Estimating Long-Term Survival of Adults with Philadelphia Chromosome-Negative Relapsed/Refractory B-Precursor Acute Lymphoblastic Leukemia Treated with Blinatumomab Using Historical Data

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ABSTRACT

Introduction: Blinatumomab is a bispecific T cell-engaging antibody construct indicated for adult patients with relapsed/refractory (R/R) Ph(−) B-precursor acute lymphoblastic leukemia (ALL), an aggressive disease with poor prognosis. A phase 2 single-arm clinical study showed that 43% of patients achieved CR/CRh within two cycles and approximately 20% of patients receiving blinatumomab were still alive after 2 years.

Methods: The objective of the current analysis was to estimate long-term survival of patients receiving blinatumomab beyond the observed time period in the clinical study using a large historical observational dataset. Conditional survival probabilities of blinatumomab-treated patients beyond month 60 were assumed to be the same as the US general population.

Results: At month 60, the estimated proportion of blinatumomab-treated patients alive was more than double that of historical patients (12.6% vs 5.4%). The mean overall survival was 76.1 months for blinatumomab patients and 39.8 months for historical patients. Sensitivity analyses including additional follow-up data from the clinical study showed consistent results.

Conclusions: These findings suggest that blinatumomab provides substantial overall survival benefit to patients with (R/R) Ph(−) B-precursor ALL compared with salvage chemotherapy.

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Keywords: Acute lymphoblastic leukemia; Blinatumomab; Hematology; Long-term survival; Oncology
INTRODUCTION

The prognosis for adult patients with Philadelphia chromosome-negative (Ph[−]) B-precursor ALL who are refractory to treatment or experience relapse (R/R) is poor: over 90% die from the disease and their survival time is short (median OS is 3–5 months) [1–4]. These patients tend to be particularly young, with a median age of 34–39 years, and die, on average, 30 years prematurely [5–7]. In general, response to salvage treatment followed by hematopoietic stem cell transplant (HSCT) offers the only potential for cure and long-term survival [1, 4]. Until recently, salvage treatment for adult patients with R/R Ph(−) B-precursor ALL was limited to highly toxic multi-agent chemotherapy regimens. Most patients who receive these regimens endure repeated and prolonged hospitalizations due to the severity of the disease and the aggressiveness of the treatment itself [8–11].

Blinatumomab is a novel bispecific T cell engager (BiTE®) antibody construct that simultaneously binds CD3-positive cytotoxic T cells and CD19-positive B cells. Blinatumomab was approved for adults with R/R Ph(−) B-cell precursor ALL by the FDA in December 2014 and subsequently by the European Medicines Agency (EMA) in November 2015 [12, 13]. Regulatory approval of blinatumomab in the US was based on results from a single-arm phase 2 study in 189 adults with R/R Ph(−) B-cell precursor ALL [14]. Historical observational data from 1139 patients with R/R Ph(−) B-cell precursor ALL who received standard of care therapy in Europe and the US provided context for the interpretation of the single-arm clinical study [15]. The single-arm clinical study and the historical observational data have been compared for rates of complete remission (42.9% vs 24%) and overall survival (OS) at 1 (32.0% vs 15.5%) and 3 years (13.8% vs 6.2%) [14, 15]. To date, the blinatumomab clinical study includes 3 years of survival follow-up data, after which time 13.8% of patients remained alive [Amgen data on file]. In comparison, 6.4% patients in the historical data were alive at 3 years. The historical observational dataset included up to 21 years of follow up, after which time between 2% and 3% of patients remained alive. Given the very small percentage of patients in the historical observational dataset who lived for at least 21 years, the proportion of patients receiving blinatumomab who were still alive after 3 years is of particular interest.

To estimate the effect of a treatment on long-term survival, the mean OS is a preferred endpoint to median OS because it captures the entire survival curve over the lifetime of a population [16]. The objective of this analysis was to estimate the long-term survival of patients with R/R Ph(−) B-cell precursor ALL receiving blinatumomab, leveraging the long duration of follow-up data available in the historical observational dataset.

METHODS

Data Sources

Blinatumomab Clinical Trial

This phase 2, open-label, single-arm study was conducted at sites across Europe and the US and enrolled patients over the period 2010–2014 (ClinicalTrials.gov NCT01466179) [14]. Eligible patients (N = 189) included men and women ≥18 years of age with Ph(−) B-cell precursor ALL, with one of the following characteristics:

- relapsed/refractory disease with first remission duration ≤12 months in first salvage;
- relapsed/refractory disease after first salvage; or
• relapsed/refractory disease within 12 months of allogeneic HSCT.

Patients with these characteristics are considered particularly difficult to treat. Overall survival was collected as a secondary endpoint.

All procedures followed were in accordance with the ethical standards of the responsible committee on human experimentation (institutional and national) and with the Helsinki Declaration of 1964, as revised in 2013. Informed consent was obtained from all patients for being included in the study.

**Historical Observational Dataset**

A retrospective analysis was performed by combining previously collected data from adult R/R ALL patients in national study groups and large treatment centers in Europe and the US. (clinicaltrials.gov NCT02003612). Patients were initially selected from study group/site databases if they had a diagnosis of R/R Ph(−) B-precursor ALL between 1990 and 2013, were aged ≥15 years at the time of initial ALL diagnosis of ALL, and had no previous treatment with blinatumomab. Patients were further selected from the historical dataset based on key eligibility criteria from the blinatumomab clinical study, including age ≥18 years at relapse, ALL that relapsed within 12 months from initial diagnosis, or relapsed after alloHSCT, or refractory to initial or subsequent treatments, or in second or later relapse. Patients with a first remission duration of >12 months in first salvage were excluded, unless they had a second relapse or a relapse within 12 months of receiving alloHSCT.

In total 1139 patients met these criteria, of whom survival data was available from 1112 patients [15]. Two-thirds of patients (67%) were diagnosed after the year 2000.

To calculate overall survival estimates, six strata were created based on known prognostic factors in R/R ALL (age, prior HSCT, and line of salvage). Combined survival outcomes were obtained by weighting the stratum-specific survival estimates according to the proportion of patients in the blinatumomab trial in each stratum [15].

**Estimation of Long-Term Overall Survival of Patients Receiving Blinatumomab**

Estimation of long-term survival was comprised of three phases. In the first phase, survival from months 0–25 was estimated using observed Kaplan–Meier survival data from the primary analysis of the blinatumomab clinical trial [14, Amgen data on file]. In the second phase, survival from months 26–60 was estimated using the historical observational dataset. Specifically, from months 26–60, we assumed that there was no additional treatment effect from blinatumomab—i.e., the survivors in the blinatumomab clinical trial at months 26–60 had the same monthly conditional survival probabilities as those in the historical observational dataset who received salvage chemotherapy [17, 18]. In the third phase, since patients who survive for more than 5 years are considered to be cured of the disease, survival from month 60 onwards was estimated by applying the age-specific annual conditional probability of survival from the life tables of the US population.

**Calculation of Mean OS**

The mean OS for patients receiving blinatumomab was estimated by calculating the area under the curve (AUC) for all three phases combined. The same AUC calculation
was also applied to calculate the mean OS for patients in the historical observational dataset.

**Validation of Estimates Using 3 Years of Observed Data From The Clinical Study**

Additional OS data from the blinatumomab clinical trial after 36 months of follow-up was used to validate the long-term OS estimation [Amgen data on file]. The three-phase method described above was replicated using the blinatumomab clinical trial data to estimate survival from months 0–36, extrapolating survival from months 37–60 using the historical observational dataset, and extrapolating from month 60 onwards using conditional survival probabilities from the general US population.

**RESULTS**

Details of the patient populations from the clinical and historical studies have been described previously, and rates of CR and OS were compared between the blinatumomab patients and the historical patients using appropriate adjusted analyses [14, 15].

**Observed OS Data**

In both the blinatumomab clinical trial and the historical observational dataset, the observed OS probability decreased quickly in the first 10 months and plateaued afterwards (Fig. 1).

Comparing the observed blinatumomab clinical trial survival curve to the historical observational dataset survival curve, there is a clear separation between the curves from month zero to month 25 (Fig. 1). At month 25, the survival probability among patients from the blinatumomab clinical trial was 18.7%, more than double the survival probability among patients from the historical observational dataset (8.0%) based on the Kaplan–Meier estimate.

**OS Extrapolation up to 60 Months**

Based on extrapolation from the historical observational data, the OS probability at month 60 was 12.6% among patients from the blinatumomab clinical study and 5.4% in the historical observational dataset (Fig. 1).

**Mean OS**

The mean OS was calculated from the AUC of all three phases of the long-term survival estimation. The mean OS for patients in the blinatumomab clinical study was 76.1 months compared with 39.8 months for patients from the historical observational dataset (Table 1).

**Validation of Estimation**

The long-term OS estimation based on 36 months of OS data from the blinatumomab clinical trial was consistent with the estimation based on 25 months of trial data plus extrapolation with survival probabilities from
the historical observational dataset (Fig. 2). Using 36 months of survival data from the blinatumomab clinical trial, the survival probability at month 60 was 12.0% and the mean OS was 74.2 months.

**DISCUSSION**

This study estimated long-term survival for R/R Ph(−) ALL patients receiving blinatumomab, using both the blinatumomab clinical trial results and long-term natural history data. Assuming no additional effects of blinatumomab beyond two years, the percentage of long-term survivors treated with blinatumomab is estimated to be more than double that of patients treated with salvage chemotherapy (12.6% vs 5.4% after 5 years). The estimation technique was validated using an additional year of observed clinical trial data. Similarly, the mean OS—a measure of overall survival over the lifetime of the population—was also approximately doubled among blinatumomab-treated patients in comparison to patients in the historical observational dataset (76.1 vs 39.8 months).

The historical data set was pooled from European national study groups and large individual sites from Europe and the United States. Given rarity of R/R Ph(−) B–precursor ALL, the historical dataset represents the largest study of its kind in adults with this disease [13, 15], spanning over 20 years (1990–2013). During this time, no new effective treatments have emerged [3]. The increased use of paediatric-inspired protocols and improvements in supportive care and transplant realisation may have improved survival over time in some groups of adult R/R ALL patients [19]. Two-thirds of patients in the historical dataset were diagnosed after the year 2000, the survival times were similar between the whole dataset and those patients diagnosed from 2000 onwards [15]. The historical data therefore appears to provide a valid reference for survival data extrapolation and comparison between novel agents and salvage chemotherapy.

It is worth noting that the assumed difference in survival between blinatumomab and salvage chemotherapy after the observed data time period in this study might be an underestimation. After the observed OS data, we assumed that the patients treated with blinatumomab had the same conditional survival probability as patients who received salvage chemotherapy. In fact, patients receiving blinatumomab can achieve a deep response, which is known to correlate with better long-term survival outcomes [20]. Among 81 patients with R/R Ph(−) B-precursor ALL who achieved complete remission (CR/CRh*), 82%
also achieved minimal residual disease (MRD)-negativity within two cycles of blinatumomab [14]. In an earlier clinical study of 36 patients with Ph(−) B-cell precursor R/R ALL, 28% of those treated with blinatumomab survived for at least 30 months, all of whom achieved an MRD-negative response [21]. These findings suggest that patients treated with blinatumomab might have higher conditional survival probability than patients receiving salvage chemotherapy. More patients in the blinatumomab clinical study proceed to receive transplants, possibly due to favourable CR rate, or higher rate of MRD negativity [14], which is highly correlated to positive HSCT outcome and better overall survival, compared to those who do not achieve MRD [22, 23].

With 12.6% of R/R ALL patients treated with blinatumomab expected to be alive after 5 years other treatment strategies are still required. Treatment with blinatumomab before relapse has been shown to produce high response rates and survival outcomes: 78% of patients with MRD-positive disease achieved a complete MRD response after 1 cycle of blinatumomab, and RFS was 58% after 18 months [24]. In a smaller study in an MRD-positive population, after a median of 33 months follow up, 61% of blinatumomab-treated patients remained relapse-free [25]. These data raise the possibility that earlier intervention with blinatumomab may allow long-term remissions and reduce the need for salvage therapy.

The blinatumomab clinical trial enrolled R/R Ph(−) B-precursor ALL patients who were selected for negative prognostic factors, which represents the majority of patients but nonetheless a subset of the total R/R Ph(−) B-precursor ALL population. Caution needs to be taken when extrapolating the results from this study to the broader patient group.

Limitations of this study are inherent in the use of historical data to compare and extrapolate survival outcomes with data from a clinical study. These include the assumption of no additional effects of blinatumomab beyond two years, potential changes in treatment practices over time in the clinical vs historical populations and differences in transplant realisation rates. Nonetheless, the superior survival outcomes with blinatumomab treatment over standard of care have recently been confirmed in a phase 3 study [26].

**CONCLUSION**

Compared to salvage chemotherapy, based on this analysis, blinatumomab provides substantial overall survival benefit to patients with R/R Ph(−) B-precursor ALL.

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**Disclosures.** Arie Barlev was an employee of Amgen at the time the study was conducted. Vincent Lin is an employee and shareholder in Amgen. Aaron Katz is an employee and
Compliance with Ethics Guidelines. All procedures followed in the blinatumomab clinical study were in accordance with the ethical standards of the responsible committee on human experimentation (institutional and national) and with the Helsinki Declaration of 1964, as revised in 2013. Informed consent was obtained from all patients for being included in the study. The historical study was based on data collected from previously-conducted studies.

Data Availability. The datasets during and/or analyzed during the current study are available from the corresponding author on reasonable request.

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