLL n = 13 for PU.1, n = 12 for GATA-2, and n = 7 for GATA-1. The displayed number of data points for GATA-1 are reduced due to several individuals containing no detectable GATA-1 protein. We were unable to perform statistical comparisons between control and CLL LMPPs and GMPs for GATA-1 MFI as few cells contained detectable GATA-1 protein: HSC/MPP (Lin⁻CD34^{hi}CD38⁻CD45RA⁻), LMPP (Lin⁻CD34^{hi}CD38⁻CD45RA⁺), CMP/MEP (Lin⁻CD34⁺CD38⁺CD45RA⁻), and GMP (Lin⁻CD34⁺CD38⁺CD45RA⁺). Data are presented as the mean ± SEM; individual points represent a unique donor and are representative of 24 individual experiments. *p < 0.05, **p < 0.01, ****p < 0.0001, and ns by Mann-Whitney U test.

(D) Representative histograms of control and CLL patient BM progenitors. The stained sample is colored red and the isotype is shown in blue (see also Figure S1). (E–H) Combined intranuclear flow cytometry data from control (n = 8) and CLL patient (n = 6) HSPCs was utilized in the cytofkit2 R package to generate UMAP plots. Each point in the UMAP represents a single cell.

(E) Contour plots of the UMAP architecture for all combined samples or separated into control or CLL patients.

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Figure 1. Continued

(F) Colored overlay of control (teal) and CLL patient (red) HSPCs. The borders of five distinct regions are indicated and individually numbered. (G–H) Expression of PU.1, GATA-2, GATA-1, CD38, and CD45RA was superimposed on the cohort-specific architectures. Note that the data are normalized and expression values are relative to each individual factor. See also Figures S1–S4.

CLL B cells (Manso et al., 2019; Sala et al., 1998; Tsopra et al., 2009). Thus, the mechanism(s) responsible for impaired BM hematopoiesis in CLL are likely multifactorial and, at present, largely unexplored.

CLL B cells constitutively produce multiple factors capable of modulating BM hematopoiesis, most prominently tumor necrosis factor alpha (TNFα) (Foa et al., 1990; Michalevicz et al., 1991). TNFα is elevated in patient plasma, and levels associate with disease progression (Bojarska-Junak et al., 2008). The role of TNFα in steady-state hematopoiesis is largely suppressive to maintain BM homeostasis and aberrant exposure can result in BM failure, cytopenias, and anemia, clinical complications often observed in late-stage CLL patients (Broxmeyer et al., 1986; Geissler et al., 1991; Means et al., 1990; Michalevicz et al., 1991; Skobin et al., 2000; Tian et al., 2014; Tsopra et al., 2009). Limited studies have shown that TNF α , in part, can alter TF expression levels in mouse and human hematopoietic progenitors to mediate these modulatory actions (Etzrodt et al., 2019; Grigorakaki et al., 2011). Our previous report was the first to assess this mechanism in the context of CLL where we showed that in vitro exposure of control human BM progenitors (agedmatched to CLL patients) to TNFa upregulated PU.1 and GATA-2 proteins, phenocopying ex vivo findings in CLL patient marrow (Manso et al., 2019). However, questions remain concerning in vivo sources of TNFa in CLL, developmental stage-specific HSPC subset sensitivity to direct CLL B cell exposure or their secreted factors, mechanisms of TNFa modulation of HSPC TF expression, and reversibility of TF alterations upon effective TNF α neutralization. This report details our new findings in relation to these questions at both a global and single-cell level as assayed in primary human BM hematopoietic progenitors from control subjects and CLL patients.

Results

Global and single-cell analysis of control and CLL patient bone marrow progenitors

GATA-2, PU.1, and GATA-1 are cell fate determinants directly modulated by TNF α in BM progenitors (Etzrodt et al., 2019; Grigorakaki et al., 2011). GATA-2 is critical for HSC maintenance and survival and has additional roles in determining erythroid, megakaryocyte, and granulocyte cell fates (Vicente et al., 2012). PU.1 and GATA-1 are master regulators of myeloid and erythroid cell fates, respectively (Burda et al., 2010). Importantly, these factors are cross-antagonistic and the relative abundance of each reinforces bifurcating lineage decisions (Walsh et al., 2002). Given the importance of these TFs in regulation of primitive hematopoiesis, and their modulation by TNF α (Etzrodt et al., 2019; Grigorakaki et al., 2011), we examined global expression levels of these TFs in developmental stage-specific hematopoietic stem and progenitor cell (HSPC) populations: HSC/multipotent (HSC/MPP, Lin⁻CD34^{hi}CD38⁻CD45RA⁻), lymphoid-biased multipotential (LMPP, Lin⁻CD34^{hi}CD38⁻CD45RA⁺), common myeloid/megakaryocyte-erythroid (CMP/MEP, Lin⁻CD34⁺CD38⁺CD45RA⁻), and granulocyte-monocyte (GMP, Lin⁻CD34⁺CD38⁺CD45RA⁺) progenitors (Figure S1).

Freshly isolated HSC/MPPs from CLL patient BM exhibited significantly increased protein levels of PU.1 and GATA-2 when evaluated by flow cytometry and compared with age-matched controls (Figures 1A and 1B), consistent with and extending our previous findings that were largely performed in cryopreserved patient samples among simplified progenitor cell fractions (Manso et al., 2019). Here, our refined analysis further revealed that CLL patient-derived LMPPs and CMP/MEPs exhibited significantly higher levels of PU.1 and GATA-2. GATA-2 was also elevated among CLL GMPs. In contrast, an increased frequency of GATA-1⁺ cells was only observed in HSC/MPPs and CMP/MEPs (Figure S2), and those populations also display elevated levels of GATA-1 protein (among total GATA-1⁺ cells, Figure 1C). Representative histograms of PU.1, GATA-2, and GATA-1 (shown in red) from control and CLL HSPCs compared with isotype controls (shown in blue) are displayed in Figure 1D.

TF expression levels exist along a dynamic spectrum at the single-cell level, and as a function of cellular differentiation, a feature global analysis of total cell fractions does not wholly represent. Therefore, we employed uniform manifold approximation and projection (UMAP) (Leland McInnes, 2018) dimensionality reduction across the totality of the four HSPC populations from controls and CLL patients to evaluate



alterations in TFs and/or phenotypic markers at the single-cell level. We reasoned that, because UMAP normalizes expression of clustering markers across the combination of each input sample, this analysis would allow for the relative relatedness of individual cells to be analyzed. This first-of-a-kind analysis of control and CLL HSPCs enabled us to define global architectures for each cohort and also capture TF alterations at the resolution of single cells (Figures 1E and S3A). As expected, control HSPCs displayed significant heterogeneity in their UMAP occupancy, reflecting a diversity of phenotypes. When overlaid, CLL HSPCs were found, differentially represented, within three distinct control phenotypes (Figure 1F, circled regions 1, 2 and 4), yet largely enrich into region 4. A subset of control HSPCs also uniquely occupied a third region (region 3), which CLL HSPCs were strikingly absent from. Interestingly, region 5 represents a unique population of HSPCs found in CLL BM only.

Using the UMAP architecture, we next visualized the relative expression of PU.1, GATA-2, and GATA-1 (normalized to each specific parameter) across control and CLL HSPCs (Figure 1G). Control HSPCs exhibited considerable PU.1 heterogeneity and moderately uniform, yet variable, GATA-2 and GATA-1 expression. CLL HSPCs exhibited increased expression heterogeneity across all three TFs. When individual regions were evaluated, region 1 displayed variable PU.1 expression and high levels of GATA-2 and GATA-1. CLL HSPCs in region 1 were found to exhibit similar PU.1 and GATA-2 expression patterns as control HSPCs but have a distinct increase in GATA-1 positivity (Figures 1G and S4). Control HSPCs in region 2, a smaller region also occupied by CLL HSPCs, was characterized by low levels of PU.1, intermediate levels of GATA-2, and low levels of GATA-1. CLL HSPCs in region 2 show comparable PU.1 and GATA-2 expression patterns compared with controls, yet there is a small population of PU.1^{hi} cells. CLL HSPCs in this region also display increased GATA-1 expression. The unique control HSPC region 3 is composed of GATA-2^{hi} cells that also express the highest control HSPC levels of PU.1 and GATA-1. Region 4, a CLL-enriched group sharing marginal overlap with controls, displayed variable expression of GATA-2 and low levels of PU.1 and GATA-1 in control HSPCs. In contrast, CLL HSPCs in region 4 exhibit increased yet bimodal expression of PU.1 and GATA-2 and very low levels of GATA-1. Finally, the CLL-enriched region 5 is comprised of cells with low levels of all three TFs.

The cell surface markers CD38 and CD45RA are utilized to distinguish human HSPC subsets (Figures 1H and S4). When assessed by UMAP analysis, significant heterogeneity of CD38 and CD45RA expression was observed in control and CLL HSPCs. Regions 1 and 2 are enriched for cells with higher expression of CD38 and low/intermediate CD45RA. Control HSPCs in region 3 are characterized by intermediate levels of CD38 and increased, although variable, CD45RA expression. Both control and CLL HSPCs exhibit variable CD38 expression in the CLL-enriched region 4. Interestingly, control HSPCs in region 4 show bimodal CD45RA expression, whereas CLL HSPCs are largely CD45RA^{low}. In contrast, the CLL-specific region 5 displays a unique signature of CD38^{low} levels and uniformly high expression of CD45RA. Together, these data demonstrate that CLL BM HSPCs are distinct from controls and express fundamentally altered TF and cell surface marker profiles when assessed globally and at the single cell level.

CLL cells modulate TF protein levels in specific BM HSPC subsets

CLL B cells constitutively produce soluble and membrane-bound factors that may impact normal hematopoiesis (Burger et al., 2009; Foa et al., 1990; Janel et al., 2014; Kay et al., 2002; Lahat et al., 1991; Lotz et al., 1994; Saulep-Easton et al., 2016; van Attekum et al., 2017). Therefore, we sought to test if CLL cells modulate PU.1, GATA-2, and GATA-1 expression levels in HSPC subsets via cell contact and/or secreted/soluble factors by comparing direct and Transwell co-cultures consisting of primary CLL cells and age-matched control Lin⁻CD34⁺ HSPCs. To establish an experimental ratio of CLL B cells:control HSPCs, we assessed the cellular ratio of CD19⁺CD5⁻ and CD19⁺CD5⁺ B cells to Lin⁻CD34⁺ HSPCs in control and CLL BM (Figure S5). As expected, the ratio was orders of magnitude higher in CLL patients compared with controls. A 10:1 (CLL cells:HSPCs) ratio was used in all *in vitro* experiments for two reasons. First, it is reasonably greater than control CD19⁺CD5⁺ B cell to HSPC ratios. Second, it allows for evaluation of CLL B-cellinduced alterations in an environment mimicking lower thresholds of leukemic marrow infiltration as would be predicted in early stage disease.

Following 24 h of short-term co-culture allowing direct contact between control HSPCs and primary CLL B cells, the HSC/MPP, LMPP, CMP/MEP, and GMP populations were evaluated for levels of PU.1, GATA-2, and GATA-1 by flow cytometry. We note that the co-culture is serum free, eliminating any contribution of serum-derived factors in TF modulation. Short-term direct exposure to CLL cells significantly increased











Figure 2. Expression of PU.1, GATA-2, and GATA-1 are modulated *in vitro* by direct co-culture with CLL B cells (A–D') Freshly isolated control BM CD34⁺ cells were cultured in media alone, (A–D) directly with peripheral blood CLL cells, or (A'–D') indirectly by use of a 1.0 μ m Transwell insert at a ratio of 10:1 (CLL cells:HSPCs) for 24 h. Flow cytometry was performed to determine the MFI (indicative of relative protein levels) of PU.1, GATA-2, and GATA-1 among (A and A') Lin⁻CD34^{hi}CD38⁻CD45RA⁻ HSC/MPPs, (B and B') Lin⁻CD34^{hi}CD38⁻CD45RA⁺ LMPPs, (C and C') Lin⁻CD34⁺CD38⁺CD45RA⁻ CMP/MEPs, and (D and D') Lin⁻CD34⁺CD38⁺CD45RA⁺ GMPs. Each point represents a unique pairing of a control HSPC donor and CLL patient, with lines connecting the pairs (n = 11 for direct co-culture and n = 7 for Transwell co-cultures, 11 individual experiments). *p < 0.05, **p < 0.01, ****p < 0.0001, and ns by paired t test. (E) Summary table of the *in vitro* co-culture results. Changes to expression of each TF is indicated by \uparrow = increased, \downarrow = decreased, and — = no change. See also Figures S5 and S8.

PU.1 and GATA-2 in HSC/MPPs without affecting levels of GATA-1 compared with HSC/MPPs cultured in media alone (Figure 2A). Similarly, PU.1 was upregulated in the CMP/MEP and GMP (Figures 2C and 2D), but not LMPP, subsets (Figure 2B). In addition to HSC/MPPs, GATA-2 was also increased among CMP/ MEPs. Interestingly, GATA-2 displayed a variable response, with some control HSC/MPPs and CMP/ MEPs not affected by the co-culture (Figures 2A and 2C). GATA-1 remained unchanged in all populations. These new findings indicate that direct short-term co-culture of CLL B cells with control HSPCs differentially alters TF expression levels across multiple transitional stages of hematopoietic differentiation.

Next, we asked whether the TF alterations required direct contact between control HSPCs and CLL B cells or if modulation could be mediated by soluble factors provided by the CLL cells. To test this possibility, we physically separated the CLL B cells from control HSPCs by using Transwell inserts during the short-term coculture. In contrast to the results obtained in the direct co-cultures, HSPCs separated from CLL cells showed no significant change in PU.1 in any population examined (Figures 2A'–2D'). GATA-2 remained unchanged in HSC/MPPs and LMPPs but significantly increased in CMP/MEPs and GMPs, indicating differential TF sensitivity of individual progenitor stages to CLL-derived soluble factors (Figures 2C'–2D'). HSC/ MPPs and CMP/MEPs, but not LMPPs or GMPs, exhibited a decline in GATA-1 levels. Together, these findings suggest differential requirements for direct CLL:HSPC cell contact versus CLL-derived soluble factors in modulation of PU.1, GATA-2, and GATA-1 within developmental stage-specific HSPC subsets (summarized in Figure 2E).

CLL-derived TNFa directly modulates PU.1 and GATA-2 expression

In our previous study we reported that exogenous TNFa rapidly increased expression of PU.1 and GATA-2 in control HSPCs (Manso et al., 2019). We have extended these findings and determined that TNFa exposure reduces levels of GATA-1, consistent with previously published data (Buck et al., 2009; Grigorakaki et al., 2011) (Figure S6A). Further, NF- κ B p65 (ReIA), a TF at the nexus of inflammatory and TNF α signaling cascades (Liu et al., 2017; Nakagawa et al., 2018), is overexpressed in CLL-derived HSC/MPPs, unchanged in LMPPs and CMP/MEPs, and reduced in GMPs (Figure S7). We also note that the effects of TNFa on human HSPCs may be long-lasting or irreversible (Dybedal et al., 2001). TNF α is constitutively produced by CLL B cells in both membrane-bound and soluble forms (Foa et al., 1990; Michalevicz et al., 1991). Furthermore, TNFRI and TNFRII are expressed on primitive human HSPCs (Wang et al., 2017). To determine if $TNF\alpha$ derived from CLL B cells contributes to the alterations in PU.1, GATA-2, and GATA-1 in control HSPCs, a specific neutralizing antibody against TNFa (anti-TNFa) (Kim et al., 2018; Rothhammer et al., 2018) was added to the short-term direct and Transwell CLL:control HSPC co-cultures. Addition of anti-TNFa to the direct co-cultures significantly reduced levels of PU.1 in HSC/MPP, CMP/MEP, and GMP populations (Figures 3A, 3B, and 3D). GATA-2 was reduced in HSC/MPPs and CMP/MEPs but not GMPs. As PU.1 and GATA-2 were not modulated in LMPPs in the direct co-culture (Figure 2B), they remained unchanged when TNF α was neutralized (Figure 3B). Interestingly, GATA-1 protein levels were increased in HSC/MPPs and LMPPs upon neutralization of TNFa.

In contrast to the direct co-culture results, anti-TNF α did not significantly alter PU.1 levels in HSC/MPPs, LMPPs, or CMP/MEPs in Transwell cultures (Figure 3A'–C'), which was expected, as CLL-derived soluble factors alone did not alter PU.1 expression. However, a reduction in PU.1 was observed in GMPs (Figure 3D'). GATA-2 expression was reduced in HSC/MPPs and CMP/MEPs in Transwell cultures supplemented with anti-TNF α (Figure 3A' and 3C'). Neutralizing TNF α did not alter GATA-2 levels in LMPPs or GMPs (Figure 3B' and 3D'). This was expected of LMPPs, as they were refractory to CLL-derived soluble factors in the Transwell culture. The unchanging levels of GATA-2 in GMPs suggest a TNF α -independent











Figure 3. CLL B-cell-derived TNF α modulates transcription factor expression levels in vitro

(A-D') Freshly isolated control BM CD34⁺ cells were cultured in the presence or absence of a neutralizing antibody against TNF α (anti-TNF α) either (A–D) directly with peripheral blood CLL cells or (A'–D') indirectly cultured by use of a 1.0 μ m Transwell insert at a ratio of 10:1 (CLL cells:HSPCs) for 24 h. Flow cytometry was performed to determine the MFI (indicative of relative protein levels) of PU.1, GATA-2, and GATA-1 among (A, A') Lin⁻CD34^{hi}CD38⁻CD45RA⁻ HSC/MPPs, (B and B') Lin⁻CD34^{hi}CD38⁻CD45RA⁺ LMPPs, (C and C') Lin⁻CD34⁺CD38⁺CD45RA⁻ CMP/MEPs, and (D and D') Lin⁻CD34⁺CD38⁺CD45RA⁺ GMPs. Each point represents a unique pairing of a control HSPC donor and CLL patient, with lines connecting the pairs (n = 8 for direct co-culture and n = 7 for Transwell co-cultures, 8 individual experiments). *p < 0.05, **p < 0.01, and ns by paired t test.

(E) Summary table of the *in vitro* co-culture results. Changes to expression of each TF is indicated by \uparrow = increased, \downarrow = decreased, and — = no change.

See also Figures S5, S6, and S8.

effect for GATA-2 modulation in this population. GATA-1 protein remained unchanged upon neutralization of TNF α in all subsets except CMP/MEPs, where it was elevated (Figure 3C'). The effects of anti-TNF α *in vitro* are summarized in Figure 3E. Importantly, complete normalization of PU.1, GATA-2, and GATA-1 was observed when anti-TNF α was added to recombinant TNF α -exposed control progenitors, confirming effective TNF α neutralization in the co-culture model (Figure S6B).

We next compared media controls with direct CLL B cell:HSPC co-cultures that were subjected to TNF α neutralization to further investigate TNF α -specific modulation (Figure S8A). Compared with media controls, neutralization of TNF α normalized PU.1 in all populations except CMP/MEPs, where PU.1 remained slightly elevated. Anti-TNF α completely restored GATA-2 to control levels in all progenitor subsets except LMPPs, where it was slightly decreased. Notably, although direct co-culture did not change GATA-1 levels, neutralization of TNF α slightly elevated GATA-1 in CMP/MEPs.

When the same comparisons were made in Transwell co-cultures (Figure S8B), no changes were observed in PU.1. However, GATA-2 was slightly decreased in HSC/MPPs and restored in LMPPs, CMP/MEPs, and GMPs. GATA-1 also demonstrated differential responses. GATA-1 was reduced among HSC/MPP and CMP/MEP, but not LMPP and GMP, progenitors. Together, these data demonstrate that CLL B-cell-derived TNF α directly modulates aberrant expression of PU.1 and GATA-2 (and to a lesser extent GATA-1) in BM HSPCs. Further, given that complete TNF α neutralization does not restore every population to media control levels, other CLL-derived factors are likely contributing to TF modulation in BM progenitor cells.

Single-cell analysis of CLL:HSPC co-cultures reveal distinct mechanisms of TF modulation in HSPCs by CLL B cells

To investigate HSPC population changes in our co-culture model at the single-cell level, we performed UMAP analysis collectively of all *in vitro* experiments (Figures 4 and S3B). The general single-cell distribution of the media controls revealed segregation into three distinct regions that were largely maintained across all experimental conditions. However, unique variations to the overall UMAP architecture were induced by individual experimental manipulations. Next, we assessed the relative expression profiles of PU.1, GATA-2, and GATA-1 across the *in vitro* UMAP architecture for each experimental condition. Overall, the media controls exhibited high heterogeneity. The *in vitro* UMAP region 1 exhibited low levels of PU.1 and GATA-1 with intermediate levels of GATA-2. The second region displayed high levels of PU.1 and GATA-2, with lower, yet varying, levels of GATA-1. The third region was comprised of high levels of GATA-2 and GATA-1 with lower levels of PU.1. Addition of recombinant TNF α shifted control progenitors into regions with high levels of PU.1 and GATA-2 with variable, yet declining, GATA-1 expression. When the recombinant TNF α was neutralized, virtually identical UMAP architecture and TF patterning was observed when compared with the media controls.

Direct co-culture with CLL cells shifted control HSPCs to a unique signature of highly variable TF expression (Figure 4). Of note, the majority of cells in this culture condition expressed high levels of PU.1 and GATA-2 while maintaining a similar GATA-1 signature compared with media controls. Interestingly, the signature adopted when TNF α was neutralized in the CLL direct co-culture maintained regions of high PU.1 and GATA-2 expression, yet with a different cellular distribution among the UMAP architecture. The bimodal expression of GATA-1 was maintained. When CLL-derived soluble factors were evaluated in Transwell co-cultures, another unique signature was observed. Compared with media controls, relative levels of PU.1 were increased while GATA-2 and GATA-1 were maintained. Neutralization of TNF α in the Transwell











Figure 4. Single-cell UMAP analysis of HSPC expression of PU.1, GATA-2, and GATA-1 induced by CLL B cell direct or Transwell co-culture The relative expression of PU.1, GATA-2, and GATA-1 was overlaid on each individual media (n = 20), TNF α (n = 10), TNF α +anti-TNF α (n = 6), CLL direct coculture (n = 12), CLL + anti-TNF α direct co-culture (n = 9), CLL Transwell (TW) co-culture (n = 7), and CLL + anti-TNF α TW co-culture (n = 7) in vitro condition UMAP. The expression values and colors of each factor are normalized and relative to themselves. Each point in the UMAP represents a single cell and the total data are comprised of 20 individual experiments.

co-cultures largely restored control HSPC TF expression to that of the media control signature, although slight differences were observed, such as an overall reduction in PU.1 levels. Collectively, the UMAP analysis of the *in vitro* co-culture conditions revealed distinct alterations in the expression of PU.1, GATA-2, and GATA-1 at the single-cell level. Furthermore, direct CLL B cell:HSPC contact versus exposure to CLL soluble factors induced distinct TF signatures within specific HSPC populations and at the single-cell level.

Partial restoration of myelopoiesis upon removal of HSCs from the leukemic bone marrow microenvironment

We previously reported reduced frequencies and functional impairment of BM HPSCs analyzed immediately ex vivo from untreated CLL patients (Manso et al., 2019). We next sought to determine if the cellintrinsic functional defect documented in the CLL HSPCs analyzed ex vivo was reversible with prolonged removal from the leukemic microenvironment. To make this determination, we utilized the long-term culture initiating cell (LTC-IC) assay (Liu et al., 2013) to test the ability of control and CLL-derived HSCs to initiate and sustain myelopoiesis for 5 weeks across serial input cell concentrations (Figure 5A). As determined by frequencies of wells containing clonal progeny, we observed that CLL myeloid-biased HSCs generated significantly fewer clonal progeny, particularly early in the culture period. This finding is likely a result of maintaining biologic alterations incurred from their *in vivo* exposure to CLL B cell contact, secreted factors, or other features of a CLL-modified microenvironment. Interestingly, some individual CLL donor HSCs recovered functional capacity and were indistinguishable from controls by weeks 4–5 of culture, although the overall response remained diminished. Notably, this effect was not a function of altered CD34⁺ isolation purity between controls and CLL patients (Figure 5B). Importantly, when limiting dilution analysis (LDA) was performed (Hu and Smyth, 2009), a near 2-fold decrease in myelopoiesis-potentiating HSC precursor frequency was observed among CLL patients (Figures 5C and 5D).

Chronic exposure to TNFa reduces myeloid-biased HSC clonality

Given the impact of TNF¢ on progenitor TF profiles that regulate myelopoiesis, we next determined the contribution of TNF¢ to myeloid dysfunction. Confirming previous reports (Broxmeyer et al., 1986; Geissler et al., 1991; Skobin et al., 2000), addition of TNF¢ to CFU assays significantly reduced colony formation, which was restored when TNF¢ was neutralized (data not shown). Given these findings, we wished to determine if chronic exposure to TNF¢ would recapitulate the functional decrease of CLL-derived myeloid HSCs. As it is difficult to experimentally mimic the exact local *in vivo* TNF¢ concentrations in CLL patient BM HSPC niches, we performed a dilution series of TNF¢ in the LTC-IC assay with control BM HSPCs. As previously reported for CD34⁺CD38⁻ hematopoietic progenitors (Dybedal et al., 2001; Maguer-Satta et al., 2000; Petzer et al., 1996), chronic TNF¢ exposure of control HSPCs resulted in a dose-dependent inhibition of myelopoiesis (Figure 5E). Interestingly, chronic low-dose exposure effects were more prominent at early culture time points and resolved over time. Importantly, the TNF¢-induced reduction in clonal progeny recapitulate the diminished ex vivo CLL functional responses shown in Figure 5A. These data further reinforce a role for TNF¢ in diminishing myelopoiesis in CLL patient BM.

Discussion

Here, we report a mechanism for diminished BM myelopoiesis in CLL patients driven by leukemic B-cellderived TNF α . Combining *ex vivo* and *in vitro* experimental findings, we show that CLL B cells directly modulate PU.1 and GATA-2 expression in BM HSCs and myelo-erythroid progenitor subsets at the single-cell level. We pursued mechanisms of TF modulation and found that TNF α , produced by the CLL clone, plays an integral role. Importantly, short-term TNF α modulation of PU.1 and GATA-2 is reversible as neutralization of TNF α normalized changes in expression of PU.1 and GATA-2 induced by CLL cells. CLL is a chronic disease and BM infiltration is found even in early stage patients. We determined that removal of HSCs from the leukemic BM microenvironment restored myelopoietic function, although numbers of myeloid-biased HSCs were significantly reduced. Building on this observation, we show that chronic *in vitro* exposure of control age-matched HSPCs to a range of TNF α concentrations, including low dose and approximating levels reported to be produced by CLL B cells *in vitro*(Rosati et al., 2005), reduced

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Figure 5. CLL HSPCs are functionally compromised, and the dysfunction is recapitulated by exposure to $TNF\alpha$

(A) CD34⁺ HSPCs were isolated from control (green) and CLL (blue) BM and plated in the long-term culture initiating cell (LTC-IC) assay under limiting dilution conditions (12 or 36 replicates per dilution). Cultures were evaluated weekly for presence of clonal progeny (as evidenced by the occurrence of at least one colony). Data are presented as the mean \pm SD frequency of wells containing clonal progeny. Weeks 1–4 represent n = 4 for both controls and CLL, with week 5 data containing n = 8 controls and n = 9 CLL donors across 18 individual experiments. *p < 0.05, **p < 0.01, ***, p < 0.001, and ****p < 0.0001 by two-way ANOVA.

(B) When available, surplus isolated CD34⁺ cells (see Transparent methods) were tested by flow cytometry for purity (Control n = 6, CLL n = 4). ns by Mann-Whitney *U* test. Due to the reduced frequency of CD34⁺ cells in CLL BM (Manso et al., 2019) and limiting volumes of fresh BM, not every donor was able to be evaluated.

(C) Limiting dilution analysis (LDA) was performed at each time point for control and CLL LTC-IC assays to estimate numbers of responding HSCs among the input CD34⁺ cells. Circles indicate individual data points, with downward-facing triangles representing a negative (zero) response. Solid lines are the dilution model's fit, with dashed lines indicating the 95% confidence interval. Each time point is represented by a unique color as indicated.

(D) LDA estimates of 1/(stem cell frequency) for control and CLL HSPCs at each time point and fold difference in estimated precursor cell frequency. (E) LTC-IC assays were performed at limited cell dilutions (12 replicates each) of control BM HSPCs with media alone or supplemented with the indicated concentrations of TNF α . Cells were scored weekly as in (A). Data are presented as the mean \pm SD frequency of wells containing clonal progeny at each time point and dilution. n = 4 and is representative of three individual experiments. Statistical comparisons between each dilution at each time point are indicated in the accompanying data tables. *p < 0.05, **p < 0.01, ****p < 0.0001, and ns by two-way ANOVA adjusted for multiple comparisons (Tukey).

the numbers and function of myeloid-biased HSCs, largely mimicking CLL patient clinical data. The experimental findings detailed herein provide new insight into the basis of BM hematopoietic dysfunction in CLL patients that likely contribute to impaired immune status.

Our study highlights CLL B-cell-derived TNFα as a major driver of impaired BM myelopoiesis in CLL patients. We focused on TNFa, as it is constitutively produced by CLL B cells (Foa et al., 1990; Michalevicz et al., 1991), has a well-described role in modulating myelopoiesis (Buck et al., 2009; Dybedal et al., 2001; Grigorakaki et al., 2011; Tian et al., 2014), and is implicated in CLL disease progression (Bojarska-Junak et al., 2008). TNF α is produced in two trimeric forms, soluble and membrane-bound, each of which preferentially binds to either TNF receptor 1 or TNF receptor 2 (TNFRI and TNFRII), respectively (Grell et al., 1995). Signaling through both TNF receptors is required for HSC regulation (Grell et al., 1995; Pronk et al., 2011), a critical consideration in Transwell cultures, where membrane-bound TNFα-TNFRII interaction is inhibited, resulting in differential responses compared with direct co-culture. Although we cannot discount the complex combinatorial effects that multiple cell-cell contacts and/or soluble mediators between CLL B cells and HSPCs may have, the reversal of PU.1 and GATA-2 protein elevation by anti-TNF α strongly implicate TNFa in mediating, at least in part, CLL-induced impaired myelopoiesis. Further, when the in vitro cultures and CFU assays were performed with interleukin-10 (IL-10), another cytokine constitutively produced by CLL B cells that is elevated in patient serum (DiLillo et al., 2013; Fayad et al., 2001) and has HSC modulating activity (Kang et al., 2007), no alterations were observed, further demonstrating the specific activity of CLL-derived TNFa (Figure S9). Lastly, the alterations in myelopoiesis we document do not require remodeling of the BM microenvironment, as the changes in expression levels of PU.1 and GATA-2 could be reproduced in short-term in vitro co-cultures comprised of CLL B cells and normal HSPCs that were devoid of stromal and other niche supportive cells.

In CLL patient BM, the hematopoietic dysfunction we show is associated with elevated levels of PU.1, GATA-2, and GATA-1 simultaneously in specific HSPC subsets at the single-cell level. Signaling pathways activated by cytokines influence cell fate trajectories by modulating the activity and/or expression levels of lineage-determining TFs. Here, we focused on TNF α , as levels of this cytokine are elevated in CLL BM (Bojarska-Junak et al., 2008) and CLL-related anemia has been shown to be strongly linked to $TNF\alpha$ (Tsopra et al., 2009). Our in vitro direct CLL B cell:control HSPC co-culture model showed increased PU.1 expression in primitive HSC/MPPs via CLL-derived TNFa. GATA-2 represses the GATA-2/GATA-1 switch critical for erythroid commitment (Bresnick et al., 2012; Moriguchi and Yamamoto, 2014) and was also elevated in progenitors exposed to CLL-derived TNFa. It is well established that overexpression of PU.1 or GATA-2 is disruptive for proper erythroid lineage commitment (Grigorakaki et al., 2011). GATA-2 can also directly antagonize PU.1 (Walsh et al., 2002; Zhang et al., 2000), suppressing myelopoiesis. We show, for the first time, that direct contact between CLL cells and HSPCs is likely required in vivo to increase HSPC expression levels of PU.1 via TNF α , as the PU.1 levels were normalized upon TNF α neutralization. Similarly, we found that CLL B cell direct contact and, to a certain extent, CLL B cell soluble factors (including $TNF\alpha$), can upregulate GATA-2. Collectively, simultaneously increased levels of PU.1, GATA-2, and GATA-1 may antagonize erythroid and myeloid cell fate programs (Arinobu et al., 2007; Bresnick et al., 2012; Burda et al., 2010; Moriguchi and Yamamoto, 2014; Walsh et al., 2002; Wolff and Humeniuk, 2013), subverting lineage





commitment and differentiation, resulting in diminished mature blood cell output from the BM resulting in cytopenias and diminished immune competence.

To further understand the biology of BM hematopoiesis in CLL we also examined ex vivo phenotypic changes among control and CLL HSPCs at the single-cell level using UMAP dimensionality reduction. Distinct global architectures were observed that demonstrated that CLL-derived BM HSPCs are distinct from healthy controls and express fundamentally altered TF and cell surface marker profiles. Intriguingly, the CLL-enriched UMAP region 4 displayed high levels of CD38 with intermediate and variable CD45RA expression, phenotypically overlapping with CMP/MEPs. CLL HSPCs in region 4 also express higher levels of PU.1 and GATA-2 globally and at the single-cell level. We hypothesize that the phenotypic CMP/MEPs may be unable to resolve developmental transitions due to cross-antagonism between TFs. Conversely, the CLL-specific UMAP region 5 displayed a unique expression pattern of low CD38 and high CD45RA, phenotypically representative of LMPPs. Interestingly, this population displays low levels of PU.1, GATA-2, and GATA-1. Given that region 5 is notably enriched in our untreated CLL cohort, and does not express TFs indicative of myeloid or erythroid fates, it is tempting to speculate that this region is enriched for cells that are a CLL stem cell, a CLL-modified LMPP population, and/or a population protected from CLL-induced cellular reductions and is therefore overrepresented.

This study provides, for the first time, a unique and targetable mechanism by which CLL cells negatively regulate BM myelopoiesis. Investigation of mechanisms set a framework for designing interventions that could ameliorate clinically important cytopenias that strongly impact immune competency and clinical outcomes. Importantly, our study challenges the existing paradigm of peripheral immunosuppression as the only mechanism inducing immune cell dysfunction in CLL patients. Together, these new findings explain, in part, the etiology of cytopenias in CLL, particularly of the myelo-erythroid lineage (Manso et al., 2019; Zent et al., 2008). Advances in understanding BM hematopoietic dysfunction in CLL patients offers targetable mechanism(s) for the treatment and reversal of CLL-induced cytopenia and accompanying immunosuppression.

Limitations of the study

A caveat of this study is that it relied on the utilization of freshly prepared, untreated CLL samples and each individual CLL patient sample to be co-cultured with unique control donor HSPCs collected on the same day. This unavoidable constraint imposed by the availability of human experimental research material impaired our ability to further stratify results by clinically relevant CLL patient features (Table S1). We did not measure soluble TNF α levels in the co-cultures. TNF α levels in BM plasma are significantly higher than blood plasma and correlate with disease stage and lymphocytosis (Bojarska-Junak et al., 2008). Regardless, it is very difficult to know precise physiologic levels of secreted or membrane-bound TNFa in the in vivo BM HSPC niche and thus the relevance to in vivo or in vitro levels cannot be determined. Multiple studies have reported that CLL B cells constitutively produce TNF α in vitro in the 20pg/mL range that can be effectively neutralized by an anti-TNF antibody (Aue et al., 2011; Djurdjevic et al., 2009; Rosati et al., 2005). This is important as the levels of TNFa production in vitro are strikingly consistent with reported serum levels of TNFa in patients, which was validated in an independent study with a different cohort of CLL patients (Ferrajoli et al., 2002). An interesting, but potentially relevant, observation by another group was that cellular release of TNF α by primary CLL B cells was higher in early stage (Rai 0–1) compared with advanced stage (Rai II-III) patients, and the majority of CLL B cells used in our in vitro co-cultures were from early stage patients (Foa et al., 1990) (Table S1). Regardless, TNFα inhibits human CFU assays at a range of 1–10ng/mL (Skobin et al., 2000). We show that as little as 0.8µg of anti-TNF antibody mediated complete neutralization of the 25ng/mL exogenous TNF α added to our co-cultures (Figure S6), providing confidence that we are achieving complete TNFa neutralization of either recombinant or CLL-derived TNFa. Importantly, our heterogeneous CLL patient and donor input samples gave a striking uniformity of results, suggesting that the modulation of HSPCs is a shared feature of CLL cells. We also note that TNF signaling promotes CLL B cell survival and that inhibition of that pathway could result in increased susceptibility to apoptosis (Cordingley et al., 1988; Digel et al., 1989; Durr et al., 2018). However, neutralization of TNFα in our short-term co-culture system did not change the viability of any cell fraction studied (CLL cells or HSPCs).

Resource availability

Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Kay Medina (medina.kay@mayo.edu).

CellPress



Materials availability

This study did not generate new unique reagents.

Data and code availability

This study did not generate or analyze datasets. Further, it did not produce any original code. The code for the R packages used in this manuscript can be found within the references provided.

Methods

All methods can be found in the accompanying Transparent methods supplemental file.

Supplemental information

Supplemental information can be found online at https://doi.org/10.1016/j.isci.2020.101994.

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Author contributions

B.A.M., K.L.M., and N.E.K. wrote the manuscript. B.A.M., K.A.G., P.K.L., and B.M.W. performed and analyzed all experiments. J.E.K. provided expertise with the R program and assisted with UMAP generation. All authors reviewed and contributed to manuscript revisions.

Declaration of interests

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Supplemental Information

Chronic lymphocytic leukemia

B-cell-derived TNFα impairs

bone marrow myelopoiesis

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Figure S1. Flow cytometry gating strategy for *ex vivo* **and** *in vitro* **HSPCs and transcription factors.** Related to Figures 1-3. The sequential gating strategy used for analysis is illustrated using representative control and CLL *ex vivo* bone marrow samples. Following gating, each population was assessed for expression of PU.1, GATA-2, and GATA-1 as shown.



Figure S2. Frequency of GATA-1⁺ cells among HSPCs from control and CLL patient bone marrow. Related to Figure 1. Freshly isolated BM was evaluated by intranuclear flow cytometry for the frequency of GATA-1 positive cells among HSC/MPP, LMPP, CMP/MEP, and GMP populations. Each point represents individual donors and Data is presented as mean +/- SEM. Control n=8 and CLL n=7. ***P<0.001 and ns by Mann-Whitney U test.

. ex vivo UMAP input (cytofkit2 R package)



Figure S3. Experimental design and input for *ex vivo* **and** *in vitro* **UMAP analysis.** Related to Figures 1 and 4. (**A**) CD34^{hi}CD38⁻ and CD34⁺CD38⁺ populations among the Lin^{low/-} compartment were concatenated in FlowJo and used as the raw data input for the *ex vivo* UMAP analysis. Data from the *ex vivo* flow cytometry analysis was exported from FlowJo and imported directly into the cytofkit2 R package with the settings and pipeline outlined. (**B**) Lin⁻CD34⁺ cells were used as the raw data input for the *in vitro* UMAP analysis. Each individual *in vitro* data file was pre-processed in FlowJo by using the DownSample plugin to obtain files of 1000 Lin⁻CD34⁺ cells. All data files from each individual *in vitro* condition was then concatenated and DownSampled to 6000 total cells prior to inputting into R using the cytofkit2 package and pipeline outlined. All UMAP graphical plots and raw data was generated and the raw UMAP data was further graphed using the ggplot2 R package or inputting the analyzed data back into FlowJo. Each UMAP figure is indicative of the combination (or specific subsets) of the total *ex vivo* or *in vitro* data.

Α



Figure S4. *ex vivo* **UMAP** marker expression patterns. Related to Figure 1. The analyzed *ex vivo* UMAP data from the cytofkit2 R package was imported back into FlowJo for further analysis. Each of the five individual regions were overlaid for the expression of the clustering markers. Each histogram represents the normalized expression data for each marker as calculated during the UMAP analysis. The histogram overlays are displayed as either the total UMAP (controls and CLL) or cells from the control or CLL cohorts only.



Figure S5. Determination of the *ex vivo* **bone marrow B cell:HSPC BM ratio.** Related to Figures 2-4. Freshly isolated control (n=32) or CLL (n=26) BM was analyzed by flow cytometry to determine the frequency of CD19⁺CD5⁺ and CD19⁺CD5⁻ B cells and Lin⁻CD34⁺ HSPCs among total BM cells. This data was generated from our previously published data set(Manso et al., 2019). The ratio for each individual sample (represented by a single point) was determined by dividing the frequency of either CD19⁺CD5⁺ or CD19⁺CD5⁻ B cells by the frequency Lin⁻CD34⁺ HSPCs. Data is presented as the mean +/- SEM.



Figure S6. Effect of TNF α **on transcription factor expression in HSPCs.** Related to Figures 3 and 4. (A) Freshly isolated control Lin⁻CD34⁺ HSPCs were exposed to TNF α as described in the Transparent Methods (n=10 or 6). Following 24 hours of culture, intranuclear flow cytometry was performed to determine the expression of GATA-1. ***P*<0.01 and ns by paired t-test. (B) Additional cultures were set up as in (A) and received a neutralizing antibody against TNF α (anti-TNF α) at the start of the experiment (n=6). Following 24 hours of culture, intranuclear flow cytometry was performed to determine the expression of PU.1, GATA-2, and GATA-1. Each point represents an individual donor with lines connecting the same donor across different experimental conditions. **P*<0.05, ***P*<0.01, ****P*<0.001, and ns by repeated measures one-way ANOVA adjusted for multiple comparisons.



Figure S7. *ex vivo* analysis of NF- κ B. Related to Figure 1. Freshly prepared BM from control and CLL patients were evaluated by intranuclear flow cytometry for expression of NF- κ B p65 among HSC/MPPs, LMPP, CMP/MEPs, and GMPs. Each point represents individual donors and Data is presented as mean +/- SEM. Control n=7 and CLL n=7. **P*<0.05, ***P*<0.01, and ns by Mann-Whitney *U* test.



Figure S8. Neutralization of TNF α normalizes transcription factor expression in CLL:HSPC cocultures. Related to Figures 2-4. The HSC/MPP, LMPP, CMP/MEP, and GMP subsets were evaluated by flow cytometry for relative levels of PU.1, GATA-2, and GATA-1 protein levels (by MFI) between media controls and CLL+anti-TNF α cultures. All CLL co-cultures at a ratio of 10:1 (CLL cells:HSPCs). Evaluation of direct (**A**) and 1.0µm Transwell (**B**) co-cultures was performed. Each point and connecting line represents a unique pairing of a control HSPC donor and CLL patient (n=11 for direct co-culture and n=7 for TW co-cultures, 11 individual experiments). **P*<0.05, ***P*<0.01, and ns by paired t-test.



Figure S9. IL-10 does not alter levels of PU.1, GATA-2, or GATA-1 in HSPCs or change colony forming unit (CFU) capacity. Related to Figures 2-4. (**A**) Freshly isolated control CD34⁺ HSPCs from BM were exposed *in vitro* to IL-10 for 24 hours. Following incubation, the HSPCs were analyzed by intranuclear flow cytometry for expression of PU.1, GATA-2, and GATA-1 as described in the Transparent Methods. Each point and connected line represents a unique donor (n=6). Some culture conditions did not contain detectable GATA-1 protein among GMPs and therefore are not represented in the graph. (**B**) Freshly isolated CD34⁺ HSPCs from control BM were plated in triplicate CFU assays. CFU assays were supplemented at assay initiation with 25 ng/mL IL-10 (n=5). Each point and connected line represents a unique donor. **P*<0.05 and ns by paired t-test.

												Experimental use of samples:			es:
CLL ID	Age	Sex	Rai Stage	Cytogenetics	ZAP- 70	CD38	CD49d	B2M	IGHV	Leukemic marrow involvement (%) – hematopathology	Sample tissue type	<i>ex viv</i> o TFs	<i>ex vivo</i> UMAP	<i>in vitro</i> co- culture	LTC-IC assay
CLL-1	66	М	0	Trisomy 12	+	-	+	+	UM	>90	BM	Х			
CLL-2	68	F	I	13q-	-	-	-	+	М	10	BM	Х			Х
CLL-3	80	М	II	Trisomy 12	+	-	+	+	UM	90	BM	Х			Х
CLL-4	68	М	I	11q-	-	-	n.d.	+	UM	90	BM	Х			Х
CLL-5	58	М	I	13q-	+	-	-	+	UM	90	BM	Х			Х
CLL-6	58	F	III	Trisomy 12	+	-	n.d.	+	UM	80	BM	Х			Х
CLL-7	75	F	0	13q-	-	-	n.d.	+	UM	80	BM	Х	Х		
CLL-8	79	F	П	Trisomy 12	-	-	+	+	М	30	BM	Х	Х		
CLL-9	40	М	П	Trisomy 12	n.d.	+	n.d.	+	М	n.d.	BM	Х	Х		
CLL-10	63	F	III	13q-	+	-	+	+	UM	90	BM	Х	Х		
CLL-11	58	М	II	Trisomy 12	-	+	+	+	М	90	BM	Х	Х		
CLL-12	63	М	IV	13q-	+	+	+	+	UM	95	BM	Х	Х		Х
CLL-13	53	М	0	13q-	+	-	-	+	UM	n.d.	BM	Х	Х		Х
CLL-14	71	М	IV	Normal	-	-	+	+	М	<1	BM				Х
CLL-15	56	М	0	17p-	+	+	-	+	UM	80	BM				Х
CLL-16	41	М	I	Normal	+	-	+	+	М	20	BM	Х	Х		
CLL-17	74	М	0	13q-	-	-	-	-	М	n.d.	Blood			Х	
CLL-18	67	М	0	Normal	-	-	-	-	М	n.d.	Blood			Х	
CLL-19	69	F	III	13q-	-	-	-	+	М	90	Blood			Х	
CLL-20	67	М	0	13q-	-	-		+	М	n.d.	Blood			Х	
CLL-21	55	F	I	13q-	-	-	-	+	М	n.d.	Blood			Х	
CLL-22	44	М	I	NA	-	-	-	+	UM	n.d.	Blood			Х	
CLL-23	80	М	0	n.d.	-	-	-	+	М	n.d.	Blood			Х	
CLL-24	71	М	0	Trisomy 12	+	-	+	+	UM	80	Blood			Х	
CLL-25	92	F	0	13q-	-	-	-	+	М	n.d.	Blood			Х	
CLL-26	80	М	0	Normal	-	-	-	+	М	n.d.	Blood			Х	
CLL-27	82	М	0	Trisomy 12	-	+	-	+	UM	n.d.	Blood			Х	

 Table S1. CLL patient characteristics.
 Related to Figures 1-5.
 Demographics for each CLL patient are listed and the tissue source and list of experiments the sample was used for is indicated.
 B2M=beta 2 microglobulin, IGHV=immunoglobulin heavy chain variable region, M=mutated, UM=unmutated, +=positive, -=negative, BM = bone marrow, TF = transcription factor, and n.d.=no data.

			Catalogue								
Reagent or antibody specificity	Fluorochrome	Vendor	Number	Clone							
Sample preparation reagents											
1X phosphate buffered saline (PBS) prepared from 10X stock	N/A	Gibco	14200-075	N/A							
Liveran Ea black	N1/A	Miller i Dietee	120.050.001	N1/A							
Human FC block	IN/A		130-059-901	IN/A	ł						
1% percentermoldobyde (PEA) properted from 10% stock	NI/A	Electron Microscopy	15710 8	N1/A							
1% paraiornaidenyde (PFA) prepared from 10% stock	IN/A	Sciences	15/12-5	IN/A	1						
	NI/A	CollGonix	20802	NI/A	1						
Person binant human stem cell factor	N/A	BoproToch	20002	N/A	ł						
	N/A	PeproTech	200-06	N/A	ł						
Recombinant human IL-3	N/A	PeproTech	200-03		ł						
Recombinant human Enthropoietin	N/A	PeproTech	100-64	N/A	ł						
Recombinant human TNFg	N/A	PeproTech	300-01A	N/A	ł						
anti-buman TNFa	N/A	R&D Systems	AF-210-NA	N/A	1						
Recombinant human II -10	Ν/Δ	Biol egend	573202	NI/A	1						
1.0 um TransWell insert/plate	N/A	Corning	CL \$3380	N/A	ł						
MethoCult H1/135 media	Ν/Δ	StemCell Technologies	04435	NI/A	ł						
MyeloCult H5100 media	N/A	StemCell Technologies	05150	N/A	1						
Hydrocortisope	N/A	StemCell Technologies	07904	N/A	1						
Cell S	Surface Markers	oternoen reennologies	07504	19/73	1						
Fixable Viability Stain 510	V510	BD Biosciences	564406	N/A							
CD3	PerCP	BD Biosciences	347344	Sk7							
CD5	PerCP	BioLegend	300618	UCHT2	⊕ ∕si –						
CD11b	PerCP	BioLegend	101230	M1/70	age						
CD14	PerCP	BD Biosciences	340660	M ₀ P9	ine						
CD19	PerCP	BD Biosciences	347544	4G7	ant						
CD56	PerCP	BioLegend	318342	HCD56							
CD34	PE-Cy7	BD Biosciences	348791	8G12							
CD38	V450	BD Biosciences	646851	HB7							
CD45RA	APC/Fire 750	BioLegend	304152	HI100							
CD19	APC/Fire 750	BioLegend	363030	SJ25C1							
CD5	PerCP-Cy5.5	BD Biosciences	341089	L17F12							
Intranuclear staining with the True-Nuclear staining buffer kit											
True-Nuclear staining buffer kit	N/A	BioLegend	424401	N/A							
PU.1	Alexa Fluor 647	BioLegend	658004	7C6B05							
Mouse IgG1, κ isotype control (for PU.1)	Alexa Fluor 647	BioLegend	400136	MOPC-21							
GATA-2	FITC	R&D Systems	IC2046F	N/A							
Goat IgG isotype control (for GATA-2)	FITC	R&D Systems	IC108F	N/A							
GATA-1	PE	Cell Signaling Technologies	13353S	D52H6							
NF-kB p65	PE	Cell Signaling Technologies	9609S	D14E12	1						
Rabbit IgG isotype control (for GATA-1 and NF-κB p65)	PE	Cell Signaling Technologies	5742S	DA1E							

Table S2. List of reagents used. Related to Figures 1-5.

Transparent Methods

Experimental model and subject details Human subjects

These studies are restricted to untreated CLL patients to eliminate confounding factors introduced by treatment. Fresh BM and/or blood samples were obtained from consenting CLL patients (Mayo Clinic IRB 1827-00, Table S1) and BM from age-matched controls either undergoing hip replacement surgery or recruited (Mayo Clinic IRBs 1062-00 and 16-006204) who signed informed consent per the Declaration of Helsinki and Mayo Clinic guidelines. BM samples were subjected to mature erythrocyte depletion by ACK lysis then used immediately. CLL blood samples underwent Ficoll separation immediately before use.

Cell lines

The murine stromal cell line M2-10B4 was obtained from ATCC and maintained in RPMI 1640 (Gibco) media supplemented with 10% fetal bovine serum (FBS, Mediatech), 1% penicillin/streptomycin (Gibco), and 1% L-glutamine (Gibco). The M2-10B4 cells were cultured in an incubator at 37°C and 5% CO₂.

Method details

Flow cytometry

Following isolation, cells were Fc blocked (Miltenyi Biotec, San Diego, CA, USA). Surface staining with monoclonal antibodies and fixable viability dye was performed on ice in 1X PBS for 30 minutes. Following washing, the cells were fixed and permeabilized for 60 minutes according to manufacturer's directions (BioLegend, San Diego, CA, USA) to allow for intranuclear staining. Cells were then stained for 30 minutes at room temperature with monoclonal antibodies against nuclear targets or appropriate isotype controls. Following staining, cells were washed and suspended in 1% paraformaldehyde (Electron Microscopy Sciences, Hatfield, PA, USA) for acquisition. When cell surface staining only was performed, cells were washed twice following staining and fixed with 1% paraformaldehyde for acquisition. Flow cytometry data was collected on a FACS-Canto (Becton Dickinson, Franklin Lakes, NJ, USA) that is standardized daily to allow for direct comparisons of mean fluorescent intensity (MFI) across separate experiments(Manso et al., 2019; Perfetto et al., 2006; Perfetto et al., 2012).

Flow cytometry analysis

For analysis of PU.1, GATA-2, and NF- κ B p65 relative protein levels, the MFI of the stained sample was subtracted against the appropriate isotype control as expression is unimodal (Figure 1D and Figure S1). The isotype-subtracted MFI is the value used in the figures. GATA-1 staining is bi-modal (Figure 1D and Figure S1). Therefore, relative protein levels of GATA-1 was determined by first gating on GATA-1 positive cells (determined by the isotype control), then calculating the MFI of GATA-1 among only the cells that are GATA-1⁺. This process results in no isotype control staining in the GATA-1 histogram analysis by virtue of the initial GATA-1⁺ determination. Flow cytometry analysis was performed with FlowJo 10.5.3 (Becton Dickinson).

In vitro cell cultures

CD34⁺ cells from freshly processed control BM were isolated by positive magnetic bead selection (Miltenyi Biotec). CLL cells were plated at 1.0x10⁶ cells per well with 1.0x10⁵ CD34⁺ cells (10:1 ratio) in a 96 well round-bottom plate. The CLL:CD34⁺ co-cultures were incubated undisturbed for 24 hours at 37°C 5%CO₂ in serum-free media (CellGenix, Portsmouth, NH, USA) supplemented with human recombinant stem-cell factor (10ng/mL), IL-6 (10ng/mL), IL-3 (1ng/mL), and erythropoietin (1ng/mL) (all from PeproTech, Rocky Hill, NJ, USA). Transwell (Corning, Corning, NY, USA) assays utilized a 1.0µm insert. CLL cells were plated in the Transwell with CD34⁺ cells in the lower chamber. Where indicated, 0.8µg of anti-TNF α antibody (R&D Systems, Minneapolis, MN, USA) was added to the cultures (in the lower chamber for Transwell experiments) at assay initiation. Recombinant human TNF α (R&D Systems) or IL-10 (BioLegend), when utilized, was added at 25ng/mL(Asano et al., 1999; Grigorakaki et al., 2011; Xiao et al., 2002). In all cases, cells were stained for flow cytometry as outlined above to allow for detection of surface and intranuclear proteins.

Uniform manifold approximation and projection (UMAP)

The R program (version 3.6.1)(R Core Team, 2019) and the cytofkit2 package (Jinmiao Chen's Lab, version 2.0.1)(Becher et al., 2014; Chen et al., 2016; Wong et al., 2015) were used to generate UMAP(Leland McInnes, 2018) data and graphics plots from *ex vivo* and *in vitro* flow cytometry data. The single/live/Lin⁻ CD34^{hi}CD38⁻ and CD34⁺CD38⁺ populations from each individual *ex vivo* sample was concatenated in FlowJo and exported whereas each *in vitro* flow cytometry data file was gated and exported on single/live/Lin⁻/CD34⁺ cells. The *ex vivo* data was imported into cytofkit2 with the parameters outlined in Figure S3A. Due to increased numbers of *in vitro* conditions/samples, additional data preprocessing was performed (Figure S3B). Some UMAP plots underwent additional graphical modifications with the ggplot2 package to allow for color selection and identical axis scaling across plots(Wickham, 2016).

Long-term culture initiating cell (LTC-IC) assay

Fresh BM CD34⁺ cells from controls and CLL patients were isolated as above and plated across 12 or 36 replicates in LTC-IC medium (H5100 MyeloCult with 10⁻⁶M hydrocortisone, StemCell Technologies, Vancouver, Canada). Cells were plated at limiting dilution (1000, 100, 50, 25, 10, 5, and 1 cell(s) per well) on irradiated feeder M2-10B4 stromal cells (12,500 cells per well) in 96 well flat-bottom plates. The cultures were maintained for five weeks at 37°C 5%CO₂ with weekly half-media exchange and scoring for the presence or absence of one or more colonies per well. In experiments containing TNF α , a limited cell dilution range was utilized (1000, 100, 50, and 10) with 12 replicates per dilution. Fresh media with the indicated concentration of TNF α was added weekly to the cultures to mimic *in vivo* chronic exposure.

Limiting dilution analysis (LDA)

Results from the LTC-IC assays containing control or CLL CD34⁺ cells were analyzed by LDA (<u>http://bioinf.wehi.edu.au/software/elda/</u>)(Hu and Smyth, 2009). The confidence interval was set at 0.95, with options for testing inequality in frequency between multiple groups and adequacy of the single-hit model selected as analysis parameters.

Colony forming unit (CFU) assays

Following isolation of CD34⁺ cells from freshly processed control BM, CFU assays were performed by plating triplicates of 500 CD34⁺ cells in semisolid MethoCult H4435 media (StemCell Technologies). This media formulation allows for differentiation and enumeration of CFU-granulocyte, monocyte (CFU-GM), CFU-granulocyte, erythrocyte, monocyte, megakaryocyte (CFU-GEMM), and CFU-erythroid (CFU-E) colonies by morphology. When utilized, recombinant human TNF α was added at 25 ng/mL(Asano et al., 1999; Grigorakaki et al., 2011; Xiao et al., 2002). Neutralization of TNF α was achieved by addition of 0.8 μ g of an anti-TNF α antibody (R&D Systems). The CFU cultures were incubated at 37°C 5% CO₂ for 11-12 days then manually scored, counted, and averaged across triplicates.

Quantification and statistical analysis

Statistical tests are indicated in figure legends and performed using GraphPad Prism version 8.1.2 (GraphPad Software, San Diego, CA, USA). For all tests, **P*<0.05, ***P*<0.01, ****P*<0.001, *****P*<0.0001, and "ns" indicates non-statistical significance. Note that, as some *ex vivo* and *in vitro* conditions had no detectable GATA-1, some paired t-tests are missing data for one or multiple conditions. When statistical analysis was performed, only values with data in each column were used for statistical comparison, as is standard practice. A power calculation was not performed owing to limited sample availability. Bar graphs of summary statistics indicate the mean ± standard error of the mean (SEM). The number of individual experiments and samples are outlined in the figure legends.

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