Somatic activating mutations in Notch1 contribute to the pathogenesis of T cell acute lymphoblastic lymphoma (T-ALL), but how activated Notch1 signaling exerts this oncogenic effect is not completely understood. Here we identify HIV-1 Rev–binding protein (Hrb), a component of the clathrin-mediated endocytosis machinery, as a critical mediator of Notch-induced T-ALL development in mice. Hrb was found to be a direct transcriptional target of Notch1, and Hrb loss reduced the incidence or delayed the onset of T-ALL in mouse models in which activated Notch1 signaling either contributes to or drives leukemogenesis. Consistent with this observation, Hrb supported survival and proliferation of hematopoietic and T cell precursor cells in vitro. We demonstrated that Hrb accelerated the uptake of transferrin, which was required for upregulation of the T cell protooncogene p21. Indeed, iron-deficient mice developed Notch1-induced T-ALL substantially more slowly than control mice, further supporting a critical role for iron uptake during leukemogenesis. Taken together, these results reveal that Hrb is a critical Notch target gene that mediates lymphoblast transformation and disease progression via its ability to satisfy the enhanced demands of transformed lymphoblasts for iron. Further, our data suggest that Hrb may be targeted to improve current treatment or design novel therapies for human T-ALL patients.

Introduction
T cell acute lymphoblastic lymphoma (T-ALL) are serious hematologic malignancies of children and young adults. Current treatments that include intensive chemotherapy and cranial radiation are unsatisfactory, as they frequently cause severe long-term toxicities. Furthermore, significant numbers of patients die from recurrent disease, in spite of therapy. Better understanding of the molecular basis of lymphogenesis will likely lead to improved therapy. The Notch receptor is implicated in the pathogenesis of T-ALL (1–3). Recent studies have demonstrated that Notch1 is activated by somatic mutations in approximately 60% of cases of pediatric T-ALL (4). Notch1 is a cell surface receptor that is activated by ligands from the DSL family. Ligand binding induces proteolytic cleavage of Notch1 (S2), which is immediately followed by further cleavage by gamma-secretase (S3). This cleavage results in the release of the soluble Notch1 intracellular domain (ICN1), which translocates to the nucleus, where it activates transcription of target genes via its interaction with the DNA-binding protein CSL.

How Notch transforms T cell precursors remains a subject of intense study. Activated Notch has multiple pleiotropic effects in T cell precursors, which include dramatic acceleration of proliferation, increased thymocyte survival, and a block in differentiation (5). Preliminary studies on Notch inhibition by gamma-secretase inhibitors (GSIs) have demonstrated the importance of this signaling pathway in T-ALL. However, systemic toxicity limits the use of these drugs, and current efforts by many investigators focus on the downstream molecular sequelae of Notch activation with the hope that they may provide useful therapeutic targets.

In previous studies, we found that mice bearing a conditional knockout allele of Creb-binding protein (CBP) developed fatal T cell lymphomas with high penetrance. The transformed cells resembled immature thymocytes and had a CD4+CD8+ double-positive (DP) phenotype (6). Microarray analyses revealed upregulation of multiple transcripts, including components of the Notch signaling pathway, c-myc, and the HIV-1 Rev–binding protein (Hrb). Hrb is an essential cofactor for HIV replication (7–9) and serves as an AP-2/EPS15 adapter required for vesicle fusion events that create the acrosome during spermatogenesis (10). Additionally, Hrb is implicated in clathrin-dependent endocytosis of VAMP7 (11) and the transferrin receptor (TfR) (12). However, it has not been implicated in oncogenesis. Here, we identify what we believe is a novel role for Hrb in T cell lymphomagenesis and find that it contributes significantly through its effects on transferrin uptake. In addition, we demonstrate what we believe is a novel function of Hrb in regulating protein levels of the T cell prosurvival factor p21 (WAF1/CIP1).

Results
Notch activation cooperates with CBP loss to accelerate T cell lymphomagenesis. We previously observed that transcriptional targets of Notch1 were upregulated in the majority of CBP-null T cell lymphomas (6), raising the possibility that activated Notch might have been a contributing factor to their development. If Notch indeed cooperates with CBP loss, then activation of Notch by use of a transgene might accelerate T cell lymphomagenesis in CBP-null mice. Thus, we generated mice that had both a conditional deletion of CBP and
that expressed the intracellular activated form of Notch1 (ICN1) (13). ICN1 transgenic mice developed T cell lymphomas around 98 weeks (data not shown). Mice with the ICN1 transgene combined with CBP loss developed T cell lymphomas much faster than littermate control animals that were singly CBP-null or ICN1-transgenic (P < 0.0001 by Mantel-Cox log-rank analysis) (Figure 1A), consistent with the notion that activated Notch could synergize with loss of CBP to generate T cell lymphoma.

**Hrb is a direct transcriptional target of Notch1.** Hrb mRNA levels were significantly elevated in T cell lymphomas from CBP-null mice (6). To verify whether this was also reflected at the protein level, we prepared lysates from 6 independent spontaneous T cell lymphomas from CBP-null mice and analyzed Hrb protein by Western blotting. Five out of 6 tumors expressed dramatically higher levels of Hrb protein compared with nontransformed thymocytes (Figure 1B).

In an independent study, Weng et al. investigated changes in gene expression in T-ALL cells following Notch1 inhibition. Among their results was the finding that Hrb transcript levels were reduced following Notch inactivation (14). In addition, microarray analyses of several murine T-ALL cell lines have consistently shown Hrb to be regulated by Notch1 (W.S. Pear, unpublished observations). To address whether Hrb is directly regulated by Notch1 signaling, we utilized the Notch-dependent murine T-ALL cell line T6E12 (15). Transfection of cells with a dominant negative MAML1 construct (DNMAML1) or treatment with GSI decreased Hrb mRNA and protein levels (Figure 2B), accompanied by progressive cell growth arrest/death. Expression of a constitutively active, GSI-insensitive form of Notch1, ICN1, restored Hrb mRNA levels even in the presence of GSI. Treatment of T-ALL cells with GSI resulted in the accumulation of a transmembrane-tethered form of Notch that can be rapidly cleaved to the intracellular activated form (ICN1) upon GSI removal. GSI washout resulted in the derepression of Hrb mRNA, even in the presence of the protein synthesis inhibitor cycloheximide, suggesting that Hrb could be a direct transcriptional target of Notch1 (Figure 2B).

**Hrb is crucial for CBP-null lymphomagenesis.** To explore whether Hrb might participate in oncogenesis, we generated cohorts of Hrb+/+ and Hrb−/− mice in addition to conditional deletion of CBP in the thymus. These mice were followed for up to 90 weeks. We observed a 70% reduction in the incidence of T cell lymphomas in the Hrb−/− mice in comparison with Hrb+/− littermate controls (15% vs. 50%, P = 0.0396 by Mantel-Cox log rank analysis) (Figure 3A), indicating that Hrb plays a critically important role in efficient tumor formation due to loss of CBP in thymocytes.

Hrb expression was highly induced above basal levels in all lymphomas that had Notch1 upregulation (Figure 3B). However, Hrb could also be induced by alternative pathways, since high-level Hrb was detected in some tumors lacking Notch1 (Figure 3B). Additionally, the upregulation of Hrb in CBP-null tumors correlated with Notch activation as demonstrated with an antibody that detects the gamma-secretase cleaved form of Notch1 (Figure 3C).

**Hrb is not required for physiologic Notch signaling.** Although the initial site of transformation in T-ALL is not known, we postulated that either a committed lymphoid precursor in the BM or an early-stage thymocyte would be a likely candidate target cell, because of the phenotype of most lymphoma cells that arise in CBP-null or Notch-transgenic mice (CD4-positive, CD8-negative). If loss of the Hrb gene would cause a defect in thymocyte development, this would be a relatively trivial mechanism underlyling its protective effect in preventing T-ALL in the absence of CBP (Figure 3A). To test this possibility, we examined thymi from Hrb−/−, Hrb+/−, and Hrb+/+ mice. These studies revealed an equal number of total thy-
Figure 2
Hrb is a direct Notch1-regulated transcriptional target. (A) A murine Notch-dependent cell line, T6E12, was utilized for Hrb mRNA and protein analysis. T6E12 cells were transduced with MigR1 (empty vector), MigR1-ICN1, or MigR1-DNMAML1 for 48 hours followed by 24 hours DMSO or GSI (1 μM) treatment as labeled. Hrb mRNA and protein levels were determined by quantitative RT-PCR and Western blot analysis, respectively. Results were obtained from 3 independent experiments. (B) T6E12 cells were treated with GSI (1 μM) for 48 hours to permit accumulation of the gamma-secretase substrate (transmembrane-Notch1 [N^TM]). Cells were then washed and replenished with medium containing GSI (mock washout) or medium lacking GSI (washout) in the absence or presence of 20 μM cycloheximide (CHX). Hrb mRNA levels were evaluated by quantitative RT-PCR after 4 hours of additional incubation. Similar results were obtained in 3 independent experiments. (C) ChIP was performed on cross-linked fragmented DNAs prepared from T6E12 cells treated with DMSO or 1 μM GSI for 24 hours. Eluted DNAs were then analyzed by quantitative RT-PCR using primers flanking putative CSL-binding sites (A and B). Amplification of hes1 CSL-binding site served as a positive control to validate ChIP efficiency. The amount of DNA amplified from immunoprecipitated DNAs was normalized to that amplified from input DNA. (D) Western blot of lysates prepared from empty vector– (MigR1) and ICN1-transduced Hrb^+/+ and Hrb^-/- T cell precursors blotted for Hrb (clone H-300) and actin (loading control). Asterisk indicates nonspecific band. Data are shown as mean ± SD.
Hrb is required for efficient CBP-null lymphomagenesis. (A) Kaplan-Meier survival curve of Hrb+/+ and Hrb−/− cohorts monitored over a period of 90 weeks. Hrb−/− mice on the background of MMTV-Cre/CBPlox/lox were intercrossed to generate Hrb+/+ and Hrb−/− mice with conditional loss of CBP. Mice were sacrificed once moribund and confirmed via dissection for the occurrence of T cell lymphoma (n = 19 for each cohort; P = 0.0396, log-rank Mantel-Cox test). (B) Western blot of lysates prepared from Hrb+/+ CBP-null tumors (n = 7) and Hrb−/− CBP-null tumors (n = 2). Membrane was blotted for Notch1 protein (clone mN1a), Hrb (clone H-300), and GAPDH (loading control). Asterisks indicate nonspecific bands.

Hrb is required for growth and survival of ICN1-transduced hematopoietic and T cell precursors. To investigate the mechanism of Hrb in oncogenic Notch signaling, we cultured hematopoietic precursors from Hrb+/+ and Hrb−/− BM on OP9 stromal cells, transduced with ICN1, and then monitored cells for growth over a period of 7 days. ICN1-transduced BM cells that were Hrb−/− had reduced growth and increased death as measured by annexin V/propidium iodide (PI) staining, compared with Hrb+/+ cells (Figure 5, A and B). Retroviral mediated reexpression of Hrb rescued both defects. Next, we isolated CD4+CD8− DN thymocytes from Hrb+/+ and Hrb−/− mice and plated these T cell precursors onto OP9DL1 stromal cells. Upon transduction with ICN1, loss of Hrb was associated with a marked reduction in cell numbers in all 3 thymocyte subsets (DN, CD8 immature SP [ISP], and DP) that developed after 5 days in these cocultures, in comparison with wild-type cells (Figure 5C). We verified that Hrb protein levels were significantly upregulated in both hematopoietic precursors and T cell precursors (Figure 5D) transduced with ICN1.

Hrb loss increases surface CD71 and is required for efficient transferrin uptake. Hrb is important for the internalization of cell surface VAMP7 and delivery to the lysosomal membrane (11) and efficient clathrin-dependent endocytosis (12). The TR (CD71) is a well-known substrate for clathrin-dependent endocytosis (19–22), and its endocytosis and trafficking are critically important for its essential role in cellular uptake of iron in the form of holo-transferrin. Intriguingly, transferrin and TR have both been implicated in thymocyte development (23–25) and TR has been found to be elevated on immature cycling thymocytes (26) and leukemias (27, 28). Therefore, we asked whether Hrb might participate in regulation of CD71 in the context of oncogenic Notch signaling. DN thymocytes cultured on OP9DL1 cells were transduced with control or ICN1-expressing retrovirus and cultured for 5 days. CD71 at the cell surface detected by flow cytometry after antibody staining revealed several important findings. First, DN cells had higher levels of CD71 than subsequent developmental

mocytes (Supplemental Figure 1A), and also completely normal CD4+CD8− DN (Supplemental Figure 1B), CD4+CD8− DP, CD4− single-positive (SP), and CD8− SP subsets (Supplemental Figure 1C). In addition, analysis of the TCR-β repertoire by an RT-PCR method (17) revealed results virtually identical to those from Hrb+/+ and Hrb−/− splenocytes. We conclude that Hrb is not required for normal thymocyte development at various stages. Finally, Hrb was not required for expansion of stimulated peripheral T cells (Supplemental Figure 4).

Hrb is required for efficient ICN1-induced leukemogenesis. Since Hrb is a direct transcriptional target of Notch and Notch activation cooperates with CBP loss to accelerate lymphomagenesis, we asked whether Hrb participates in Notch-induced tumorigenesis using a well-characterized BM transduction and transplantation model (18). Hrb+/+ and littermate control mice were treated with 5-FU (Plate 20) to enrich for hematopoietic stem cells, which were transduced with control virus (MigR1) or ICN1-expressing retrovirus. Equal transduction efficiencies were verified by flow cytometric detection of GFP (Figure 4A) and semiquantitative RT-PCR of ICN1 mRNA (data not shown). Transduced BM cells were transferred to lethally irradiated wild-type recipients, which were then observed for onset of T-ALL over the next several months. These experiments revealed a clear requirement for Hrb in efficient ICN1-induced leukemogenesis (Figure 4B). The median survival of mice receiving Hrb+/+ ICN1–transduced BM was 27.5 days after BM transplant (BMT). Median survival of mice receiving Hrb−/− ICN1–transduced BM was approximately 2-fold longer at 54.0 days after transplant (P < 0.0001 by Mantel-Cox log rank analysis). Mice that received control retrovirus-transduced BM cells of either WT or Hrb−/− genotype did not develop leukemia and had normal engraftment and survival. Analysis of the peripheral blood from transplant recipients 15 days after BMT revealed an accumulation of GFP+ DP cells in both Hrb+/+ ICN1 and Hrb−/− ICN1 recipients (Figure 4C). Additionally, splenocyte (Figure 4D) and BM (Figure 4E) analyses performed 26 days after BMT also showed an equal accumulation of GFP+ DP cells in both Hrb+/+ and Hrb−/− ICN1–transduced BM recipients. Thus, we do not suspect that the significant delay in development of T-ALL in Hrb−/− cells was a result of failed engraftment. BM isolated from Hrb−/− ICN1 transplants did show reduced numbers of GFP+ cells and an increased percentage of GFP+CD8− SP cells (Supplemental Figure 5).

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Hrb is required for efficient ICN1-induced leukemogenesis. (A) Analysis of GFP expression in Hrb<sup>+/+</sup> and Hrb<sup>−/−</sup> BM after transduction with ICN1-expressing retrovirus and before transplantation into lethally irradiated recipients. Numbers in each GFP histogram refer to the GFP-positive percentage. (B) Kaplan-Meier survival curve of mice receiving Hrb<sup>−/−</sup> ICN1–transduced BM (n = 10), Hrb<sup>−/−</sup> ICN1–transduced BM (n = 13), Hrb<sup>−/−</sup> MigR1–transduced BM (n = 4), and Hrb<sup>−/−</sup> MigR1–transduced BM (n = 4). Statistical significance was established using the Mantel-Cox log rank analysis and showed a significant increase in survival in mice receiving Hrb<sup>−/−</sup> ICN1–transduced BM (P < 0.0001). Median time of survival after BMT for mice receiving Hrb<sup>−/−</sup> ICN1–transduced BM was 27.5 days. Median time of survival after BMT for mice receiving Hrb<sup>−/−</sup> ICN1–transduced BM was 54 days. Mice receiving MigR1-transduced BM did not succumb to disease. (C) GFP analysis and CD4<sup>+</sup>CD8<sup>+</sup> surface staining of peripheral blood cells isolated 15 days after BMT from mice receiving Hrb<sup>−/−</sup> and Hrb<sup>−/−</sup> MigR1– and ICN1–transduced BM cells. (D–E) GFP analysis and CD4<sup>+</sup>CD8<sup>+</sup> surface staining of splenocytes (D) and BM cells (E) isolated from mice receiving Hrb<sup>−/−</sup> and Hrb<sup>−/−</sup> ICN1–transduced BM cells. Numbers in GFP histograms refer to the percentage of GFP-positive cells. Numbers in each dot plot refer to the percentages in each quadrant.

Stages, consistent with their high rates of proliferation (29). Furthermore, ICN1 expression induced higher levels of CD71 at each developmental stage (Figure 6A). The most striking finding was that Hrb<sup>−/−</sup> cells, either with or without ICN1, had higher surface CD71 than Hrb<sup>+/+</sup> cells (Figure 6A). Hrb<sup>−/−</sup> BM hematopoietic precursors transduced with ICN1 also expressed significantly higher surface CD71 compared with WT controls (Figure 6B). The high levels of surface CD71 in knockout cells was reduced to WT levels with reexpression of Hrb.

To test the function of CD71, cells were incubated with fluorescently labeled transferrin for various periods of time, washed, and then analyzed by flow cytometry to detect uptake. As expected from the increased surface CD71, expression of ICN1 caused a more rapid and greater final uptake of transferrin both in WT and in Hrb<sup>−/−</sup> cells compared with MigR1-transduced cells (Figure 6, C and D). Loss of Hrb in ICN1-expressing DP cells caused a relative reduction in the uptake of transferrin, in spite of their slightly higher CD71 levels. In the case of ICN1-transduced DN and CD8 ISP cells, transferrin uptake was equal between Hrb<sup>−/−</sup> and Hrb<sup>+/+</sup> cells (Supplemental Figure 2, A and B), even though there were significantly higher CD71 levels on the Hrb<sup>−/−</sup> cells. Together, these studies suggest that transferrin uptake is less efficient in cells lacking Hrb if they also express ICN1.

Efficient transferrin uptake and subsequent utilization of iron are essential for rapidly growing cells in order to support cell division. We wondered, therefore, whether the defect in proliferation and survival seen in ICN1-expressing Hrb<sup>−/−</sup> cells could, in part, be explained by insufficient availability of transferrin. Although the mutant cells had inefficient uptake, we found that increasing the extracellular transferrin did lead to increased uptake (Figure 6D). To bypass the Hrb<sup>−/−</sup> defect, we cultured hematopoietic precursors transduced with ICN1 in the presence of excess transferrin. Under these conditions, mutant cells recovered their ability to proliferate in vitro, while the addition of excess transferrin had little effect on WT cells (Figure 6E). This was accompanied by a significant reduction in cell surface CD71 as well (Figure 6F). To further confirm that iron supplementation can rescue the defect in Hrb<sup>−/−</sup> ICN1–transduced cells, we incubated DN T cell precursors with ferric ammonium citrate (FAC), a compound that stimulates a transferrin-independent uptake of iron (30). Similar to the results of holo-transferrin addition, FAC addition rescued the Hrb<sup>−/−</sup> defect in proliferation, but had little effect on WT cells (Supplemental Figure 3). These results strongly suggested that inefficient iron utilization was a major factor limiting the proliferation and survival of ICN1-expressing Hrb<sup>−/−</sup> cells.

Consistent with the increased surface expression of CD71 following ICN1 transduction, total cellular levels of CD71 were also induced compared with MigR1 controls (Figure 6G). Poor transferrin uptake and/or trafficking in the absence of Hrb might explain why CD71 levels were higher on mutant cells, since translation of the TfR mRNA is increased when intracellular levels of iron are low. Western blotting indicated that total cellular levels of CD71 were indeed elevated in Hrb<sup>−/−</sup> MigR1– and ICN1–transduced cells (Figure 6G). As predicted, total cellular CD71 in Hrb<sup>−/−</sup> ICN1–transduced cells decreased in the presence of excess holo-transferrin.
Moreover, there is evidence that p21 can inhibit apoptosis and mitosis cycle progression and survival. One such component that is dependent upon iron is the p21 (CIP1/WAF1) protein, a cdk inhibitor that has been shown in various systems to halt cell cycle progression (33–37). However, there is also evidence to suggest that p21 is required for leukemia cell proliferation and survival. To determine whether the observed interplay among Notch, Hrb, and iron metabolism may function in human T-ALL as well, we utilized the Notch-dependent T-ALL cell lines DND41, T-ALL1, KOPT1, and SUPT1. GSI treatment of DND41 and T-ALL1 cells decreased the transcript levels of Hrb (Figure 8A), as was observed with the mouse T6E cell line (Figure 2A). Knockdown of Hrb resulted in a decreased accumulation of human T-ALL cells as well as decreased survival, as assessed by annexin V/PI staining (Figure 8, B–D). Our data suggest that Hrb plays a role in the efficient utilization of iron via transferrin. Thus, inefficient use of iron in Hrb-depleted cells should confer greater sensitivity to the iron chelator desferrioxamine (DFO). Indeed, reducing Hrb levels in DND41 cells resulted in greater sensitivity to iron chelation (Figure 8E). Consistent with the increased surface levels of CD71 in DN/Hrb−/− cells, treatment with DFO at 2.0 μM in KOPT1 cells resulted in significantly greater surface CD71 (Figure 8F).

Hrb regulates p21 protein levels via transferrin uptake. Iron is important in several cellular processes (31, 32) and particularly in cell cycle progression and survival. One such component that is dependent upon iron is the p21 (CIP1/WAF1) protein, a cdk inhibitor that has been shown in various systems to halt cell cycle progression (33–37). However, there is also evidence to suggest that p21 is required for leukemia cell proliferation and survival. To determine whether the observed interplay among Notch, Hrb, and iron metabolism may function in human T-ALL as well, we utilized the Notch-dependent T-ALL cell lines DND41, T-ALL1, KOPT1, and SUPT1. GSI treatment of DND41 and T-ALL1 cells decreased the transcript levels of Hrb (Figure 8A), as was observed with the mouse T6E cell line (Figure 2A). Knockdown of Hrb resulted in a decreased accumulation of human T-ALL cells as well as decreased survival, as assessed by annexin V/PI staining (Figure 8, B–D). Our data suggest that Hrb plays a role in the efficient utilization of iron via transferrin. Thus, inefficient use of iron in Hrb-depleted cells should confer greater sensitivity to the iron chelator desferrioxamine (DFO). Indeed, reducing Hrb levels in DND41 cells resulted in greater sensitivity to iron chelation (Figure 8E). Consistent with the increased surface levels of CD71 in DN/Hrb−/− cells, treatment with DFO at 2.0 μM in KOPT1 cells resulted in significantly greater surface CD71 (Figure 8F).

Iron is critical for T cell precursor development and for ICN1-induced leukemogenesis. To determine whether iron itself is required for thymocyte development, we depleted thymocyte-OP9DL1 cocultures of iron by addition of DFO. This caused a dramatic loss of thymocyte numbers (Figure 9A). Interestingly, Hrb−/− DN cells were more sensitive to depletion of iron with DFO, consistent with their reduced ability to utilize iron-loaded transferrin.

Together, the results above indicate a critical role for Hrb in mediating in vitro survival of ICN1-transduced T cell precursors and human T-ALL cell lines, by driving efficient utilization of iron. Reduced iron uptake might similarly explain the decreased incidence (Figure 3A) or delay in development (Figure 4B) of T-ALL in mice lacking Hrb. If so, suppression of iron uptake in WT mice might delay their development of T-ALL. We reasoned that placing...
recipient mice on a low-iron diet would reduce their total body iron stores and decrease availability of the iron to developing leukemic blasts. Cohorts of WT mice transplanted with WT ICN1–transduced BM cells were placed on either a low-iron diet or normal mouse chow for up to 48 days. Mice that were transplanted with control (MigR1-transduced) BM cells on the low-iron diet engrafted, survived, and appeared well, although they did develop iron-deficiency anemia (hemoglobin 7.8 g/dl and mean corpuscular volume [MCV] 43.9 femtoliters [fL], compared with control mice with hemoglobin of 11.7 g/dl and MCV 47.9 fL). Remarkably, mice transplanted with ICN1-expressing BM demonstrated significantly delayed death from ICN1-induced leukemogenesis if they were maintained on the iron-restricted diet (P = 0.0001 by Mantel-Cox log rank analysis) (Figure 9B).

**Discussion**

We report an important role for Hrb in T-ALL by using mice with conditional deletion of CBP and mice transplanted with ICN1-transduced BM. Our data demonstrate that Hrb upregulation occurs in the vast majority of CBP-null T cell lymphomas, as a
downstream effect of activated Notch. Several important Notch target genes have been identified since its discovery as a potent oncogene (44). The majority of these target genes play crucial roles in physiologic Notch signaling. Here, we identify Hrb as a direct transcriptional target of Notch. Hrb is not required for physiologic Notch signaling, since thymocyte development is completely normal in \( Hrb^{-/-} \) mice. On the other hand, Hrb is clearly required for efficient Notch-induced lymphomagenesis, as evidenced by a delay in ICN1-induced T-ALL and a reduction in tumor incidence in the CBP-null mouse model. Clues from in vitro proliferation and development clearly suggested a failure to achieve optimal cell survival under the influence of ICN1 expression. Indeed, Hrb was required for survival of Notch-dependent human T-ALL cell lines in culture. Oncogenes (such as myc) that are upregulated by Notch are known to activate the cellular suicide program (45). ICN1 itself simultaneously offsets some of these effects through inhibition of p53 (46). However, lacking Hrb expression, the drive to apoptosis was inadequately suppressed, and we suspect this is a major cellular mechanism by which Hrb supports Notch-mediated lymphomagenesis.

Our data argue that Hrb facilitates oncogenesis by enhancing the ability of cells to take up transferrin. According to this model, in \( Hrb^{-/-} \) cells, particularly under the conditions of enforced proliferation driven by ICN1, iron uptake does not keep pace with the increased metabolic requirements of transformed cells, resulting in a state of iron depletion. This in turn induces the synthesis of CD71 in a failed attempt to acquire more iron from the cell surroundings. If this is unsuccessful, cell survival is compromised, leading to apoptosis (Figure 9C). In such a situation, leukemic progenitor cells cannot survive the stress of transformation in the face of relative iron starvation, which could explain the reduced incidence of T-ALL in CBP-null mice. In the case of ICN1-transduced BM, the oncogene is of such high potency that eventually even the severe defect of Hrb loss in transformed cell viability would ultimately be overcome by a rare secondary event that could bypass the apoptotic response. However, in the presence of supra-normal levels of transferrin in culture, Hrb is no longer required, as cells can compensate for inefficient uptake. It is likely that competition for transferrin within the BM is much more intense than that in vitro, thus amplifying the effect in vivo.

Notch activation has previously been shown to upregulate surface transferrin receptor (TIR) (47), and TIR has recently been shown to be a direct transcriptional target of c-myc (48), itself a direct target of Notch (14). Our work provides the first description at the molecular level of how oncogenic Notch signaling directs the transformed cell to meet increased metabolic demands for needed iron through upregulation of Hrb to accelerate transferrin uptake. Hrb was previously shown to be essential for efficient transferrin uptake by the use of knockdown experiments (12). Additionally, a role for Hrb in clathrin-mediated relocalization of VAMP7 from the cell surface to the endocytic compartment was demonstrated by Pryor et al. (11). Here, using \( Hrb^{-/-} \) mice, we provide an important context to Hrb’s role in clathrin-dependent endocytosis of cell surface TIR, downstream of oncogenic Notch signaling. Also, by utilizing a knockdown approach in Notch-dependent T-ALL cell lines, we demonstrate the importance of Hrb in the viability of human leukemia cells.

How does the failure to efficiently take up holo-transferrin lead to the death of ICN1-expressing thymocytes? Although it could result from a general metabolic failure as iron-containing enzymes become depleted, we suspect a more direct cause may be the failure to upregulate the p21 protein after ICN1 transduction. Much

![Figure 7](image-url)

Addition of excess holo-transferrin restores p21 protein levels in ICN1-transduced \( Hrb^{-/-} \) T cell precursors. (A) Western blot of lysates prepared from \( Hrb^{+/+} \) and \( Hrb^{-/-} \) MigR1− and ICN1− transduced thymocytes grown on OP9DL1 cocultures and harvested 5 days after transduction. Lysates blotted for p21 and actin (loading control). (B) Quantitative RT-PCR of mRNA isolated from the \( Hrb^{+/+} \) and \( Hrb^{-/-} \) MigR1− and ICN1− transduced thymocytes grown as in panel A. (C) Western blot of lysates from Figure 6G blotted for p21 and actin (loading control). (D) GFP-positive cell number of ICN1- and ICN1+p21−transduced WT hematopoietic precursors cocultured on OP9 stromal beds for 10 days. Data are shown as mean ± SD. *** \( P < 0.001 \).
The focus has been placed on the role of p21 as a tumor-suppressor protein due to its cell cycle inhibitory functions, but there is considerable evidence for a prosurvival function of p21 (49). p21 appears to have a special prooncogenic function in thymocytes because the incidence of T cell lymphomas is significantly decreased in p21 knockout mice (40). How p21 is regulated by activated Notch in T cell precursors is not known, but in cells of the renal glomerulus, it was shown to be due to p53-dependent upregulation of the p21 mRNA (50), and in keratinocytes, p21 is a direct transcriptional target of Notch1 activation (51). Interestingly, we found the transcript levels of p21 to be equal in ICN1-transduced cells, regardless of their Hrb status. These results suggest a posttranscriptional or posttranslational mechanism of regulation of p21. Fu et al. demonstrated that iron chelation leads to the ubiquitin-independent proteasomal degradation of p21 protein and a concomitant decrease in nucleocytoplasmic transport of p21 mRNA (43). It is likely that deficiency in iron utilization in Hrb−/− cells similarly prevented the ICN1-induced upregulation of p21 that occurs at the posttranscriptional level in WT cells, since addition of excess transferrin restored p21 accumulation. What role p21 plays in ICN1-induced leukemogenesis is not yet clear, but it is likely that p21 stabilization serves a prosurvival function, as has been shown in other T cell lymphomas (40). Indeed, coexpression of p21 with ICN1 led to a significant increase in cell accumulation.

Figure 8
Hrb is regulated by Notch signaling in human T-ALL cell lines and is required for cell proliferation and survival. (A) Quantitative RT-PCR analysis demonstrating the downregulation of Hrb following treatment with GSI (1 μM). Human T-ALL cell lines DND41 and T-ALL1 were treated with DMSO or GSI for 24 hours and 72 hours, respectively. Total RNA was prepared for reverse transcription and human Hrb mRNA abundance was measured by quantitative RT-PCR. (B) Nontarget control (NTC) and Hrb knockdown (HO22/HO23) stably transduced T-ALL cell lines were seeded at 100,000 cells/well and cultured for 7 days. Total cell numbers were counted based on forward and side scatter. (C) Western blot of lysates prepared from KOPTK1 and SUPT1 cell lines stably transduced with nontarget control and HO22/HO23 Hrb knockdown lentiviral vectors. Membranes were blotted with Hrb (clone H-300) and GAPDH (loading control). (D) Nontarget control and Hrb knockdown (HO22/HO23) stably-transduced T-ALL cell lines were seeded at 100,000 cells/well and cultured for 7 days. Cell survival was assessed using annexin V/PI staining. (E) DFO sensitivity of DND41 cells stably transduced with nontarget control or HO22. nontarget control or HO22 DND41 T-ALL cells were treated with 0, 25, or 50 μM DFO for 4 days. Following treatment, total cell numbers were acquired based on forward/side scatter and normalized to untreated. (F) Surface CD71 levels on NTC and HO23 stably transduced KOPTK1 cells treated with 0, 1.0, and 2.0 μM DFO. Data are shown as mean ± SD. **P < 0.01; ***P < 0.001.
The concept of nononcogene addiction (NOA) (52, 53) is attractive in this case, as we suspect that Notch-transformed cells are "addicted" to their high levels of Hrb. Identification of gene products that in and of themselves are not oncogenic, but that are required for efficient tumor growth, underlies this concept. Hrb overexpression is not known to be tumorigenic but is evidently required for efficient T cell malignancy. It has been proposed that discovering pathways that allow cancer cells to more efficiently metabolize nutrients may ultimately result in better treatments for cancer. Indeed, depriving cancer cells of iron with the use of an iron chelator (54, 55) or limiting the iron intake of mice with cancer (56) has previously been shown to inhibit cancer cell growth. This work provides what we believe is new evidence for the role of iron in the development of Notch-induced leukemias and raises the possibility that Hrb could be an attractive molecular target for T-ALL therapy, as it is not required for normal cell development, has a significant role in efficient iron uptake in Notch-transformed cells, and its loss sensitizes cells to iron depletion.

Additionally, our data suggest that the iron status of leukemia patients may be a contributory factor in the disease outcome. Intensified chemotherapeutic protocols for childhood leukemia have necessitated more frequent blood transfusions, which can predispose patients to iron overload (57). It may thus be detrimental to transfuse packed red blood cells into these patients, with little regard for the resulting iron overload that ensues. Although not thought to cause direct tissue damage as does occur in more chronically transfused patients, such as those with thalassemia and sickle cell anemia (58, 59), administration of a high iron load concurrently with chemotherapy may serve to enhance the survival properties of residual malignant cells. Future clinical studies should explore...
whether appropriate timing of blood transfusions may improve outcomes in leukemia or other cancer therapeutic trials.

Methods
Mice. CBP conditional knockout mice and MMTV-Cre mice were previously described (6). Hrb$^{-/-}$ mice on the background of MMTV-Cre/CBPlox/lox were intercrossed to generate Hrb$^{-/-}$ and Hrb$^{+/+}$ cohorts in the background of CBP deletion. ICN1/CBP-null mice were generated by mating CBPlox/lox mice to mice with ICN1 downstream of a floxed STOP sequence, which have been previously described (13). These mice were, in turn, mated to MMTV-Cre mice to activate ICN1 expression and delete CBP. Recipients for the BMT experiments were 4- to 8-week-old female C57BL/6 mice (Jackson Laboratory). Mice were observed daily for any signs of poor health. Sick mice were sacrificed and screened for tumors. All mouse studies were approved by the Mayo Clinic Animal Care and Use Committee.

Flow cytometry. Antibodies were from BD Biosciences — Pharmingen. Cells were stained for surface expression of the following markers using the antibodies in parentheses: CD4 (RM4-5), CD8 (53-6.7), CD25 (7D4), CD44 (IM7), and CD71 (C2). Cell survival was assessed using annexin V–Cy5 and PI staining (BD Biosciences — Pharmingen). Viable cells were gated according to their forward scatter and side scatter profiles.

Western blotting. Lysates from CBP-null lymphomas were prepared by snap-freezing lymphoma tissue in liquid N₂. Snap-frozen lymphoma tissue or PBS-washed T cell precursors and hematopoietic precursors grown on OP9/OPO9DL1 stromal beds were lysed in NP-40 lysis buffer (1% NP-40, 0.5% sodium deoxycholate, 5 mM EDTA, 50 mM Tris, pH 7.4, 150 mM NaCl, and a protease inhibitor cocktail from Roche). Blots were probed with antibodies that recognize Notch1 (m11a; Santa Cruz Biotechnology Inc.), gamma-secretase cleaved Notch1 (VAL1744; Cell Signaling), Hrb (H-300; Santa Cruz Biotechnology Inc.), p21 (M-19, Santa Cruz Biotechnology Inc.), CD71 (Invitrogen), Actin (Millipore), and GAPDH (Trevenig). The affinity-purified Hrb antibody (clone 368–562) was previously described (10).

Real-time RT-PCR. 1 μg total RNA isolated from T6E12 cells or KOPTK1 and TALL1 T-ALL cell lines was reverse-transcribed with SuperScript II (Invitrogen) according to the manual. All quantitative RT-PCR reactions were performed in triplicate. Primer sequences for real-time PCRs were as follows: human Hrb forward, 5′-GGCGGGTCATGGGAATAAC-3′; human Hrb reverse, 5′-TTCATCAAGTGTGCCTGCTC; human p21 forward, 5′-CCGGCCAAAGAGTCCCAAGGCAACAGCTGAACATCCTGAGGAAATGGTTTATCTGCCTGCTCTGCA; and human p21 reverse, 5′-CCGGCCAAAGAGTCCCAAGGCAACAGCTGAACATCCTGAGGAAATGGTTTATCTGCCTGCTCTGCA. Where appropriate, all statistical analyses were carried out using GraphPad Prism software using a 2-tailed Student’s t test or 2-way ANOVA with Bonferroni’s post-tests. P ≤ 0.05 was considered significant. All experiments were performed in triplicate and are presented as mean ± SD.

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