Myeloid Dysregulation in a Human Induced Pluripotent Stem Cell Model of PTPN11-Associated Juvenile Myelomonocytic Leukemia

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In Brief
Using hiPSCs, Mulero-Navarro et al. recapitulate the principal features of JMML and show dysregulation in myeloid cells in the context of NS. Moreover, they identify upregulation of two microRNAs as potential biomarkers in JMML mononuclear cells, and this upregulation distinguishes JMML caused by PTPN11 mutations from other genetic forms of this disease.

Highlights
- hiPSCs from NS/JMML and NS patients with PTPN11 mutations model JMML
- The hiPSC model reveals features of early oncogenesis
- CD33+ myeloid cells from NS/JMML hiPSCs exhibit miR-223 dysregulation
- MicroRNA dysregulation provides a genotype-phenotype association for JMML

Mutations model JMML.

The hiPSC model reveals features of early oncogenesis.

CD33+ myeloid cells from NS/JMML hiPSCs exhibit miR-223 dysregulation.

MicroRNA dysregulation provides a genotype-phenotype association for JMML.
SUMMARY

Somatic \( \textit{PTPN11} \) mutations cause juvenile myelomonocytic leukemia (JMML). Germline \( \textit{PTPN11} \) defects cause Noonan syndrome (NS), and specific inherited mutations cause NS/JMML. Here, we report that hematopoietic cells differentiated from human induced pluripotent stem cells (hiPSCs) harboring NS/JMML-causing \( \textit{PTPN11} \) mutations recapitulated JMML features. hiPSC-derived NS/JMML myeloid cells exhibited increased signaling through STAT5 and upregulation of miR-223 and miR-15a. Similarly, miR-223 and miR-15a were upregulated in 11/19 JMML bone marrow mononuclear cells harboring \( \textit{PTPN11} \) mutations, but not those without \( \textit{PTPN11} \) defects. Reducing miR-223’s function in NS/JMML hiPSCs normalized myelogenesis. MicroRNA target gene expression levels were reduced in hiPSC-derived myeloid cells as well as in JMML cells with \( \textit{PTPN11} \) mutations. Thus, studying an inherited human cancer syndrome with hiPSCs illuminated early oncogenesis prior to the accumulation of secondary genomic alterations, enabling us to discover microRNA dysregulation, establishing a genotype-phenotype association for JMML and providing therapeutic targets.

INTRODUCTION

Juvenile myelomonocytic leukemia (JMML), an aggressive, difficult-to-treat myelodysplastic and myeloproliferative neoplasm of early childhood, is characterized by excessive proliferation of monocytic and granulocytic cells along with dysplastic features. Most JMML cases are associated with somatic gain-of-function (GOF) mutations in components of the RAS/MAPK signal transduction pathway (Yoshida et al., 2012). A minority of cases arise in young children with Noonan syndrome (NS; OMIM: 163950), a genetic disorder with increased RAS/MAPK signaling (Tartaglia and Gelb, 2008). 50% of NS patients and 35% of JMML cases carry GOF \( \textit{PTPN11} \) mutations altering SHP-2. However, the molecular mechanisms through which \( \textit{PTPN11} \) mutations result in deranged myelopoiesis are not well understood.

SHP-2 is a member of the tyrosine phosphatase family and regulates several biological processes, particularly embryogenesis and hematopoietic cell development (Tartaglia and Gelb, 2008). For NS, particular germline mutations in \( \textit{PTPN11} \) underlie JMML, which has clinical features similar to those in children with
Figure 1. Clinical Features of NS/JMML hiPSC-Derived Hematopoietic Cells

(A) Hematopoietic growth factor dose-response curves to GM-CSF. Values are the mean ± SEM for three independent experiments. At 0.01 ng/ml GM-CSF, **p < 0.01 in NS/JMML patient #4 versus CTRL. At 0.1 ng/ml GM-CSF, **p < 0.01 in NS/JMML patients #3 and #4 versus CTRL.

(B) Representative pictures of CFU-GM from study of hypersensitivity to GM-CSF. Scale bars represent 200 μm.

(C) Colony-forming capacity. Bar graphs represent mean ± SEM of three independent experiments. The CTRL group consisted of hiPSC lines, each derived from different healthy individuals.

(D) Representative images of CFU-GM and BFU-E colonies and Wright-Giemsa staining after cytospin. Scale bars represent 200 μm for colonies and 20 μm for cytospin analysis.

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JMML arising from somatic mutations in PTPN11, although with generally better outcomes. Mutations in PTPN11, RAS genes, NFKB1, and CBL are mutually exclusive in JMML, suggesting that one hit in this pathway is sufficient for leukemogenesis (Tartaglia and Gelb, 2008; Pérez et al., 2010; Yoshida et al., 2012).

The ability to induce human induced pluripotent stem cells (hiPSCs) from terminally differentiated cell types such as skin fibroblasts (Takahashi et al., 2007) provides an opportunity to study disease pathogenesis. For hereditary disorders associated with cancer such as NS/JMML, hiPSCs derived from non-cancerous cells permit investigation of the role of the inherited mutations per se, free of secondary or additional genomic alterations that frequently occur during cancer evolution and that might obscure studies performed with hiPSCs derived from the cancer cells (Kotini et al., 2015). Here, we report our findings from modeling JMML using hiPSCs derived from subjects with NS/JMML and NS without leukemia harboring PTPN11 mutations.

RESULTS

Generation of hiPSC Lines and Clinical Manifestations in NS/JMML-Derived Hematopoietic Cells

Using somatic cell reprogramming (Takahashi et al., 2007), we obtained hiPSC lines from skin fibroblasts harboring mutations in PTPN11 from two subjects with NS/JMML. As controls, we used five hiPSC lines derived from skin fibroblasts of unrelated healthy individuals and two subjects with NS, the latter to clarify which perturbations were attributable to the PTPN11 GOF mutations generally versus those specific to JMML pathogenesis (Tables S1 and S2). All PTPN11 mutations resided in the N-SH2 domain, destabilizing SHP-2’s inactive conformation. Pluripotency was established based on in vitro characteristics and in vivo teratoma formation (Figure S1C). All hiPSC lines had a normal karyotype (Figure S1C).

Pluripotency was established based on in vitro characteristics and in vivo teratoma formation (Figure S1C). All hiPSC lines had a normal karyotype (Figure S1C). The NS/JMML megakaryocyte population was similar to the control and NS samples (Figure 1E). The NS/JMML megakaryocyte-macrophage colonies were notably larger (Figure 1D). Cytospin analysis confirmed the granulocyte/monocyte and erythroid cell morphology in the CFU-GM and BFU-E colonies, respectively (Figure 1D).

To characterize the types of myeloid cells differentiating from hiPSC lines, cell populations were analyzed using flow cytometry. We observed a significantly greater number of CD33+ myeloid cells at day 14 in NS/JMML lines compared to those from comparable numbers of control or NS cells, which were attributable to increases among the CFU-erythroid (CFU-E), burst-forming unit-erythroid (BFU-E), and CFU-granulocyte/macrophage (CFU-GM) subtypes (Figure 1C). Moreover the NS/JMML granulocyte-macrophage colonies were notably larger (Figure 1D). The NS/JMML derived cultures compared to controls and NS groups. The NS/JMML megakaryocyte population was similar to the control and NS samples (Figure 1E).

To analyze seminal features of JMML, we used differentiating cells at day 14 after embryoid body (EB) formation. GM-CSF hypersensitivity was observed in the number of colonies formed from NS/JMML hemapoietic cells relative to controls (Figure 1A). Exposure of NS/JMML hematopoietic cells in semi-solid media to low concentrations of GM-CSF (0.01 or 0.1 ng/ml) produced larger colonies compared to controls or NS (Figure 1B). Fluorescence in situ hybridization with a dual probe showed an absence of the BCR-ABL fusion (Figure S3A).

Next, we performed colony-forming unit (CFU) assays in methylcellulose and a cocktail of cytokines to determine the proliferative capacity of the different hematopoietic progenitors. We observed a 10-fold increase in the total number of colonies forming from 1.5 × 10⁴ NS/JMML cells compared to those from comparable numbers of control or NS cells, which were attributable to increases among the CFU-erythroid (CFU-E), burst-forming unit-erythroid (BFU-E), and CFU-granulocyte/macrophage (CFU-GM) subtypes (Figure 1C). Moreover the NS/JMML granulocyte-macrophage colonies were notably larger (Figure 1D). Cytospin analysis confirmed the granulocyte/monocyte and erythroid cell morphology in the CFU-GM and BFU-E colonies, respectively (Figure 1D).

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Genotype-specific dysregulation of genes related to biological processes such as metabolism, development, and proliferation were observed in CD33+ myeloid cells using Gene Ontology (GO) classification (Figure 2E). Gene set enrichment analysis (GSEA) comparing control and NS/JMML CD33+ myeloid cells also revealed significant differences among gene lists related to receptor tyrosine kinase signaling pathways, cell cycle, and myeloid leukemias (Figure 2F). NS and NS/JMML samples differed for gene lists relevant for cell cycle and myeloid leukemia (Figure S4). At the protein level, we found that total and activated (phosphorylated) STAT5, a critical component of the JAK-STAT signaling pathway implicated in JMML pathogenesis (Kotecha et al., 2008), were vastly increased in the NS/JMML CD33+ myeloid cells as compared to control CD33+ myeloid cells (Figure 2G). STAT5 activation in NS myeloid cells was intermediate. ERK activation was increased basally and after GM-CSF stimulation in the PTPN11-mutant CD33+ myeloid cells, being the highest in NS/JMML CD33+ myeloid cells (Figure 2H). GSEA also revealed upregulation of CDKN1A (p21) and MYC in NS/JMML CD33+ myeloid cells. Although their levels (Figure 2I) were comparable between NS and NS/JMML, contributions from others cell-cycle regulators were suggested (Figure S4).

We assessed apoptosis and proliferation rates in the CD33+ myeloid cells using Annexin V and Ki67 staining, respectively. As compared to controls, NS/JMML CD33+ myeloid cells showed similar early-stage apoptosis but ~8-fold increased proliferation (Figures 2J and 2K). Proliferation of NS CD33+ myeloid cells was increased, but not to the level observed in NS/JMML.

miR-223 and miR-15a Are Dysregulated in PTPN11 Mutant JMML Myeloid Cells

We assessed expression levels of mature microRNAs (miRNAs) relevant to hematopoiesis and cancer in the hiPSC-derived CD33+ myeloid cells (Chen et al., 2004; Ramkissoon et al., 2006; Vasilatou et al., 2010). While the levels of several miRNAs were not different, miR-223 and miR-15a expression levels were increased in NS/JMML 50- and 10-fold, respectively (Figures 3A and 3B). This miRNA upregulation was cell-type specific as levels of those two miRNAs were at most subtly increased in the parental hiPSC lines (Figure 3C).

To confirm the relevance of our in vitro findings for JMML in vivo, miR-223 and miR-15a expression levels were measured in mononuclear cells isolated from the bone marrow of children with JMML harboring PTPN11, KRAS, NTRAS, or CBL mutations, with or without NS, as well as from controls (children in remission from acute leukemia) (Table S3). miR-223 and miR-15a levels were significantly increased among patients with germline or somatic PTPN11 mutations compared to controls (Figure 3D). 11 of 19 JMML bone marrow mononuclear cells with PTPN11 mutations (~58%) showed increased expression of miR-223 and/or miR-15a, eight being increased for both. In contrast, miR-223 and miR-15a levels were not increased in the bone marrow mononuclear cells from patients with JMML with other RAS pathway somatic mutations.

Upregulation of miR-223 Is Necessary and Sufficient for JMML-Associated Abnormal Myelopoiesis

To determine if upregulation of miR-223 and miR-15a were necessary for the abnormal myelopoiesis observed in NS/JMML, we reduced their function using molecular decoys (Mullokandov et al., 2012). hiPSCs infected with decoy lentivirus expressing GFP retained pluripotency (Figure 4A). The majority of cells after hematopoietic differentiation continued to express GFP (Figure 4A, bottom row), indicating the presence of the decoy. In the miR-223 decoy-treated cells, the percentage of CD33+/GFP+ myeloid cells was significantly decreased in NS/JMML, achieving a level comparable to that observed with decoy-untreated control hiPSCs (Figure 4B). miR-15a decoy treatment did not alter the production of CD33+ cells (Figure 4C).

Next, we sought to determine whether reduction of miR-223 rescued the GM-CSF hypersensitivity and hyper-proliferation of the NS/JMML myeloid progenitors. Using a low concentration of GM-CSF (0.01 ng/ml), we observed a significant decrease in the number of colonies arising from the miR-223 decoy-treated NS/JMML cells as compared to untreated ones (Figure 4D). Compared to the control hiPSC-derived cells, the rescue of the GM-CSF hypersensitivity was partial. Using colony-formation assays, we observed a decrease of the CFU-GM colony subtype in miR-223 decoy-treated NS/JMML cells (Figure 4E).

To determine whether upregulation of miR-223 was sufficient to perturb myelopoiesis, control hiPSCs were infected with lentivirus expressing miR-223 (Baccarini et al., 2011). This vector expresses a pri-miR-223 transcript within an intron, which is upstream of a reporter encoding a truncated nerve growth factor
receptor (NGFR). Pluripotency of the successfully infected hiPSCs was confirmed (Figure 4F). Overexpression of miR-223 in control and NS hiPSCs resulted in a 10%–12% increase of the CD33+/NGFR+ population compared to those cells infected with the same lentiviral construct without miR-223 (Figure 4G). Moreover, we observed increased GM-CSF sensitivity and increased numbers of CFU-GM colonies in control hematopoietic cells overexpressing miR-223 (Figures 4H and 4I). Since overexpression of miR-223 alone upregulated myelopoiesis in control hiPSCs, we concluded that PTPN11 mutation-driven upregulation of miR-223 is sufficient to drive this aspect of JMML.

Figure 3. Increased Expression Levels of miR-223 and miR-15a in NS/JMML CD33+ Cells
(A) Expression level of miR-181, miR-128a, miR-17, miR-20a, and miR-106 in CD33+ myeloid cells derived from hiPSCs.
(B) Expression level of miR-223 and miR-15a in CD33+ myeloid cells derived from hiPSCs.
(C) qRT-PCR analysis of miR-223 and miR-25a in hiPSC lines. Expression levels were normalized to U6 and relative to CTRL sample. Bar graphs represent mean ± SEM. Data represent a composite of three separate experiments.

In (A)–(C), the CTRL group consisted of hiPSC lines, each derived from different healthy individuals.

(D) qRT-PCR showed the upregulated expression pattern of miR-223 and miR-15a in bone marrow cells from six controls and 26 JMML patients harboring PTPN11, KRAS, NRAS or CBL mutations, with or without NS (Table S3). The red squares represent JMML patients harboring PTPN11 mutations with NS and black squares represent JMML harboring PTPN11 mutations patients without NS. Dot plots represent mean ± SEM.
Effects of miR-223 and miR-15a Upregulation on mRNA Levels in JMML

One of the major mechanisms by which miRNAs’ reduce protein expression is targeted degradation of mRNAs. To determine relevant targets of miR-223 and miR-15a in JMML, we performed an integrated analysis to look for reduced steady-state expression levels among genes predicted to be their targets in hiPSC-derived CD33⁺ myeloid cells (Figure S5A). Comparisons of expression among control and NS/JMML CD33⁺ myeloid cells revealed a significant number of potential miR-223 and miR-15a mRNA targets with reduced expression levels (Figure S5B). Permutation testing revealed that the number of putative miR-15a mRNA targets with reduced expression level was significantly greater than expected by chance (Figure S5C). Reduced levels of putative miR-223 targets were also observed but did not reach statistical significance. The set of genes predicted to be targets for both miRNAs was the most strikingly enriched.

To choose genes with the highest probability of being true targets of one or both of these miRNAs, calculations of thermodynamic accessibility and gene function were combined (Table S4). The lower steady-state levels of six putative target genes, two each from the miR-223 only, miR-15a only, and miR-223/miR-15 groups, were confirmed using RT-PCR (Figure 5A).

To establish genes that were bona fide targets of miR-223, we leveraged an in vitro system using HEK293 cells stably expressing miR-223 constitutively (called HEK293-miR223 hereafter) (Baccarini et al., 2011). Of note, native HEK293 cells do not express miR-223. We generated constructs containing the 3’ UTR of miR-223 putative targets, RAB12 and FOXO3, including their putative miRNA target and mutated sites, inserted downstream of the firefly luciferase gene (Table SS). Expression of miR-223 with these 3’ UTRs reduced luciferase expression ~50%, which was statistically significant (Figure 5B) and comparable to that observed in a previous study that established LMO2 and CEBP-β as miR-223 targets involved in myeloid cell development (Sun et al., 2010). We also confirmed that LMO2 and CEBP-β steady-state mRNA levels were reduced in NS/JMML CD33+ cells (Figure S5D). Mutations in the miR223-binding sites resulted in significantly less degradation of the 3’ UTR gene targets and sequence-specific suppression (Figure 5B). Taken as a whole, these results indicated that RAB12 and FOXO3 are true targets of miR-223.

We leveraged the identification of bona fide targets of miR-223 to document the functional effects of the miR-223 decoy. Reduction of miR-223 function with the decoy would be predicted to increase the steady-state mRNA levels of miR-223 target genes, which is what was observed for RAB12 and FOXO3 in NS/JMML CD33⁺ myeloid cells (Figure S5E).

Finally, we examined the steady-state expression levels of 13 putative and proven miR-223 and miR-15 target genes in PTPN11 and non-PTPN11 mutant JMML and control bone marrow mononuclear cells. Among those, nine (FOXO3, SPTB, NPM1, WHSC1K1, DICER1, MTL5, RAB12, SMAP1, and SHROOM4) were significantly reduced in PTPN11 mutant JMML (Figures 5C and 5E). Of note, these results suggest that the dysregulation of these genes are specific to PTPN11-associated JMML. Moreover, gene expression was similarly reduced among samples from patients with inherited and somatic PTPN11 mutations, suggesting similar JMML pathogenesis.

DISCUSSION

In general, JMML remains a lethal disorder, and progress in treatment has been hindered by the lack of a robust model with which to study it. Taking this into account, our study represents an important advance in our understanding of JMML pathophysiology, providing a valuable model in the context of RASopathies, for several reasons. First, we used hiPSC technology as an alternative model for studying a myeloproliferative and myelodysplastic disorder, and our study had the key feature that it began with a non-hematopoietic cell type, skin fibroblasts. Although the hiPSCs used for our study harbored an activating RAS/ERK pathway mutation, they should not have selected for secondary genomic alternations contributing to hematopoietic disease. A previous hiPSC study used circulating cancerous cells from two JMML patients (Gandre-Babbe et al., 2013). While valuable, this approach may not capture the earliest stages of JMML leukemogenesis. Here, we were able to establish hiPSC lines from NS and NS/JMML patients with germline mutations in specific residues of PTPN11 and then differentiate them into hematopoietic lineages to recapitulate seminal features of JMML. This study demonstrated that the pre-existing PTPN11 GOF mutations were sufficient to drive key features of this leukemogenesis. Moreover, NS- and JMML-associated PTPN11 missense mutations result in increased SHP-2 phosphatase activity, leading to increased ERK activation (Araki et al., 2004; Fragale et al., 2004). Similarly, ERK hyperactivation was present in our patient-derived hiPSCs as well as in the CD33⁺ myeloid cells differentiated from them. JMML-associated somatic PTPN11 mutations (E76K, D61V, and D61Y) induce hematopoietic progenitor hypersensitivity to GM-CSF (Chan et al., 2005; Mohri et al., 2005). Similarly, our study showed that inherited PTPN11 NS/JMML-associated mutations (D61H and G503R) induced GM-CSF hypersensitivity. In contrast, the two PTPN11 mutations associated only with NS, Y63C and E76D, caused at most minimal GM-CSF hypersensitivity. In addition, the observation of STAT5 hyper-activation in the NS/JMML hiPSC-derived CD33⁺ myeloid cells suggested that increased STAT5 signaling may play a role in NS/JMML as it does in somatically acquired JMML (Kotecha et al., 2008). Our findings were also consistent with the observation that bone marrow cells from children with NS and transient myeloproliferative disorder are less responsive to STAT5 than similar cells from children with JMML (Kotecha et al., 2008). Taken together, our results provide further evidence that there is biological specificity and a severity gradient for the hematological defects underlying NS/JMML, arising from the degree of SHP-2’s destabilization, its interactions with signaling partners, or PTPase-independent roles of SHP-2 (Tartaglia and Gelb, 2008). From a clinical point of view, this finding would suggest that genetic differences in PTPN11 encoded in the germline may explain why only a small percentage of NS patients get JMML.

Second, this study elaborated on the roles of miR-223 and miR-15a and their targets in mediating the JMML phenotype. Over the past decade, the study of miRNAs in leukemias has...
established the importance of these regulators for diagnosis, as prognostic factors, and as therapeutic targets (Mian and Zeleznik-Le, 2010). It is worth noting that the upregulation of miR-223 and miR-15a in nearly 60% of the JMML patients harboring PTPN11 mutations may facilitate subclassification of this disorder. Moreover, JMML outcomes have previously been shown to be worse when somatic PTPN11 mutations were the RAS pathway drivers (Yoshida et al., 2009). Our work provides a specific molecular mechanism that might explain that clinical difference and creates opportunities for developing targeted therapies. Specifically, inhibitors and regulators of miR-223 could be used to treat JMML involving dysregulation of the RAS/MAPK pathway. In addition, miR-223 and/or miR-15a target genes, which were downregulated in the NS/JMML could be used to treat JMML involving dysregulation of the specific molecular mechanism that might explain that clinical difference and creates opportunities for developing targeted therapies. Specifically, inhibitors and regulators of miR-223 could be used to treat JMML involving dysregulation of the RAS/MAPK pathway. In addition, miR-223 and/or miR-15a target genes, which were downregulated in the NS/JMML could be used to treat JMML involving dysregulation of the RAS/MAPK pathway.

In summary, our present study showed dysregulation in myeloid cells derived from NS/JMML hiPSCs and elucidated a molecular mechanism underlying aspects of JMML in the context of NS, identifying upregulation of two miRNAs as potential biomarkers in JMML bone marrow mononuclear cells. Their upregulation distinguished JMML caused by PTPN11 mutations from other genetic forms of the disease.

EXPERIMENTAL PROCEDURES

Generation of hiPSC Lines and Myeloid Differentiation

Dermal biopsies and bone marrow aspirates were taken with informed consent and institutional review board approval. Fibroblasts were reprogrammed based on a protocol using retroviruses expressing separately human c-MYC, OCT3/4, SOX2, and KLF4 (Figure S1; Table S1) (Carvajal-Vergara et al., 2010). To induce differentiation, EBs were generated in six-well, ultra-low-attachment plates in differentiation media consisting of a basic media with different cytokines as described previously (Figure S2) (Strigoiadis et al., 2010; Kennedy et al., 2007). See also Supplemental Experimental Procedures for more details.

Flow Cytometric, Cell Sorting, Apoptosis, and Proliferation Assays

Flow cytometric analyses were performed at day 14 of differentiation on hematopoietic cells harvested from dissociated EBs (Table S2). Cells were stained in PBS and 2% fetal bovine serum (FBS) with primary antibodies against CD45, CD14, CD33, CD11b, CD71, CD235a, and CD41 (BD Pharmingen), and analyzed on an LSR-II (BD Biosciences). Apoptosis and proliferation were quantified as CD33+ cells that were Annexin V+/Death dye- (PerCP-Cy; BD Pharmingen and Blue Pacific Invitrogen) and Ki67+ (BD Pharmingen), respectively. Data were analyzed using FlowJo software, version 7.6.

CFU Assay and Cytospin

For cytokine sensitivity assays, hematopoietic cells (Table S2) were plated at 2 × 10^4 in methylcellulose (MethoCult H4531, STEMCELL Technologies) with escalating concentration of GM-CSF (0.01, 0.1, 1, and 10 ng/ml) (R&D Systems) or 1.5 × 10^5 in methylcellulose containing agar leukocyte conditioned medium (Agar-LCM), formulated to support the growth of granulocyte and macrophage progenitors. The hypersensitivity to GM-CSF was observed in a dose-response curve, represented as percentage of maximal colony formation and calculated by dividing the number of colonies at each GM-CSF concentration by the average number of colonies at GM-CSF 10 ng/ml (Table S2). For decay and overexpression miR-223 assays, the treated hematopoietic cells were plated at 2 × 10^4 with 0.01 ng/ml GM-CSF (Table S2). After 14 days at 37°C in humidified 5% CO2, myeloid colonies were counted. For clonogenic myeloid progenitor assays, MethoCult H4435 Enrich (STEMCELL Technologies) was used. Fourteen days after plating, myeloid and erythroid colonies were counted based on standard guidelines previously described (Nissen-Druey et al., 2003). Cell morphology was assessed using a Cytoscan 2 cyto centrifuge (Thermo Fisher Scientific) and staining with Giemsa (Accustain, Sigma-Aldrich).

Treatment with GM-CSF and Immunoblotting

Flow-sorted CD33+ cells were treated with GM-CSF (1 ng/ml) for 15 min at 37°C (Table S2). The 937 cell line was used as a control for STAT5 phosphorylation assays (10 ng/ml GM-CSF). Cells were lysed with RIPA buffer containing protease and phosphatase inhibitors. Blots were probed with antibodies to p-STAT5 (1:250, BD Pharmingen), STAT5 (1:1,000, Cell Signaling Technology), p-ERK1/2 T202/Y204 (1:2,000, Cell Signaling), ERK1/2 (1:2,500, Santa Cruz Biotechnology), and β-actin (1:10,000, Sigma-Aldrich) (loading control). To measure ERK levels in undifferentiated hiPSCs, the cells were treated with 10 ng/ml hEGF for 0, 5, 15, and 60 min at 37°C.

miRNA qRT-PCR

miRNA expression levels were quantified in total RNA from sorted CD33+ cells isolated using Trizol and MiRnase kit (QIAGEN) (Table S2). TaqMan human miRNA assays were used to quantify mature miRNA expression levels as previously described (Chen et al., 2005). U6 (Applied Biosystems) was used as the endogenous control. Reverse-transcribe reactions contained 10 ng of total RNA, 50 nM reverse transcription (RT) primer, 1x RT buffer, 0.25 mM each of dNTPs, 3.33 U/L MultiScribe reverse transcriptase and 0.25 U/L RNase inhibitor. The 7.5 μl reactions were incubated in an Applied Biosystems 7900 Thermocycler in a 96-well plate for 30 min at 16°C, 30 min at 42°C, and 5 min at 85°C and then held at 4°C. All reverse-transcribe reactions were run in triplicate. Real-time PCR was performed using a standard TaqMan PCR kit protocol on an Applied Biosystems 7900 Thermocycler. The 10 μl PCR included 0.67 μl RT product, 1x TaqMan Universal PC Master Mix, and 0.2 mM TaqMan probe. The reactions were incubated in a 96-well plate at 95°C for 10 min, followed by 40 cycles of 95°C for 15 s and 60°C for 1 min. All reactions were run in triplicate. All TaqMan miRNA assays are available through Applied Biosystems.
Figure 5. miR-223 and miR-15a Target Genes: Validation in hiPSC-Derived CD33+ Cells and Expression Levels in Bone Marrow Cells from JMML Patients

(A) Confirmation of the expression level of selected miR-223 and miR-15a target genes in CD33+ cells using qRT-PCR. Bar graphs represent mean ± SEM (p < 0.05 for all).

(B) Luciferase activity of HEK293T cells or HEK293T cells overexpressing miR-223 transfected with a plasmid encoding luciferase without any 3' UTR sequence (pMIR.Report,Vector) and luciferase upstream of the RAB12 and FOXO3 3' UTR normal and mutated sequences (p < 0.05 for all) (Table S5). Bar graphs represent mean ± SEM.

(C) Gene expression levels of targets for miR-223, miR-15a, and both miRNAs in bone marrow mononuclear cells from JMML patients harboring PTPN11, KRAS, NRAS, and CBL mutations (p < 0.05 for all). Red and black squares represent JMML samples with or without NS, respectively. Dot plots represent mean ± SEM. See also Figure S5.
Decoys and Overexpression Assays for miR-223 and miR-15a

For the decoy assay, we used the “tough decoy” (TuD) design previously described (Mulkokandov et al., 2012). Briefly, AB.pCC.Lsin.PPT.U6.hPGK.GFP.wpre was used as a backbone vector with two target sites for miR-223 or miR-15a expressing EGFP to mark infected hPSCs. Lentivirous production and titration were performed as described previously (Mulkokandov et al., 2012). For the miR-223 overexpression, we used the third-generation self-inactivating lentiviral vector platform. The lentiviral vector (LV) encoded hsa-miR-223 and a truncated nerve growth factor receptor (NGFR) as previously described (Baccarini et al., 2011). hiPSCs were infected for 24 hr and sorted 3 days later for GFP+ and NGFR-PE+ expression for the decoy and overexpression functional assays, respectively. These cells were cultured using the same conditions as the original hiPSCs. The infected cells were differentiated, and flow cytometry, GM-CSF sensitivity, and colony-formation assays were performed and analyzed as described above. In sorted CD33+ cells treated with decoys, RNA was isolated and RT-PCR was performed for target genes as previously described in Supplemental Experimental Procedures (“Reverse Transcription Quantitative and Conventional PCR Analysis” section).

Luciferase Assay in HEK293-miR-223 Cells

To analyze the regulation of the putative target FOXO3 and RAB12 by miR-223, we generated constructs using pMIR-Report-Vector. These vectors express Firefly luciferase and the 3′ UTR sequences of FOXO3 and RAB12 by miR-223 (see sequence details in Table S9). We used the same vector without a 3′ UTR as a negative control. The mutated binding site was generated by directed mutagenesis as indicated the manufacturer’s provider (Invitrogen). 1.5 x 10⁵ HEK293T cells or HEK293T expressing constitutively miR-223 were transfected with these constructs with Lipofectamine 2000 (Invitrogen). For normalization, a plasmid encoding Renilla luciferase was included in the transfection in each condition. Luciferase activity was detected with a Dual Luciferase Reporter kit (Promega) 48 hr after transfection.

Statistical Analysis

For overlapping gene sets and correlation patterns between miRNAs and their targets genes, p values were calculated from different permutation and statistical approaches (n = ≥ 10,000) to assess the overlap significance between different gene sets. Additional statistical analyses were performed with Prism 5.0 (Graph Pad Software) using ANOVA and t test. p values < 0.05 were considered statistically significant. The number of stars indicates the significance level: *p < 0.05, **p < 0.01, ***p < 0.001, and ****p < 0.0001.

SUPPLEMENTAL INFORMATION

Supplemental information includes Supplemental Experimental Procedures, five figures, and five tables and can be found with this article online at http://dx.doi.org/10.1016/j.celrep.2015.09.019.

AUTHOR CONTRIBUTIONS

S.M.-N. established and characterized hPSC lines, designed and performed the experiments, and prepared the manuscript; A.S. performed microarray analysis and miRNA expression; A.C.R. performed microarray analysis and miRNA target prediction; G.Y. performed microarray analysis; D.-F.L. performed GSEA analysis and Southern blotting; J.S. performed Southern blotting; S.P. established and partially characterized hPSC lines; S.L.D. provided technical support and established and characterized the CTRL2 hiPSC line; I.K., K.A.M., C.Y.P., S.Y., and D.P. provided technical support; N.C. performed karyotype and BCR-ABL analyses; L.E. and S.B. performed karyotype analysis and miRNA expression; A.C.R. performed microarray analysis and functional validation; and N.C. performed karyotype and BCR-ABL analyses; L.E. and S.B. performed karyotype analysis and miRNA expression; A.C.R. performed microarray analysis and functional validation.

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