Hypoxic, glycolytic metabolism is a vulnerability of B-acute lymphoblastic leukemia-initiating cells

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SUMMARY

High-risk forms of B-acute lymphoblastic leukemia (B-ALL) remain a therapeutic challenge. Leukemia-initiating cells (LICs) self-renew and spark relapse and therefore have been the subject of intensive investigation; however, the properties of LICs in high-risk B-ALL are not well understood. Here, we use single-cell transcriptomics and quantitative xenotransplantation to understand LICs in MLL-rearranged (MLL-r) B-ALL. Compared with reported LIC frequencies in acute myeloid leukemia (AML), engraftable LICs in MLL-r B-ALL are abundant. Although we find that multipotent, self-renewing LICs are enriched among phenotypically undifferentiated B-ALL cells, LICs with the capacity to replenish the leukemic cellular diversity can emerge from more mature fractions. While inhibiting oxidative phosphorylation blunts blast proliferation, this intervention promotes LIC emergence. Conversely, inhibiting hypoxia and glycolysis impairs MLL-r B-ALL LICs, providing a therapeutic benefit in xenotransplantation systems. These findings provide insight into the aggressive nature of MLL-r B-ALL and provide a rationale for therapeutic targeting of hypoxia and glycolysis.
In brief

Morris et al. use single-cell transcriptomics to identify a candidate-initiating cell in B-acute lymphoblastic leukemia (B-ALL) with rearrangement of the KMT2A/MLL1 locus (MLL-r), finding that this population is plastic and exists in a hypoxic state that can be pharmacologically targeted.

INTRODUCTION

B-acute lymphoblastic leukemia (B-ALL) is heterogeneous, with certain unfavorable genetic subtypes continuing to bear poor prognosis despite the remarkable progress made in improving overall outcomes. B-ALL with rearrangement of the KMT2A/MLL1 locus (MLL-r) constitutes about 80% of B-ALL of infancy, with incidence diminishing in childhood but later increasing in adulthood (Boissel and Baruchel, 2018). MLL-r B-ALL is a very high-risk form of B-ALL, often presenting with high leukemic burden, corticosteroid resistance, and central nervous system infiltration (Pieters et al., 2007). This aggressive clinical behavior is associated with distinctive underlying biology, including coexpression of myeloid markers and a capacity to undergo a B-lymphoid-to-myeloid lineage switch in response to therapy (Pieters et al., 2007). Based on these clinical observations, we hypothesized that the aggressive nature of human MLL-r B-ALL is underpinned by a high content of leukemia-initiating cells (LICs) retaining hematopoietic stem cell (HSC)-and multipotent progenitor (MPP)-like differentiation states and multipotency programs.
LICs possess the capability to fully reconstitute fulminant leukemia at relapse or, experimentally, upon xenotransplantation. This definition was initially formed based on the foundational observation that only a small, specific minority of human acute myeloid leukemia (AML) cells could sustainably engraft immunodeficient mice, establishing engraftment in xenotransplantation as the gold standard to detect LICs (Lapidot et al., 1994). Although LICs have been previously accepted to be phenotypically primitive and rare, recent advances suggest that such early paradigms of LIC biology might be subject to revision. The use of improved immunodeficient mouse strains with heightened sensitivity for human stem cell engraftment has revealed that LICs are heterogeneous in their oncogenetics and can be detected in multiple differentiation states in both AML and ALL, although the common factors that maintain diverse LIC states are not well understood (Anderson et al., 2011; Aoki et al., 2015; McKenzie et al., 2019; Sarry et al., 2011). These findings could explain the limited clinical success of interventions targeting LICs despite intensive investigation of their molecular vulnerabilities (Pollyea and Jordan, 2017). Much of this investigation has focused on metabolic vulnerabilities, particularly oxidative phosphorylation as an LIC dependency in AML (Lagadinou et al., 2013).

Although findings illustrating LIC heterogeneity have led to reassessment of classical models of leukemic ontogeny, it remains generally accepted that leukemia cells are functionally heterogeneous with variable xenotransplantation capacities. Since the frequency of engraftable cells within individual leukemias—a readout of LIC content—is reproducibly of prognostic importance, better understanding of determinants of the LIC state has therapeutic potential (Monaco et al., 2004; Paczulla et al., 2017). Here, we use single-cell RNA sequencing (scRNA-seq) combined with xenotransplantation to define LICs in MLL-r B-ALL. We demonstrate that, although LICs are enriched in the most phenotypically primitive fraction of MLL-r B-ALL, they can emerge from blast populations with more differentiated immunophenotypes to reconstitute the full leukemic cellular diversity. Although most blasts utilize oxidative phosphorylation for expansion, primitive MLL-r B-ALL LICs can emerge from more differentiated populations by acquisition and maintenance of a relatively quiescent, hypoxic, glycolytic phenotype that can be therapeutically targeted. Consistent with its global effects on cancer cell metabolism, MYC plays a dual role in regulating both metabolic pathways in multiple strata of leukemic ontogeny (Fang et al., 2019; Lee et al., 2017). Together, our findings define mechanisms of LIC integrity and advance a candidate approach to therapy for high-risk forms of B-ALL.

**RESULTS**

**Single-cell RNA sequencing identifies candidate LICs in MLL-r B-ALL**

The ability to switch lineage from B-lymphoid to myeloid at relapse suggests that MLL-r B-ALL LICs possess multipotency (Pieters et al., 2007; Rossi et al., 2012). To determine whether we can capture this property of normal HSCs and MPPs in MLL-r B-ALL, we aimed to elicit both B-lymphoid and myeloid differentiation at the single-cell level. To this end, we employed early-passage patient-derived xenografts (PDXs) and primary patient cells. Upon xenotransplantation, human B-ALL cells infiltrated the liver, spleen, lymph nodes, and central nervous system of unconditioned non-obese diabetic (NOD).Cg-
Prkdc<sup>scid</sup>L2rg<sup>tm1Wjl</sup> (NSG) mice, recapitulating the clinical behavior of MLL-r B-ALL in humans (Figure S1A). MLL-r B-ALL cells nearly uniformly expressed the B-cell marker CD19, with rare cells expressing the myeloid marker CD33 while maintaining the morphology of immature lymphoblasts (Figure S1B). Following culture in a pro-myeloid and lymphoid MS5 stromal system, single MLL-r B-ALL cells showed B-lymphoid and myeloid multipotent differentiation outcomes with detection of MLL fusions in all populations (Figures S1C–S1E; Issaad et al., 1993; Doulatov et al., 2010). In this system, MLL-r B-ALL cells were highly clonogenic (Figure S1F). These findings were consistent with a previously reported dependence on stromal co-culture for B-ALL expansion (Pal et al., 2016). These results demonstrate single-cell-level B-lymphoid and myeloid potential consistent with multipotent HSC- and MPP-like programs in MLL-r B-ALL.

With the aim of identifying candidate LICs, we performed scRNA-seq using two human MLL-r B-ALL specimens (MLL-AF4 and MLL-ENL) with tracking of multipotency signatures. We obtained 11,709 viable, human CD45<sup>+</sup> single cells for analysis after performing quality controls. Unsupervised graph-based clustering of the combined datasets identified 12 transcriptional groups with unique signatures, with partial overlap of the two specimens (Figures 1A and 1B). Using the SingleCellNet algorithm to compare each cluster with normal adult bone marrow hematopoietic benchmarks, we found that most cells classified strongly as pro-B cells (Figure 1C; Tan and Cahan, 2019; van Galen et al., 2019). By tracking reported hematopoietic stem and progenitor cell (HSPC) signatures, transcriptional clusters 5 and 10 were advanced as candidate LIC-enriched populations; however, only cluster 10 had consistent enrichment of both HSPC signatures and contributions from both specimens (Figure 1D; Ivanova et al., 2002; Jaatinen et al., 2006). Compared with other populations, this population showed relatively low pre-B classification, concomitant classical dendritic and promonocyte signatures, and progenitor classification suggestive of relative dedifferentiation and potential multipotency (Figure 1C).

Furthermore, to further define differentiation state, we classified these MLL-r B-ALL cells against known HSPC benchmarks in another published scRNA-seq dataset, finding that cluster 10 showed relatively weak pre-B/NK classification— indicative of a lesser degree of lineage commitment— with strong multi-lymphoid progenitor (MLP) classification relative to other clusters, consistent with a less differentiated, multipotent-like state, as supported by consistent expression of HSPC signatures (Figures 1D and 1E; Pellin et al., 2019).

With the aim of isolating LIC-enriched clusters, we analyzed the expression of cell surface markers. We found that the HSPC marker CD34 was widely expressed, but the candidate LIC-enriched cluster 10 showed low expression of the committed progenitor marker CD38, further suggestive of a multipotent-like state, along with its relatively low B-cell commitment and consistent HSPC classification (Figure 1F). These cells expressed the B-cell markers CD86, CD79A, and CD24 and the monocyte marker CD58, suggestive of mixed lineage potential (Figure 1F). Given these findings and considering the B-lymphoid and myeloid multipotency programs of MLL-r B-ALL, we analyzed expression of CD34 and CD38 as well as the additional primitive human HSPC markers CD90 and CD45RA by flow cytometry. We found that MLL-r B-ALLs generally contained three distinguishable populations that we designated based on their surface immunophenotype: rare, undifferentiated CD34<sup>+</sup> CD38<sup>−</sup> CD90<sup>−</sup> CD45RA<sup>+</sup> cells (34+38−) with an MLP-like
surface profile consistent with the observed scRNA-seq classification; more abundant, progenitor-like CD34+ CD38+ cells (34+38+); and the remaining CD34− blasts, all of which could be readily sorted from multiple leukemias (34−; Figures 1G, 1H, and S1G). Using additional scRNA-seq datasets, we confirmed evidence of similar hierarchical organization of MLL-r B-ALL and possibly BCR-ABL-driven B-ALL, although with differing ontogeny in the latter case (Figures S2A–S2I; Candelli et al., 2022; Chen et al., 2021; Witkowski et al., 2020). In line with this finding, using RNA-seq of fluorescence-activated cell sorting (FACS)-purified populations, we found that the 34+38− fraction bore the strongest primitive HSPC signature (Figures S2J and S2K). These findings are suggestive of a phenotypically primitive, LIC-enriched population in MLL-r B-ALL.

**Functional cellular heterogeneity in MLL-r B-ALL**

Since LICs are typically quiescent relative to other blasts, we first analyzed cell cycle status (Guan et al., 2003; Saito et al., 2010; Terpstra et al., 1996). First, using PDX cells, we found that the MLP-like, 34+38− fraction contained the highest proportion of cells in G0 phase (Figures 2A and 2B). Next, we used MS5 assays to quantify leukemic progenitors, finding that more actively cycling 34+38+ cells possessed the highest frequency of short-term clonogenic cells (Figure 2C). We next used limiting dilution xenotransplantation to quantify bona fide LICs (Hu and Smyth, 2009). Using the onset of terminal leukemia as our endpoint, as we initially hypothesized, we first observed that unfractionated human MLL-r B-ALL possessed a remarkably high frequency of LICs (1/426 engrafting cells [1/1,417–1/128 95% confidence interval]) compared with AML (347-fold more frequent than the most efficiently engrafting AML reported previously; Sarry et al., 2011). Upon fractionation, we found that the MLP-like 34+38− population caused leukemia with the shortest latency and contained the highest frequency of engrafting LICs (Figures 2D–2F; Table S2). To exclude an artifactual effect of the PDX system, we corroborated this finding using primary cells sorted from patient peripheral blood (Figures S1H and S1I). These findings indicate that the 34+38− fraction of human MLL-r B-ALL is enriched in engraftable LICs.

**Latent MLL-r B-ALL LICs in 34− and 34+38+ fractions**

Although the 34+38− fraction contained abundant LICs, we observed that terminal leukemia also developed in recipients of the 34+38+ and 34− fractions, albeit at lower efficiency (Figures 2D–2F, S1H, and S1I). When we compared the cellular content of leukemia derived from 34+38−, 34+38+, and 34− PDX cells, we found that the full diversity emerged from each source in the terminal disease (Figures 3A and 3B). Moreover, by secondary xenotransplantation, we found that leukemia derived from each source contained serially transplantable LICs (Figure 3C). To directly observe this effect during leukemic reconstitution in vivo, we transplanted the least phenotypically primitive 34− cells into NSG mice and isolated bone marrow at defined time points prior to the expected onset of terminal leukemia, finding that 34− cells gradually replenished the CD34+ compartment, suggestive of latent LIC potential (Figures 3D and 3E). Consistent with these xenotransplantation findings, following 4 weeks of culture on MS5 stroma, we found that all input populations could initiate colonies at similar frequencies (Figures S3A and S3B).
To further understand these findings, we cultured sorted populations of 34+38−, 34+38+, and 34− cells on MS5 stroma and monitored population flux. We observed that the content of cultures derived from 34+38+, 34+38−, and 34− cells showed similar CD34+ content by 15 days (Figures S3C and S3D). We observed similar cell state plasticity in primary patient blasts in vivo and in culture (Figures S4A–S4D). To control for possible effects of sorting impurities impacting plasticity readouts, we observed cell state plasticity down to the single-cell level using primary patient cells, where we also observed latent myeloid potential in single-cell clones from all phenotypic populations (Figures S4E–S4H).

Finally, to track surface-marker profile and primitive stem cell signatures, we used single-cell cellular indexing of transcriptomes by sequencing (CITE-seq). With this approach, we found that cells with primitive HSPC signatures were present throughout ontogeny, suggesting that cells with the least primitive differentiation states can bear latent stem cell signatures (Figure 3F). Together, these data form a basis for the observed emergence of LICs from more mature populations.

**MYC-mediated heterogeneity and plasticity of LICs**

We aimed to understand the mechanistic determinants of the MLL-r B-ALL LIC state. To this end, we queried the transcriptional profile of phenotypically immature, LIC-enriched 34+38− cells relative to other populations. Moreover, to uncover signatures associated with leukemic engraftment and reconstitution, we also evaluated the profile of mature 34− blasts as they were actively reconstituting the cellular diversity following xenotransplantation (“34−r” cells) versus 34− cells— with the same apparent differentiation state—engrafted at steady state in the presence of other populations (Figure 4A). We found that 34+38−, 34−, and 34−r cells showed markedly divergent transcriptional profiles by RNA-seq, despite the identical immunophenotypes of 34− and 34−r cells (Figure 4B). We noticed that the divergent transcriptional profiles were indicative of variability in metabolic state, with LIC-enriched 34+38− cells showing signatures of hypoxia and glycolysis, and actively reconstituting 34−r cells showing signatures of oxidative phosphorylation and MYC target gene expression (Figure 4C). By performing chromatin immunoprecipitation with sequencing (ChIP-seq) in the B-lymphoid and myeloid MLL-r cell line MV4;11 and the MLL-r B-ALL cell line RS4;11, we found that transcripts encoded by genes bound by MYC were enriched in 34−r cells, suggestive of a MYC-driven engraftment and reconstitution process (Figure 4D; Lange et al., 1987). Moreover, in a publicly available dataset, we observed that MLL-AF4 bound the MYC locus in human HSPCs (Figure S5A).

Since MYC is a central regulator of cancer cell metabolism, we treated cells with IBET-151, a BET bromodomain inhibitor that decreases MYC activity known to impair leukemic growth and monitored B-ALL reconstitution (Dawson et al., 2011; Fang et al., 2019; Lee et al., 2017). As expected, we found that IBET-151 diminished MYC levels (Figure 4E). Treatment of 34− cells either in culture or in vivo with IBET-151 blocked the acquisition of the CD34+ state, consistent with a role for MYC in mediating the reconstitution process (Figures 4F–4I). We next generated a doxycycline-inducible, GFP-tagged lentiviral vector for ectopic MYC expression in MLL-r B-ALL cells (Figure 4J). Following transduction and sorting of GFP+ B-ALL PDX cells, culture in the presence of doxycycline for 14 days...
preserved CD34+ content (Figure 4K). Together, these data suggest that MYC promotes the reconstitution of MLL-r B-ALL from CD34− cells.

Inhibition of oxidative metabolism promotes emergence of primitive LICs

Our transcriptional profiling suggested that engrafting MLL-r B-ALL cells are metabolically heterogeneous, with 34−r cells recruiting latent LIC properties possibly through activating oxidative phosphorylation to engraft and reconstitute the B-ALL cellular diversity (Figure 4C). We hypothesized that, by targeting oxidative phosphorylation, we could impede emergence of LICs from 34− blasts. This hypothesis is supported by findings in AML, where LICs have been reported to depend on mitochondrial oxidative phosphorylation (Jones et al., 2018; Lagadinou et al., 2013). Using the Mitotracker dye to quantify mitochondrial content in MLL-r B-ALL cells, we confirmed that 34−r cells contained a higher level of mitochondrial content compared with bulk cells, with a trend toward higher mitochondrial numbers (Figures 5A, 5B, S5B, and S5C). Using the Seahorse Mito Stress assay, we observed that 34−r cells possessed higher rates of mitochondrial respiration compared with steady-state MLL-r B-ALL blasts (Figures 5C and 5D). We found that MYC activity modulated mitochondrial metabolism in MLL-r cells using both IBET-151 and the direct MYC inhibitor KI-MS2-008 (MS2) (Figures S5D–S5T; Struntz et al., 2019). To determine whether targeting oxidative metabolism affected reconstitution of MLL-r B-ALL, we used tigecycline, an inhibitor of mitochondrial translation, which has been advanced as a candidate therapy targeting LICs in AML and chronic myeloid leukemia (Kuntz et al., 2017; Skrtic et al., 2011). We found that tigecycline decreased viability of MLL-r B-ALL cells in culture and diminished mitochondrial content (Figures S5U–S5X). We next treated 34− cells reconstituting in vivo with a course of tigecycline starting at 3 weeks post-engraftment, when we would expect the reconstitution process to commence (Figure 3D). We found that this decreased overall disease burden (Figures 5E–5H). However, at completion of the treatment course, tigecycline promoted emergence of CD34+ cells, suggestive of induction of an LIC-enriched state (Figures 5I and 5J). We found that, despite the early effect of tigecycline on chimerism, there was no benefit in survival studies (Figures 5K and 5L). Secondary transplantation of vehicle- or tigecycline-treated leukemia confirmed that tigecycline treatment promoted LIC emergence from 34− cells in vivo (Figure 5M). These findings indicate that inhibition of oxidative phosphorylation could enforce an LIC-enriched state and that oxidative phosphorylation is not an LIC dependency.

34+38− B-ALL LICs maintain a glycolytic state

Since blockade of mitochondrial translation facilitates LIC emergence, this further supported the notion that MLL-r B-ALL LICs maintain a relative glycolytic state. Indeed, we observed that treatment of MLL-r B-ALL cells with tigecycline not only decreased the oxygen consumption rate (OCR) but also the OCR/extracellular acidification rate (ECAR) ratio, consistent with a shift toward glycolysis (Figures 6A–6C; Potter et al., 2016). Therefore, we next sought to better understand the energy state of 34+38− LIC-enriched cells. 34+38− cells showed lower levels of reactive oxygen species and mitochondrial content compared with 34− cells (Figures 6D–6F, S6A, and S6B). Ex vivo culture of MLL-r B-ALL cells under hypoxic conditions preserved the 34+38− state more effectively than normoxic culture (Figure 6G). By analyzing our ChIP-seq data, we found that MYC-bound
peaks were enriched in gene ontology terms related to both oxidative phosphorylation and the hypoxic response and that MYC directly bound glycolysis-associated loci (Figures 6H and 6I; Fang et al., 2019). We did not observe consistent enrichment of HIF-1α protein in 34+38− cells, suggesting that the relative dependence on glycolysis is due to lower levels of oxidative phosphorylation (Figure S6C). Consistent with this, in an independent dataset, we observed that predicted corticosteroid-resistant LICs showed a lower oxidative phosphorylation signature (Candelli et al., 2022; Figure S6D). Inhibiting MYC with IBET-151 impaired glycolytic activity in MV4;11 cells (Figures 6J and 6K). Together, these data indicate that the LIC-enriched 34+38− fraction maintains a relative MYC-maintained glycolytic state.

**Inhibiting hypoxic glycolysis to target MLL-r B-ALL LICs**

Inhibition of glycolysis has been pursued as an approach to cancer therapy as a means of targeting the Warburg effect, and so we next examined glycolysis as a target in MLL-r B-ALL. Dichloroacetate (DCA) is an inhibitor of pyruvate dehydrogenase kinase that shunts pyruvate into the tricarboxylic acid (TCA) cycle and away from lactate production (Bonnet et al., 2007). As expected, DCA increased oxidative metabolism in MV4;11 cells and MLL-r B-ALL PDX cells (Figures S6E–S6H). Treatment of MV4;11 cells with either DCA or the direct glycolysis inhibitor 3-BP diminished leukemic progenitor cells in colony formation assays (Figures S6I). DCA decreased the viability of B-ALL cells selectively under hypoxic conditions, where cells would be more reliant on glycolysis (Figure S6J). Treatment of mice xenografted with 34+38− B-ALL cells with DCA resulted in a modest but significant survival benefit (Figure S6K).

Since glycolysis is preferentially utilized under hypoxic conditions, we treated MLL-r cell lines and xenografts with the HIF-1α inhibitor echinomycin (Wang et al., 2011). Echinomycin treatment decreased the viability of a panel of human MLL-r B-ALL specimens while relatively preserving healthy human HSPCs, suggestive of a degree of HIF-1α dependence in all blasts (Figures 7A and S6L). Echinomycin treatment ablated the viability and colony forming capacity of MV4;11 cells and the viability of RS4;11 cells at low nM concentrations (Figures 7B and 7C). Moreover, we found that echinomycin treatment impaired leukemic progenitors in a mouse model of MLL-AF9-driven mixed B-lymphoid and myeloid leukemia (Figure S6M; Rowe et al., 2019). We found that, following 2 weeks of echinomycin treatment, mice bearing MLL-r B-ALL xenografts showed diminished overall disease burden as well as CD34+ content in vivo (Figures 7D–7F). By monitoring for survival following an upfront 2-week treatment course, we found that echinomycin prolonged survival of mice bearing B-ALL xenografts (Figure 7G). Secondary transplantation indicated LIC depletion (Figure 7H). Finally, we tested the potential of combined treatment with echinomycin and DCA, finding that this combination promoted survival (Figures 7I and S6N). Echinomycin did not cause overt toxicity and effected only subtle changes on hematologic parameters, with myeloid skewing observed in the peripheral blood and at the HSPC level (Figures S6O and S6P). Together, these findings indicate the feasibility of inhibiting glycolysis and hypoxic signaling as a means of targeting LICs in B-ALL.
DISCUSSION

In this study, we aimed to gain understanding of LIC biology using LIC-rich, human B-lymphoid and myeloid multipotent MLL-r B-ALL as a model. Although the heterogeneous cells of MLL-r B-ALL appear to follow a hierarchical organization, in that LICs are enriched in the most undifferentiated fraction, engraftable, self-renewing cells emerge from all levels of this hierarchy. We find that LICs in MLL-r B-ALL bear hallmarks of cancer stem cells—some of which were until recently thought to be specific to solid tumors—including cell state plasticity (Gupta et al., 2019; Jones et al., 2018; Lee et al., 2017). Our findings uncover aspects of the biology underlying the aggressive clinical behavior of this high-risk form of B-ALL.

Since the initial supposition of LICs, models of leukemic ontogeny have been built upon paradigms of normal hematopoiesis (Bonnet and Dick, 1997; Kreso and Dick, 2014; Pollyea and Jordan, 2017). Based on seminal studies performed in AML, the “classic” conclusions that LICs are both uniform and rare akin to normal HSCs have gradually been revised (Pollyea and Jordan, 2017). At first glance, our data would appear to be consistent with this classic model in that we found distinct populations stratified from least to most primitive based on transcriptional profiles, immunophenotypes, and engraftment capacities. In B-ALL, though engrafting LICs have been reported in multiple populations, suggestive of a stochastic rather than hierarchical organization, their relative frequency in each of these apparent differentiation states has not previously been quantified (Prieto et al., 2018; Aoki et al., 2015; Bardini et al., 2015; Kong et al., 2008; Lapidot et al., 1994; le Viseur et al., 2008). Our findings indicate that LICs are quantitatively enriched in relatively dedifferentiated fractions but can also emerge from more differentiated fractions, accounting for this apparent stochasticity (Elder et al., 2017). Use of more immunodeficient mouse models with heightened sensitivity for human engraftment has suggested plasticity of AML cells like our findings, although the molecular mechanisms underpinning this effect were unclear (Ng et al., 2016; Sarry et al., 2011). The consensus is emerging that, across many forms of leukemia, leukemic ontogeny departs from the strict stratification and unidirectional differentiation of healthy hematopoiesis and that LICs are not uniform in their marker profiles (Ng et al., 2016). Despite LIC paradigms undergoing revision, the notion that engraftable LIC content predicts disease outcome highlights the importance of LICs in leukemic pathobiology and provides an ongoing impetus for investigation of the determinants of their state (Monaco et al., 2004; Paczulla et al., 2017). Our findings expand understanding of B-ALL LICs by demonstrating latency in phenotypically mature (i.e., 34−) populations in addition to the known genetic and immunophenotypic heterogeneity (Anderson et al., 2011; Aoki et al., 2015).

Our data reinforce the idea that LIC identity can be fluid, as has been described in stem cells of solid tumors and sporadically in LICs (Fumagalli et al., 2020; Gupta et al., 2019). Use of NSG mice led to the evidence that AML LICs within CD34− fractions could reconstitute the full cellular diversity of the original AML specimen, although the mechanism of this cell state interconversion was unknown (Sarry et al., 2011). A recent study reported that acute promyelocytic leukemia cells undergoing differentiation driven by retinoic acid could re-acquire the LIC state following drug withdrawal (McKenzie et al., 2019). In B-ALL, we
detected LIC emergence from all immunophenotypic populations—even the least primitive CD34− blasts—with the capability of reconstituting the full cellular diversity of the original leukemia. These findings are likely relevant to alterations in the LIC state occurring as an adaptation to therapy at relapse of B-ALL (Dobson et al., 2020).

Our efforts to uncover determinants of the LIC state led us to MYC signaling. Oncogenic MYC activity is a downstream effector of transforming MLL translocations, and mitochondrial turnover is important for LIC homeostasis (Dawson et al., 2011; Pei et al., 2018). MYC can drive both mitochondrial oxidative metabolism and glycolysis in cancer (Fang et al., 2019; Lee et al., 2017). We impaired MYC activity, finding that this intervention perturbed cellular metabolism and blocked the conversion of CD34− to CD34+ cells (Bardini et al., 2018; Dawson et al., 2011; Delmore et al., 2011). Although we found that direct inhibition of mitochondrial translation transiently impeded growth of cells in vivo, this intervention promoted the emergence of CD34+ cells enriched in engrafting LICs, suggesting that inhibition of oxidative phosphorylation facilitates transition to the quiescent, glycolytic LIC state. Our data indicate that LICs are quiescent and glycolytic with low levels of oxidative phosphorylation and reactive oxygen species compared with other populations. In AML, LICs also contain relatively low levels of reactive oxygen species; however, in contrast to our findings, AML LICs seem to have low glycolytic reserve and are therefore susceptible to inhibition of oxidative phosphorylation (Jones et al., 2018; Lagadinou et al., 2013). The metabolic state of MLL-r B-ALL LICs seems to be co-opted from normal long-term HSCs and could explain the concomitant retention of multipotency programs in this form of leukemia (Nakamura-Ishizu et al., 2020; Qiu et al., 2021; Spencer et al., 2014; Takubo et al., 2010). MYC seems to fulfill context-specific roles in B-ALL cells both as an activator of oxidative phosphorylation to promote leukemic reconstitution by 34− blasts as well as to maintain glycolysis to support the primitive LIC state. Although MLL-r B-ALL typically contains relatively few somatic mutations, future study will investigate the role of clonal heterogeneity and DNA methylation alterations in establishing LIC ontogeny (Andersson et al., 2015; Tejedor et al., 2021).

Our conclusion that the latent LICs can emerge from relatively differentiated blasts can be connected to the clinical behavior of this form of high-risk B-ALL and may have therapeutic implications. We find latent myeloid transdifferentiation potential within single-cell-derived clones from each stratum of the leukemic hierarchy, consistent with a multipotent cell state (Rossi et al., 2012). The coexisting LIC and lineage plasticity in MLL-r B-ALL provides two distinct mechanisms of chemotherapy evasion. We conclude that the high functional LIC content and plasticity in both lineage and the LIC state likely contribute to the poor outcomes of MLL-r B-ALL (Pieters et al., 2019). Our approach to understanding the mechanisms of LIC variation uncovered glycolysis as a potential vulnerability in MLL-r B-ALL. A recent report used scRNA-seq to characterize corticosteroid-resistant, putative relapse-initiating cells, confirming that these cells seemed to be largely quiescent (Candelli et al., 2022). Better understanding of the molecular basis of LICs in other forms of leukemia relative to normal hematopoiesis could lead to candidate therapies.

Overall, we show that LICs in high-risk B-ALL (1) maintain an HSC- and MPP-like state and retain multipotency programs; (2) are frequent compared with reported LIC

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frequencies in AML; (3) are relatively glycolytic, with the LIC state implemented upon inhibition of oxidative phosphorylation, contrasting with dependency of LICs on oxidative phosphorylation in myeloid leukemias (Kuntz et al., 2017; Lagadinou et al., 2013; Skrtic et al., 2011); and (4) the glycolytic state of MLL-r B-ALL LICs is a therapeutic vulnerability. Our findings in MLL-r B-ALL reinforce the notion that LICs are adaptable, providing possible explanations as to why therapies targeting LICs have yet to prove widespread efficacy despite being the object of intense investigation for over 2 decades (Pollyea and Jordan, 2017; Saygin et al., 2019). Our data would suggest that a hybrid hierarchical and stochastic LIC ontogeny exists in MLL-r B-ALL, with a highly engraftable population detected but considerable flux through all populations likely manifesting as stochasticity seen in prior studies and, to a degree, in our transplants (Aoki et al., 2015; le Viseur et al., 2008). Further studies will determine the relative cell-intrinsic and extrinsic contributions to the relatively glycolytic LIC state in B-ALL, as well as the mechanisms relating metabolism and stemness in B-ALL, which remain elusive (Hayashi et al., 2018).

Limitations of the study

Our study was performed using human-patient-derived leukemia cells, which provides translational relevance compared with studies reliant exclusively on inbred mouse models. However, this human system comes with the limitation that only relatively few donor specimens could be used for quantitative xenotransplantation and in-depth analysis of LIC function (seven used in this study), with inter-patient variability challenging to mitigate using such a design. Our use of FACS-sorted populations in xenotransplantation assays could confound results due to incomplete purity, in particular underestimating differences in LIC content between populations, though we aimed to mitigate this using single-cell-level studies. We also narrowed our focus to MLL-r B-ALL, raising the questions as to generalizability to other forms of high-risk B-ALL. Indeed, our analysis of BCR-ABL-driven B-ALL would suggest alternative LIC ontogeny that should be investigated in future study.

STAR METHODS

RESOURCE AVAILABILITY

Lead contact—Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Grant Rowe (grant_rowe@dfci.harvard.edu).

Materials availability—Plasmids generated in this study will be deposited to Addgene prior to the date of publication or are available upon request from the lead contact.

Data and code availability

- scRNA-seq, CITE-seq, ChIP-seq, and bulk RNA-seq (processed human data) have been deposited in Gene Expression Omnibus and are publicly available as of the date of publication. All raw sequencing data derived from human samples have been deposited in the Sequence Read Archive and are available through
request to the NIH Database of Genotypes and Phenotypes as of the date of publication. Accession numbers are listed in the key resources table.

- This paper does not report original code.
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

**Animal models**—All studies were approved by the Boston Children’s Hospital Institutional Animal Care and Use Committee. Eight-week-old unconditioned female NSG mice (Jackson Laboratory stock 005557) were transplanted with the indicated cell sources at the indicated doses by tail vein injection. Mice were randomly allotted to each experimental group. Mice were followed to the onset of terminal leukemia as assessed by the onset of morbidity, judged by investigators blinded to the experimental conditions. Mice were housed in autoclaved cages with sterilized food and water.

For tigecycline treatment, mice were treated for two days with 25 mg/kg tigecycline (LKT labs) dissolved in 10% DMSO/90% (5% Kleptose HPB in PBS). The dose was then escalated to 50 mg/kg for three days and then 100 mg/kg for 7 days. After 48 hours, mice were euthanized for endpoint analysis. For IBET-151 treatment, mice were treated with IBET-151 (R and D Systems) at a dose of 30 mg/kg in 10% DMSO/90% (5% Kleptose HPB in PBS) for 5 days per week for two weeks followed by euthanization for analysis. For *in vivo* dichloroacetate treatment, mice were engrafted with the indicated leukemias and dichloroacetate added to the drinking water to target a dose of 100 mg/kg/day based on average daily water consumption (Bonnet et al., 2007). Echinomycin was given at 10 μg/kg per day for five days on, two days off, five days on (10 total treatments per mouse) (Wang et al., 2011).

**Human subjects**—Use of primary human cells, PDX cells, and cell lines was reviewed and approved by the Boston Children’s Hospital Institutional Review Board. All primary patient and PDX specimens were procured following informed consent. Specimens of both sexes were included. Sex and age of specimens included in this study is provided in Table S1. Human cell lines were obtained from the ATCC.

**Cell culture**—Patient derived leukemia cells were cultured on MS5 stromal cells in the presence of 50 ng/mL recombinant human stem cell factor (SCF), 50 ng/mL recombinant human thrombopoietin (TPO), 10 ng/mL recombinant human FLT3 ligand (FLT3L), and 10 ng/mL recombinant human interleukin-7 (IL-7, all from R and D Systems) in Myelocult H5100 (Stem Cell Technologies) supplemented with 1% penicillin/streptomycin (Issaad et al., 1993) with modifications based on (Doulatov et al., 2010) to facilitate lymphoid cell growth; i.e., without horse serum or hydrocortisone (Issaad et al., 1993). Leukemia 1/MLL-AF4 was most amenable to *in vitro* culture and so was used for most culture-based experiments. For *in vitro* LDA experiments, 5,000 MS5 cells were plated in wells of gelatin-coated Nunc 96 well plates (Fisher Scientific) with cytokines 48 hours prior to FACS-based sorting of leukemia cells directly into the wells.
Human MV4; 11 and RS4; 11 cells (ATCC) were cultured in RPMI with 10% fetal calf serum supplemented with penicillin and streptomycin.

As indicated in Table S1, some specimens were subjected to focused somatic mutation analysis (Rapid Heme Panel (Kluk et al., 2016)).

METHOD DETAILS

Flow cytometry and cell sorting—Data were acquired on either BD LSR Fortessa or LSR II Instruments (BS Biosciences). Cells were sorted on a BD FACS Aria (BD Biosciences) with a 100 μm nozzle. Mitotracker Green and CellRox were purchased from Thermo. Mitotracker Green was used at 300 nM and all experiments included verapamil at 50 μM with incubation at 37°C for 30 minutes prior to analysis (de Almeida et al., 2017). Antibodies used are listed in the key resources table.

Morphology—For morphologic analysis, leukemia cells were spun onto slides and stained with May-Grunwald and Giemsa stains (Sigma) sequentially. Transmission electron microscopy was performed at the electron microscopy core at Harvard Medical School.

Recombinant DNA—The human MYC cDNA was purchased from Addgene (pDONR223_MYC_WT, a gift from Jesse Boehm and Matthew Meyerson and David Root (plasmid # 82927; http://n2t.net/addgene:82927; RRID: Addgene_82927). The MYC cDNA was cloned into the pCW57.1 vector (gift from David Root (Addgene plasmid # 41393; http://n2t.net/addgene:41393; RRID: Addgene_41393) using LR clonase (Thermo). The purified plasmid was used to generate lentivirus in HEK-293T cells, which was used to transduce MV4; 11 cells. A stable transduced polyclonal line was selected with 0.5 μg/mL puromycin (Thermo). The MYC cDNA was also cloned into pINDUCER21 (ORF-EG) which was a gift from Stephen Elledge and Thomas Westbrook (Addgene plasmid # 46948; http://n2t.net/addgene:46948; RRID: Addgene_46948) to generate lentivirus for transduction of B-ALL patient derived xenografts. Gene expression was confirmed by Western blotting following doxycycline exposure.

Western blotting—Antibodies are listed in the key resources table. After lysis in SDS sample buffer, proteins were resolved on 4–20% gradient polyacrylamide gels (BioRad) using the Tris buffer system. Proteins were transferred to PVDF membranes, and membranes were blocked and probed per the manufacturer’s protocol. Signal was developed using the ECL system (Pierce) and detected by exposure to radiography film. Western blot band intensities were quantified with ImageJ.

Seahorse Mito Stress assay—The Seahorse assays were performed using the Agilent Seahorse XF Cell Mito Stress Test kit using injections of 1 μM oligomycin, 1 μM carbonyl cyanide-4 (trifluoromethoxy) phenylhydrazone (FCCP) and 0.5 μM each of antimycin A and rotenone at the intervals indicated. Prior to analysis, cells were equilibrated for one hour in Mito Stress medium supplemented with 1 mM pyruvate, 10 mM glucose, and 2 mM/L-glutamine. The instrument is located at the Seahorse Core at Brigham and Women’s Hospital. All assays were normalized based on the number of viable cells included following drug pre-treatment to control for effects of drug treatment on viability. For assays using
primary leukemia cells, cells were cultured on MS5 stroma for 72 hours prior to B-ALL cell purification by human CD45 bead enrichment (Miltenyi) followed by Seahorse Mito Stress analysis.

**Long-term culture initiating cell assay**—The indicated populations were sorted by FACS and placed onto MS5 stroma with cytokines either under normoxia (21%) or hypoxia (5%) for four weeks. At that time, viable, human CD45+ cells were sorted into 96-well plates at the indicated doses. After two weeks, outgrowths were blindly scored.

**Reverse transcription PCR**—RNA was isolated from the indicated cell populations using Trizol reagent. cDNA was prepared using the miScript II kit (Qiagen). Primers are listed in the key resources table. MLL-AF4 primers were reported previously (Jansen et al., 2005). Band were visualized following gel electrophoresis and ethidium bromide staining. Product specificity was confirmed by Sanger sequencing.

**Single cell RNA sequencing**—For scRNA-seq using the Seq-Well platform, 20,000 cells were applied to one Seq-Well array pre-loaded with mRNA capture beads (ChemGenes) as previously described (Gierahn et al., 2017). The arrays were sealed with a polycarbonate membrane with a 0.01 μm pore size permissive of exchange of buffers but not macromolecules. Following cell lysis, transcript hybridization, and bead recovery by buffer exchange, reverse transcription was performed using Maxima H Minus Reverse Transcriptase (ThermoFisher) with random second-strand synthesis was performed to generate double stranded cDNA. Exonuclease I treatment (New England Biolabs) was performed to remove excess primers. PCR was performed using the following primer sequence 5′-AAGCAGTGGTATCAACGCAGAGT–3′ and the KAPAHiFi PCR Mastermix (Kapa Biosystems) with 2,000 beads per 50 μL reaction volume. Sequencing libraries were generated using the Illumina Nextera XT protocol using custom N700 sequencing indices. Libraries were sequenced using Next-Seq 75 cycle high output sequencing kits with 20 base read 1 sequence and 50 base read 2 sequence.

For inDrop-seq, FACS-sorted cells at 100,000 cells/mL were encapsulated in 2–3 nL droplets using a microfluidic device (located at the Harvard Medical School Single Cell Core) and the libraries were made following a previously described protocol (Klein et al., 2015; Zilionis et al., 2017), with the following modifications in the primer sequences. RT primers on hydrogel beads-5′ CGATTGATCAACGTAATACGACTCACTATAGGTGTGCGGGTGCGCAG[bc1,8nt ]GTCTCGTGGGCTCGAGATGTGTATAAGAGACAG[bc2,8nt]NNNNNNNTTTTTTTTTTTTTTTTTT- 3′. 

R1-N6 primer sequence (step 151 in the library prep protocol in (Zilionis et al., 2017))- 5′ TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGNNNNNNN-3′.

PCR primer sequences (steps 157 and 160 in the library prep protocol in (Zilionis et al., 2017))- 5′-AATGATACGGCGACCACCGA GATCTACACXXXXXXXTCGTCGGCAGCGTC-3′, where XXXXXX is an index sequence for multiplexing libraries.
5′-CAAGCAGAGACGGCATACGAGATGGGTGTCGGGTGCAG-3′.

With these modifications in the primer sequences, custom sequencing primers are no longer required. Reverse transcription was performed at 50°C for two hours followed by 15 minutes at 70°C. scRNA-seq was library preparation was performed by the Single Cell Core at Harvard Medical School, Boston, MA using the NEBNext mRNA Second Strand Synthesis Module with random hexamer priming.

For CITE-seq, cells were pre-stained with barcoded antibodies against CD38 and CD34 (Biolegend) and inDrop libraries were prepared as above.

**Bulk RNA sequencing library preparation**—Cells were sorted by FACS and lysed in Trizol reagent (Thermo). RNA was isolated using RNAeasy columns (Qiagen) and low input libraries prepared in collaboration with the Molecular Biology Core at the Dana-Farber Cancer Institute using the NEBNext Single Cell/ Low Input RNA Library Prep Kit for Illumina (New England Biolabs).

**Data analysis**

**InDrop single-cell RNA-sequencing**: Raw sequencing reads were processed using the inDrop pipeline ([https://github.com/indrops/indrops](https://github.com/indrops/indrops)) using default parameters (Klein et al., 2015) The hg38 reference genome was used for alignment of sequencing reads. We used scImpute to account for dropout rates in single-cell RNA-seq data and obtain an imputed count matrix that was used for all downstream analysis described (Li and Li, 2018). We used scImpute with the parameter ‘Kcluster = 10’. To analyze imputed single cell inDrop data we performed quality control, dimensionality reduction, clustering and differential expression analysis using CellRouter (Lummertz da Rocha et al., 2018). For this leukemia, we applied the following quality control metrics: all genes that were not detected in at least 20 cells were excluded. All cells with less than 200 genes detected were also excluded. As expression of ribosomal or mitochondrial genes is indicative of technical variation in single-cell RNA-seq data we also removed cells where the proportion of the transcript counts derived from mitochondrial genes was greater than 10% (Ilicic et al., 2016). After such quality control of the imputed count matrix, we retained 5,153 cells with a median of 15,214 genes detected per cell.

**Integrated analysis of leukemia samples**: We used Seurat 4.0 to integrate our leukemia samples, which were generated using the inDrop and SeqWell technologies (Stuart et al., 2019). Briefly, we individually normalized each sample using the LogNormalize() function, and identified the 3000 most variable genes in each dataset based on a variance stabilizing transformation. Then, we used the FindIntegrationAnchors() function followed by the IntegrateData() function from Seurat to integrate the datasets. Following data integration, we performed a cell cycle regression of S and G2/M phase scores, which preserves the cell cycle signal associated with cell differentiation. We then performed UMAP and clustering analysis using the first 20 principal components. For clustering, we used 20 principal components in the function Find-Neighbors(), and defined the parameter ‘resolution’ of the FindClusters function as 0.25. Differential expression analysis was used to identify cluster-specific genes.
by selecting those more highly expressed in each cluster based on a fold change cutoff of 0.2.

**Seq-Well single-cell RNA-sequencing:** Read alignment was performed as described (Macosko et al., 2015). Briefly, for each NextSeq sequencing run, raw sequencing data was converted to FASTQ files using bcl2fastq2 that were demultiplexed by Nextera N700 indices corresponding to individual samples. Reads were first aligned to hg19, and individual reads were tagged according to the 12-bp barcode sequence and the 8-bp UMI contained in read 1 of each fragment. Following alignment, reads were binned and collapsed onto 12-bp cell barcodes that corresponded to individual beads using Drop-seq tools (http://mccarrollab.com/dropseq). Barcodes were collapsed with a single-base error tolerance (Hamming distance = 1), with additional provisions for single insertions or deletions. An identical collapsing scheme (Hamming distance = 1) was then applied to UMIs to obtain quantitative counts of individual mRNA molecules. We also used scImpute to impute the raw counts matrix obtained, with the parameter of ‘Kcluster = 6’. For this leukemia, we removed all genes expressed in less than 10 cells and also removed all cells expressing less than 500 genes. Cells with transcript counts derived from mitochondrial genes larger than 10% were also removed. After QC, we retained 6,320 cells with a median of 2,588.5 genes detected per cell.

**Normalization:** Both inDrop and Seq-Well data were analyzed with CellRouter. In CellRouter, transcript counts are normalized using a global scaling normalization method that normalizes expression measurements for each cell by the total expression, multiplied by a scale factor of 10,000, and log-transformed the result.

**Re-analysis publicly available leukemia samples:** We re-analyzed scRNA-seq data from four Ph+ samples at diagnosis generated by Witkowski et al., 2020, GEO accession number GSE134759. Briefly, we subset CD19 + cells by selecting individual cells whose normalized expression level is higher than the mean expression of all cells expressing CD19 above zero. This approach resulted in 1,452 CD19 + single cells from four Ph + samples, which were integrated using Seurat as described above. UMAP and clustering analysis were performed using 20 principal components and the parameter “resolution” of the FindClusters function was set to 0.5.

We also re-analyzed data generated by Candelli et al. (2022), accession number EGAS00001003986. From this dataset, we analyzed 13,250 individual cells after quality control (genes expressed in less than 25 cells and cells expressing less than 300 genes were removed from all downstream analysis). These samples were integrated with Seurat as previously described. UMAP and clustering analysis were performed using 17 principal components and the parameter “resolution” of the FindClusters function was set to 0.75.

Finally, we reanalyzed a dataset generated by Chen et al. (2021), composed of 88,228 single cells from 18 MLL-r B-ALL patient samples for which a Seurat object was downloaded from Synapse at https://www.synapse.org/. To integrate these samples we used the algorithm Harmony (Korsunsky et al., 2019). After integration, UMAP and clustering analysis were performed using 20 principal components and the parameter “resolution” of the FindClusters
function was set to 0.5. Regression of the difference between cell cycle scores for the 
S-phase and G₂/M-phase was performed for the three studies mentioned above.

**SingleCellNet analysis:** We downloaded scRNA-seq data from the GEO accession number 
GSE116256 (van Galen et al., 2019). This study performed a random sampling of 
hematopoietic cells in the normal and leukemic bone marrow (BM) ecosystem. We 
reanalyzed five healthy BM samples published with this study and used cell types identified 
by the authors to train machine learning models of cell type identity of BM cells using 
SingleCellNet. After training, we classified each single cell in our leukemia samples as 
belonging to any of the classes in our training dataset. Additionally, we also used the dataset 
generated by Pellin et al., (2019) as training dataset to classify leukemia samples.

**Signature scores:** We downloaded gene lists from 
https://www.gsea-msigdb.org/gsea/msigdb. Specifically, we downloaded 
the following gene sets: HALLMARK_MYC_TARGETS_V1.txt, 
HALLMARK_OXIDATIVE_PHOSPHORYLATION.txt, 
JAATINEN_HEMATOPOIETIC_STEM_CELL_UP, and 
IVANOV_A_HEMATOPOIESIS_STEM_CELL.txt. Then, we used CellRouter to calculate 
signature scores for each cell and plotted the distribution of these scores.

For the signature analysis of Candelli et al. (2022), we used data 
from 15 peripheral blood samples of infants with MLL-rearranged ALL 
(EGA accession number EGAS00001003986). We considered GSEA gene sets 
‘HALLMARK_OXIDATIVE_PHOSPHORYLATION’, ‘HALLMARK_GLYCOLYSIS’, 
and ‘HALLMARK_HYPOXIA’ and used them to calculate a module score for each gene 
set as described in Candelli et al., (2022). Annotation of resistant (LIC) and sensitive 
(differentiated blasts) populations is as in Candelli et al., (2022).

**CITE-seq:** We converted binary files from the Illumina run to fastq files corresponding 
to 4 reads of indrops3 library with bcl2fastq/2.18.0.12 without read trimming. https:// 
support.illumina.com/sequencing/sequencing_software/bcl2fastq-conversion-software.html. 
We created a single fastq file containing cell and sample barcode information in the header 
of the read with umis fastqtransform https://github.com/vals/umis. We then wrote a custom 
python script to count antibody barcodes and assign the counts to a proper cell/sample pair. 
The input parameters of the script were antibody barcodes (tags), a fastq file from step 2, 
and a list of sample barcodes. We ran the script using the wrapper https://github.com/hbc/ 
rowe2020_indrop_citeseq_hbc03948/blob/master/04.cite_seq_count.sh.

**Bulk RNA-sequencing:** Fastq files containing single-end RNA-seq reads were aligned with 
Tophat 2.0.12 against the UCSC hg38 reference genome using Bowtie 2.2.4 with default 
settings (Kim et al., 2013; Langmead and Salzberg, 2012). Gene level counts were obtained 
using the sub-Read featureCounts program (v1.5.1) using the parameter “--primary” and 
gene models from the UCSC hg38 Illumina iGenomes annotation package (Liao et al., 
2014). Differential expression analysis was performed with the DESeq2 package using 
‘ashr’ as a shrinkage estimator for the log₂ fold changes (Love et al., 2014). Genes with log₂ 
fold changes > 0.5 and P_{adj} < 0.05 were considered significantly differentially expressed.
For visualization of gene expression patterns and hierarchical clustering, read counts were normalized using size factors as available by the DESeq2 package and visualized using the pheatmap package (Love et al., 2014).

**ChIP-seq**—ChIP-seq was performed as previously described and in accordance with the Encode guidelines (Chapuy et al., 2013; Landt et al., 2012). We used $100 \times 10^6$ exponentially growing MV4; 11 or RS4; 11 cells. Cells were fixed with 1% formaldehyde for 10 minutes at room temperature and then quenched with 125 mM glycine for 5 minutes and washed with PBS. Nuclei were isolated using the Nucleo EZ isolation buffer (Sigma) and resuspended in 10 mM Tris-HCl, pH 8.0, 1 mM EDTA, 0.1% SDS with 1x HALT protease inhibitor (ThermoFisher). Chromatin was fragmented by sonication on an E220 Covaris focused sonication machine using the parameters of 140 mV, 5% duty factor, 200 cycles/burst for 14 minutes. The following antibodies were used for ChIP: MYC (Cell Signaling 9402). ChIP-seq libraries were prepared using Swift S2 Acel reagents (Swift 21096) on a Beckman Coulter Biomek i7 liquid handling platform from approximately 1 ng of DNA according to manufacturer’s protocol and using 14 cycles of PCR amplification. Sequencing libraries were quantified by Qubit fluorometer and Agilent TapeStation 2200. Library pooling and indexing was evaluated by shallow sequencing on Illumina MiSeq. Subsequently, libraries were sequenced on NovaSeq targeting 40 million 100 bp read pairs by the Molecular Biology Core facilities at the Dana-Farber Cancer Institute. Sequencing reads were processed using the AQUAS pipeline (https://github.com/kundajelab/chipseq_pipeline) with minor modifications and according to the ENCODE3 guidelines. Reads were aligned to the hg38 genome build using BWA-ALN and peaks were called using MACS2.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

Statistical testing was performed as described in the figure legends using either Prism (Graphpad) or Excel (Microsoft). For each experiment, n and precision measures are provided in the figure legends. Power calculations were used to estimate cohort sizes for *in vivo* experiments. P-values are shown in the figures or are presented in the figure legends.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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**Highlights**

- *MLL-r* B-ALL cells are varied, with a defined population enriched in initiating cells
- *MLL-r* B-ALL-initiating cells are plastic, emerging from more differentiated states
- MYC-driven metabolic heterogeneity may provide a therapeutic opportunity
Figure 1. Single-cell heterogeneity of MLL-r B-ALL
(A) Human CD45+ MLL-r B-ALL cells were isolated from peripheral blood and analyzed by scRNA-seq. A combined uniform manifold approximation and projection (UMAP) including both specimens was used to visualize 12 populations with clustering (top) and contribution by specimen (bottom) shown.
(B) Heatmap showing differential transcript expression across each cluster in both combined specimens.
(C) Single cells were classified against a benchmark human bone marrow scRNA-seq dataset (van Galen et al., 2019).
(D) Enrichment of the indicated HSPC signatures was quantified across each cell and visualized using UMAP, with violin plots showing expression of the signature-associated scores.
(E) Single-cell transcriptomes were classified against benchmark HSPC and differentiated populations from a published dataset (Pellin et al., 2019).
(F) The expression of the indicated cell surface markers in each population is shown.
(G) Representative flow cytometry distributions of cells using the indicated markers.
(H) The indicated populations were isolated from representative leukemic marrow (MLL-AF4) by fluorescence-activated cell sorting (FACS) and morphology examined following May-Grunwald-Giemsa staining (scale bars represent 10 μm).
Figure 2. Functional hierarchy in MLL-r B-ALL

(A and B) The indicated populations were isolated by FACS and cell cycle analyzed by flow cytometry. Cell cycle distribution of each population was quantified (results aggregated over two independent experiments; n = 11 xenografted mice tested; *p < 0.05 by one-way ANOVA, results show mean ± SEM).

(C) The indicated populations were sorted onto MS5 stromal layers at doses of 1, 2, or 4 cells per well. After 4 weeks, outgrowths at each dose were tabulated and progenitor cell content quantified by limiting dilution analysis. Estimated progenitor cell frequency is presented. Results are aggregated over two independent experiments (for 34+38− versus 34+38+, X^2 = 64.3, p = 1 × 10^{-15}; for 34+38− versus 34−, X^2 = 14.9, p = 0.0001; for 34+38+ versus 34−, X^2 = 18.1, p = 0.00002).

(D) Four hundred cells of the indicated fractions were xenotransplanted into unconditioned NSG recipients and the incidence of terminal leukemia monitored (results aggregated over two independent transplantation experiments; p = 0.008 for 34+38− versus 34+38+, p = 0.005 for 34+38− versus 34−, and p = 0.9 for 34+38+ versus 34− comparisons by log rank test).

(E) NSG mice were transplanted with the indicated doses of cells from each population and the incidence of terminal leukemia quantified at each dose aggregated over four experiments including two patient donors (see Table S2).

(F) LIC content of each population was quantified by in vivo limiting dilution analysis. Results are aggregated over four independent xenotransplantation experiments including two leukemias (for 34+38− versus 34+38+, X^2 = 7.7, p = 0.006; for 34+38− versus 34−, X^2 = 6, p = 0.01; for 34+38+ versus 34−, X^2 = 0.1, p = 0.7).
Figure 3. LIC emergence from all hierarchical levels

(A) Representative flow cytometry profiles of terminal leukemias derived from the indicated transplanted cell populations gated on viable and human CD45+ cells within the bone marrow.

(B) Proportions of each cell type in outcome leukemias derived from the indicated transplanted cell sources are presented (p = NS by ANOVA comparing each outcome population between leukemia sources except in MLL-ENL leukemia, where p < 0.05 comparing the 34+38+ and 34− outcomes of leukemias derived from 34+38+ to both 34+38− and 34− derived leukemia).

(C) Terminal leukemias derived from the indicated cell populations of MLL-AF4 primary transplants were transplanted into secondary recipients at the indicated doses and the onset of terminal leukemia monitored.

(D and E) 34− cells were purified from MLL-AF4 leukemia and transplanted into recipient mice. Reconstitution of CD34+ populations was quantified over time.

(F) CITE-seq was performed on MLL-AF4 leukemia. Cells were scored for enrichment of HSPC signatures (signature score above 95th percentile cutoff defining “HSPC”) and plotted versus CD34 and CD38 expression determined from barcoded antibodies (Ivanova et al., 2002; Jaatinen et al., 2006). In all experiments, results are presented as mean ± SEM and results compared by Student’s t test, with p value shown.
Figure 4. MYC regulates MLL-r B-ALL LIC state

(A) Schematic showing the input populations sorted for RNA-seq, including 34+38− cells and 34− cells steady state (left) and from regenerating 34− cells (right) in mouse bone marrow.

(B) RNA-seq was performed on the indicated FACS-sorted populations and differential gene expression analyzed (cutoff log₂ fold change > 0.5 and padj < 0.05).

(C) Gene set enrichment analysis (GSEA) was used to identify differentially enriched signatures (circle size indicates normalized enrichment score [NES] with shading representing false discovery rate [FDR] q value).

(D) MYC ChIP-seq was performed in MV4;11 and RS4;11 cells and peaks associated with transcriptional start sites identified. GSEA was performed using preranked lists comparing transcript expression of 34−r versus 34− cells.

(E) MV4;11 or RS4;11 cells were treated for 48 h with the indicated concentrations of IBET-151, at which time MYC protein was measured by western blot with bands quantified (relative to β-actin).

(F and G) FACS-sorted CD34− cells were cultured on MS5 stroma for 14 days with DMSO vehicle or 50 nM IBET-151, at which time the CD34+ content of the cultures was quantified (n = 5 independent donors).
(H and I) CD34−r cells were engrafted into NSG mice for 3 weeks, at which time a 2-week treatment course with IBET-151 began, after which the human CD45+ content of the bone marrow was analyzed by flow cytometry with the indicated markers (n = 5 mice per group).

(J) K562 cells were transduced with a doxycycline-inducible MYC vector and treated with or without doxycycline for 72 h, at which time MYC levels were measured by western blotting.

(K) MLL-r B-ALL cells were transduced or not transduced with the doxycycline-inducible MYC vector, sorted for GFP+ cells, and cultured on MS5 stroma with cytokines with or without doxycycline for 14 days, at which time CD34+ content of the cultures was measured by flow cytometry. NS, not significant.

In all experiments, results are presented as mean ± SEM and results compared by Student’s t test, with p value shown.
Figure 5. Targeting oxidative phosphorylation drives LIC emergence

(A and B) Active mitochondria were quantified in bulk leukemia or 34–r cells using Mitotracker green, with representative flow cytometry results presented compared with background of unstained cells (n = 5 biologic replicates each over two independent experiments).

(C and D) The indicated B-ALL cell populations were cultured on MS5 stroma for 72 h prior to analysis of oxygen consumption rate (OCR) by Seahorse Mito Stress Assay (results normalized to 10^6 viable cells and aggregated over two independent experiments, four biologic replicates each).

(E and F) NSG mice were engrafted with MLL-r B-ALL for 3 weeks, at which time treatment with tigecycline commenced and continued for 2 weeks, when the mice were euthanized and mouse and human chimerism within the bone marrow analyzed (n = 8 subjects in vehicle and tigecycline groups).

(G and H) Spleens were isolated and weighed at the endpoint from (C) (scale bar represents 1 cm).

(I and J) CD34+ content of the xenografts isolated in (E) was analyzed by flow cytometry (n = 8 subjects in vehicle and tigecycline groups).
(K and L) Xenotransplanted NSG mice were treated with the same course of tigecycline as in (E) but followed to the onset of terminal leukemia (n = 7 subjects in vehicle and tigecycline groups; survival compared with log rank test).

(M) NSG mice were secondarily transplanted with primary grafts from (E) and survival compared by log rank test (n = 5 subjects in each group).

In all experiments, unless otherwise stated, results are presented as mean ± SEM and results compared by Student’s t test, with p value shown.
Figure 6. Primitive MLL-r B-ALL LICs maintain a glycolytic state

(A–C) Human MLL-r B-ALL cells were cultured in the presence of vehicle or tigecycline (1 μM) for 72 h, at which time OCR and ECAR were measured by Seahorse assay (results aggregated over three biologic replicates).

(D) Relative Mitotracker signal was quantified in the indicated populations from two independent leukemia sources (n = 6 biologic replicates over two experiments compared by one-way ANOVA with p values shown).

(E and F) CellRox was used to stain for reactive oxygen species in MLL-r B-ALL cells and relative signal quantified (n = 6 biologic replicates comparisons by ANOVA with p values shown).

(G) PDX cells were cultured for 10 days under either 21% oxygen or 5% oxygen, and the composition of the cultures was measured by flow cytometry (result combined across three independent experiments).

(H) Gene ontology analysis was used to query for terms enriched in MYC ChIP-seq peaks in RS4;11 cells.

(I) Visualization of MYC ChIP-seq peaks associated with the indicated glycolytic genes in MV4;11 and RS4;11, with control input for each shown.
(J and K) MV4;11 cells were cultured in the presence or absence of 50 nM IBET-151 for 72 h, at which time the extracellular acidification rate (ECAR) was measured using the Seahorse assay.

Unless otherwise indicated, in all experiments, results are presented as mean ± SEM and results compared by Student’s t test, with p value shown.
Figure 7. Targeting glycolysis and hypoxic signaling

(A) The indicated MLL-1 B-ALL specimens or normal human bone marrow (BM) CD34+ HSPCs were cultured in the presence of the indicated concentrations of echinomycin for 72 h, at which time cell viability was quantified.

(B) One thousand MV4;11 cells were cultured in the presence or absence of the indicated concentrations of echinomycin for 48 h and then placed in methylcellulose culture. After 5 days, colony formation was quantified (compared by t test with p values shown).

(C) MV4;11 or RS4;11 cells were cultured with the indicated concentration of echinomycin for 48 h, at which time viability was measured (n = 3 independent experiments).

(D–F) NSG mice were transplanted with MLL-AF4 B-ALL cells. Three weeks following engraftment, mice were treated with echinomycin for 2 weeks, at which time human CD45 and CD34 chimerism were analyzed.

(G) Mice xenotransplanted with MLL-AF4 B-ALL were treated for 2 weeks with echinomycin or vehicle (treatment window indicated) and survival monitored following completion of treatment.

(H) Cells from mice in (G) were secondarily transplanted and survival of recipient mice monitored.

(I) Mice xenotransplanted with MLL-AF4 leukemia were treated for echinomycin or vehicle for 2 weeks and then with or without DCA and survival monitored. In all experiments,
unless otherwise stated, results are presented as mean ± SEM and results compared by Student’s t test or log rank test for survival analysis, with p value shown.
**KEY RESOURCES TABLE**

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| Anti-human CD19 PE (4G7) | Biolegend | Cat#392505; RRID: AB_2750096 |
| Anti-human CD45 FITC (HI30) | BD Biosciences | Cat#561865; RRID: AB_10896120 |
| Anti-human CD33 APC (P67.6) | BD Biosciences | Cat#340474; RRID: AB_400518 |
| Anti-human CD34 PE-Cy7 (8G12) | BD Biosciences | Cat#348791; RRID: AB_400381 |
| Anti-human CD38 PE | BD Biosciences | Cat#347687; RRID: AB_400341 |
| Anti-human CD45 APC | BD Biosciences | Cat#555485; RRID: AB_398600 |
| Anti-human CD45RA FITC | BD Biosciences | Cat#347723; RRID: AB_400343 |
| Anti-human CD90 BV605 | BD Biosciences | Cat#562685; RRID: AB_2744468 |
| Anti-mouse CD45.1-APC-Cy7 (A20) | BioLegend | Cat#110716; RRID: AB_313505 |
| Anti-Ki67 APC | Biolegend | Cat#350513; RRID: AB_10959326 |
| Anti-c-MYC | Cell Signal | Cat#9402; RRID: AB_2151827 |
| Anti-beta-actin (8H10D10) | Cell Signal | Cat#3700; RRID: AB_2242334 |
| **Bacterial and virus strains** |        |            |
| Sthl3 E. coli | Thermo Fisher | C737303 |
| **Biological samples** |        |            |
| Patient derived xenografts | This paper | N/A |
| Primary patient leukemia cells | This paper | N/A |
| **Chemicals, peptides, and recombinant proteins** |        |            |
| May-Grünwald stain | Sigma-Aldrich | MG500 |
| Giemsa stain | Sigma-Aldrich | R03055 |
| Doxycycline hyclate | Sigma-Aldrich | D9891 |
| Recombinant human stem cell factor | R and D Systems | 255-SC-010 |
| Recombinant human FLT3-ligand | R and D Systems | 308-FK-005 |
| Recombinant human thrombopoietin | R and D Systems | 288-TP-005 |
| Recombinant human interleukin-7 | R and D Systems | 207-IL-005 |
| Mitotracker green | Thermo Fisher | CAS: 201860-17-5 |
| IBET-151 | SelleckChem | CAS: 1300031-49-5 |
| 3-bromopyruvate | Sigma | CAS: 113-59-3 |
| Dichloroacetate | Sigma | CAS: 2156-56-1 |
| Echinomycin | Tocris | CAS: 512-64-1 |
| Tgecycline | LKT Labs | CAS: 220620-09-7 |
| CellRox Deep Red | Thermo Fisher | C10422 |
| **Critical commercial assays** |        |            |
| Seahorse Mito Stress Test | Agilent | 103015-100 |
| CellTiter Glo | Promega | G7570 |
| **Deposited data** |        |            |
| Raw data | This paper | Sequence Read Archive SRA: phs002492 |
| Analyzed data | This paper | Gene Expression Omnibus GEO: GSE147862 |

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| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Experimental models: Cell lines** |
| MV4; 11             | ATCC   | ATCC CRL-9591; RRID:CVCL_0064 |
| RS4; 11             | ATCC   | ATCC CRL-1873; RRID:CVCL_0093 |
| HEK293             | ATCC   | ATCC CRL-3216; RRID:CVCL_0045 |
| MS5 mouse bone marrow stroma | N/A   | RRID:CVCL_2128 |
| **Experimental models: Organisms/strains** |
| NSG mice             | Jackson Laboratory | Strain #:005557; RRID:BCBC_4142 |
| **Oligonucleotides** |
| Human **MLL-AF4** forward: MLL-F1 5′-CGCCTCAGCCAACCTACAG-3′ | Jansen et al. (2005) | N/A |
| Human **MLL-AF4** reverse: AF4-R1 5′-AGGTCGTCTTGAGCATGGA-3′ | Jansen et al. (2005) | N/A |
| Human **GAPDH** forward: GAPDH-F 5′-GTCTCTCTGAACGTCAACAGCG-3′ | This paper | N/A |
| Human **GAPDH** reverse: GAPDH-R 5′-ACCACCCCTGTTGCTGTAGCCAA-3′ | This paper | N/A |
| **Recombinant DNA** |
| pPAX2               | Addgene | 12260 |
| pCMV-VSV-G           | Addgene | 8454 |
| pDONR223_MYC_WT      | Addgene | 82927 |
| pCW57.1             | Addgene | 41393 |
| **Software and algorithms** |
| SingleCellNet       | Tan and Cahan (2019) | https://github.com/pcahan1/singleCellNet/ |
| Bowtie2             | Langmead and Salzberg (2012) | http://bowtie-bio.sourceforge.net/bowtie2/index.shtml |
| TopHat2             | Kim et al. (2013) | https://ccb.jhu.edu/software/tophat/index.shtml |
| Indrops pipeline    | Klein et al. (2015) | https://github.com/indrops/indrops |
| Drop-seq pipeline   | Macosko et al. (2015) | http://mccarrolllab.com/dropseq |
| GSEA                | Subramanian et al. (2005) | https://www.gsea-msigdb.org/gsea/index.jsp |
| AQUAS               | N/A | https://github.com/kundajelab/chipseq_pipeline |
| CellRouter          | Lammertz da Rocha et al., 2018 | https://github.com/edroaldo/cellrouter |
| Seurat              | Stuart et al. (2019) | https://github.com/satjialab/seurat |
| Bcl2fastq           | Illumina | https://support.illumina.com/sequencing-sequencing_software/bcl2fastq-conversion-software.html |
| Umis                | N/A | https://github.com/vals/umis |