



Tarafdar, A. et al. (2017) CML cells actively evade host immune surveillance through cytokine-mediated downregulation of MHC-II expression. *Blood*, 129(2), pp. 199-208. (doi: [10.1182/blood-2016-09-742049](https://doi.org/10.1182/blood-2016-09-742049))

This is the author's final accepted version.

There may be differences between this version and the published version. You are advised to consult the publisher's version if you wish to cite from it.

<http://eprints.gla.ac.uk/130643/>

Deposited on: 28 October 2016

CML cells actively evade host immune surveillance through cytokine-mediated downregulation of MHC-II expression

Anuradha Tarafdar,¹ Lisa E.M. Hopcroft,¹ Paolo Gallipoli,^{1,2} Francesca Pellicano,¹
Jennifer Cassels,¹ Alan Hair,¹ Koorosh Korfi,¹ Heather G. Jørgensen,¹ David
Vetrie,¹ Tessa L. Holyoake,^{1,†} Alison M. Michie^{1,†}

¹Institute of Cancer Sciences, College of Medicine, Veterinary and Life Sciences, University of Glasgow, UK; ²Department of Haematology, Cambridge Institute for Medical Research and Wellcome Trust-MRC Cambridge Stem Cell Institute, University of Cambridge, UK;

†Co-corresponding authors: Alison M. Michie, Paul O’Gorman Leukaemia Research Centre, Gartnavel General Hospital, 21 Shelley Road, Glasgow, G12 0ZD; Tel: +44 (0)141 301 7885; E-mail: Alison.Michie@glasgow.ac.uk

Tessa L. Holyoake, Paul O’Gorman Leukaemia Research Centre, Gartnavel General Hospital, 21 Shelley Road, Glasgow, G12 0ZD; Tel: +44 (0)141 301 7880; E-mail: Tessa.Holyoake@glasgow.ac.uk

Running Title: Immune evasion by CML cells

Word number: 3650; References: 49; Tables: 1; Figures: 5.

Key Words: CML, MHC-II, cytokines, immune evasion

Abstract

Targeting the fusion oncoprotein BCR-ABL with tyrosine kinase inhibitors has significantly impacted on chronic myeloid leukemia (CML) treatment, transforming the life expectancy of patients; however the risk of relapse remains, due to persistence of leukemic stem cells (LSCs). Therefore it is imperative to explore the mechanisms that result in LSC survival and develop new therapeutic approaches. We now show that MHC-II and its master regulator class II transactivator (*CIITA*) are downregulated in CML compared to non-CML stem/progenitor cells in a BCR-ABL kinase independent manner. IFN γ stimulation resulted in an upregulation of *CIITA* and MHC-II in CML stem/progenitor cells, however the extent of IFN γ -induced MHC-II upregulation was significantly lower than when compared with non-CML CD34⁺ cells. Interestingly, the expression levels of *CIITA* and MHC-II significantly increased when CML stem/progenitor cells were treated with the JAK1/2 inhibitor ruxolitinib (RUX). Moreover, mixed lymphocyte reactions (MLRs) revealed that exposure of CD34⁺ CML cells to IFN γ or RUX significantly enhanced proliferation of the responder CD4⁺CD69⁺ T cells. Taken together, these data suggest that cytokine-driven JAK-mediated signals, provided by CML cells and/or the microenvironment, antagonize MHC-II expression, highlighting the potential for developing novel immunomodulatory-based therapies to enable host-mediated immunity to assist in the detection and eradication of CML stem/progenitor cells.

Key Points

- MHC-II and its master regulator *CIITA* are downregulated in CML stem/progenitor cells in a BCR-ABL kinase independent manner.
- JAK1/2 inhibition increased MHC-II expression, suggesting elevation of CML immunogenicity may provide a way to reduce CML persistence.

Introduction

The development of tyrosine kinase inhibitors (TKIs) to target BCR-ABL kinase has revolutionized the management of chronic phase CML with many patients now predicted to have a normal life expectancy.^{1, 2} Remission is maintained by continuous administration of TKI and assessed by quantification of *BCR-ABL* transcripts in the blood. For the 10-20% of patients who achieve deep and durable molecular responses, discontinuation studies have been conducted.^{3, 4} Approximately 60% of patients maintain a major molecular response over time.⁵

Prior to TKI introduction, CML was a common indication for allogeneic stem cell transplantation (alloSCT). In this setting disease remission was achieved by the combination of anti-leukemic chemo-radiotherapy and active graft versus leukemia (GVL) effect. The degree of immune recognition of leukemic cells by the donor immune system was such that disease relapse, if it occurred, could be managed successfully by administration of donor lymphocytes.⁶ Whilst it is well recognised that the effect of alloSCT and GVL is mainly an allo-immune effect mediated through non-disease-specific minor histocompatibility antigens, it is likely that CML cells express disease-specific antigens recognizable by the donor immune system. The role of the patient's own immune system in recognizing BCR-ABL-expressing cells, and whether this can be boosted for beneficial effect, is currently under investigation in vaccination studies, although no convincing results have been reported.^{7, 8} Similarly, it is not known whether immune recognition by the patient's immune system is playing a part in maintaining remission of non-relapsing patients in whom TKI treatment is discontinued.

While CD8⁺ cytotoxic T lymphocytes (CTL) are considered to play a major role in tumor immunity, CD4⁺ T helper cells are also important for mediating anti-tumor associated immune responses, possibly through optimal induction and maintenance of CTL responses, interactions with effector cells and production of anti-tumor-associated cytokines such as IL2 and IFN γ .^{9, 10} As such, solid tumors (e.g. non small cell lung cancer, mammary adenocarcinoma, colorectal and gastric) and hematological cancers (B cell lymphomas) display MHC class II (MHC-II) downregulation, reducing the host immune response towards the tumor; correlations have been found between higher MHC-II expression and better prognosis.^{11, 12} Our microarray datasets comparing the expression of genes between normal and CML stem/progenitor revealed a significant

downregulation in the antigen presentation (exogenous antigen) pathway in quiescent and dividing CD34⁺ CML cells.¹³ Here, we investigate the biological relevance of this finding, determining the mechanisms that underlie MHC-II downregulation in CML stem/progenitor cells and examining whether its induction could render these cells more immunogenic.

Materials and Methods

Primary samples cell culture: CD34⁺ cells were enriched, after informed consent, from either CML patient chronic-phase samples at diagnosis (fresh or cryopreserved; Table 1) or allograft donors/lymphoma patients without bone marrow (BM) involvement as “non-CML” controls. The studies were approved by the West of Scotland Research Ethics Committee 4, NHS Greater Glasgow and Clyde (UK). Primary CML cells were cultured in serum-free medium, supplemented with Flt-3 ligand and SCF (each 100 ng/mL), IL3 and IL6 (each 20 ng/mL; StemCell Technologies, Cambridge, UK) and G-CSF (Chugai Pharma, London, UK) overnight. Thereafter for experimental conditions, CD34⁺-enriched CML cells were cultured in SCF, GM-CSF, and MIP- α (all 0.2 ng/mL), G-CSF and IL6 (both 1.0 ng/mL) and 0.05 ng/mL leukemia inhibitory factor (StemCell Technologies). IFN γ and TGF β were purchased from Peprotech EC Ltd., (London, UK), nilotinib (NIL) from Stratech Scientific Ltd., (Newmarket, UK) and imatinib mesylate (IM), dasatinib (DAS), SB-505124 (SB) and ruxolitinib (RUX) from Selleckchem (Houston, TX). Pan-MHC-II antibody (Ab; purified, clone Tü39) used for blocking experiments was purchased from BD Biosciences (Oxford, UK).

Microarray analysis: The microarray analysis leading to the initial identification of the MHC-II gene family (Figure 1A) was described previously.¹³ The experimental details for the second, larger quiescent/dividing microarray dataset (Figure 1B&C) and the CML primitive/progenitor microarray dataset (Figure 1C, Supplementary Figure 1A&B) are described previously.^{14, 15} The processing and normalization procedures for all datasets was carried out as described.¹⁶ All microarray datasets are summarized with respect to sample size, sorting strategy and the relevant figure in Supplemental Table 1; by combining these datasets, transcriptional profiles of 19 independent CML samples and 10 independent normal samples were analyzed. Where genes were represented by multiple probes, expression was summarized using the median value. Any genes not represented on all microarray chips were removed.

Gene set enrichment analysis (GSEA): Gene set enrichment was performed using GSEA¹⁷ (gsea2-2.2.2.jar) as obtained from the Broad Institute (<http://software.broadinstitute.org/gsea/index.jsp>); q values were calculated using

10,000 permutations of the phenotype label. The MHC-II geneset analysed by GSEA is shown in Figure 1C and is comprised of the Ingenuity pathway shown in Figure 1A and “MHC class II” molecules identified by searching Metacore KB (<https://portal.genego.com>).

Flow cytometry: Cells were harvested and resuspended in 1×10^6 cells/100 μ L FACS Buffer (PBS with 2% FCS and 0.02% sodium azide). All antibodies (Abs) were purchased from BD Biosciences unless otherwise stated. Cell staining was performed using 0.5 μ g of each flurochrome-conjugated Ab for 30 min, 4°C in dark (anti-CD34, -CD38, -HLA-DP, -DQ, -DR, -CD4, -CD69 Ab). Anti-CD8 and -CD107a Ab were purchased from eBioscience (Hatfield, UK). Events were acquired using a FACSCanto™ II cytometer and FACSDiva software (BD Biosciences) and data was analyzed using FlowJo software (Treestar Inc, Ashland, OR). To isolate CD34⁺CD38⁺ (bulk) and CD34⁺CD38⁻ (leukemic stem cell (LSC)) populations, cells were sorted using a FACS Aria™ cytometer (BD Biosciences).

RT-PCR and Fluidigm analysis: Total RNA was prepared using RNAeasy Plus extraction kit (Qiagen, Valencia, CA). RNA (1 μ g) was reverse-transcribed using SuperScript reverse transcriptase and oligo dT primers (Invitrogen Life Technologies, Paisley, UK); or where mentioned, cDNA was prepared from 300 cells using the SuperScript™ III One-Step RT-PCR System with Platinum® Taq High Fidelity (Invitrogen). PCR analysis was performed on a 48.48 dynamic array using the BioMark™ HD System (Fluidigm, San Francisco, CA) or Applied Biosystems Prism 7900HT system (Applied Biosystems, Paisley, UK) as per manufacturers' instructions. Relative gene expression was calculated using the $2^{-\Delta\Delta CT}$ method.

Mixed lymphocyte reactions (MLRs) and CD107a expression assay: Peripheral blood mononuclear cells (PBMCs; responder cells) were isolated from healthy donors by density centrifugation using Histopaque (Sigma-Aldrich, Poole, UK). The PBMCs were incubated with Dynabeads® Human T-Activator CD3/CD28 (Life Technologies) as per manufacturer's instructions for 48h in RPMI-1640 medium supplemented with 10% heat inactivated FCS, penicillin (100 U/mL), streptomycin (100 U/mL) and L-glutamine (2 mM) (Life Technologies). Activated PBMCs were sequestered from the beads using a magnet and labelled with Cell Trace Violet (CTV) (Life Technologies) prior to co-culture

with stimulator/target (bulk CD34⁺ CML) cells. For MLRs, activated PBMCs were co-cultured with stimulator cells for up to 6 days at a ratio of 3:1 responder:stimulator. Proliferation was measured as a loss of CTV mean fluorescence intensity (MFI) upon analysis by flow cytometry. For CD107a expression, activated PBMCs were co-cultured with bulk CD34⁺ CML target cells for up to 6 days at a ratio of 1.5:1 CD4⁺ T cells:target.

Statistical analysis: Average responses from at least 3 individual CML donors are shown (mean \pm SEM). Statistical analysis was performed using GraphPad Prism 4 (GraphPad Software Inc., CA), using students paired or unpaired *t*-test (* $p < 0.05$; ** $p < 0.005$; *** $p < 0.001$), as appropriate.

Results:

MHC-II expression is downregulated in CML CD34⁺ stem/progenitor cells, in a BCR-ABL kinase independent manner

Microarray datasets analyzing the expression of genes in either normal or CML CD34⁺ quiescent (Hoechst^{lo}Pyronin Y^{lo}) or dividing (Hst⁺Py⁺) cells revealed that the antigen presentation pathway was the top deregulated pathway in the CML versus normal comparison ($p=5.93 \times 10^{-9}$; Figure 1A; significantly downregulated components highlighted in green).¹³ To validate this observation *in silico*, and assess more broadly whether the MHC-II gene family is suppressed in CML cells, we extended this genelist to include other relevant MHC-II molecules and identified a second, complementary dataset¹⁴ in which we could perform Gene Set Enrichment Analysis (GSEA)¹⁷ of this MHC-II gene family. Limited numbers of normal samples prevented such an analysis in the original dataset. This analysis confirmed significant downregulation of the MHC-II family in both quiescent and dividing CML cells ($FDR \leq 0.15$; Figure 1B). The same analysis was then performed in another dataset transcriptionally profiling primitive (Lin⁻CD34⁺CD38⁻CD90⁺) CML cells against three progressively more mature cell populations all from a single cohort of CML patients and non-CML controls.¹⁵ While some genes are not downregulated individually, there is a trend for downregulation of the MHC-II genes in the primitive (Lin⁻CD34⁺CD38⁻CD90⁺) and the three more mature (Lin⁻CD34⁺CD38⁺CD123⁺CD45RA⁻; Lin⁻CD34⁺CD38⁺CD123⁻CD45RA⁻; Lin⁻CD34⁺CD38⁺CD123⁺CD45RA^{lo}) cell populations (Figure 1C and Supplementary Figure 1A respectively). This represents a significant downregulation of the MHC-II gene family overall in all four cell populations, as confirmed by GSEA (Supplementary Figure 1B). Overall, this *in silico* validation process considered 19 independent CML samples and 10 independent normal samples (Supplementary Table 1). Analysis of co-stimulatory molecules indicated that while there was some variation in their expression across the populations, TNFSF4 (OX-40L) and CD40 appeared to be downregulated in LSC-enriched populations compared with non-CML populations (Supplementary Figure 1C).

At the protein level we demonstrated a significant reduction in MHC-II surface expression on CML stem/progenitor cells, both in the bulk CD34⁺ population and in the CD34⁺CD38⁻ LSC population, compared with normal hematopoietic stem cell (HSC) populations (Figures 1D&E; Supplementary Figure 2). IFN γ , an established activator of

MHC-II expression¹⁸ which signals through the JAK/STAT pathway, upregulated MHC-II expression in CML stem/progenitor cells (Figures 1D&E; Supplementary Figure 2). However, the level of IFN γ -induced MHC-II upregulation in CML cells was significantly reduced compared with non-CML HSCs (Figures 1D&E; Supplementary Figure 2). Analysis of the master regulator of MHC-II expression, class II transactivator (*CIITA*),¹⁹ revealed a downregulation of this gene in CML stem/progenitor cells compared to non-CML cells (Figure 1F). Supporting the MHC-II data, IFN γ increased *CIITA* transcription in CML stem/progenitor cells, but not to the level observed in IFN γ -treated non-CML cells (Figure 1F). These data indicate that CML stem/progenitor cells exhibit MHC-II downregulation, which may assist in their evasion from host immunity.

Treatment of bulk CD34⁺ and CD34⁺CD38⁻ CML cells for 48h with TKIs (NIL, DAS, IM) had no effect on either MHC-II or *CIITA* expression, demonstrating that BCR-ABL activity did not significantly influence *CIITA* or MHC-II expression on CML stem/progenitor cells, in the absence or presence of IFN γ (Figure 2; Supplementary Figure 3). This suggests that MHC-II regulation in CML is BCR-ABL kinase independent.

To determine whether MHC-II expression remains downregulated upon extended treatment with TKI, bulk CD34⁺ CML stem/progenitor cells were cultured in the presence or absence of IM for 7 days prior to analysis of the residual cells *in vitro*. Over this time, IM may be expected to act either by inducing reversible changes in expression of genes that are kinase dependent, or by selecting for a population of cells expressing higher or lower levels of certain genes as the result of killing more mature CD34⁺ cells that are sensitive to apoptosis. Surface protein and gene expression of MHC-II in d7 cultured CML cells was normalized to untreated (UT) d0 expression levels. Although no significant differences in MHC-II expression were observed between d7 IM treated and UT CML cells, either on the surface of CD34⁺CD38⁺ or CD34⁺CD38⁻ LSC populations, or in the expression of MHC-II genes in bulk CD34⁺ CML cells (*HLA-DR*, *HLA-DP*, *HLA-DQ*; Figure 3A-C; Supplementary Figure 4), there was a trend towards further downregulation of MHC-II expression in 7 day IM treated CML cells. This was supported by a downregulation in *CIITA* in IM treated cells when comparing to UT on d7 (Figure 3D). When IM treated cells at d7 were compared to UT cells at d0, the observed trends reached significance for surface expression of MHC-II on CD34⁺CD38⁻ LSC CML cells, and *CIITA* expression (Figure 3A, B&D). These data suggest that the regulation of MHC-II expression in CD34⁺CD38⁺ or CD34⁺CD38⁻ LSC

CML populations is BCR-ABL kinase independent, as it is not normalized after 48h (Figure 2) or 7 day of IM treatment (Figure 3). These data also suggest that by killing more mature cells within the CD34 compartment IM enriches for cells with lower expression of MHC-II and *CIITA*, inferring that these surviving cells would be even less susceptible to immune attack.

MHC-II/CIITA expression is not modulated by aberrant methylation or TGF β -mediated signaling in CML stem/progenitor cells

Aberrant DNA hypermethylation of the *CIITA* promoter has been identified as a mechanism for diminished IFN γ -induced *CIITA* and MHC-II expression in both hematopoietic and solid tumors.^{20, 21} Analysis of *CIITA* promoter methylation did not reveal any statistical differences between CML and normal stem/progenitor cells (Supplementary Figure 5). Moreover, treatment of CML cells with methyltransferase inhibitor 5-azacytidine, did not lead to increased MHC-II expression (Supplementary Figure 6).

TGF β has previously been shown to antagonize *CIITA*^{18, 22, 23} and MHC-II expression in the presence and absence of IFN γ .¹⁹ Indeed, previous work has shown that TGF β -mediated signals are activated in CML cells,²⁴ and CML has been demonstrated to drive deregulated cytokine expression in mouse models.²⁵ Initially, to analyse the impact of TGF β on *CIITA*/MHC-II, we treated CML and non-CML cells with SB (TGF β RI, ALK4 and ALK7 inhibitor²⁶). While CML stem/progenitor cells were slightly more responsive to TGF β -mediated downregulation of MHC-II levels than non-CML cells, SB treatment did not effect MHC-II/*CIITA* expression on CML stem/progenitor cells (Supplementary Figure 7).

Inhibition of JAK elevates MHC-II expression in CML stem/progenitor cells and increases their immunogenicity.

The cytokine IL4 is also known to antagonise MHC-II and *CIITA* expression^{18, 22} suggesting that aberrant alterations in cytokine production within the niche may be responsible for their deregulation. Moreover, IL4 has been shown to maintain survival of Ph⁺ cells upon TKI inhibition.²⁷ We demonstrate elevated transcript levels of IL4 in CML compared with non-CML stem/progenitor cells (Figure 4A). As IL4 and many cytokines signal via the JAK/STAT pathway we analyzed the impact of cytokine-mediated signaling on *CIITA*/MHC-II, treating CML cells with RUX (a broad spectrum JAK1/2

inhibitor²⁸). RUX treatment of CML stem/progenitor cells significantly increased MHC-II and *C/ITA* expression both on bulk CD34⁺ cells (Figures 4B, D&E) and CD34⁺CD38⁻ LSC populations (Figures 4C,D&F). As IFN γ activates the JAK/STAT pathway and RUX inhibits the same pathway, it was anticipated that RUX treatment combined with IFN γ might reduce the level of MHC-II upregulation compared to that of IFN γ alone. However this reduction was not significant, suggesting that IFN γ signaling was not fully inhibited in the presence of a JAK inhibitor (Figure 4B-F). Supporting this, analyzing genes associated with the IFN response, only the *GBP1* response was partially modulated downwards in CML cells when IFN γ and RUX treatments were combined (Supplementary Figure 8). These data indicate that RUX-mediated elevation in MHC-II may result from an inhibition of JAKs downstream of IL4-receptors, thus arresting antagonistic signals and enhancing MHC-II expression.²²

To determine whether the IFN γ - or RUX-mediated elevation in MHC-II expression on CML cells could induce an immune response from alloreactive CD4⁺ T cells we co-cultured previously treated or control (UT) CML CD34⁺ stem/progenitor cells with CD3/CD28 activated PBMCs. No increase in expression of the cytolytic marker CD107a (LAMP1) was observed on CD4⁺ T cells after exposure to the CML targets, suggesting that there was no induction of cytotoxic CD4⁺ T cells (Supplementary Figure 9). However, when MLRs were conducted, IFN γ and RUX treatments enhanced proliferation of responder CD4⁺CD69⁺ T cells, as indicated by decreased mean fluorescence intensity of CTV (Figure 5A). Importantly, the induction of proliferation was MHC-II dependent, as addition of anti-MHC-II blocking Ab blocked cellular proliferation in the cultures containing IFN γ and RUX treated CML cells (Figure 5A).

Discussion:

Eradicating CML LSCs represents a significant therapeutic challenge that will not be addressed with TKI therapy alone, and requires novel treatments to enable removal of the disease reservoir to pave the way to a cure. One potential mechanism utilized by CML stem/progenitor cells to maintain a reservoir of LSCs is to evade the immune system by downregulating MHC-II expression. Our studies demonstrate that modulation of MHC-II is associated with an increase in CML cell immunogenicity in a JAK-dependent manner, leading us to propose a cytokine-mediated pathway for MHC-II deregulation in CML cells (Figure 5B). MHC-II and *CIITA* expression was independent of BCR-ABL kinase activity, as TKI treatment of CML cells did not significantly alter expression levels. Indeed, during IM exposure over 7d the levels of MHC-II and *CIITA* were even further downregulated, rather than normalized. Of note, *in silico* analysis of co-stimulatory molecules revealed a mixed pattern of expression depending on the dataset, although it does appear that OX40L is consistently downregulated in more quiescent CML cell subsets, compared with non-CML cells. These findings suggest that in addition to downregulating MHC-II genes, CML stem/progenitor cells may downregulate additional co-stimulatory molecules, further enhancing immune evasion. Exploitation of therapies that enhance CML stem cell immunogenicity to host immune cells could intensify immune responses against CML LSCs, thereby assisting in the elimination of this population, and ultimately the disease.

While a number of mechanisms regulate MHC-II expression, one main cause for the loss of constitutive or inducible MHC-II expression, in both hematopoietic and non-hematopoietic tumor cells, is epigenetic silencing of the gene encoding *CIITA*.^{29, 30} *CIITA* acts as a scaffold, interacting with DNA-binding transcription factors, regulatory factor X family, NF-Y and CREB, which interact with and induce MHC-II expression.^{31, 32} In addition to recruiting transcription factors, *CIITA* orchestrates the recruitment of histone acetyltransferases and histone methyltransferases to MHC-II gene promoters to regulate MHC-II expression.³³ While aberrant DNA hypermethylation of the *CIITA* promoter and breaks (non-random chromosomal translocations) can aid in the downregulation of MHC-II in hematopoietic malignancies,^{21, 29, 31} analyses of methylation marks on the *CIITA* promoter of CML LSCs revealed no difference in the CpG methylation profiles compared with non-CML stem/progenitor cells. Supporting this, 5-azacytidine treatment did not lead to increased MHC-II expression on CML cells.

These data indicate that MHC-II downregulation in CML does not occur as the result of CIITA promoter hypermethylation, suggesting other molecular mechanisms are responsible for regulating CIITA/MHC-II expression in CML.

MHC-II and CIITA expression can be modulated by a number of cytokines with TGF β , TNF α , IL4 and IL10 known to antagonise their expression,^{18, 22, 23} while IFN γ upregulates expression (Figure 5B). Therefore it is likely that the cytokine profile of the BM microenvironment and/or of the leukemic cells themselves would have a significant impact on surface MHC-II expression of CML stem/progenitor cells. Indeed CML LSCs have been demonstrated to produce higher levels of growth factors, via an autocrine loop, including elevated levels of TNF α ³⁴ and *IL4* in CD34⁺ CML cells. Furthermore, analysis of the BM stroma in CML patients and CML mouse models revealed elevated IL1, IL6, GCSF and TNF α , which together confer a growth advantage of CML LSCs over normal HSCs.^{25, 35} These findings indicate that modulation of the cytokine profile in the CML BM will have a significant impact not only on CML versus normal stem/progenitor cell survival, but also on CML LSC immunogenicity.

One family of protein kinases responsible for relaying signals downstream of cytokine receptors are the JAKs. BCR-ABL⁺ cells exhibit constitutively active JAK/STAT signaling, with JAK2 playing a central role in BCR-ABL-induced leukemogenesis.³⁶ However STAT5 activation is not dependent on JAK2 activity,³⁷ and JAK2 can also act independently of STAT5 by activating Myc,³⁸ β -catenin,³⁹ or acting directly as a histone modifier,⁴⁰ indicating that JAK2 may have pleiotropic roles in CML cells. Moreover one of the main roles of JAK signalling is in immune cell activation/regulation.⁴¹ Here, we showed that treatment of cells with the JAK1/2 inhibitor RUX enhanced the expression of MHC-II on CML stem/progenitor cells, enhancing CD4⁺ T cell proliferation. While we demonstrated the induction of a helper T cell response through RUX-mediated induction of MHC-II expression, CD4⁺ T cells have also been demonstrated to function as cytotoxic effectors against leukemic cells.^{42, 43} Upon assessment of the cytolytic marker CD107a *in vitro*, we were unable to detect an elevation in expression, indicating that under our experimental conditions cytotoxic CD4⁺ T cells were not induced. These results may reflect the limitations of the assay regarding the polyclonality of the CD4⁺ T cell population, or a requirement for optimization at the level of effector/target ratio or co-culture duration prior to assessing for the presence of cytotoxic CD4⁺ T cells.^{42, 43}

Recently RUX has been shown to modulate immune responses in mouse models of graft versus host disease (GVHD) and hemophagocytic lymphohistiocytosis

by dampening the hyperactive immune responses typical of these conditions.^{44, 45} However, despite its anti-inflammatory properties, RUX was able to preserve the GVL effect in GVHD mouse models comprising *in vivo* tumour inoculation.^{46, 47} Some of these selective effects were explained by differential responses of distinct T cell subsets to RUX particularly when used at different concentrations.⁴⁵ It is therefore possible to speculate that the modulation of the immune system by RUX is cell context dependent and potentially affected by the intensity of the inflammatory responses. Moreover using different doses of RUX might enable titration of its effects, thus maximising its ability to enhance immunogenicity of CML cells while sparing T cell immune function.

In conclusion our studies suggest a key role for cytokines, such as IL4, in reducing CML immunogenicity, and offer a novel therapeutic route to aid in the elimination of CML LSCs. Due to the BCR-ABL-independent nature of MHC-II downregulation, combination therapies of JAK inhibitors with TKI could target the disease bulk and the LSCs.^{39, 48} Indeed recent studies demonstrate synergy between RUX and NIL, inducing apoptosis in CML LSCs.⁴⁹ Our studies could suggest that LSCs remaining after dual RUX/TKI treatment may have a reduced ability to evade host immunity, due to re-expression of MHC-II. Analysis of MHC-II surface expression on LSCs of patients treated with RUX in combination with NIL in current clinical trials (clinicaltrials.gov NCT 01751425; 02253277; 01702064; 01914484) would be of great interest to test this *in vivo*. Boosting MHC-II expression on CML stem/progenitor cells may represent a valid therapeutic approach to reduce disease persistence and our findings should promote the development of novel immunomodulatory-based therapies for CML patients. These therapeutic strategies could enable CML stem cells to be more readily detected and eradicated by the host immune response, which in turn will deliver long-term remission or cure for patients.

Acknowledgements: The authors thank all CML and non-CML patients and United Kingdom (UK) haematology departments including those patients on the UK SPIRIT2 Trial Management Group for access to CML samples, Karen Stewart for collecting clinical information on patients. This study was funded by project grants from Leuka and Tenovus-Scotland (Ref. S12/21). This study was supported by the Glasgow Experimental Cancer Medicine Centre, which is funded by Cancer Research UK and the Chief Scientist's Office, Scotland. Cell sorting facilities were funded by the Kay Kendall Leukaemia Fund (KKL501) and the Howat Foundation. A.T. was funded by a Bloodwise project grant (13012), P.G. was funded by a Medical Research Council (MRC) UK clinical research training fellowship grant (G1000288), H.G.J. was funded by the Friends of Paul O'Gorman Leukemia Research Centre, F.P., L.H. and T.L.H. were supported by Cancer Research UK Programme grant (C11074/A11008), D.V. by LLR project grant (14005) and A.M.M was supported by an MRC project grant (MR/K014854/1).

Authorship Contributions:

AT designed/performed the majority of experiments, analyzed and interpreted the data, carried out statistical analysis and drafted the manuscript; PG, KK and FP performed some experiments and carried out data analysis; LEMH carried out microarray and pathway analyses; JC carried out the cell sorting for all the experiments and AH processed all of the primary samples; HGJ supervised the studies and performed some of the experiments; DV, TLH, AMM obtained funding for the study, designed the research, supervised the studies, analyzed and interpreted the data and wrote the manuscript. All authors reviewed the manuscript.

Disclosure of Conflicts of Interest:

The authors have no disclosures to declare.

References:

1. Druker BJ, Guilhot F, O'Brien SG, et al. Five-year follow-up of patients receiving imatinib for chronic myeloid leukemia. *N Engl J Med*. 2006;355(23):2408-2417.
2. Bower H, Björkholm M, Dickman PW, Höglund M, Lambert PC, Andersson TM. Life Expectancy of Patients With Chronic Myeloid Leukemia Approaches the Life Expectancy of the General Population. *J Clin Oncol*. 2016;34(24):2851-2857.
3. Mahon FX, Réa D, Guilhot J, et al. Discontinuation of imatinib in patients with chronic myeloid leukaemia who have maintained complete molecular remission for at least 2 years: the prospective, multicentre Stop Imatinib (STIM) trial. *Lancet Oncol*. 2010;11(11):1029-1035.
4. Ross DM, Branford S, Seymour JF, et al. Safety and efficacy of imatinib cessation for CML patients with stable undetectable minimal residual disease: results from the TWISTER study. *Blood*. 2013;122(4):515-522.
5. Rousselot P, Charbonnier A, Cony-Makhoul P, et al. Loss of major molecular response as a trigger for restarting tyrosine kinase inhibitor therapy in patients with chronic-phase chronic myelogenous leukemia who have stopped imatinib after durable undetectable disease. *J Clin Oncol*. 2014;32(5):424-430.
6. Chalandon Y, Passweg JR, Schmid C, et al. Outcome of patients developing GVHD after DLI given to treat CML relapse: a study by the Chronic Leukemia Working Party of the EBMT. *Bone Marrow Transplant*. 2010;45(3):558-564.
7. Bocchia M, Defina M, Aprile L, et al. Complete molecular response in CML after p210 BCR-ABL1-derived peptide vaccination. *Nat Rev Clin Oncol*. 2010;7(10):600-603.
8. Biernacki MA, Marina O, Zhang W, et al. Efficacious immune therapy in chronic myelogenous leukemia (CML) recognizes antigens that are expressed on CML progenitor cells. *Cancer Res*. 2010;70(3):906-915.
9. Hung K, Hayashi R, Lafond-Walker A, Lowenstein C, Pardoll D, Levitsky H. The central role of CD4(+) T cells in the antitumor immune response. *J Exp Med*. 1998;188(12):2357-2368.
10. Ossendorp F, Mengedé E, Camps M, Filius R, Melief CJ. Specific T helper cell requirement for optimal induction of cytotoxic T lymphocytes against major histocompatibility complex class II negative tumors. *J Exp Med*. 1998;187(5):693-702.
11. Løvig T, Andersen SN, Thorstensen L, et al. Strong HLA-DR expression in microsatellite stable carcinomas of the large bowel is associated with good prognosis. *Br J Cancer*. 2002;87(7):756-762.
12. Rimsza LM, Roberts RA, Miller TP, et al. Loss of MHC class II gene and protein expression in diffuse large B-cell lymphoma is related to decreased tumor immunosurveillance and poor patient survival regardless of other prognostic factors: a follow-up study from the Leukemia and Lymphoma Molecular Profiling Project. *Blood*. 2004;103(11):4251-4258.

13. Graham SM, Vass JK, Holyoake TL, Graham GJ. Transcriptional analysis of quiescent and proliferating CD34+ human hemopoietic cells from normal and chronic myeloid leukemia sources. *Stem Cells*. 2007;25(12):3111-3120.
14. Affer M, Dao S, Liu C, et al. Gene Expression Differences between Enriched Normal and Chronic Myelogenous Leukemia Quiescent Stem/Progenitor Cells and Correlations with Biological Abnormalities. *J Oncol*. 2011;2011:798592.
15. Cramer-Morales K, Nieborowska-Skorska M, Scheibner K, et al. Personalized synthetic lethality induced by targeting RAD52 in leukemias identified by gene mutation and expression profile. *Blood*. 2013;122(7):1293-1304.
16. Abraham SA, Hopcroft LE, Carrick E, et al. Dual targeting of p53 and c-MYC selectively eliminates leukaemic stem cells. *Nature*. 2016;534(7607):341-346.
17. Subramanian A, Tamayo P, Mootha VK, et al. Gene set enrichment analysis: a knowledge-based approach for interpreting genome-wide expression profiles. *Proc Natl Acad Sci U S A*. 2005;102(43):15545-15550.
18. Reith W, LeibundGut-Landmann S, Waldburger JM. Regulation of MHC class II gene expression by the class II transactivator. *Nat Rev Immunol*. 2005;5(10):793-806.
19. Silacci P, Mottet A, Steimle V, Reith W, Mach B. Developmental extinction of major histocompatibility complex class II gene expression in plasmacytes is mediated by silencing of the transactivator gene CIITA. *J Exp Med*. 1994;180(4):1329-1336.
20. Holling TM, Schooten E, Langerak AW, van den Elsen PJ. Regulation of MHC class II expression in human T-cell malignancies. *Blood*. 2004;103(4):1438-1444.
21. Satoh A, Toyota M, Ikeda H, et al. Epigenetic inactivation of class II transactivator (CIITA) is associated with the absence of interferon-gamma-induced HLA-DR expression in colorectal and gastric cancer cells. *Oncogene*. 2004;23(55):8876-8886.
22. O'Keefe GM, Nguyen VT, Benveniste EN. Class II transactivator and class II MHC gene expression in microglia: modulation by the cytokines TGF-beta, IL-4, IL-13 and IL-10. *Eur J Immunol*. 1999;29(4):1275-1285.
23. Lee YJ, Han Y, Lu HT, et al. TGF-beta suppresses IFN-gamma induction of class II MHC gene expression by inhibiting class II transactivator messenger RNA expression. *J Immunol*. 1997;158(5):2065-2075.
24. Naka K, Hoshii T, Muraguchi T, et al. TGF-beta-FOXO signalling maintains leukaemia-initiating cells in chronic myeloid leukaemia. *Nature*. 2010;463(7281):676-680.
25. Zhang B, Ho YW, Huang Q, et al. Altered microenvironmental regulation of leukemic and normal stem cells in chronic myelogenous leukemia. *Cancer Cell*. 2012;21(4):577-592.
26. DaCosta Byfield S, Major C, Laping NJ, Roberts AB. SB-505124 is a selective inhibitor of transforming growth factor-beta type I receptors ALK4, ALK5, and ALK7. *Mol Pharmacol*. 2004;65(3):744-752.
27. Gregory MA, Phang TL, Neviani P, et al. Wnt/Ca2+/NFAT signaling maintains survival of Ph+ leukemia cells upon inhibition of Bcr-Abl. *Cancer Cell*. 2010;18(1):74-87.

28. Quintás-Cardama A, Vaddi K, Liu P, et al. Preclinical characterization of the selective JAK1/2 inhibitor INCB018424: therapeutic implications for the treatment of myeloproliferative neoplasms. *Blood*. 2010;115(15):3109-3117.
29. Steidl C, Shah SP, Woolcock BW, Rui L, Kawahara M, Farinha P, et al. MHC class II transactivator CIITA is a recurrent gene fusion partner in lymphoid cancers. *Nature*. 2011;471(7338):377-381.
30. Wright KL, Ting JP. Epigenetic regulation of MHC-II and CIITA genes. *Trends Immunol*. 2006;27(9):405-412.
31. Morimoto Y, Toyota M, Satoh A, et al. Inactivation of class II transactivator by DNA methylation and histone deacetylation associated with absence of HLA-DR induction by interferon-gamma in haematopoietic tumour cells. *Br J Cancer*. 2004;90(4):844-852.
32. Zhu XS, Linhoff MW, Li G, Chin KC, Maity SN, Ting JP. Transcriptional scaffold: CIITA interacts with NF-Y, RFX, and CREB to cause stereospecific regulation of the class II major histocompatibility complex promoter. *Mol Cell Biol*. 2000;20(16):6051-6061.
33. Beresford GW, Boss JM. CIITA coordinates multiple histone acetylation modifications at the HLA-DRA promoter. *Nat Immunol*. 2001;2(7):652-657.
34. Gallipoli P, Pellicano F, Morrison H, et al. Autocrine TNF- α production supports CML stem and progenitor cell survival and enhances their proliferation. *Blood*. 2013(19);122:3335-3339.
35. Zhang B, Chu S, Agarwal P, et al. Inhibition of interleukin-1 signaling enhances elimination of tyrosine kinase inhibitor treated CML stem cells. *Blood*. Prepublished on Sept 12, 2016, as DOI 10-1182/blood-2015-11-679928.
36. Xie S, Wang Y, Liu J, Sun T, et al. Involvement of Jak2 tyrosine phosphorylation in Bcr-Abl transformation. *Oncogene*. 2001;20(43):6188-6195.
37. Ilaria RLJ, Van Etten RA. P210 and P190(BCR/ABL) induce the tyrosine phosphorylation and DNA binding activity of multiple specific STAT family members. *J Biol Chem*. 1996;271(49):31704-31710.
38. Xie S, Lin H, Sun T, Arlinghaus RB. Jak2 is involved in c-Myc induction by Bcr-Abl. *Oncogene*. 2002;21(47):7137-7146.
39. Neviani P, Harb JG, Oaks JJ, et al. PP2A-activating drugs selectively eradicate TKI-resistant chronic myeloid leukemic stem cells. *J Clin Invest*. 2013;123(10):4144-4157.
40. Dawson MA, Bannister AJ, Göttgens B, et al. JAK2 phosphorylates histone H3Y41 and excludes HP1 α from chromatin. *Nature*. 2009;461(7265):819-822.
41. Ghoreschi K, Laurence A, O'Shea JJ. Janus kinases in immune cell signaling. *Immunol Rev*. 2009;228(1):273-287.
42. Fujiwara H, Melenhorst JJ, El Ouriaghli F, et al. In vitro induction of myeloid leukemia-specific CD4 and CD8 T cells by CD40 ligand-activated B cells gene modified to express primary granule proteins. *Clin Cancer Res*. 2005;11(12):4495-4503.
43. Jiang YZ, Mavroudis D, Dermime S, et al. Alloreactive CD4+ T lymphocytes can exert cytotoxicity to chronic myeloid leukaemia cells processing and presenting exogenous antigen. *Br J Haematol*. 1996;93(3):606-612.

44. Das R, Guan P, Sprague L, et al. Janus kinase inhibition lessens inflammation and ameliorates disease in murine models of hemophagocytic lymphohistiocytosis. *Blood*. 2016;127: 1666-1675.
45. Spoerl S, Mathew NR, Bscheider M, et al. Activity of therapeutic JAK 1/2 blockade in graft-versus-host disease. *Blood*. 2014;123: 3832-3842.
46. Carniti C, Gimondi S, Vendramin A, et al. Pharmacologic Inhibition of JAK1/JAK2 Signaling Reduces Experimental Murine Acute GVHD While Preserving GVT Effects. *Clin Cancer Res*. 2015;21: 3740-3749.
47. Choi J, Cooper ML, Alahmari B, et al. Pharmacologic blockade of JAK1/JAK2 reduces GvHD and preserves the graft-versus-leukemia effect. *PLoS One*. 2014;9: e109799.
48. Chen M, Gallipoli P, DeGeer D, et al. Targeting primitive chronic myeloid leukemia cells by effective inhibition of a new AHI-1-BCR-ABL-JAK2 complex. *J Natl Cancer Inst*. 2013;105: 405-423.
49. Gallipoli P, Cook A, Rhodes S, et al. JAK2/STAT5 inhibition by nilotinib with ruxolitinib contributes to the elimination of CML CD34+ cells in vitro and in vivo. *Blood*. 2014;124: 1492-1501.

Sample No.	Sample ID	Source	Gender	Stage	FISH/t(9:22)	Figures
1	Spirit CML 0762	PB	M	CP	ND	1, 2, 5
2	CML 339	Leukapheresis	F	CP	+	2, 4, 5
3	CML 340	Leukapheresis	M	CP	+*	4, 5
4	CML 341 (Fresh)	Leukapheresis	M	CP	+*	1, 2
5	CML 378	Leukapheresis	F	CP	+	1, 2
6	CML 381	Leukapheresis	F	CP	+	1, 2
7	CML 385	Leukapheresis	M	CP	+*	1, 2
8	CML 388	Leukapheresis	F	CP	+*	1, 2
9	CML 411 (Fresh)	PB	F	CP	ND	4, 5
10	CML 412 (Fresh)	Leukapheresis	M	CP	ND	2, 4, 5
11	CML 441	Leukapheresis	M	CP	+	2
12	CML 442	PB	F	CP	+	4, 5
13	CML 450	PB	M	CP	+	5
14	CML 452	Leukapheresis	M	CP	+	2
15	CML 454 (Fresh)	Leukapheresis	M	CP	ND	3
16	CML 456	Leukapheresis	F	CP	ND	3
17	CML 457 (Fresh)	PB	M	CP	ND	3,4
18	CML 459	Leukapheresis	M	CP	ND	3
19	CML 460 (Fresh)	Leukapheresis	F	CP	ND	1, 3, 4
20	CML 461 (Fresh)	Leukapheresis	F	CP	ND	3, 5

Table 1: Source of clinical samples

PB, peripheral blood; CP, chronic phase; * indicates that the CD34⁺CD38⁻ CML LSCs were also confirmed to be Ph⁺ by FISH after sorting; ND, not determined.

Figure Legends:

Figure 1. MHC-II and CIITA are selectively downregulated in CML stem/progenitor cells.

(A) Ingenuity's Antigen Presentation Pathway overlaid with deregulation in CML versus normal G₀ CD34⁺ cells (n=5 CML, n=2 non-CML). Deregulated members for MHC II- α and II- β (including HLA-DP, DQ, DR) are shown expanded in the bottom left. Significantly downregulated components highlighted in green; (B) GSEA analysis demonstrating significant (FDR \leq 0.15) downregulation of the extended MHC-II gene family in a larger, complementary microarray dataset of G₀/quiescent (left) and dividing (right) CML and normal cells;¹⁴ (C) Heat map showing deregulation (as represented by logFC, see colour scale below heatmap) of the MHC-II gene family across multiple CML populations as compared to corresponding non-CML cells from distinct microarray datasets as indicated.^{14,15} The data derived from the original dataset¹³ is highlighted by the orange bar above the corresponding columns. Average normalized MFI of surface MHC-II expression in primary (D) CD34⁺CD38⁺ CML and non-CML stem/progenitor cells or (E) CD34⁺CD38⁻ CML LSCs and non-CML HSCs cultured for 48h in the presence or absence (untreated (UT)) of IFN γ (100U/mL; n=6 for CD34⁺CD38⁺, n=4 for LSC; \pm SEM). CD34⁺CD38⁻ cells, when stringently gated, represent ~1-5% of bulk CD34⁺ cells and overlap considerably with either Hst^{lo}/Py^{lo} or CD34⁺ CFSE^{max} populations described previously.¹³ (F) Average gene expression of *CIITA* in bulk CD34⁺ CML and non-CML stem/progenitor cells cultured for 48h \pm IFN γ (mean fold change; n=6; \pm SEM). Statistical significance was calculated between UT CML sample and all other samples, and if significant, it is indicated by asterisks above the bars. Additional comparisons between samples are indicated by lines.

Figure 2. MHC-II and CIITA downregulation occurs independently of BCR-ABL kinase activity in CML stem/progenitor cells.

Primary (A) CD34⁺CD38⁺ CML cells and (B) CD34⁺CD38⁻ CML LSCs were cultured for 48h with TKIs (5 μ M NIL, 150 nM DAS, 5 μ M IM) or no drug control (NDC) in the presence or absence of IFN γ . Average normalized MFI of MHC-II expression was determined using flow cytometry (n=5; \pm SEM); (C) *CIITA* expression levels in CD34⁺CD38⁺ CML cells were analyzed by qRT-PCR (n=5; \pm SEM; calibrated to UT NDC sample). Statistical significance was calculated between UT CML sample and all other samples, and if significant, it is indicated by asterisks above the bars.

Figure 3. Extended treatment of CML cells with IM does not normalise MHC-II or CIITA expression. Primary CD34⁺ CML were treated with 5 μ M IM for 7d. Average surface MHC-II expression levels were determined by flow cytometry, normalized to CML UT (d0) sample for (A) CD34⁺CD38⁺ CML cells and (B) CD34⁺CD38⁻ CML LSCs; (C) The average expression of MHC-II encoding genes *HLA-DR*, *HLA-DP* and *HLA-DQ* was determined in CD34⁺ CML cells by qRT-PCR (n=7, calibrated to UT (d0) CML sample). (D) The average gene expression of *CIITA* was determined in CD34⁺ CML cells by qRT-PCR. Statistical significance was calculated between d0 UT CML sample and all other samples, and if significant, it is indicated by asterisks above the bars. Additional comparisons between samples are indicated by lines. NS – not significant.

Figure 4. JAK inhibition elevates MHC-II expression in CML stem/progenitor cells. (A) qRT-PCR of RNA/cDNA generated from either non-CML or CML stem/progenitor samples for IL4 transcripts revealed higher expression levels of IL4 in CML cells. Data is expressed relative to the reference gene HPRT1; Primary (B) CD34⁺CD38⁺ CML cells and (C) CD34⁺CD38⁻ CML LSCs were treated with IFN γ and/or JAK inhibitor (RUX (200 nM)), as indicated. The average MHC-II expression levels were determined by flow cytometry, normalized to CML UT sample. NS – not significant; (D) Representative histograms showing MHC-II expression on CD34⁺CD38⁺ and CD34⁺CD38⁻ CML cells treated with IFN γ and/or RUX as indicated; Primary CD34⁺ CML stem/progenitor cells were treated with IFN γ and/or RUX, as indicated above. Thereafter, 300 cells were sorted for either (E) CD34⁺CD38⁺ CML and (F) CD34⁺CD38⁻ CML LSC populations and the average gene expression of *CIITA* was determined by qRT-PCR (n=6 for CD34⁺CD38⁺, n=3 for LSC; calibrated to UT CML sample). Statistical significance was calculated between UT CML sample and all other samples, and if significant, it is indicated by asterisks above the bars.

Figure 5. Elevated MHC-II expression is associated with an increase in CD34⁺ CML cell immunogenicity. (A) Activated and CTV labelled T cells (MNCs from healthy donors) were co-cultured with bulk CD34⁺ CML cells for 72h (\pm treatment as indicated; Block – 10 μ g/mL anti-pan HLA-class II blocking Ab). Proliferation of the responder cells was measured as a reduction in the MFI of CTV-labelled CD4⁺CD69⁺ T cells (n=6 \pm SEM; n=3 for MHC-II blocking assays); (B) The autocrine or paracrine growth

factor/cytokine signaling pathways that regulate MHC-II expression. Statistical significance was calculated between UT CML sample and all other samples, and if significant, it is indicated by asterisks above the bars.

Figure 1 - Tarafdar et al.

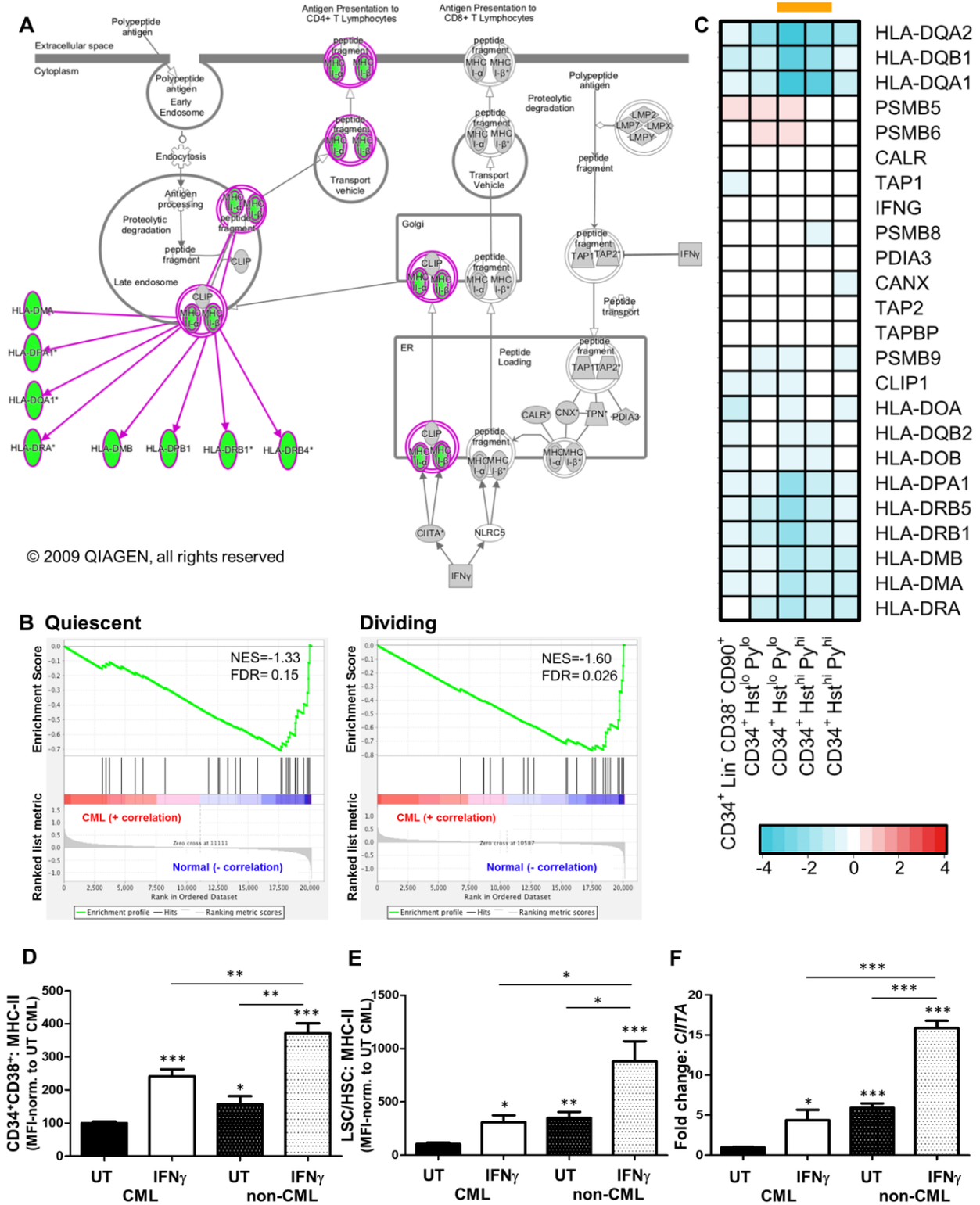


Figure 2 - Tarafdar et al.

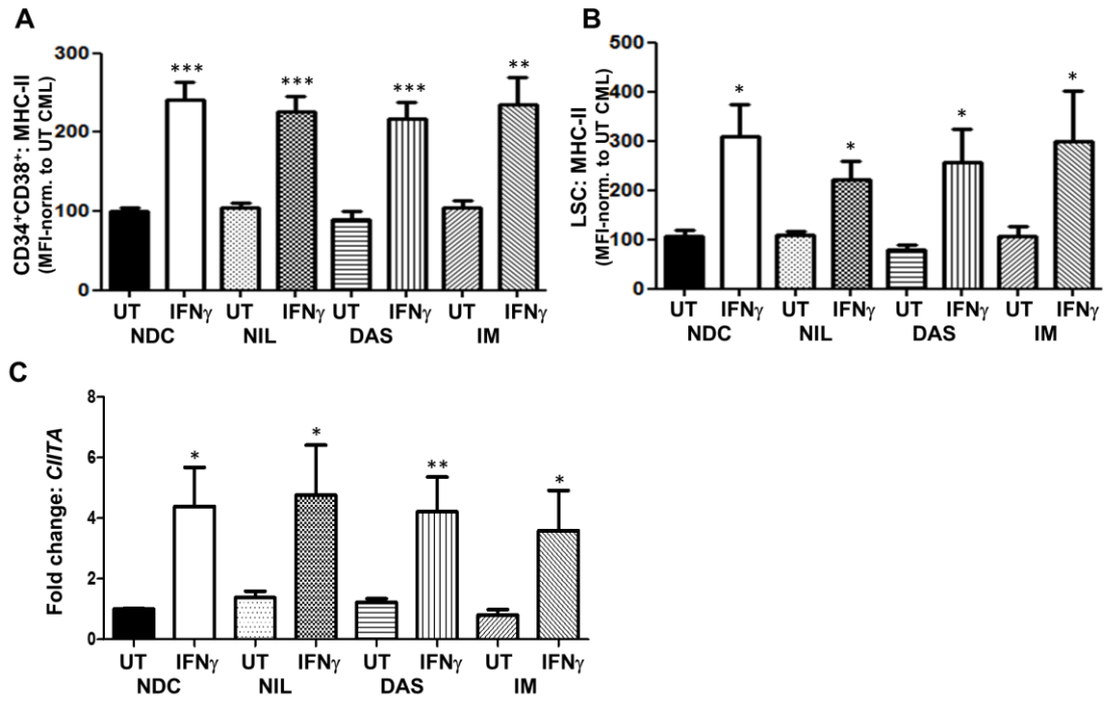


Figure 3 - Tarafdar et al.

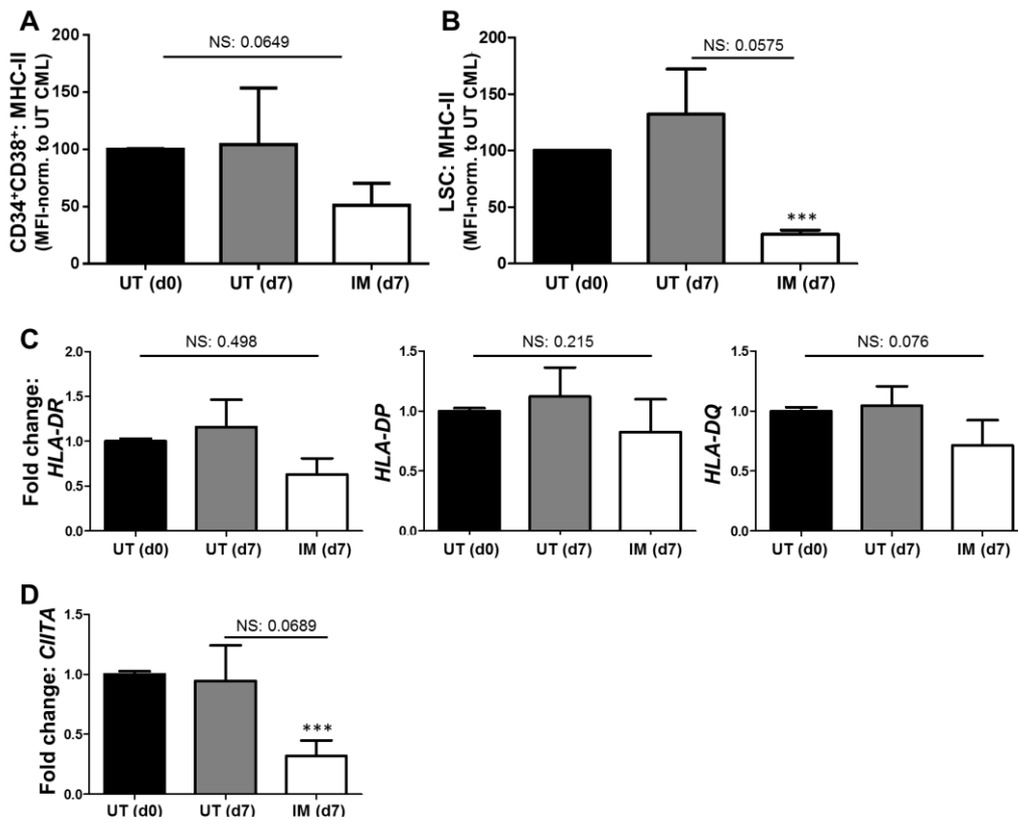


Figure 4 - Tarafdar et al.

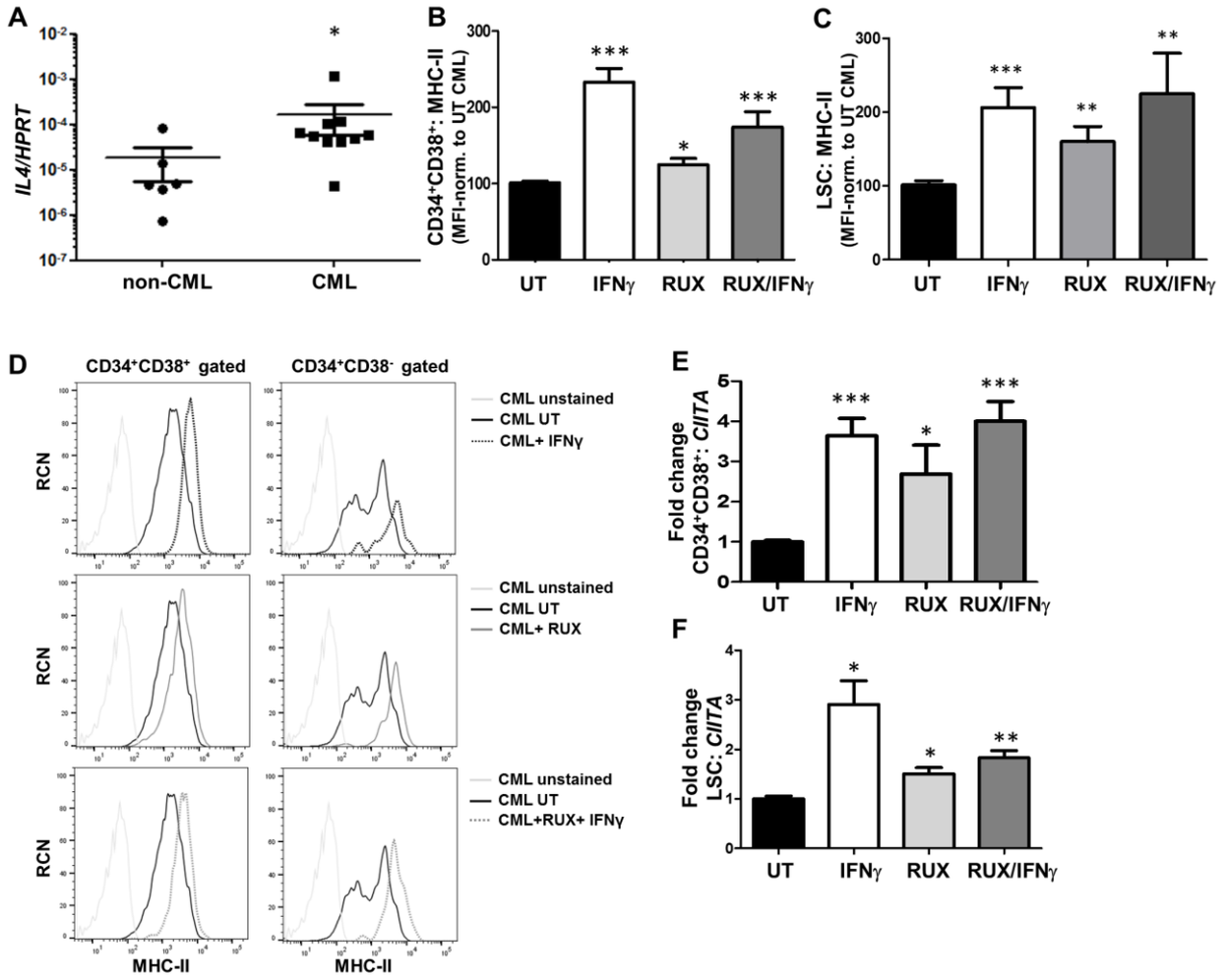
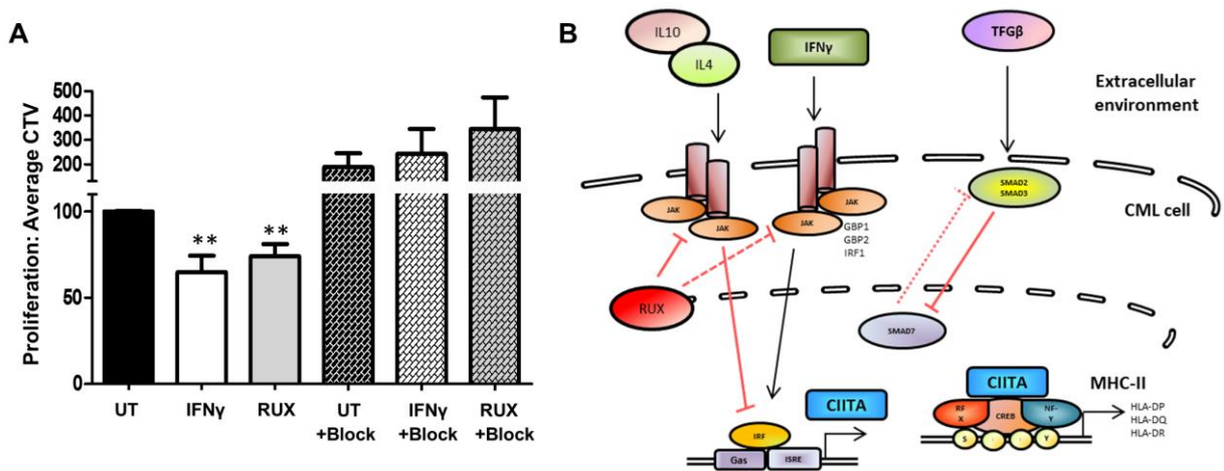


Figure 5 - Tarafdar et al.

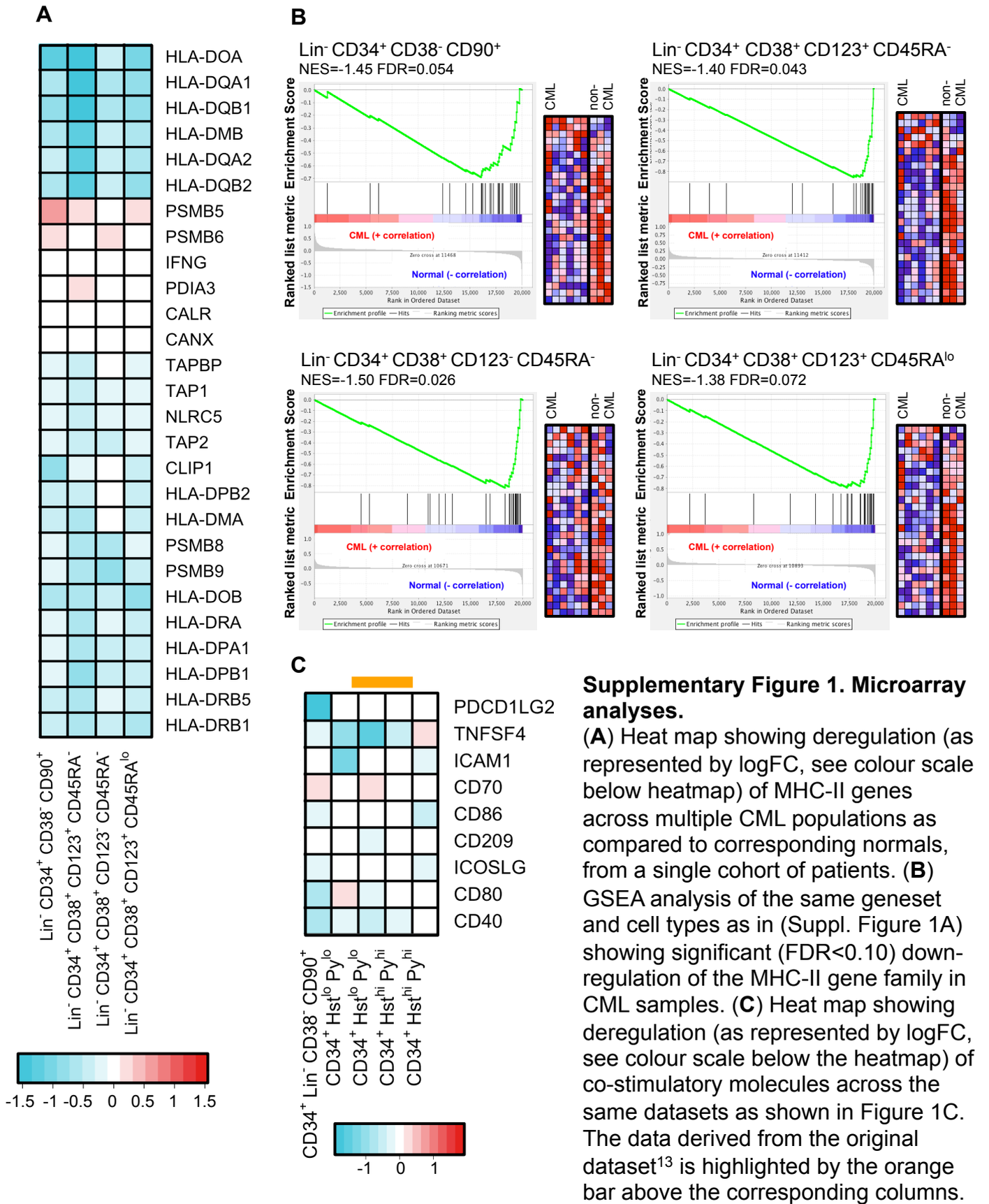


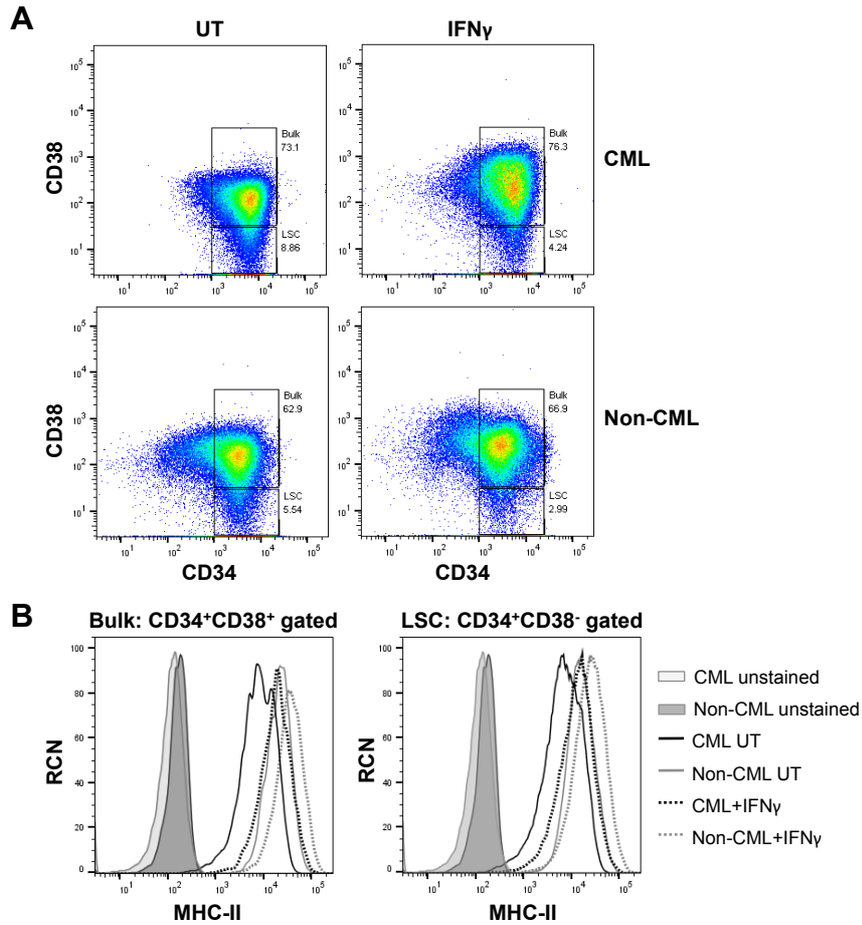
Supplementary Table 1 - Tarafdar et al.

Ref	Accession	Repository	# CML	# normal	Cell type	Figures
15	GSE47927	GEO	6	3	Lin ⁻ CD34 ⁺ CD38 ⁻ Leuk-Norm CD90 ⁺	1C Suppl Fig.1
15	GSE47927	GEO	6	3	Lin ⁻ CD34 ⁺ CD38 ⁺ Leuk-Norm CD123 ⁺ CD45RA ⁻	Suppl Fig.1
15	GSE47927	GEO	6	3	Lin ⁻ CD34 ⁺ CD38 ⁺ Leuk-Norm CD123 ⁻ CD45RA ⁻	Suppl Fig.1
15	GSE47927	GEO	6	3	Lin ⁻ CD34 ⁺ CD38 ⁺ Leuk-Norm CD123 ⁺ CD45RA ^{lo}	Suppl Fig.1
13	E-MTAB-2508	ArrayExpress	5	2	G0/quiescent Leuk-Norm	1C
13	E-MTAB-2508	ArrayExpress	5	3	dividing Leuk-Norm	1C
14	GSE24739	GEO	8	4	G0/quiescent BM-Norm	1C 1B (left)
14	GSE24739	GEO	8	4	dividing BM-Norm	1C 1B (right)
			19	10		

Supplemental Table 1: Summary of the microarray datasets analysed.

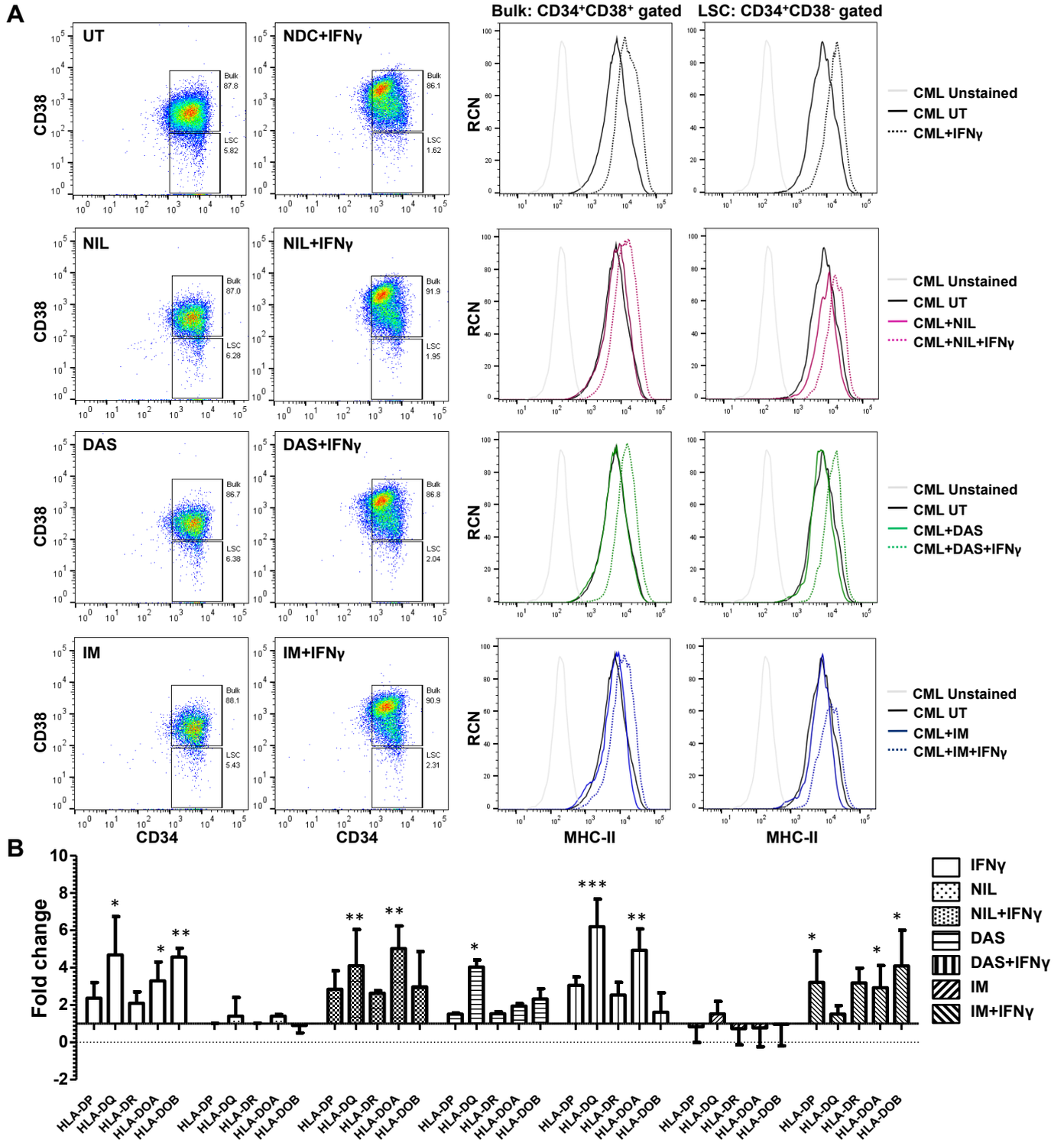
All microarray datasets are summarised with respect to sample size, sorting strategy, source of normal reference cells and the relevant figure, as indicated. Row colour indicates shared samples within a dataset. Leuk-Norm, leukapheresis product of normal donors; BM-Norm, bone marrow sample from normal donors.

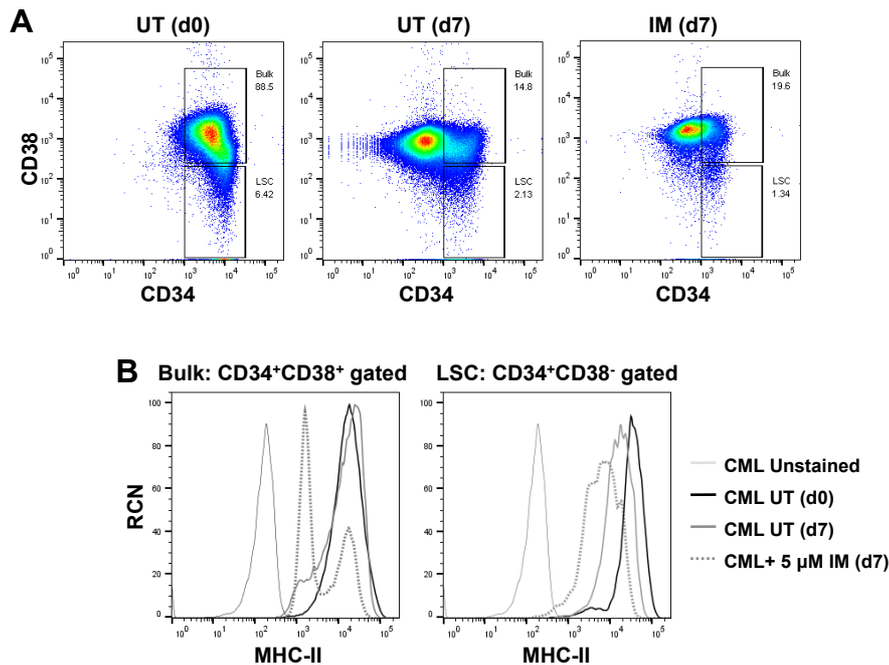




Supplementary Figure 2. Representative flow cytometry plot of MHC-II expression of CML cells and non-CML cells.

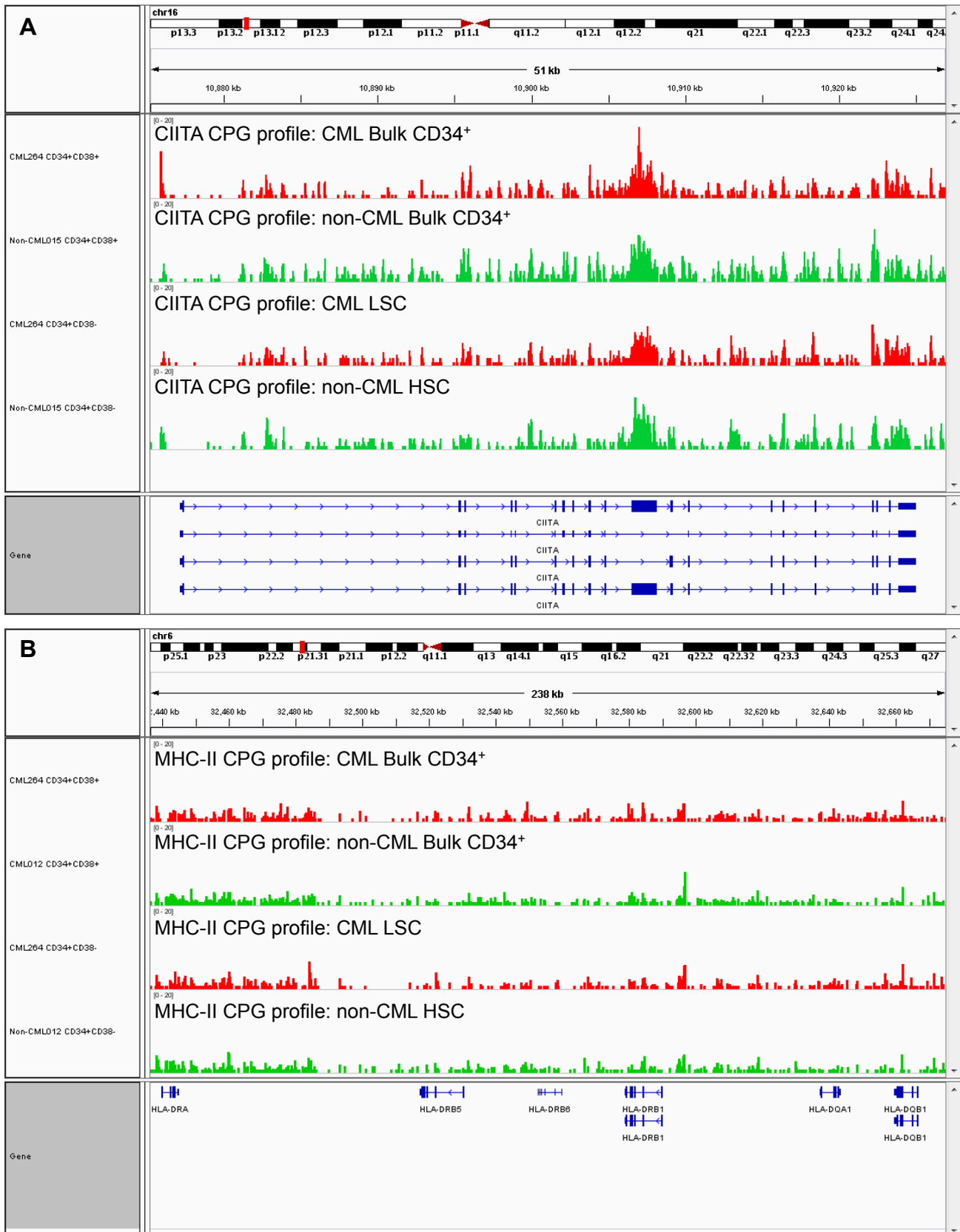
Primary CD34⁺ CML and non-CML stem/progenitor cells were cultured for 48h in the presence or absence (untreated (UT)) of IFN γ (100U/mL), as indicated. **(A)** A representative flow cytometry dot plot is shown for CD34 vs. CD38 staining in CML cells (top) and non-CML stem/progenitor cells (bottom) after culture; **(B)** To assess MHC-II expression, the populations were gated on CD34⁺CD38⁺ CML/non-CML cells or CD34⁺CD38⁻ CML LSCs/HSCs as indicated, compared with unstained cell subsets.



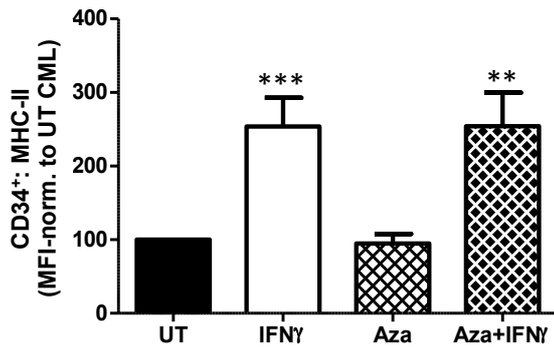


Supplementary Figure 4. Representative flow cytometry plot of MHC-II expression of CML cells in response to 7 days treatment with imatinib.

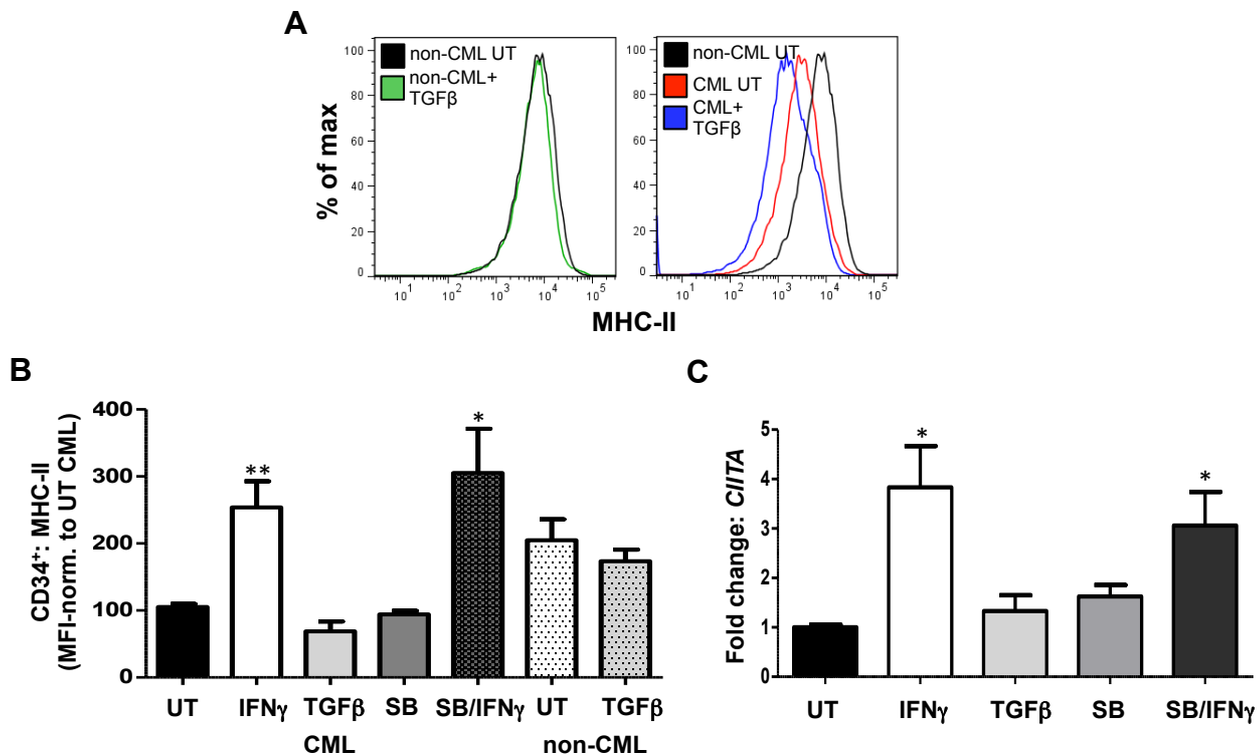
Primary CD34⁺ CML were cultured for 7 days in the presence or absence (untreated (UT)) of 5 μM IM, as indicated. **(A)** A representative flow cytometry dot plot is shown for CD34 vs. CD38 staining in CML cells after culture; **(B)** To assess MHC-II expression, the populations were gated on CD34⁺CD38⁺ (bulk) CML or CD34⁺CD38⁻ CML LSCs as indicated, compared with unstained cell subsets.



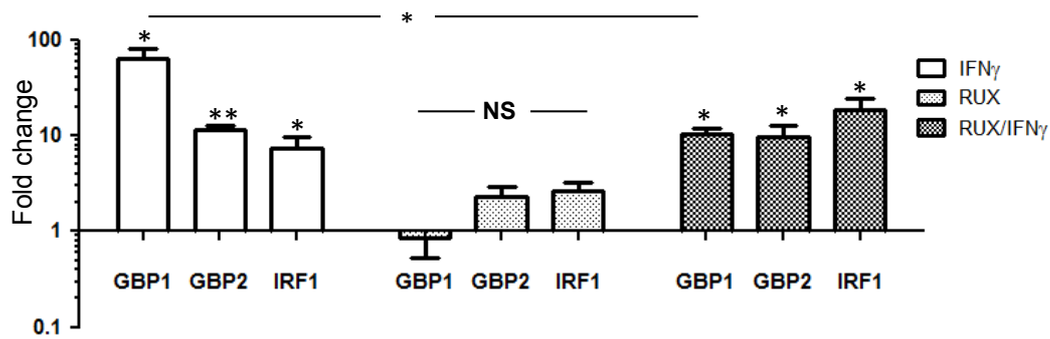
Supplementary Figure 5. Promoter methylation profiling: We profiled the promoters of *CIITA* (A) and *MHC-II* (B), comparing CML to non-CML. We found no differences in the methylation levels in either LSC or bulk populations. Representative profile shown (n=3).



Supplementary Figure 6. MHC-II expression profiles on CD34⁺ CML cells: CML cells were left UT or cultured with IFN γ in the presence or absence of 1 μ M 5-azacytidine. MHC-II expression was profiled using flow cytometry. (Mean fluorescence intensity (MFI); n=4; \pm SEM. All p values generated by paired t test. *, p < 0.05). Statistical significance was calculated between UT CML sample and all other samples, and if significant, it is indicated by asterisks above the bars.

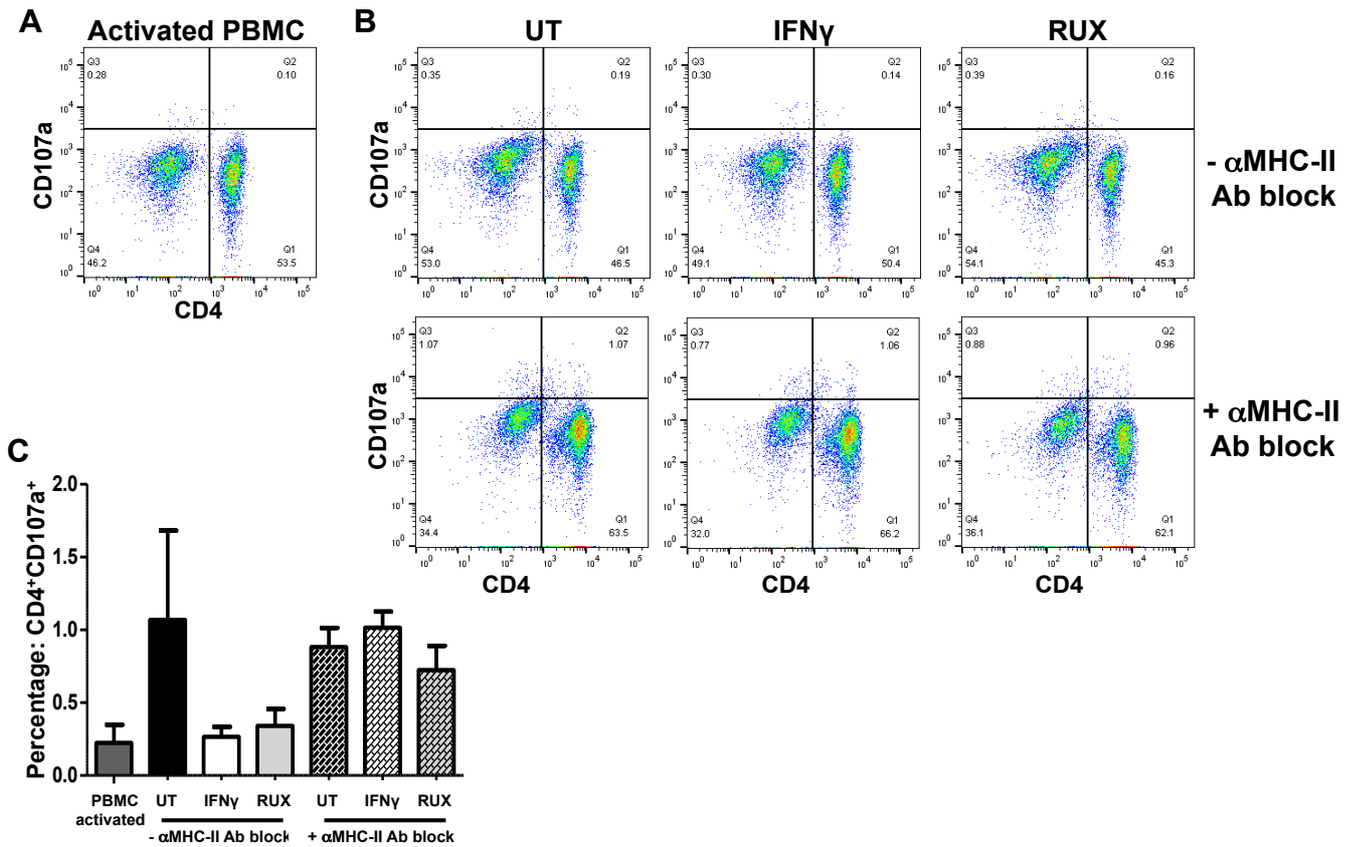


Supplementary Figure 7. Regulation of MHC-II and *CIITA* in the presence of TGF β -modulating agents. (A) MHC-II expression levels were determined by flow cytometry after treatment of CD34⁺ CML and non-CML stem/progenitor samples with 5 ng/mL TGF β for 48h. Cells were co-cultured with murine OP9 stromal cells that provide the extracellular proteins required to activate TGF β . Left - non-CML stem/progenitor sample either UT or TGF β -treated, as indicated. Right – UT non-CML stem/progenitor sample, CD34⁺ CML sample either UT or TGF β -treated, as indicated. Representative overlay histograms are shown; (B) CML stem/progenitor cells were stimulated with IFN γ , TGF β and the TGF β inhibitor SB (10 μ M) as indicated. Average MHC-II expression levels shown determined by flow cytometry, normalized to CML UT sample. (C) CML stem/progenitor cells were treated with IFN γ , TGF β and the TGF β inhibitor (SB) as indicated above. The average gene expression of *CIITA* was determined in CML cells by qRT-PCR (n=3, calibrated to UT CML sample). Statistical significance was calculated between UT CML sample and all other samples, and if significant, it is indicated by asterisks above the bars.



Supplementary Figure 8. Gene expression showing IFN response.

Gene expression profiling showed that the IFN pathway was active at the transcriptional level as the downstream genes *GBP1*, *2* and *IRF1* were upregulated following IFN γ stimulation. There were no significant changes observed with RUX only, however, combination of RUX with IFN γ resulted in a reduced fold change for *GBP1* compared to IFN γ alone as RUX inhibited JAKs downstream of the IFN γ pathway. (Mean fold change relative to UT cells; n=3 CML samples; \pm SEM). Statistical significance was calculated between UT CML sample and all other samples, and if significant, it is indicated by asterisks above the bars. NS – not significant.



Supplementary Figure 9. Analysis of CD107a expression on responder T cells.

(A) FACS plots showing percentage of CD107a on CD4⁺ T cells when activated with beads alone; (B & C) FACS plots showing percentage of CD107a⁺CD4⁺ T cells (B) and a graph showing a percentage average of CD107a⁺CD4⁺ T cells (C; n=3 ± SEM) when co-cultured in the presence of UT, IFN γ or RUX treated CML cells ± anti-MHC-II Ab block for 3 days, as indicated.