Calcium and Calcineurin-NFAT Signaling Regulate Granulocyte-Monocyte Progenitor Cell Cycle via Flt3-L

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Key Words. Flt3 signaling • Cyclosporine A • Hematopoiesis • Myeloid differentiation • Tacrolimus

ABSTRACT

Maintenance of myeloid progenitor cells is controlled by complex regulatory mechanisms and is orchestrated by multiple different transcription factors. Here, we report that the activation of the transcription factor nuclear factor of activated T cells (NFAT) by calcium-sensing protein calcineurin inhibits the proliferation of myeloid granulocyte–monocyte progenitors (GMPs). Myeloid progenitor subtypes exhibit variable sensitivity to induced Ca\textsuperscript{2+} entry and consequently display differential engagement of the calcineurin-NFAT pathway. This study shows that inhibition of the calcineurin-NFAT pathway enhances the proliferation of GMPs both in vitro and in vivo and demonstrates that calcineurin-NFAT signaling in GMPs is initiated by Flt3-L. Inhibition of the calcineurin-NFAT pathway modified expression of the cell cycle regulation genes Cdk4, Cdk6, and Cdkn1a (p21), thus enabling rapid cell cycle progression specifically in GMPs. NFAT inhibitor drugs are extensively used in the clinic to restrict the pathological activation of lymphoid cells, and our data reveal for the first time that these therapies also exert potent effects on the maintenance of the myeloid cell compartment through specific regulation of GMP proliferation.

INTRODUCTION

Regulation of myeloid hematopoiesis plays a key role in the maintenance of innate immune responses. The nuclear factor of activated T cells (NFAT) family of transcription factors has been recently been identified as an important player in the renewal of various myeloid cell subsets [1, 2]. The NFAT family has five members, of which the activation cascade of NFAT1-4 is driven by increased levels of intracellular Ca\textsuperscript{2+}. Ca\textsuperscript{2+} is sensed by calmodulin, which activates calcineurin-mediated dephosphorylation of NFAT resulting in translocation of NFAT into the nucleus [3–5]. Apart from its crucial and well-described role in embryogenesis [6, 7] and T cells [3], the calcineurin-NFAT pathway controls several innate immune functions of dendritic cells (DCs), macrophages, mast cells, megakaryocytes, and osteoclasts [2, 8]. NFAT signaling also regulates apoptosis of terminally differentiated DCs [9], further promoting maintenance of the steady-state. In contrast, during infection, the calcineurin-NFAT pathway is required for effective neutrophil responses to Candida [10] and effective macrophage responses to Leishmania [11].

As well as being involved in myeloid cell functions, NFAT now appears to be important for myeloid compartment development [1] and megakaryopoiesis [12]. Although myeloid cells are present in mice lacking calcineurin-NFAT signaling [13], NFAT deficiency leads to progressive abnormalities including extramedullary hematopoiesis in the spleen and reduced numbers of hematopoietic stem cells (HSCs) in bone marrow (BM) [14]. In humans, there is parallel evidence for involvement of NFAT in the differentiation of immunomobilized CD34\textsuperscript{+} HSCs [15]. However, in each case, the mechanisms underlying these effects of NFAT are unknown. Myeloid hematopoiesis proceeds from HSCs, through multipotent progenitors (MPPs), common myeloid progenitors (CMPs), and finally to GMPs that give rise to fully committed myeloid cells [16]. Within the myeloid lineage, NFAT negatively regulates the differentiation of megakaryocytes [12, 17] and induces development of osteoclasts [18]. Genes regulating the cell cycle have been identified as NFAT targets in T cells, [19] stem cells [20], and in embryonic development and lineage specification [21], but not in the myeloid compartment. Indeed, the main networks...
regulating proliferation and differentiation in hematopoietic progenitors have also been identified [22], but a role for NFAT in the maintenance of myeloid progenitor cells has not previously been reported.

The hematopoietic process is controlled by growth factors and cytokines, including SCF, IL-3, and IL-6 [23], and specifically, in the case of myeloid cells by G-CSF, M-CSF, GM-CSF [24, 25], and Flt3-L [16, 26, 27]. Interestingly, SCF, IL-3, IL-6, and GM-CSF signaling all increase the levels of intracellular Ca$^{2+}$ in HSCs [28–30]. Furthermore, IL-3 and GM-CSF signaling are associated with phospholipase C$\gamma$ (PLC$\gamma$), the main driver of increases in intracellular Ca$^{2+}$ levels [30], and M-CSF [25] and G-CSF [31] phosphorylate PLC$\gamma$2 in BM progenitors, while also being critical determinants of cell lineage commitment. Whether induction of Ca$^{2+}$ release in any of these instances results in NFAT activation is unknown, as are the potential downstream effects on cell function. It has been shown that Flt3 ligation activates PLC$\gamma$2 [26, 32], but the association with Ca$^{2+}$ release, calcineurin and NFAT translocation are currently unknown. The possible link between Flt3/Flt3-L signaling and NFAT induction is particularly intriguing since Flt3-L is a key growth factor for hematopoietic progenitors and also initiates the main signaling pathway responsible for in vivo steady-state differentiation of DCs [33–37]. Flt3 is expressed on MPPs, CMPs, and GMPs [16, 36, 37], while also sustaining progenitor expansion [38, 39], and promoting the growth of colony-forming units (CFU) [40]. Flt3-L is a key cytokine responsible for both development of myeloid cells [33, 34, 41] and promotion of inflammatory immune responses [42]. Furthermore, expression of Flt3-L has been reported on GMPs [36, 37] and Flt3 signaling is known to directly impact on GMP development [43]; however, the mechanism underlying this process remains poorly defined.

Here, we report that NFAT is both present and functional within myeloid progenitors, and directly inhibits the proliferation of GMPs. In addition, we reveal that NFAT mobilization can be triggered by Flt3-L signaling specifically in GMPs, thus providing compelling evidence of a role for NFAT in myeloid hematopoiesis, which has direct implications for the therapeutic inhibition of NFAT in the clinic.

**Flow Cytometry and Sorting of BM Progenitor Populations**

Lineage marker positive BM cells were depleted using biotinylated antibodies (CD45R, CD3e, Gr-1, CD19, NK1.1, TER-119, CD127, and CD11b) and streptavidin-microbeads using a cell separator (AutoMACS-Milenyi, http://www.milenyi-biotec.com). In some experiments, progenitors were further enriched with cKIt beads. Progenitors were labeled with anti-mouse antibodies: PB-c-KIT (CD117), FITC-CD16/32, PE-Flt3 (CD135), AF647-Sca-1 (Ly6A/E), AF700-CD34, and PE-Cy7 streptavidin. The lineage-depleted cells were sorted into progenitor subsets: HSCs (Lin$^-$, c-Kit$, Sca-1$, CD34$, Flt3$), MPPs (Lin$^-$, c-Kit$, Sca-1$, CD34$, Flt3$), CMPs (Lin$^-$, Sca-1$, c-Kit$, CD34$), CD34$, CD16/CD32$), and GMPs (Lin$^-$, Sca-1$, c-Kit$, CD34$, CD16/CD32$), using an Aria II cell sorter.

**PLC$\gamma$ Phosphorylation Analysis by Flow Cytometry**

Lineage-negative cells were cultured for 4 hours and then stimulated for 2 or 10 minutes with rmFlt3-L (1μg/ml). Progenitor subsets were labeled with antibodies, cells were fixed and incubated with anti-pPLC$\gamma$-FITC and total PLC$\gamma$-PE and analyzed using the FlowCellect PLC$\gamma$1 Activation Dual Detection Kit (Merck Millipore, Billerica, MA, www.emdmillipore.com) according to manufacturer’s instructions. Alternatively, cells were fixed and incubated with anti-pPLC$\gamma$-FITC (BD Phosphoflow, BD Bioscience, Mississauga ON, www.bdbiosciences.com), according to the kit instructions, using Phosphoflow Perm Buffer III.

**Intracellular Ca$^{2+}$ Mobilization Assay**

Sorted GMP, CMP, and Lin$^-$ Sca-1$^+$ c-KIT$^+$ BM cells (LSKs), pooled HSCs and MPPs) progenitors (1.5 × 10$^6$ cells per well) were plated in a black 384-well-plate (Perkin Elmer, Waltham, MA, www.perkinelmer.com) and rested at 37°C with 5% CO$_2$ for 3 hours. The cells were then incubated for 45 minutes in dark condition with 20 μl of Hanks’ balanced salt solution (HBSS) containing HEPEs 20 mM, probenecid 2.5 mM and Fluo4-NW (Life Technologies, Invitrogen, Carlsbad, CA, www.lifetechnologies.com). Fluorescence were measured with spectrophotomter Victor$^4$ (Perkin Elmer) (excitation, 485 nm; emission, 535 nm) every second for 180 seconds after injection of the stimuli (diluted in HBSS): Flt3L (4 μg/ml), ionomycin (500 ng/ml), thapsigargin (2 μM). The intracellular calcium chelator BAPTA (10–20 μM, Invitrogen) was added to the culture 1 hour before the incubation with Fluor4-AM. The entire experiments were performed at 37°C. The values (F) were normalized by the first point (Fo) after the injection of the stimuli and the percentage (F/Fo*100) is shown. Alternatively, lineage-depleted BM cells were loaded with Indo-1 AM (2 μM; Life Technologies, Molecular Probes, Carlsbad, CA, www.lifetechnologies.com) by incubation at 37°C for 20 minutes, and Ca$^{2+}$ release monitored by measuring the ratio of signal from Indo-1 (Violet) BP 525/50 versus Indo-1 (Blue) BP 450/50 on a LSR II for 1 minute followed by 4 minutes after adding the trigger Flt3-L (2 μg/ml), ionomycin (500 ng/ml), thapsigargin (2 μM). Cells were kept in a at a temperature of 37°C until the measurement. Kinetics was analyzed using FlowJo software.

**Luciferase Assay**

Lineage-depleted cKIt$^+$ cells or the HSC line were transduced using Cignal Lenti NFAT Reporter (luc) Kit (Quiagen, www.StemCells.com) ©2014 The Authors. STEM CELLS Published by Wiley Periodicals, Inc. on behalf of AlphaMed Press
SAB Biosciences, Venlo, Netherlands, www.sabiosciences.com), using Transcriptional Regulatory Element Sequence GGA GGA AAA ACT GTT TCA TAC AGA AGG CGT according to protocol. Luciferase activity was detected 4 hours after trigger using ONE-Glo Luciferase Assay System (Promega, Madison, WI, www.promega.com).

Cell Culture
BM cells were cultured in IMDM (containing 200 ng/ml mrf3-L (Stem cells), 10% heat-inactivated FCS (Life Technologies), streptomycin 100 mg/ml, and penicillin 100 U/ml) at 3 × 10^6 cells per milliliter. For culture of sorted progenitors, stem cell factor (SCF; 50 ng/ml), IL-6 (20 ng/ml), and IL-3 (10 ng/ml; all R&D Systems, Minneapolis, MN, http://www.rndsystems.com) were added (referred to as HSC medium). Two micrograms per milliliter CsA or 0.2 μg/ml FK506 (Cell Signaling Technology, Danvers, MA, www.cellsignal.com) were added to the cell culture for 30 minutes preceding the addition of Flt3-L/GM-CSF and were maintained throughout the culture.

Isolation and Culture of Primary Human CD34+ Cells
CD34+ umbilical cord blood cells (CB) were obtained from full-term healthy deliveries after informed consent and purified as described previously [44, 45]. Cells were cultured in 24-well plates (2 × 10^5 cells per well) in serum-free X-vivo 15 medium (BioWhittaker, Lonza, Walkersville, MA, www.lonza.com) supplemented with 100 ng/ml Flt3-L in presence or absence of CsA (2 μg/ml). Approval was obtained from the Medical University of Vienna Institutional Review Board.

Immunofluorescence Labeling
GMPs were fixed after sorting or cultured for 24 hours in HSC medium followed by stimulation with ionomycin (500 ng/ml) for 15 minutes. Cells were fixed with paraformaldehyde (2%) before permeabilization in 0.5% Saponin, and blocking for 1 hour with 3% bovine serum albumin. Cells were incubated for 1 hour (37°C) with anti-NFAT2 antibody (10 μg/ml) (Thermo Scientific, Waltham, MA, www.thermoscientific.com) followed by secondary antibody (AF633 goat anti-mouse IgG, Life Technologies, Molecular Probes) at 1 μg/ml and DAPI at 2 μg/ml. Cellular localization of NFAT2 was visualized using an Olympus FV1000 confocal microscope.

CFU Assay
Mouse BM cells were cultured at 4 × 10^4 cells per milliliter in MethoCult M3534 (StemCell Technologies, Vancouver, BC, www.stemcell.com). MethoCult (500 μl per well) were plated in a 12-well suspension culture plate (Greiner, Sigma Aldrich, St. Louis, MO, www.sigmaaldrich.com) and incubated at 37°C in 5% CO2. Colonies were counted under the light microscope after 5–7 days.

Cell Proliferation Analysis
For in vitro bromodeoxyuridine (BrdU) assays, cells were pulsed with 1 mM BrdU for 1 hour. For in vivo BrdU assays, mice were injected with CsA (4 mg/mouse), FK506 (0.4 mg/mouse), or vehicle. At the 48 hours time-point, mice were killed and BM analyzed. BrdU (1.5 mg/mouse) was injected 1 or 18 hours before analysis. BM was lineage-depleted and progenitors populations labeled and gated as described in the sorting strategy above. Cells were fixed and labeled using a BrdU flow cytometry kit (BD Biosciences). Proliferation was also assessed by CFSE dilution (Life Technologies, Molecular Probes), cells were labeled with 2 μM CFSE following the manufacturer’s protocol.

Quantitative Real-Time PCR
Total cellular RNA was extracted by Trizol (Invitrogen) phase separation followed by purification using RNasy Mini/Mini kit (Qiagen), or by using the Arcturus PicoPure RNA Isolation Kit. Reverse transcription was carried out using high-capacity cDNA Reverse Transcription Kit with RNase Inhibitor (Applied Biosystems), or with SuperScript III First Strand Synthesis System for RT-polymerase chain reaction (PCR) (Invitrogen). Real-time PCR was carried out with primers listed in the Supporting Information using GoTaq qPCR Master Mix (Promega).

Microarray Hybridization and Analysis
Total RNA was extracted using a double extraction protocol. ssDNA was prepared, fragmented, and labeled according to the Affymetrix protocol. Fragmented ssDNAs were hybridized to the standard arrays for 17 hours at 45°C; the arrays were then washed and stained using the fluids station and then scanned using GeneChip Scanner 3000. The gene expression data were then filtered for only probes where the associated gene had a valid NCBI Entrez Gene ID to restrict data to well annotated genes. Gene ontology terms were used to identify genes involved in regulation of cell cycle and transcriptional regulation of differentiation and hematopoiesis. These genes were then tested using a series of two-way analysis of variance (ANOVA) to identify genes that differed in their expression levels due to time or treatment. Processing of the data used Accelrys Pipeline Pilot with visualizations in TIBCO Spotfire. All microarray data files are available for free download at the Gene Expression Omnibus (GEO accession number: GSE47208, http://www.ncbi.nlm.nih.gov/geo. Detailed procedure is described in Supporting Information Methods.

Statistical Analysis
Unless specified differently in the legend, all values are shown as means ± SEM. Student’s t-test was used to identify significant differences between groups. For all tests, the 0.05 confidence level was considered statistically significant. In figures, *denotes p < .05, **denotes p < .01, and ***denotes p < .001 in an unpaired Student’s t-test.

RESULTS
Calcineurin-NFAT Inhibitors Cyclosporin A and FK506 Selectively Increase Proliferation and Numbers of GMPs In Vivo
We have previously observed that calcineurin inhibitors enhance myelopoiesis [1]. To identify which hematopoetic progenitors are regulated by calcineurin-NFAT signaling, mice were treated with the calcineurin-NFAT inhibitor drugs CsA or Tacrolimus (FK506). After 48 hours of CsA treatment, we detected no change in the percentage of HSCs, MPPs and CMPs among lineage-negative BM cells, whereas percentages of GMPs were significantly increased (Fig. 1A, 1E). The proliferation rate of GMPs was then assessed by BrdU incorporation and DNA content analysis (Fig. 1B). The percentage and total number of proliferating BrdU-positive...
GMPs (Fig. 1B) was significantly increased in CsA-treated mice. Significant effects of calcineurin-NFAT inhibition on GMP proliferation were also observed with the alternative drug inhibitor FK506 (Fig. 1C–1E). Supporting Information Figure 1A shows the gating strategy used. Increased cell cycle progression upon in vivo administration of both CsA and FK506 was shown. The percentage of hematopoietic progenitors in bone marrow (BM) upon in vivo treatment with Cyclosporine A (CsA) and FK506 is quantified in Figure 1A–1D. Progenitor populations identified as LSKs (lin−, cKit+, Sca-1+), multipotent progenitors (MPPs) (lin−, cKit+, Sca-1+, CD34+, FLt3+), common myeloid progenitors (CMPs) (lin−, cKit+, Sca-1−, CD34+, CD16/32int), and GMPs (lin−, cKit+, Sca-1−, CD34−, CD16/32high). Percentage of progenitors among lineage negative cells from mice treated with CsA (A) and FK506 (C). Detailed analysis of percentage and cell numbers of GMPs from mice treated with CsA (B) and FK506 (D). Mean ± SE from two independent experiments out of five is shown. At least three mice per group (vehicle, CsA, or FK506) were analyzed in each experiment, *p < .05; **, p < .01; and ***, p < .001 in an unpaired Student’s t-test. Figure 1E: Cell cycle analysis in CMPs (lin−, cKit+, Sca-1−, CD34−, CD16/32int) and GMPs (lin−, cKit−, Sca-1−, CD34−, CD16/32high) isolated from mice injected with CsA, FK506, or vehicle and bromodeoxyuridine (BrdU). Representative plot of cell cycle analysis gated for CMPs and GMPs is shown. Dividing cells in S phase (BrdU+) and M phase (DNA content – DAPI+) were further analyzed. Representative experiment is shown, five independent experiments were performed with 3–5 mice per experimental group. Figure 1F: Number of myeloid progenitors in BM. BM cells were plated in methylcellulose and resulting colonies counted 5–7 days later. Total number of colony-forming units (CFUs) (myeloid progenitors) per 10⁴ BM cells is plotted. One representative experiment of three is shown. Mean ± SE is plotted, n = 5; *, p < .05 in an unpaired Student’s t-test. Abbreviations: BrdU, bromodeoxyuridine; CFU, colony-forming units; CMP, common myeloid progenitors; CsA, Cyclosporine A; DAPI, 4',6-diamidino-2-phenylindole; GMP, granulocyte–monocyte progenitor; LSK, lin− Sca1− cKit+ bone marrow cells; MPP, multipotent progenitors; NT, non-treated; PBS, phosphate-buffered saline; PE, phycoerythrin.
and FK506 was specifically observed in GMPs, whereas CMPs were not affected (Fig. 1A, 1C, 1E). Treatment with calcineurin-NFAT inhibitors did not significantly change the total numbers of BM cells or splenocytes (Supporting Information Fig. 1D). Ratio of each subpopulation from individual mice is shown. Representative of two independent experiments, with at least five mice in each is shown. Supporting Information Fig. 1A depicts gating strategy used. Mann-Whitney test (**, p < .01) was used to compare changes in ratios of GMPs and CMPs. Abbreviations: BM, bone marrow; CMP, common myeloid progenitors; GMP, granulocyte–monocyte progenitor.

These data indicate that GMPs but not LSKs, MPPs, or CMPs exhibit increased numbers of CD11b+ myeloid cells, CD4+ T cells, and CD19+ B cells in black were gated and changes in the donor ratio depicted as logarithm of Cnb1flox/floxMx1-cre frequency divided by Cnb1flox/fox control frequency. Log of ratio for common myeloid progenitors (CMPs) and GMPs is shown in red indicating specific change in GMPs. Original injected ratio 1:1 is 0 (black line). Ratio of each subpopulation from individual mice is shown. Representative of two independent experiments, with at least five mice in each is shown. Supporting Information Fig. 3A illustrates the gating strategy used. Figure 2B reveals the ratio between Cnb1flox/floxMx1-cre (CD45.2) and control donor (CD45.1) cells in each gated subset (CD11b+ Gr1+ myeloid/granulocytic cells, CD4+ T cells, CD19+ B cells, CMPs, and GMPs). We observed significant increases in the frequency of GMPs originating from Cnb1flox/floxMx1-cre BM. In contrast, CMPs and B cells remained at the original injected ratio of 1:1 cells derived from the Cnb1-knockout and control BM. Reconstituted mice also exhibited increased numbers of CD11b+ Gr1+ and CD11c+ MHCII+ cells in calcineurin-NFAT-impaired animals, while in contrast the majority of T cells were recruited from the control donor. Similar changes were also reflected in total cell numbers counts (Supporting Information Fig. 1D). Ratio of engrafted donor cells was 1:1 (black line, Fig. 2B). Poly I:C-induced knockout of Cnb1 was confirmed in sorted Cnb1flox/flox Mx1-cre (CD45.2) DCs and CD11b+ Gr1+ myeloid cells by comparing expression levels with control (CD45.1) cells (Supporting Information Fig. 3B). Furthermore, we validated knockout of Cnb1 in magnetic bead-enriched Ly6G+ cells and CD4+ cells obtained from Cnb1flox/floxMx1-cre mice compared with cells isolated from littermate controls (Cnb1flox/floxMx1-wt) (Supporting Information Fig. 3C).

These results suggest that impaired calcineurin signaling confers GMPs with a proliferation advantage that impacts downstream myeloid differentiation.

**CsA Promotes the Proliferation of Flt3-L Stimulated Human CD34+ Umbilical Cord Blood Cells In Vitro**

Flt3-L promotes myeloid and DC differentiation when added to serum-free suspension cultures of human CD34+ hematopoietic progenitor cells [49]. We studied the effects of calcineurin inhibitor addition on the in vitro proliferation of Flt3-

**Calcineurin (Cnb1) Deficiency in HSCs Increases the Frequency of GMPs In Vivo**

To confirm the effects of calcineurin-NFAT signaling on GMP proliferation in vivo, we next generated mice that harbor a conditional knockout of the regulatory subunit of calcineurin (Cnb1, Ppp3r1) [46] in all hematopoietic cells. Cnb1flox/flox animals [47] were crossed with Mx1-cre mice expressing inducible cre-recombinase [48]. Cnb1 knockout in HSCs in Cnb1flox/flox Mx1-cre mice was then induced using poly(I:C) injections. Irradiated recipient mice (heterozygote CD45.1/CD45.2) underwent hematopoietic reconstitution with a 1:1 mix of BM cells from Cnb1flox/floxMx1-cre (CD45.2) and control (CD45.1) donors (Fig. 2A; Supporting Information Fig. 3A, 3B). After full engraftment, Cnb1 knockout was induced with poly(I:C) treatment, as depicted in Figure 2A. The origin and frequency of different progenitors and differentiated cells from Cnb1 knockout and control donors were measured using CD45.1 and CD45.2 expression. The ratio of Cnb1 knockout versus control origin within different cell populations was assessed by flow cytometry. Supporting Information Figure 3A illustrates the gating strategy used. Figure 2B reveals the ratio between Cnb1flox/floxMx1-cre (CD45.2) and control donor (CD45.1) cells in each gated subset (CD11b+ Gr1+ myeloid/granulocytic cells, CD4+ T cells, CD19+ B cells, CMPs, and GMPs). We observed significant increases in the frequency of GMPs originating from Cnb1flox/floxMx1-cre BM. In contrast, CMPs and B cells remained at the original injected ratio of 1:1 cells derived from the Cnb1-knockout and control BM. Reconstituted mice also exhibited increased numbers of CD11b+ Gr1+ and CD11c+ MHCII+ cells in calcineurin-NFAT-impaired animals, while in contrast the majority of T cells were recruited from the control donor. Similar changes were also reflected in total cell numbers counts (Supporting Information Fig. 1D). Ratio of engrafted donor cells was 1:1 (black line, Fig. 2B). Poly I:C-induced knockout of Cnb1 was confirmed in sorted Cnb1flox/flox Mx1-cre (CD45.2) DCs and CD11b+ Gr1+ myeloid cells by comparing expression levels with control (CD45.1) cells (Supporting Information Fig. 3B). Furthermore, we validated knockout of Cnb1 in magnetic bead-enriched Ly6G+ cells and CD4+ cells obtained from Cnb1flox/floxMx1-cre mice compared with cells isolated from littermate controls (Cnb1flox/floxMx1-wt) (Supporting Information Fig. 3C).

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Figure 2. Granulocyte–monocyte progenitors (GMPs) deficient in calcineurin specifically expand in vivo and over control counterparts. (A–B): Bone marrow (BM) chimeric mice were generated using CD45.1 control and Cnb1flox/floxMx1-cre (CD45.2) BM cells mixed in 1:1 ratio as BM graft. After engraftment (8 weeks) the knock out was induced by Poly(I:C) injection. (A): Experimental schema showing BM from Cnb1flox/floxMx1-cre (CD45.2) and control (CD45.1) donors mixed in 1:1 ratio and injected into irradiated hosts (heterozygotes CD45.1/CD45.2). (B): Different cell subpopulations (CD11b+ Gr1+—myeloid cells, CD4+ T cells, and CD19+ B cells) in black were gated and changes in the donor ratio depicted as logarithm of Cnb1flox/floxMx1-cre frequency divided by CD45.1 control frequency. Log of ratio for common myeloid progenitors (CMPs) and GMPs is shown in red indicating specific change in GMPs. Original injected ratio 1:1 is 0 (black line). Ratio of each subpopulation from individual mice is shown. Representative of two independent experiments, with at least five mice in each is shown. Supporting Information Fig. 3A depicts gating strategy. Mann-Whitney test (**, p < .01) was used to compare changes in ratios of GMPs and CMPs. Abbreviations: BM, bone marrow; CMP, common myeloid progenitors; GMP, granulocyte–monocyte progenitor.

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Figure 3. Cyclosporine A (CsA) promotes the proliferation of Flt3-L dependent human hematopoietic progenitors cells. CD34+ cells were loaded with CFSE dye and cultured for 3 days with Flt3-L in presence or absence of CsA (2 μg/ml). Total cell numbers (A), percentage of divided cells (B), and mean of CFSE (C) are shown. Data from four donors are shown, mean ± SE and individual value for each donor are plotted, *, p < .05 in an unpaired Student’s t-test. Abbreviations: CFSE, carboxyfluorescein succinimidyl ester; CsA, Cyclosporine A.
L-dependent human hematopoietic precursors. CD34⁺ progenitors were cultured in serum-free medium supplemented with Flt3-L in presence or absence of CsA for 3 days. CsA addition increased total cell number (Fig. 3A), percentage of dividing cells (Fig. 3B) as well as increased the rate of CFSE dye dilution (Fig. 3C). Therefore, CsA promotes the Flt3-dependent proliferation initiation of hematopoietic progenitor cells.
Calcineurin-NFAT Inhibitors Selectively Increase GMP Proliferation In Vitro and Global Expression Profiling Reveals NFAT as a Cell Cycle Regulator in Progenitor Cells

To identify the downstream processes activated by calcineurin-NFAT signaling during Flt3-L-driven differentiation of myeloid progenitors, we performed a global gene expression analysis on cKIT⁺-enriched lineage-negative primary BM cells. Pooled progenitor populations (predominantly HSCs, MPPs, CMPs, and GMPs) were cultured for 24 or 48 hours in HSC medium with Flt3-L, in the presence or absence of CsA or FK506. Differentially expressed genes (DEG) were then identified within the groups: regulation of cell cycle (GO:0007049), regulation of cell differentiation (GO:0045595), and hematopoiesis (GO:0030097) (Fig. 4A; Supporting Information Table 1). The GO processes and pathway enrichment analysis are shown (Supporting Information Fig. 4A, 4B). We detected increased expression of genes controlling the main cell cycle check points, as well as up-regulation of several genes responsible for myeloid cell differentiation (Fig. 4A).

To confirm the relevance of these trends in CsA treated patients, we have reanalyzed gene expression data comparing PBMCs collected of healthy donors and from stable kidney recipients under immunosuppressant monotherapy [50]. The analysis showed significant activation of hematopoiesis and proliferation (Supporting Information Fig. 5A–5C). Furthermore, we compared our mouse array with the patient obtained data. Supporting Information Figure 6 shows that significant IPA processes induced with CsA treatment are similar in both mice and human cells.

Lineage-determining transcription factors were downregulated, suggesting a lower rate of differentiation in the presence of the inhibitors. In contrast, kinases including Cdk4 and Cdk6 were expressed at increased levels when the calcineurin-NFAT pathway was inhibited. To determine how calcineurin-NFAT inhibitor treatment affected transcription in different progenitor subpopulations, the expression of the DEGs identified by microarray analysis was measured in sorted HSCs, MPPs, CMPs, and GMPs cultured for 24 hours in HSC medium with Flt3-L in the presence or absence of CsA or FK506. The expression of the main kinases regulating the cell cycle G0 checkpoint, Cdk4 and Cdk6, was significantly downregulated during differentiation toward GMPs (Fig. 4B, 4C). Conversely, expression of key inhibitors of cell cycle progression, including Cdkn1a (p21), increased with differentiation toward GMPs (Fig. 4D). Comparable changes in Cdk4, Cdk6, and Cdkn1a expression were observed in the progenitors analyzed immediately after sorting (Supporting Information Fig. 7A–7C). The sorting strategy used and purity achieved is shown in Supporting Information Figure 1A, 1B. These data clearly suggested a decrease in the self-renewal rate of progenitors during the process of differentiation. Figure 4E, 4F show the relative changes in expression of Cdk4 and Cdk6 mRNAs in different progenitor populations following calcineurin-NFAT inhibition. Cdk4 and Cdk6 expression in GMPs remained significantly higher in the presence of inhibitors, and accordingly, expression of Cdkn1a (p21) was downregulated (Fig. 4G). This finding again indicated that GMPs are the sole progenitor target affected by CsA or FK506 treatment.

We next sorted HSCs, MPPs, CMPs, and GMPs from untreated mice and stimulated these cells with Flt3-L in vitro in the presence or absence of CsA or FK506 before assessing their proliferation by flow cytometry 1-2 days later using CFSE staining (Fig. 4H, 4I) or BrdU incorporation and DNA content analysis (Fig. 4J). In vitro, the progenitor populations showed different proliferation rates, with GMPs replicating the least when inhibitors were absent (Fig. 4H). In contrast, when CsA or FK506 was added, GMPs substantially increased their proliferation rate (Fig. 4I). GMPs exclusively responded to CsA and FK506 treatment by significantly increasing their proliferation rate relative to GMPs in control cultures (Fig. 4I). Total cell numbers from these cultured progenitors show similar trends (Supporting Information Fig. 2E).

**Myeloid Progenitors Express Functional NFAT**

We next determined whether Nfat1-4 were expressed at the mRNA level in sorted hematopoietic progenitor cell...
populations of HSCs, MPPs, CMPs, and GMPs. Progenitors were isolated from lineage-depleted BM cells according to the gating strategy shown in Supporting Information Fig. 1. mRNA expression levels of NFAT family members were measured after 24 hours of culture in HSC medium (Fig. 4A). Each progenitor population expressed Nfat1, 2, 4, and Cnb1 (Fig. 5A), which was confirmed in cells analyzed immediately after sorting (Supporting Information Fig. 7A, 7B). Expression of Nfat2 protein in GMPs was confirmed by confocal microscopy (Fig. 5B); Partial nuclear translocation of Nfat2 protein was visualized using an Olympus FV1000 confocal microscope.

**Figure 5.** The calcineurin-nuclear factor of activated T cells (NFAT) pathway is present and functional in myeloid progenitors. (A): Quantitative polymerase chain reaction analysis of Nfat1-4 and calcineurin Cnb1 mRNA expression in hematopoietic stem cells (HSCs), multipotent progenitors (MPPs), common myeloid progenitors (CMPs), and granulocyte–monocyte progenitors (GMPs). Data are presented as mean ± SE from at least three independent experiments, in which bone marrow (BM) cells from at least 15 mice were pooled for sorting. Gating strategy used for flow cytometric sorting of subpopulations of hematopoietic progenitors identified as HSCs (Lin<sup>-</sup>, cKit<sup>+</sup>, Sca-1<sup>+</sup>, CD34<sup>-</sup>, Flt3<sup>-</sup>), MPPs (Lin<sup>-</sup>, cKit<sup>+</sup>, Sca-1<sup>+</sup>, CD34<sup>-</sup>, Flt3<sup>-</sup>), CMPs (Lin<sup>-</sup>, cKit<sup>+</sup>, Sca-1<sup>+</sup>, CD34<sup>-</sup>, CD16/32<sup>-</sup>), and GMPs (Lin<sup>-</sup>, cKit<sup>+</sup>, Sca-1<sup>+</sup>, CD34<sup>-</sup>, CD16/32<sup>+</sup>) is shown at Supporting Information Figure 1. (B): Representative confocal images of sorted GMPs. GMPs were incubated for 1 hour (37°C) with anti-NFAT2 antibody (Thermo Scientific) diluted to 10 μg/ml in the blocking solution, washed three times in phosphate-buffered saline and incubated for 1 hour (37°C) with the secondary antibody (AlexaFluor 633 goat anti-mouse IgG, Molecular Probes) and nuclei were stained with DAPI (original magnification, x200). Images were captured from cells labeled immediately after sorting (first line) or after 24 hours culture in HSC medium (lines 2 and 3). Ionomycin (1 μg/ml for 15 minutes) was used to induce NFAT nuclear translocation. Cellular localization of NFAT2 was visualized using an Olympus FV1000 confocal microscope. (C): Representative graph of fluorometric Ca<sup>2+</sup> mobilization analysis in sorted BM progenitors (LSKs, CMPs, and GMPs). Ionomycin (500 ng/ml) or HBSS were added after 20 seconds of measurement. BM cells from at least 15 mice were pooled for sorting. (D, E): Cells were transduced with a NFAT responsive element luciferase reporter construct. NFAT translocated to the nucleus and started transcription of the reporter gene upon ionomycin stimulation of the HSC line (D) and in lineage-depleted, cKIT<sup>+</sup> purified progenitor cells (E). Data are presented as mean ±SE from one of three independent experiments. *, p < .05; **, p < .01 and ***, p < .001 in an unpaired Student’s t-test. Abbreviations: CMP, common myeloid progenitors; DAPI, 4’,6-diamidino-2-phenylindole; GMP, granulocyte–monocyte progenitor; HBSS, Hanks’ balanced salt solution; HSC, hematopoietic stem cell; IONO, ionomycin; LSK, lin<sup>-</sup> Sca1<sup>+</sup> cKIT<sup>+</sup> bone marrow cells; MPP, multipotent progenitors; NFAT, nuclear factor of activated T cells; NT, non-treated.

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Figure 6. Flt3-L triggers signaling leading to nuclear factor of activated T cells (NFAT) translocation in granulocyte–monocyte progenitors (GMPs). (A): Levels of intracellular Ca\(^{2+}\) measured as Fluo4 fluorescence in sorted Lin\(^{-}\)Sca-1\(^{+}\)cKit\(^{+}\) bone marrow (BM) cells (LSKs), common myeloid progenitors (CMPs), and GMPs. After 20 seconds of measurements cells were triggered with FLT3-L, ionomycin, thapsigargin, or Hanks’ balanced salt solution (HBSS), and Ca\(^{2+}\) levels were assessed for another 5 minutes. (B): Intracellular Ca\(^{2+}\) chelator BAPTA was used to block Ca\(^{2+}\) release induced by FLT3-L in sorted GMPs. (C): Flt3-L induces Ca\(^{2+}\) release in GMPs sorted as Lin\(^{-}\), cKit\(^{+}\), Sca-1\(^{+}\), CD34\(^{+}\), CD16/32\(^{+}\). Graph of Ca\(^{2+}\) release analysis in GMPs from lineage-depleted BM cells. Flt3-L (1\(\mu\)g/ml) was added after 1 minute of measurement followed by 4 minutes Ca\(^{2+}\) release measurement. Representative of three independent experiments, where BM cells from five mice were pooled. (D): Different levels of phospholipase C\(_{\beta}\) (PLC\(_{\beta}\)) phosphorylation (pPLC\(_{\beta}\)) in LSKs (pool of hematopoietic stem cell [HSCs] and multipotent progenitors [MPPs]; Lin\(^{-}\), cKit\(^{+}\), Sca-1\(^{+}\)), CMPs, and GMPs in freshly isolated BM. (E–G): Different levels of pPLC\(_{\beta}\) in progenitor populations before and after Flt3-L administration. Lineage-depleted BM cells were cultured for 4 hours in HSC medium, Flt3-L (1\(\mu\)g/ml) was added 15 minutes before fixing and labeling with antibodies for progenitor markers, PLC\(_{\beta}\) and pPLC\(_{\beta}\). Cells were gated as LSKs (pool of HSCs and MPPs; Lin\(^{-}\), cKit\(^{+}\), Sca-1\(^{+}\)), CMPs (Lin\(^{-}\), cKit\(^{+}\), Sca-1\(^{+}\), CD34\(^{+}\), CD16/32\(^{-}\)), and GMPs (Lin\(^{-}\), cKit\(^{+}\), Sca-1\(^{+}\), CD34\(^{+}\), CD16/32\(^{-}\)). (E, F): Percentage of pPLC\(_{\beta}\) cells and Gm of fluorescence of pPLC\(_{\beta}\) and double labeling of total PLC\(_{\beta}\) with pPLC\(_{\beta}\) upon trigger with Flt3-L are shown. Data are presented as mean ± SE, *, p < .05 and **, p < .01 in an unpaired Student’s t-test. (G): Histogram overlay of pPLC\(_{\beta}\) intensity in GMPs after the Flt3-L trigger. Representative experiment from three independent experiments is shown. (H): Flt3-L induces NFAT translocation in cKit\(^{+}\)-enriched BM cells. BM cells were depleted of lineage cells and enriched with cKit\(^{+}\) beads, maintained in HSC medium and transduced with NFAT reporter constructs. Transfected cells were kept in HSC medium for 48 hours, and 1\(\mu\)g/ml of Flt3-L was added for last 4 hours of culture before nuclear translocation of NFAT reflecting activity of luciferase reporter gene was measured. Data are presented as mean ± SE from one representative of three independent experiments, *, p < .05 in an unpaired Student’s t-test. BM pooled from 10 mice was used for each experiment. (I): Proposed graphical scheme of cell cycle regulation in GMPs. Flt3-L activates PLC\(_{\beta}\) and induce increase in intracellular Ca\(^{2+}\) levels, this further activates calcineurin and NFAT translocation. When calcineurin-NFAT interaction is blocked, expression of cell cycle regulation genes in GMPs is changed to promote further proliferation. Abbreviations: CMP, common myeloid progenitors; CsA, Cyclosporine A; FITC, fluorescein isothiocyanate; GMP, granulocyte–monocyte progenitor; HBSS, Hanks’ balanced salt solution; LSK, Lin\(^{-}\)Sca-1\(^{+}\)cKit\(^{+}\) bone marrow cells; NFAT, nuclear factor of activated T cells; NT, nontreated; PLC, phospholipase C\(_{\beta}\).
occurred after ionomycin-triggered Ca\(^{2+}\) release, indicating that Nfat2 was able to respond functionally in these cells (Fig. 5B). However, treatment with ionomycin (which mobilizes calcium from intracellular stores) resulted in variable levels of intracellular Ca\(^{2+}\) increase across the different progenitor populations (Fig. 5C; Supporting Information Fig. 9A): increases in Ca\(^{2+}\) levels in GMPs were substantially higher than in CMPs and LSKs. This led us to investigate NFAT functionality in more detail within the myeloid lineage using a NFAT luciferase reporter prepared from a previously established HSC line [51]. Treating HSCs with ionomycin or thapsigargin (which inhibits intracellular Ca\(^{2+}\) clearance) confirmed NFAT expression and translocation to the nuclei with consequent dose-dependent induction of luciferase transcription (Fig. 5D). These results were replicated in primary cKIT\(^{+}\)-enriched, lineage-negative cells isolated from BM and transduced with the NFAT reporter construct (Fig. 5E). Taken together these data demonstrate the presence and functionality of the Ca\(^{2+}\)-calcineurin-NFAT pathway at early stages of differentiation of myeloid progenitors.

**Flt3-L Mediates Activation of the Calcineurin-NFAT Pathway in GMPs**

Since calcineurin and NFAT members are expressed in multiple different hematopoietic progenitors, we next assessed whether the main myeloid growth factor Flt3-L might be involved in triggering calcineurin-NFAT signaling to regulate the cell cycle and proliferation rate of these populations. LSKs, CMPs, and GMPs were sorted from BM and loaded with Fluor4-NW, before being stimulated with Flt3-L, ionomycin, or thapsigargin and assessed for changes in intracellular Ca\(^{2+}\) levels using a spectrophotometer (Fig. 6A). We observed that Flt3-L-induced Ca\(^{2+}\) release was effectively blocked by addition of Ca\(^{2+}\) chelator BAPTA (Fig. 6B). We also confirmed our findings by using flow-cytometry to identify changes in intracellular Ca\(^{2+}\) levels in different populations of BM progenitors stained, loaded with Indo-1 and exposed to Flt3-L (Fig. 6C). In agreement with the high sensitivity of GMPs to intracellular Ca\(^{2+}\) release induced by ionomycin treatment (Fig. 5C; Supporting Information Fig. 9A), GMPs also displayed marked increases in intracellular Ca\(^{2+}\) levels upon Flt3-L stimulation (Fig. 6A, 6C), while LSKs and CMPs did not increase the levels of intracellular Ca\(^{2+}\) upon Flt3-L stimulation (Supporting Information Fig. 9B). Flt3-L, which is expressed on GMPs as well as MPPs and CMPs (Supporting Information Fig. 9C), is associated with PLC\(_{7}\) as a possible inducer of Ca\(^{2+}\) entry; so we next measured expression and phosphorylation of PLC\(_{7}\) and PLC\(_{7}\) in myeloid cell progenitors (Fig. 6D–6G). Steady-state phosphorylation of PLC\(_{7}\)(pPLC\(_{7}\)) was observed to increase in parallel with cell differentiation from Lin\(^{-}\), Sca-1\(^{-}\), c-KIT\(^{-}\) progenitors (pooled HSCs and MPPs, referred to as LSK) to GMPs (Fig. 6E–6G) and was further significantly elevated in GMPs by 10 minutes stimulation with Flt3-L (Fig. 6E–6G; Supporting Information Fig. 9D). On contrary, PLC\(_{7}\) seems to be heavily phosphorylated in steady state in freshly isolated GMPs (Fig. 6D), and we have not observed increase in phosphorylation after Flt3-L trigger (Supporting Information Fig. 9E, 9F). To confirm that Flt3-L stimulation results in NFAT translocation in myeloid progenitors, lineage-depleted, cKIT\(^{+}\)-enriched cells were isolated from BM and immediately transduced with the NFAT luciferase reporter construct. Treatment with Flt3-L resulted in significantly increased luciferase activity, thereby reflecting functional NFAT stimulation (Fig. 6H).

In summary, we demonstrate that Flt3-L mobilized Ca\(^{2+}\) in GMPs and increased phosphorylation of PLC\(_{7}\). Moreover, primary cKIT\(^{+}\)-enriched BM cells transduced with an NFAT reporter gene confirmed functional translocation of NFAT upon Flt3-L stimulation. Taken together, these findings show that GMP cell cycle regulation is regulated by Flt3-L activation of PLC\(_{7}\) and induction of Ca\(^{2+}\) entry, which in-turn activates calcineurin and NFAT translocation. Calcineurin-NFAT signaling subsequently modulates the expression of genes that affect the cell cycle progression of GMPs so that these cells can proliferate more rapidly than other progenitor populations (Fig. 6I).

**DISCUSSION**

In the current report, we identified and characterized a new pathway that regulates the cell cycle specifically in GMPs (Fig. 6I). In vivo treatment with CsA or FK506 facilitated the cycling of GMPs, leading to a rapid increase in GMP numbers in the BM. Conditional Cnb1-knockout mice were used to confirm these findings by mixing BM cells from control and Cnb1-knockout mice and engraving them into irradiated recipients. Soon after knockout induction with polyI:C, the majority of GMPs was observed to originate from the calcineurin-impaired donor cells, while the ratio of CMPs remained as injected. These data indicate a specific in vivo effect of calcineurin-NFAT signaling on GMPs proliferative potential. Consistent with these data, sorted, GMPs proliferated less than HSCs, MPPs and CMPs when left untreated during in vitro culture, but addition of CsA or FK506 to these cultures was sufficient to significantly increase GMP proliferation.

To dissect the mechanism of enhanced GMP proliferation under NFAT inhibition, we performed a microarray analysis of gene expression on progenitor cells following brief differentiation with Flt3-L in the presence of calcineurin-NFAT inhibitors. We observed similar changes in PBMCs from CsA treated patients. Global gene expression analysis from mouse BM progenitors revealed marked changes in expression of cell cycle regulation genes as a result of CsA treatment, later validated specifically in GMPs. An important observation was the gradual downregulation of Cdk4 and Cdk6 and upregulation of Cdkn1a (p21) gene expression with the differentiation of cells from HSCs through MPPs, CMPs, and GMPs. This suggests a link between cell differentiation and progression through the cell cycle, mediated by NFAT and triggered by Flt3-L. Our results indicate that the process of slowing down the cell cycle during differentiation is perturbed by calcineurin-NFAT inhibition, particularly at the level of GMPs, both in vitro and in vivo. Therefore, we conclude that the calcineurin-NFAT pathway plays a key role in inhibiting GMP proliferation to regulate myeloid cell differentiation. Our results are supported by other studies in which cell cycle gene regulation was suggested to be important for HSC quiescence [52]. Furthermore, excessive progenitor proliferation led to the exhaustion of HSCs [53, 54], which has recently been linked to NFAT.
signaling [14]. Several cell cycle-regulating genes including cyclin D1 [55], cyclin A2 [56], Cdk4 [20], and Cdk6 [55, 57] are repressed by NFAT in multiple different cell types, and NFAT1-deficient mice exhibit increased levels of Cdk4 [58]. Furthermore, inhibitors of the cell cycle such as Cdk1a (p21) and Cdk2 are also NFAT dependent [59, 60]. These master cell cycle regulators are the main controllers of differentiation during early hematopoietic events as well as in the progression from CMPs to GMPs [22, 52, 61].

Here, we provide evidence that Nfat1, 2, 4, and calcineurin (Cnb1) are ubiquitously expressed in murine HSCs, MPPs, CMPs, and GMPs. Similarly, others have reported NFAT expression in human CD34⁺ immunomobilized progenitors, which hinted at a potential role in myeloid differentiation [15]. We show for the first time that GMPs are exquisitely sensitive to induced Ca²⁺ release which is required for NFAT activation. Furthermore, NFAT was efficiently translocated to the nucleus following activation in ckit⁺-enriched progenitors. Thus, the calcineurin-NFAT pathway is both present and functional in hematopoietic progenitors, especially in GMPs. Several studies have aimed to assess the role of calcineurin-NFAT signaling in myeloid development: Gallo et al. observed small and nonsignificant increases in myeloid cell numbers upon conditional KO induction of Cnb1 in HSCs, concluding that Cnb1 is not necessary for development of the myeloid compartment [13]. In a different experimental setting, we found that progenitors expressing the VIVIT peptide inhibitor of calcineurin-NFAT [62], give rise to increased numbers of myeloid cells. DCs and CD11b⁺Gr1⁺ myeloid cells derived from calcineurin-NFAT-impaired progenitors possessed a substantial developmental advantage over their control counterparts when engrafted into irradiated mice [1]. Similarly, increased numbers of GM-CFU were shown when enriched human CD34⁺ progenitors were treated with FK506 in vitro [63]. Congruent with these observations, we here demonstrated that CsA promotes Flt3-L-dependent initial cell proliferation of purified human CD34⁺ progenitor cells.

Notably, we illustrated that the calcineurin-NFAT pathway in GMPs is triggered by Flt3-L through pPLCγ1 and Ca²⁺ release, thus revealing another mechanism by which this growth factor regulates myeloid development. PLCγ1 phosphorylation followed by NFAT signaling has been shown to be essential in T cell development, activation, and survival [64], while other growth factors such as G-CSF, GM-CSF, and GM-CSF regulate lineage commitment through PLCγ2 [25, 30, 31]. The source of Ca²⁺ driving these processes remains to be elucidated, since PLCγ can induce both exogenous flux as well as endogenous release [65]. The relative contributions made by PLCγ1 and PLCγ2 to intracellular or extracellular Ca²⁺ entry will also require further analysis, since cooperation of Ca²⁺ entry from both compartments is particularly important for NFAT translocation [4, 66].

Flt3-L-induced NFAT signaling leads to cell cycle progression in GMPs that is controlled via a coordinated program of NFAT-regulated changes in expression of cell cycle control genes. Possible roles for other NFAT binding partners have yet to be investigated. In this study, we show that activation of GMPs with Flt3-L induces phosphorylation of PLCγ1 and consequently stimulates intracellular Ca²⁺ release, which does not occur in HSCs, MPPs or CMPs. This Ca²⁺ release initiates translocation of Nfat2 to the nucleus and the transcription of target genes. In addition, we showed that the frequency of cells expressing pPLCγ1, the main regulator of store-operated Ca²⁺ release, is increased in parallel with progenitor differentiation. The role played by Flt3 signaling in normal hematopoiesis was previously thought to be mediated solely by activation of Stat5, Ras/Mapk and PI3K [67]. However, deregulation of Flt3 signaling by activating mutation is present in one third of acute myeloid leukemia cases where expansion of GMPs occurs [26, 32]. The role of pPLCγ1, Ca²⁺, and NFAT signaling with respect to Flt3 signaling in this disease is currently unknown.

Ca²⁺ signaling in immune cells is known to have two qualitatively different outcomes: a short peak of Ca²⁺ release results in immunological synapse formation and granule exocytosis, while the type of prolonged Ca²⁺ signaling induced by growth factors or other cytokines has been shown to enhance NFAT-dependent transcription. This has been reported in T cells [68, 69] and also during embryonic development [70]. The regulatory role of Flt3 in steady state development of GMPs has been suggested [43], leading us to hypothesize that this regulation might involve the NFAT pathway. Flt3 directly regulates HSC quiescence and homeostasis, [67, 70] as well as DC development [33–36]. Flt3 is also expressed in highly proliferating MPPs [22, 71, 72] and the more differentiated CMPs and GMPs [36, 37, 43]. The role of Flt3 signaling is clearly linked to regulation of hematopoietic progenitor numbers, as shown in both Flt3 and Flt3-L KO models [34, 39], but to date, the involvement of Flt3 in the early events of steady state hematopoiesis has not been fully appreciated. Further support for our results comes from the finding that an activating mutation in Flt3 leads to the development of a myeloproliferative disorder [73] that is characterized by increased numbers of GMPs and an accumulation of mature myeloid cells [74–76].

CONCLUSIONS

Our data reveal a novel role for Flt3 signaling in NFAT activation and regulation of myelopoiesis. Investigation of the underlying mechanisms by which NFAT inhibition can increase myelopoiesis uncovered direct regulation of cell cycle control genes Cdk4, Cdk6, and Cdkn1a (p21) by NFAT, specifically in GMPs (Fig. S1). The calcineurin-NFAT pathway is a therapeutic target in multiple conditions including donor organ rejection, graft-versus-host disease, and autoimmune disorders. While T cells are the main cell type known to be subject to CsA and FK506 immunosuppression, here we provide evidence of an important influence of these drugs on myeloid cell hematopoiesis via direct effects on GMP proliferation. Clearly, the full range of effects these drugs exert on the immune system is not fully appreciated, though the prevalence of side effects following their administration is driving new research into their mechanisms of action. Improved knowledge of the roles played by NFAT during myeloid hematopoiesis will provide insight into clinical studies aiming to better understand homeostatic regulation.

ACKNOWLEDGMENTS

We thank J. Lum for microarray processing. C. Phua and S. Nabi for animal handling; I. Low and N. Shadan from SigN flow cytometry facility for cell sorting; H.S. Tay for technical assistance; K. Karjalainen for HSC lines; and L. Robinson with N. McCarthy from Insight Editing London for review of the
paper. A. Wong for the graphical abstract. This work was supported by the BMRC, A*STAR, Singapore.

**AUTHOR CONTRIBUTIONS**

J.F.: designed and conducted the majority of the experiments, analyzed data and wrote the manuscript; C.X.F.L., A.M., and E.V.: designed, performed, and analyzed some experiments; B.T.K.L. and J.C.: performed microarray data analysis; T.Z.: designed and analyzed some experiments and revised the manuscript; A.L.: supervised flow cytometry and sorting experiments; F.Z. and M.P.: supervised microarray processing and analysis; H.S.: supervised some experiments and revised the manuscript; P.R.C.: supervised the project and revised the manuscript. A.M. and C.X.F.L. are joint second authors.

**DISCLOSURE OF POTENTIAL CONFLICTS OF INTEREST**

The authors indicate no potential conflicts of interest.

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