Using patient-derived iPSCs to develop humanized mouse models for chronic myelomonocytic leukemia and therapeutic drug identification, including liposomal clodronate

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Chronic myelomonocytic leukemia (CMML) is an entity of myelodysplastic syndrome/myeloproliferative neoplasm. Although CMML can be cured with allogeneic stem cell transplantation, its prognosis is generally very poor due to the limited efficacy of chemotherapy and to the patient’s age, which is usually not eligible for transplantation. Comprehensive analysis of CMML pathophysiology and the development of therapeutic agents have been limited partly due to the lack of cell lines in CMML and the limited developments of mouse models. After successfully establishing patient’s derived disease-specific induced pluripotent stem cells (iPSCs) derived from a patient with CMML, we utilized these CMML-iPSCs to achieve hematopoietic re-differentiation in vitro, created a humanized CMML mouse model via teratomas, and developed a drug-testing system. The clinical characteristics of CMML were recapitulated following hematopoietic re-differentiation in vitro and a humanized CMML mouse model in vivo. The drug-testing system using CMML-iPSCs identified a MEK inhibitor, a Ras inhibitor, and liposomal clodronate as potential drugs for treating CMML. Clodronate is a drug commonly used as a bisphosphonate for osteoporosis. In this study, the liposomalization of clodronate enhanced its effectiveness in these assays, suggesting that this variation of clodronate may be adopted as a repositioned drug for CMML therapy.

Induced pluripotent stem cells (iPSC) technologies have been applied increasingly across clinical medicine and medical research over the last decade1. iPSCs for the purposes of disease-modeling and regenerative medicine are two applications that have been widely explored. In disease-modeling studies, cells from patients...
are reprogrammed to create disease-specific iPSCs and to recapitulate disease-associated phenotypes. Such disease-specific iPSCs can be used as a model in studies on various diseases, and these iPSCs provide a platform for investigating pathophysiology as well as a method for drug testing.

Hematological, disease-specific iPSCs are divided into two groups: those derived from patients with hereditary diseases and those derived from patients with non-hereditary diseases, including acute myeloid leukemia (AML) and myelodysplastic syndrome (MDS). AML and MDS are difficult to reprogram, although recent studies have reported successful establishment of iPSCs specific to AML and MDS. Applications of these AML and MDS iPSCs demonstrated the possibility of producing disease models and analyzing genetic composition, drug sensitivity, and clonal progression.

Chronic myelomonocytic leukemia (CMML) is a clonal hematopoietic stem cell disorder characterized by absolute monocytosis in the peripheral blood (PB) and myelodysplastic along with myeloproliferative properties in the bone marrow (BM). The disease primarily affects the elderly population, with a median age of diagnosis between 65 and 75 years. Prognosis is generally very poor due to the limited efficacy of chemotherapy and to the fact that for many elderly patients, allogeneic hematopoietic stem cell transplantation is not an applicable option. Recently, the hypomethylating drugs azacitidine and decitabine were reported as treatment options for CMML. However, the effectiveness of these drugs such as azacitidine and decitabine remains limited; with overall responses rates of 30–40% and complete remission rate of only 7–17%

Results

Generation of iPSCs from a CMML patient sample. Primary samples from one CMML patient and two healthy donors were used after obtaining informed consent. A 75-year-old male, without any past medical history, presented himself with a chief complaint of weight loss, and he was referred to our institution for anemia, monocytosis, and identification of blast cells in the peripheral blood. The bone marrow analysis diagnosed him as CMML-1 with proliferative neoplasms, and his chromosomal study revealed a 46 XY, +1, der (1;7) (q10; p10) karyotype. The collected bone marrow samples in the experiment included CD13+, CD14+ monocytes and 4% CD34+ immature cells in his chronic phase (WBC 43,200/μL, Blast 3.0%). These samples were found with EZH2, NRAS, and RUNX1 gene mutations (Supplemental Fig. 1(a)).

CD34+ cells from primary samples of the CMML patient and two healthy control donors were isolated from BM mononuclear cells. OCT3/4, SOX2, KLF4, L-MYC, LIN 28, and shP53 were transduced using episomal vectors (Supplemental Table 1). Eight clones of CMML-iPSCs from this patient with CMML were established using episomal vectors (Fig. 1a). Three stable clones of CMML-iPSCs were obtained. The remaining five clones had differentiation propensity, and they failed to maintain the stable passage cultures of their iPSCs. All three stable clones of CMML-iPSCs were found with 46 XY, +1, der (1;7) (q10; p10), the identical chromosomal abnormality of translocation found in the patient's original cells. Therefore, these three stable clones of CMML iPSCs were selected for further analyses. Three stable clones of CMML-iPSCs and four Normal-iPSCs clones were obtained from the two healthy donors. CMML-iPSCs displayed the same morphology as that of Normal-iPSCs and expressed pluripotency markers, including SSEA-4 and Tra-1-60 (Fig. 1b and Supplemental Fig. 2a). The endogenous expression of ESC-related transcription factors (OCT3/4, SOX2, KLF4, C-MYC, NANOG, REX1, and TERT) was confirmed by reverse transcriptase PCR (RT-PCR) (Fig. 1c and Supplemental Fig. 2b).

The CMML patient sample and the CMML-iPSCs were screened for genetic mutations in CBL, KRAS, NRAS, IDH1, IDH2, DNMT3A, TET2, ASXL1, RUNX1, JAK2, SRSF2, SF3B1, U2AF35, and EZH2. Both the primary patient sample and all CMML-iPSCs had the same mutations in EZH2, RUNX1, and NRAS (Fig. 1d and Supplemental Fig. 1a). Three Normal-iPSCs were characterized by a 46 XY (Fig. 1e and Supplemental Fig. 1d), and Normal-iPSCs derived from another healthy donor were characterized by a 46XX (Supplemental Fig. 2c). Both CMML and Normal-iPSCs developed into teratomas, containing three germ layers (Fig. 1f and Supplemental Fig. 2d). It has been reported that the cytostatic guanine dinucleotides (CpG) in promoter regions, such as NANOG and OCT3/4, are unmethylated when iPSCs are induced. Based on the study by Takahashi and Yamanaka, we attempted to prove whether the established clones had been reprogrammed as iPSCs (Fig. 1g).

The patient's pathogenesis of CMML was recapitulated in vitro in CMML iPSC-derived HPCs. Using the previously reported “iPS-sac” method to induce the differentiation of iPSCs into hematopoietic cells, we generated hematopoietic cells from iPSCs. Small clumps, 1 × 10^5 iPSCs, were transferred to
Figure 1. Generation of CMML patient-derived iPSCs. (a) Protocol for the generation of CMML patient-derived iPSCs. CD34+ cells from patient samples were isolated from BM mononuclear cells. OCT3/4, SOX2, KLF4, L-MYC, LIN28, and shP53 were transduced using episomal vectors under hypoxic conditions in the presence of a Rho kinase (ROCK) inhibitor and butyrate acid. Three clones of CMML iPSCs from one patient with CMML-1 were established. (b) Immunofluorescence staining of pluripotency marker antigens (SSEA-4 and Tra-1-60) in Normal and CMML iPSCs. (c) Semi-quantitative RT-PCR of pluripotency markers. The endogenous expression of pluripotent stem cell-specific genes (OCT3/4, SOX2, KLF4, C-MYC, NANOG, REX1, and TERT) was confirmed. Each of the images is cropped from different gels. (d) EZH2, RUNX1, and NRAS mutations were identified in CMML iPSCs. (e) Representative karyotypes of CMML iPSCs showing derivative chromosome (1;7)(q10;p10), an unbalanced translocation, and Normal-iPSCs. (f) Histological analyses of teratomas from CMML iPSCs. A teratoma with three germ layers, the ectoderm (neural tube), mesoderm (cartilage), and endoderm (intestinal tract), was observed following H&E staining. (g) Bisulfite sequence
analysis of the NANOG gene promoter; the black circles represent methylated CpG, while the white circles represent unmethylated CpG. (b) CMML iPSCs grew rapidly and displayed a five-fold higher proliferation rate compared to control iPSCs (n = 3 independent experiments, \( p < 0.001 \), CMML-iPSCs 2 clones and control iPSCs 2 clones derived from same donor samples, paired two-sided t-test). (i–j) Cell cycle analysis showed a relative increase in CMML iPSCs in the G2/M phase. Statistical analysis was performed (n = 3 independent experiments, \( p < 0.01 \), CMML-iPSCs 1 clones and control iPSCs 1 clones derived from same donor samples, Student’s t test). Horizontal lines represent the means ± s.d.

a dish containing irradiated C3H10T1/2 cells. iPSCs with the C3H10T1/2 cells were cultured in differentiation medium with VEGF, which was refreshed every 3 days for 2 weeks. After 2 weeks, these round, hematopoietic-like CD34+CD43+ HPCs were harvested and sorted by flow cytometry (Fig. 2a). Notably, more CD34+CD43+ HPCs were generated when using CMML-iPSCs as compared to Normal-iPSCs (Fig. 2b and Supplemental Fig. 2f). The CD34+ fraction from the primary BM of the CMML patient was merely 2.3%. However, re-differentiated CD34+CD38-CD90+ HPCs could be expanded effectively in the differentiated system (Fig. 2c). The CD34+CD38-CD90- fraction re-differentiated from CMML iPSCs increased compared to Normal-iPSCs (Fig. 2d).

To investigate the differentiation capacity, 1 × 10^5 normal and CMML iPSC-derived CD34+CD43+ HPCs were seeded on a semi-solid culture medium. The semi-solid culture of CD34+CD43+ HPCs derived from CMML-iPSCs yielded many more large-sized colonies on day 14, which mainly consisted of CFU-GM and CFU-GEMM colonies, compared to their normal counterparts (Fig. 2e). Colonies derived from CMML iPSC-derived HPCs mainly consisted of monoblasts, whereas those from Normal iPSC-derived HPCs mainly consisted of macrophages (Fig. 2f).

Colonies from each of the three iPSCs clones from one CMML patient and the three clones of Normal-iPSCs from two healthy donors were performed in triplicate (Fig. 2e and Supplemental Fig. 2g). Flow cytometric analysis showed an increase in CD34+ HPCs in CMML iPSC-derived hematopoietic colonies (Fig. 2g). Substantial numbers of CD13+CD14+ and CD13+CD56+ monocytic cells were generated by CMML iPSC-derived HPCs through granulocyte/macrocyte terminal differentiation, which recapitulated the representative CD13+ and CD14+ marker phenotype of CMML (Fig. 2h,i).

**Serial re-plating capacity of hematopoietic progenitors derived from CMML iPSCs.** Subsequently, colony re-plating assays using CMML iPSC-derived HPCs and control iPSC-derived HPCs were performed. CMML iPSC-derived HPCs retained their serial re-plating capacity and generated colonies even after the fourth plating, whereas control iPSC-derived HPCs lost the ability to generate colonies after the second plating (Fig. 3a). Therefore, CMML iPSC-derived HPCs possess the ability of self-renewal. Even when cultured without cytokines, CMML iPSC-derived HPCs formed colonies. In contrast, Normal iPSC-derived HPCs yielded almost no detectable colonies (Fig. 3b). Taken together, these results suggest that CMML iPSC-derived HPCs retained multiple biological properties of CMML.

**Clinical characteristic of increased STAT5 phosphorylation in CMML iPSC-derived HPCs.** It has been reported that JMML and CMML cells exhibit hyper-phosphorylation of STAT5 induced by GM-CSF using phospho-STAT5 (pSTAT5) intracellular staining followed by flow cytometric analysis15. CD34+CD43+ iPSCs-derived hematopoietic progenitors from Normal and CMML iPSCs were analyzed for STAT5 hyper-phosphorylation by flow cytometric analysis. Increased STAT5 activation in the CD34+CD43+ iPSCs-derived hematopoietic progenitors was observed in CMML iPSC-derived HPCs (Fig. 3c,d).

**Up-regulation of NF1 and EZH2 target gene signaling in CMML iPSCs and CMML iPSC-derived HPCs.** We performed a hierarchical clustering analysis using clustering software (cluster 3.0). This clustering software can analyze the clustering based on gene expression data. Using this software, the hierarchical clustering separated iPSCs from iPSC-derived HPCs. Furthermore, the categories of CMML-iPSCs were different from those of Normal-iPSCs. Similarly, the categories of CMML-iPSC-derived HPCs were different from Normal-iPSC-derived HPCs (Fig. 3e).

Following, we used gene set enrichment analysis (GSEA) to determine whether specific gene sets correlated with CMML. We performed comprehensive gene analysis to select the gene set that showed significant differences in gene expressions between CMML iPSCs-derived HPCs and normal iPSCs-derived HPCs. GSEA analysis revealed enrichment of many gene sets including NF1 and EZH2 pathways-associated ones in CMML-derived samples. The gene sets commonly enriched in CMML-iPSCs and CMML iPSC-derived HPCs we identified are listed in revised supplemental data (Supplementary LIST S1). We focused on NF1 and EZH2 pathways-associated gene sets, because these genes were known to be involved in the pathogenesis of CMML.

CMML-iPSCs were predominantly correlated with the EZH2- and NF1-related genes compared with Normal-iPSCs (Fig. 3f,g-1). Even after the re-differentiation of hematopoietic cells from iPSCs, CMML iPSC-derived HPCs correlated with the EZH2- and NF1-related genes compared with Normal iPSC-derived HPCs (Fig. 3f,g-2).

**Hematopoietic cells generated through in vivo teratoma formation from CMML iPSCs.** We used the in vivo differentiation system to generate transplantable HPCs through teratoma formation16,17. A mixture of CMML iPSCs and OP9 cells was injected subcutaneously into 7-week-old NSG mice. Teratomas in the NSG mice were generated 3 months after transplantation (Fig. 4a). The teratomas from CMML iPSCs were larger in volume than those from Normal iPSCs (Fig. 4b). These teratomas contained all three germ layers.
Figure 2. The patient’s pathogenesis of CMML was recapitulated in vitro in CMML-iPSC-derived hematopoietic progenitor cells (HPCs). (a) Scheme for inducing CMML and normal iPSC-derived HPCs. We obtained CD34+ CD43+ hematopoietic progenitor cells in CMML-iPSC sac on day 17 of the co-culture system. We evaluated two different lines of one normal and unique one CMML iPSC clone in triplicate tests. (b) CMML iPSCs generated more CD34+ CD43+ HPCs than normal-iPSCs (n = 3, independent experiments, ***p < 0.001, Normal-iPSCs1 and 2 from a healthy donor, and CMML-iPSCs1 and 2 from a CMML patient, paired two-sided t-test). (c,d) The CD34+ CD38− CD90− fraction re-differentiated from CMML-iPSCs increased more than normal-iPSCs (n = 3, independent experiments, ***p < 0.001, Normal-iPSCs1 and 2 from a healthy donor, and CMML-iPSCs1 and 2 from a CMML patient, Experiments were performed in triplicate. Statistical analyses were performed with ANOVA and the Dunnett post-test for multiple comparisons. ***p < 0.001). (e) The semi-solid culture of CD34+ CD43+ HPCs derived from CMML-iPSCs yielded many more CFU-GM and CFU-GEMM colonies, compared to their normal counterparts (n = 3, independent experiments, ***p < 0.001, Normal-iPSCs1 and 2 from a healthy donor, and CMML-iPSCs1 and 2 from a CMML patient, Statistical analyses were performed...
Immunohistostaining of human CD45 revealed the presence of human CD45+ blood cells, particularly around the small blood vessels in the teratomas (Fig. 4c). Flow cytometric analysis using dissociated teratoma tissues also revealed the presence of human CD45+ CD34+ HPC fractions. Moreover, in the BM of teratoma-bearing mice, human CD45 + CMML cells were detected, which were few CD34+ and CD14+ monocytes (Fig. 4d).

We estimated the number of CMML cells in the BM of the engrafted mice in which teratomas formed from the CMML iPSCs. Human CD45 positive CMML cells were reproduced in the BM and were more abundant than in the normal controls (Fig. 4e).

**Second transplantation of CMML progenitor cells generated in vivo.** To determine whether these HPCs from CMML iPSC-derived teratomas were transplantable, they were transplanted a second time into the BM of immunodeficient mice (Fig. 4a). Hematopoietic progenitors (HPCs) stained with both human CD45 and human CD34 were sorted from the teratoma and BM using a FACS Aria Cell Sorter. Subsequently, 1 × 10^6 human CD45+ CD34+ HPCs harvested from the teratoma were injected into the bone marrows of irradiated NSG mice (2 Gy). After 6 months, bone marrow cells from NSG mice were analyzed (Fig. 4a).

CMML-iPSC-derived blood in vivo reproduced 23.4 ± 0.437% of the bone marrow and recapitulated the profiles of CMML surface antigens, CD13+ or CD34+. Normal iPSC-derived blood in vivo reproduced 0.83 ± 0.437% of the bone marrow in small amounts (Fig. 4f,g). Immunohistochemical staining revealed that human CD45+ cells were engrafted in the BM of recipient mice that received CMML-iPSCs (Fig. 4h). The sorted human CD45+ cells consisted of monocytes (Fig. 4i-a) and a few monoblastic cells (Fig. 4i-1,2). Collectively, the CMML marker expression profile was reproduced in this in vivo model. EZH2, RUNX1, and NRAS mutations were also identified in the BM of secondarily transplanted NSG mice (Fig. 4i).

**Drug discovery using CMML iPSCs.** We tested the effect of several candidate drugs in order to identify therapeutic agents for CMML. The drug testing method involved three phenotype-screening systems with a colony assay, pERK, and re-plating assay using CMML iPSCs. To evaluate our CMML iPSCs as a tool for drug discovery, we performed colony assays on CD34+ HPCs harvested from the teratoma and BM using a FACS Aria Cell Sorter. Subsequently, 1 × 10^6 human CD45+ CD34+ HPCs harvested from the teratoma were injected into the bone marrows of irradiated NSG mice (2 Gy). After 6 months, bone marrow cells from NSG mice were analyzed (Fig. 4a).

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**Discussion**

Due to the lack of cell lines in CMML and the limited developments of mouse models thus far, it has remained difficult to analyze the pathology of CMML and to develop therapeutic agents for the disease. In this study, we developed and utilized CMML-iPSCs to model the disease for purposes of pathophysiological analysis and drug testing to search for potential treatments.

First, we reproduced the hematopoietic system of CMML using CMML-iPSCs. The CMML-iPSCs retained several important disease characteristics of CMML. The blood cells, which were re-differentiated from CMML-iPSCs, recapitulated the pathogenesis of the CMML patient in their myeloproliferation, marker phenotypes, and extensive self-renewal, even after the reprogramming processes. CMML-iPSCs-derived HPCs from this patient reproduced proliferative monocytosis and demonstrated a serial re-plating ability mimicking a leukemic phase. It has been reported that juvenile myelomonocytic leukemia (JMML) iPSCs exhibit similar characteristics. JMML-derived HPCs demonstrated the proliferative capacities of monocytic cells and activation of the JAK/STAT or MEK/ERK pathway. As JMML and CMML may have a common pathophysiology, it is possible that our understanding of these diseases will be advanced by using both CMML-iPSCs and JMML-iPSCs.
Figure 3. Characteristics of colony formation and genetic analysis of CMML iPSC-derived HPCs. (a) CMML iPSC-derived HPCs retained their serial re-plating capacity and generated colonies even after the fourth plating, whereas Normal iPSC-derived HPCs lost the ability to generate colonies after the second plating. CMML iPSC-derived HPCs showed an enhanced self-renewal capacity (n = 3, independent experiments, *p < 0.05, Normal-iPSCs1 and 2 from a healthy donor, and CMML-iPSCs1 and 2 from a CMML patient). Statistical analyses were performed with ANOVA and the Dunnett post-test for multiple comparisons. *p < 0.05). (b) The CMML iPSC-derived HPCs generated hematopoietic colonies under cytokine-deprived conditions. The CMML iPSC-derived HPCs could form hematopoietic colonies, while Normal iPSC-derived cells gave rise to few detectable colonies (n = 3, independent experiments, ***p < 0.001, Normal-iPSCs1 and 2 from a healthy donor, and CMML-iPSCs1 and 2 from a CMML patient, paired two-sided t-test). (c) Using pStat5 intracellular flow cytometric analysis, we observed increased phosphorylation of STAT5 in CMML iPSC-derived CD34+CD43+ HPCs. (d) The percentage of pSTAT5 on CMML iPSC-derived CD34+CD43+ HPCs was significantly higher than Normal iPSC-derived HPCs (n = 3, independent experiments, ***p < 0.001, Normal-iPSCs1 from a healthy donor, and CMML-iPSCs1 from a CMML patient, paired two-sided t-test). (e) Hierarchical clustering analysis of gene expression patterns. Using hierarchical clustering, these cells were distinguished into two groups: the Normal/CMML iPSC group and Normal/CMML iPSC-derived HPC group. (f-1 and f-2) Expression data were...
analyzed to identify genes in CMML-iPSC-derived HPCs and CMML-iPSCs that positively affected NF1 using GSEA. \( p < 0.001 \). (g-1 and g-2) Expression data were analyzed to identify genes in CMML-iPSC-derived HPCs and CMML-iPSCs that positively affect EZH2 using GSEA. \( p < 0.001 \). Horizontal lines represent the means ± s.d.

Second, we established a humanized CMML mouse model via teratomas using CMML iPSCs in vivo. This mouse model makes it possible to analyze the pathogenesis of CMML, explore treatment options for individual patients, and discover other useful insights about the disease’s characteristics. With the development of an in vivo transplantation system using teratomas, the formation of vascular niches makes it possible to reconstruct CMML iPSCs-derived from HPCs\(^8\). Immunohistostaining of human CD45 revealed the presence of human CD45+ blood cells, particularly around the small blood vessels in the teratomas. It was recently reported that primary samples of CMML and JMML can be transplanted into NSGS mice that trans-genetically express human GM-CSF, interleukin-3, and stem cell factor to develop remarkably humanized mouse models\(^21-23\). Our particular technique of developing humanized CMML mouse models using CMML-iPSCs offers an alternative approach that boasts an added advantage: because CMML-iPSCs can be used repeatedly, the mouse models can be produced in infinitely large quantities with limited initial resources.

Furthermore, several drug tests were conducted using CMML-iPSCs, which were established from a CMML patient with genetic mutations in NRAS, RUNX1, and EZH2. Ras and MEK inhibitors effectively suppressed the proliferative capacity of CMML iPSC-derived HPCs. Notably, these drug tests also revealed that liposomal clodronate is a possible agent for treating CMML. Clodronate is a drug used clinically for treatment of osteoporosis, but it has other clinical properties. In particular, clodronate covered with liposomes (liposomal clodronate) can be used to deplete monocytes in monocyte/macrophage systems\(^3\). In addition, several reports have indicated that injection of liposomal clodronate reduced or depleted monocytes in mice and rats\(^24-26\). The reduction of monocytes by liposomal clodronate may indicate that liposomal clodronate may be effective in treating CMML and monocytic leukemia. However, this study is the first to attempt administration of liposomal clodronate to the human CMML patient samples. Clodronate has been widely used for the treatment of osteoporosis, and its liposomalization will enable clodronate to be applied to the treatment of CMML as a drug-repositioning treatment.

We acknowledge the limitation of this study in that we obtained and used three clones of CMML iPSCs from only one patient, as it was difficult to program them from CMML patient samples. Our experiment is not the first to suggest that not all samples can be reprogrammed and that samples with chromosome 7 abnormalities are more easily reprogrammed. In the cases of past experiments, samples from MDS patients successfully reprogrammed into iPSC cells were MDS samples with t(1;7) derivatives or del(7q), or samples from AML patients with del(7q) complex karyotypes\(^4\). Past studies have linked successful iPSC reprogramming to the methylation of samples\(^27\). Additionally, it has been shown that samples with chromosome 7 abnormalities are more easily methylated\(^28\). Therefore, it may be possible that the methylation of the two t(1;7) derivative samples in our experiment was what enabled their successful reprogramming into iPSC cells, whereas our eight other samples, which did not have chromosome 7 abnormalities, could not be reprogrammed.

Therefore, the analysis may not fully reflect all cases of CMML. In order to compensate for this limitation, we utilized patient-derived samples for some parts of this investigation.

Conclusion
In summary, we established patient-derived CMML-iPSCs and developed various associated systems for disease-modeling and drug-testing purposes: hematopoietic re-differentiation in vitro, a humanized CMML mouse models via teratomas, and a drug-screening system using CMML-iPSCs. The clinical characteristics of CMML were recapitulated following hematopoietic re-differentiation in vitro and in humanized CMML mouse models in vivo. In addition, the well-established drug-testing system indicated the possibility of repositioning liposomal clodronate as a clinically applicable candidate drug for CMML therapy. CMML-iPSCs provide a powerful platform for dissecting the pathology of CMML, and clodronate is a potential repositioned drug for the treatment of CMML.

Materials and Methods
Protocol for the generation of patient-derived CMML iPSCs. CD34+ cells from patient samples and a healthy control donor were isolated from BM mononuclear cells using FACS Aria. CD34+ cells from patient samples were cultured in DMEM, with SCF, TPO, Flat3, IL3, and IL6. DMEM contained 10% FBS, 10 ng/mL IL-3, 10 ng/mL IL-6, and 10 ng/mL SCF. The cultured cells were exposed to episomal vectors expressing five factors (Oct3/4, Sox2, KLF4, L-myc, LIN28, and shTP53). The transfected cells were cultured in a MEF feeder layer. The medium was changed from DMEM to iP5 medium on day 0. The iP5 medium containing 1 mM butyrate and 10 mM Y-27632 (ROCK inhibitor) was changed every other day. Throughout the process, we used 5% O2, hypoxic conditions\(^12\). After 30 days, ES-like colonies developed in the dish. The colonies were moved to another dish.

Episomal vector method. This study (approval number 2771) was approved by the ethics committee of the University of Tokyo, and written informed consent was obtained. In addition, all methods in the study were performed in accordance with the relevant stem cell guidelines and regulations established by the University of Tokyo ethics committee. Okita K. and his colleagues have developed efficient methods to generate integration-free human iPSCs\(^11\). Three episomal plasmid vectors were used: pCXLE-hOCT3/4-shp53-F, pCXLE-hSK (SOX2/KLF4), and pCXLE-hUL (L-MYC/LIN28). The established efficiency of iPSC cells can be dramatically enhanced by using an episomal vector encoding EBNA-1 together with three episomal vectors. The episomal vectors...
Figure 4. In vivo CMML xenograft model recapitulate its pathogenesis. (a) Experimental scheme for the first transplantation. We co-injected CMML-iPSCs along with OP9 stromal cells into immunodeficient mice using an in vivo system in which human CMML iPSCs develop into HPCs within teratomas. Second transplantation of CMML progenitor cells generated in vivo. (b) CMML iPS-induced teratomas had larger volumes than the induced teratomas from Normal-iPSCs (n = 3, independent experiments, Normal-iPSCs1 from a healthy donor, and CMML-iPSCs 1 from a CMML patient, paired two-sided t-test). (c) Immunostaining of CD45+ blood cells around the small blood vessels generated in the teratoma. (d) CD45+CD34+ HPCs were isolated from the teratoma. Leakage of the CD45+ CMML cells from the teratoma into the BM was observed. (e) Estimated number of CMML cells in the BM of engrafted mice in which teratomas formed from tCMML iPS cells (n = 4, independent experiments, *p < 0.05, Normal-iPSCs1 from a healthy donor, and CMML-iPSC1 from a CMML patient, paired two-sided t-test). (f) CMML-iPSC-derived blood in vivo reproduced and recapitulated the profiles of CMML surface antigens, CD13+ or CD34+. (g) Flow cytometry revealed human chimerism of CD45+ cells in the BM of the secondarily transplanted NSG mice. Statistical analysis was performed (n = 6,
disappeared after iPSCs passed several times. Therefore, analysis of integration-free human iPSCs can be performed without any affects from the transduction of foreign genes in this episomal vector method\(^1\).

**Mutation analysis of CMML iPSCs.** We performed direct genomic sequencing using standard techniques and an ABI 3130 Genetic Analyzer for the patient samples and CMML iPSC samples to screen for the following mutations: **CB1** (exons 8–9); **KRAS** and **NRAS** (exons 1–2); **IDH1** and **IDH2** (exons 4); **DNMT3A** (exons 18–23); **TET2** (exons 3–12); **ASXL1** (exon 12); **RUNX1** (exons 3–8); **JAK2** (V617F); and **SRSF2**, **SF3B1**, **U2AF35**, and **EZH2** (all coding exons)\(^1\)\(^9\).

**Bisulfite sequencing assay.** One microgram of genomic DNA was converted using the EpiTect Bisulfite Kit (Qiagen, Hilden, Germany) according to the manufacturer's recommendations. The NANOG promoter region in the converted DNA was amplified by PCR using previously reported primers. The PCR products were cloned into bacteria using the TA PCR Cloning Kit (DynaExpress, Tokyo, Japan). Ten clones from each sample were analyzed by sequencing with the M13 universal primers\(^1\).

**“iPS-sac” method to induce iPSCs differentiation into hematopoietic cells.** Small clumps (<100 cells) of iPSCs were transferred to a dish containing irradiated C3H10T1/2 cells. iPSCs with the C3H10T1/2 cells were cultured in a differentiation medium containing VEGF, which was refreshed every 3 days for 2 weeks. After 2 weeks, the round, hematopoietic-like cells were harvested and then sorted by flow cytometry\(^1\)\(^2\).

**Colony forming assay.** To investigate differentiation capacity, 1 × 10\(^5\) control cells and CMML iPSC-derived CD34\(^+\)CD43\(^+\) HPC (hematopoietic progenitor cells) were seeded on a semi-solid classical medium (H4434) (Stem cell Technologies) containing recombinant (rh) human stem cell factors, rh GM-CSF, rh IL3, and rh erythropoietin.

**Flow cytometric and morphological analyses.** The following antibodies were used for FACS analysis and immunocytochemistry: anti-human CD34 APC-conjugated (Beckman Coulter, Brea, CA, USA), anti-human CD43 PE-conjugated (Beckman Coulter), CD13 APC-conjugated (Beckman Coulter, Brea, CA, USA), CD14 PE-conjugated (Beckman Coulter, Brea, CA, USA), CD24 APC-conjugated (Beckman Coulter, Brea, CA, USA), anti-human CD5 FITC-conjugated (Beckman Coulter), CD7 PE-conjugated (Beckman Coulter), and CD56 PE-conjugated (Beckman Coulter). The cells were analyzed and sorted using a FACS Aria II (Becton Dickinson).

**Microarray analysis.** Total RNA was extracted from CD34\(^+\) cells from a primary CMML patient's BM and a healthy donor's BM and their derived iPSCs and HPCs. Samples were analyzed using the Whole Human Genome DNA microarray (Agilent Technologies, Santa Clara, CA, USA). Raw array signals were extracted from the CEL files, and background correction and normalization were performed using RMA. A heat map for selected probes associated with hematopoiesis was produced.

Using the t-test, standardized signal values were compared between all pairs of the four groups: normal iPSCs, CMML iPSCs, normal iPSC-derived HPCs, and CMML iPSC-derived HPCs. Any probe sets for which the p-values were not <0.05 for any comparisons were excluded from further analysis. Any probe sets for which the maximum/minimum ratio was <4 were also excluded. Using these criteria, 3372 probes were identified. Subsequently, unsupervised hierarchical clustering was performed with Cluster 3.0 (Stanford University), and the results were visualized with Java TreeView. GSEA (www.broadinstitute.org/gsea) was then performed by comparing normal and CMML-derived samples both for iPSCs and HPCs against MSigDB c2.all.v4.0. Significant gene sets enriched in CMML samples were identified using an FDR q-value method (Benjamini-Hochberg). Significant gene sets enriched in CMML samples were identified using an FDR q-value method (Benjamini-Hochberg). The study to analyze hematological mouse models using immunodeficient mice (approval number P09-104) was approved by the animal care committee.

**Intracellular phospho-specific flow cytometry in phosphorylation of Stat5 and MEK/ERK pathways.** CD34 + CD43 + iPSCs-derived hematopoietic progenitors from Normal and CMML iPSCs were analyzed in hyper-phosphorylation of Stat5 by flow cytometric analysis using the Fixation/Methanol Protocol. The cells were fixed with Cytofix/Cytoperm solution (Becton Dickinson; BD) for 20 min at 4 °C and permeabilized with Perm buffer III (BD Perm/Wash™ buffer) for 30 min on ice. The cells were immunostained for pSTAT5 at room temperature for 1 h.

To evaluate our CMML iPSCs as the tools for cell-based assays, we performed colony assays on CD34 + CD43 + HPCs differentiated from iPSCs by culturing them with and without the MEK inhibitor (PD0325901) or the Ras inhibitor (salirasib). After treatment with the MEK or Ras inhibitor, the cells from these colonies were harvested on day 14 and analyzed by pERK intracellular phospho-specific flow cytometry. These cells were intracellularly stained with Phospho-ERK1/2 monoclonal Antibody (APC, ebioscience) using the Fixation/Methanol Protocol. In this protocol, fixation was followed by treatment of cells with methanol. These treated cells were analyzed by pERK intracellular phospho-specific flow cytometry.
Figure 5. Drug discovery using CMML-iPSCs. (a) An MEK inhibitor (PD0325901) reduced the colony-forming capacity of CMML iPSC-derived HPCs (n = 3, independent experiments, Normal-iPSCs1 from a healthy donor, and CMML-iPSCs1 from a CMML patients, Experiments were performed in triplicate. Statistical analyses were performed with ANOVA and the Dunnett post-test for multiple comparisons. ***p < 0.001)). (b) RAS inhibitor (salirasib) reduced the colony-forming capacity of CMML iPSC-derived HPCs (n = 3, independent experiments, Normal-iPSCs1 from a healthy donor, and CMML-iPSCs1 from a CMML patients, Experiments were performed in triplicate. Statistical analyses were performed with ANOVA and the Dunnett post-test for multiple comparisons. **p < 0.01). (c) pERK was constitutively activated only in the colonies of CMML iPSC-derived HPCs. Furthermore, ERK phosphorylation was inhibited by MEK and RAS inhibitors in these colonies. (d) The expression level of pERK was significantly decreased in CMML-iPSC-derived hematopoietic colonies by MEK or Ras inhibitor (n = 3, independent experiments, *p < 0.05, Normal-iPSCs1 from a healthy donor, and CMML-iPSCs1 from a CMML patients, paired two-sided t-test). (e) Liposomal clodronate suppressed the growth of colonies from CMML iPSCs. Statistical analysis was performed (n = 3,
independent experiments, Normal-iPSCs1 from a healthy donor, and CMML-iPSCs1 from a CMML patients. Experiments were performed in triplicate. Statistical analyses were performed with ANOVA and the Dunnott post-test for multiple comparisons. \( * * p < 0.001, \) \( * p < 0.01. \) (f) Liposomal clodronate reduced re-plating colonies formed by CMML-iPSC-derived HPCs \( (n = 3, \) independent experiments, Normal-iPSCs1 from a healthy donor, and CMML-iPSCs1 from a CMML patients. Experiments were performed in triplicate. Statistical analyses were performed with ANOVA and the Dunnott post-test for multiple comparisons. \( * * p < 0.001. \) (g) Colony formation by three other primary samples from CMML patients was reduced by liposomal clodronate. Horizontal lines represent the means ± s.d.

experiment committee of the University of Tokyo. In addition, all methods in the study were performed in accordance with the relevant animal studies guidelines and regulations established by the University of Tokyo ethics committee. A total of \( 5 \times 10^6 \) CMML-iPSCs and \( 1 \times 10^6 \) OP9 cells were injected subcutaneously into the abdomen of 7-week-old NSG mice. Three months after injection, the developed teratomas and BM were harvested. We used four NSG mice for Normal-iPSC- derived teratomas and four mice for CMML-iPSCs via teratomas\(^{14,15}\).}
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**Author Contributions**
K.T. mainly performed the experiments. K.K., M.H., M.M., S.Y., A.H., W.A. and K.K. performed parts of experiments. S.A. and A.Y. analyzed the array data. N.T., M.O., K.E., A.N., T.N. and H.N. contributed to the novel in vivo experiments. K.T. mainly wrote the manuscript, T.K., S.A., A.Y., F.N. and M.K. revised the manuscript.

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