Discovery of Dihydroartemisinin and Dasatinib Drug Combination to Cure Poor outcome BCR-ABL+ Acute Lymphoblastic Leukemia

Harpreet Singh

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Discovery of Dihydroartemisinin and Dasatinib Drug Combination to Cure Poor-outcome BCR-ABL+ Acute Lymphoblastic Leukemia

A Dissertation
Presented for
The Graduate Studies Council
The University of Tennessee
Health Science Center

In Partial Fulfillment
Of the Requirements for the Degree
Doctor of Philosophy
From The University of Tennessee

By
Harpreet Singh
August 2012
DEDICATION

I dedicate this dissertation to my parents, family members and close friends for their endless love, support and guidance.
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ABSTRACT

Oncogenic signaling by the Philadelphia chromosome-encoded BCR-ABL fusion kinase initiates and drives both Philadelphia chromosome-positive acute lymphoblastic leukemia (Ph+ ALL) and chronic myelogenous leukemia (CML). Food and Drug Administration (FDA)-approved BCR-ABL-specific kinase inhibitors (BCR-ABL–KIs) imatinib, dasatinib and nilotinib induce prolonged remissions in CML but poor leukemia-reduction and relapse-control in Ph+ ALL. The relative primary BCR-ABL–KI-resistance in Ph+ ALL patients carrying predominantly BCR-ABLWT disease cannot be attributed to drug-resistant BCR-ABL mutations (BCR-ABL MUTANTS), and remains poorly understood.

We established a cell-based platform to evaluate the modulation of anti-Ph+ ALL activity of drugs by both tumor-extrinsic cytokines normally present in the leukemia microenvironment and tumor-intrinsic vector-mediated alteration of candidate genes. We identified that BCR-ABLWT-driven Ph+ ALL cells are rendered significantly resistant against all FDA-approved BCR-ABL–KIs by 1) Several host-cytokines, but most dominantly by Interleukin7 (IL7), and 2) Cell-intrinsic functional-loss of IKAROS. Utilizing IL7-deficient recipient mice, we demonstrated that physiological levels of IL7 significantly attenuate the survival benefit derived from dasatinib monotherapy against Ph+ ALL. Follow-up mechanistic studies using cell signaling and gene expression comparisons indicated that IL7 imparts chemo-refractoriness by initiating IL7-pSTAT5-c-MYC signaling. Interestingly, IKAROS haploinsufficiency, which has been previously associated with poor clinical prognosis in patients with Ph+ ALL, was recently demonstrated to directly de-repress c-MYC expression. This suggested that both cell-extrinsic IL7 and cell-intrinsic IKAROS loss converge on c-MYC, which may act as node of leukemia BCR-ABL–KI-refractoriness. We confirmed that vector-mediated modifications imitating IL7 induction of STAT5 activity, functional-loss of IKAROS, and c-MYC over-expression, all selectively enrich Ph+ ALL cells during prolonged exposures to imatinib and dasatinib, thus revealing new chemo-refractory phenotypes.

Contemporary medicine advocates co-treatment with agents of complementary mechanism of drug-action to tackle drug resistance. To identify combination agents that could overcome the dasatinib-resistant phenotypes of Ph+ ALL, we screened a library of 3200 agents including known anti-infective and chemotherapy drugs. We discovered that a well known anti-malarial drug dihydroartemisinin (DHA) killed host-IL7-protected BCR-ABLWT, c-MYC-overexpressing BCR-ABLWT and BCR-ABL MUTANT Ph+ ALL cells in vitro. In vivo, DHA displayed weak activity as a single agent but its addition synergistically augmented the leukemia reduction by dasatinib, relative to either of the two drugs alone. Remarkably, DHA and dasatinib combination regimen eliminated host-protected dasatinib-refractory persistent leukemia and improved long-term survival from 0 to >90% in a murine model that faithfully captures the BCR-ABL–KI drug-refractoriness of human Ph+ ALL.

This study: 1) Uncovers novel mechanisms of clinical drug-resistance against BCR-ABL–KIs, 2) Identifies increased levels of IL7, pSTAT5 and c-MYC protein, and
IKAROS haploinsufficiency as potential biomarkers of BCR-ABL-targeted drug-resistance, 3) Strongly supports clinical exploration of the BCR-ABL–KI and DHA combinations for treating patients with Ph+ ALL, and 4) Establishes a paradigm for investigating frequently overlooked host-tumor-drug interactions.
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CHAPTER 1. INTRODUCTION

Target Focused Therapies in New Era Oncology and Road to Cure

Ongoing efforts to understand critical signaling pathways that undergo deregulation during tumorigenesis identify specific oncogenes and driver mutations integral for the sustenance and proliferation of various cancer types [1-2]. The development of agents that modulate such key molecular targets should improve patient tolerance and cure rates over conventional non-specific chemotherapy by selectively killing cancer cells [1-3].

Trends of Drug Discovery Industry in Post-imatinib Era

Imatinib (Gleevec by Novartis) is the first example of a cancer-specific small molecule inhibitor that was ever approved for clinical use, in particular to treat human CML. Initial clinical success of imatinib in dramatically controlling the previously poor outcome CML marked a paradigm shift in our approach to anti-cancer therapy and jump-started drug discovery efforts against an ever-growing list of validated molecular targets in other cancers [4-8]. Since then, remarkable progress has been made in the field of target focused drug discovery [9-11] especially for agents that particularly inhibit deregulated receptor or non-receptor kinase proteins [8, 10-14], similar to imatinib, and are commonly referred to as kinase inhibitors (KIs). The fast-pace development of KIs is facilitated by the chemically druggable structure of kinase proteins, advances in computational modeling, medicinal chemistry and high throughput biophysical assays that compare binding of chemicals against normal and deregulated kinases [5, 15-20]. Current automated drug pipelines against diverse kinases and non-kinase targets generate hundreds of potential leads, providing tremendous hope to patients suffering from diverse leukemias and solid tumors worldwide [5, 9-10, 14, 21].

Current Clinical Status of Target Focused Drugs in Oncology

Rational target focused drug discovery has tremendously decreased the total time required between the initial identification of a potential agent and its actual testing in cancer patients. Within the last decade, multiple cancer-selective drugs and biological agents have been formally approved and many more are being evaluated in clinical trials, namely against CML, Ph+ ALL, gastro-intestinal stromal tumors, non-small cell lung carcinoma, melanoma, sarcoma, renal cancers etc. However, the majority of cancer-selective agents often induce suboptimal primary responses and offer meaningful benefit in only a sub-set of cancer patients, which frequently lasts only for a short period of time [7, 9-10, 12, 14, 22-23]. Most frustratingly, all current agents which kill the respective cancer cells with potent selective activities in vitro, invariably fail to eradicate the same cancer cells expressing intact molecular targets in human patients [7, 9-10, 12, 14, 22-24]. As a consequence, many pre-clinical agents have failed to produce
meaningful long-term clinical benefit to the majority of patients despite expected pharmacodynamic changes in the intended molecular targets in vivo [7, 9-10, 12, 14, 22-24], and as a consequence, a few agents have been completely dropped. Since negative results are often not reported, the number of such failed agents is bound to be much higher.

Paradoxical clinical (in vivo) drug-resistance observed against available cancer-selective agents in general, remains poorly understood. It likely stems from our limited experience with this modern class of oncology agents and has to be addressed to improve long term cancer outcomes.

Targeted Therapy against Philadelphia Chromosome-positive (Ph+) Leukemias

From among the available cancer targeting drugs, the most extensive preclinical and clinical data is available for imatinib and imatinib-like drugs, primarily for their use against a set of leukemias defined by a unique cytogenetic abnormality – Philadelphia Chromosome [25-26].

The Philadelphia Chromosome

The Philadelphia chromosome is the hallmark of CML and a high-risk subset of acute lymphoblastic leukemia (Ph+ ALL). The Philadelphia chromosome arises from a balanced translocation between chromosomes 9 and 22 [25-26]. It encodes an abnormal fusion protein, BCR-ABL, which plays an essential role in initiating and driving both CML and Ph+ ALL [25-26]. The oncogenic potential of BCR-ABL protein arises from the constitutive activation of the tyrosine kinase domain (KD) of ABL in the fused protein configuration.

In CML, mainly the p210\textsuperscript{BCR-ABL} isoform is expressed in hematopoietic stem cells (HSCs) which are conceptually referred to as leukemia stem cells (LSCs) [27], and in both myeloid and lymphoid progeny. Whereas in Ph+ ALL, expression of mainly p185\textsuperscript{BCR-ABL} or alternative p210\textsuperscript{BCR-ABL} isoform is restricted to the lymphoid B cell lineage, which display robust LSC-like function to both initiate and propagate leukemia during serial transplantation assays.

Rationale and Agents for BCR-ABL Targeted Therapy

The required role for BCR-ABL in sustaining Ph+ leukemias, an attribute referred to as ‘oncogene addiction’, rationalizes the inhibition of BCR-ABL as an attractive therapeutic strategy in patients with CML and Ph+ ALL [25-26].
The design of imatinib allows it to selectively bind to the KD of ABL and inhibit the constitutive tyrosine kinase activity of BCR-ABL fusion protein. Imatinib was initially approved against CML alone [28]. Currently, three BCR-ABL–specific kinase inhibitors (BCR-ABL–KIs) namely imatinib (first generation), nilotinib and dasatinib (second generation) are FDA-approved for use against both CML and Ph+ ALL. First and second generation BCR-ABL–KIs differ in their chemical structures, pharmacokinetic properties, binding sites on the ABL, potency to inhibit ABL kinase activity, spectrum of off-target inhibition of other kinases and associated adverse effects [20].

Outcomes of BCR-ABL Targeted Therapy in CML

At the time of diagnosis, CML most commonly is detected in its chronic (early) phase and less frequently in accelerated or blast (advanced) phase. In most of the CML patients suffering from chronic phase of the disease, continuous BCR-ABL–KI monotherapy (using imatinib and more recently dasatinib and nilotinib) induces dramatic and long-lasting remissions [29-30]. The majority of patients who have received long-term treatments with BCR-ABL–KIs show strong leukemia control by cytogenetic evaluation [26, 29, 31-32], and similar results are observed in animal models [33]. However, almost invariably patients test positive for mRNA transcripts of BCR-ABL (approx 97%) [34], indicating that leukemia has not been completely eliminated [31]. The dormant leukemia cells which evade eradication by BCR-ABL–KIs in CML have been shown to express the characteristic surface markers of LSCs, and have been proposed to sustain drug resistant disease, in a fashion similar to HSCs [35-36] and independent of BCR-ABL kinase [37]. Discontinuation of BCR-ABL–KI treatment often results in clinical relapse and hematopoietic stem cell transplantation (HSCT) still remains important modality for treating those patients who have accelerated phase and blast crisis CML or those who either do not tolerate or fail/relapse during continuous BCR-ABL–KI treatments [38-41].

Outcomes of BCR-ABL Targeted Therapy in Ph+ ALL

In contrast to the robust cytogenetic and molecular responses in CML patients, BCR-ABL–KI monotherapy evokes much weaker and short lasting responses in patients with Ph+ ALL [42]. These differences are also appreciated in animal leukemia models [33, 43]. Despite receiving continuous BCR-ABL–KI therapy, human patients with Ph+ ALL typically carry significantly higher levels of persistent leukemia and inevitably relapse within a few months [42]. The relative primary drug-resistance in human Ph+ ALL necessitates the addition of several chemotherapeutic agents and HSCT in combination with BCR-ABL–KIs [44]. Multi-modality BCR-ABL–KI-combined regimens, the current standard of clinical care in Ph+ ALL, improve the remission rates but relapses remain common [44-47] with an 18- to 24-month survival rate of 64% and an even lower relapse-free survival rate [44].
The leukemia cells in Ph+ ALL which display poor initial response and evade eradication by BCR-ABL–KIs [45-47] express cell surface markers of lymphoid B cell lineage [48]. No distinct population of LSCs has been characterized to drive drug-resistant populations, unlike CML [35-36, 48].

**Deficiencies of Current BCR-ABL Inhibition Therapy**

**BCR-ABL Kinase Domain Mutations Mediate BCR-ABL–KI Drug Resistance**

The most frequently studied mechanism of drug-resistance against BCR-ABL-KIs is point mutations that occur within the KD of ABL (BCR-ABL$^{\text{MUTANTS}}$), which is common to both CML and Ph+ ALL [49-51]. Mechanistically, the mutations in BCR-ABL, especially in the KD, deteriorate the drug-binding and rescue the oncogenic kinase from drug-induced inhibition [49-51]. The total number of the distinct ABL KD mutations that have been characterized from refractory Ph+ leukemia samples now approaches 80 [53]. The spectrum of BCR-ABL mutations that each of the BCR-ABL-KIs select in Ph+ leukemias is distinct, but overlapping, with the notable exception of the T315I (Threonine to Isoleucine) mutation which confers profound cell-intrinsic resistance to all three approved BCR-ABL–KIs.

Rigorous research efforts are being made to design the next generation agents which can mitigate the effect of BCR-ABL mutations on drug-binding and thus achieve sustained BCR-ABL inhibition *in vivo*, as a critical step towards improvement in the clinical efficacy of BCR-ABL targeted therapy [17-18].

**Primary Drug Resistance against BCR-ABL–KIs Is Independent of BCR-ABL Mutations**

According to the long-term and metadata studies, drug-resistant BCR-ABL mutations are predominantly absent in: i) Ph+ ALL at the time of diagnosis which responds poorly to BCR-ABL–KIs [50-51, 54], and ii) persistent Ph+ leukemias (both CML and Ph+ ALL) which evade eradication during the first few months of targeted therapy [49-52]. Periodic testing reveals that as high as 50% of the Ph+ leukemia patients never test positive for drug resistant BCR-ABL mutations. Mutations in the ABL KD when detected for the first time are quantitatively present as a minority leukemia sub-population. Frequency and population size of mutations increases after prolonged period of treatment with BCR-ABL–KIs [50-52, 54]. On the other hand, ABL KD mutations can spontaneously regress in patients when the BCR-ABL–KI is changed, implicating BCR-ABL inhibition in the selective evolution of drug-resistant mutations.

Thus, primary clinical drug resistance against BCR-ABL–KIs occurs in the absence of well characterized ABL KD mutations, and likely plays an instrumental role in overall therapeutic failures [49-52].
Ph+ ALL Faithfully Captures the Deficiencies of Targeted Therapy

Approved BCR-ABL–KIs potently inhibit the KD of ABL protein in biochemical assays and in vivo [55] and commit Ph+ leukemia cells to death in vitro [56]. In stark contrast, due to unknown reasons, BCR-ABL–KIs fail to completely eliminate leukemia in patients with Ph+ ALL [42, 44] and CML [31]. While the drug refractoriness mediated by ABL KD mutations is common to both CML and Ph+ ALL, poor responses to BCR-ABL–KIs during early therapy, when patients carry predominantly non-mutated BCR-ABL (BCR-ABL\textsuperscript{WT}) leukemia, are clearly more evident in patients with Ph+ ALL compared to CML.

Thus, Ph+ ALL faithfully captures both the primary and target-acquired (ABL KD mutations) mechanisms of therapeutic failures against BCR-ABL–KIs. These features qualify Ph+ ALL as an ideal disease system to study and delineate the deficiencies of imatinib, nilotinib and dasatinib – the drugs which have played an instrumental role in qualifying the use of target focused cancer therapy in clinical settings.

Elaboration of the deficiencies of BCR-ABL–KIs may hold general lessons for further optimization and development of alternative targeted therapies in other cancers.

An Accurate Murine Model System of Human Ph+ ALL

In generating and validating an accurate model of human Ph+ ALL that recapitulates key features of the human disease, we previously demonstrated that p185\textsuperscript{BCR-ABL} expression potently synergized with inactivation of the \textit{Arf} tumor suppressor gene (\textit{Arf}\textsuperscript{−/−}); these two engineered genetic events are sufficient to fully transform murine pre-B cells into leukemia-initiating cells (LICs) [43]. The BCR-ABL oncogenic kinase potently ‘drives’ the proliferation of primary \textit{Arf}\textsuperscript{−/−} p185\textsuperscript{+} pre-B cells in a growth factor independent manner, while the loss of \textit{Arf} function dramatically attenuates oncogene-induced, p53-dependent apoptosis, confers cellular immortalization and facilitates exponential, high-viability growth. In this leukemia model system, the \textit{Arf}-loss dependent phenotype closely matches that associated with engineered loss of the p53 tumor suppressor, thus \textit{Arf} and p53 do essentially function in a ‘linear’ pathway [57-58]. The process of BCR-ABL mediated transformation of pre-B cells is dependent upon cooperation of transcription factors – signal transducer and activator of transcription (STAT5) [59] and c-MYC [60-62], a known oncogene. BCR-ABL fusion protein phosphorylates the STAT5 protein (pSTAT5) in the cytoplasm which results in its dimerization and transport to the nucleus where it binds to specific STAT5 response elements in the promoter region of target genes, including c-MYC.

More recently, this murine Ph+ ALL model was adapted for in vivo luminescent imaging to quantitatively monitor leukemic cell dissemination and therapeutic responses, using \textit{Arf}\textsuperscript{−/−} p185\textsuperscript{BCR-ABL} luciferase-expressing (Luc+) pre-B or LICs [57, 63]. As described in human Ph+ ALL clinical trials, single-agent dasatinib therapy in this murine
model induces transient remissions in recipient mice harboring very high leukemic burdens, only to be followed by clinical relapse a few weeks later (i.e. acquired dasatinib resistance) which is frequently associated with the emergence of BCR-ABL kinase domain mutations in cells recovered from leukemic-infiltrated tissues [57, 63]. However, dasatinib-refractory leukemic cells recovered from leukemic recipients at the end of ‘induction’ (i.e. at the nadir of leukemic burden, after maximal drug response had been achieved) infrequently harbored BCR-ABL kinase domain mutations [63], strongly implicating that BCR-ABL <sup>MUTANT</sup>-independent mechanisms initially attenuate dasatinib’s ability to fully eradicate this minimal residual disease (MRD).

**Etiology of Clinical BCR-ABL<sup>MUTANT</sup>-independent BCR-ABL-KI-resistance**

While rigorous studies has been overtaken to study BCR-ABL mutation mediated drug resistance [51, 53, 64-65], sparse efforts have been made to address and overcome the primary BCR-ABL-KI-resistance which is predominantly BCR-ABL<sup>MUTANT</sup>-independent. Primary BCR-ABL-KI-resistance can originate from factors that can broadly be classified as either leukemia-extrinsic or leukemia-intrinsic.

Among leukemia-extrinsic influences, tumor microenvironment [36, 66-67] and poor availability of drug in the target tissues can play a role. Though the tumor microenvironment can profoundly alter the drug-action against the tumor in vivo, its influence is conventionally not addressed during in vitro drug-evaluations [66]. Cell-extrinsic soluble factors such as GM-CSF and GM-CSF plus G-CSF in human CML, IL3 in myeloid and lymphoid systems, common gamma chain cytokine signaling and its family member IL7 in lymphoid pre-B cells, significantly attenuate BCR-ABL-KI induced leukemia cell killing [48, 62, 66, 68-69]. This suggests that signaling pathways activated by host growth factors can play a role in drug resistance in BCR-ABL driven leukemias in vivo. On the other hand, primary chemo-refractoriness may also arise from poor drug-availability in specific target tissues that harbor leukemic cells [70-71]. Both CML and Ph+ ALL are primarily disorders of hematopoietic tissues and share a majority of the sites of spread, namely, bone marrow, lymph nodes, spleen, liver and thymus [26]. Unfavorable pharmacokinetic properties, which are inherent to the chemical nature of drug, are thus less likely to play an important role in the varied responses to BCR-ABL-KI agents in Ph+ ALL and CML.

Cell-intrinsic drug resistance mechanisms that function independent of BCR-ABL mutations include genetic factors, epigenetic changes or altered dynamics of drug import and drug efflux that modulate intracellular inhibitor concentrations. Studies by others have established that genetic deletions involving the tumor suppressor CDKN2A/B [72-74], lymphoid transcriptional regulator IKAROS (also called IKZF1) [75-76], PAX5 [73], EBF1 [73], cooperate with BCR-ABL in leukemogenesis. These genetic events, which are involved in the pathological development of disease, are present at the time of diagnosis in a majority of the patients with Ph+ ALL [74-76] but not CML patients. They
may also play a role in poor primary responses to BCR-ABL–KIs in Ph+ ALL, but this hypothesis remains to be ascertained.
CHAPTER 2. HYPOTHESIS AND SPECIFIC AIMS

Problem at Hand

Elucidation of the mechanisms of resistance against clinically used drugs, especially the new era target-selective agents, is integral to improve their efficacy and achieve long-term survival in cancer patients. Significantly, primary drug resistance against BCR-ABL–KIs in human Ph+ ALL very well represents the clinical dilemma shared by a majority of current drugs that selectively target and kill cancer cells in vitro but not in vivo. This disparity strongly implicates the modulation of therapeutic response of tumor cells within the host microenvironment.

In contrast to the remarkable susceptibility of BCR-ABL WT cells to BCR-ABL-KIs in vitro, patients suffering from Ph+ ALL receive poor and transient benefit from BCR-ABL–KI therapy despite the predominant absence of ABL-KD mutations at diagnosis and during initial few months of therapy. The poor primary responses evoked by BCR-ABL–KI monotherapy necessitates clinical management of Ph+ ALL patients with a multi-modality approach, which is often complicated by the frequent unavailability of HSCT donors and the toxicity/intolerability of conventional chemotherapy and HSCT, especially in the elderly who form majority of the Ph+ ALL population. The performance of BCR-ABL–KIs deteriorates even further later during the therapy when ABL KD mutations appear in leukemia lymphoblasts, imparting molecular drug resistance.

While wide scientific attention is paid to improve the drug design for tackling molecular drug resistance, it appears clinically more relevant to mechanistically understand why BCR-ABL WT Ph+ ALL responds poorly and evades eradication by BCR-ABL–KIs prior to the development of molecular resistance mediated by KD mutations. The unforeseen primary (BCR-ABL mutation-independent) resistance is most likely imparted by the tumor microenvironment (leukemia-extrinsic) and/or alternatively by changes within the tumor itself (leukemia-intrinsic). We previously reported a role for host cytokines of the common gamma chain receptor family in mediating primary imatinib resistance [48], but many alternative host-derived growth factors have similarly been proposed in BCR-ABL–KI drug resistance [62, 66-69]. Currently, no consensus exists for the relative importance of cytokines that control the cellular responses to BCR-ABL targeted agents.

Goal of the Study

The central goal of this scientific work was to advance the performance of BCR-ABL targeted therapy in Ph+ ALL – a human leukemia that currently has poor long-term survival outcomes.
Hypothesis

Defining and countering the factors that confer the poor response to BCR-ABL–KIs in patients with non-mutated BCR-ABL (BCR-ABL\textsuperscript{WT}) Ph+ ALL is critical to harness the full therapeutic potential of targeting BCR-ABL for improving the long-term survival. We hypothesized that – ‘Combination therapy with drugs that are active against the Ph+ ALL phenotypes displaying mechanisms of primary drug resistance can eradicate leukemias that are refractory to BCR-ABL–KIs \textit{in vivo}.’

Specific Aims of the Study

To maximize the likelihood of discovering combination agents that may overcome multi-faceted BCR-ABL–KI drug resistance in Ph+ ALL \textit{in vivo}, we aimed to:

i. Create a rank-order of leukemia microenvironment-based host-cytokine/s for imparting primary drug resistance (cell-extrinsic) against current FDA-approved BCR-ABL–KI agents \textit{in vitro}.

ii. Test the most relevant host-cytokine/s from (i) for conferring BCR-ABL–KI-refractory phenotype \textit{in vivo} and characterize its associated cell-intrinsic changes.

iii. Screen drug-rich chemical libraries to identify agents that possess \textit{in vitro} activity against Ph+ ALL cells protected by cell-extrinsic host-cytokine(s) (from i and ii) and previously known molecular mechanisms (T315I) of BCR-ABL–KI drug-resistance.

iv. Test agents (from iii) alone and in combination with dasatinib in our murine model of Ph+ ALL. Assess agent’s ability to improve the primary responses to dasatinib and eradicate dasatinib refractory persistent disease, as a critical step towards their evaluation in patients with Ph+ ALL clinics.

Experimental Methodology

Here, we report efforts to establish a high throughput (HT) cell-based platform that captures the dominant cell – extrinsic (host environment imparted) and intrinsic (BCR-ABL mutation-dependent) mechanisms of BCR-ABL–KI-refractoriness in Ph+ ALL, in order to discover therapeutic agents capable of improving \textit{in vivo} BCR-ABL–KI performance.

Utilizing Arf-null (Arf\textsuperscript{−/−}) BCR-ABL transformed LICs, which model key biological and genetic features of human Ph+ ALL, we developed an assay that evaluates drug performance against BCR-ABL\textsuperscript{WT} LICs co-cultured with candidate cell-extrinsic cytokines and growth factors.
Following an unbiased screening approach, we identified IL7 as a specific host-cytokine that conferred primary chemo-refractoriness against all FDA-approved BCR-ABL–KIs in vitro, and subsequently confirmed these clinically-relevant results in vivo. Next, we assayed this drug-resistant phenotype against a library of FDA-approved drugs, natural products, and chemical tool compounds. In addition to the identification of many clinical anti-leukemic and other chemotherapeutic agents, several novel agents overcame the BCR-ABL–KI-resistant phenotype. Testing of combinations of dasatinib with one of our most compelling screening hits, the FDA-approved antimalarial drug dihydroartemisinin (DHA), eradicated the residual pool of dasatinib-refractory leukemia in vivo. This treatment protocol durably cured 90% of treated animals. This opens up a promising therapeutic avenue for treating human Ph+ ALL patients by adding DHA to current dasatinib regimens, at the time of diagnosis.
CHAPTER 3. MATERIALS AND METHODS

Production and Culture of Murine Leukemia-initiating Cells (LICs)

Replication-defective mouse stem cell virus (MSCV) retroviral vectors coexpressing either wild-type (WT) allele of human p185BCR-ABL or mutant BCR-ABL alleles, namely p185T315I or p185F317L, and either green fluorescent protein (GFP) or luciferase, were packaged into replication-incompetent ecotropic virions [43, 48, 63]. Whole bone marrow cell suspensions from Arf−/− young adult mice were transduced to produce Arf−/− p185+ pre-B cells [43, 63]. After initial establishment on autologous stromal cell layers over 7 days, transformed pre-B cells were briefly expanded in the absence of a stromal layer for 2 days in liquid culture in BCM10 media (RPMI1640 supplemented with 10% HyClone fetal calf serum, 4 mM glutamine, 100 Units/mL penicillin, 100 μg/mL streptomycin, and 55 μM beta-mercaptoethanol) [48], and cryopreserved as 40 identical vials per cell line, generating a master cell bank. Arf−/− p185WT and Arf−/− p185T315I pre-B cells (BCR-ABLWT LICs and BCR-ABL T315I LICs, respectively) from the cell bank were thawed and allowed to recover in BCM10 for 3 days, prior to their use in any in vitro or in vivo assays.

MSCV-IRES-GFP retroviral vectors expressing human c-Myc or empty ‘control’, and MSCV-IRES-GFP or MSCV-IRES-RFP retroviral vectors expressing previously-validated mutants of IKAROS (either IK6 or IK1) [75] or constitutively active (CA)-STAT5 mutants [77], were utilized to transduce BCR-ABL-expressing LICs, and cultured for 48 in BCM10 media before use in drug competition assays.

In Vivo Adoptive Leukemia Transfer Model of Ph+ ALL

Mice were housed in an American Association of Laboratory Animal Care (AALAC)-accredited facility and treated on Institutional Animal Care and Use Committee (IACUC)-approved protocols in accordance with NIH guidelines. LICs were injected into tail veins of healthy, non-conditioned, immune-competent 10- to 12-week-old IL7+/+ or IL7−/− C57Bl/6J mice (Jackson Labs, Bar Harbor, ME). Animals were observed daily and were sacrificed when moribund (dehydration, ruffled fur, poor mobility, respiratory distress). Survival curves were generated using GraphPad Prism Version 5.0 (La Jolla, CA). Mantel-Cox test was applied to pair-wise comparisons of survival data.

FACS Assessment of Viability

Viability of LICs was estimated after counterstaining approximately 5 x 10^5 LICs (in 0.5 mL volume) with 35 μL of a propidium iodide (PI) solution (0.25 mg/mL in PBS). The percentage of viable cells was calculated by electronic gating on PI-negative cells on PI versus forward scatter dot plots, with forward scatter signals consistent with LICs, and
comparison of this number with total cell number. Data were collected and analyzed on a BD Biosciences LSR II flow cytometer (San Jose, CA), using BD Biosciences FACS DiVa software.

**FACS Assessment of Cell Cycle Percentages**

For determination of percentages in cell cycle phases, 5 x 10^5 LICs were centrifuged, washed once with PBS, recentrifuged, and cell pellets were resuspended in 0.5 mL PI staining solution (0.05 mg/mL PI, 0.1% sodium citrate, 0.1% Triton X100), which lysed the cells and stained nuclear DNA with PI. Samples were then treated with DNase-free RNase (0.2 mg/mL in 10 M Tris-HCl/15 mM NaCl, pH 7.5) for 30 min at room temperature, filtered, and analyzed for red fluorescence from a PI-labeled DNA on BD Biosciences FACS Calibur flow cytometer (San Jose, CA) by using BD Biosciences CellQuest Pro software. The percentages of cells within each cell cycle phase were computed using the computer program ModFit (Verity Software House, Topsham, ME).

**Development of LIC-based Multimodality Screening Assay**

Before use in any assay, LICs from the master cell bank had to pass the following quality control parameters: test thawing (benchmark: 95% viability at 24 h); PCR genotyping to confirm Arf^-/- status [78]; mycoplasma testing (negative by Takara Bio Mycoplasma PCR assay, Clontech, Madison WI); sequence verification of BCR-ABL allele [79]; and in vitro dasatinib sensitivity.

Maximum DMSO concentration tolerated by LICs in culture was determined to be 0.2% (by volume) and was always maintained ≤0.1 % (by volume). LICs plated in 384-well plates at a cellular density of 5 x 10^4 per mL in BCM10 were confirmed to give exponential and reproducible growth during the first 72 h (Figure 3.1). To select assay reagents, 3 high throughput screening (HTS) relevant cellular assays were evaluated according to manufacturer’s instructions (Figure 3.2), and flow cytometry analysis was performed in parallel (as a reference control) using a high-throughput sampling device (Hypercyte by Intellicyte Corporation, Albuquerque, NM) and BD Biosciences LSR II flow cytometer (San Jose, CA). Relative to accuracy of high throughput flow cytometry (HT-flow) to detect low number of viable LICs, especially at high drug concentrations, the luminescent CellTiter-Glo (CTG)-assay by Promega Madison, WI, outperformed a common spectrophotometry-based methane-thiosulfonate-based (MTS) reagent (CellTiter 96® AQeuous One Solution Cell Proliferation assay; Promega Madison, WI), and alamar blue (not shown). Since CTG-assay monitors LIC proliferation by using total ATP content as a surrogate marker, it inherently cannot distinguish between genetically distinct sub-populations within the same well. To adapt the evaluation of cell-intrinsic genetic factors into our drug screening platform, cell-mixtures of Arf^-/- BCR-ABL^WT-RFP LICs and Arf^-/- BCR-ABL^T315I-GFP LICs were treated with dasatinib (Figure 3.3) and HT-flow assay performed after 72 hr drug exposures. HT-flow (drug-challenge competition assay) correctly detected the expected dasatinib-induced selective
Figure 3.1. Comparison of the growth properties of \(Arf^{+/−}\) p185\(^{WT}\) (filled symbols) and \(Arf^{-/-}\) p185\(^{T315I}\) pre-B (empty symbols) populations during \textit{in vitro} cell culture.

Both cell types were plated at multiple (eight) cell dilutions (\(n=3\) for each condition, time=0 h) in 6-well plates and serially followed daily for 5 days to evaluate absolute viable cell count, total viability, doubling times, and total population doublings, using the Beckman Coulter cell counter (trypan blue staining) and fluorescence-activated cell sorting (FACS) analysis in parallel. Both cell types had identical growth properties. Only the comparison for population doubling time in hours (\(+/-\ sd\)) for 3 relevant plating densities is presented. FACS cell-cycle analysis confirmed that a population doubling time less than 14 h was sufficient to maintain cells in exponential growth (>50% cells in S-phase).
Figure 3.2. Optimization of the high-throughput LIC-based phenotypic assay.

Arf\(^{-/-}\) BCR-ABL\(^{WT}\) LICs plated at 5 \times 10^4 per mL were challenged with 6 log-fold concentration range of dasatinib or DMSO in 384-well microplates. After a 72 h drug challenge, LIC growth was measured by 3 cellular readouts: CellTiter-Glo (CTG)-assay (ATP content), MTS assay (viability), and flow cytometry (total viability and viable cell counts by DAPI exclusion) in high-throughput (HT-flow) mode. Measured assay readouts at various drug concentrations (x-axis) were normalized to DMSO-treated LICs and graphed (+/− sd) on y-axis. Total viable LIC counts were determined by FACS, reflecting dasatinib-induced phenotypic changes across a 6 log-fold drug-concentration, were closely paralleled by the CTG cellular assay output but not by the commonly used MTS assay.
Figure 3.3. Optimization of HT-flow output to perform drug competition assays.

(a) *Arf*⁻/⁻ BCR-ABL<sup>WT</sup> LICs and *Arf*⁻/⁻ BCR-ABL<sup>T315I</sup> LICs were mixed in 1:1 ratio at time *t*=0 hrs and treated with dasatinib under optimized assay conditions. Schematic of BCR-ABL alleles in LICs is presented. (b) Schematic of set-up of flow cytometry analysis at *t*=72 hrs. (c) Change in ratio of BCR-ABL<sup>WT</sup> LICs to BCR-ABL<sup>T315I</sup> LICs in response to increasing drug concentration at *t*=72 hrs. Real experimental data are plotted for dasatinib (n=3) and a hypothetical drug X that is T315I selective is depicted for comparison.
enrichment of Arf−/− BCR-ABL^T315I^ LICs within the same well. Thus, HT-flow assay complements the CTG-assay readout by quantifying the impact of ‘engineered’ cell-intrinsic change towards drug refractoriness within the same well. Of the two, CTG-assay is easy to use in HTS process (add and read one-step assay with no prolonged incubation step required between reagent addition and assay readout), has 10000 fold dynamic range, measures IC_{50} shifts of ≤ 2 fold for dasatinib, displays minimal drift or edge effects and robust inter-plate and inter-day reproducibility.

**Forward Cytokine Phenotypic Screen and Other *In Vitro* Cellular Assays**

Lyophilized murine and human recombinant cytokines (R&D Systems, Minneapolis, MN) were diluted with 1 mg/mL BSA in sterile PBS to a final concentration 20 μg/mL and stored at 4°C. Anti-leukemia drugs, BCR-ABL–KIs and non-BCR-ABL-specific compounds (LC labs, MA and Sigma, MO), were solubilized in DMSO to obtain 10 mM stocks concentrations. Subsequent serial titrations were prepared using DMSO in 0.5 mL vials or 384-well drug master plates, and stored at –20°C.

To study the impact of murine cytokines on dasatinib action, LIC stocks containing different concentrations of candidate cytokines were prepared in multi-channel sterile reservoirs and plated into 384-well plates manually using handheld Finnpipette® Novus (Thermo Scientific, Waltham, MA) electronic 16 channel pipette. Drug delivery to cell suspensions with a V & P Scientific (San Diego, CA) pin tool and all subsequent steps of the CTG-assay were performed under optimized automated conditions, as described in Table 3.1. In a forward candidate approach, evaluation of IL7’s ability to attenuate drug-induced cell killing and growth inhibition was later extended to all BCR-ABL–KIs and non-BCR-ABL specific drugs using the CTG-assay in semi-automated mode (non-automated cell-plating as described above).

Human SUP-B15 Ph+ ALL cells (DSMZ cell culture collection, Germany) in McCoys media containing 20% fetal calf serum and penicillin/streptomycin (Invitrogen, Carlsbad, CA), were plated at 12,500 cells in 25 μL per well in black clear-bottom 384-well microplates. Drugs were transferred by pin-transfer. After 72 h incubation at 37°C, 20% (by vol) MTS assay reagent was added. Microplates were incubated for 75 min at 37°C, and read colorimetrically at 490 nm.

**Drug/Cytokine Washout and Cytokine Recovery Studies**

For washout studies, BCR-ABL^WT^ LICs were diluted to a density of 1 × 10^5 LICs/mL in BCM10 containing no or 5 mg/mL IL7, and treatment with either dasatinib or 0.1% DMSO by volume (non-drug treated controls) was initiated at time 0 h in triplicate 100 mm petri dishes. Daily monitoring for expected drug-induced changes was performed by naked-eye microscopy from 0 to 72 h. At the 72 h time point, after harvesting under identical conditions, total LIC density and viability percentage for all samples were determined using Beckman Coulter Vi-cell (trypan blue) in triplicates.
Table 3.1. Steps in optimized high-throughput screening assay (CTG readout).

<table>
<thead>
<tr>
<th>Step</th>
<th>Parameter</th>
<th>Value</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Plate cells</td>
<td>25 μL</td>
<td>Plate LICs 5 x 10⁴ per mL (25 μL/well) in BCM10 + 0.85 ng/mL IL7 into 384-well opaque bottom white microplates using Wellmate (Matrix)</td>
</tr>
</tbody>
</table>
| 2    | Primary drug screening controls in columns 1-2 and 13-14 | 25 nL | 1A-1I: Nine dasatinib concentrations (100 nM, 30 nM, 10 nM, 3 nM, 1 nM, 0.3 nM, 0.1 nM, 0.03 nM and 0.01 nM) – control for IL7- and BCR-ABL mutation-imparted resistance against BCR-ABL–KIs  
2A-2I: Nine staurosporine concentrations (300 nM, 100 nM, 60 nM, 30 nM, 10 nM, 6 nM, 3 nM, 1 nM, 0.6 nM) – non-BCR-ABL-specific reference control in the presence of IL7  
13A-13P, 1J, 2J: DMSO 0.1% (Max Signal OR Negative Control); 14A-14P: staurosporine at 300 nM (Min Signal OR Positive Control); 1K-1P, 2K-2P: 40 μM dasatinib  |
|      |                                               |       | 23A-23J: Staurosporine – 10 log-fold dilutions (3 μM to 0.3 pM) – non-BCR-ABL-specific reference control in the presence of IL7  
24A-24J: Dasatinib – 10 log-fold dilutions (1 μM to 0.1 pM) – control for IL7- and BCR-ABL mutation-imparted BCR-ABL–KI drug-resistance                                                                 |
Table 3.1. Continued.

<table>
<thead>
<tr>
<th>Step</th>
<th>Parameter</th>
<th>Value</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>Add test compounds in columns 3-12 and 15-24 in primary; columns 1-20 in secondary screen</td>
<td>25 nL</td>
<td><strong>Primary screening:</strong> Single final drug concentration of 10 μM <strong>Secondary screening:</strong> Triplicate ten half-log serial dilutions (5 μM to 2 nM final concentration) of primary screening hits to determine their half-maximal inhibitory concentrations (IC₅₀) against LICs. 384-well master drug-stock plates prepared at 1000X in DMSO. Drug delivered to assay plates by pin transfer (using V&amp;P Scientific pin tool, 10H pins), giving a final drug concentration of 1× and 0.1% DMSO in all control and test wells of assay plates.</td>
</tr>
<tr>
<td>4</td>
<td>Incubation 1</td>
<td>72 h</td>
<td>Liconics incubator, 8%CO₂, 37°C (standard LIC culture conditions)</td>
</tr>
<tr>
<td>5</td>
<td>Incubation 2</td>
<td>20 min</td>
<td>Room temp, low light (instructions as per Promega)</td>
</tr>
<tr>
<td>6</td>
<td>Add reagent</td>
<td>25 μL</td>
<td>CellTiter Glo (Promega) pre-equilibrated to RT before use</td>
</tr>
<tr>
<td>7</td>
<td>Incubation 3</td>
<td>25 min</td>
<td>Room temp, low light (per Promega)</td>
</tr>
<tr>
<td>8</td>
<td>Assay readout</td>
<td>ATP levels</td>
<td>Envision, Luminescent mode – ATP levels as a measure of LIC number and viable growth in 72 h – <strong>Signal</strong> measured in RLUs</td>
</tr>
<tr>
<td>9</td>
<td>Data processing</td>
<td>Pipeline Pilot</td>
<td>% inhibition by test compound = 100 * (Log₁₀ Test Signal mean – Log₁₀ Min Signal mean)/(Log₁₀ Min Signal mean – Log₁₀ Max Signal mean) [Z′ = 1−\frac{(3\times\text{stdev of Log₁₀ Max Signal} + 3\times\text{stdev of Log₁₀ Min Signal})/\text{(Log₁₀ Max Signal mean – Log₁₀ Min Signal mean)}}{\text{(Log₁₀ Max Signal mean – Log₁₀ Min Signal mean)}}] [Z′] is a dimensionless metric to assess assay quality – for a good assay (Z′\geq0.5), and for a perfect assay (Z′=1)</td>
</tr>
</tbody>
</table>
Each sample was spun down in 50 mL BD Falcon™ tubes; the supernatant was discarded and LIC pellet resuspended in freshly prepared BCM10 (not containing any cytokine or drug). This washing step was repeated 4 times to ensure complete removal of the drug and/or cytokine. After the last wash, each pellet was resuspended to $1 \times 10^6$ total LICs per mL master stocks. Aliquots from master stocks were taken for (i) flow assessment of viability and cell cycle percentages (as described above); (ii) manual CTG-assay using 25 μL sample per well in 384-well microtiter plates (in triplicates), as an independent determination for LIC growth. For assessment of extracellular cytokine on recovery of LICs that were previously challenged with dasatinib (0 to 72 h), washed master stocks were diluted to $5 \times 10^5$ total LICs per mL in BCM10 with or without 5 ng/mL IL7, plated in 6 well plates (10 cm² approx.), and incubated at 37°C from 72 to 144 h for recovery, during which time daily naked-eye microscopic examinations were performed. At 144 h, for each sample total LIC density and viability percentage assessment by Beckman, viability and cell cycle percentages by flow assessment and independent CTG evaluations were performed as described for the 72 h time point. Trends in cell were also measured by CTG readout and used to independently corroborate the findings from trypan blue (Beckman) and DAPI staining (FACS). Total viable LICs in a sample were calculated as a product of absolute LIC count determined by Beckman and absolute sample viability determined by FACS analysis.

**Dasatinib Re-challenge Studies**

Further studies were performed to evaluate the possibility of a genetically drug-resistant subpopulation that could undergo selection with dasatinib/IL7 co-treatment during 0-72 h of dasatinib/cytokine washout studies above. To this aim, BCR-ABL<sup>WT</sup> LICs were treated with 100 nM dasatinib or 0.1% DMSO in the absence or presence of 5 ng/mL IL7 in 100 mm dishes from 0-72 h, as described above; in addition, a dasatinib challenge (first-time challenge) across 6 log-fold concentration in the presence or absence of 5 ng/mL IL7 was also performed using CTG-assay during this time period (0-72 h). At 72 h, dasatinib/IL7 co-treated LICs from 100 mm dishes were washed (as described above) and immediately challenged (second challenge or re-challenge) with six log-fold concentration range of dasatinib in 384-well microtiter plates either in the presence or absence of 5 ng/mL IL7 from 72 to 144 h. At 144 h, CTG-assay was performed.

**High-throughput (HT) Phenotypic Drug Screening**

Minimal concentration of IL7 that conferred near-maximal drug resistance against dasatinib, nilotinib and imatinib in BCR-ABL<sup>WT</sup> LICs was first determined by performing cytokine titration assays (0.85 ng/mL IL7) under semi-automated mode and later confirmed to reproduce results under fully-automated HT screening conditions (Table 3.1). Primary single-point (10 μM) HT evaluation of 5600 compounds (3200 unique) and subsequent secondary drug screens across 10 two-fold drug dilutions were performed. In addition to BCR-ABL<sup>WT</sup> and BCR-ABL<sup>T315I</sup> LICs, secondary drug-screens were also performed against BCR-ABL<sup>T317L</sup> LICs, another clinically-known dasatinib-
resistant ABL KD mutation which gave identical results to that for BCR-ABL^{T315I} (not shown). To track the quality of screening experiments, multiple control assay plates were set-up during all screening runs to allow for inter-plate and inter-day comparisons at 0, 24, 48 and 72 hrs.

**Description of Chemical Library Screened**

The screening library consisted of 5600 (approximately 3200 unique) approved drugs and chemicals with known biological activity (bioactives). The library was assembled from 3 commercial suppliers: Microsource, Prestwick, and Sigma. The Microsource compounds included the following: (a) the Spectrum collection, which contains 2000 biologically active and structurally diverse compounds, including known drugs, experimental bioactives, and pure natural products [80-81]; (b) the US Drug Collection, which contains 1040 drugs that have reached clinical trials in the US and have been assigned USAN or US Pharmacopeia status; and (c) the Killer collection, which contains a reference set of 160 synthetic and natural toxic substances (http://www.msdiscovery.com/index.html). The Prestwick compounds include 1120 small molecules selected for high chemical and pharmacologic diversity. Ninety percent of the collection is composed of known marketed drugs, and the remainder includes bioactive alkaloids or related substances. Human bioavailability and human toxicity data are available for most compounds (http://www.prestwickchemical.fr/index.php?pa=26). The Sigma LOPAC1280 (Library of Pharmacologically Active Compounds) collection reflects the most commonly screened targets in the drug discovery community, including marketed drugs, failed development candidates, and "gold standards" that have well-characterized activities (http://www.sigmaaldrich.com/chemistry/drug-discovery/validation-libraries.html).

**Data Processing, Quality Control and Hit Scoring Criteria for HT Drug Screens**

All data processing and visualization was performed using custom programs written in the Pipeline Pilot platform (Accelrys, v.7.5) and the R program 6.6 [82-83]. The R drc package was used to fit sigmoidal curves [83]. ROC statistics were computed using the ROCR package [84].

The quality of the drug-screening studies against BCR-ABL^{WT} and BCR-ABL^{T315I} LICs was evaluated by multiple methods including, but not limited to, the identification of known antileukemic agents, identification of multiple compound replicates intentionally included in the 5600 collection, evaluation of control assay plates, and Z' (see Table 3.1) and other screening quality metrics. In the screening experiments, all assay plates demonstrated attenuation of dasatinib potency in the presence of IL7 or BCR-ABL mutation. An empirically determined, statistically significant but liberal cut-off of >10% activity was chosen to include agents of relatively lower activity considering the high-risk nature of Ph+ ALL, the drug-rich nature of the library as well as
to allow for a more complete structure-activity relationship and therapeutic class analysis after subsequent potency determination through secondary screening.

The discriminatory power of the phenotypic assay was assessed using receiver operating characteristic (ROC) statistics. A total of 165 compounds were selected to sample the primary assay according to the distribution of observed activities. ROC curves were generated and area under curves (AUCs) calculated to determine the discriminatory power of the assay for BCR-ABL<sup>WT</sup> and BCR-ABL<sup>T315I</sup> LICs, respectively.

The activity class designations were assigned based on the curve score and visual inspection of dose-response curves (see supplemental data in Appendix). The curve score is a 4 character code derived from the following heuristic:

- First character: letter (A,B,C or D) indicating curve quality
  - D = less than 2 points above the noise (defined as outliers from the negative control population)
  - C = \( r^2 < 0.50 \) or range \( > 200\% \) or \( EC_{50} > \) maximum concentration tested
  - A = \( r^2 > 0.80 \) and difference between two last points \( < 10\% \) (indicating saturation)
  - B = not A,C, or D
  - N prefix = activity decreases with increase concentration
- Second character: number (1-6) indicating potency
  - 1 = \( EC_{50} < 0.1 \) \( \mu \)M
  - 2 = \( 0.1 \) \( \mu \)M \( \geq \) \( EC_{50} < 1.0 \) \( \mu \)M
  - 3 = \( 1.0 \) \( \mu \)M \( \geq \) \( EC_{50} < 10.0 \) \( \mu \)M
  - 4 = \( EC_{50} > 10.0 \) \( \mu \)M
  - 5 = all C and D curves are assigned 5
- Third character: number (1-6) indicating efficacy
  - 1 = 75\% \( \geq \) range \( < 200\% \)
  - 2 = 50\% \( \geq \) range \( < 75\% \)
  - 3 = 25\% \( \geq \) range \( < 50 \)
  - 4 = range \( < 25\% \)
  - 5 = all C and D curves are assigned 5
- Fourth character: number (1-6) indicating hill slope
  - 1 = hill \( \leq 2 \)
  - 2 = 2 \( < \) hill \( \leq 4 \)
  - 3 = 4 \( < \) hill \( \leq 8 \)
  - 4 = hill \( > 8 \)
  - 5 = all C and D curves are assigned 5

A complete list of all compounds evaluated in the secondary drug-screening experiments with their structure, supplier information, synonyms, \( EC_{50} \) values, and therapeutic classification is reported in the supplemental data in Appendix.
**Cluster Analysis of Hits Using Therapeutic Classes**

The therapeutic cluster graph was generated by first assigning each compound an Anatomical Therapeutic Chemical Classification System (ATC). This classification system groups drugs into a 5 level hierarchy according to therapeutic indication and chemical properties. The first level indicates anatomical main group and consists of 1 letter; the second level indicates therapeutic main group and consists of 2 digits; the third level indicates therapeutic or pharmacological subgroup and consists of 1 letter; the fourth level indicates chemical, therapeutic or pharmacological subgroup and consists of one letter; and the fifth level indicates the chemical substance and consists of 2 digits. The resulting 7 characters constitute an ATC code, and a compound can be assigned multiple ATC codes. ATC codes can be depicted as a network graph by defining a node to represent each level in the ATC code and then connecting nodes according to parent-child relationships (e.g., first-level node A is the parent of second level nodes A01, A02, etc.). Compounds without an ATC code were assigned based on similarity to known agents, or were assigned to custom codes located under the parent V99 (labeled as “Not defined in ATC”). The frequency of ATC Level 1 and Level 2 codes among the 706 compounds submitted to dose-response analysis was determined.

The resulting network was visualized in Cytoscape (v 2.8.1) using the yFiles circular layout algorithm [85].

**Evaluation of Hits for Tolerance in Human Patients and Selectivity for MYC Gene Expression in Their Mechanism of Drug Action Using Public Databases**

Searches in Pubmed and MetaDrug™ software (by Genego, CA) [86] were performed for validated phenotypic hits to document approved use/uses in human disease/s if any, and known/predictive toxicity data in human and animal models, especially lymphotoxicity. Hits with favorable toxicity profiles were prioritized using a previously described statistical analysis [87-88], similar to the COMPARE algorithm [89-91], that evaluates National Cancer Institute (NCI) anti-cancer drug screening data. Briefly, the publically available databases containing half-maximal growth-inhibitory concentrations (GI50) of approximately 4444 common chemicals and FDA-approved drugs against NCI60 panel of 60 human cancer cell-lines (of 9 different tissues of origin) and complete gene expression datasets of all NCI60 cell lines, were accessed from the developmental Therapeutics Program (DTP) website (http://www.dtp.nci.nih.gov) and (http://discover.nci.nih.gov/cellminer/), respectively. A correlation analysis was performed between GI50 values and c-MYC expression across NCI60 cell lines for i) phenotypic hits demonstrating significant potency in a secondary screen, (whenever available in the DTP database, using NSC number as chemical identifier), ii) dasatinib, as a control for BCR-ABL–KI drug family, and iii) 4444 commonly used non-BCR-ABL–KI DTP agents (to achieve statistical power and compute p values).

To extend the mechanistic understanding on DHA drug action against human cancer cells, a subset of DHA-sensitive NCI60 cancer cell lines was identified. Using
complete affymetrix data set from Cellminer™ website, the expression of each gene across the DHA-sensitive cell lines was subjected to correlative analyses with paired GI50 values of DHA. Pearson coefficient of correlation computed for individual genes was used to generate a rank order for all genes. A probeset of top 40 positively correlated genes was selected, representing a distinctive drug-sensitivity signature for DHA across human cancer cell lines. The expression level of this gene-set was then compared between murine LICs and primary Ph+ ALL leukemic samples, obtained from seven patients at the time of diagnosis (GSE17459, Pubmed). Cross-species correlative analysis was performed for each of the 40 genes by both expression values and rank order.

Gene expression signature of IL7-induced dasatinib resistance in LICs was compared with DHA sensitivity signature across NCI60 panel to select overlapping genes which plausibly may play a role in overcoming dasatinib resistance.

**In Vivo Preclinical Therapeutic Studies**

For *in vivo* use, dasatinib (LC Labs, Woburn, MA) in citric acid (pH 3.1) [92], and DHA (Avachem, San Antonio, TX) in 0.5% carboxy-methylcellulose/0.5% Tween 80/0.5% benzyl alcohol, were administered by oral gavage. In toxicity studies (data not shown), DHA up to 300 mg/kg given as a single or split dose for ≤ 5 days/week over a six week time period induced no significant weight loss, lethargy, seizures or deaths corroborating published findings [93]. During one or two agent therapeutic studies, animal weights were monitored daily (5 days/week) and drug dosing was adjusted accordingly.

**Bioluminescent Imaging to Track Therapeutic Responses In Vivo**

Bioluminescent imaging and analysis was performed using a Xenogen IVIS-200 system and Living Image software 3.01 (Caliper Life Sciences, Hopkinton, MA), as described previously [63]. Total bioluminescent flux measurements (photons/second) were quantified over the whole animal body. Recipient mice that remained clinically healthy 12 months after terminating therapy and had undetectable bioluminescent signal *in vivo* were designated “long-term survivors.”

**Analysis of BCR-ABL KD Mutations**

Genomic DNA was extracted with phenol-chloroform from cultured cells, hematopoietic tissues, and leukemic-infiltrated meninges from moribund mice during therapy. Polymerase chain reactions (PCR) to selectively amplify only the human (and not murine) ABL KD were performed using primers as described previously [63, 79]. Sequence analysis was performed from 10 to 20 independent bacterial colonies, while identified mutations were considered significant if present at ≥ 20% occurrence in individually sequenced clones.
CHAPTER 4. RESULTS

A BCR-ABL Kinase Domain Mutation Does Not Impart Any Advantage in Leukemia Progression In Vitro or In Vivo

First, to model the clinically most dominant BCR-ABL mutation-dependent (molecular) resistance against all BCR-ABL–KI drugs in Ph+ ALL, we generated Arf–/– pre-B cells driven with p185BCR-ABLWT or p185BCR-ABLT315I mutant allele, referred as Arf–/– BCR-ABLWT LICs and Arf–/– BCR-ABL T315I LICs, respectively. Both LICs performed similarly during growth assays in vitro (Figure 3.1) and displayed a potent and highly comparable leukemia-initiating capacity in vivo when injected in serial limiting cellular dilutions (Figure 4.1).

Thus, T315I BCR-ABL mutation, which imparts remarkable cell-intrinsic molecular drug-resistance, has no impact on leukemia progression in vivo in the absence of BCR-ABL–KI-treatment.

Cytokine Screen Identifies IL7 as the Main Host Factor That Mediates Primary Dasatinib Resistance, Independent of BCR-ABL Mutations

To address the role of cell extrinsic mechanisms in modulating primary responses to BCR-ABL inhibitory therapy in Ph+ ALL, fifteen cytokines normally present in the leukemia microenvironment were evaluated for their ability to protect murine Arf–/– BCR-ABLWT LICs from dasatinib, the most potent clinical BCR-ABL inhibiting agent (Figure 4.2a). Arf–/– BCR-ABLWT LICs were confirmed beforehand to not harbor any drug-resistant mutations in the ABL KD. It is important to note that genetically-defined Arf–/– BCR-ABLWT LICs are transformed primary pre-B cells that retain phenotypic and signaling properties essential for suspended leukemic blasts to respond to the surrounding host-environment [48]. These properties are typically not present in established leukemic cell lines that have been extensively cultured in vitro.

Among the cytokines tested, low-level (< 1 ng/mL) IL7 provided the most potent and robust protection against dasatinib-induced growth arrest and killing of Arf–/– BCR-ABLWT LICs (Figure 4.2a). In addition, Interleukin3 (IL3) [62, 69, 94], thymic stromal lymphopoietin (TSLP, an IL7-like cytokine that shares a receptor subunit with IL7) [95] and Interleukin4 (IL4) conferred resistance, but much more weakly. In contrast, TGFβ enhanced dasatinib activity.

IL7 Mediates Primary Drug Resistance against All Approved BCR-ABL–KIs in Murine and Human Ph+ ALL Cells, Which Is Completely Reversible

Next, we extended the evaluation of IL7, our most potent candidate from the cytokine screen, to assess its impact on the performance of other drugs, including
Figure 4.1. Leukemia stem cell (LSC) function assay by serial dilution and implantation comparing the *in vivo* leukemia-initiating capacity of $Arf^{-/-} \ p185^{WT}$ (left) and $Arf^{-/-} \ p185^{T315I}$ pre-B (right) populations. $Arf^{-/-} \ p185$-transformed pre-B murine cells are referred as LICs. Kaplan-Meier curves represent the overall survival of nonconditioned immunocompetent C57Bl/6J recipient mice that received serial log-fold dilutions (n=4 per arm) of LIC-number (as indicated in the inset legend) on day 0.
Figure 4.2. IL7 confers BCR-ABL<sub>MUTANT</sub>-independent primary drug-resistance against BCR-ABL–KIs in Ph+ ALL.

(a) Radar plot representing the effect of 15 hematopoietic cytokines on dasatinib potency (EC<sub>50</sub>) against <i>Arf</i>–/– BCR-ABL<sup>WT</sup> LICs. CTG-assay measured LIC killing after 72 h dasatinib exposures in presence of increasing concentrations of indicated cytokines. Each spoke represents an individual cytokine; each line graph represents dasatinib EC<sub>50</sub> (nM) at indicated titration of the parent cytokine (see supplemental data in Appendix).

(b) <i>Arf</i>–/– BCR-ABL<sup>WT</sup> (triangles) and <i>Arf</i>–/– BCR-ABL<sup>T315I</sup> (circles) LICs were treated with drug or DMSO for 72 h in the absence (blue) or presence (red) of 1 ng/mL mIL7, in triplicates. For each drug concentration, LIC growth measured by CTG-assay was normalized to DMSO-treated LICs and fit using nonlinear regression (+/- sd).

(c) Viability of human Ph+ ALL SUP-B15 cells after 72 h drug-treatment in absence or presence of 25 ng/mL hIL7, was measured by MTS assay. Values were normalized to DMSO-treated cells. (d, e) Kaplan-Meier survival curves showing overall survival in IL7<sup>+/+</sup> and IL7<sup>–/–</sup> Bl6 host-mice receiving day 0 tail vein injection of (d) 2×10<sup>2</sup> <i>Arf</i>–/– BCR-ABL<sup>WT</sup> LICs (vehicle treated), with median survival of 20 days in IL7<sup>+/+</sup> (n=4) versus 33 days in IL7<sup>–/–</sup> mice (n=4), P=0.0069, or (e) 2×10<sup>5</sup> <i>Arf</i>–/– BCR-ABL<sup>WT</sup> LICs. Data are an average of 2 independent experiments in which vehicle or dasatinib therapy was initiated on day 7 or 9. Survivals of vehicle-treated IL7<sup>+/+</sup> and IL7<sup>–/–</sup> mice were comparable. Dasatinib treatment caused a median survival of 33 days in IL7<sup>+/+</sup> (n=12) versus 41 days in IL7<sup>–/–</sup> (n=11) mice, P=0.0061.
BCR-ABL–KIs and non-BCR-ABL inhibiting drugs. IL7 significantly attenuated the potency of all FDA-approved BCR-ABL–KIs against murine Arf−/− BCR-ABLWT LICs (Figure 4.2b) and the human Ph+ ALL cell line SUP-B15 (Figure 4.2c). In contrast, IL7 did not affect the potency of tested non-BCR-ABL drugs underscoring the specificity of this cytokine protection (Figure 4.3).

The specificity and dynamic nature of IL7 in mediating primary resistance against BCR-ABL-targeting drugs was further confirmed by four in vitro findings. First, mechanistic evaluations by FACS analysis revealed dasatinib-induced phenotypic effects of G0-G1 growth arrest and apoptosis of Arf−/− BCR-ABLWT LICs were significantly blunted by the presence of cell-extrinsic IL7 during drug exposure (Figure 4.4a and b). Second, transient pharmacological BCR-ABL-inhibition, achieved by washing out drug/cytokine after a period of dasatinib/IL7 co-treatment (see ‘Drug/cytokine washout and cytokine recovery studies’ in Chapter 3), dramatically increased the responsiveness of Arf−/− BCR-ABLWT LICs to cell-extrinsic IL7 (Figure 4.4a, c and d) [96]. Third, removing (washing out) IL7 completely reversed cytokine protection and restored dasatinib’s potent activity to kill Arf−/− BCR-ABLWT LICs (see ‘dasatinib re-challenge studies’ in Chapter 3, data not shown). Fourth, IL7 significantly rescued the clonal outgrowth capacity of single flow-sorted Arf−/− BCR-ABLWT LICs during prolonged treatments with dasatinib (Figure 4.5).

Thus, IL7 imparts primary refractoriness against approved BCR-ABL–KIs in murine and human Ph+ ALL cells, which exists as a completely reversible phenomenon.

Physiological Levels of Host-expressed IL7 Contributes to an Aggressive, Dasatinib-resistant Phenotype of Ph+ ALL In Vivo

We studied the effect of physiologic levels of IL7 on leukemia progression and dasatinib responsiveness in vivo. Host IL7+/+ (i.e. wild-type; WT) mice injected with 2×10² Arf−/− BCR-ABLWT LICs showed more rapid leukemia onset compared to strain-matched IL7−/− (null) hosts (Figure 4.2d), suggesting that murine Ph+ ALL responds to host-IL7 with accelerated progression. Injecting 2×10⁵ Arf−/− BCR-ABLWT LICs overcame this trend (Figure 4.2e), suggesting the phenomena is most important during the phase of disease establishment. Host-expressed physiologic level of IL7 significantly shortened the survival benefit from dasatinib mono-therapy of BCR-ABLWT leukemias (Figure 4.2e). This finding is the first definitive demonstration of a single host-cytokine in inducing BCR-ABL–KI drug resistance in vivo. Of note, terminal clinical presentation in IL7-WT host-mice was either with signs of CNS disease (bulging skull, uneven gait, erratic behavior) or frank relapse in rest of the body, whereas the majority of IL7−/− host-mice presented with signs of CNS disease necessitating their sacrifice, consistent with reports on low bioavailability of BCR-ABL–KIs in CNS [70-71].

From a therapeutic perspective, these findings in combination with our in vitro observations support that in vivo responses to dasatinib therapy in Ph+ ALL are modulated by cell-extrinsic IL7 in the absence of BCR-ABL KD mutations.
Figure 4.3. Comparative evaluation of IL7-mediated protection conferred against non-BCR-ABL-specific targeting drugs (top panel) and conventional cytotoxic antileukemic clinical drugs (bottom panel).

Arf\textsuperscript{+} BCR-ABL\textsuperscript{WT} LICs were treated with indicated drugs or DMSO (no drug) for 72 h in the absence (blue) or presence (red) of high (10 ng/mL) murine IL7, in triplicates. The antibiotic drug Cefaclor (often used in leukemia clinics) was used as a non-antileukemic drug control. For each drug concentration, LIC growth was measured by the CTG-assay and average values were normalized to DMSO-treated LICs and graphed (+/− sd).
Figure 4.4. Protection of $Arf^{-/-}$ BCR-ABL$^{WT}$ LICs by IL7 during dasatinib exposures (0 to 72 h) and in the post-exposure recovery period (72 to 144 h, no dasatinib).

See Chapter 3 for details. (a) Total sample viabilities (DAPI staining, FACS) at the end of exposure period (dark gray=IL7; light gray=No IL7) and recovery period (red=IL7; blue=No IL7). Light gray versus dark gray comparison depicts significant IL7 protection during dasatinib exposures. Few LICs surviving dasatinib exposures in the absence of IL7 (light gray), later deteriorated and died during recovery period, irrespective of IL7, confirming that BCR-ABL inhibitor exposures commit BCR-ABL+ cells to death [56]. IL7 significantly improved recovery of dasatinib/IL7 co-treated LICs (red vs blue), clearly evident at 1000 nM but not at 100 nM and 10 nM dasatinib concentration because of over-confluent growth in presence of IL7 (see (c) and (d)). (b) Cell cycle changes (FACS) at the end of exposure period (corresponding to light and gray bars in (a)). Incremental dasatinib doses (which cause incremental BCR-ABL inhibition [55]) induced cell cycle arrest and apoptosis; IL7 significantly protected LICs from these changes (c) Cell cycle analysis at the end of recovery period (corresponding to blue and red in (a)). (d) Conditions showing positive LIC growth during recovery period, normalized for equal number of LICs plated at start of recovery (dark grey). Ratio of fold LIC-change in presence (face-down bracket) to that in absence (face-up bracket) of IL7 increased in LICs with higher previous BCR-ABL inhibition from higher dasatinib concentrations during exposure period [96].
Figure 4.5. *In vitro* leukemia clonal outgrowth efficiency assays.

Either 1 or 10 viable \( Arf^{−/−} \) BCR-ABL\(^{WT} \) LICs per well sorted by single-cell flow cytometry into 3 clear-bottom 384-well plates for each condition were treated (t=0) with DMSO (0.1 % by volume) or dasatinib in the absence (dark grey) or presence (light grey) of IL7 (5 ng/mL). After 8 days of incubation, naked-eye microscopic examination was performed. Wells showing significant positive growth (approximately \( ≥ 50 \) LICs per well) were scored positive and percentage of positive wells for each condition indicated on the x-axis was calculated.
IL7 Protects against Dasatinib by Strongly Counteracting the Decrease in pSTAT5 Protein and c-MYC Expression

What intra-cellular changes initiated by the host-IL7 are critical for imparting primary dasatinib resistance? Arf\textsuperscript{−/−} BCR-ABL\textsuperscript{WT} LICs were briefly (4 h) challenged with 100nM dasatinib in the presence or absence of IL7 and then evaluated using a cell-signaling kit (Luminex® 8-plex kit by Millipore, MA) which detects level of phosphorylated (active) form of eight different cell-signaling proteins relevant to cancer biology (Figure 4.6a) and gene expression arrays (Figure 4.6b). Comparative assessments in the global gene expression and various signaling pathways allowed us to detect the intra-cellular changes crucial for mediating IL7-induced dasatinib-refractoriness.

Among the eight phosphorylated signaling proteins evaluated, short dasatinib exposure in the absence of IL7 decreased the level of mainly pSTAT5 protein (Figure 4.6a). STAT5 is an established phosphorylation target of uninhibited BCR-ABL kinase activity [97-98], and decrease in the level of pSTAT5 is commonly used as a surrogate marker of cellular BCR-ABL-inhibition [99-100]. Interestingly, the presence of IL7 significantly maintained high levels of pSTAT5 in Arf\textsuperscript{−/−} BCR-ABL\textsuperscript{WT} LICs despite pharmacological BCR-ABL inhibition by dasatinib (Figure 4.6a) [101]. These findings were confirmed using conventional western blotting (Singh and Opferman, unpublished). A positive trend was observed between the levels of pSTAT5 protein in Arf\textsuperscript{−/−} BCR-ABL\textsuperscript{WT} LIC samples at 4 hours and their respective total sample viability and total viable cell counts after 72 hours, as measured by CTG and FACS (compare with Figure 4.2b and Figure 4.4a, respectively). This suggested a new mechanistic role for pSTAT5 in IL7 mediated rescue of Ph+ ALL from BCR-ABL inhibitory therapy.

From gene expression data, we dissected a set of mRNAs that underwent the most significant quantitative change in cellular expression due to brief dasatinib exposure in the absence of IL7, relative to untreated LICs, to determine a ‘dasatinib response signature’ in Ph+ ALL. Using this signature as a reference, we identified a subset of mRNAs which were most significantly protected from dasatinib-induced change in cellular expression due to the presence of IL7 during drug exposure, referred as ‘IL7-imparted dasatinib resistance signature.’ Enrichment analysis (Figure 4.6b) of the cytokine-mediated dasatinib resistance signature demonstrated that IL7 strongly protects cellular expression of c-MYC and c-MYC target genes, with c-MYC being the second highest protected mRNA by IL7 (see supplemental data in Appendix).

Cellular binding of growth factor IL7 activates the receptor-associated JAK3-STAT5 signaling pathway [101], resulting in the direct upregulation of c-MYC by activated STAT5 (pSTAT5) [102]. Thus, IL7 counteracts the decrease in both pSTAT5 and c-MYC resulting from pharmacological inhibition of BCR-ABL by dasatinib, and rescues Arf\textsuperscript{−/−} BCR-ABL\textsuperscript{WT} LICs by initiating IL7-JAK3-pSTAT5-cMYC signaling.
**Figure 4.6.** Mechanistic evaluation of host-IL7 imparted primary resistance against dasatinib.

*Arf*<sup>−/−</sup> BCR-ABL<sup>WT</sup> LICs were challenged with dasatinib or DMSO for 4 hrs in presence or absence of IL7. (a) Levels of pSTAT5 measured at 4 hrs by Luminex 8-plex multipathway kit are presented. Change in level of pErk/MAP kinase 1/2, pSTAT3, pJNK, p70 S6 kinase, IkBalpha, CREB, and p38 was statistically insignificant and is not shown. (b) Cartoon shows enrichment analysis of ‘IL7 protection gene signature against dasatinib’. Increase or decrease in expression of genes by dasatinib alone (no IL7) is presented by red (decrease) and blue (increase) color, respectively. Size of the circle presents fold-protection against dasatinib-induced change in gene expression due to the presence of IL7. Only top representative genes undergoing significant change in the presence of IL7 are shown. See supplemental data in **Appendix** for complete data set.
High pSTAT5 and IKAROS Haploinsufficiency Are Candidate Resistance Biomarkers with c-MYC as a Convergence Node of Resistance against BCR-ABL Inhibitor Therapy in Ph+ ALL

Our mechanistic studies implicated c-MYC expression in cytokine mediated BCR-ABL–KI-resistance. Because IKAROS directly represses c-MYC expression in pre-B cells and IKAROS haploinsufficiency relieves such c-MYC repression [103], we hypothesized that IKAROS haploinsufficiency by derepressing c-MYC expression may also impart cell-intrinsic BCR-ABL–KI-resistance in Ph+ pre-B cells.

To confirm the role of pSTAT5, c-MYC and IKAROS haploinsufficiency in ABL KD mutation-independent drug-resistance, we utilized fluorescent vectors expressing constitutively active STAT5 (CA-STAT5) single or double mutants, human c-MYC or dominant negative (DN) IKAROS (IK6), imitating constitutive signaling by pSTAT5, c-MYC-overexpression and IKAROS haploinsufficiency, respectively (Figure 4.7). Analogous to IKAROS-WT, the IK1 mutant binds to the c-MYC promoter and inhibits its expression, whereas the IK6 mutant derepresses c-MYC expression [75, 103]. Arf−/− BCR-ABLWT LICs transduced (florescent) with these vectors were mixed with parent non-transduced (non-florescent) LICs and subjected to drug-competition assays in the absence of IL7. HT-flow assay (please refer Figure 3.3) performed on day 2, 3, 5 and 9 relative to day 0 allowed a temporal assessment of ‘engineered’ cell-intrinsic genetic changes on growth and cellular response to dasatinib and imatinib.

In the absence of BCR-ABL–KI treatment (Figure 4.7a), proliferation of Arf−/− BCR-ABLWT LICs expressing GFP-only (‘empty’ vector), IK6 or c-MYC was comparable to non-transduced parent LICs, whereas fluorescent Arf−/− BCR-ABLWT LICs expressing pre-validated single or double CA-STAT5 mutants or the IK1 mutant [75, 103], were quickly lost in culture. The later observation corroborated the known essential role of c-MYC expression in sustaining Ph+ LICs [60-61].

Treatment (Figure 4.7b) of control LIC mixtures with increasing imatinib and dasatinib concentrations did not alter the ratio between non-transduced LICs and daughter LICs transduced with GFP-only vectors. The total mixture viability (by DAPI exclusion) of control LIC mixtures decreased in a dose and time proportionate fashion. This corroborated our previous flow-cytometry results (Figure 4.4a), and confirmed the overall quality of HT-flow drug-screening runs. In stark contrast to control LIC mixtures, drug exposures temporally selected the Arf−/− BCR-ABLWT LICs transduced with vectors expressing c-MYC or IK6 or single mutant CA-STAT5 or double mutant CA-STAT5. Remarkably, HT-flow counter genetic-screen demonstrated that the degree of LIC-selection to be quantitatively proportional to BCR-ABL-inhibition achieved by incremental BCR-ABL–KI doses (Figure 4.7b and c) [55]. These results were later confirmed with western blots (unpublished, Singh and Opferman). Thus, pSTAT5, c-MYC and functional-loss of IKAROS, all mediate BCR-ABL mutation-independent drug resistance according to proposed model in Figure 4.7d. In a direct test of c-MYC as a node for dasatinib-resistance in Ph+ ALL, dasatinib-challenges were performed against Arf−/− BCR-ABLWT LICs expressing either an empty vector or c-MYC protein.
Figure 4.7. Results of counter genetic screen by HT-flow confirming the role of pSTAT5 activity, *IKAROS* haploinsufficiency and c-MYC over-expression in primary drug resistance against BCR-ABL-inhibiting drugs.

Mixtures composed of non-transduced (non-fluorescent) parent p185<sup>WT</sup>LICs and transduced (fluorescent) daughter p185<sup>WT</sup>LICs expressing protein of interest (as per legends) were exposed to (a) no drug (controls) or (b,c) wide-range of imatinib and dasatinib concentrations in the absence of IL7, in triplicate wells. Percentage of fluorescent (transduced) LICs on day 0, 2, 5 and 9 was measured by HT-flow assay. (b) Ratio of average (n=3) number of viable fluorescent (transduced) LICs (by DAPI) on day 2, 5 and 9 relative to day 0, at each indicated experimental condition, are shown for dasatinib and imatinib. Change in ratio is presented using a three-color graded scale (red>1, yellow=1, green<1; dark (red/green) color implies higher change and light (red/green) color implies lower change. (c) X-Y line graph representing temporal selection (x-axis) of CA-STAT5 double mutant (DM) expressing LICs (relative to day=0) due to exposure to dasatinib concentrations, in a dose-response fashion. (d) Proposed mechanistic model of BCR-ABL mutation-independent drug-resistance against BCR-ABL inhibiting drugs in Ph+ ALL.

* Zero values (darkest green) indicate that total sample viability is below the detection-threshold by flow cytometry.

** Relatively low number of LICs transduced by single and double CA-STAT5 mutants on day=0 in (a) due to low transduction efficiency (from day –2 to 0) later allowed much higher fold selection within the same wells (from day 0 to 9) in (b).
Over-expression of c-MYC resulted in a significant deterioration (100 fold) of dasatinib potency (EC$_{50}$) in a BCR-ABL mutation-independent fashion (data shown later for convenience).

As an independent test of our hypothesis, we examined published biochemical profiles of agents for their binding affinity to kinase proteins [20], especially against BCR-ABL and JAK3 kinases, the latter being specific for mediating the effects of extracellular IL7 to intracellular compartment. Indeed, both dasatinib and nilotinib inhibit BCR-ABL at nanomolar concentrations, but were found to completely lack the ability to bind and inhibit JAK3 kinase.

**HT Drug Screens to Identify Agents with Activity against Ph+ ALL Cells Protected by Extracellular IL7 and T315I Mutation**

To search for novel agents with potent cellular activity against IL7-induced BCR-ABL–KI-resistant phenotypes, we screened 3200 well-annotated and mechanistically diverse bioactive agents, which included FDA-approved oncology and anti-infective drugs. In the primary screen, Arf$^{−/−}$ BCR-ABL$^{WT}$ LICs and Arf$^{−/−}$ BCR-ABL$^{T315I}$ LICs were treated with a single concentration (10 μM) of agents in the presence of IL7 (0.85 ng/mL). The quality of the screen was confirmed by receiver operating characteristic (ROC) analysis and other statistical metrics (Figures 4.8 to 4.12). The 627 compounds showing statistically significant activity (≥ 10% growth inhibition) were further augmented with available structural analogs and drugs of the same pharmacologic class to yield 706 active compounds. Concentration–response relationships were established for these compounds against Arf$^{−/−}$ BCR-ABL$^{WT}$ LICs and drug-resistant Arf$^{−/−}$ BCR-ABL$^{T315I}$ LICs in the presence of IL7, giving 229 validated hits with reasonable potency. All validated hits were equipotent in inhibiting the growth of Arf$^{−/−}$ BCR-ABL$^{WT}$ LICs and Arf$^{−/−}$ BCR-ABL$^{T315I}$ LICs. This set included 15 known cytotoxic antileukemic drugs (See supplemental data in Appendix), which enhance BCR-ABL–KI therapy in vivo [44-45, 63], thus validating the detection of appropriate phenotypic responses. Furthermore, cluster analysis (Figure 4.13) indicated that this set effectively sampled the space of all existing therapeutic classes (Figures 4.12 and 4.13). See supplemental data in Appendix for detailed annotation of all validated hits.

**A Common Anti-malarial Agent DHA Overcomes Multi-factorial BCR-ABL–KI Drug Resistance by Cell-extrinsic IL7 and Cell-intrinsic c-MYC Over-expression**

Given that the validated hits that overrode the IL7-protected leukemia phenotype contained compounds with varying mechanisms of action (Figure 4.13), and that increased c-MYC expression acts as a common node for primary BCR-ABL–KI-resistance in ‘high-risk’ Ph+ ALL (Figures 4.6 and 4.7), we aimed to select those validated hits which may demonstrate selectivity to kill cancer cells with increased c-MYC expression and a known favorable toxicity profile in humans.
Figure 4.8. Quality control for the primary screen of 5600 agents against *Arf*–/– BCR-ABL<sup>WT</sup> LICs performed in the presence of 0.85 ng/mL IL7.

See Table 3.1 for well locations. (a) Distribution of EC<sub>50</sub> of non-BCR-ABL-inhibiting reference compound staurosporine (calculated from wells 2A-2J in (d)). (b) Scatter plot of percent activity of the test compounds relative to controls. Good separation is seen between negative controls (red, DMSO-treated, max signal, column 13 in (d)) and positive controls (green, 300 nM staurosporine, min signal, column 14 in (d)). Test compounds showing statistically significant activity relative to background noise (negative controls, red) are depicted in blue and inactive test compounds in black. Densities for positive, negative, and test compounds are shown on the right. (c) Distribution of Z’ values for each plate in the primary screen. All Z’ values throughout the run – average=0.82, lowest=0.71 and highest=0.9 (d) Heat map comparison of well activity averaged across all plates. Wells 1A-1J and 2A-2J show serial dilutions of dasatinib and staurosporine. In presence of IL7, average LIC growth inhibition of approx. 45% was achieved by 100 nM dasatinib (well 1A), the later approaching maximum concentration achieved in human plasma.
Figure 4.9. Quality control for primary screen of 5600 agents against $Arf^+$ BCR-ABL$^{T315I}$ LICs performed in the presence of 0.85 ng/mL IL7.

(a) Distribution of EC$_{50}$ of non-BCR-ABL-inhibiting reference compound staurosporine (calculated from wells 2A-2J in (d)). (b) Scatterplot of percent activity of the test compounds relative to controls. Good separation is seen between negative controls (red, DMSO-treated, max signal, column 13 in (d)) and positive controls (green, 300 nM staurosporine, min signal, column 14 in (d)). Test compounds showing statistically significant activity relative to background noise (negative controls, red) are depicted in blue; inactive test compounds are black. (c) Distribution of Z’ values for each plate in the primary screen. All Z’ values throughout the run – average=0.85, lowest=0.77, and highest=0.92 (d) Heat map comparison of well activity averaged across all plates. Wells 1A-1J and 2A-2J show serial dilutions of dasatinib and staurosporine tested against BCR-ABL$^{T315I}$ LICs. Although 100 nM dasatinib (Well 1A) was totally ineffective, well 2A with highest staurosporine concentration (300 nM) showed average LIC growth inhibition of approximately 100%.
Figure 4.10. Primary high-throughput drug screening experiments captured BCR-ABL–KI drug-resistant phenotypes with high accuracy.

Distributions of independent dose response curves of dasatinib and staurosporine against Arf–/– BCR-ABLWT and Arf–/– BCR-ABL T315I LICs in the presence of 0.85 ng/mL IL7 calculated separately from all assay plates (wells 1A-1J and 2A-2J) of primary drug screening experiments (also see Figures 4.8d and 4.9d) are presented. Both IL7-imparted dasatinib–resistant phenotype in BCR-ABLWT LICs (EC50 in nM: 95%CI = 86-116, average >100) and BCR-ABL-mutation-imparted BCR-ABL–KI-resistant phenotype in BCR-ABL T315I LICs, similar to Figure 4.2b, were consistently captured across all plates. Low interplate variability in calculated EC50 values was observed for the non-BCR-ABL-targeting reference compound staurosporine (95%CI for EC50 in nM: WT = 11-12, T315I = 15-16) and majority of the validated hits (refer the supplemental data in Appendix).
Figure 4.11. Receiver Operating Characteristic (ROC) analysis demonstrates that both (a) \( \text{Arf}^{--} \) BCR-ABL\(^{WT}\) and (b) \( \text{Arf}^{--} \) BCR-ABL\(^{T315I}\) LIC-based assays have high discriminatory power (AUC 0.83 and 0.85, respectively).

The ROC curve is plotted as a function of % activity, and is color coded according to the right Y-axis. The ROC curves in gray are calculated from 200 bootstrap simulations. The dashed line corresponds to a random assay.
Figure 4.12. Frequency of ATC Level 1 (top) and ATC Level 2 (bottom) compounds for the 706 compounds submitted to secondary analysis.
Figure 4.13. High-throughput drug screen to identify novel agents possessing activity against IL7-mediated early (primary) and BCR-ABL$^{\text{MUTANT}}$-mediated late (secondary) BCR-ABL–KI drug-resistant phenotypes.

Clustering of validated hits using ATC therapeutic classification established by World Health Organization (WHO) (http://www.whocc.no/atc/structure_and_principles/). A single compound can be represented more than once because of multiple therapeutic indications, as is the case for the corticosteroids (highly active clusters in A, C, D, H, R, and S). This demonstrates that the active compounds come from drugs used to treat many indications. For details on clustering methodology, please refer to Chapter 3, and for details on clustering analysis, please refer to supplemental data in Appendix and Figure 4.12.
A correlative statistical analysis of NCI60 screening databases was performed. While a few validated hits demonstrated a significant positive (+) correlation between their anti-cancer activity and c-MYC expression across 60 human cancer cell lines, the phenotypic sensitivity to dasatinib was found not to be correlated (R = –0.11, where R is Pearson’s coefficient of correlation) with c-MYC expression (Table 4.1). From the pool of validated hits demonstrating positive correlation with cellular c-MYC expression for their drug activity, we selected the artemisinin class of FDA-approved and very well-tolerated antimalarial drugs (ARTs) which included artemisinin, artesunate, and dihydroartemisinin (DHA) [104]. As a class, the ARTs were equipotent against $\text{Arf}^{-/-}$ BCR-ABL$^\text{WT}$ and $\text{Arf}^{-/-}$ BCR-ABL$^{T315I}$ LICs (Figure 4.14a) but the anti-LIC potency of artemisinin was much lower than that of artesunate or DHA. DHA and artesunate were also active against the human Ph+ ALL cell line SUP-B15 (Figure 4.14b) and had similar anti-Ph+ ALL potency in the presence of cell-extrinsic IL7 and cell-intrinsic over-expression of c-MYC, in stark contrast to dasatinib (Figure 4.14c). Studies have shown that DHA kills other genotypes of leukemic cells while selectively sparing normal human lymphocytes [105-106]. DHA, an active metabolite of many ARTs family members including artesunate, was chosen for further analysis because of its positive correlation for activity against tumor cells with relatively higher MYC expression ($R = +0.5996, p = 0.0000433$), in vitro potency, cellular selectivity, favorable pharmacokinetics, known bioavailability in hematopoietic system, and tolerability in all age groups.

**Dihydroartemisinin Possesses Single-agent In Vivo Activity against Ph+ ALL Phenotypes Demonstrating Primary and Acquired Dasatinib Resistance**

DHA monotherapy gave significant dose-dependent survival benefit in IL7-WT host-mice bearing low-burden $\text{Arf}^{-/-}$ BCR-ABL$^\text{WT}$ LICs (Figure 4.15a). Leukemic samples collected from the relapsing mice at the time of sacrifice revealed the complete absence of ABL KD mutations, which are often selected with dasatinib treatment in this model. DHA monotherapy also gave a similar response against low-burden $\text{Arf}^{-/-}$ BCR-ABL$^{T315I}$ leukemia, a disease that is completely refractory to dasatinib (Figure 4.15b). Thus, DHA is active in vivo against murine Ph+ ALL through a mechanism independent of the BCR-ABL kinase.

**Cotreatment with Dihydroartemisinin Overcomes Host-mediated Primary Drug-resistance against Dasatinib and Eliminates Ph+ ALL In Vivo**

Given that $\text{Arf}^{-/-}$ BCR-ABL$^\text{WT}$ leukemias in IL7-deficient mice were more sensitive to dasatinib (Figure 4.2e), and that DHA circumvented IL7 protection in vitro (Figure 4.14a and b) and displayed anti-Ph+ ALL activity in IL7-WT host mice (Figure 4.15a and b), the potential for synergy between the two drugs was evaluated. This evaluation was performed in cohorts of IL7$^{+/+}$ immuno-competent mice bearing high burdens of $\text{Arf}^{-/-}$ BCR-ABL$^\text{WT}$ Luc+ LIC-driven leukemia (analogous to IL7$^{+/+}$ [WT] [WT]
Table 4.1. Priortization of validated hits according to known human tolerability and predicted involvement of c-MYC expression in their anti-cancer drug action.

<table>
<thead>
<tr>
<th>Hits Active against Dasatinib Resistant Ph+ ALL Phenotypes*</th>
<th>NSC No. for Hits in DTP Database</th>
<th>R (Coefficient of Correlation)**</th>
</tr>
</thead>
<tbody>
<tr>
<td>GAMBOGIC ACID</td>
<td>693702</td>
<td>+0.59</td>
</tr>
<tr>
<td>ARTESUNTE</td>
<td>712571</td>
<td>+0.59</td>
</tr>
<tr>
<td>PATULIN</td>
<td>8120</td>
<td>+0.59</td>
</tr>
<tr>
<td>DIHYDRO-ARTEMISINYL ETHER</td>
<td>685989</td>
<td>+0.57</td>
</tr>
<tr>
<td>DIHYDRO-ARTEMISINYL ETHER</td>
<td>685988</td>
<td>+0.56</td>
</tr>
<tr>
<td>ARTEMISININE</td>
<td>369397</td>
<td>+0.48</td>
</tr>
<tr>
<td>MYCOPHENOLIC ACID</td>
<td>129185</td>
<td>+0.44</td>
</tr>
<tr>
<td>CELASTROL</td>
<td>70931</td>
<td>+0.42</td>
</tr>
<tr>
<td>THIRAM</td>
<td>1771</td>
<td>+0.40</td>
</tr>
<tr>
<td>AKLAVINE HYDROCHLORIDE</td>
<td>100290</td>
<td>+0.40</td>
</tr>
<tr>
<td>CHELERYTHRINE</td>
<td>36405</td>
<td>+0.36</td>
</tr>
<tr>
<td>METHOTREXATE</td>
<td>740</td>
<td>+0.35</td>
</tr>
<tr>
<td>EMETINE DIHYDROCHLORIDE</td>
<td>33669</td>
<td>+0.25</td>
</tr>
<tr>
<td>ANTIMYCIN A</td>
<td>186</td>
<td>+0.06</td>
</tr>
<tr>
<td>MITOXANTHRONE HYDROCHLORIDE</td>
<td>301739</td>
<td>+0.04</td>
</tr>
<tr>
<td>MITOMYCIN C</td>
<td>26980</td>
<td>+0.04</td>
</tr>
<tr>
<td>DASATINIB</td>
<td>732517</td>
<td>-0.11</td>
</tr>
</tbody>
</table>

* Examplary screening hits are presented for which preclinical and clinical toxicity data is available.

** R was calculated between drug activity and c-MYC expression across NCI60 human cancer cell lines, using NSC numbers as chemical identifiers for identified hits. Total sample size of 4461 (4444 + 17) chemicals and drugs was used to calculate p values; see Chapter 3 for details.
Figure 4.14. Dihydroartemisinin of Artemisinin family of antimalarials circumvents dasatinib resistance induced by IL7, c-MYC overexpression and T315I mutation in vitro.

(a) $Arf^{-/-}$ BCR-ABL$^{WT}$ (triangles) and $Arf^{-/-}$ BCR-ABL$^{T315I}$ (circles) LICs were treated with artemisinin, artesunate, dihydroartemisinin, or DMSO for 72 h in the absence (blue) or presence (red) of 1 ng/mL murine IL7, in triplicates. For each drug concentration, LIC growth (measured by the CTG-assay) was normalized to DMSO-treated LICs and fit using nonlinear regression (+/- sd).

(b) Human Ph+ ALL SUP-B15 cells were treated with artesunate, dihydroartemisinin, or DMSO (no drug) for 72 h in the absence (blue) or presence (red) of 25 ng/mL human IL7. Viability was measured by the MTS assay.

(c) Heads-up comparison of potencies of dihydroartemisin and dasatinib against (i) $Arf^{-/-}$ BCR-ABL$^{WT}$ LICs +/- IL7, (ii) $Arf^{-/-}$ BCR-ABL$^{WT}$ LICs +/- c-MYC overexpression +/- IL7, and (iii) $Arf^{-/-}$ BCR-ABL$^{WT}$ LICs, using CTG-assay.
Figure 4.15. Dihydroartemisinin is active as a single agent against dasatinib-resistant Ph+ ALL in IL7+/+ host mice.

Kaplan-Meier survival curves showing the overall survival in response to therapy (survival benefit) in IL7+/+ Bl6 host mice that received day 0 tail vein injection of (a) $2 \times 10^3$ Arf–/– BCR-ABLWT Luc+ LICs; on day 3, once daily for 5 days of each week, vehicle, 100 mg/kg DHA, 200 mg/kg DHA, 300 mg/kg DHA, or 300 mg/kg DHA (for 3 weeks only) therapy was initiated in mice bearing low burdens of BCR-ABLWT-driven leukemia, or (b) $2 \times 10^3$ Arf–/– BCR-ABLT315I Luc+ LICs; on day 3, once daily for 5 days of each week, vehicle, 300 mg/kg DHA, or 10 mg/kg dasatinib was given to mice bearing low burdens of BCR-ABL T315I-driven leukemia.
human patients carrying high burdens of Arf-deficient p185BCR-ABL pre-B cell leukemia at the time of diagnosis). Separate cohorts of animals were treated with dasatinib (10 mg/kg), DHA alone (200 or 300 mg/kg) or binary combinations of dasatinib and DHA as shown in Figure 4.16a. Despite continuous monotherapy, dasatinib induced weak, transient primary responses. As a result, all animals harbored dasatinib-refractory persistent leukemia that inevitably led to relapse (Figure 4.16b) and caused death of all recipients within 6 weeks of initiating therapy (Figure 4.16a). DHA alone was completely ineffective (200 mg/kg) or weakly effective (300 mg/kg) in the high-burden leukemia model (Figure 4.16a). The combination of DHA and dasatinib significantly increased overall survival (Figure 4.16a), induced more rapid and deeper initial responses to dasatinib, and sensitized host-protected persistent leukemia to dasatinib (Figure 4.16b), in comparison to the treatment with either of the two drugs alone. Of the 12 mice given 200 mg/kg DHA and 10 mg/kg dasatinib, 3 survived long term, apparently free of all leukemia despite cessation of therapy. All mice (n=12) given 300 mg/kg DHA and 10 mg/kg dasatinib survived the treatment period. After 4 weeks from the termination of combined therapy, 1 mouse succumbed to an isolated CNS relapse, which may be related to poor CNS penetration of dasatinib.[70-71]. Remarkably, the 14 surviving mice taken off the dasatinib and DHA combination regimens (3/12 at 200 mg/kg and 11/12 at 300 mg/kg DHA) for one year show no trace of leukemia. This data strongly suggested that drug combination cured the disease because in this leukemia model all recipient animals left untreated invariably succumb to fatal, full-blown leukemia within 4 weeks of injecting a very low numbers of LICs, as shown in Figure 4.1. Dasatinib-DHA regimens were well tolerated in leukemic and non-leukemic mice, in congruence with prior work utilizing similar dosages of DHA [93] and other ART family members [107-108] in rodents [93, 107] and sheep [108].

To extend our mechanistic understanding on DHA drug action, we identified genes whose expression levels were positively correlated with drug-sensitivity to DHA across 60 different human cancer cell lines, using public databases. We determined that the top 40 genes, which represent the drug-sensitivity signature for DHA in diverse human tumors, have a high and comparable expression in murine LICs and primary human Ph+ ALL leukemic samples with a positive correlation of 0.592 (p < 0.001) as shown in Figure 4.17. This data predicts that DHA may maintain its anti-leukemic drug action against primary Ph+ ALL in human patients, similar to its efficacious results in our genetically relevant murine leukemia model (Figure 4.16a).

Next, to identify candidate genes targeted by DHA that plausibly resulted in overcoming host-cytokine mediated refractoriness against dasatinib, we compared gene signatures of DHA drug sensitivity (represented in Figure 4.17) and IL7-mediated dasatinib resistance (represented in Figure 4.6b). Three genes were found to overlap, namely MYC (probeset 1424942_a_at), Cabc1 (probeset 1417066_at, chaperone, ABC1 activity of bc1 complex like (S. pombe)) and Akap1 (probeset 1449019_at, A kinase (PRKA) anchor protein 1) using a 2-fold change as a cut-off in the IL7-resistance signature, whereas setting a 4-fold cut-off left MYC as the only overlapping gene between the two signatures. This provided an alternative confirmation for our results.
Figure 4.16. Dihydroartemisinin works synergistically with dasatinib to cure Ph+ ALL by improving primary responses to dasatinib.

(a) Kaplan-Meier survival curves showing the survival benefit in response to therapy in IL7+/+ Bl6 host mice that received day 0 tail vein injection of $2 \times 10^5$ Arf−/− BCR-ABLWT Luc+ LICs. On day 7, six mouse groups bearing high burdens of BCR-ABLWT-driven leukemia were started on daily treatment for 5 days of each week, using two-dose regimens – 2 vehicles or 1 vehicle and 1 agent or 2 agents (see label), with 2 doses of each regimen given separately, 8 h apart, by oral gavage. On day 52, all therapy was terminated. (b) For 6 mouse groups in (a), separate graphs represent quantitative assessment of serial whole-animal luminescent signals (p/s/cm²/sr) acquired weekly. Solid lines represent imaging intensity of signal (y-axis) as a measure of leukemia burden from individual mice in the group, which is traced in real time on the x-axis. On day 7, before initiating therapy, all 6 groups had comparable and high burden of leukemia. The horizontal dotted line represents the empirically determined lower limit of sensitivity for luminescent signal as an experimental cut-off for detectable levels of persistent leukemia. Rectangular gray box quantifies experimental primary therapeutic response to indicated regimens; box-width depicts approximate time (days) to induce maximal initial response and box-depth represents maximal decrease achieved in leukemia burden signal. Increasing depth and decreasing width indicate stronger and quicker primary responses, respectively.
Figure 4.17. Statistical analyses to predict the activity of dihydroartemisinin against human Ph+ ALL.

‘Global anti-tumor activity signature’ i.e. top 40 genes positively correlated with sensitivity to DHA across 60 human cancer cell lines, of dihydroartemisinin was statistically determined (y-axis) and then its normalized cellular expression (x-axis) was compared in LICs (mouse, black) and primary Ph+ ALL samples collected at the time of diagnosis (human, white). Calculated $R = +0.592$ where $R$ is the Pearson’s coefficient of correlation and $p<0.001$. 
CHAPTER 5. DISCUSSION AND CONCLUSIONS

The findings of this study reinforce a key therapeutic paradigm that *in vivo* responsiveness to host-derived growth factors may shield autonomous cancer cells from chemotherapy-induced cell killing. Host-derived signaling likely contributes to the frustratingly poor clinical performances of some targeting agents with potent *in vitro* activities. We designed a multi-modality platform that interrogates host-tumor-drug interactions in an unbiased manner. In this forward-building HT method, a logical battery of cell-based screens performed across collections of host-factors, short hairpin (sh)-RNAs and chemical libraries accelerates the identification of host-factor(s) and associated cell-intrinsic molecular change(s) that can impart drug-resistant phenotype(s) and leverages it for the development of improved therapeutic strategies. Applying this HT method to Ph+ ALL, we identified i) IL7 as the dominant cytokine from the tumor-microenvironment that confers remarkable resistance to all FDA-approved BCR-ABL-inhibitors, and ii) that anti-malarial drug DHA can reverse such host-IL7 mediated protection against dasatinib *in vivo*. This opens a novel potential avenue for clinical evaluation to treat human Ph+ ALL patients.

IL7 is essential for human T cell and post-fetal B cell development [109-110] and has a major contribution in the regulation and maintenance of mature T cells [111]. IL7 is constitutively produced by stromal cells in bone marrow, spleen, lymph nodes and thymus, among other tissues [112]. We propose a model in which IL7-rich niches harbor early persistent drug-refractory leukemias in humans [42, 44, 67, 113] and mice [63], with the presence of host-derived IL7 as the key driving factor behind the location of sequestration and the acquisition of resistance.

Results from cytokine-screens (Figure 4.2a) suggested that dasatinib resistance may be a more commonly prevalent host-dependent phenomenon, with IL7 [48, 114-115] as a predominant contributor in Ph+ ALL and other host-cytokines like TSLP [95], IL3 [62, 69, 94], and IL4 playing a minor role. We show that although murine Arf–/– BCR-ABL<sup>WT</sup> LICs bearing, analogous to human Ph+ ALL at diagnosis, are irreversibly committed to cell death on treatment with dasatinib alone [56], cell-extrinsic IL7 reversibly sustains Ph+ ALL, despite therapy that effectively inhibits BCR-ABL *in vivo* [55] by at least temporarily over-riding their dependence on BCR-ABL kinase for sustenance (Figure 4.4) [37, 67-68]. Our demonstration of poor dasatinib responsiveness against BCR-ABL<sup>WT</sup> Ph+ ALL in IL7<sup>+/+</sup> versus IL7<sup>−/−</sup> host-mice (Figure 4.2e) supports a novel role for physiological IL7 in drug resistance *in vivo*. The primary protection by host-IL7, coupled with higher leukemia burden in Ph+ ALL compared to CML in the clinical settings, likely directly contributes to the greater BCR-ABL–KI refractoriness in Ph+ ALL relative to CML.

Leukemogenesis by BCR-ABL oncprotein is a complex process, and depends on constitutive expression of STAT5 [59] and c-MYC [60-62] for initial transformation and sustenance of Ph+ ALL. BCR-ABL either directly or indirectly through JAK2 drives the phosphorylation of STAT5 (pSTAT5) in the cytoplasm. Activated STAT5 (pSTAT5)
translocates to the nucleus and directly upregulates the expression of c-MYC [102]. This was confirmed by our failure to transform cellular bone marrow of STAT5-null mice with p185 isoform of BCR-ABL (data not shown). As demonstrated by others in human Ph+ cells [56, 62, 116], brief BCR-ABL inhibition in BCR-ABLWT LICs by dasatinib (no IL7) significantly decreased the levels of both pSTAT5 protein and expression of MYC and MYC target genes, committing leukemia blasts to death. The specificity of IL7 to attenuate the potency of only BCR-ABL inhibitors in both murine and human Ph+ ALL cells but not of those agents which inhibit targets other than BCR-ABL suggested that IL7 likely rescues a particular signaling pathway downstream of BCR-ABL to impart primary drug resistance. Further mechanistic studies revealed a model in which dasatinib blocks the essential BCR-ABL-JAK2-pSTAT5-c-MYC signaling but presence of cell-extrinsic IL7 rescues the critical expression of c-MYC and its target genes by phosphorylating STAT5 through parallel IL7-JAK3-pSTAT5-cMYC signaling, in a BCR-ABL-independent fashion [102]. The primary IL7-protection against dasatinib is relatively lower than the molecular drug resistance conferred by T315I mutation (Figure 4.2b). Dasatinib does not inhibit BCR-ABL kinase or decrease the cellular level of pSTAT5 or c-MYC in BCR-ABL^{T315I} human cells [117]. Interestingly, STAT activity [117-120] and expression of c-MYC [121] has recently been used as an explanation for resistance to imatinib. c-MYC has also been implicated by others in stroma-mediated drug resistance [66, 122]. Reinforcing our primary drug resistance model (Figure 4.7d), we show that overexpression of c-MYC imparts >100 fold resistance against dasatinib in the absence of both leukemia-extrinsic IL7 and leukemia-intrinsic ABL KD mutations. We propose c-MYC as a cellular node for primary drug resistance against BCR-ABL targeted therapy in Ph+ ALL.

A recent mechanistic study demonstrated direct de-repression of c-MYC by \textit{IKAROS} haploinsufficiency in pre-B cells [103]. Our preliminary experiments depict that modeling the functional loss of \textit{IKAROS} in BCR-ABL^{WT} pre-B LICs by expression of DN \textit{IKAROS} mutant, which increases c-MYC expression, imparts cell-intrinsic refractoriness against dasatinib and imatinib. \textit{IKAROS} haploinsufficiency is a poor human prognostic marker [123] and is detected in leukemia samples obtained (at the time of diagnosis) from a large percentage of Ph+ ALL patients [75, 123] but almost never in chronic phase CML patients [75, 124]. Thus, \textit{IKAROS} haploinsufficiency may also play a role in relative primary BCR-ABL–KI-resistance in Ph+ ALL, similar and in addition to that by host-IL7, which is a non-redundant cytokine present in all human patients.

In summary, repeated BCR-ABL–KI exposures in host-IL7 rich niches result in ‘darwinian’ enrichment of leukemic blasts with relatively higher levels of pSTAT5 and c-MYC, and/or genetic mutations in \textit{IKAROS} and \textit{Arf} loci, accelerating the progression of disease. Selective multi-clonal clonal expansions follow [72, 125], leading to variance in therapeutic responses among human patients, similarly, BCR-ABL-mutations also accumulate with continuous therapy [50-51]. Acquisition of these complex changes impart niche-independent cell-intrinsic drug-resistance and clinically manifest as frank leukemic relapses [72, 125]. In lieu of our findings, high pSTAT5 protein and c-MYC expression level and loss of \textit{IKAROS} in Ph+ lymphoblasts, and elevated levels of host IL7 can potentially be used as surrogates of primary BCR-ABL–KI drug resistance,
pending further preclinical and clinical confirmation. These markers may facilitate personalized medicine in future.

In keeping with critical homeostatic role of IL7 in immune-reconstitution [111], IL7 levels dramatically vary, inverse to individual patient’s absolute lymphocyte counts [126-127]. Current management of Ph+ ALL patients with BCR-ABL–KI-combined protocols deploy maximum tolerated dose (MTD)-guided conventional chemotherapy agents, high-dose corticosteroids, allo-HSCT and immune-suppressants, all of which induce profound iatrogenic-lymphopenias [126-127], and activate the immune IL7-axis [126-127] which can decrease the efficacy of BCR-ABL targeted therapy. Dasatinib therapy induces robust survival-benefit in human Ph+ ALL patients with prolonged lymphocytosis, which may arise from their inherent low levels of IL7, and vice versa [128]. Also, human Ph+ ALL patients with increased levels of pSTAT5, whose phosphorylation is triggered by IL7, respond poorly to imatinib therapy [129]. Collectively, these data strongly support the proposition that host-IL7 protection phenomenon contributes to BCR-ABL–KI-resistance in human patients. This suggests strategic amendments to current leukemia management to avoid prolonged lymphopenias.

From therapeutic perspective, BCR-ABL focused drug discovery aims at achieving more potent BCR-ABL inhibition, and views the escaped kinase activity due to ABL KD mutations, especially T315I, as an integral part of the problem [17-18]. In contrast, our host-tumor-drug interaction studies underscore the necessity for ‘complementing’ pharmacological BCR-ABL-inhibition with strategies/drugs that circumvent primary host-cytokine protection and its associated cellular changes (pSTAT5 levels and c-MYC expression) to accomplish long-lasting responses in Ph+ ALL. A clinically-validated strategy to overcome drug resistance is to concomitantly use multiple agents whose mechanism of action, and therefore of drug resistance, vary significantly, as illustrated by the success of the combination chemotherapy regimens against various cancers and the highly active anti-retroviral therapy (HAART) against HIV [130]. Likewise, we adapted drug resistant phenotypes, mediated by host-IL7 and BCR-ABL T315I mutation, into our flexible cell-based platform and deployed a HT screen across a library of 3200 known drugs and natural products to select active agents, ideal for combined use with BCR-ABL–KIs in vivo.

Our approach accurately confirmed activities of various chemotherapeutic agents and majority of clinically used anti-leukemic drugs, many of which enhance the clinical and preclinical performances of BCR-ABL–KIs [44-45, 63]. In addition, several novel chemotypes were identified to be active against primary host-IL7 and late BCR-ABL mutation mediated drug-resistant phenotypes of Ph+ ALL. These chemically-diverse hits are non-BCR-ABL specific and inhibit cellular targets ‘downstream’ of BCR-ABL or outside its ATP-binding or allosteric site. By definition, such agents cannot be identified by biochemical kinase binding assays or target-drug structural modeling, which form the backbone of target-focused approach. Our results predict that an ideal BCR-ABL–KI should additionally inhibit JAK1/JAK3 kinase activity to prevent relay of drug-refractory signal from the activated IL7 surface receptor to the inside of leukemic lymphoblast. A review of public kinase-binding profiles of imatinib and dasatinib [20] confirms their
inability to inhibit JAK1/JAK3 kinase activity. Until BCR-ABL–KIs with additional JAK3-inhibiting activity become available, empirically determined drug combinations of BCR-ABL–KIs are bound to play an important bridging role in Ph+ ALL therapeutics.

In support of our phenotypic complementation strategy, we characterized the anti-leukemia activities of one screening hit, the orally-bioavailable, well-tolerated antimalarial drug DHA [104], in detail. The biggest challenges in curing Ph+ ALL using BCR-ABL–KIs are poor primary responses, persistence of disease, invariant relapses, and late BCR-ABL mutational resistance. DHA, acting mechanistically independent of BCR-ABL–KIs (Figures 4.14 and 4.15), synergistically corrects the deficiencies associated with dasatinib monotherapy (Figure 4.16b), and achieves near-complete long-term survival in a robust murine model of human Ph+ ALL. Distinct from its anti-malarial drug action [131-132], drug sensitivity to the ARTs family across various cancer types appears to correlate with relative c-MYC expression. While it has been shown that DHA induces apoptosis in cancer cells by selectively accelerating the degradation of c-MYC [133], our preliminary bio-physical studies (Singh and Kriwacki, unpublished), using previously validated assays, indicate that DHA directly binds to basic-helix-loop-helix domain of c-MYC. Most importantly, as described earlier (Figures 4.6 and 4.7), c-MYC is the converging node of primary dasatinib resistance, and targeting c-MYC, using dominant negative approaches, has been validated as a BCR-ABL-independent therapeutic intervention in pre-B Ph+ ALL [60-61]. Thus, DHA might augment BCR-ABL–KI therapy by accelerating the degradation of c-MYC, thereby countering the protective effects of IL7.

The accelerated characterization of anti-malarial drug DHA for a new anti-leukemic use was linked to our original decision of testing only a small but focused collection of very well-characterized bioactive agents and drugs, and their mechanistic evaluation in DTP screening drug databases [89, 134-135]. In past, this strategy has made immense and rapid contributions in the field of mechanistic drug discovery [90-91, 134, 136-137]. In addition to ARTs family, anti-tumor activity of gambogic acid and mycophenolic acid was also found to be statistically correlated with high MYC expression in NCI60 human cancer cell lines (Table 4.1), and could be considered to target dasatinib-resistant Ph+ ALL. However, direct involvement of c-MYC in the mechanism of action of these two alternative agents remains to be experimentally proven [138-140]. Further, Gambogic acid, a natural resin product, has never been tested in humans, and mycophenolic acid is an FDA-approved agent but induces immunosuppression and lymphopenia [138, 141]. In contrast, DHA selectively kills leukemic cells while sparing normal lymphocytes [105-106], therefore is less likely to induce host-IL7. We determined that gene signature predicting sensitivity to DHA across human tumors of 9 different tissue lineages (Figure 4.17) is expressed with a high degree of similarity in murine LICs and primary samples obtained from human Ph+ ALL patients. It is noteworthy that DHA-dasatinib combination is significantly more efficacious than our previous experience with triple combination between dasatinib, dexamethasone and asparaginase [63]. Collectively, these data strongly support the testing of efficacy of BCR-ABL–KI/DHA co-therapy for the clinical management of human patients with high-risk Ph+ ALL.
Complementing BCR-ABL and alternative target-focused drug pipelines [15-16, 19-20, 23] with our host-tumor-drug screening approach can timely predict mechanisms and biomarkers of clinical drug-resistance. A holistic approach encompassing these features can improve the prioritization of hits and facilitate improvement in design of agents against Ph+ ALL and other cancers in future [15-16, 19-20, 23]. Broader implementation of reported screening model will also address two important and inter-related issues which remain pending. First, whether the overly-specific targeting nature of the modern era oncology drugs, relative to the non-specific drugs, renders them more vulnerable to compensatory (cell extrinsic/intrinsic) parallel pathways in the tumor cells, and second, what is the extent of this problem among the current clinically approved agents.
LIST OF REFERENCES


70. Lagas, J.S., et al., Brain accumulation of dasatinib is restricted by P-glycoprotein (ABCB1) and breast cancer resistance protein (ABCG2) and can be enhanced by elacridar treatment. Clin Cancer Res, 2009. 15(7): p. 2344-51.


APPENDIX. SUPPLEMENTAL DATA

Supplemental data is made available online in the form of a Microsoft Excel file which contains eight (i through viii) separate worksheets – i) table of contents (comprised of description, legends and experimental notes for the subsequent worksheets ii through viii which contain real experimental data), ii) host factors, iii) gene expression 1, iv) gene expression 2, v) structure, vi) drug screen summary 1, vii) drug screen summary 2, and viii) ATC classification. The Excel file is approximately 3.8Mbytes and can be downloaded from http://etd.uthsc.edu/World-Access/Singh/Supplementaldata.xlsx.
VITA

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