Using Pharmacology to Squeeze the Life Out of Childhood Leukemia, and Potential Strategies to Achieve Breakthroughs in Medulloblastoma Treatment

Juwina Wijaya, Tomoka Gose, and John D. Schuetz
Department of Pharmaceutical Sciences, St. Jude Children’s Research Hospital, Memphis, Tennessee

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Address correspondence to: John D. Schuetz, Department of Pharmaceutical Sciences, MS 313, St. Jude Children’s Research Hospital, 262 Danny Thomas Place, Memphis, TN 38105. E-mail: John.schuetz@stjude.org
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Abstract—Eliminating cancer was once thought of as a war. This analogy is still apt today; however, we now realize that cancer is a much more formidable enemy than scientists originally perceived, and in some cases, it harbors a profound ability to thwart our best efforts to defeat it. However, before we were aware of the complexity of cancer, chemotherapy against childhood acute lymphoblastic leukemia (ALL) was successful because it applied the principles of pharmacology. Herein, we provide a historic perspective of the experience at St. Jude Children’s Research Hospital. In 1962, when the hospital opened, fewer than 3% of patients experienced durable cure. Through judicious application of pharmacologic principles (e.g., combination therapy with agents using different mechanisms of action) plus appropriate drug scheduling, dosing, and pharmacodynamics, the survival of patients with ALL now exceeds 90%. We contrast this approach to treating ALL with the contemporary approach to treating medulloblastoma, in which genetics and molecular signatures are being used to guide the development of more- efficacious treatment strategies with minimal toxicity. Finally, we highlight the emerging technologies that can sustain and propel the collaborative efforts to squeeze the life out of these cancers.

Significance Statement—Up until the early 1960s, chemotherapy for childhood acute lymphoblastic leukemia was mostly ineffective. This changed with the knowledge and implementation of rational approaches to combination therapy. Although the therapeutics of brain cancers such as medulloblastoma are not as refined (in part because of the blood-brain barrier obstacle), recent extraordinary advances in knowledge of medulloblastoma pathobiology has led to innovations in disease classification accompanied with strategies to improve therapeutic outcomes. Undoubtedly, additional novel approaches, such as immunological therapeutics, will open new avenues to further the goal of taming cancer.
research and created the National Cancer Institute in Frederick, Maryland. Nixon hedged his bets: the initial funding for cancer research was substantial ($1.5 billion), but it was available for only 3 years, and scientific understanding of the disease was still nascent.

Because our knowledge of cancer was so elementary at the time, there was substantial resistance in the scientific community; people considered the investment and promise of a cure to be premature. One vocal critic was Professor Sol Spiegelman, the director of the Institute of Cancer Research at Columbia College of Physicians and Surgeons (Manhattan, NY), who pointedly mocked the campaign by saying, “an all-out effort at this time [to find a cure for cancer] would be like trying to land a man on the moon without knowing Newton’s laws of gravity.” For certain cancers, such as MB, this simile remains apt today. However, for childhood leukemia, cure rates that were in the single digits in 1969 are exceeding 90% for some ALL subtypes today. This achievement was only possible through the concerted and cooperative efforts of many researchers.

This review profiles the advances in therapy, highlighting how key principles of pharmacology were used for more than 55 years, by researchers at St. Jude Children’s Research Hospital (St. Jude), to develop ALL protocols that have proven to be beneficial to many children. The second part of the review highlights the continuing challenges of improving outcomes for MB, the most frequent CNS tumor in children. We selected MB to not only underscore the problems of treating a cancer that arises behind the blood-brain barrier (BBB) but also illustrate how advances in MB treatment can coevolve with advances in our understanding of the pathogenesis of the disease. Contemporary omics studies of MB have propelled the field and led to a rapid expansion in knowledge of MB subgroups, understanding of their oncogenic drivers, and engineering of model systems to aid in the development of new, more effective therapies.

II. ALL: Development and Evolution of Pharmacotherapy to Cure Childhood Leukemias

A. Introduction to ALL

ALL is the most common childhood cancer, accounting for 25% of all cases (Pui, 1995). Despite cure rates exceeding 90% in children, ALL remains a leading cause of morbidity and mortality among them (Pui and Evans, 2013). Patients with ALL typically present with symptoms related to anemia, thrombocytopenia, and neutropenia as the tumor replaces the bone marrow. The malignancy also causes uncontrolled proliferation of abnormal, immature lymphocytes and their progenitors in either the B- or T-cell lineage (Terwilliger and Abdul-Hay, 2017). Symptoms can include frequent infections, fever, easy bruising, bleeding that is hard to stop, and flat, dark-red spots (petechiae) caused by bleeding under the skin. Involvement of the CNS remains a major challenge in clinical management of ALL. The molecular mechanisms and pathways mediating leukemic cell entry into the CNS have not been completely elucidated.

Here, we provide a historic perspective and evolving conceptual basis for curative chemotherapy for ALL, as an example of a potential pharmacologic paradigm: successive improvements in survival, driven mostly by pharmacologic modifications, and tested in randomized clinical trials.

B. Focus on Curing ALL

1. The Drugs. Prior to the use of chemotherapy, blood transfusion (used until the mid-1940s) was the only therapy for ALL, and all it achieved was to delay the inevitable—death caused by hemorrhage and infection. The first blood transfusion administered to a patient with leukemia was performed by George Callender at St. Bartholomew’s Hospital in London in 1873; it was understood that leukemia was a disease of the blood (Piller, 2001). The transfusion provided about 6 weeks of relief for the patient. Blood transfusion as a treatment of acute leukemia was first attempted in the mid-19th century, and the first complete blood transfusion to treat leukemia was given by Bessis and Bernard (1947). The result was a temporary remission that lasted a few months (remission of acute leukemia was almost unheard of at that time). However, ALL was uniformly fatal; most children survived only 2 to 3 months after diagnosis.

Insight into the role of folic acid in hematopoiesis was gained through the efforts of Lucy Wills, a British hematologist, who found that extracts of yeast and liver reduce anemia and leukopenia in pregnant women. She also found that removing this antianemic factor (the so-called Wills factor) from the diet of monkeys severely reduced their red and white blood cell counts (Wills et al., 1937; Jukes, 1987). Knowledge of the Wills factor led to the discovery of folic acid. Because “folic acid concentrate” caused regression of mammary tumors in mice (Leuchtenberger et al., 1945), Sidney Farber, Louis Diamond, and their research assistants administered folic acid to 90 patients with late-stage cancers, including acute leukemia. Unexpectedly, folic acid accelerated the leukemic process to an unprecedented degree, leading to the sudden mortality of all patients. Farber concluded that folic acid conjugates should not be used to treat patients with cancer (Farber et al., 1947). Importantly, he inferred from the observed “acceleration phenomenon” that reducing folic acid levels might suppress the proliferation of malignant cells and restore normal bone marrow function. Farber proposed that a drug that chemically blocks folic acid would inhibit the abnormal marrow production associated with leukemia. In 1947, the antifolate aminopterin was synthesized (Seeger et al., 1947). In 1948, Farber
and Diamond reported that aminopterin induced temporary remission of acute leukemia in children (Farber and Diamond, 1948). Subsequent studies noted that a deficiency of folate acid impairs hematopoiesis (Heinle and Welch, 1948; Farber, 1949). This work offered a new direction for further research on the nature and treatment of acute leukemia (Piller, 2001).

Because manufacturing aminopterin was difficult, the compound was modified to make it easier to synthesize; that modified compound was methotrexate (MTX) (Weinblatt, 2013). MTX interferes with the metabolism of folic acid. The enzyme dihydrofolate reductase (DHFR) converts dihydrofolate into tetrahydrofolate, which is a methyl-group shuttle required for the de novo synthesis of purines, thymidylate, and certain amino acids. After MTX enters a cell by using multiple uptake transporters [reviewed in Lima et al. (2014)], it is polyglutamated, binds DHFR with an affinity 1000-fold greater than that of folate, and competitively inhibits the conversion of dihydrofolate to tetrahydrofolate. Thus, MTX deprives the cell of a necessary cofactor required for the synthesis of thymidylate, a precursor to an essential building block of DNA synthesis (Fig. 1). A healthy dividing cell uses large amounts of reduced folate to maintain the ongoing synthesis of purine and thymidine; for a rapidly dividing malignant cell, the demand is even greater.

The discovery of the benefits of glucocorticoids for treating ALL built upon the findings of Dougherty and White, who demonstrated that administration of adrenocorticotropic hormone (ACTH) reduces the size of most lymphoid tissue (Pufall, 2015). In 1949, cortisone and ACTH were shown to significantly improve rheumatoid arthritis symptoms, which led to the prediction that the combination might be an effective treatment of cancer. Either ACTH or cortisone acetate dramatically reduced the size of lymphoid tumors and leukemias but not other carcinomas (Pearson and Eliel, 1949). Despite their effective reduction of lymphoid disease, glucocorticoids given as single agents did not produce durable remissions, much less a cure. The glucocorticoid prednisone (Pred) is metabolized to prednisolone, which binds to the cytoplasmic glucocorticoid receptor and subsequently translocates to the nucleus (Fig. 1). The mechanism of cell death appears to require glucocorticoid-induced gene transcription, as its effects are blocked by inhibition of transcription by actinomycin D or cycloheximide, indicating a requirement for de novo transcription and translation in the execution of the apoptotic cascade (Smith and Cidlowski, 2010).

One beneficial finding from the development of chemical weapons during World War I was that exposure to mustard gas (i.e., sulfur mustard) depletes cells from the bone marrow and lymph nodes. Goodman and Gilman confirmed the antineoplastic action of nitrogen mustards against a murine lymphoma (Gilman et al., 1946). Drugs were then developed that had the same alkylating properties as the gases, and those agents

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**Fig. 1.** Overview of the mechanisms of action of 6-mercaptopurine, methotrexate, prednisolone, cyclophosphamide, vincristine, and L-asparaginase. HGPRT, hypoxanthine-guanine phosphoribosyl transferase; P450, cytochrome P450.
proved to be potent antitumor agents in clinical trials. Cyclophosphamide (CYCLO) was synthesized in 1959 to release mustard via enzymatic degradation. CYCLO causes DNA alkylation and cross-links to proteins after cytochrome P450-mediated oxidative metabolism to 4-hydroxy-CYCLO (Fig. 1).

Nobel prize winners Gertrude Elion and George Hitchings synthesized another antimetabolite, 6-mercaptopurine (6-MP), in 1951 (Elion et al., 1951) and found that purine nucleotide synthesis was inhibited because of the inhibition of phosphoribosyl pyrophosphate amidotransferase, a rate-limiting step in purine synthesis (Fig. 1). Inhibition of nucleotide synthesis was effective; 6-MP showed promise in acute leukemias and other cancers (DeVita and Chu, 2008). Despite having these unique drugs to target ALL, their effectiveness in inducing remission in patients with leukemia was only temporary when given singly. For the more than 600 patients studied between 1943 and 1952, the median survival was approximately 6 weeks; 95% of the patients did not survive 12 months past diagnosis (MacMahon and Forman, 1957).

In the 1950s, more than 120 alkaloids were discovered in the rosy periwinkle, Catharanthus roseus, including vincristine (VCR) and vinblastine. Vinca alkaloids block cell growth by interfering with microtubule polymerization, which in turn, stops mitosis (Fig. 1). The discovery that vinca alkaloids cause myelosuppression led to a study in mice with leukemia; the lifespan of the animals was dramatically prolonged by the use of a vinca alkaloïd preparation (Johnson, 1968; Gidding et al., 1999).

2. The Total Therapy Approach. From the early 1950s to the early 1960s, Frei and Freireich conducted a series of protocols that evaluated the value of drug and dose combinations. They compared the efficacy of three therapeutic programs for acute leukemia: 1) MTX (phase I) followed by 6-MP (phase II), 2) 6-MP (phase I) followed by MTX (phase II), and 3) 6-MP given with MTX (combination therapy). In children with ALL, combination therapy achieved a superior remission rate (Frei et al., 1961). Donald Pinkel and James Holland, both at the Roswell Park Cancer Institute (Buffalo, NY), and Joe Burchenal at Memorial Sloan-Kettering Cancer Center (New York, NY) also conducted the first systematic combination chemotherapy trials for the treatment of leukemia, primarily in children (Rivera et al., 1993; Schilsky et al., 2006). Their results suggested several new ideas that changed our approach to the disease: 1) Combinations of active drugs with different modes of actions can destroy leukemia tissue in an additive manner that leaves fewer residual leukemic cells, from which a drug-resistant cell population could emerge. 2) Interruption of leukemic cell proliferation at multiple points in the cell cycle reduces the opportunity for drug-resistant cells to form. 3) The combination of three or four drugs might be lethal for leukemic cells that are only partially sensitive to individual drugs. 4) Chemotherapy sustained for 2 to 3 years might more effectively eradicate leukemic cells. This approach would be particularly effective if nonproliferating leukemic cells were triggered to proliferate during remission. 5) The minority (about 20%) of children with lymphocytic leukemia have disease that is sensitive to all antileukemic drugs when they are used sequentially. Thus, simultaneous administration of all effective drugs to these children might destroy all leukemic cells and cure the disease. 6) However, one major challenge to curing ALL was relapse. Relapse resulting in ALL infiltrating the CNS was a major reason for termination of complete remission status (Frei et al., 1961). Additional complications during relapse that posed a risk of morbidity and mortality were bleeding and infections due to a compromised immune system.

In 1962, Donald Pinkel became the first Director at St. Jude and initiated “Total therapy,” a multipronged, curative approach to treating childhood ALL (Simone, 2003). Total therapy consisted of four components: remission induction, CNS-directed therapy with cranial irradiation and intrathecal MTX, intensification (or consolidation) therapy, and continuation treatment. These four components have formed the “backbone” of ALL treatments over the 58-year evolution of Total therapy and remain a mainstay today. Here, we highlight some milestones in that evolution.

a. Era 1 (1962–1967) Total I–IV: increasing survival by reducing myelosuppression. The first era of Total studies (Total I–IV) accrued patients from 1962 to 1967 and empirically employed a variety of chemotherapeutic combinations and dosages of MTX, 6-MP, VCR, Pred, and CYCLO. The multiple mechanisms of action were thought essential to achieve additive or synergistic effects and prevent the development of cells resistant to any one agent (George et al., 1968; Pinkel, 1971; Pinkel et al., 1971, 1972; Simone et al., 1972). One key to the success of this approach was the randomization included in the experimental design. Randomization systematically determines whether new innovations improve established therapies. Total I–IV demonstrated the power of combination chemotherapy, which was far superior to the sequential administration of single agents (Pinkel, 1971; Pinkel et al., 1971). The primary treatment plan consisted of three chemotherapy phases: 1) chemotherapy to induce remission; 2) short-term, high-dosage intensive chemotherapy; and 3) long-term continuation chemotherapy. The first phase was designed to induce remission of leukemia while concurrently minimizing the hazards of gastrointestinal toxicity and bone marrow depression. This was accomplished by a 4-week regimen of Pred and VCR, which had rapid antileukemic effects without the untoward side effects (e.g., vomiting, mucosal ulceration, leukopenia, or thrombocytopenia) of other chemotherapeutics. The second phase was intensive to achieve
complete bone marrow remission; this phase included intravenous MTX followed by intravenous 6-MP and, finally, CYCLO. The third phase was analogous to the approach used to successfully eradicate tuberculosis, which was achieved by prolonged treatment with effective drug combinations. This phase of leukemia treatment consisted of a 2- to 3-year interval of MTX, 6-MP, CYCLO, and VCR.

Giving four agents (MTX, 6-MP, CYCLO, and VCR) in full dosage (Total I–III) often caused dangerous myelosuppression; thus, Total IV (1965–1966) investigated whether such high dosages were necessary. Upon completing the intensive phase and a 2-week rest period, patients were randomized to one of two treatment schedules: half of the children received continuation chemotherapy as maximum dosages of oral 6-MP and a weekly intravenous injection of VCR, MTX, and CYCLO; the other half received the same drugs but at half the dose. In Total IV, the median durations of complete remission and hematologic remission were longer in the full-dosage group than the half-dosage group. The full-dosage chemotherapy was also superior in preventing CNS leukemia. This suggested that higher dosages of drugs might penetrate the BBB and produce drug levels sufficient to delay or block the development of CNS leukemia. Despite this promising result, relapse occurred, usually within 10 months. CNS leukemia was ultimately the major reason for the re-emergence of disease.

b. Eras 2 and 3 (1967–1983) Total V–X: complete remission prolonged by intensive CNS therapy and combination chemotherapy. Overcoming CNS leukemia (despite hematologic remission) was a goal of the Total V study (1967–1968). The therapy included an increased dose of irradiation that was confined to the cranium to avoid bone marrow damage caused by spinal irradiation. To treat the spinal bone marrow and annihilate the lurking leukemia, intrathecal MTX was administered. Like its predecessors, Total V’s induction therapy consisted of Pred and VCR, followed by maintenance with high-dose 6-MP, MTX, and CYCLO. These adjustments to reduce CNS relapse appeared effective; the CNS relapse rate remained significantly lower than that in Total I–IV. Thus, combination chemotherapy and CNS-directed therapy cured approximately 50% of the patients (Aur et al., 1971).

Total VI (1968–1970) aimed to determine, in a controlled manner, the value of “prophylactic” craniospinal irradiation (CSI) to prevent CNS leukemia. Intrathecal MTX was omitted to avoid uncontrolled systemic toxicity posed by the additional chemotherapy. After entering remission, patients were randomized to either receive CSI or not. The majority (95%) of patients who received CSI did not relapse because of CNS leukemia, whereas 65% of those who did not receive CSI did relapse, indicating that prophylactic irradiation was effective in reducing CNS leukemia (Aur et al., 1972).

An additional objective of Total VI was to determine whether an intensive course of chemotherapy early in remission would be beneficial. After patients entered remission, they were randomized to receive either 1 week of high-dosage intravenous chemotherapy composed of 6-MP, MTX, and CYCLO or no additional treatment. All subsequent chemotherapy was the same in all patients. Unfortunately, this modification did not reduce the frequency of relapse or increase the number of survivors and was eliminated from subsequent Total therapy protocols.

Although the primary goal of ALL therapy is to cure the disease, it is also crucial to achieve this by using the least-toxic methods. In Total VII (1970–1971), the efficacy and toxicity of the two forms of effective CNS prophylactic therapy were compared. Patients were randomized to receive either CSI and five doses of intrathecal MTX (as in Total V) or CSI (as in Total VI). Both forms of therapy were equally effective in preventing CNS leukemia; however, during the first 3 months of remission, the frequency of leukopenia and temporary interruption of chemotherapy was significantly greater in those who received CSI. Thus, toxic side effects and complications were associated with both therapies, and intrathecal MTX appeared ineffective at preventing CNS ALL relapse (Aur et al., 1973).

Although Pred and VCR induced remission, whether additional periodic (or pulse) reinduction treatments during remission would be of further therapeutic benefit was unknown. Patients were randomized to receive either three weekly doses of VCR and 15 days of Pred every 12 weeks or standard treatment (i.e., daily 6-MP and MTX and weekly CYCLO). The added pulses of VCR and Pred were of no significant value (Simone et al., 1972).

By the end of Total VII, the probability of long-term survival had increased to 50%. However, treatment of ALL still left much to be desired: it was largely empirical; it did not prevent hematologic relapse in as many as 66% of patients; and its acute side effects ranged from merely annoying to fatal. Furthermore, the treatment had latent, long-term detrimental effects on multiple organs (e.g., gonads, brain).

Total VIII (1972–1975) was aimed at further refining the antileukemogenic effectiveness of combination therapy. The rationale was that a combination of agents might target different inherent vulnerabilities in a population of leukemic cells. However, a caveat was that the toxicity might be additive and therefore limit the dose of each agent. The question asked by this study was, “Should one give only the most effective agent(s) in relatively high dosage or try to achieve a broader spectrum of action by using a combination of agents, even though the tolerable doses of each would reduce toxicity?” The efficacy of 1 (MTX), 2 (MTX plus 6-MP), 3 (same as group 2 plus CYCLO), and 4 (same as group 3 plus arabinosyl cytosine) therapies during remission...
was evaluated. The results showed the relapse rate in patients given MTX alone was significantly greater than that of the other groups receiving combination therapy. At the conclusion of era 2, therapy modifications that reduced CNS leukemia also significantly increased event-free survival.

The beginning of era 3 in the late 1960s was highlighted by studies showing that high dosages of MTX (HDMTX) promote remission among patients with relapsed ALL (Djerassi et al., 1967). In support of this, Wang et al. (1976) demonstrated that high levels of MTX in cerebrospinal fluid (CSF) suppress CNS leukemia. Total X (1979–1983) showed that by administering HDMTX, cranial irradiation could be reduced (Abromowitch et al., 1988). To ameliorate the hematopoietic and gastrointestinal toxicity of HDMTX, leucovorin (5-formyl-tetrahydrofolate) was administered; this treatment was termed “leucovorin rescue.” However, leucovorin rescue was delayed many hours after MTX was given because concurrent administration might be antagonistic (MTX and leucovorin enter the cell by the same transporter, the reduced folate carrier). The success of the delay strategy is based on leukemia cells inherently forming more MTX polyglutamates (MTX-PGs), which are more potent inhibitors of DHFR and thymidylate synthase than MTX. In contrast, susceptible host tissues (e.g., the bone marrow and gastrointestinal tract) accumulate fewer MTX-PGs, which are readily displaced by the flood of the leucovorin-derived natural folate polyglutamates. Leucovorin rescue remains a key element of HDMTX therapy today.

Although HDMTX is an effective treatment of ALL, it can cause significant toxicity, especially acute kidney injury. MTX is primarily cleared by renal excretion. MTX and its metabolites can precipitate within the renal tubules, causing crystal nephropathy and, subsequently, HDMTX-associated nephrotoxicity. Because MTX is acidic, it does not form crystals when the urine has an alkaline pH; thus, alkalinization greatly increases MTX solubility and excretion. To minimize renal toxicity during HDMTX therapy, supportive care (e.g., intense hydration and urinary alkalinization) are employed to enhance the drug’s solubility.

**c. Era 4 (1984–1991) Total XI–XII: dose intensification and individualized pharmacokinetic analysis to improve outcome.** The eradication of drug-resistant ALL was hypothesized to require an increased intensity of induction therapy (i.e., higher doses, resulting in fewer surviving treatment-resistant leukemic cells). The premise of this study was based on the Goldie-Coldman hypothesis (Goldie et al., 1982), which related the development of drug-resistant cells to the tumor size and its mutation rate. Thus, chemotherapeutic elimination of a tumor required agents with nonoverlapping mechanisms of action, given at their maximum cytotoxic dose. The addition of teniposide (TEN) and cytarabine (Ara-C) to the arsenal was based on the effective cytoreduction achieved in patients with high-risk leukemia (Dahl et al., 1987). Thus, in Total XI (1984–1988), early treatment intensification improved outcome and increased the 5-year event-free survival to 70% (Rivera et al., 1991).

In Total XII (1988–1991), the benefits of therapy guided by individualized pharmacokinetics was tested. The rate of clearance of some antileukemic agents (e.g., MTX, TEN, and Ara-C) varies as much as 10-fold among children with ALL. Patients (postremission) were treated by either conventional therapy (i.e., dosing based on body surface area) or individualized therapy (i.e., dosing based on the patient’s rate of drug clearance). In the individualized-therapy group, those with rapid clearance received higher doses, and those with very slow clearance received lower doses. The study showed no significant difference between the two treatments for patients with T-lineage leukemia. However, patients with B-lineage leukemia who received individualized therapy had significantly better outcomes than those who received conventional therapy. Thus, for childhood B-lineage ALL, increasing MTX dosage in patients with rapid clearance improved their outcome without increasing toxicity (Evans et al., 1998).

Long-term anticonvulsant therapy increases the systemic clearance of several antileukemic agents and is associated with lower efficacy of chemotherapy. Thus, to prevent reduced therapeutic efficacy, alternatives to enzyme-inducing anticonvulsants are recommended for patients receiving chemotherapy for ALL (Relling et al., 2000). The failure of clearance-adjusted therapy to improve the outcome in T-lineage ALL cases might have been due to inadequate MTX levels. Several studies have shown that T-lineage blasts accumulate MTX and MTX-PGs less avidly than do B-lineage blasts and suggested that higher doses might increase MTX accumulation (Barredo et al., 1994; Synold et al., 1994; Galpin et al., 1997). Indeed, the responsiveness of B-lineage ALL to MTX, compared with that of T-lineage ALL, appears to be related, in part, to folypolyglutamate synthetase (FPGS), an enzyme that metabolizes natural folates and a broad range of folate antagonists to polyglutamate derivatives. FPGS activity in blasts from children with newly diagnosed ALL is significantly higher in B-lineage ALL (188%) than in T-lineage ALL (37%) (Galpin et al., 1997). Higher FPGS activity in B-lineage blasts could also explain the superior outcome in children with B-lineage ALL treated with antimetabolite therapy. Notably, in B-lineage ALL blasts, the level of MTX-PGs is significantly related to factors beyond MTX dose, and such factors might account for the increased accumulation of MTX-PGs. These factors could be lymphoblast ploidy (hyperdiploid > nonhyperdiploid) and the percentage of cells in S-phase (Synold et al., 1994) with higher FPGS and lower DHFR (Galpin et al., 1997).
d. Eras 5 and 6 (1991–2007) Total XIII–XV: reducing CNS relapse and individualizing treatment with pharmacogenomic analysis. Total XIIIIA (1991–1994) and Total XIIIIB (1994–1998) studies showed reduced relapse rates by further individualizing therapy based on disease stratification and pharmacogenetics. Total XIIIIA employed early intensification of intrathecal chemotherapy in patients with high-risk ALL (i.e., 22% of the total study population). This approach reduced the risk of CNS relapse to about 1% and boosted the 5-year event-free survival to 80%. However, although cranial irradiation reduces CNS relapse, it causes secondary side effects, including neurocognitive deficits and endocrinopathies (Pui et al., 2003).

In Total XIIIIB, treatment was modified to improve clinical outcome and reduce untoward long-term side effects. Instead of giving Pred during postremission therapy, dexamethasone (Dex) was substituted. The substitution was because Dex is superior to Pred in relation to reduced frequency of recurrence of CNS and systemic disease (Bostrom et al., 2003). Total XIIIIB was at least as effective as Total XIIIIA (5-year event-free survival, 80.8% ± 2.6% vs. 77.6% ± 3.2%). Only 12% of the patients in Total XIIIIB received CSI, compared with 22% of those in Total XIIIIA. The efficacy of early intensification of intrathecal therapy (simultaneously administered MTX, hydrocortisone, and Ara-C) was confirmed, thereby paving the way to eliminating prophylactic cranial irradiation.

Total XIIIIB also demonstrated that host pharmacogenetics affects the pharmacodynamics of chemotherapy and antileukemic outcome. It was the first leukemia trial to apply pharmacogenetics during therapy to guide dosing. 6-MP is an immunosuppressant and antineoplastic agent. To reduce hematopoietic toxicity and decrease risk of secondary cancers (Pui et al., 2004), 6-MP was carefully dosed in Total XIIIIB. The dosage of 6-MP given during continuation therapy was based on polymorphisms of the gene encoding thionine methyltransferase (TPMT). TPMT inactivates 6-MP and circumvents the formation of thioguanine nucleotides, the major active metabolites. Among patients who inherit defective TPMT alleles, severe myelosuppression was frequently observed, even among patients receiving standard doses of 6-MP, because of the high levels of thionine nucleotides. Patients with defective TPMT alleles received lower doses of 6-MP to reduce risk of secondary cancer in these patients (Rocha et al., 2005; Relling et al., 2006; Abaji and Krajnovic, 2017).

Total XIV (1998–1999) attempted to find optimal doses of MTX for various subtypes of ALL while also determining pharmacogenetic factors that affect MTX sensitivity. Methylenetetrahydrofolate reductase (MTHFR) catalyzes the reduction of 5,10-methylenetetrahydrofolate to 5-methyltetrahydrofolate. This study hypothesized that in patients with MTHFR deficiency, hyperhomocysteinuria occurs more frequently after MTX treatment. Additionally, genetic polymorphisms in the reduced folate carrier, the transporter for cellular uptake of reduced folates (e.g., leucovorin), and MTX might cause MTX toxicity or poor response to therapy. CSF levels of homocysteine were transiently elevated in children treated with HDMTX, suggesting that HDMTX is related to seizure risk in children with leukemia. However, the plasma and CSF concentrations of homocysteine did not differ by MTHFR or reduced folate carrier genotypes, suggesting that these gene variations are unrelated to MTX sensitivity (Kishi et al., 2003).

The St. Jude Total therapy program for childhood ALL spans seven eras (Table 1). Between 1958 and 1962, before Pinkel opened the first Total therapy study, only 3% of patients with ALL survived 5 years. After continued investigation of the optimal use of antileukemic agents, improved supportive care, and precise risk assessment in several contemporary trials, the Total XV (2000–2007) study achieved a 5-year event-free survival for childhood ALL of more than 85% and 5-year overall survival of more than 90% (Fig. 2). Notably, steps to prevent CNS leukemia were revealed during era 2, when craniospinal irradiation and therapeutic intensification after remission induction boosted cure rates to near 70% (Pui and Evans, 2013). Further efforts revealed that the early intensification of intrathecal therapy substantially reduced CNS relapse hazard (era 5). Finally, during era 6, with effective risk-adjusted chemotherapy, prophylactic cranial irradiation could be safely omitted from the treatment of childhood ALL. Taken together, these studies have transformed this once uniformly fatal cancer into one with a cure rate approaching 90% (Fig. 2). This is one of the pivotal success stories of modern medicine.

![Fig. 2. Kaplan-Meier analysis of survival of 2852 children with newly diagnosed ALL who were enrolled in 1 of 15 consecutive Total therapy studies at St. Jude from 1962 to 2007. Ten-year survival estimates are shown. The results demonstrate steady improvement in outcome over the past half century. The mean (± S.E.) 5-year survival probabilities are shown. Adapted from Pui and Evans (2013).](image-url)
<table>
<thead>
<tr>
<th>Era (y)</th>
<th>Total Therapy</th>
<th>Remission Induction</th>
<th>Intensive Chemotherapy</th>
<th>Continuation</th>
<th>CNS Therapy</th>
<th>Reinduction</th>
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<td>I</td>
<td>Pred, VCR</td>
<td>None</td>
<td>MTX or 6-MP, CYCLO, VCR</td>
<td>CSI</td>
<td>None</td>
<td></td>
</tr>
<tr>
<td>II</td>
<td>Pred, VCR</td>
<td>None</td>
<td>MTX, 6-MP, CYCLO, VCR</td>
<td>CSI</td>
<td>None</td>
<td></td>
</tr>
<tr>
<td>III</td>
<td>Pred, VCR</td>
<td>MTX, 6-MP, CYCLO</td>
<td>MTX, 6-MP CYCLO, VCR</td>
<td>CSI</td>
<td>None</td>
<td></td>
</tr>
<tr>
<td>IV</td>
<td>Pred, VCR</td>
<td>MTX, 6-MP, CYCLO, Pred, VCR</td>
<td>MTX, 6-MP CYCLO, Pred, VCR</td>
<td>CSI</td>
<td>Pred, VCR</td>
<td></td>
</tr>
<tr>
<td>2 (1967–1979)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>V</td>
<td>Pred, VCR</td>
<td>MTX, 6-MP, CYCLO</td>
<td>MTX, 6-MP CYCLO, Pred, VCR</td>
<td>CSI</td>
<td>Pred, VCR</td>
<td></td>
</tr>
<tr>
<td>VI</td>
<td>Pred, VCR, DNR</td>
<td>MTX, 6-MP, CYCLO vs. none</td>
<td>MTX, 6-MP CYCLO, Pred, VCR</td>
<td>CSI vs. none</td>
<td>None</td>
<td></td>
</tr>
<tr>
<td>VII</td>
<td>Pred, VCR</td>
<td>None</td>
<td>MTX, 6-MP, CYCLO vs. Pred, VCR</td>
<td>CSI</td>
<td>None</td>
<td></td>
</tr>
<tr>
<td>VIII</td>
<td>Pred, VCR, ASNase</td>
<td>None</td>
<td>MTX, 6-MP, CYCLO, Pred, VCR</td>
<td>CSI, MTX vs. CSI</td>
<td>None</td>
<td></td>
</tr>
<tr>
<td>3 (1979–1983)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>IX</td>
<td>Three groups</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1. Pred, VCR, DNR</td>
<td>MTX vs. CSI, MTX</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. Pred, VCR, DNR, ASNase, TEN, Ara-C</td>
<td>MTX, 6-MP vs. MTX, 6-MP, CYCLO, Dox, TEN, ara-C</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. Pred, VCR, DNR, ASNase, TEN, Ara-C, MTX</td>
<td>MTX, 6-MP vs. MTX, 6-MP, CYCLO, Dox, TEN, ara-C</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>XI</td>
<td>Pred, VCR, DNR, ASNase, TEN, Ara-C</td>
<td>MTX</td>
<td>MTX, Hyd, Ara-C (+ cranial irradiation in higher risk)</td>
<td>CSI</td>
<td>None</td>
<td></td>
</tr>
<tr>
<td>XII</td>
<td>Pred, VCR, DNR, ASNase, TEN, Ara-C</td>
<td>MTX, 6-MP, TEN, Ara-C</td>
<td>MTX, Hyd, Ara-C (+ cranial irradiation in higher risk)</td>
<td>CSI</td>
<td>None</td>
<td></td>
</tr>
<tr>
<td>XIIIA</td>
<td>MTX (preinduction), Pred, VCR, DNR, ASNase, EP, Ara-C</td>
<td>MTX, 6-MP</td>
<td>MTX, Hyd, Ara-C (+ cranial irradiation in higher risk)</td>
<td>CSI, MTX</td>
<td>None</td>
<td></td>
</tr>
<tr>
<td>XIIIB</td>
<td>MTX (preinduction), Pred, VCR, DNR, ASNase, EP, Ara-C</td>
<td>MTX, 6-MP</td>
<td>MTX, Hyd, Ara-C (+ cranial irradiation in higher risk)</td>
<td>CSI, MTX</td>
<td>None</td>
<td></td>
</tr>
<tr>
<td>6 (2000–2007)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>XV</td>
<td>MTX (preinduction), Pred, VCR, DNR, ASNase, Ara-C, CYCLO, 6-MP (Imatinib; Ph-positive)</td>
<td>MTX, 6-MP</td>
<td>MTX, Hyd, Ara-C</td>
<td>CSI, MTX</td>
<td>None</td>
<td></td>
</tr>
<tr>
<td>7 (2007–)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>XVI</td>
<td>Pred, VCR, DNR, ASNase, CYCLO, Ara-C, TG</td>
<td>MTX, 6-MP</td>
<td>MTX, Hyd, Ara-C</td>
<td>CSI, MTX</td>
<td>None</td>
<td></td>
</tr>
<tr>
<td>XVII*</td>
<td>Pred, VCR, DNR, ASNase, CYCLO, Ara-C, 6-MP</td>
<td>MTX, 6-MP, Ara-C, VCR, ASNase, CYCLO</td>
<td>MTX, Hyd, Ara-C</td>
<td>CSI, MTX</td>
<td>None</td>
<td></td>
</tr>
</tbody>
</table>

ASNase, asparaginase; DNR, daunorubicin; DOX, doxorubicin; EP, etoposide; Hyd, hydrocortisone; 6-MP, 6-mercaptopurine; TG, thioguanine.

*Additional targeted therapy including tyrosine kinase inhibitors, immunotherapies, and CAR T-cell therapy.
3. Improving Therapy by Understanding the Underlying ALL Biology.

a. ALL and minimal residual disease. Advances in therapy have been facilitated by both technology and an improved understanding of leukemia biology. ALL can be stratified into two main immunophenotypes identified by distinct hematopoietic lineage markers: B-ALL and T-ALL (Huntly and Gilliland, 2005). These immunophenotypes affect response to therapy, which is notable because pediatric B-ALL comprises as many as 85% of cases; T-ALL comprises the remaining 15% and is more common among male patients (Hunger and Mullighan, 2015). One crucial finding was that a submicroscopic amount of leukemia remains after remission. In 1981, immunodetection of lymphoid surface antigens by indirect immunofluorescence and antisera to human differentiation-linked antigens was used to identify residual leukemic blasts in bone marrow samples from some patients with ALL. Ultimately, multiparameter flow cytometry capable of detecting 1 in 10,000 leukemic cells revealed that this “residual” leukemia at any point after remission is predictive of relapse. This condition is referred to as minimal residual disease (MRD). Importantly, genetic abnormalities have been correlated with detectable disease at the end of remission, and the detection of MRD has become an important diagnostic assessment tool to determine the likelihood of relapse and risk stratification (Ravandi et al., 2018).

b. Genetic alterations influence ALL outcome. To determine whether the genome is altered in leukemia, it was crucial to know the accurate number of chromosomes. The ability to stratify ALL by the DNA content (i.e., ploidy) is an important feature in risk assessment. In 1978, Secker-Walker et al. (1978) classified leukemia cases into five categories based on the chromosomal characteristics of the major proportion of leukemic cells present: hyperdiploid, pseudodiploid, diploid, hypodiploid, and mixed. Patients in the hyperdiploid category had a significantly longer duration of first remission than did those in all of the other categories, and those in the pseudodiploid category had the shortest, suggesting that the proportion of hyperdiploid cells, as determined by conventional chromosomal-staining methods, can be used as an additional prognostic feature in childhood ALL.

The Philadelphia (Ph) chromosome was the first genomic alteration in leukemia that was reported; the Ph chromosome was detected in myeloid leukemia (Nowell et al., 1960). Reciprocal translocation of genetic material between chromosomes 9 and 22 produces the fusion gene breakpoint cluster region-Abelson (BCR-ABL) 1 (Rowley, 1973). The fusion of the ABL kinase with BCR produces a cytosolic kinase that is an oncoprotein with constitutive kinase activity that promotes leukemogenesis by driving cell proliferation and blocking cell death (Kurzrock et al., 1988). For many years, BCR-ABL was thought to occur only in myeloid leukemias; however, Propp and Lizzi (1970) reported an adult with ALL whose marrow cells expressed a high percentage of Ph chromosomes.

Cytogenetic analysis was crucial to identifying gross chromosomal alterations in many childhood ALL cases. In 1984, cytogenetic analysis of leukemic cells obtained from 122 pediatric patients with ALL identified chromosomal translocations in 36 cases. Among these, two new immunophenotype-specific chromosomal translocations, t(11; 14) in T-cell ALL and t(1; 19) in pre-B–ALL, were associated with specific immunophenotypes of the disease. Consistent with translocations in Burkitt lymphoma, B-cell ALL, and Ph+ ALL, the new chromosomal rearrangements identified in pre-B–cell ALL and T-cell ALL represented specific anomalies involving defined cytogenetic loci. In cases with either t(1; 19) or t(11; 14), specific break points were consistently found, suggesting that genes at or near these sites have a role in leukemogenesis, leukemic cell proliferation, or both (Williams et al., 1984).

Today, we recognize multiple ALL subtypes with distinct somatic genetic alterations, including aneuploidy (changes in chromosome number); chromosomal rearrangements that deregulate gene expression or result in expression of chimeric fusion proteins, deletions, and gains of DNA; and DNA-sequence mutations (Fig. 3) (Harrison, 2009). The advent of high-resolution whole-genome and whole-transcriptome sequencing permits further classification of ALL according to its specific genetic abnormalities. Prior to these approaches, key surface proteins were identified to aid ALL stratification and classification. Importantly, a child’s ALL genome harbors 10–20 nonsilent coding mutations at diagnosis, but this number increases almost 2-fold at relapse (Ma et al., 2015). Many of these mutations perturb key cellular processes, including the transcriptional regulation of lymphoid development and differentiation; cell cycle regulation; the TP53-retinoblastoma protein tumor-suppressor pathway; growth factor receptor, Ras, phosphatidylinositol 3-kinase, and Janus kinase-signaling transducers and activators of transcription (JAK-STAT) signaling; nucleoside metabolism; and epigenetic modification. Perturbation of the latter two processes is common at relapse (Hunger and Mullighan, 2015).

Applying genomics to improve therapy. An ALL with a gene-expression profile similar to that of ALL with the characteristic BCR-ABL translocation is referred to as Philadelphia chromosome–like acute lymphoblastic leukemia (Ph-like ALL); this form of the disease represents about 3% to 4% of pediatric ALL cases and about 25% of adult ALL cases. Ph-like ALL is characterized by genetic alterations that activate cytochrome receptor and kinase signaling. Before the advent of tyrosine kinase inhibitors (TKIs), Ph-like ALL was associated with a very poor prognosis, despite intensive
chemotherapy and frequent hematopoietic stem cell transplantation (Aricò et al., 2010). The development of TKIs revolutionized the therapy of Ph-like ALL. Early studies using the first-generation ABL1-class TKI, imatinib, as monotherapy in Ph-like ALL were disappointing; initial responses rapidly progressed to TKI-resistant disease. When imatinib was added to intensive chemotherapy, the survival of children with Ph-like ALL dramatically increased. In the years since the initial success of imatinib, second-generation (i.e., nilotinib, dasatinib, bosutinib) and third-generation (ponatinib) ABL1-class TKIs have been developed that are active against multiple imatinib-resistant BCR-ABL mutants (Bernt and Hunger, 2014). The standard of care for the frontline treatment of Ph+ ALL consists of combinations of cytotoxic chemotherapy with TKI. Examples of clinical trials of TKI-based therapies for children, adolescents, and adults with Ph-like ALL are shown in Table 2.

One strategy to improving therapeutic efficacy is to consider how the genetic variation in host genes influences host toxicity. This area of pharmacogenomics has dramatically expanded over the past 20 years. For example, polymorphism in the TPMT, NTP diphosphatase, or ABCC4 (MRP4) gene accounts for why some patients suffer hematopoietic toxicity caused by the thiopurine regimen and others do not (Relling et al., 1999; Krishnamurthy et al., 2008; Moriyama et al., 2016). Some of these genes are considered when determining the dose of 6-MP to administer to patients in the Total studies.

d. Immunotherapy, an emerging treatment modality. Although the Total studies have proven that small molecules are efficacious in treating leukemia, immunotherapy is emerging as a promising modality. There are several immunotherapy-based approaches: bispecific T-cell engager (BiTE), antibody-drug conjugates, and most recently approved for leukemia, chimeric antigen receptor (CAR) T-cell therapy. Blinatumomab (Blincyto; Amgen), a bispecific monoclonal antibody, is a first-in-class targeted immunotherapy agent for the treatment of B-cell malignancies. Blinatumomab simultaneously binds CD19 expressed on the surface of ALL cells and CD3 expressed on T cells. The bridge formed by blinatumomab binding brings the T cells close enough to the ALL cells to recognize and kill them. Clinical trials have demonstrated its efficacy in relapsed B-cell ALL and B-cell non-Hodgkin lymphoma, including in patients who are refractory to chemotherapy (Kantarjian et al., 2017; Gökbuget et al., 2018).

Fig. 3. Cytogenetic and molecular genetic abnormalities in childhood ALL. Abnormalities that are exclusively seen in T-cell–lineage ALL are shown in pink. Abnormalities that are commonly seen in B-cell–lineage ALL are shown in blue. ABL, Abelson; BCR, breakpoint cluster region; CRLF2, cytokine receptor-like factor 2; ERG, ETS transcription factor ERG; ETP, early T-cell precursor; ETV6, ETS variant transcription factor 6; iAMP21, intrachromosomal amplification of chromosome 21; LYL1, lymphoblastic leukemia derived sequence 1; MLL, mixed lineage leukemia; PBX1, pre-B-cell leukemia transcription factor 1; RUNX1, runt-related transcription factor 1; TAL1, T-cell acute lymphocytic leukemia protein 1; TCF3, transcription factor 3; TLX1, T cell leukemia homeobox 1; TLX3, T cell leukemia homeobox 3.
Another strategy is drug delivery by antibody. Inotuzumab ozogamicin (InO; Besponsa), is a CD22 monoclonal antibody conjugated with calicheamicin, a natural product derivative that targets and produces double-stranded breaks in DNA (Kantarjian et al., 2012, 2013). CD22 is detected on leukemic blasts in more than 90% of patients with ALL. After the antibody portion binds to CD22 receptors on the surface of leukemic cells, the entire InO-calicheamicin conjugate enters the cell. The antibody-drug conjugate is transported through endosomes to the lysosomes, where the acidic environment hydrolyzes the inactive inotuzumab into ozogamicin. Ozogamicin is then activated by intracellular glutathione and travels to the nucleus to induce cell cycle arrest or cell death. InO was recently introduced as a targeted therapy for relapsed or refractory CD22+ B-cell precursor ALL (DeAngelo et al., 2017; Yurkiewicz et al., 2018).

CAR T-cell therapy is an emerging molecular hack of the immune system. Killer T cells are obtained from a patient and engineered to express a chimeric receptor, which in simple terms has two parts: a targeting domain that recognizes a protein on the surface of the patient’s cancer cells and an activation domain. When CAR T cells bind their targeted antigen, they elicit a cytotoxic effect to trigger cancer cell death (Sadelaín et al., 2016). Tisagenlecleucel (Kymriah;Novartis) is an autologous CD19-targeted CAR T-cell product approved for treatment of relapsed or refractory B-cell ALL and lymphoma (Maus and Levine, 2016). In a global study of CAR T-cell therapy, a single infusion of tisagenlecleucel provided durable remission and long-term persistence in pediatric and young adult patients with relapsed or refractory B-cell ALL, with transient high-grade toxic effects (Maude et al., 2018). The SJCAR19 trial (NCT03573700) for ALL in children and young adults is currently underway at St. Jude. These novel immunotherapies have opened an exciting new avenue of anticancer therapy with the potential to transform the lives of many patients whose disease was hitherto incurable.

### Future outlook

With increasing knowledge of biology, improved methods to detect and monitor disease, and the discovery of new therapies, it does not come as a surprise that the next generation of clinical trials and therapy combines all of these advances, and St. Jude appears at the forefront. The evolution of Total therapy over the years exemplifies St. Jude’s ongoing commitment to improving patient outcomes, and it continues to be reflected in the latest protocols, including Total XVI and XVII. In Total XVI, to improve systemic and CNS disease control, the protocol included delivering chemotherapeutic drugs to the cerebrospinal fluid earlier in the treatment regimen. This resulted in reduction of CNS relapse from 5.7% in Total XV to 1.8% in Total XVI (Jeha et al., 2019). Showing St. Jude’s commitment to improving patients’ quality of life, this regimen spared children of prophylactic cranial radiation, even for patients with high-risk disease. In Total XVII, patients with standard or high-risk disease with more than 1% MRD at the end of remission will receive additional targeted therapy based on the genetics of the disease. Patients with refractory B-ALL will receive the immunotherapy blinatumomab or CAR T-cell therapy and proteasome inhibitor bortezomib, whereas those with alterations in the tyrosine kinase pathway will have TKIs added to their treatment regimen (NCT03117751). Importantly, results from prospective trials continue to be leveraged to improve patients’ quality of life. In Total XVII, patients with high-risk centrosomal protein 72 (CEP72) TT genotype (Diouf et al., 2015) will receive reduced doses of VCR in an effort to reduce both the incidence and severity of VCR-induced peripheral neuropathy. Total XVII represents the next generation of therapy, as it not only builds on the past advances but also incorporates modern strategies, including genomic, biologic, and pharmacologic understanding of disease status to customize treatments that will improve survival and patients’ quality of lives.

Looking forward, several questions arise: What is the optimal sequence to treat patients; should the usually less toxic immunotherapy be used first and the more toxic chemotherapy be reserved? Can multiple monoclonal antibodies be incorporated into one regimen? Moreover, in the current treatment era, which is evolving toward precision medicine, can we replace the adverse effects of chemotherapy with a single CAR T-cell infusion? Moreover, with the speed at which technology is complementing human intelligence, can
we exploit it to detect disease earlier, predict treatment response, and give patients the best chance for a good outcome? Without a doubt, we will have hints to answer these questions in the upcoming years thanks to numerous collaborative efforts across disciplines and institutions. Ultimately, all researchers are committed to serving and contributing what is best for the patients.

III. Medulloblastoma

A. Introduction to Medulloblastoma

Medulloblastoma (MB) is an embryonic tumor found in the posterior fossa (Rutka and Hoffman, 1996). It predominantly arises in the cerebellum and spreads to the fourth ventricle. Patients typically experience cerebellar symptoms, such as headache and incoordination, for 2 to 3 months before diagnosis (Raffel, 2004). Approximately 300–400 new cases of childhood MB occur in the United States each year, representing 9.2% of pediatric tumors. Thus, MB is one of the most common malignant brain tumors in children (Johnson et al., 2014). However, until the late 1970s, the World Health Organization's classification of childhood brain tumors was incapable of accurately discriminating between various childhood brain tumors (Rorke et al., 1985).

Here, we will describe how our understanding of MB has expanded over the years and how accurate classification of MB subtypes was essential to the development of preclinical MB disease models in which new therapeutic strategies for MB could be tested (Table 3). The identification of MB subgroups and origins of the disease has rapidly progressed recently. Although this knowledge has the potential to help identify features that are important to therapeutic response, progress in that area lags. This lag might be due to a combination of factors, such as fewer patients, more disease heterogeneity, and accessing the CNS tumor with effective agents that can cross the BBB. Nonetheless, therapy for MB is advancing.

B. MB Classification to Understand Disease and Improve Therapy

In 1969, Chu Chang, Edgar Housepian, and Charles Herbert introduced the Chang staging system, which used tumor size and metastatic status to categorize MBs (Chang et al., 1969). Chang stage T1 and T2 tumors are smaller than 3 cm in diameter and are seen infrequently, whereas T3 and T4 tumors are larger lesions that invade adjacent structures or have spread. Additionally, the team introduced the M0–M4 metastasis grading system (Chang et al., 1969), which was invaluable because MB has the propensity to metastasize; approximately 30% of children with MB present with CSF metastasis at diagnosis (Johnson et al., 2014). For CNS tumors, the Chang staging system has been important for stratifying therapy. However, in the 1980s, MB tumors were still not clearly understood by neuropathologists, as evidenced by the following comment: "In fact, they were classified together with primitive neuroectodermal tumors because they were primarily in the cerebellum with similar histological features" (Rorke et al., 1985). This unclear MB classification probably affected the disease's response to therapy. By 2007, all CNS tumors were reclassified by the World Health Organization. MB tumors were stratified into four histologic subtypes: classic, desmoplastic, MB with extensive nodularity, and large-cell/anaplastic (Louis et al., 2007). Tumor histology has been associated with clinical outcome. For example, patients with large-cell/anaplastic tumors have significantly poorer survival than those whose MB has classic histology (Lamont et al., 2004).

C. Evolution of Therapy and Impact on Mortality

1. Surgery and Radiation Therapy. In 1925, Cushing began surgically removing MBs. Despite a staggering surgical mortality rate of more than 30%, the patients who survived the procedure experienced increased survival to 17 months, compared with 6 months without tumor resection (Rutka and Hoffman, 1996). In 1953, Patterson and Farr reported a breakthrough in MB therapy: 3-year survival of 65% in patients who received CSI (Jenkin, 1996). Although the long-term cognitive side effects can be devastating, radiation therapy remains the standard of care for MB today.

As was shown for ALL, risk stratification of MB improved therapeutic outcomes. Which MB tumors must be treated aggressively? Important prognostic factors (i.e., age, amount of tumor remaining post-surgery, and the extent of metastasis, as classified by the Chang M stage) are used to assign patients to either standard- or high-risk therapy (Fig. 4A). Patients with standard-risk MB have residual tumor smaller than 1.5 cm² and no metastases; these patients receive a reduced dose (23.4 Gy) of radiation, a localized irradiation boost to the posterior fossa region, and adjuvant chemotherapy. Patients with high-risk MB have residual tumors larger than 1.5 cm² and evidence of metastases; these patients receive 36 Gy radiation, a localized irradiation boost to the posterior fossa, and more-aggressive adjuvant chemotherapy (Gilbertson, 2004). Patients in the standard-risk group have an average 5-year survival of 75%, compared with less than 50% for those in the high-risk group (Gilbertson, 2004). What are the therapeutic implications of such a simple binary system of tumor stratification? This scheme could jeopardize the survival of some patients by under-treating those who need more-intensive treatment. Conversely, patients might unnecessarily receive high doses of radiation if their disease is misclassified as high-risk. This binary stratification does not represent the complexity of MB; thus, by developing
TABLE 3
History/timeline of MB research and impact on treatment

<table>
<thead>
<tr>
<th>Year</th>
<th>Landmark Event/Research</th>
<th>Impact on Patient Therapy</th>
</tr>
</thead>
<tbody>
<tr>
<td>1925</td>
<td>Cushing failed to establish an MB model in monkeys</td>
<td>Frontline therapy was the surgical resection of tumor.</td>
</tr>
<tr>
<td>1953–today</td>
<td>Patterson and Farr reported improved 3-y survival of 65% in patients receiving CSI</td>
<td>CSI included as a frontline therapy for many patients, but not for infants.</td>
</tr>
<tr>
<td>1969–today</td>
<td>Chang classification system introduced to classify tumors based on size and metastatic status</td>
<td>Chang M stage determined whether patients had standard- or high-risk MB. Radiation dose was adjusted based on risk.</td>
</tr>
<tr>
<td>1970–1990s</td>
<td>Establishment of MB cell lines DAOY, D283, D341, and others that form tumor upon implantation in immunocompetent recipients</td>
<td>Multimodality treatment: chemotherapy was added as adjuvant and neoadjuvant treatment.</td>
</tr>
<tr>
<td>1997</td>
<td>Ptc(^{1/2}) mouse model of SHH-MB was established. Many other SHH-MB mouse models were established thereafter.</td>
<td>Enabled testing of specific SHH inhibitors in many different preclinical models. Many clinical trials are underway to test the efficacy of SHH inhibitors.</td>
</tr>
<tr>
<td>2007–today</td>
<td>World Health Organization reclassification of MBs based on histology</td>
<td>Certain tumor histologic findings indicate worse survival that may warrant different therapy.</td>
</tr>
<tr>
<td>2010–today</td>
<td>Molecular subgrouping of MB</td>
<td>Targeted therapies based on MB subgroup.</td>
</tr>
<tr>
<td>2010</td>
<td>WNT-MB was found to arise outside of the cerebellum. Establishment of the first WNT-MB mouse model. Patients with WNT-MB have the best prognosis.</td>
<td>Clinical trials conducted to determine whether patients can benefit from reduced-intensity radiation therapy.</td>
</tr>
<tr>
<td>2012</td>
<td>Establishment of the first two group 3 MB mouse models; many other group 3 MB models were developed thereafter.</td>
<td>Provided more understanding of the most aggressive MB subgroup, including molecular drivers and pathways. Clinical trials evaluated the inclusion of gemcitabine and pemetrexed and a PI3K inhibitor in the current therapy.</td>
</tr>
<tr>
<td>2012–today</td>
<td>Bioinformatics platforms developed to analyze and visualize large data sets</td>
<td>Empower researchers to translate these data into biologic insights and clinical applications.</td>
</tr>
<tr>
<td>2016</td>
<td>ChIP-seq reveals distinct enhancer signatures in MB tumors and helps elucidate cellular origins of MB</td>
<td>Provided additional layer to distinguish group 3 and group 4 MB tumors to guide therapy selection.</td>
</tr>
<tr>
<td>2018</td>
<td>Integration of methylation status and molecular signature further stratified MB into different subtypes</td>
<td>Allowed for a more refined targeted therapy to minimize long-term side effects of radiation and chemotherapy. Guide the design of clinical trials.</td>
</tr>
<tr>
<td>2018</td>
<td>Proteomics and phosphoproteomics analyses highlight aberrant ERBB4-SRC pathway in group 4 MB tumors</td>
<td>Provided further insights into MB biology, including cell of origin identification and drug-resistant populations. Enabled studies to improve patients' clinical outcomes.</td>
</tr>
<tr>
<td>2018</td>
<td>scRNA-seq of developing murine cerebellum and MB tumors</td>
<td>Provided further insights into MB biology and cell of origin, especially for group 4 MB.</td>
</tr>
<tr>
<td>2018</td>
<td>Brain tumor resource biobank established</td>
<td>Immunotherapies are being evaluated in clinical trials.</td>
</tr>
<tr>
<td>2018</td>
<td>CAR T cells showed preclinical efficacy in MB models</td>
<td>Provided further insights into MB biology and cell of origin, especially for group 4 MB.</td>
</tr>
<tr>
<td>2019</td>
<td>scRNA-seq of MBs from patients</td>
<td></td>
</tr>
</tbody>
</table>

ChIP-seq, chromatin immunoprecipitation sequencing; ERBB4, Erb-b2 receptor tyrosine kinase 4; scRNA-seq, single cell RNA sequencing.
a more refined classification system, the therapeutic outcome and patients’ quality of life should improve.

Epidemiologic studies suggest progress in therapy, as evidenced by the increased overall survival of patients with MB. For example, the National Cancer Institute’s Surveillance, Epidemiology, and End Results program reported a 5-year survival of 45% during 1973–1986 (Roberts et al., 1991). Increased chemotherapy implementation during 1978–1986 translated into a significant decrease in the hazard ratio and increased overall survival (Roberts et al., 1991). A 1988 report from the Children’s Hospital of Philadelphia showed a 2-year disease-free survival of 96% for patients receiving radiation in combination with VCR, cisplatin, and lomustine, compared with 59% for patients who were treated with radiation only (Packer et al., 1988). Despite the progress, the prognosis for infants was dismal (5-year survival was 20%), with chemotherapy often prescribed to reduce the level of radiation therapy and minimize the devastating neurologic side effects of CSI (Baram et al., 1987; Loeffler et al., 1988). Some of the failure to achieve improved survival was probably due to a poor understanding of MB subtypes. In retrospect, it was fortuitous that some chemotherapy regimens were successful.

2. The Challenge of Treating CNS Tumors. In the late 1960s, when our understanding of cancer was naïve, one oncologist suggested that chemotherapy of brain tumors was limited by their “peculiar environment,” (i.e., the BBB). The author’s intended meaning was that drugs successfully used for non-CNS tumors can be effective if the BBB is circumvented. In reality, bypassing the BBB remains a formidable impediment to treating CNS disease. An additional challenge was that the pharmacologic properties required for a drug to bypass the BBB were mostly unknown (a “known unknown”). Today, this remains a challenge, but avoiding high lipophilicity is a main criterion for the development of such drugs (Rankovic, 2015).

Although the BBB was first described in the early 1900s, its molecular basis was not accurately described until the late 1980s and not functionally confirmed until the 1990s (Obermeier et al., 2013). One approach to bypass the BBB is direct delivery. For example, in the late 1960s, MTX was promoted as a candidate for treating brain tumors because it had been used successfully to treat childhood ALL. Because MTX is a substrate of ABCG2, a major drug-efflux transporter at the BBB, it does not readily penetrate the BBB (Jacus et al., 2016; Wijaya et al., 2017). In the early days,
researchers developed a system using shunts and a brain-implanted pump that enabled intrathecal delivery of MTX (Norrell and Wilson, 1967). These same authors reported in 1969 that MTX was successfully delivered to patients with tumors of the posterior fossa (i.e., gliomas and MBs) by intrathecal delivery (Wilson and Norrell, 1969). Although this treatment was initially successful, relapse ultimately occurred. The authors recognized that the challenge at the time was their lack of basic knowledge of the growth characteristics of brain tumors. What is the tumor’s generation time? How long is one cell cycle? The view of the time was that for chemotherapy to be effective, it had to be present for an entire generation period and kill a certain portion of the tumor. For brain tumors, this information was grossly lacking. As the 1970s progressed, chemotherapy supplemented surgery and radiation, but the drugs used were mostly based on successful therapies for other systemic cancers (e.g., leukemia). In fact, the approach was very empirical. As recently as 2000, the chemotherapeutic approach to MB used so-called non-targeted drugs (i.e., those that did not exploit potential tumor liabilities). These included combinations of cisplatin (a DNA cross linker/etoposide (a topoisomerase inhibitor) and CYCLO (DNA-alkylating agent)/VCR (microtubule inhibitor) or the “Packer protocol,” which included VCR, cisplatin, and lomustine (Cohen and Packer, 1996).

3. Using Molecular Subgrouping and Omics Studies to Improve Therapy. The emergence of “omics” technologies refined MB classification by using features such as a tumor’s molecular gene-expression pattern. In 2010, MB experts agreed on four principle transcriptional subgroups of MB: wingless (WNT), sonic hedgehog (SHH), group 3, and group 4 (Fig. 4B) (Taylor et al., 2012). WNT and SHH are characterized by mutations in key WNT- and SHH-signaling proteins, respectively, resulting in aberrant signaling. MYC proto-oncogene, bHLH transcription factor (MYC) amplification is one of the key drivers of group 3 MB, and MYCN and CDK6 (cyclin-dependent kinase) are often amplified in group 4 tumors (Northcott et al., 2012). In addition to molecular signatures, the subgrouping displays distinct demographics, histology, cytogenetics, and clinical outcome. The molecular subgroupings are valuable prognostic tools that can serve as guides toward better therapies. In 2017, such a multimomic approach integrated DNA methylation, gene expression, somatic copy-number alterations, and clinical features to further stratify MB; this study revealed two subtypes within the WNT subgroup: four within SHH and three each within group 3 and group 4 (Cavalli et al., 2017). The relation between this in-depth omic-based stratification and therapeutic outcome is actively investigated.

The recent molecular subgrouping of MB revealed differences in response to therapy. For example, the cure rate of patients with WNT-MB treated with conventional therapy exceeds 90% (Northcott et al., 2012). Given the excellent prognosis with current therapy, a trial is underway to determine whether reduced-intensity radiotherapy for WNT-MB is feasible. If so, implementation of such a regimen would improve long-term quality of life (NCT02724579).

Among patients with SHH-MB, infants and children have a poorer prognosis than adults (Cavalli et al., 2017). Given the aberrant SHH signaling in these tumors, including a Smoothened (SMO) inhibitor, a regulator of the SHH pathway is beneficial against adult recurrent SHH-MB (Robinson et al., 2015). Nonetheless, molecular and clinical studies showed that the SMO inhibitor was effective only in a subset of SHH-MB that harbors mutations upstream of SMO. This strategy of “drugging SMO” to treat SHH-MB revealed three unexpected challenges: 1) The treatment causes developmental defects in the young, especially in skeletal development (Robinson et al., 2017). 2) It also results in acquired resistance, which produces tumors that harbor constitutively active SMO (due to mutation) and are aggressive and therapeutically refractory (Taipale et al., 2000; Atwood et al., 2015). 3) Patients with defects in the SHH pathway downstream of SMO (e.g., Suppressor of Fused (SUFU) mutations or Glioma associated oncogene family zinc finger 2 (GLI2) amplification) do not benefit from inclusion of an SMO inhibitor in their treatment (Robinson et al., 2018). However, other treatment options are available. An SHH-specific treatment that antagonizes GLI (Glioma associated oncogene family zinc finger) proteins, such as arsenic trioxide, is being clinically evaluated (Kim et al., 2013). The TP53 status in SHH-MB influences risk, as those with mutated TP53 are considered at high risk (41% overall survival vs. 81% for those with wild-type TP53) (Fig. 4B).

Approaches that identify other potential vulnerabilities in MB subtypes might be therapeutically exploited, or additional MB strata might improve our knowledge of responders. Recent examples are MB phosphoproteome and epigenetic-methylation status coupled with gene-expression profiles, which appear to be valuable. For instance, a recently published phase II clinical trial by Robinson et al. (2018) identified two distinct methylation subtypes in infants with SHH-MB: iSHH-I and iSHH-II. The iSHH-II subtype showed favorable progression-free survival in the absence of radiation and chemotheraphy compared with that of the iSHH-I subtype (Robinson et al., 2018). A preclinical study in mice used phosphoproteomics to monitor SHH-MB progression and discovered that casein kinase 2 is a promising target for SHH-MB therapy, including SMO-resistant tumors (Purzner et al., 2018). Casein kinase 2 modifies the phosphorylation status of many key SHH proteins. A clinical trial (NCT03904862) is underway to test the safety and tolerability of a casein kinase 2 inhibitor in patients with SHH-MB. The
further integration of molecular signatures into treatment might enable reduced-intensity therapy and minimize the long-term side effects of current MB treatment.

Molecular subgrouping has also uncovered pathways that drive disease and may be therapeutically tractable. For instance, the phosphatidylinositol-3-kinase (PI3K) pathway is upregulated in group 3 MB (Pei et al., 2012, 2016). Given the promising results from preclinical models treated with PI3K inhibitors, downregulation of this pathway might improve MB outcome. Most recently, Erb-b2 receptor tyrosine kinase 3-Src proto-oncogene (ERBB4-SRC) signaling was found to be a hallmark of group 4 MB, as revealed by phosphoproteomic profiling (Forget et al., 2018). Although no specific drug has been proposed, this study uncovered a promising target for treating group 4 MB.

Insights into the oncogenesis and pathways of MB that have been gleaned over the past 10 years would not have been possible without the major technological advances that have been made. In addition to the development of faithful model systems (described in the next section), the omics tools (proteome, transcriptome, phosphoproteome, and epigenome) have been invaluable, not only in MB disease classification and stratification but also in defining the biologic properties of the disease. Genome-editing technology, especially through CRISPR, has enabled functional genomic screens in relevant model systems that identified factors affecting the SHH pathway. For example, two independent groups uncovered the critical roles of cilia in SHH signaling (Breslow et al., 2018; Pusapati et al., 2018). Because proteins are trafficked in and out of primary cilia by microtubule-based motors, these studies may validate the use of vinca alkaloids, such as VCR. The functional genomics approach can be applied to other MB subgroups to gain insight into the genes and pathways that promote their cell proliferation and drug resistance.

Recent advances in transcriptomics technology, including single-cell RNA sequencing (scRNA-seq), are most likely to help advance our knowledge of which progenitors trigger MB development (Hedlund and Deng, 2018). The ability to measure gene expression in hundreds to thousands of individual cells generates unique cell-specific signatures. In aggregate, these signatures might provide a “fingerprint” of unknown factors affecting developmental timing, which could aid in determining where development goes awry to create MB. The scRNA-seq data on developing murine cerebellum are now available, and they have revealed novel transcriptional regulators for distinct cell populations (Carter et al., 2018). Furthermore, single-cell transcriptomics will help us understand MB heterogeneity and identify specific cell populations that drive oncogenesis or drug resistance (El-Hamamy et al., 2018). Most recently, an integration of scRNA-seq performed in 25 patients with MB and mouse cerebellar scRNA-seq data sets revealed that WNT-MB, SHH-MB, and group 3 MB tumors consist of specific undifferentiated cells and differentiated neuron-like malignant cells, whereas group 4 MB tumors are mostly differentiated neuron-like neoplastic cells (Hovestadt et al., 2019). This study suggests that glutaminergic cerebellar nuclei and unipolar brush cells within the cerebellum are the cells of origin for group 4 MB; this finding will aid in the development of group 4 MB models and understanding the biology of this subgroup (Hovestadt et al., 2019).

D. Development of Models to Better Understand MB Biology and Develop New Therapies

After Cushing began removing MB tumors in 1925, he attempted to establish an animal model by implanting human MB tumor fragments into the cisterna magna of monkeys (Rutka and Hoffman, 1996). Unfortunately, his efforts were largely unsuccessful. Animal models of brain tumors are advantageous from a drug-development perspective because salient features, like the amount of drug reaching the brain, the effects of route and schedule of administration, and efficacy, can be investigated. Without animal models, the strategy in the clinic was empirical—did a patient respond or not? It was challenging to determine whether brain tumors responded differently to chemotherapy than did extracerebral tumors. Since 2000, our understanding of MB has advanced, and the challenges of developing effective (and most likely tumor-tailored) MB therapy have been advanced by the development of in vitro cell model systems, genetically engineered mouse models (GEMMs), orthotopic patient-derived xenografts (O-PDXs), and other sophisticated models, including brain organoids, especially when paired with high-throughput sequencing and genome-editing technologies.

1. Immortalized Cell Lines and Those Derived from GEMMs. Many human MB cell lines, including DAOY, D283, UW228, D341, and HD-MB03, have been established since the 1980s (Friedman et al., 1985, 1988; Jacobsen et al., 1985; Keles et al., 1995; Milde et al., 2012). These lines represent different MB subgroups, thereby enabling researchers to not only study specific tumor subgroups but also use a cost-effective means to determine whether MB subgroups exhibit differences in drug response. Such differences might provide insight into the molecular signature of targetable drug pathways. One study found differential sensitivity of cells to the flavonoid quercetin as a radiosensitizer in DAOY cells, which represent SHH-MB with TP53 mutation (Jacobsen et al., 1985), compared with D458-med cells (He X et al., 1991), which represent group 3 MB (Lagerweij et al., 2016). Although the number of MB cell lines has increased, in vitro models for WNT-MB and group 4 MB are underrepresented compared with SHH-MB and group 3 MB.
Immortalized cell lines are amenable to genetic manipulation and drug screening. However, these studies have caveats. For instance, cell lines are maintained in environments that are nonphysiologic (e.g., 20% O₂, whereas many organs function in the range of 2%–8% O₂); the lines are prone to harboring genetically unstable genomes and can respond differently to drugs in vitro versus in vivo. For example, MYC overexpression sensitizes some MB cell lines to drugs in vitro, but this is not recapitulated in patients’ response (von Bueren et al., 2011). To overcome the limitations of established cell lines, primary tumor cells from patients or cells derived for GEMMs are viable alternatives. Tumors isolated from GEMMs can grow three-dimensionally as tumorspheres, are tumorigenic, and retain MB markers. For example, tumorspheres from Ptc+/– mice were used to determine that the activation of the Ras-type GTPase (RAS/mitogen-activated-protein kinase (MAPK) pathway promotes resistance to SMO inhibition (Zhao et al., 2015). Cell lines derived from group 3 MB GEMMs have shown great promise in identifying translationally important therapeutics. A recent study from group 3 MB GEMMs has enabled biologic discoveries. For example, tumorspheres derived from Ptc+/– mice were used to determine that the activation of the Ras-type GTPase (RAS/mitogen-activated-protein kinase (MAPK) pathway promotes resistance to SMO inhibition (Zhao et al., 2015). Cell lines derived from group 3 MB GEMMs have shown great promise in identifying translationally important therapeutics. A clinical trial (NCT01878617) evaluating the efficacy of combining the nucleoside gemcitabine (targets DNA synthesis by inhibiting ribonucleotide reductase) and the antifolate pemetrexed with standard chemotherapy for group 3 and group 4 MB was based on a successful high-throughput screen of a Food and Drug Administration–approved library using mouse group 3 tumorspheres (derived from Trp53–/+; Cdkn2c–/– mice, with enforced expression of MYC) (Morfoiuace et al., 2014). Similarly, a screen using a group 3–derived GEMM cell line identified a PI3K/mTOR (mammalian target of rapamycin) inhibitor as a good therapeutic prospect, which has also been taken into a clinical trial (NCT03213678) (Pei et al., 2016).

The ease of genetic manipulation of GEMM-derived cell lines has enabled biologic discoveries. For example, Vo et al. (2016) found that the interaction of MYC and MIZ (MIZ-type zinc finger) is crucial for maintaining tumor stemness, which appears to affect the development of group 3 MB in vivo. The loss of this interaction changes the disease pathology entirely, further highlighting the role of MYC and MIZ in defining subgroup identity (Vo et al., 2016). Our group found that the expression of the ATP-binding cassette transporter subfamily G member 2 (ABCG2) significantly influences group 3 MB response to chemotherapy (Morfoiuace et al., 2015; Wijaya et al., 2017). Using a group 3 GEMM–derived model, we found that ABCG2 is highly expressed at the plasma membrane. Inhibition of ABCG2 function in vitro by using the ABCG2-specific inhibitor fumitremorgin C increases the number of apoptotic tumor cells. Importantly, ABCG2 inhibition in vivo increases topotecan cytotoxicity and extends the survival of tumor-bearing mice (Morfoiuace et al., 2015).

2. GEMMs.

a. SHH-MB GEMMs. The first MB GEMM was for the SHH subgroup. The Ptc+/- mouse was engineered in 1997 (Goodrich et al., 1997). By homologous recombination, part of the first and the entire second exon of Ptc was replaced with the marker gene LacZ (encoding β-galactosidase) and neomycin genes (Goodrich et al., 1997). The authors discovered that Ptc is an essential gene because homozygous deletion of Ptc was embryonically lethal (E9.0–E10.5). In contrast, the loss of one allele of Ptc produced phenotypes commonly observed in basal cell nevus syndrome (also known as Gorlin syndrome), which includes formation of extra digits in the hindlimb and an increased propensity to develop brain tumor/MB (14%), with peak incidence between 16 and 24 weeks (Goodrich et al., 1997). Subsequent studies found that the loss of Trp53 and inactivation of one Ptc allele resulted in more than 95% of mice developing MB about 12 weeks after birth (Wetmore et al., 2000, 2001). Loss of the cyclin-dependent kinase inhibitor p18^kip^ in the Ptc+/- background also increased the incidence of MB, highlighting the role of Retinoblastoma protein in MB pathogenesis (Uziel et al., 2005). Similarly, the loss of p27^kip1^ cyclin-dependent kinase inhibitor in the Ptc+/- background resulted in MB with increased latency and penetrance compared with the loss of Ptc alone (Ayrault et al., 2009). These latter two GEMMs are particularly useful to model SHH-MB without the loss of the tumor suppressor TP53, which is the subtype that develops in most patients with SHH-MB (Cavalli et al., 2017). Further developments in SHH-MB GEMMs include the Cre-LoxP system, which enables the inducible deletion of Ptc in specific cell types (Yang et al., 2008); Sufu+/- (Suppressor of Fused); Trp53+/- mice, 60% of which develop MB within 4 months (Lee et al., 2007); and oncogenic SMO mutants, which are accurate models of drug-resistant MB tumors and those that produce leptomeningeal metastases (Hallahan et al., 2004; Hatton et al., 2008).

b. WNT-MB GEMM. For years, MB tumors were thought to arise only within the cerebellum and from granule neuron progenitors as the cell of origin (Rutka and Hoffman, 1996). However, such a monolithic perspective was changed by GEMMs. Specifically, Gibson et al. (2010) discovered that WNT-MB tumors arise outside the cerebellum from progenitor cells in the dorsal brainstem. Unlike SHH tumors, which are located in the cerebellar hemisphere, WNT tumors are Zic+ and found in the dorsal brainstem (Gibson et al., 2010). This finding enabled the development of mouse models of the WNT-MB subgroup, which accurately resemble human WNT-MB.

Under current standard MB chemotherapeutic regimens, the WNT-MB subtype has the best overall survival, suggesting that these tumors are more responsive than other MB subtypes (Northcott et al., 2012). One well-known clue was that the human WNT-MB tumors

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appeared hemorrhagic (Phoenix et al., 2016). The development of a WNT-MB GEMM provides a foundation for understanding both the hemorrhagic nature of WNT-MB tumors and their overall responsiveness to therapy in vivo. Researchers found that WNT-MB tumors increase the permeability of the BBB by increasing paracrine WNT signaling (Phoenix et al., 2016). WNT-MB tumors also secrete WNT inhibitor factor 1 and Dickkopf 1 as a part of a negative feedback loop of a constitutively active WNT-signaling pathway. The high level of WNT agonists silences WNT signaling in surrounding epithelial cells, which contribute to BBB formation. This results in tumors with compromised BBBs, which allows for increased exposure of systemic chemotherapy, such as VCR, and most likely contributes to excellent prognosis (Phoenix et al., 2016).

c. **Group 3 MB GEMMs.** In 2012, two group 3 MB mouse models were independently developed (Kawauchi et al., 2012; Pei et al., 2012). One group enforced Myc expression in Trp53-null cerebellar progenitor cells (Kawauchi et al., 2012), whereas the other overexpressed Myc in Prominin1+; Lineage1− (Prom1+; Lin1+); cerebellar stem cells, along with a dominant-negative Trp53 (Pei et al., 2012). These models suggested that group 3 MB can arise from different progenitors but nonetheless results in mice that developed tumors that closely resembled the most aggressive type of MB, MYC-driven MB. Tumors from these GEMMs exhibit large-cell/anaplastic histology, with a gene-expression signature closely resembling the human group 3, MYC-overexpressing cohort, which is associated with the poorest prognosis (Wang et al., 2018). These GEMMs have advantages, in that the tumors isolated from these mice grow in culture indefinitely, maintain their similarity to human group 3 MB, and have been used to identify new chemotherapeutics that are now in clinical trials, as mentioned previously. Other accurate group 3 MB preclinical models have since emerged, including those driven by activation of growth factor independent 1 and 1B (Gfi1 and Gfi1B) in Myc-expressing tumors (Northcott et al., 2014), inactivation of Ezh2 (enhancer of zeste homolog 2) (Vo et al., 2017), and Myc upregulation by CRISPR technologies (Vo et al., 2018). These models also highlight the heterogeneity in group 3 MBs and may be used accordingly to study different drivers of this subtype of disease.

d. **Group 4 MB GEMMs.** To date, there are only a few reports of group 4 MB models, and their drivers might be heterogeneous. Targeted expression of partially stabilized Mycn (T28A) in neural stem cells isolated from postnatal day 0 cerebella produced tumors with a group 4 signature (Swartling et al., 2012). One study identified regulatory elements and core transcription factors and proposed deep cerebellar nuclei in the cerebellar nuclear transition zone or earlier precursors from the upper rhombic lip as the putative cell of origin for a subset of group 4 MB (Lin et al., 2016). Inactivating mutations in lysine (K)-specific demethylase 6A (KDMA6), tandem duplication of synuclein alpha interacting protein (SNCAIP), and PR domain 6 (PRDM6)-enhancer hijacking have also been implicated in driving group 4 MB tumors. Until recently, molecular signatures of MB had been restricted to the genome, epigenome, and transcriptome; however, proteomic and phosphoproteomic studies suggest that post-transcriptional pathway regulation markedly differs among MB subgroups (Forget et al., 2018; Zomerman et al., 2018). Importantly, an analysis of the global correlation between transcript levels and protein levels from 41 flash-frozen primary MBs revealed a poor mRNA-protein correlation in group 4 samples, with overall higher levels of protein than mRNA (Forget et al., 2018). This study determined that aberrant receptor tyrosine kinase signaling is a hallmark of group 4 MB, suggesting that clinically approved TKIs, such as lapatinib or dasatinib, may be effective in this cohort. Phosphoproteomics also enabled the development of the second group 4 MB GEMM. Enforced expression of constitutively active SRC kinase and dominant-negative Trp53 produced tumors that do not resemble MYC-driven group 3 MB but rather yield tumor with the molecular signature of group 4 (Forget et al., 2018).

3. **O-PDXs.** Although GEMMs have been essential to gaining insight into the MB cell(s) of origin, underlying drivers of MB subtypes, and candidate therapeutics, they have an inherent limitation—GEMMs do not capture the underlying diversity of MB subtypes seen in humans. To overcome this challenge, fresh tumors resected from patients have been implanted into the cortices of immunodeficient mice to produce O-PDXs. One advantage of this system over human cell lines in culture is that the tumors expand in vivo, in a normal physiologic environment, without ever encountering the artificial environment of a culture dish. Although expensive, O-PDX models are clinically relevant because they replicate the interindividual heterogeneity of the disease. This trait will be useful for designing individualized therapy, which has been applied for some tumors (Kawauchi et al., 2012; Brabetz et al., 2018). However, until that time, this diversity is important for revealing new therapeutic vulnerabilities of some MBs. For example, one group used different O-PDX models (i.e., MYC-amplified, MYC-amplified classic morphology group 3, and desmoplastic morphology SHH subgroup) to examine the efficacy of CDK4/6 inhibition and found that palbociclib suppresses tumor growth and significantly extends survival of tumor-bearing mice (Cook Sangar et al., 2017). As an extension of this study, a clinical trial (NCT03709680) is evaluating palbociclib in combination with other agents for treatment of pediatric patients with recurrent/refractory solid tumors, including MB. Another group has
used O-PDXs to study CSI dose and volume (Smith et al., 2019). Successful delivery of radiation therapy to the O-PDX model will provide a better tool for testing the timing and dosing of chemotherapeutics that closely resemble patients’ treatment regimen.

4. Three-Dimensional Organoids and Other Recent Advances. In addition to immortalized cell lines and tumorspheres derived from GEMMs, an emerging model that might capture the complexity of the brain is three-dimensional organoids. Using human embryonic stem cells and the appropriate culture conditions, researchers have generated brain organoids containing functional cell types and architecture that resemble the developing fetal brain (Lancaster et al., 2013; Muguruma et al., 2015). Disease-context organoids derived from healthy donors or patients provide a novel brain tumor resource biobank (Brabetz et al., 2018) for MB based on molecular signatures of the disease. The therapeutic approaches described above are further enhanced by the following approaches: antibody-drug conjugates display selectivity because the antibody recognizes a unique or tumor-overexpressed antigen and eradicates the tumor by delivering a toxic molecule to it, and antisense oligonucleotides alter the level of disease-modifying or protective mRNA. The most recent addition to the cancer therapeutic armamentarium is proteolysis-targeting chimeras, which recruit the tumor’s degradation machinery to degrade disease-modifying proteins (Pettersson and Crews, 2019).

The therapeutic approaches described above are further enhanced by the following approaches: antibody-drug conjugates display selectivity because the antibody recognizes a unique or tumor-overexpressed antigen and eradicates the tumor by delivering a toxic molecule to it, and antisense oligonucleotides alter the level of disease-modifying or protective mRNA. The most recent addition to the cancer therapeutic armamentarium is proteolysis-targeting chimeras, which recruit the tumor’s degradation machinery to degrade disease-modifying proteins (Pettersson and Crews, 2019).

All the advances described herein would not be possible without interdisciplinary collaborations. The teams acquire large-scale cancer genomic data, which are analyzed by computational biologists, validated by biologists, and then translated into information used by clinicians to design better clinical trials. Such cooperative effort requires data sharing from each functional area. Fortunately, our scientific community is open to collaboration. Websites such as cBioportal (Gao et al., 2013), R2: genomics analysis and visualization platforms (r2.amc.nl), and many others aid in data sharing and visualization of large-scale genomics data. Bioinformatics tools, such as DiSCOVER (Hanaford et al., 2016), can guide predictions for therapeutic strategy for MB based on molecular signatures of the disease. The brain tumor resource biobank (Brabetz et al., 2018) for O-PDXs enables researchers to investigate and test new
targets and therapeutic modalities to treat MB. Studies across diverse scientific disciplines and collaborations between academic institutions and biotechnology/pharmaceutical industries will be the key to discovering the best combination of therapies for the appropriate tumor types and significantly advancing patient care and survival.

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