

A novel tumour-suppressor function for the Notch pathway in myeloid leukaemia

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Notch signalling is a central regulator of differentiation in a variety of organisms and tissue types¹. Its activity is controlled by the multi-subunit γ -secretase (γ SE) complex². Although Notch signalling can play both oncogenic and tumour-suppressor roles in solid tumours, in the haematopoietic system it is exclusively oncogenic, notably in T-cell acute lymphoblastic leukaemia, a disease characterized by Notch1-activating mutations³. Here we identify novel somatic-inactivating Notch pathway mutations in a fraction of patients with chronic myelomonocytic leukaemia (CMML). Inactivation of Notch signalling in mouse haematopoietic stem cells (HSCs) results in an aberrant accumulation of granulocyte/monocyte progenitors (GMPs), extramedullary haematopoiesis and the induction of CMML-like disease. Transcriptome analysis revealed that Notch signalling regulates an extensive myelomonocytic-specific gene signature, through the direct suppression of gene transcription by the Notch target *Hes1*. Our studies identify a novel role for Notch signalling during early haematopoietic stem cell differentiation and suggest that the Notch pathway can play both tumour-promoting and -suppressive roles within the same tissue.

Notch is essential for the emergence of definitive haematopoiesis⁴, controls HSC differentiation to the T-cell lineage^{5,6} and is a major oncogene, as most patients with T-cell acute lymphoblastic leukaemia harbour activating *Notch1* mutations⁷. To study haematopoiesis in the absence of any Notch-derived signal (as mammals express four different Notch receptors), we targeted Nicastrin (*Ncstn*), a member of the γ SE complex and one of the few non-redundant members of the pathway (Supplementary Fig. 1). We crossed the *Ncstn*^{fl/fl} mice to both an inducible (*Mx1-cre*)⁸ and a haematopoietic-specific (*Vav-cre*)⁹ recombinase strain. Both modes of deletion (referred to herein as *Ncstn*^{-/-}) produced identical phenotypes. Unexpectedly, none of the *Ncstn*^{-/-} mice survived longer than 20 weeks. Further analysis revealed a striking peripheral blood leukocytosis and monocytosis with enlargement of the spleen (Fig. 1a, b and Supplementary Figs 2 and 3). Histological analysis of the spleen showed a marked expansion of the red pulp with diffuse infiltration by myeloid and monocytic cells. The infiltrating myeloid cells were partly myeloperoxidase positive and CD11b⁺ and/or Gr1⁺ (Fig. 1b and Supplementary Figs 2 and 3). The increase in monocyte numbers was also observed in the bone marrow and liver (Fig. 1a and data not shown). Taken together, these findings were diagnostic for a myeloproliferative/myelodysplastic process and reminiscent of human CMML. CMML is a myeloid malignancy, classified as a myeloproliferative disorder/myelodysplastic syndrome overlap, which is characterized by monocytosis, myeloproliferation, variable bone marrow dysplasia and a high rate of progression to acute myeloid leukaemia¹⁰.

Because monocytes and granulocytes originate from the GMP subset, we examined the stem and progenitor cell populations in the bone marrow. *Ncstn* deletion led to an enlargement of the Lineage^{neg}Sca1⁺c-Kit⁺ (LSK), specifically the LSK CD150⁺CD48⁺ subset, a population shown to have a myeloid commitment bias¹¹. This differentiation bias was coupled to a significant reduction of the lymphoid-biased multipotential progenitor population (L-MPP)¹² (Supplementary Fig. 4). Moreover, there was a striking increase in the absolute numbers of both bone marrow and spleen GMP cells (Fig. 1d) coupled to a decrease of the megakaryocyte-erythrocyte progenitor population. This apparent predisposition towards GMP-derived lineages was also evident *in vitro*, as *Ncstn*^{-/-} progenitors generated more granulocyte-macrophage and macrophage colonies (Supplementary Fig. 5). Further studies also revealed a striking ability of the *Ncstn*^{-/-} progenitors to replate serially (Supplementary Fig. 6), suggesting an increase in their self-renewal potential. Consistent with this idea, whole transcriptome profiling of *Ncstn*^{-/-} GMP progenitors revealed enrichment of a 'leukemic self-renewal' signature¹³. Finally, bone marrow transplantation assays demonstrated that the effects of *Ncstn* deletion were cell autonomous (Supplementary Fig. 7)¹⁴.

Although the γ SE complex has other substrates², we focused on Notch signalling because of our finding that *Ncstn* deletion led to known Notch^{-/-} phenotypes, including a block in T-cell differentiation (Supplementary Fig. 8)¹⁵. To prove a connection to Notch signalling, we generated animals that conditionally lack the expression of three out of four Notch receptors (*Mx1-cre*⁺*Notch1*^{fl/fl}*Notch2*^{fl/fl}*Notch3*^{-/-})¹⁶. Strikingly, triple *Notch1/2/3* deletion copied the *Ncstn*^{-/-} phenotypes (Supplementary Fig. 9). *Notch1-3*^{-/-} mice developed both CMML-like symptoms, and significant enlargement of the GMP population. *Notch3* expression was dispensable, as simultaneous deletion of only *Notch1* and *Notch2* led to an identical CMML-like pathology. However, introduction of a single wild-type (WT) *Notch1* or *Notch2* allele was able to suppress the disease phenotype. Consistent with the importance of Notch1–2 receptor signalling in these stages, quantitative PCR studies revealed expression of *Notch1* and *Notch2*, but not *Notch3*, in WT stem and progenitor cells (Supplementary Fig. 10).

We next sought to delineate the mechanism by which Notch directs the regulation of early haematopoiesis. Deletion of *Ncstn* did not lead to any alterations in the GMP cell-cycle status or cell death rate (not shown). We hypothesized that γ SE complex/Notch signals actively suppress a GMP-specific gene expression program. We sorted LSK and GMP cells and studied their transcriptome. This analysis revealed a statistically significant de-repression of an extended myeloid-specific program¹² in *Ncstn*^{-/-} LSK cells (Fig. 2a and Supplementary Fig. 11).

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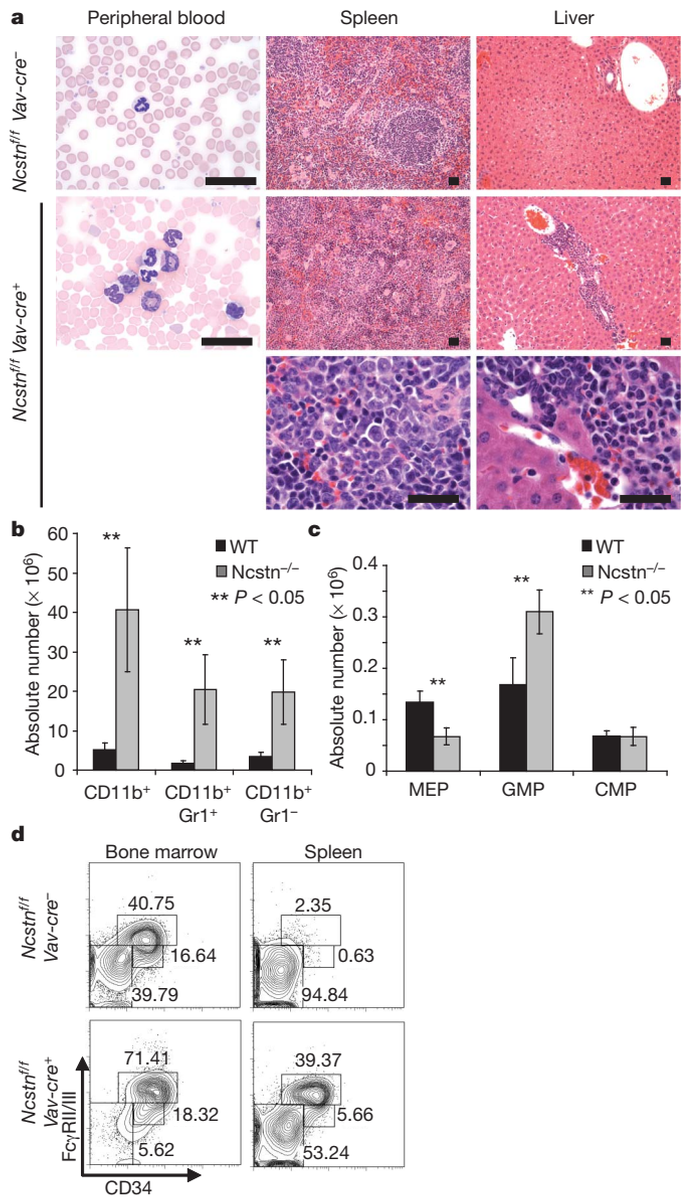


Figure 1 | *Ncstn* deficiency leads to CMML-like disease and a significant enlargement of the GMP progenitor population. **a**, Histological analysis showing accumulation of monocytes and granulocytes in peripheral blood (Wright–Giemsa staining), spleen and liver (haematoxylin and eosin staining). A magnification of each infiltrant is shown in the lowest panels. **b**, Absolute numbers of each monocytic/granulocytic subset from the spleen of control and *Ncstn^{fl/fl} Vav-cre⁺* littermate animals (12 weeks of age, mean \pm s.d., $n = 10$). **c**, Absolute numbers of each progenitor subpopulation in the bone marrow (mean \pm s.d., $n = 10$). **d**, Detailed FACS analysis of bone marrow and spleen myeloid progenitor (myeloid progenitors: Lin⁻/c-Kit⁺/Sca-1⁻) populations of *Ncstn^{fl/fl} Vav-cre⁺* and *Ncstn^{fl/fl} Vav-cre^{-/-}* littermates showing a significant enlargement of the GMP (Fc γ RII/III⁺, CD34⁺) subset.

Gene-set enrichment analysis demonstrated a significant enrichment of myeloid-specific gene-sets within the *Ncstn^{-/-}* LSK gene signature (Fig. 2b and Supplementary Table 1). Further dissection of the LSK subset showed that the GMP gene expression program was initiated as early as the CD150⁺ HSC stage of differentiation, and persisted at the CD150⁻ subset, which included MPPs (Supplementary Fig. 11). Furthermore, we were able to show that gene expression in LSKs purified from an inducible Notch1 gain-of-function genetic model (*Ef1 α 1-lsl-Notch1^{IC}Mx1-cre⁺*)¹⁷ was inversely correlated with the loss-of-function signature seen in *Ncstn^{-/-}* LSK progenitors (Fig. 2a). Notch1^{IC} expression led to suppression of myeloid-specific genes, suggesting that activation of Notch

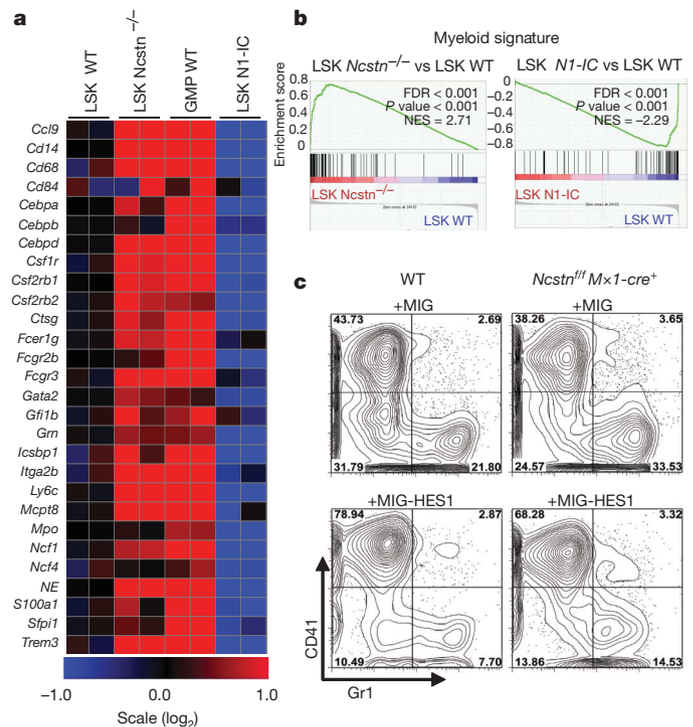


Figure 2 | Notch signalling suppresses an extensive myeloid gene expression program through the induction of the transcriptional repressor Hes1.

a, Heat map showing regulation of genes representative of the myeloid signature from the indicated cell populations and mice. **b**, Expression data were analysed for lists of genes positively involved in myelopoiesis using gene-set enrichment analysis. Enrichment plots show upregulation of myeloid-specific genes in *Ncstn^{fl/fl}Mx1-cre⁺* and downregulation of Notch1^{IC} *Mx1-cre⁺* LSK cells (compared with WT counterparts). **c**, Purified cKit⁺ progenitors from WT and *Ncstn^{fl/fl}Mx1-cre⁺* mice were transduced with retroviruses encoding Hes-1 or empty vector, subsequently plated on methylcellulose for 7 days and analysed for expression of myeloid or megakaryocyte differentiation markers (Gr1, CD41). A representative of four experiments is shown.

signalling can alter the transcription and differentiation of uncommitted HSC and MPP cells. Gene-set enrichment analysis further supported these findings as it showed a negative correlation between the Notch1^{IC} LSKs and previously reported myeloid gene expression signatures (Fig. 2a, b, Supplementary Table 2 and Supplementary Fig. 11).

As Notch is thought to function primarily as a transcriptional activator, we hypothesized that its suppressive effects on GMP-specific gene expression could be explained by the induction of a transcriptional repressor. A search for such a molecule revealed *Hes1*, a known Notch target and a transcriptional repressor (Supplementary Fig. 11a). Interestingly, *Hes1* was previously suggested to play a role in myeloid leukaemia, as a downstream effector of the *Junb* tumour suppressor in an animal model of chronic myeloid leukemia¹⁸. To test the potential involvement of *Hes1* directly, we used *in vitro* differentiation assays and showed that *Hes1* ectopic expression was sufficient to direct differentiation away from the granulocyte/monocyte lineage (Fig. 2c). Furthermore, *Hes1* expression suppressed the expression of key granulocyte/monocyte commitment genes such as *Cebpa* and *Pu.1* (Supplementary Fig. 12a). In agreement with this finding, we identified putative *Hes* (N-box) binding sites on the promoters of both genes. Reporter and chromatin immunoprecipitation assays proved direct binding of *Hes1* on these promoters and suppression of transcription (Supplementary Fig. 12b–e).

These findings suggested that Notch (or *Hes1*) hyper-activation could suppress CMML-like disease developing in *Ncstn^{-/-}* animals. To test this, we used the previously described *Ef1 α 1-lsl-Notch1^{IC}* mice¹⁷. We generated *Notch1^{IC}Ncstn^{fl/fl}Mx1-Cre⁺* animals and analysed both GMP accumulation and disease progression. Notch1^{IC} expression was

sufficient to suppress both GMP expansion and disease development significantly (Fig. 3a, b and Supplementary Fig. 13). Interestingly, *Notch1^{IC}* expression drove progenitor commitment towards the lymphoid (T cell) and megakaryocyte–erythrocyte progenitor lineages¹⁹. However, *Notch1^{IC}* expression did not affect cell-cycle kinetics within the GMP subset (Supplementary Fig. 13). To uncouple differentiation to the GMP subset from effects on GMP homeostasis, we purified WT GMP and plated them on stroma in the presence or absence of Notch ligands (Dll1 and Dll4). A striking increase of apoptosis rate was noted in the presence of Dll1–4 (Fig. 3c). These experiments suggest that Notch ectopic expression can affect both the commitment to the GMP subset and the survival of already committed GMP progenitors.

Our studies so far demonstrate that Notch controls murine myelopoiesis and that its deletion leads to GMP expansion and monocytic disease. To prove that Notch is also important in human haematopoiesis and leukemogenesis, we initially cultured purified human *CD34⁺CD38⁻Lin⁻* bone marrow and cord blood stem and progenitor cells on stroma expressing different Jagged and Delta-like Notch ligands²⁰. We found that expression of Notch ligands efficiently suppressed differentiation of human multipotential progenitors towards both the granulocyte (*CD15⁺*) and monocyte (*CD14⁺*) lineages (Supplementary Figs 14 and 15). This suppression was prevented when the activity of the γ SE complex was suppressed using either small molecule inhibitors or by the expression of a dominant negative *MAML1* mutant.

To gain further insights into the role of Notch in human CMML, we extensively sequenced many γ SE/Notch pathway genes. Exon resequencing of specimens from patients with CMML (Supplementary Table 3) identified a substantial fraction (six novel mutations in 5 out of 42 patients) harbouring somatic heterozygous mutations in

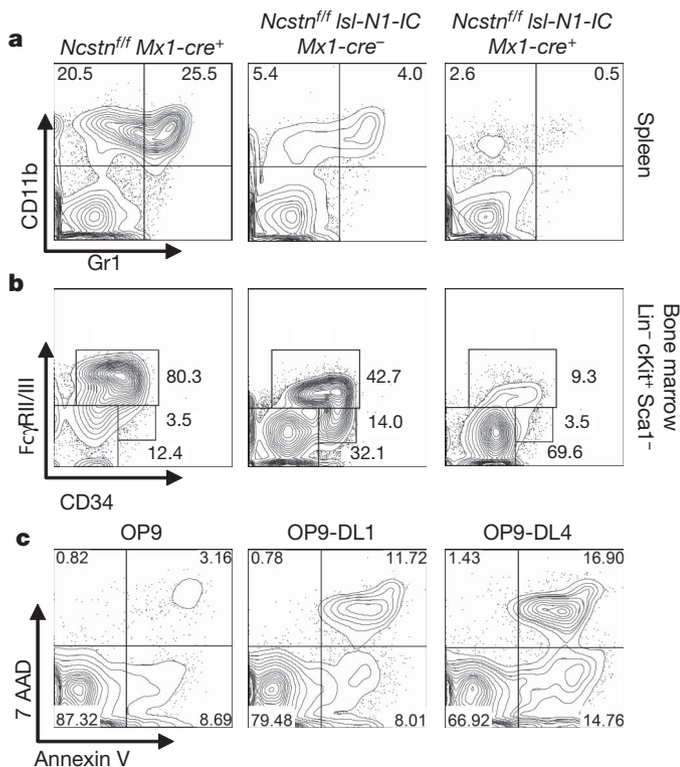


Figure 3 | Ectopic expression of Notch1-IC is able to prevent CMML-like disease in *Ncstn^{-/-}* mice. **a**, PolyI:polyC-induced Notch1-IC expression in *Ncstn^{fl/fl}Isl-N1-IC Mx1-Cre⁺* animals suppresses myeloid cells in spleen. **b**, Notch1-IC expression suppresses GMP progenitor population in bone marrow. **c**, Induction of cell death in WT GMP cells cultured in the presence (OP9-DL1–4) or absence (OP9) of Notch ligands. Cell death was measured by the combination of 7AAD and annexin V staining 48 h after co-culture initiation. For **a–c**, a representative of more than three experiments is shown.

multiple Notch pathway genes including *NCSTN*, *APH1*, *MAML1* and *NOTCH2* (Fig. 4 and Supplementary Table 4). In addition, several other putative mutations (single nucleotide variants) were detected, which are not annotated as known germline single nucleotide polymorphisms but for which we could not prove a somatic origin (Supplementary Table 5). The validated somatic mutations were only observed in CMML, as re-sequencing of 47 samples from patients with myeloproliferative disorders (polycythemia vera or myelofibrosis) did not reveal somatic mutations in the Notch pathway (Fig. 4b). Importantly, CMML specimens with Notch mutations also had somatic alterations in well-characterized myeloid oncogenic lesions²¹,

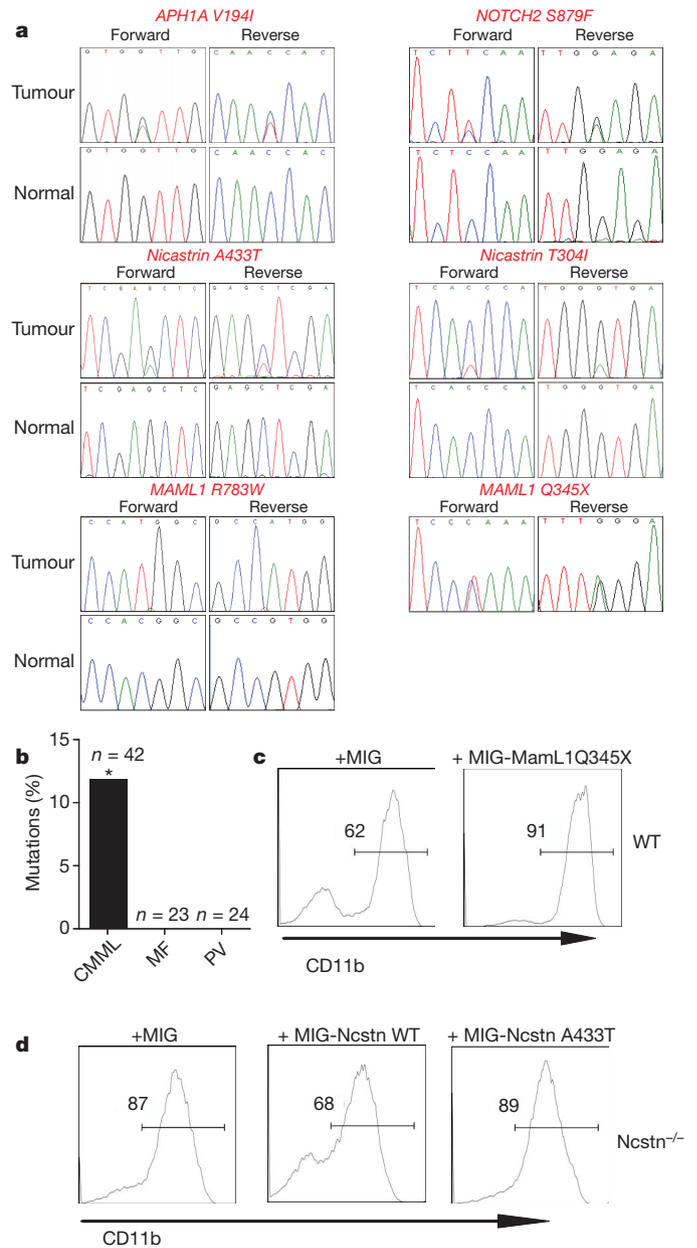


Figure 4 | Novel, loss-of-function Notch pathway mutations in human CMML. **a**, Sequence traces of identified Notch pathway mutations in tumours from patients with CMML but not in normal tissues show somatic origin. **b**, Comparison of the percentage of Notch pathway mutations in specimens from patients with CMML, myelofibrosis and polycythemia vera. The asterisk denotes that only verified somatic CMML mutations are included. **c**, OP9-DL1 co-culture of WT LSK cells infected with specified constructs. Analysis of the *CD11b⁺* population was studied 14 days after the initiation of the culture. **d**, A similar experiment as in **c**, using LSK *Ncstn^{-/-}* progenitors infected with the specified constructs. In all cases, a representative of more than three experiments is shown.

including *JAK2*, *KRAS*, *TET2* and *ASXL1*, suggesting mutational cooperation between Notch signalling and other oncogenic pathways in CMML (Supplementary Table 4). We then asked if those mutations were causally related with the disease by using transcriptional reporter and *in vitro* differentiation assays. We demonstrated that selected mutations had the ability to affect Notch activity negatively either as dominant negative (MAML1Q345X) or null (NCSTNA433T) alleles (Fig. 4c, d and Supplementary Fig. 16). This is the first description, to our knowledge, of somatic Notch pathway loss-of-function mutations in human cancer.

The presented studies identify novel inactivating Notch pathway mutations and suggest that γ SE complex/Notch signalling controls early HSC/MPP commitment decisions in bone marrow. A significant portion of this regulation is controlled by the Hes family of transcriptional repressors. Most importantly, our studies suggest that silencing Notch activity leads to the development of myeloid leukaemia, implying a novel tumour-suppressor function for the Notch pathway in haematopoiesis. Although mutation-mediated pathway silencing can be found in CMML, it is conceivable that there are additional control mechanisms, including epigenetic silencing of Notch pathway target genes. Whatever the mode of regulation, our observations suggest that reversible activation of the Notch pathway may represent an attractive future therapy, targeting specifically the progression and relapse of granulocytic and monocytic neoplasms.

METHODS SUMMARY

Animals. All mice were kept in specific pathogen-free animal facilities at the New York University School of Medicine. Mx1-Cre⁺ animals were injected with 20 μ g polyI:polyC per gram of body weight for a total of six injections. The injections were initiated 14 days after birth and done every 2 days. Animals were analysed 4–6 weeks after the last injection unless indicated otherwise. All animal experiments were done in accordance to the guidelines of the New York University School of Medicine Institutional Animal Care and Use Committee.

Antibodies and fluorescence-activated cell sorting analysis. Antibody staining and fluorescence-activated cell sorting (FACS) were performed as previously described²². All antibodies were purchased from BD-Pharmingen or e-Bioscience. We used the following antibodies: c-kit (2B8), Sca-1 (D7), Mac-1 (M1/70), Gr-1 (RB6-8C5), NK1.1 (PK136), TER-119, CD3 (145-2C11), CD19 (1D3), IL7R α (A7R34), CD34 (RAM34), Fc γ RII/III (2.4G2), Flk-2/Flt-3 (A2F10.1), CD4 (RM4-5), CD4 (H129.19), CD8 (53-6.7), CD45.1 (A20), CD45.2 (104), CD150 (9D1), CD48 (HM481). Bone marrow lineage antibody cocktail included Mac-1, Gr-1, NK1.1, TER-119, CD3 and CD19.

OP9-DL1/DL4 *in vitro* co-culture. OP9-DL1/DL4 cells were maintained in MEM with 20% fetal bovine serum. Ten thousand purified infected (green fluorescent protein (GFP⁺)) LSK or cKit⁺ bone marrow progenitors were seeded into a six-well plate with confluent OP9 cells in the presence of 10 ng ml⁻¹ SCF, 10 ng ml⁻¹ interleukin-3 (IL-3), 5 ng ml⁻¹ Flt3-L, 10 ng ml⁻¹ IL6 and 5 ng ml⁻¹ IL-7. Flow cytometric analysis was performed on an LSRII (BDIS). Haematopoietic cells were gated using CD45 cell-surface expression.

Microarray analysis. LSK or GMP cells from individual mice were used. To generate sufficient sample quantities for oligonucleotide gene chip hybridization, we used the Ovation RNA Amplification System V2 (Nugen) for antisense RNA (cRNA) amplification and labelling. The amplified cRNA was labelled and hybridized to the Mouse 430.2 microarrays (Affymetrix). The data were normalized using the previously published robust multi-array average algorithm using the GeneSpring GX software (Agilent).

Full Methods and any associated references are available in the online version of the paper at www.nature.com/nature.

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Supplementary Information is linked to the online version of the paper at www.nature.com/nature.

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Author Contributions I.A., C.L. and A.K. conceived the study and designed all experiments. A.K. and A.E. helped with experimental planning and generated the Ncstn^{fl/fl} mice. C.L. performed most of the mouse experiments and *in vitro* studies aided by P.O., S.B., S.C., T.T. and E.A. R.L.L., O.A.-W. and M.B. performed and analysed human leukaemia sample exon sequencing. H.H. and F.M. helped with disease modelling and computational analysis of disease progression. I.v.D.W. and T.T. performed the human stem cell differentiation assays. C.L. and S.I. analysed mouse disease pathology. J.Z. processed and analysed gene expression data.

Author Information The microarray data are deposited in Gene Expression Omnibus of the National Center for Biotechnical Information under accession numbers GSE27794, GSE27799 and GSE27811. Reprints and permissions information is available at www.nature.com/reprints. The authors declare no competing financial interests. Readers are welcome to comment on the online version of this article at www.nature.com/nature. Correspondence and requests for materials should be addressed to I.A. (iannis.aifantis@nyumc.org) or A.K. (aklinakis@bioacademy.gr).

METHODS

Animals. Genotyping of $N1^{flf}N2^{flf}N3^{-/-}$ mice^{23–26} was performed as previously reported. $Ncstn^{flf}Mx1-Cre^{+}$ and $N1^{flf}N2^{flf}N3^{-/-}Mx1-Cre^{+}$ animals were injected with 20 μ g polyI:polyC per gram of body weight for a total of six injections. The injections were initiated 14 days after birth and done every 2 days. Animals were analysed 4–6 weeks after the last injection unless indicated otherwise. All animal experiments were done in accordance to the guidelines of the New York University School of Medicine Institutional Animal Care and Use Committee.

Generation of the $Ncstn^{flf}$ mice. To generate a conditional $Ncstn$ allele we used standard ES cell targeting approaches. Exons 5–7 of the $Ncstn$ locus were targeted for Cre-dependent recombination. Removal of these exons generates an in-frame stop codon that prematurely terminates translation. As the homology region we used a 7 kb BsrGI–EcoRI genomic fragment spanning introns 2–11. A PGK–NEO cassette and a loxP site were cloned into an EcoRV site located in intron 4, whereas the second loxP site was subcloned into a unique PmlI site of intron 7. Embryonic stem cell colonies (129/sj) were screened by Southern blot of genomic DNA digested with BamHI (Supplementary Fig. 1b). Correctly targeted embryonic stem cells were injected in C57BL6 blastocysts and chimaeras were generated. After verification of germline transmission, the PGK–NEO cassette was removed using a germline Flp mouse²⁷. $Ncstn^{flf}Flp^{+}$ mice were crossed to the Mx-cre strain. Genotyping primers and probes are available upon request.

Antibodies and FACS analysis. Antibody staining and FACS analysis was performed as previously described²². All antibodies were purchased from BD-Pharmingen or e-Bioscience. We used the following antibodies: c-kit (2B8), Sca-1 (D7), Mac-1 (M1/70), Gr-1 (RB6-8C5), NK1.1 (PK136), TER-119, CD3 (145-2C11), CD19 (1D3), IL-7 α (A7R34), CD34 (RAM34), Fc γ RII/III (2.4G2), Flk-2/Flt-3 (A2F10.1), CD4 (RM4-5), CD4 (H129.19), CD8 (53-6.7), CD45.1 (A20), CD45.2 (104), CD150 (9D1), CD48 (HM481). Bone marrow lineage antibody cocktail included Mac-1, Gr-1, NK1.1, TER-119, CD3, CD19. For western blotting, goat polyclonal anti-Nicastrin antibody (N19, sc-14369, Santa Cruz) and mouse monoclonal anti-Actin antibody (Clone C4, MAB1501R, Millipore) were used. Annexin V/7AAD staining was done using an annexin V PE detection kit (BD Pharmingen, 559763) following the manufacturer's protocol.

PCR with reverse transcription. Total RNA was isolated using the RNeasy Plus Mini Kit (Qiagen) and cDNA was synthesized using the SuperScript First-Strand Kit (Invitrogen). Quantitative PCR was performed using SYBR green iMaster and a LightCycler 480 (Roche) with the following primer sequences (melting temperature = 60 °C for all primers): *Cebpa* forward TTACAACAGGCCAGGTTCCT, *Cebpa* reverse CTCTGGGATGGATCGATTGT; *Pu.1* forward ATGGAAGGGTTTTCCCTCACCGCC, *Pu.1* reverse GTCCACGCTCTGCAGCTCTGTGAA; *Gata2* forward AACGCTGTGGCCCTACTA, *Gata2* reverse TCTCTTGCACTGCACTTGGAG; *Cebpd* forward ATCGCTGCAGCTTCTATGT, *Cebpd* reverse AGTCATGCTTTCCCGTGTTC; *Hes1* forward TCCAAGCTAGAGAAGGCAGAC, *Hes1* reverse TGATCTGGGTCATGCAGTTG; *Gata1* forward ACTGTGGAGCAACGGCTACT, *Gata1* reverse TCCGCCAGAGTGTTGTAGTG; *Ncstn* forward CTGGCGGTGCAGCTGTATGAG, *Ncstn* reverse GGAGACGGCGATGTAGTGTGAAG.

Bone marrow transplantation assays. Five hundred thousand bone marrow cells ($Ly5.2^{+}$) were transplanted by retro-orbital intravenous injections into lethally irradiated (960 cGy) BL6SJL ($Ly5.1^{+}$) recipient mice. Four weeks after transplantation, mice were injected with 20 μ g polyI:polyC per gram of body weight for a total of six injections. Peripheral blood of recipient mice was collected at 4, 7, 9 and 12 weeks after transplantation. Recipient mice were killed 16 weeks after transplantation for analysis.

Microarray analysis. LSK or GMP cells from individual mice were used. Microarray analysis was performed as previously described²⁸. Briefly, freshly isolated cells were sorted by surface marker expression, and total RNA was extracted using the RNeasy kit (Qiagen). To generate sufficient sample quantities for oligonucleotide gene chip hybridization, we used the Ovation RNA Amplification System V2 (Nugen) for cRNA amplification and labelling. The amplified cRNA was labelled and hybridized to the Mouse 430.2 microarrays (Affymetrix). The Affymetrix gene expression profiling data were normalized using the previously published robust multi-array average algorithm using the GeneSpring GX software (Agilent). The gene-expression intensity presentations were generated with Matrix2png software (<http://chibi.ubc.ca/matrix2png/bin/matrix2png.cgi>) or Multi Experiment Viewer software (<http://www.tm4.org/mev/>).

Gene set enrichment analysis. Gene set enrichment analysis was performed using Gene Set Enrichment Analysis software^{29,30} (<http://www.broadinstitute.org/gsea/>) using gene set as permutation type, 1,000 permutations and \log_2 ratio of classes as metric for ranking genes.

The 'myeloid signature' gene set was generated using a systematic approach based on the comparison of gene expression arrays from WT LSK and WT GMP. Genes that were significantly upregulated in GMP compared with LSK

(over 1.5-fold induction $P < 0.05$) were used to define myeloid signature genes. The list was trimmed of genes of unknown function and those related to metabolism.

Other myeloid-specific gene sets used in the analysis were taken from gene sets already present in the MSig database of the Broad Institute.

Retroviral infection of Lineage^{neg}cKit⁺ bone marrow cells or LSK cells. Bone marrow cells were enriched for cKit-positive cells using either the EasySep kit (StemCell Technology) or the Dynabead kit (DYNAL, Invitrogen). LSK cells were flow sorted using lineage markers Mac-1, Gr-1, NK1.1, TER-119, CD3, CD19, Sca1 and cKit. Cells were subsequently cultured in OPTI-MEM supplemented with 10% fetal bovine serum, 100 μ g ml⁻¹ penicillin, 100 μ g ml⁻¹ streptomycin, 50 ng ml⁻¹ SCF and Flt3l, and 10 ng ml⁻¹ IL-6 and IL-7. For retroviral production, Plat-E cells were transfected with appropriate retroviral expression constructs by the calcium phosphate method. Virus supernatant was collected 48 h after transfection and used directly for spin infection of cKit positive-enriched bone marrow cells or sorted LSK cells at 660g for 90 min. Forty-eight hours after infection, lineage-negative GFP-positive cells were sorted for RT-PCR analysis, methylcellulose plating assays or OP9 co-culture assay.

In vitro differentiation assays. Total bone marrow (15,000), sorted LSK (500) and GMP (500) cells were plated in triplicates into cytokine-supplemented methylcellulose medium (MethoCult 3434, Stem Cell Technologies). Colony type was scored after 10 days of culture. Replating was performed after 8 days of culture. For experiments involving Hes1 overexpression, 10,000 GFP⁺ infected cells were plated in duplicate on 35 mm cytokine-supplemented methylcellulose medium (MethoCult 3434, Stem Cell Technologies). Cells were recovered 8 days later, stained and analysed by FACS as described.

Human progenitor/OP9 co-cultures. Bone marrow samples were obtained and used according to the guidelines of the Medical Ethical Commission of Ghent University Hospital (Belgium). Bone marrow mononuclear cells were isolated through Lymphoprep density-gradient centrifugation and CD45⁺ glycoprotein-A positive (Gly-A⁺) cells were enriched by micro-magnetic beads (Miltenyi Biotec). Subsequently, CD45⁺Gly-A⁺ cells were labelled with CD34-FITC, CD38-PE, CD19-APC and CD14-APC to sort CD34⁺CD38⁻Lin⁻ cells on a FACSaria to a purity greater than 98%. OP9-control and OP9 cells expressing human ligands Dll1, Dll4, Jag1 and Jag2 were generated by retroviral transduction of OP9 cells (provided by J. C. Zúñiga-Pflücker), followed by sorting for eGFP⁺ cells, as described previously³¹. Purified bone-marrow progenitors ($2.0 \times 10^3 - 2.6 \times 10^3$) were seeded into a 24-well plate with confluent OP9 cells in the presence of 20 ng ml⁻¹ SCF, 20 ng ml⁻¹ Flt-3L, 20 ng ml⁻¹ TPO, 10 ng ml⁻¹ GM-CSF and 10 ng ml⁻¹ G-CSF. Co-cultures were performed and prepared for analysis as described previously³². Flow cytometric analysis was performed on an LSRII (BDIS). Human cells were gated using a CD45 monoclonal antibody and dead cells were excluded using propidium iodide.

OP9-DL1/DL4 in vitro co-culture. OP9 cells were maintained in MEM with 20% fetal bovine serum. Ten thousand purified infected (GFP⁺) LSK or cKit⁺ bone marrow progenitors were seeded into a six-well plate with confluent OP9 cells in the presence of 10 ng ml⁻¹ SCF, 10 ng ml⁻¹ IL-3, 10 ng ml⁻¹ IL-6, 5 ng ml⁻¹ Flt3-L and 5 ng ml⁻¹ IL-7. Flow cytometric analysis was performed on an LSRII (BDIS). Haematopoietic cells were gated using CD45 cell-surface expression and GFP expression.

Chromatin immunoprecipitation assays. Chromatin immunoprecipitation assays were performed with 80 μ g of genomic DNA from 32D-MIG- or 32D-HES1-infected cells following standard procedures. Briefly, chromatin was cross-linked with 1% formaldehyde, and sheared by sonication. Immunoprecipitation was performed with immunoglobulin-G or anti-HES1, followed by incubation with protein A magnetic beads/salmon sperm DNA (Invitrogen Dynal). DNA isolated from antibody bound fraction was eluted after washing, extracted with phenol/chloroform and precipitated with ethanol. Real-time PCR quantification of immunoprecipitated DNA was performed with the SYBR Green PCR Master Mix (Roche) and primers designed to amplify regions covering each putative HES1 binding site in the *Pu.1* and *Cebpa* promoters: *Pu.1-1* forward ACGTTCAAGGGTTGGAGAAA, *Pu.1-1* reverse GCCAATTAGGGCCAACAGTA; *Pu.1-2* forward GACCAAAGTCCCTCCACCTGA, *Pu.1-2* reverse CTGGGAGGGAGAAAGGCTA; *Cebpa-1* forward CCAAAGCAGTCTCCAACCTC, *Cebpa-1* reverse CCACTCCAGCCAACACTA; *Cebpa-2* forward CGCCTAACCACGGACCAC, *Cebpa-2* reverse AGTAGGATGGTGCCTGCTG.

Histological analyses. Mice were killed and autopsied, then dissected tissue samples or tumours were fixed for 24 h in 10% buffered formalin, dehydrated and embedded in paraffin. Paraffin blocks were sectioned at 5 μ m and stained with haematoxylin and eosin.

Luciferase reporter assays. pGL3 promoter reporter plasmid (Promega) was used to clone *Pu.1*, *Cebpa* or *Hes1* promoter sequences. pGL3 promoter empty vector or the vector containing *Pu.1*, *Cebpa* or *Hes1* promoter sequences, HA-HES1 expression plasmid or pCDNA3.1 control plasmid and *Renilla* expression vector

were co-transfected into HEK293T cells using regular calcium phosphate transfection protocol. Luciferase activities were examined 24 h after transfection using the Dual Luciferase reporter assay system (Promega) following the manufacturer's instructions, and normalized to *Renilla* activity.

Statistical analysis. The means of each data set were analysed using Student's *t*-test, with a two-tailed distribution and assuming equal sample variance.

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