Targeting CD38 is lethal to Breg-like chronic lymphocytic leukemia cells and Tregs, but restores CD8+ T-cell responses

Alak Manna,1 Timothy Kellett,1 Sonikpreet Aulakh,2 Laura J. Lewis-Tuffin,1 Navnita Dutta,3 Keith Knutson,3 Eduardo Chini,4 Javier Pinilla-Ibarz,6 Nicole Lamanna,9 Rami Manochakian,9 Fabio Malavasi,7 Taimur Sher,9 Asher A. Chanan-Khan,1,2,8 Sikander Ailawadhi,2 and Aneel Paulus1,2
1Department of Cancer Biology, Mayo Clinic, Jacksonville, FL; 2Division of Hematology and Oncology, Mayo Clinic, Jacksonville, FL; 3Department of Immunology, Mayo Clinic, Jacksonville, FL; 4Signal Transduction Laboratory, Kogod Aging Center, Department of Anesthesiology, Mayo Clinic, Rochester, MN; 5Department of Malignant Hematology, Moffitt Cancer Center, Tampa, FL; 6Columbia University Medical Center, New York, NY; 7Laboratory of Immunogenetics, Department of Medical Science, University of Turin, Turin, Italy; and 8Mayo Clinic Cancer Center at St. Vincent’s Medical Center Riverside, Jacksonville, FL

Key Points
• CD38hi Breg-like CLL cells promote conversion of naive T cells into Tregs in an IL-10/TGF-β-dependent manner.
• CD38 targeting drugs are lethal to both IL-10–producing Breg-like CLL cells and Tregs, but lead to improved CD8+ T-cell cytolytic responses.

Patients with chronic lymphocytic leukemia (CLL) are characterized by monoclonal expansion of CD5−CD23−CD27−CD19+/k− B lymphocytes and are clinically noted to have profound immune suppression. In these patients, it has been recently shown that a subset of B cells possesses regulatory functions and secretes high levels of interleukin 10 (IL-10). Our investigation identified that CLL cells with a CD19−CD24+CD38hi immunophenotype (B regulatory cell [Breg]–like CLL cells) produce high amounts of IL-10 and transforming growth factor β (TGF-β) and are capable of transforming naive T helper cells into CD4+ CD25+FoxP3+ T regulatory cells (Tregs) in an IL-10/TGF-β-dependent manner. A strong correlation between the percentage of CD38hi CLL cells and Tregs was observed. CD38hi Tregs comprised more than 50% of Tregs in peripheral blood mononuclear cells (PBMCs) in patients with CLL. Anti-CD38 targeting agents resulted in lethality of both Breg-like CLL and Treg cells via apoptosis. Ex vivo, use of anti-CD38 monoclonal antibody (mAb) therapy was associated with a reduction in IL-10 and CLL patient-derived Tregs, but an increase in interferon-γ and proliferation of cytotoxic CD8+ T cells with an activated phenotype, which showed an improved ability to lyse patient-autologous CLL cells. Finally, effects of anti-CD38 mAb therapy were validated in a CLL–patient-derived xenograft model in vivo, which showed decreased percentage of Bregs, Tregs, and PD1+CD38hiCD8+ T cells, but increased Th17 and CD8+ T cells (vs vehicle). Altogether, our results demonstrate that targeting CD38 in CLL can modulate the tumor microenvironment; skewing T-cell populations from an immunosuppressive to immune-reactive milieu, thus promoting immune reconstitution for enhanced anti-CLL response.

Introduction

Chronic lymphocytic leukemia (CLL) is a B-cell cancer in which there is a concomitant dysregulation of the nonmalignant T-cell compartment and immune cytokine milieu.1-5 T cells from patients with CLL are often impaired, with a notable increase in T regulatory cells (Tregs) and compromised CD8+ cytotoxic T-cell (CTL) functionality.6,7 These cellular deviations are accompanied with dysregulation of immunosuppressive cytokines such as interleukin 10 (IL-10) and transforming growth factor β (TGF-β) secretion.2,8 Together, these changes contribute to clinical progression of disease.9

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Abnormalities in the T-cell population are present at early stages of disease in patients with CLL, suggesting the ability of malignant B-CLL clones (even in low numbers) to exert a dominant effect on their microenvironment. It has recently been shown that within the overall CLL cell compartment, a subset of B-CLL cells phenotypically resemble and function as B regulatory cells (Bregs). Bregs constitute a newly designated group of B cells that have the capability to exert suppressive effects on a variety of immune cell types, mediated in part by IL-10 secretion. Several types of Bregs have been identified, with 1 subset having a CD19 CD24 CD38 immunophenotype and an enhanced capacity to secrete IL-10 (termed B10 Bregs). B10 Bregs are highly immunosuppressive and can dampen effector CD4+ T helper cells, specifically Th1 and Th17 cell responses, and also impair cTL activity.

Patients with CLL with at least 30% CD38+ B-CLL cells are designated as having CD38+ disease, which is associated with an unfavorable clinical prognosis. We have recently demonstrated that targeting CD38 with the anti-CD38 human monoclonal antibody (mAb) daratumumab downregulates B-cell receptor signaling and enhances the antitumor activity of ibrutinib in CLL cells and Waldenstrom macroglobulinemia tumor cells. These investigations revealed that daratumumab induces antibody-dependent cell-mediated cytotoxicity, complement-dependent cytotoxicity, antibody-dependent cellular phagocytosis, and direct apoptosis of CLL cells in vitro, and together these undergird its ant-CLL activity in vivo.

As CLL cells are highly dependent on their interaction with neighboring immune cells, we investigated the effects of CD38-targeting agents on Breg-like CLL cells, T-cell subsets, their associated immune cytokine environment, and the downstream effect on functionality of cTls from patients with CLL.

Materials and methods

Human samples, T-cell assays, mouse model, and statistical analysis

Peripheral blood mononuclear cells (PBMCs) from patients with CLL (n = 22 with ≥90% tumor B cells; clinical/biological data in Table 1) and healthy donors were isolated under a protocol approved by the Mayo Clinic Institutional Review Board. CD19 CD5 CD24 CD38lo CLL cells and CD4+ CD25 CD127lo Tregs were sorted out using magnetic beads/flow-sorter (sorting and gating strategy in supplemental Materials and methods). A CLL–patient-derived xenograft (PDX) model was established, using PBMCs isolated from a patient with CLL with CD38+ disease, injected into NSG mice (The Jackson Laboratory). Treg formation assays were carried out using naive Th cells prestimulated with anti-CD3 (5 μg/ml)/CD28 (5 μg/ml) antibodies followed by coculture with either autologous Breg-like (CD19 CD24 CD38+) or non-Breg (CD19 CD24 CD38+) CLL cells. cTL proliferation was determined using CellTrace carboxyfluorescein succinimidyl ester (Thermo Scientific) on a flow cytometer. cTL cytolytic activity was measured by co-culturing cTls with calcine-AM–labeled autologous/allogeneic CLL cells for 6 hours. For experiments in which CD38 expression was assessed in cells treated with daratumumab or kuromanin, a multiplate fluorescent isothiocyanate-conjugated anti-CD38 antibody (Cytognos, Salamanca, Spain) was used as previously described by us. Daratumumab was acquired through clinical sources; small molecule CD38 inhibitors, kuromanin and 78c, were purchased from Selleckem and provided as a gift from Eduardo Chini (Mayo Clinic, Rochester, MN), respectively. All resulting values are expressed as mean ± standard error of the mean (SEM). Data were analyzed for significance in GraphPad Prism by paired Student t test, Mann-Whitney U test, or 1-way analysis of variance with Bonferroni adjustment as specified. Correlation coefficients and their significance were calculated by 2-tailed Spearman’s rank correlation. A confidence interval of 95% or P < .05 was considered statistically significant.

Results

CD38-targeting agents decrease IL-10 and increase IFN-γ in the CLL-immune milieu

The immune system of patients with CLL is highly dysfunctional and poised toward an immunosuppressive disposition, which leads to compromised antitumor immune-effector activity and poor clinical outcome. Increased IL-10 and TGF-β (anti-inflammatory), along with decreased IL-2 and interferon-γ (IFN-γ) (proinflammatory), are known to promote tumor growth and dampen immune functionality. Cytokine analysis from the plasma of patients with CLL revealed IFN-γ and IL-2 levels to be 3.6- and 1.5-fold lower vs healthy donors (P = .0022 and P = .0065, respectively; Figure 1A-B). Conversely, plasma levels of IL-10 and TGF-β were two- and 6.4-fold higher in patients with CLL vs healthy donors (P = .001 and P = .0012, respectively; Figure 1C-D). As increased expression of CD38 on leukemic cells is a well-established parameter that confers poor clinical prognosis, we performed subset analysis of cytokine levels in CD38+ or CD38– patients with CLL. Expression of IFN-γ (Figure 1E; P = .0281), IL-2 (Figure 1F; P = .0006), IL-10 (Figure 1G; P = .0003), and TGF-β (Figure 1H; P = .0002) in the plasma of CD38+ patients was significantly higher than in CD38– patients. It has been previously reported that CD38 activation can increase IL-10 levels in monocytes. Conversely, inhibition of cADPR, a substrate that is biosynthesized by CD38 NADase activity, results in decreased expression of TGF-β. We hypothesized that targeting CD38 on CLL cells would decrease IL-10 and TGF-β while raising IFN-γ and IL-2 levels. Exposure of PBMCs from patients with CLL to daratumumab raised the extracellular concentration of IFN-γ (Figure 1I; P = .022), as well as intracellular expression of IL-10, in gated CLL cells (Figure 1J; P = .0286) by nearly twofold compared with untreated cells. With kuromanin, a significant increase in intracellular IFN-γ was noted in gated CLL cells (P = .018); however, this was not evident in the extracellular fraction. Examination of extracellular IL-10 in daratumumab-treated (P = .0274) or kuromanin-treated PBMCs from patients with CLL (P = .032) was significantly lower than in their untreated counterparts (Figure 1K). A similar decrease in intracellular IL-10 (and IL-10 mean fluorescence intensity [MFI]; supplemental Figure 1) was noted in the CD19+ gated fraction of daratumumab (P = .0125) or kuromanin-treated PBMCs (P = .0158; Figure 1L). Altogether, these results demonstrate that targeting CD38 (by either therapeutic mAb or small molecule) can alter cytokines (IL-10, IFN-γ) involved in regulating the interaction between CLL cells and neighboring immune cells.
A subset of CLL cells resembles B10 Bregs, produces IL-10, and can be eradicated by CD38-targeting agents

A subset of CLL cells resembles Bregs and shares IL-10 competence. We noted that IL-10 was significantly higher within the CD19+CD5+CD24+CD38+ fraction of CLL cells (Breg-like CLL cells) relative to the CD19+CD5−CD24+CD38+ fraction (non-Breg CLL cells) by nearly twofold (P = .006 and 0.0012; Figure 2A-C). We hypothesized that pharmacologic interference of CD38 would decrease the percentage of IL-10+ Breg-like CLL cells. Compared with vehicle, intracellular IL-10 in Breg-like CLL cells was significantly decreased by daratumumab (from 31.58% to 17.79%; P = .019), but not with either kuromanin or 78c (Figure 2D). In parallel experiments and in line with our recently published data, we noted significant apoptosis of Breg-like CLL cells (Figure 2D). Taking together, our work supports CD38 as a novel target for the elimination of Breg-like cells, which may be a result of soluble IL-10 and/or TGF-β secreted by daratumumab, which is associated with an overall decrease in IL-10 levels.

CLL cells promote conversion of naive T-helper cells into Tregs in an IL-10/TGF-β-dependent manner

Tregs can dampen antitumor CD4+ or CD8+ T-cell responses and promote neoplastic B-cell growth. We noted a significantly higher proportion of Tregs present within PBMCs from patients with CLL vs healthy donors (11.86% ± 2.03% vs 1.94% ± 0.32%; P = .026; Figure 3A) and which was significantly higher in CD38+ (16.10% ± 2.17%) vs CD38− patients with CLL (9.92% ± 1.38%; P = .04; supplemental Figure 2A-B). Absolute cell counts and percentage Tregs assessed from 19 patients with CLL are provided in supplemental Table 1. We hypothesized that Tregs in patients with CLL may be increased because of the influence of CLL cells on naive T helper cells. In a coculture of CD19+CD5− CLL cells with patient-autologous naive T helper cells (lower chamber), and observed significantly more induction of FoxP3+ Tregs (iTregs) in the coculture containing Breg-like CLL clones express CD38. When our cell line and cell-free supernatant from Breg-like CLL cells were cultured with autologous naive T helper cells from healthy donors (Figure 3B). Next, in a transwell assay, we placed magnetically isolated CLL Bregs or, separately, those independently isolated from healthy donors, we noted a 7.5-fold induction of FoxP3+ Tregs (iTregs) in the coculture containing naïve T helper cells from healthy donors (P = .0014). Coculture between CLL cells and autologous naive T helper cells did not induce increased iTreg formation (Figure 3B). Next, in a transwell assay, we placed magnetically isolated CLL Bregs or, separately, non-Breg CLL cells (upper chamber) in proximity with patient-autologous naïve T helper cells (lower chamber), and observed significantly more induction of iTregs (23.65% ± 1.15%) in the transwell containing CLL Bregs relative to the transwell containing non-Breg CLL cells (9.91% ± 1.61%; P = .011; Figure 3C, top and bottom red dot plots). We reasoned that increased iTreg formation may be a result of soluble IL-10 and/or TGF-β secreted mainly from Breg-like CLL cells, inferred from the proportion of IL-10+ or TGF-β+ cells being higher in CD38+ patients with CLL (Figure 1G; supplemental Figure 3). Indeed, in the transwell containing Breg-like

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**Table 1. Clinical characteristics of patients with CLL**

| Pt # | Sex | Age, y | Rai stage | Tx status | IgVH | ZAP70+ | Del 17P+ | Complex karyotype | Cytogenetics | CD38 | % | MFI | sAbc |
|------|-----|-------|-----------|-----------|------|--------|---------|------------------|-------------|------|----|-----|-----|
| 1*   | F   | 74    | I         | TN        | Unmutated | NA     | N       | N              | NA           | 38.01 | 313 | 3528 |
| 2*   | M   | 68    | IV        | R/R       | Unmutated | N      | N       | N              | del 11q, del 13q | 32.31 | 253 | 3548 |
| 3*   | M   | 66    | III       | TN        | Unmutated | Y      | N       | del 13q       | 36.46 | 270 | 3801 |
| 4    | M   | 65    | III       | R/R       | NA     | Y      | Y       | del 13q, del 2q, loss of 11p | 20.98 | 109 | 1445 |
| 5    | M   | 64    | II        | TN        | NA     | N      | N       | del 13q       | 24.55 | 208 | 2818 |
| 6*   | M   | 66    | I         | R/R       | Unmutated | Y      | N       | del 13q       | 37.43 | 180 | 3489 |
| 7*   | F   | 43    | 0         | TN        | Mutated | N      | N       | Normal        | 6.3   | 59  | 776  |
| 8    | F   | 78    | 0         | TN        | Mutated | N      | N       | Normal        | 10.54 | 132 | 954  |
| 9    | F   | 68    | R/R       | NA        | NA     | Y      | Y       | Normal        | 46.71 | 568 | 8317 |
| 10*  | F   | 69    | I         | TN        | Unmutated | Y      | Y       | del 11q, del 13q | 11.86 | 144 | 3025 |
| 11   | M   | 42    | I         | TN        | Unmutated | Y      | Y       | del 13q, del 6q, del 11q | 13.39 | 180 | 2454 |
| 12   | M   | 64    | 0         | TN        | Mutated | Y      | N       | del 13q       | 13.12 | 182 | 2511 |
| 13   | M   | 53    | I         | PT        | Unmutated | Y      | N       | Normal        | 38.81 | 243 | 3988 |
| 14   | M   | 59    | I         | TN        | Mutated | Y      | N       | del 11q       | 18.11 | 144 | 3025 |
| 15*  | M   | 71    | IV        | PT        | Unmutated | Y      | N       | del 13q       | 36.1  | 270 | 3801 |
| 16*  | F   | 69    | I         | TN        | Unmutated | Y      | N       | del 13q       | 38.54 | 347 | 4142 |
| 17*  | F   | 50    | I         | TN        | NA     | N      | N       | Normal        | 45.95 | 401 | 5498 |
| 18   | F   | 74    | 0         | R/R       | Unmutated | NA     | N       | Normal        | 7.58  | 127 | 1011 |
| 19   | M   | 66    | 0         | TN        | Mutated | NA     | N       | del 13q       | 15.98 | 137 | 2544 |
| 20   | M   | 58    | 0         | TN        | Mutated | N      | N       | Normal        | 18.3  | 50  | 630  |
| 21   | M   | 60    | I         | TN        | NA     | N      | N       | Normal        | 13.99 | 172 | 2358 |
| 22*+ | M   | 65    | III       | R/R       | Mutated | N      | N       | del 13q       | 37.49 | 431 | NA   |

*Patients who were designated as CD38 positive using the standardized clinical definition in which more than 30% of CD19+CD5− CLL clones express CD38.
CLL cells and naive T helper cells, an anti-IL-10 or anti-TGF-β neutralizing Ab significantly reduced iTreg formation by nearly five- or 4.3-fold, respectively (compare top red scatter dot plot points with adjacent blue and brown points; \( P = .0068 \)). The combination of the 2 neutralizing antibodies did not result in a greater decrease of iTreg formation than that noted with singular use of anti-IL-10 or anti-TGF-β Ab.
anti-TGF-β neutralizing Abs. Contrastingly, in the transwell with non-Breg CLL cells, insignificant reductions in iTreg formation were noted with anti-IL-10- and anti-TGF-β neutralizing Abs (Figure 3C).

These novel findings prompted us to conclude that Breg-like CLL cells are capable of promoting iTreg formation in an IL-10- and TGF-β-dependent manner.
Figure 3. Breg-like CLL cells promote conversion of naive Th cells into Tregs that are mediated by IL-10 and TGF-β. (A) Baseline proportion of CD4+CD25−CD127loFoxP3+ Tregs was compared in PBMCs isolated from healthy donors (n=6) or patients with CLL (n=12). (B) CD19+CD5+CD4−T cells and autologous naive Th cells (CD4+CD25−) were sorted from patients with CLL (n=13). CD4+ T cells were also isolated from healthy donors (n=6). Next, CLL cells were cocultured with either autologous or allogeneic (healthy donor-derived) naive Th cells in a 1:1 ratio that were prestimulated with anti-CD3 and anti-CD28 antibodies. After 3 days of incubation, cells were labeled for Treg markers and analyzed by flow cytometry. (C) To identify the mediators of conversion of naive Th cells to Tregs, a transwell coculture assay was conducted. The upper chamber contained either flow-sorted CD19+CD5+CD24+CD38hi (autologous Breg-like CLL cells) or CD19+CD5+CD24+CD38− (autologous non-Breg) CLL cells from 8 patients with CLL, whereas the lower chamber contained autologous flow-sorted naive Th cells prestimulated with anti-CD3 and anti-CD28 antibodies. In addition, IL-10 or TGF-β neutralizing Ab (alone or in combination with one another) were added to the upper chamber and incubated for another 72 hours. The proportion (%) of Tregs was subsequently analyzed by flow cytometry. Contour or density plots are representative, and compiled data are presented as mean ± SEM, with individual data points overlaid. Each experiment was performed at least twice in duplicate. *P<.05; **P<.001.

Anti-CD38 agents are lethal to Tregs

CD38hi Tregs have impressive immunosuppressive capabilities.30,31 We observed a significantly higher proportion of CD38hi Tregs in patients with CLL (54.27% ± 3.91%; MFI, 616.8 ± 36.27) compared with healthy donors (9.07% ± 1.12%; *P<.011; MFI, 234.0 ± 10.73; **P=0.12; Figure 4A). Moreover, a positive correlation between the percentage of CD38hi Tregs and CD38hi B-CELL cells was also observed (r = 0.1; P = .0042; supplemental Figure 2C). As blocking CD38 induces apoptosis in Bregs, we assessed whether Tregs were equally sensitive to CD38-targeting agents. In PBMCs from patients with CLL, ex vivo treatment with daratumumab reduced Tregs by approximately 49% (*P = .0286; Figure 4B). To determine whether CD38-targeting agents decreased the formation of iTregs, cocultures containing CLL cells and naive T helper cells from healthy donors were treated with daratumumab or kuromanin. A significant reduction in iTreg formation after exposure to daratumumab (~50% decrease) or kuromanin (30% decrease; *P = .022, Figure 4C) was observed. We also tested whether anti-CD38 agents reduced iTreg...
Figure 4. CD38-targeting agents deplete Tregs

(A) CD38 expression was measured as percentage expression and MFI on CD4+CD25+CD127lo gated Tregs in PBMCs isolated from healthy donors (n = 6) or patients with CLL (n = 12). Gating strategy is shown, and compiled data of percentage CD38+ cells are presented as mean ± SEM.

(B) Using a similar gating strategy, percentage of Tregs was measured after treatment with daratumumab (1 μg/mL; 72 hours) in PBMCs from 13 patients with CLL.

(C) The effect of cytokine induction on CD4+CD25+ cells was assessed in PBMCs from healthy donors.

(D) The effect of cytokine induction on CD4+CD25+ cells was assessed in PBMCs from healthy donors.
Separately, CD4+ daratumumab (D, 1 mg/mL) or kuromanin (K, 30 μM) were added to a mixture of PBMCs from CLL patients and healthy donors and incubated for 24 hours. The effect of daratumumab (D) or kuromanin (K) was assessed on cytokine-induced Tregs (without any CLL cells). (E) The direct cytotoxic effects of daratumumab (Dara) or kuromanin (Kuro) on CLL cells were tested by coculturing CD4+ T cells from healthy donors with or without daratumumab (1 μg/ml) or kuromanin (30 μM) for 24 hours. The effect of anti-CD38 agents and observed that ex vivo exposure of PBMCs from patients with CLL to daratumumab induced cell death in 38.36% ± 6.06% of FoxP3-gated Tregs (P = .011; Figure 4E). Treg cell death was also confirmed in flow-sorted CD4+ CD25+ T cells from PBMCs of patients with CLL, in which daratumumab or kuromanin induced apoptosis in 33.14% ± 7.73% (P = .0009) or 18.04% ± 1.94% (P = .0041) of cells, respectively (Figure 4F). Collectively, our results demonstrate that Tregs from patients with CLL have high CD38 expression and are depleted from the PBMCs of patients with CLL on exposure to daratumumab.

Daratumumab increases the proportion of Th17 cells and promotes expansion of activated cTLs that shows patient tumor-specific cytolytic activity

As anti-CD38 agents depleted Tregs and Breg-like CLL cells, we deduced that changes in the proportion of other T-cell subsets may also be evident (workflow/methodology shown in supplemental Figure 4). Assessment of basal T effector levels were found to be lower in PBMCs from patients with CLL (50.05% ± 5.41%) vs healthy donors (86.94% ± 11.19%; P = .014; Figure 5A). After treatment of PBMCs from patients with CLL with daratumumab, a significant increase was noted in the proportion of T effector cells (64.53% ± 2.12%; P = .012; Figure 5B). Notably, significant T effector proliferation was also seen in CD4+ T cells isolated from the PBMCs of patients with CLL and treated ex vivo with daratumumab, but not in isolated CD4+ T cells that were depleted of Tregs (supplemental Figure 5). We also profiled Th1, Th2, and Th17 subsets and found the percentage of Th17 cells to be 11-fold higher in patients with CLL vs healthy donors (P = .004). On treatment with daratumumab, the proportion of Th17 cells from patients with CLL further increased (P = .015), with no changes in Th1 and Th2 subsets seen (Figure 5C-D). In PBMCs depleted of CLL cells and treated with daratumumab, the proportion of Th17 cells remained unchanged, suggesting that daratumumab did not directly increase Th17 cells, but perhaps indirectly through acting on (and killing) CLL cells in the PBMC mixture (supplemental Figure 6). Overall, exposure to daratumumab also increased the proportion of CD4+IL17+ and CD4+IFN-γ+ T cells (supplemental Figure 7). As such, we reasoned that depletion of Tregs and tandem increases in the proportion of Th17 and CD4+IL17+ cells by daratumumab may lead to enhanced cTL antitumor function.32-35 CD38 expression on sorted cTLs from patients with CLL did not differ drastically from that of healthy donors (11.29% ± 1.36% vs 8.27% ± 0.56%; Figure 6A-B), and thus we tested what direct effect daratumumab would have on these cells. Intriguingly, sorted and carboxyfluorescein succinimidyl ester-labeled cTLs from healthy donors treated with daratumumab ex vivo showed a twofold increase in proliferation (P = .026) that was not observed in cTLs from patients with CLL.
isolated from patients with CLL (Figure 6C). Despite lack of proliferation, it remained plausible that cTLs may be activated as a result of daratumumab-mediated reduction in the proportion of Bregs and Tregs, decreased IL-10 and TGF-β, and increased extracellular IFN-γ. Thus, we treated PBMCs from patients with CLL with daratumumab, and noted a modest but significant increase in the proportion of cTLs in the PBMC mixture exposed to daratumumab (28% cTLs) vs control Ab (17% cTLs; P = .03; Figure 6D). In line with this observation, a significant increase in CD137⁺ cTLs (P = .022; Figure 6E) and decrease in PD-1⁺ cTL subsets (P = .011; Figure 6F) was also evident in PBMCs from patients with CLL exposed to anti-CD38 agents, indicating an activated cTL phenotype. It has recently been shown that PD1⁺ CD38⁺CD8⁺ T cells are immunosuppressive and mediate resistance to immune checkpoint therapy. Assessment of PD1⁺ CD38⁺ cTLs in PBMCs from patients with CLL treated with daratumumab or kuromanin ex vivo showed a significant decrease in this cTL subtype (P = .001 relative to control; Figure 6G). We also noted that daratumumab exposure resulted in an increased proportion of IL17⁺ and IFN-γ⁺ cTLs (supplemental Figure 8), which prompted us to test the hypothesis that these activated/daratumumab-primed cTLs would be highly effective in lysing CLL cells (schema of experiment in Figure 6H). As expected, daratumumab-primed cTLs had greater cytolytic activity toward autologous sorted CLL cells as compared with cTLs that were exposed to an immunoglobin G1 (IgG1) control Ab (P = .019; Figure 6I, compare left side blue dots with left side red squares). To further understand whether these effects were patient-tumor–specific, we exposed daratumumab-primed or IgG1 Ab-exposed cTLs from patients with CLL to either autologous or allogeneic (different patient) CLL cells. Although significant cytolysis was observed in the autologous cocultures that contained daratumumab-primed cTLs vs the IgG-exposed cTLs (P = .026; Figure 6J, compare left side blue dots with left side red squares), nearly equivalent specific lysis between daratumumab-primed cTLs or the IgG-exposed cTLs was observed in the allogeneic coculture (supplemental Figure 6J, compare right side black dots with right side red dots). Altogether, these results indicate to us that daratumumab, through an undefined mechanism, influences the activation state of cTLs and enhances their cytolytic activity toward CLL cells (in a patient tumor-specific manner).

**Anti-CD38–targeted therapy alters Breg, Treg, and cTL levels in vivo in a CLL-PDX model**

A CLL-PDX mouse model was established to determine the effect of daratumumab or kuromanin on B- and T-cell subsets. Compared with mice that received only vehicle, a significant reduction in the absolute number of CD19⁺ CD5⁺ CLL cells was noted in mice treated with daratumumab or kuromanin (Figure 7A). Notably, Breg-like CLL cells in both daratumumab-treated (P = 0.0078) and kuromanin-treated mice (P = .0045) were reduced (Figure 7B). Contrastingly, total Tregs were decreased in daratumumab-treated mice.
Figure 6. Daratumumab induces expansion of activated and patient-tumor specific cTLs. (A-B) cTLs were sorted from PBMCs of healthy donors (n = 6) and patients with CLL (n = 6), with CD38 levels determined as percentage expression (A) and MFI (B). (C) Flow-sorted cTLs were labeled with carboxyfluorescein succinimidyl ester followed by treatment with daratumumab for 72 hours. cTL proliferation was measured and data were represented as multicolor cell proliferation histograms. (D) PBMCs from patients with CLL were incubated with IgG1 control Ab or daratumumab (1 µg/mL) for 72 hours and CD3+CD8+ T cells were gated to determine percentage of cTLs. (E) Cell surface markers of T-cell activation (CD69, GITR, CD137) and (F) exhaustion (TIGIT, Lag3, CD244, PD1, CTLA4) were measured on CD3+CD8+ gated cTLs from PBMCs of patients with CLL, which were ex vivo treated with daratumumab (D, 1 µg/mL) or kuromanin (30 µM) for 72 hours. Contour plots are representative and compiled data are presented as mean ± SEM with individual data points overlaid. (G) The presence of CD38+PD1+ co-expressing cTLs was measured in CD3+CD8+ gated T cells from PBMCs from patients with CLL treated with daratumumab (1 µg/mL, 72 hours) or kuromanin (30 µM, 72 hours). Data are represented as scatter dot plots of percentage...
mice ($P = .0008$), but not in mice treated with kuromanin (Figure 7C). CD38$^+$ Tregs were significantly lower in mice treated with either CD38-targeting agent, but more remarkably so in daratumumab-treated mice (Figure 7D). Examination of T-helper subsets showed that both daratumumab and kuromanin decreased Th1 populations in vivo (Figure 7E); however, changes in Th2 and Th17 subsets were more variable. The proportional change in the percentage of Tregs, Th1, Th2, and Th17 cells and cTLs in spleens isolated from mice treated with CD38-targeted agents was mostly mirrored in the absolute cell count/spleen for these cell types (supplemental Table 1). Although minimal in change, a statistically significant ($P = .022$) increase in cTLs was noted in mice treated with daratumumab, but not kuromanin (Figure 7F). Notably, PD1$^+$ cTLs and PD1$^+$ CD38$^+$ cTLs were significantly lower in mice treated with either CD38-targeting agent compared with vehicle-treated mice (Figure 7G), but it was only in the daratumumab-treated cohort that we also observed an increase in Granzyme B$^+$ or LAMP1$^+$ cTLs, suggesting increased cytolytic capacity of these cells ($P = .03$ or $P = .012$, respectively, vs vehicle; Figure 7H-I).

**Discussion**

Physiologically, CD38 plays a central role in various immune functions, such as T-cell activation, neutrophil chemotaxis, dendritic cell migration, and monocyte chemokine production. In patients with CLL, these immune processes are highly dysregulated and underlie the immense degree of overall immune failure observed clinically. In the current study, we investigated the immune cytokine milieu as well as B- and T-cell subsets in patients with CLL, within the context of targeting CD38. Subset analysis in CD38$^+$ vs CD38$^-$ patients with CLL revealed not only higher levels of IL-10 and TGF-β but also IFN-γ and IL-2 in plasma from CD38$^+$ patients. IFN-γ$^+$ B cells have been described to promote the innate immune response against intracellular infections by activating dendritic cells via IFN-γ secretion at levels similar to those produced by NK...
On day 9, mice were sacrificed, with the spleen and blood harvested, followed by human immune cell analysis carried out showing the absolute number of CD19 subsets (including cTLs) via enhanced secretion of IFN-γ.

Increased levels postexposure to anti-CD38 agents is unclear, it is possible that these cells are able to also activate various T-cell subsets (including cTLs) via enhanced secretion of IFN-γ and IL-2.

Studies into B cells with regulatory properties demonstrate that a subset of CLL cells shares remarkable similarity with B10 Bregs. We observed that the CD19⁺CD5⁺CD24⁺CD38hi subset of Breg-like CLL cells has a distinct IL-10 secretion pattern.

Figure 7. Effects of anti-CD38 agents on Breg-like CLL cells, Tregs, T helper, and cTLs in a CLL PDX model. A short-term PDX-CLL clinical trial concept mouse model was developed via injecting (IV) PBMCs isolated from a patient with CLL (clinical characteristics in Table 1, Pt. 22) into NSG mice. After 24 hours, mice were treated with vehicle (V), daratumumab (D, 20 µg/mL loading dose followed by 10 µg/mL), or kuromanin (K, 20 µg/mL) by IV tail vein injection on postimplantation days 2, 5, and 8. On day 9, mice were sacrificed, with the spleen and blood harvested, followed by human immune cell analysis carried out showing the absolute number of CD19⁺CD5⁺ CLL cells (A); Breg-like CLL cells (CD19⁺CD5⁺CD24⁺CD38⁺) (B); Tregs (CD4⁺CD25⁺FoxP3⁺) cells (C); CD38⁺Tregs (CD4⁺CD25⁺CD127lo/FoxP3⁺) gated cells (D); T-helper cells Th1, Th2, and Th17 subsets (E); CD8⁺ cTLs (F); PD1⁺CD38⁺CD8⁺ cTLs (G); Granzyme B⁺CD8⁺ cTLs (H); and LAMP1⁺CD8⁺ cTLs (I) were probed and gated for using human antibodies in mouse splenocytes cells. Absolute cell counts for Bregs, Tregs, Th1, Th2, Th17, and cTLs were also determined and are shown in supplemental Table 2. Contour plots are representative and compiled data are presented as mean ± SEM with individual data points overlaid. Each experiment was performed at least twice in duplicate. *P < .05; **P < .001.
compared with CD38lo/non-Breg CLL cells. Indeed, anti-CD38 agents induced apoptosis in Breg-like CLL cells, which was associated with a decrease in IL-10 secretion. Mechanistically, IL-10 and TGF-β secreted by CD38hi CLL-Bregs promotes the expansion/outgrowth of Tregs, more so in comparison with CD38lo CLL cells, which we demonstrated via co-culturing CD38hi or CD38lo CLL cells with patient-autologous naïve T cells, and used neutralizing antibodies for IL-10/TGF-β. Jak et al have reported a CD27-CD70-dependent mechanism for iTRG-mediated iTRG expansion; however, our observation that iTRG formation is dependent on IL-10/TGF-β is novel (Figure 3C). In line with this, we observed that daratumumab (and to a lesser extent kuromanin) significantly reduced iTRG levels. We questioned whether a reduction in Tregs was due to the direct effect of CD38-targeting agents or because CD38-targeting agents deplete Bregs, reduce IL-10/TGF-B levels, which in turn decrease Tregs. During this investigation, we made a novel observation that nearly 50% of Tregs from patients with CLL have high CD38 expression compared with Tregs from healthy donors. Furthermore, a significant correlation between the percentage of Tregs and percentage of CD38+ CLL cells present in the PBMCs of patients with CLL was also noted (r = 0.5; P = .012; supplemental Figure 2B). Krejcik et al36 first reported that treatment of patients with multiple myeloma with daratumumab rapidly depletes CD38+ Tregs, and subsequent studies by Feng et al44 showed that blocking CD38 with isatuximab induces Treg apoptosis. Our observations mirror these findings and show that exposure of PBMCs from patients with CLL to daratumumab, ex vivo, induces death of Tregs, a substantial number of which express high levels of CD38.

Among other T-cell subsets, we noted low T effector cell levels in PBMCs from patients with CLL, which increased significantly after exposure to daratumumab. We surmise that an increase in these cells was a result of daratumumab-mediated death of Breg-like CLL cells and Tregs (with concomitant changes in cytokine levels), which act as a stimulus for effector T-cell proliferation, and was not a direct stimulatory effect of the drugs itself on T effector cells. Indeed, in PBMCs from patients with CLL, where CD4+ T cells were isolated and Treg depletion was performed, CD38-targeting agents did not increase T effector cell proliferation. In contrast, when all CD4+ T cells (which included Tregs) isolated from PBMCs from patients with CLL were treated with daratumumab or kuromanin, a significant increase in T effector cell proliferation was observed (supplemental Figure 5). Dissection of the various T helper subsets revealed increased Th17 cells in patients with CLL compared with healthy donors, which has previously been reported.1 In CLL, Th17 cells have been suggested to play an antitumor role and are garnering interest in cancer immunotherapy as a whole.47 Chatterjee et al have described the similarities between artificially generated hybrid Th1/Th17 cells and CD4+ T cells from CD38 knockout mice. These 2 T-cell populations display superior antitumor capabilities compared with standard Th1 and Th17 cells or CD4+ T cells from CD38 wild-type mice.47 We noted that daratumumab treatment of CLL PBMCs increased the proportion of Th17 cells; however, the functional aspect of this remains to be defined. Given the changes that were induced by exposure to daratumumab in Tregs and Th17 cells, we also examined cTLs. The functional capacity of cTLs in patients with CLL is severely impaired as a result of defects in immune synapse formation and dysregulated expression of activation or immune checkpoint molecules.1,48 Direct exposure of sorted CLL patient-derived cTLs treated ex vivo with daratumumab did not result in their proliferation, but rather death, in ~11% (P = .007; supplemental Figure 9), an effect most likely manifested in CD38+ or CD38+ PD1+ cTLs (Figure 6G). However, in the context of a PBMC mixture, exposure to daratumumab resulted in the relative expansion of activated cTLs, with increased cytolytic (and patient-tumor-specific) activity.

Alterations of B- and T-cell subsets in patients with multiple myeloma receiving treatment with daratumumab have been described, and our ex vivo as well as in vivo observations in a short-term CLL-PDX model echo these observations.30 As our goal was to study the effect of anti-CD38 treatment on T-cell subsets, establishment of a long-term PDX model was not optimal because of the possibility of xenograft vs host-like disease known to occur in mice injected with CD4+ naïve T cells.49,50 Results from our in vivo PDX model were similar to those from our ex vivo CLL patient PBMC analyses, in which daratumumab depleted Breg-like CLL cells and Tregs along with increasing Th17 cells and cTLs with enhanced cytolytic capacity (increased Granzyme B+ or LAMP1+ cTLs).

It is noteworthy that in the majority of experiments, we used kuromanin as a comparator to daratumumab, which inhibits CD38 enzymatic function. The effects of kuromanin were more variable compared with daratumumab in both in vitro/ex vivo as well as in vivo assays. In PBMCs from both CD38+ and CD38+ patients with CLL, we observed that kuromanin treatment led to a significant reduction in IL-10+ CLL cells. However, when interrogation of IL-10 was specifically performed in Breg-like CLL cells, kuromanin and even 78c (more potent CD38 enzymatic inhibitor) did not reduce intracellular IL-10 (Figure 2D). This finding warrants further investigation, but suggests that the link between IL-10 expression and CD38 is not linear, and enzymatic blockage of CD38 in Breg-like CLL cells specifically may not be sufficient to downregulate IL-10. Aside from this observation, we noted in our PDX model that kuromanin did not decrease the percentage of Tregs. We surmise that in vivo, kuromanin is a weak CD38 enzymatic inhibitor, and ongoing experiments with 78c will help resolve this.

Although the B-cell compartment is undoubtedly aberrant in patients with CLL, a remarkable degree of dysfunction is also noted in the various T-cell subsets. Our present investigation establishes that therapies aimed at CD38 promote both direct killing of Breg-like CLL cells along with Tregs, with a subsequent enhancement of antitumor T-cell responses. Collectively our observations demonstrate that anti-CD38 agents have an immunomodulatory effect, reprogramming the CLL microenvironment and restoring antitumor immune functionality.

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Authorship
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