

Identification of pre-leukemic hematopoietic stem cells in acute leukemia

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Summary

In acute myeloid leukemia (AML), the cell of origin, nature and biological consequences of initiating lesions and order of subsequent mutations remain poorly understood, as AML is typically diagnosed without observation of a pre-leukemic phase. Here, highly purified hematopoietic stem cells (HSC), progenitor and mature cell fractions from the blood of AML patients were found to contain recurrent *DNMT3a* mutations (*DNMT3a^{mut}*) at high allele frequency, but without coincident *NPM1* mutations (*NPM1c*) present in AML blasts.

DNMT3a^{mut}-bearing HSC exhibited multilineage repopulation advantage over non-mutated HSC in xenografts, establishing their identity as pre-leukemic-HSC (preL-HSC). preL-HSC were found in remission samples indicating that they survive chemotherapy. Thus *DNMT3a^{mut}* arises early in AML evolution, likely in HSC, leading to a clonally expanded pool of preL-HSC from which

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Author Contributions

Authors L.I.S., S.Z. contributed equally to this work; L.I.S., S.Z. designed and performed experiments, analyzed data and wrote the manuscript; A.M., W.C.C screened AML engraftment in xenotransplantation assays; J.M.B., V.G., J.A.K., A.D.S., A.C.S., K.W.Y., M.D.M. collected AML samples and assembled clinical information; J.A.K. correlated xenotransplantation engraftment data with clinical information; J.L.M., M.D. performed xenotransplantation experiments; J.J.F.M., R.M. performed ddPCR; H.J.K., K.L. performed Sanger sequencing; J.D.M., T.J.H., supervised the targeted sequencing; A.M.K.B performed targeted sequencing; Q.M.T., L.D.S. performed *DNMT3a* data mining. M.D.M. designed the study; J.C.Y.W. supervised AML xenotransplantation screening experiments, designed the study and wrote the manuscript; J.E.D. supervised the study and wrote the manuscript.

AML evolves. Our findings provide a paradigm for the detection and treatment of pre-leukemic clones before the acquisition of additional genetic lesions engenders greater therapeutic resistance.

Introduction

There is overwhelming evidence that virtually all cancers are clonal and represent the progeny of a single cell¹⁻³. However, the evolutionary trajectory that leads from the first somatic mutation to the eventual development of cancer is not well mapped. The simplest models predict that each newly acquired somatic mutation confers selective advantage to drive successive waves of clonal expansion, with the fittest clone becoming dominant. However the modern era of cancer genomics has exposed a more complex clonal architecture in many tumor types⁴, where multiple genetically distinct subclones co-exist with the dominant clone^{5,6}. Comparison of diagnostic and recurrent/metastatic samples obtained from the same patient has established that the latter frequently do not evolve from the dominant clone, but instead can be traced either to a minor subclone present at diagnosis, or to a putative, undetected ancestral clone⁷⁻¹⁵. Thus, a clear understanding of the genomic landscape of tumors is required in order to devise targeting strategies that eliminate not only the dominant clone but also the subclonal reservoirs from which recurrence can arise.

Although the clonal composition of cancer lineages within individual tumors is coming into focus, the very first steps in cancer development remain poorly defined. Early and possibly initiating mutations have been identified from analysis of pre-neoplastic lesions in breast¹⁶, lung¹⁷, skin¹⁸, and colon cancer¹⁹, as well as from studies of AML cases that evolved from a prior myelodysplastic syndrome (MDS)²⁰. However, key questions remain unanswered. In particular, can clinically relevant clones be traced back to a non-tumorigenic cell? Do pre-cancerous ancestral clones persist after tumor development? If so, are they present in the diagnostic sample, and do they survive treatment and persist in remission samples?

Human leukemia is a disease model particularly suited to addressing these fundamental questions, due to the depth of our understanding of normal hematopoiesis and the availability of functional assays and analytic tools that allow examination of phenotypically defined populations at the single cell level²¹. In AML, a subset of cases evolve from a preceding clinically overt phase such as MDS or chronic myeloid leukemia (CML), characterized by clonal expansion of one or more blood lineages^{22,23}. The founder mutations present in pre-leukemic cells are retained in the AML blasts, implicating them as putative initiating events and establishing clonal expansion as the first step in leukemogenesis. Interestingly, somatic mutations in some leukemia-associated genes such as *TET2* have also been linked to multilineage clonal hematopoiesis in aging healthy individuals²⁴. Insight into the phenotype of the normal cell from which clonal expansion can initiate was first provided by the pioneering studies of Fialkow in CML, which demonstrated that *BCR-ABL1* arises in a multipotential HSC²⁵. However, for the majority of AML cases that arise de novo without any prior clinical perturbations, insight into the cellular context and functional consequences of the earliest genetic lesions requires identification and examination of ancestral cells within the diagnostic sample. Recent studies have found that only a subset of mutations contained in AML blasts were present in HSC-enriched cell fractions isolated from AML

patient samples, and that these cells were capable of non-leukemic differentiation^{26,27}. Here we establish that these ancestral preL-HSC present at diagnosis are able to regenerate the entire hematopoietic hierarchy while possessing competitive repopulation advantage over non-leukemic HSC leading to clonal expansion. These preL-HSC are found in a high proportion of AML patients that carry mutations in *DNMT3a* and *IDH2*, and unlike AML blasts, they survive induction chemotherapy and persist in the bone marrow at remission, providing a potential reservoir for leukemic progression.

Results

During studies to examine intra-tumoral genetic heterogeneity in AML, deep targeted sequencing (Tar-seq, read depth ~250×) of 103 commonly mutated leukemia genes (Fig. 1a, Supplementary Table 1) was carried out on peripheral blood (PB) samples from 12 patients at diagnosis (blasts >80%) (Supplementary Table 2, 3). Normal T-cells were expanded *in vitro* to provide non-leukemic tissue for genetic comparison. Consistent with mutant allele frequencies reported in recent studies²⁸ *DNMT3a^{mut}* was found in 4 of 12 samples (mutant allele frequency ~50%) (Fig. 1a). Unexpectedly, in 3 of these 4 patients, *DNMT3a^{mut}* was detected in T-cells at a low allele frequency (1–20%). Other genetic alterations including *NPM1c* were found only in PB but not T-cell samples, ruling out AML cell contamination of cultured T-cells. To estimate the proportion of AML cases with *DNMT3a^{mut}*-bearing T-cells, 71 additional samples, taken at diagnosis from patients with normal cytogenetics, were screened by Sanger sequencing for *DNMT3a^{mut}* along with other common AML mutations (Supplementary Table 4). Consistent with published data^{29–31}, 17 of 71 AML samples (24%) carried mutations in *DNMT3a*, and 15 of these 17 (88%) also carried *NPM1c*. For these 17 patients, the allele frequency of *DNMT3a^{mut}* and *NPM1c* in CD33+ blasts, as well as corresponding freshly isolated T-cell controls, was measured by droplet digital PCR (ddPCR) at a sensitivity of 1 mutated allele in 1000 reference alleles. Whereas both *DNMT3a^{mut}* and *NPM1c* were always present in blasts at similar allele frequency, *DNMT3a^{mut}* with no evidence of *NPM1c* was detected in T-cells from 12 of these 17 patients (70.5%) (Fig. 1b). In addition, *FLT3*-ITD was detected in blasts but not the T-cells of 2 patients bearing this mutation (Extended Data Fig. 1). These data reveal the sequential order of mutation acquisition in these patients, with *DNMT3a^{mut}* arising earlier in leukemogenesis than *NPM1c* and *FLT3*-ITD, a conclusion predicted from recent studies on bulk AML blasts showing that *NPM1c* and *FLT3*-ITD occur late and are the only genes recurrently mutated in *DNMT3a^{mut}* AML^{15,28,32}. Moreover, our findings establish that *DNMT3a^{mut}* occurs in an ancestral cell that gives rise to both T-cells and the dominant AML clone present at diagnosis.

To gain insight into the properties of the ancestral cell within which *DNMT3a^{mut}* first arises, we examined additional non-leukemic hematopoietic cell populations from 11 *DNMT3a^{mut}*/*NPM1c* AML patients. A high resolution 12-parameter sorting strategy^{33–35} was employed to isolate non-leukemic hematopoietic stem and progenitor populations, including hematopoietic stem cells/multipotent progenitors (HSC/MPP), multilymphoid progenitors (MLP), common myeloid progenitors (CMP), granulocyte monocyte progenitors (GMP), and megakaryocyte erythroid progenitors (MEP), as well as mature B, T and natural killer (NK) cells within the CD33– cell fraction. Together with CD45dimCD33+ AML blasts,

these highly purified, phenotypically defined normal cell populations were assessed by ddPCR for *DNMT3a^{mut}* and *NPM1c* (Fig. 2, Extended Data Fig. 2, 3). *DNMT3a^{mut}* was found together with *NPM1c* in CD33+ blasts from all patients. By contrast, we found *DNMT3a^{mut}* at variable allele frequency without *NPM1c* across the spectrum of mature and progenitor cell populations. Results for a representative patient (#11) are shown in Fig. 2a–b. In this patient, *DNMT3a^{mut}* was present in HSC/MPP at an allele frequency of 12–30% without detectable *NPM1c*. Although the clonal contribution that an individual normal HSC makes in humans is unknown, studies in higher primates estimated that single HSC provide approximately 0.5% clonal contribution during steady-state hematopoiesis³⁶. Thus, the high *DNMT3a^{mut}* allele frequency in HSC/MPP points to their clonal expansion as compared to non-mutated HSC. In Patient #11, *DNMT3a^{mut}* was also present in all downstream progenitors at variable frequencies. The mean allele frequency among HSC/MPP, MLP and CMP was 24.6% across all patients analyzed (Extended Data Fig. 2). Importantly, even for patients in whom *DNMT3a^{mut}* was not detected in mature cells, *DNMT3a^{mut}* without *NPM1c* was found in stem/progenitor populations (Fig. 2c), providing further strong evidence that *DNMT3a^{mut}* precedes *NPM1c* during leukemogenesis. Interestingly, in 3 patients (#10, 14, 16), *DNMT3a^{mut}* was detected in CMP but not HSC/MPP, an outcome consistent with the existence of *DNMT3a^{mut}*-bearing HSC below our detection limit that generated a clonally expanded CMP population, or possibly the existence of a preceding lesion in HSC/MPP with later acquisition of *DNMT3a^{mut}* in CMP. Resolution of this question will require whole genome sequencing of sorted blast and phenotypically normal populations from patient samples. Our analysis also showed that in 6 of 11 patients, both *DNMT3a^{mut}* and *NPM1c* were found together in MLP and/or GMP populations, pointing to the likely progenitor cell types where overt AML driven by *NPM1c* arises. Collectively, our findings provide key insights into the leukemogenic process in human AML and confirm historical predictions from early clonality studies of the existence of a pre-leukemic state^{3,37}.

To examine how *DNMT3a^{mut}* affects population dynamics during leukemic progression, we undertook temporal analysis of mature and progenitor cells from 5 patients (#11, 28, 35, 55, 57) sampled at diagnosis, remission (3 months) or relapse (Fig. 2b, d, Extended Data Fig. 3, 4). Compared to diagnosis, the allele frequency of *DNMT3a^{mut}* alone was similar or higher at remission (Patients #11, 28, 35, 57) and relapse (Patient #11, 55). Although CD33+ leukemic blasts at diagnosis always carried both mutations, CD33+ myeloid cells at remission bore only *DNMT3a^{mut}*, suggesting that they are not AML blasts but the progeny of *DNMT3a^{mut}*-bearing progenitors with preserved myeloid differentiation capacity. In the relapse sample of Patient #11, both mutations were present in the majority of cells, with the exception of HSC/MPP in which a proportion carried only *DNMT3a^{mut}*. Patient #57 was a long-surviving patient that allowed a comparison of early and late (36 months) remission samples and showed a striking increase in *DNMT3a^{mut}* allele frequency in most cell populations over time (Fig. 2d). In addition, a small proportion of CD33+ myeloid cells in the late remission PB sample contained both *DNMT3a^{mut}* and the *NPM1c* mutation found at diagnosis, suggesting either regrowth of the diagnostic leukemic clone or emergence of a new clone following an independent *NPM1c* mutation event within the preleukemic pool. Collectively, our data indicate that the ancestral cell that bears *DNMT3a^{mut}* without *NPM1c* is an HSC/MPP capable of multilineage differentiation. Moreover, these ancestral HSC/MPP

survive chemotherapy, expand during remission, and might serve as a reservoir for clonal evolution leading to recurrent disease.

To establish conclusively whether phenotypically defined *DNMT3a^{mut}*-bearing HSC/MPPs are functional HSC and whether competitive repopulation advantage underlies their *in vivo* clonal expansion, we undertook xenograft repopulation assays. Mononuclear cells from the PB of 2 patients at diagnosis (#11, 55) with *DNMT3a^{mut}* allele frequency in HSC/MPP of 30% and 20% respectively were transplanted into cohorts of immune-deficient mice using a limiting dilution approach and analyzed after 8 and 16 weeks (Fig. 3, Extended Data Fig. 5). In this xenograft model, leukemic engraftment is characteristically seen as a dominant myeloid (CD45dimCD33+) graft, whereas non-leukemic grafts are multilineage and contain both lymphoid (predominantly CD19+ B cells) and myeloid (CD33+) cells (Extended Data Fig. 6). For Patient #11, multilineage engraftment was seen in 24 of 35 mice, giving a calculated frequency of 1 repopulating HSC in 7.3×10^5 cells (Extended Data Fig. 5a). Only a single graft contained more than 50% CD33+ myeloid cells, suggesting co-engraftment by a leukemia stem cell (LSC) that was present at low frequency³⁸. We analyzed by ddPCR 12 of the multilineage xenografts following 16 weeks of repopulation. Ten of these contained a high proportion of cells bearing *DNMT3a^{mut}* without *NPM1c* (mean allele frequency 57%), whereas both *DNMT3a^{mut}* and *NPM1c* were present in the single mouse with significant myeloid engraftment (Fig. 3b). Kinetic analysis demonstrated increasing *DNMT3a^{mut}* allele frequency in multilineage grafts over time (Fig 3c). Similar results were found for Patient #55 (Extended Data Fig. 5 and data not shown). In contrast, cells from the relapse sample of both patients generated leukemic grafts and no multilineage grafts (Fig. 3a and data not shown), consistent with a higher LSC frequency at relapse compared to diagnosis³⁸. Together, these data provide evidence that *DNMT3a^{mut}* occurs in HSC/MPP capable of generating a long-term multilineage lympho-myeloid graft, confirming their designation as pre-leukemic HSC²⁶ (preL-HSC). *DNMT3a^{mut}* also endows preL-HSC with a competitive repopulation advantage over non-mutated HSC explaining the clonal expansion of preL-HSC in patients at the time of diagnosis and during remission.

Our xenograft results indicate that when preL-HSC exists at higher frequency than LSC, non-leukemic multilineage grafts, rather than leukemic grafts, are frequently generated. Examination of our historical xenograft data from 264 diagnostic AML samples revealed that 37% did not generate any graft, 40% generated leukemia, and 23% gave rise to non-leukemic multilineage grafts (Extended Data Fig. 6). Sanger sequencing data was available for 25 samples that generated multilineage grafts (Supplementary Table 4) revealing that 10 of 25 (40%) came from patients bearing *DNMT3a^{mut}*; *IDH1/2* mutations were present in 12 patients, including 3 who had both *DNMT3a* and *IDH1/2* mutations (Fig. 4a, 4b). It was not possible to determine conclusively whether these xenografts were generated by preL-HSC, as they were not available for mutation testing. To examine whether pre-leukemic cells also exist in patients with *IDH1/2* mutations, we analyzed samples from 3 patients with *IDH1* and 3 patients with *IDH2* mutation by high resolution cell sorting and ddPCR. In 4 patients, no pattern of preceding mutation was detected in non-leukemic cell populations. However, in 2 patients we found *IDH2* mutation without *NPM1c* in a number of progenitor and mature populations (Fig. 4c), suggesting that *IDH2* mutation might also occur as a pre-leukemic event.

Our data predict that *DNMT3a* mutation may occur in healthy adults and pre-date AML diagnosis by months or even years. Through searches of exome sequence databases derived from PB (<https://esp.gs.washington.edu/drupal/>) we found that the frequency of the *DNMT3a* R882H variant (rs147001633) was 0.066% (3 in 4545). Although this was considered to be a germline variant in this healthy adult cohort, our findings raise the possibility that the mutations detected in these studies may have originated from an HSC/MPP containing an acquired somatic *DNMT3a* mutation that underwent clonal expansion.

Discussion

Our study provides a number of key insights into the leukemogenic process in human AML. Our findings establish the sequential order of mutation acquisition for the patients reported here: *DNMT3a^{mut}* occurs before *NPM1c* and *FLT3-ITD*. Additionally, we provide strong evidence for the presence, at diagnosis, of preL-HSCs that are ancestral to the dominant AML clone. Based on our data, preL-HSC are prevalent among patients with *DNMT3a^{mut}*, which account for 25% of adult AML cases; additionally, our multilineage engraftment data suggest that preL-HSC may also exist in a proportion of AML patients with *IDH2* mutations. Pre-leukemic progenitors of varying phenotypes have been reported in other types of hematologic malignancies^{25,39–41}, although functional studies were limited. Our work supports prior studies identifying phenotypic primitive cells that bear only a subset of mutations found in AML blasts^{26,27,42}. The more precise analysis of highly resolved HSC and progenitor populations that we have undertaken provides novel insight into the identity and proportional contribution of the stem/progenitor populations that acquire pre-leukemic lesions. Furthermore, our work demonstrates that *DNMT3a^{mut}* confers a functional repopulation advantage to preL-HSC over wild type HSC in xenograft assays, which likely underlies the clonal expansion of preL-HSC observed in patients at the time of diagnosis. Our study is consistent with mouse studies showing that HSC lacking *DNMT3a* have a competitive growth advantage^{43,44}, and with a recent report predicting that the human *DNMT3a^{mut}* results in loss of function⁴⁵.

Collectively, our results support a model wherein the cell of origin for *DNMT3a^{mut}* AML is an HSC and the initiating *DNMT3a* mutation results in the generation of an expanded pool of HSC and downstream progenitors, within which additional mutations such as *NPM1c* are acquired, driving progression to AML. In the samples studied here, our findings point to GMP and/or MLP as the likely populations in which *NPM1c* was acquired. Our experimental design provides a framework for the future identification of other early events in leukemogenesis, using generation of multilineage xenografts as a surrogate assay to identify AML samples that may contain preL-HSC, and for examining how these changes disrupt normal HSC function, cause clonal expansion and initiate leukemic development.

Our results have broad clinical implications. Prior studies in T-ALL⁴⁶ and B-ALL^{7,11,14} revealed the existence of genetically diverse subclones at diagnosis. As found originally in these diseases⁴⁷ and now in AML, in approximately 50% of patients the relapse clone is not related to the predominant clone at diagnosis but rather to a minor leukemic subclone^{10,15} or to a predicted ancestral clone⁴⁷. Our direct demonstration that ancestral clones persist at

remission suggests that preL-HSCs are resistant to induction chemotherapy and for some patients they might represent a reservoir from which relapse arises. If future phylogenetic single cell lineage analysis establishes this possibility, then preL-HSC should be directly targeted to prevent relapse. As new drugs are developed that effectively target mutations in *DNMT3a* or other genes (e.g. AG-221, an *IDH2* inhibitor) that give rise to preL-HSC, there may be an opportunity to eradicate these preL-HSC clones before the acquisition of additional mutations renders them more resistant to therapy. Our findings also support broadening the definition of minimal residual disease to include not only the post-therapy survival of AML blasts and LSCs but also preL-HSC. Practically, this suggests that for patients with both *DNMT3a^{mut}* and *NPM1c*, the residual level of both mutations and not *NPM1c* alone should be monitored. Finally, our database analysis showing that the *DNMT3a* R882H variant is present in blood samples from normal adults should stimulate new studies investigating whether preL-HSC are present in healthy individuals and determining the risk of progression to AML; these may in turn enable earlier diagnosis for those patients who present without prior overt hematologic disturbances. Of note, a recent report describes the development of donor-derived leukemia in a patient 27 months after receiving allogeneic HSC from an HLA-matched sibling, whose donated PB cells were later found to carry mutations in *IDH2* and *DNMT3a* at low frequency⁴⁸. Interestingly, the donor remains free of leukemia 10 years after his stem cell donation, suggesting that the mechanisms underlying the progression from subclinical pre-leukemic hematopoiesis to overt leukemia are complex and context dependent.

Methods

Targeted sequencing of leukemia-associated genes

Genomic DNA (gDNA) was subjected to limited whole genome amplification (RepliG, Qiagen) to obtain the required amount of input DNA for the SureSelect protocol. Amplified gDNA was mechanically sheared using the Covaris M220 Focused-ultrasonicator, and Illumina sequencing adaptors were ligated to fragments to make a sequencing library, which was then hybridized with 120mer biotinylated RNA library baits to capture the regions of interest. Baits were designed to capture the coding sequence of the 103 leukemia-associated genes listed in Supplementary Table 1 (total target size ~370 Kb). The targeted regions were pulled out using magnetic streptavidin beads and amplified. The resulting amplified library was quantified and sequenced on the Illumina HiSeq 2000 platform to an average on target coverage of 250×. Reads were aligned to the reference human genome build hg19 using Novoalign (Novocraft Inc.) and on-target single nucleotide variants (SNVs) and indels were called using the genome analysis tool kit (GATK). Somatic SNVs were called in AML blasts with a read depth of at least 30×. The list of contributors to the gene list is provided in Supplementary Note 1.

T cell isolation and expansion from primary AML samples

CD3+ cells were isolated from peripheral blood (PB) AML patient samples using EasySep (Stem Cell Technologies) and re-suspended at a concentration of 1×10^7 cells/2 mL in RPMI + 10% FBS-HI + rhIL-2 (250 IU/mL, Proleukin, Chiron) + anti-CD28 antibody (5 ug/mL, clone CD28.2, eBioscience). Cells were then added to one well of a 24-well plate that had

been pre-coated for 2 hours with anti-CD3 antibody (Clone OKT3, eBioscience) and cultured for 4 days at 37°C with 5% CO₂. Cells were harvested on day 4, resuspended in fresh RPMI + 10% FBS-HI + rhIL2 (250 IU/ml) and replated into one well of a 6-well plate. Cells were further cultured and expanded for 14–20 days, feeding with fresh full medium containing rhIL-2 (250 IU/ml) every 3–4 days. At the end of T cell expansion, the purity of CD3⁺ T cells was checked by flow cytometry. DNA from the cultured T cells was extracted by PureGene Cell kit (Qiagen).

Droplet digital PCR (ddPCR)

Genomic DNA (25ng) or amplified DNA (2 ul from a 1:20 dilution of a 16 hour RepliG whole genome amplification) was subjected to ddPCR in a 96-well plate according to the manufacturer's protocol. Each sample was tested in duplicate. The plate was then loaded onto a droplet reader with a two color FAM/VIC fluorescence detector. The mutant allele frequency was calculated as the fraction of positive droplets divided by total droplets containing a target. The TaqMan probes and primers used for ddPCR are listed in Supplementary Table 5. To evaluate the detection limits of the ddPCR assay, a standard curve was generated using serial dilutions of DNA with a known mutation frequency mixed with non-mutated DNA. For probes listed in Supplementary Table 5, the minimum detection level was 1:1000 (0.1%).

Fluorescence activated cell sorting of human stem/progenitor and mature cell populations

Mononuclear cells ($1 \times 10^6/100 \mu\text{l}$) from peripheral blood or bone marrow of AML patients were stained with the following antibodies (all from BD unless stated otherwise, dilution used and catalogue number in parentheses): anti-CD45RA-FITC (1:25, 555488), anti-CD90-APC (1:50, 561971), anti-CD135-Biotin (1:10, 624008), anti-CD38-PE-Cy7 (1:200, 335790), anti-CD10-Alexa-700 (1:10, 624040), anti-CD7-Pacific Blue (1:50, 642916), anti-CD45-V500 (1:200, 560777), anti-CD34-APC-Cy7 (1:100, custom made by BD, CD34 clone 581), anti-CD34-PerCP-Efluor 710 (1:100, e-Bioscience 46-0344-42), anti-CD33-PE-Cy5 (1:100, Beckman Coulter PNIM2647U), anti-CD19-PE (1:200, 349204), anti-CD3-FITC (1:100, 349201), anti-CD56-Alexafluor 647 (1:100, 557711), and Streptavidin-QD605 (1:200, Invitrogen Q10101MP). Samples from Patients #1, 10, 11 (remission sample only), 32, 35, and 55 were enriched for CD34⁺ cells using a Miltenyi CD34 MicroBead kit according to the manufacturer's protocol prior to antibody staining. Cells were sorted on a FACS AriaIII to a post-sort purity of >95%.

Xenotransplantation assays

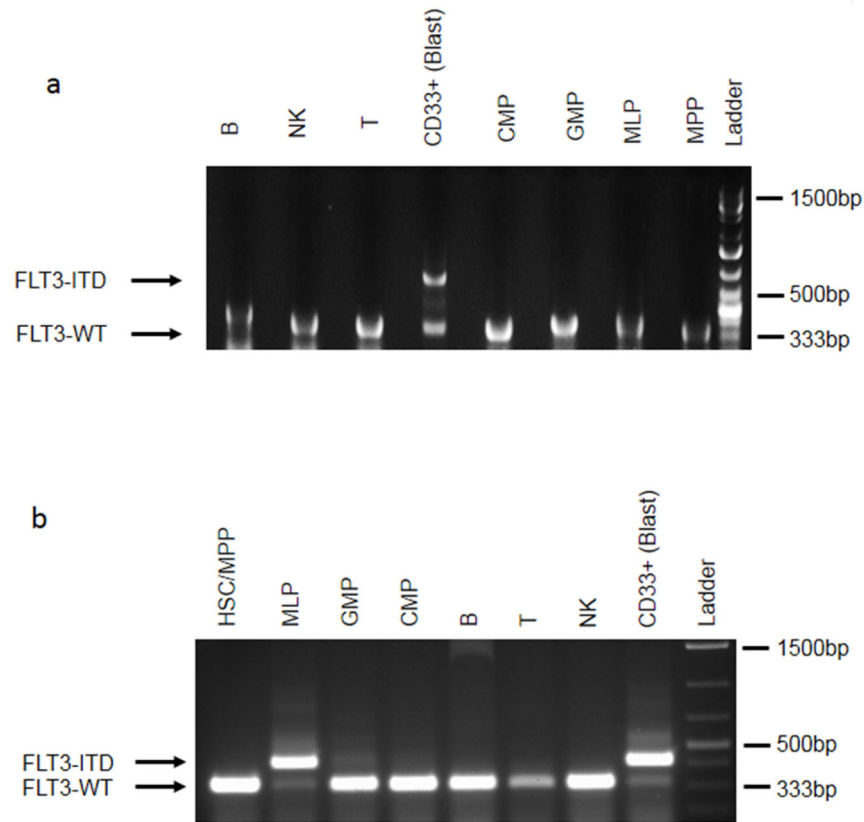
Animal experiments were performed in accordance with institutional guidelines approved by the UHN Animal Care Committee. 8 to 12-week-old female NOD/SCID/IL-2R_g^c-null (NSG) mice were sublethally irradiated (225 cGy) 6–24 hours before transplantation. Mononuclear cells from AML patients were depleted of CD3⁺ cells by EasySep (Stem Cell Technologies) prior to intrafemoral transplantation. Mice were sacrificed 8 or 16 weeks after transplantation and human engraftment in the injected femur and non-injected bone marrow was evaluated by flow cytometry using the following human-specific antibodies (all used at 1:200, all from BD unless stated otherwise, catalogue number in parentheses): anti-CD45-APC (340943), anti-CD19-PE, anti-CD33-PE-Cy5, anti-CD3-FITC, anti-CD14-PE Texas

Red (Beckman Coulter PNIM2707U), anti-CD15-Pacific Blue (642917), anti-CD38-PE-Cy7, and anti-CD34-APC-Cy7. The threshold for detection of human engraftment was 0.1% CD45+ cells. All flow cytometric analysis was performed on the LSRII (BD Biosciences). For limiting dilution assays, the frequency of repopulating cells was calculated using ELDA software⁴⁹.

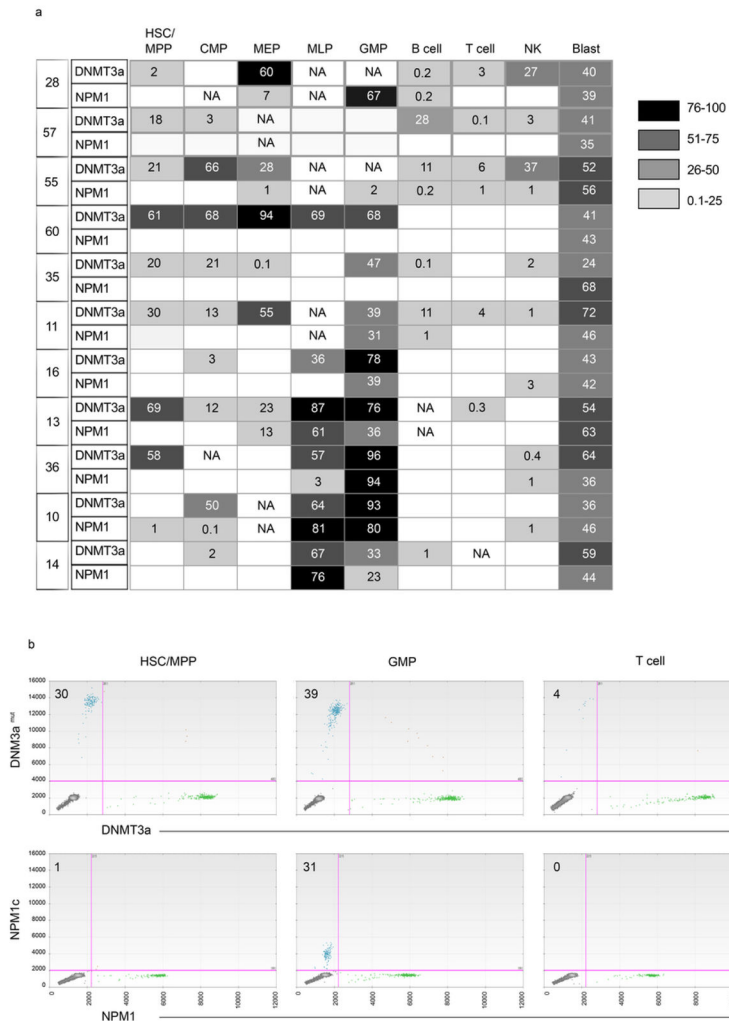
Statistical analysis

For the initial targeted sequencing analysis, 12 independent patient samples were studied to capture the biologic diversity of AML. For validation of the *DNMT3a* findings, 71 samples were screened in order to identify at least 15 with *DNMT3a* mutations, as predicted by the known prevalence of *DNMT3a* mutation in AML. For limiting dilution analyses, at least 25 xenografts were analyzed for each patient sample to ensure a large enough sample for statistical comparison. No animals or samples were excluded from any analysis. No formal randomization method was applied when assigning animals to different experimental groups. Group allocation and outcome assessment was not done in a blinded manner, including for animal studies. Frequency estimations were generated using the ELDA software, which takes into account whether the assumptions for LDA are met (<http://bioinf.wehi.edu.au/software/elda/index.html>, provided by the Walter and Eliza Hall Institute)⁴⁹. P-values were derived using two-tailed Student's t-tests. In each group of data, estimate variation was taken into account and is indicated as standard deviation. For all graphs, * p=0.01–0.05, ** p=0.001–0.01, and *** p<0.001.

Extended Data

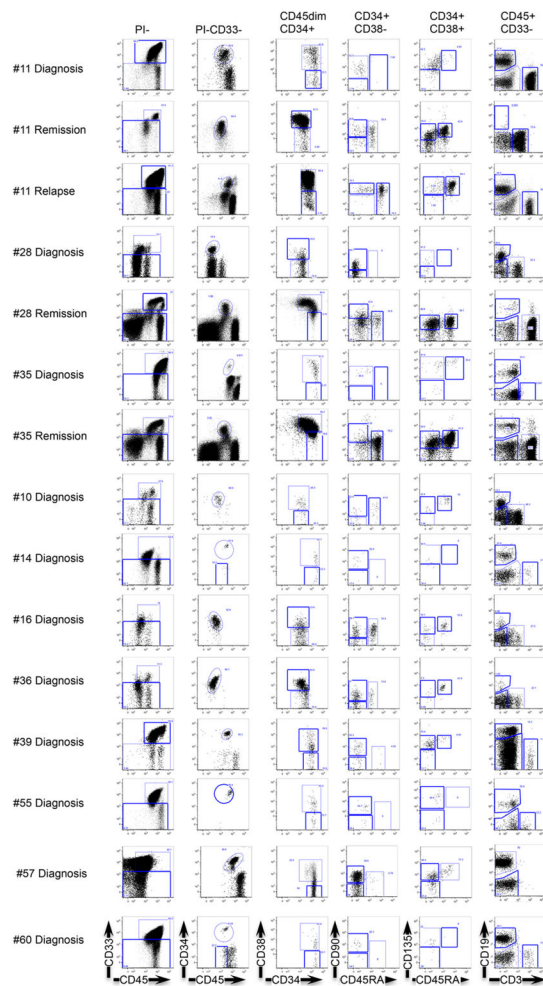


Extended Data Figure 1. *FLT3*-ITD is a late event in patients carrying *DNMT3A* mutation PCR analysis of *FLT3*-ITD⁵⁰ in stem/progenitor, mature lymphoid and blast (CD45^{dim} CD33⁺) cell populations from patient no. 13 (a) and no. 14 (b). *FLT3*-ITD was present in the blasts from both patients, and also in MLPs from patient no. 14. In contrast, *DNMT3A*^{mut} without *FLT3*-ITD was detected in multiple non-blast cell populations (see Extended Data Fig. 2). HSC, haematopoietic stem cell; MPP, multipotent progenitor; CMP, common myeloid progenitor; MLP, multilymphoid progenitor; GMP, granulocyte monocyte progenitor; NK, natural killer cells.



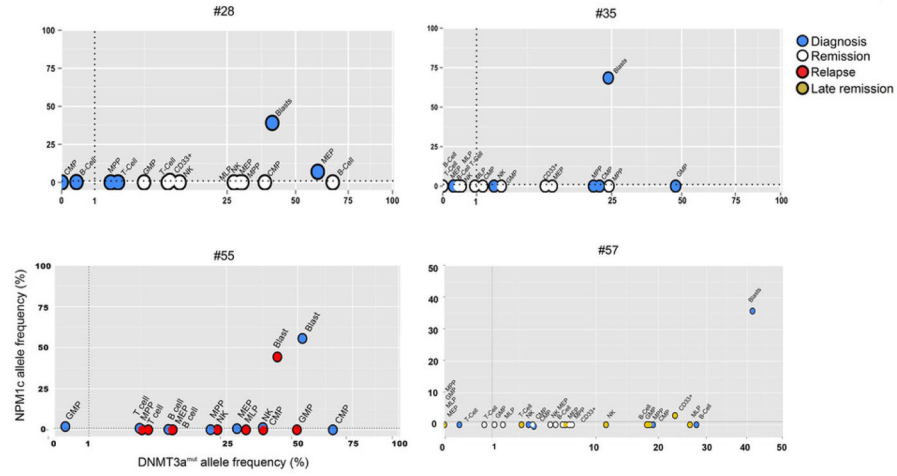
Extended Data Figure 2. Frequent occurrence of *DNMT3A* mutation without *NPM1* mutation in stem/progenitor and mature lymphoid cells in AML patients at diagnosis

a. Summary of the allele frequency (%) of *DNMT3A* and *NPM1* mutations in stem/progenitor, mature lymphoid, and blast ($CD45^{dim} CD33^{+}$) cell populations from 11 AML patient peripheral blood samples obtained at diagnosis, as determined by droplet digital PCR (ddPCR). Phenotypically normal cell populations were isolated by fluorescence activated cell sorting according to the strategy depicted in Fig. 2a. Mutant allele frequency ~50% is consistent with a heterozygous cell population. Departures from 50% mutant allele frequency may be stochastic⁵¹, related to clonal heterogeneity, or due to the presence of copy number variations, for example loss of the wild type allele (loss of heterozygosity) or amplification of the mutant allele. NA, no cell population detected; HSC, haematopoietic stem cell; MPP, multipotent progenitor; CMP, common myeloid progenitor; MEP, megakaryocyte erythroid progenitor; MLP, multilymphoid progenitor; GMP, granulocyte monocyte progenitor; NK, natural killer cells. Blank boxes indicate no *DNMT3A* or *NPM1* mutation detected. **b.** Representative plots showing ddPCR analysis of *DNMT3A*^{mut} and *NPM1c* allele frequency in sorted cell populations from patient no. 11. The mutant allele frequency (%) is indicated on each plot.

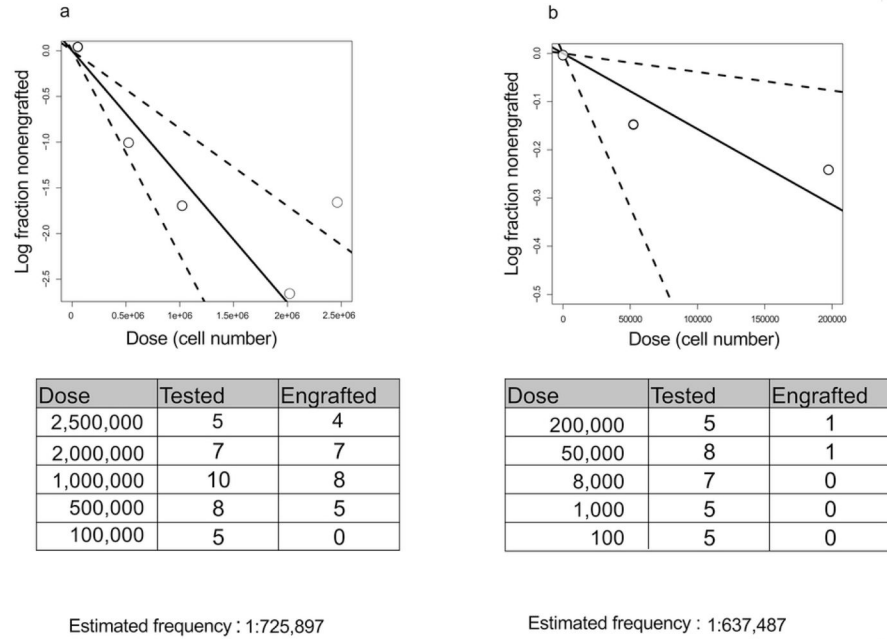


Extended Data Figure 3. Phenotypically normal stem/progenitor and mature cell populations are present in AML patient samples at diagnosis, remission and relapse

Flow cytometric analysis showing the gating strategy used to isolate phenotypically normal stem/progenitor and mature lymphoid cell populations from AML patient samples. Diagnosis and relapse samples are from peripheral blood; remission samples are from bone marrow.

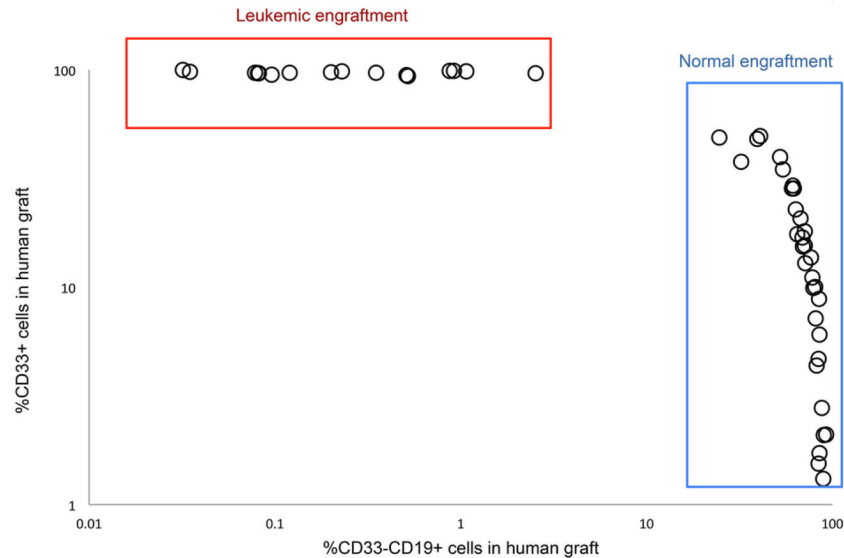


Extended Data Figure 4. Cells bearing mutations in *DNMT3A* but not *NPM1* are present at diagnosis in AML patients and persist at remission and relapse
 Allele frequency of *DNMT3A* and *NPM1* mutations of patients no. 28, 35, 55, and 57 in stem/progenitor, mature and blast ($CD45^{dim} CD33^{+}$) cell populations, as determined by droplet digital PCR (ddPCR). Cells were isolated from diagnosis (blue), early remission (white), relapse (red) or late remission (yellow) samples. At remission, $CD33^{+}$ myeloid cells were also analysed. HSC, haematopoietic stem cell; MPP, multipotent progenitor; MLP, multilymphoid progenitor; CMP, common myeloid progenitor; GMP, granulocyte monocyte progenitor; MEP, megakaryocyte erythroid progenitor; NK, natural killer cells.



Extended Data Figure 5. PreL-HSCs in the peripheral blood of AML patients generate multilineage human grafts in immunodeficient mice
 Summary of results of limiting dilution experiments to assess frequency of pre-leukaemic HSCs generating multilineage grafts after xenotransplantation. Cohorts of NSG mice were

transplanted intrafemorally with varying numbers of peripheral blood mononuclear cells from diagnostic samples of AML patient no. 11 (a) and no. 55 (b) and analysed after 8 or 16 weeks by flow cytometry. Engraftment was defined as $>0.1\%$ human $CD45^+$ cells in the injected right femur. Shown is the number of mice with multilineage human grafts containing both $CD33^+$ myeloid cells and $CD33^-CD19^+$ cells. The frequency of pre-leukaemic HSCs was calculated using the ELDA platform⁴⁹.



Extended Data Figure 6. Frequent generation of non-leukaemic multilineage human grafts following xenotransplantation of peripheral blood cells from AML patients

Summary of xenograft characteristics in 123 sublethally irradiated NSG mice transplanted intrafemorally with mononuclear peripheral blood cells from 20 AML patients at diagnosis and analysed after 8 weeks by flow cytometry. The proportion of myeloid ($CD33^+$) and B-lymphoid ($CD33^-CD19^+$) cells in the human ($CD45^+$) graft is shown. Leukaemic (AML) engraftment is characterized by a dominant myeloid ($CD45^{dim}CD33^+$) graft, whereas non-leukaemic multilineage grafts contain both lymphoid (predominantly $CD33^-CD19^+$ B cells) and myeloid ($CD33^+$) cells. No leukaemic or multilineage graft could be detected in 65/123 mice (53%) in this cohort. Red box indicates AML grafts (27 mice, 22%); blue box indicates multilineage grafts (31 mice, 25%).

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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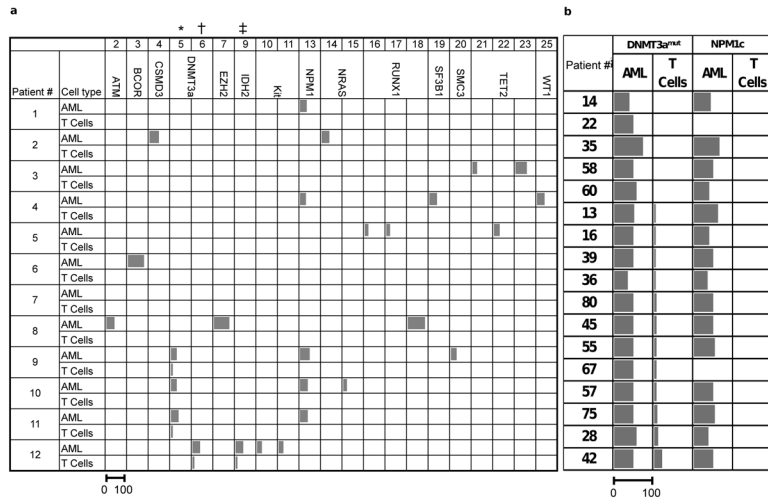


Figure 1. Recurrent somatic *DNMT3a* mutations are common in T-cells from AML patients
a, Summary of the allele frequency (%) of missense and frameshift somatic single nucleotide variants (sSNV) in AML-related genes assessed by deep targeted sequencing (read depth 250×) in AML blasts and T cells from the peripheral blood of 12 AML patients. The sSNV numbers indicated at the top of the table correspond to the numbers in Supplementary Table 3. Somatic mutations in *DNMT3a* (*, R882H; †, R137C) were found in both T-cells and AML blasts in Patients #9, 11 and 12. Patient #12 also had a low frequency *IDH2* mutation (‡, R140L) in T cells. **b**, Frequency (%) of *DNMT3a*^{mut} and *NPM1c* in freshly isolated CD33+ blasts (AML) and matched T-cell controls from 17 patients with normal karyotype AML, as determined by droplet digital PCR. For **a** and **b**, the length of the bars is proportional to the mutant allele frequency (the scale bar under the first column applies to all columns).

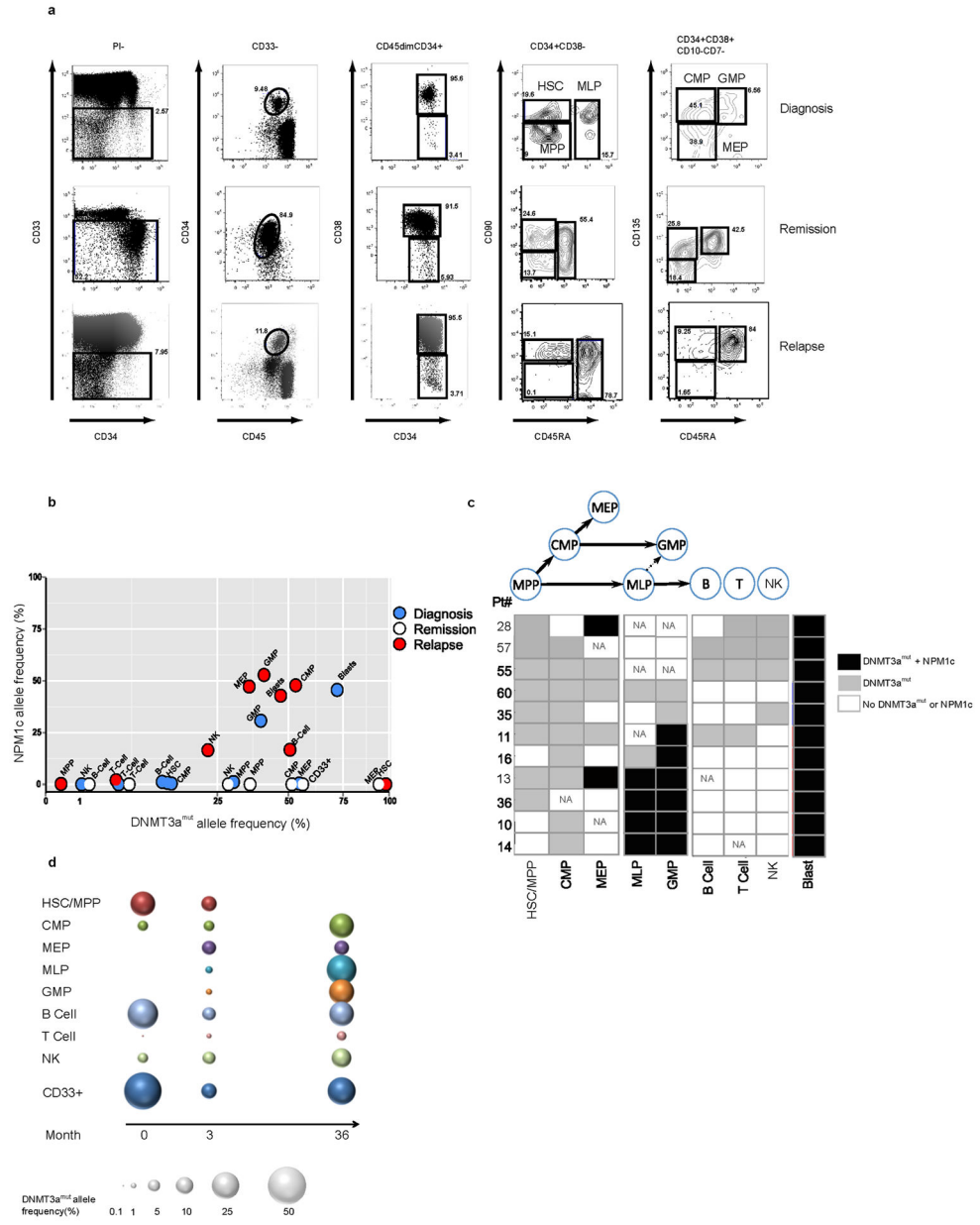


Figure 2. DNMT3a mutation precedes NPM1 mutation in human AML and is present in stem/progenitor cells at diagnosis and remission

a, Flow cytometric analysis showing the gating strategy used to isolate phenotypically normal stem and progenitor cell populations from AML patient samples. Plots show analysis of samples from Patient #11: diagnosis (day 0, peripheral blood mononuclear cells), remission (day 62, CD34+ enriched bone marrow) and relapse (day 379, peripheral blood mononuclear cells). **b**, Allele frequency of *DNMT3a* and *NPM1* mutations in stem/progenitor, mature lymphoid and blast (CD45dim CD33+) cell populations, as indicated, isolated from diagnosis (blue), remission (white) and relapse (red) samples of Patient #11 as determined by droplet digital PCR (ddPCR). At remission, CD33+ myeloid cells were also analyzed. **c**, Summary of the occurrence of *DNMT3a^{mut}* and *NPM1c* in isolated stem/

progenitor, mature and blast cell populations from 11 AML patient peripheral blood samples as determined by ddPCR. White, *DNMT3a^{mut}* or *NPM1c* not detected; gray, *DNMT3a^{mut}* alone; black, *DNMT3a^{mut}* + *NPM1c*. NA, no population detected; HSC, hematopoietic stem cell; MPP, multipotent progenitor; MLP, multilymphoid progenitor; CMP, common myeloid progenitor; GMP, granulocyte monocyte progenitor; MEP, megakaryocyte erythroid progenitor; NK, natural killer cells. **d**, Graphic representation of *DNMT3a^{mut}* allele frequency in sorted cell populations isolated from diagnosis (0 months), early (3) and late (36) remission samples of Patient #57.

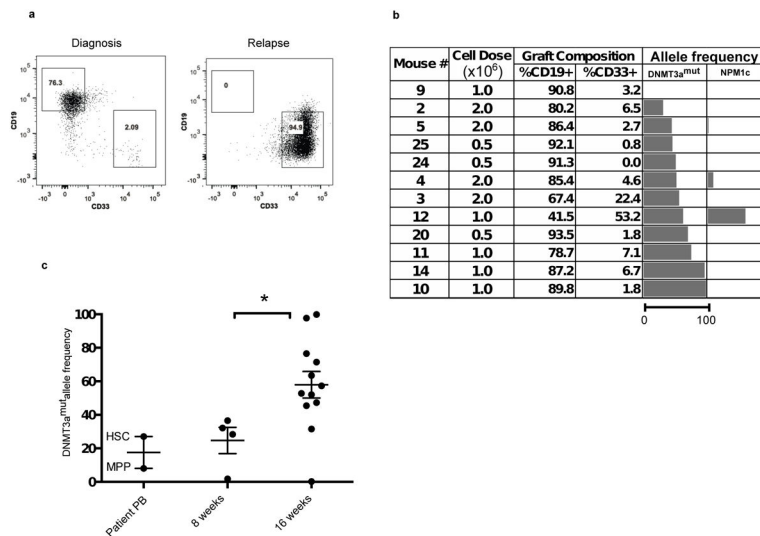


Figure 3. Pre-leukemic HSC bearing *DNMT3a^{mut}* generate multilineage engraftment and have a competitive advantage in xenograft repopulation assays

a, Representative flow cytometric analysis of engrafted human cells harvested from NSG mouse bone marrow (BM) 16 weeks after intrafemoral (i.f.) transplantation of peripheral blood mononuclear cells (PB MNC) from diagnosis and relapse samples of Patient #11. **b**, Analysis of human graft composition in NSG mouse BM 16 weeks after i.f. transplantation of PB MNC from the diagnosis sample of Patient #11 across a range of cells doses. The percentage of human (CD45+) B (CD19+) and myeloid (CD33+) cells was determined by flow cytometry. Mutant allele frequency (%) in the human graft was determined by droplet digital PCR (ddPCR) analysis of sorted human cells. The length of the bars is proportional to the mutant allele frequency (the scale bar under the first column applies to all columns). **c**, Summary of *DNMT3a^{mut}* allele frequency in the human graft from mice analyzed by ddPCR 8 and 16 weeks after transplantation of PB MNC from Patient #11, compared to isolated hematopoietic stem cell/multipotent progenitors (HSC/MPP) from the patient's PB at diagnosis. *, P<0.05. Bars indicate mean and standard deviation.

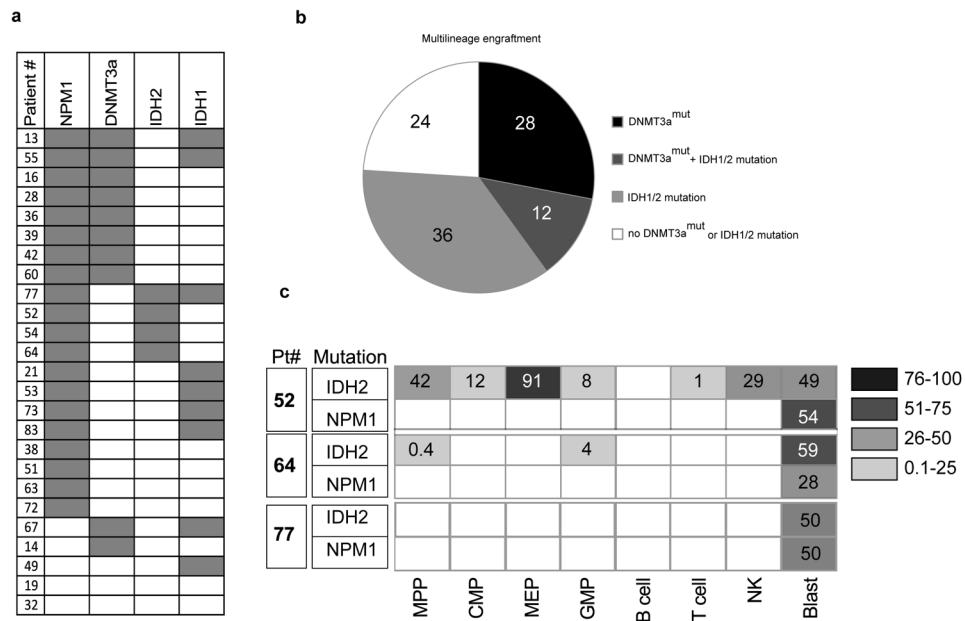


Figure 4. Identification of preL-HSC with *IDH2* mutation

a, Summary of the occurrence of mutations in *NPM1*, *DNMT3a*, and *IDH1/2* determined by Sanger sequencing, in AML patient peripheral blood samples (n=25) that generated a non-leukemic multilineage graft after transplantation into immune-deficient mice. **b**, Representation of the proportion (%) of AML patient samples with *DNMT3a* and/or *IDH1/2* mutations among samples that generated a non-leukemic multilineage graft in xenotransplanted mice. **c**, *IDH2* and *NPM1* mutant allele frequency (%) in stem/progenitor, mature lymphoid and blast (CD45dimCD33+) cell populations isolated from the peripheral blood of Patients #52, 64 and 77 at diagnosis, as determined by droplet digital PCR (ddPCR). Blank boxes indicate no mutation detected.