Safety and efficacy of second-line bosutinib for chronic phase chronic myeloid leukemia over a five-year period: final results of a phase I/II study

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Introduction

With the success of BCR-ABL1 tyrosine kinase inhibitors (TKIs), patients with Philadelphia chromosome positive (Ph+) chronic myeloid leukemia (CML) can potentially enjoy a normal life expectancy. Therefore, information regarding long-term efficacy and tolerability of TKIs is important for informing treatment selection.
Several TKIs are approved for treatment of newly diagnosed chronic phase (CP) CML, each associated with a distinct safety profile. First-line TKI therapies include imatinib and the second-generation TKIs, nilotinib, dasatinib, and, most recently, bosutinib. Although response rates are high with TKI therapy, patients develop treatment resistance or experience intolerable toxicities. Determining the most appropriate therapy following treatment failure is critical to achieve optimal outcomes and prevent disease progression. Second- and third-line treatment decisions are based on the prior therapy, the reason for failure (primary or secondary resistance, intolerance), BCR-ABL1 mutation status, comorbidities, and prior toxicities.

Bosutinib was initially approved in 2012 for the treatment of patients with Ph+ CP, accelerated-phase (AP), or blast-phase (BP) CML resistant or intolerant to prior TKI therapy. This approval was based primarily on the results of a pivotal phase I/II trial of bosutinib in CP CML patients following failure of imatinib. Results of a preliminary analysis approximately 15 months after the last enrolled patient demonstrated potent activity with second-line bosutinib across a spectrum of BCR-ABL1 mutations and a toxicity profile distinct from those of other TKIs. The durability of response was confirmed with subsequent analyses 24 and 48 months after the last enrolled patient. The final results presented here from the phase I/II trial of bosutinib for imatinib-resistant or imatinib-intolerant CP CML are assessed after at least five years from the time the last patient was enrolled.

Methods

Study design and patients

This phase I/II, open-label, multicenter study was initiated in January 2006; the design has been described previously. Part 1 (dose-escalation phase) determined the recommended Part 2 starting dose of bosutinib 500 mg/day; in Part 2, the efficacy, safety, and tolerability of bosutinib were evaluated. Patients without a complete hematologic response (CHR) by week 8 or complete cytogenetic response (CCyR) by week 12 were offered enrollment on an extension study. As for the final database lock for this study (2nd October 2015), the time from the last second-line patient’s first dose was 60 months or more. The median (range) duration of OS follow up was 54.8 (0.6-96.3) months, and the median treatment duration was 25.6 (0.2-96.3) months (Table 1). At year 2, 153 (54%) patients were receiving bosutinib and at year 5, 115 (40%); IM-R, n=81 and IM-I, n=34) patients still remained on bosutinib treatment (1 year=48 weeks). Discontinuation from treatment was most common within the first two years, with 131 (46%) patients discontinuing by the end of year 2, and 38 (13%) patients discontinuing treatment in years 3 through 5. Discontinuation from treatment through year 5 were lack of efficacy [categorized by the investigator separately as PD and unsatisfactory response; n=47 (17%) and n=21 (7%), respectively], AE [n=64 (23%)], and patient request [n=19 (7%)]. The most common reasons for discontinuation from treatment through year 5 were lack of efficacy [categorized by the investigator separately as PD and unsatisfactory response; n=47 (17%) and n=21 (7%), respectively], AE [n=64 (23%)], and patient request [n=19 (7%)].

Safety and efficacy analyses

Cytogenetic response was assessed as previously described and defined as newly-achieved on study or maintained from baseline for four weeks or more (earliest time point for assessment). Evaluable patients received at least one dose and had a valid baseline assessment for the respective end point. Molecular response data were assessed at a central laboratory; however, the International Scale (IS) was not used. Patients from sites in China, India, Russia, and South Africa were not assessed for molecular response due to logistical constraints. For the purpose of this study, responders for major molecular response (MMR) had a ≥3 log reduction from standardized baseline, a detectable BCR-ABL1 transcript at baseline or postbaseline, and must have maintained or attained a CCyR. Duration of response (date of first response until confirmed response loss or PD/death) was evaluated through 30 days after last dose using the Kaplan-Meier method; patients without events were censored at their last assessment visit. See the Online Supplementary Methods for further details of the statistical methods used for this report.

Results

Patients and treatment

A total of 284 patients with CP CML (IM-R, n=195; IM-I, n=89) were enrolled and treated with second-line bosutinib (Online Supplementary Table S1). The study was closed as of August, 2015; patients still on study were offered enrollment on an extension study. As for the final database lock for this study (2nd October 2015), the time from the last second-line patient’s first dose was 60 months or more. The median (range) duration of OS follow up was 54.8 (0.6-96.3) months, and the median treatment duration was 25.6 (0.2-96.3) months (Table 1). Discontinuation from treatment was most common within the first two years, with 131 (46%) patients discontinuing by the end of year 2, and 38 (13%) patients discontinuing treatment in years 3 through 5. Discontinuation from treatment through year 5 were lack of efficacy [categorized by the investigator separately as PD and unsatisfactory response; n=47 (17%) and n=21 (7%), respectively], AE [n=64 (23%)], and patient request [n=19 (7%)]. The most common reasons for discontinuation from treatment through year 5 were lack of efficacy [categorized by the investigator separately as PD and unsatisfactory response; n=47 (17%) and n=21 (7%), respectively], AE [n=64 (23%)], and patient request [n=19 (7%)].
8%), and more likely to enroll in the extension study (52% vs. 19%) (Online Supplementary Table S3). Ninety-nine (55%) patients completed the 2-year follow up after treatment discontinuation (IM-R, n=60; IM-I, n=39). Thirty-two (11%) patients discontinued treatment after year 5, and 83 (29%) patients continued treatment in the extension study.

**Efficacy**

Most cytogenetic responses to bosutinib occurred within two years of initiating treatment (Table 2). By week 12, the cumulative major cytogenetic response (MCyR) rate was 85% (n=93 of 262 evaluable patients), including 22% (n=57) who attained/maintained a CCyR. Of 246 evaluable patients without a CCyR at baseline, 76 (31%) attained an MCyR and 45 (18%) attained a CCyR. The cumulative MCyR rate observed by year 2 was 58%, including 46% with a CCyR. Patients continued to attain a CCyR after two years, with 10 patients having initial on-treatment CCyR during years 3-5. By year 5, the cumulative MCyR CCyR rates were 60% (n=156 of 262 evaluable patients) and 50% (n=130), respectively; 57% (n=141 of 246 evaluable patients) newly-attained an MCyR and 47% (n=116) newly-attained a CCyR. The cumulative MMR rate was 42% (n=82 of 197 evaluable patients). Cytogenetic responses by both two and five years were similar in the IM-R and IM-I subgroups, whereas MMR rates were higher among IM-R patients at both time points (Table 2). Younger patients were more likely to have at least an MCyR (61% vs. 54%); however, rates of CCyR were similar among patients aged under 65 years and 65 years or over (Online Supplementary Table S3).

Among responders, the Kaplan-Meier estimated probability of maintaining a response was similar at years 5 and 2: 71% and 76%, respectively, for an MCyR; 69% and 75% for a CCyR; and 68% and 70% for MMR (Table 2 and Figure 1). Overall, 41 of 156 (26%) responders lost MCyR, 37 of 150 (25%) lost CCyR, and 25 of 82 (30%) lost MMR. Few patients lost response after year 2 (7 lost MCyR, 10 lost CCyR, and 2 lost MMR). At the last assessment prior to discontinuation, 81% (n=127) of all 156 responders had an MCyR and 68% (n=106) had a CCyR of 141 responders without a baseline CCyR, 111 attained an MCyR and 98 attained a CCyR.

Among 132 patients (IM-R, n=81; IM-I, n=51) who required a dose reduction to 400 mg/day due to an AE, 81 (61%; IM-R, 63%; IM-I, 59%) had an MCyR, including 67 of 110 (61%) who did not have a CCyR at baseline (Online Supplementary Table S4). Fifty-seven (43%) patients first achieved an MCyR after dose reduction, 19 (14%) achieved an MCyR before and maintained it after dose reduction, and 5 (4%) lost their MCyR after dose reduction. Among 50 patients (IM-R, n=32; IM-I, n=18) who had a dose reduction to 300 mg/day due to an AE, 29 (58%; IM-R, 69%; IM-I, 39%) had an MCyR, including 25 of 43 (58%) without a CCyR at baseline. Twenty (40%) patients achieved an MCyR before and maintained it after dose reduction, 8 (16%) first achieved an MCyR after a dose reduction.

### Table 1. Treatment summary.*

| Parameter | IM-R (n=195) | IM-I (n=89) | Total (n=284) |
|-----------|--------------|-------------|---------------|
| Median (range) duration of follow up, mo | 46.8 (0.6–96.3) | 58.8 (0.6–93.2) | 54.8 (0.6–96.3) |
| Median (range) duration of treatment, mo | 27.6 (0.2–96.3) | 24.2 (0.3–84.3) | 25.6 (0.2–96.3) |
| Patients with ≥1 dose interruption due to AEs, n (%) | 133 (68) | 76 (85) | 209 (74) |
| Median (range) duration of events of dose interruptions, d | 8 (1–981) | 14 (1–280) | 10 (1–981) |
| Median (range) cumulative duration of dose interruptions, d | 24 (1–983) | 24 (1–429) | 24 (1–983) |
| Patients with ≥1 dose reduction due to AEs, n (%) | 89 (46) | 52 (58) | 141 (50) |
| Median (range) time to first dose reduction due to AEs, d | 48 (8–2166) | 52 (7–1875) | 49 (7–2166) |
| Reduction to 400 mg/d | 49 (8–1800) | 55 (11–1875) | 52 (8–1875) |
| Reduction to 300 mg/d | 194 (58–2166) | 107 (29–1286) | 162 (29–2166) |
| Patients with ≥1 dose reduction due to AEs, n (%) | 349 (3–2881) | 283 (2–2881) | 346 (2–2881) |
| Median (range) cumulative duration of dose reduction due to AEs, d | 238 (3–2667) | 84 (1–2683) | 198 (1–2667) |
| Discontinued treatment, n (%) | 195 (100) | 89 (100) | 284 (100) |
| Enrolled in extension study | 61 (31) | 22 (25) | 83 (29) |
| AE | 32 (16) | 35 (39) | 67 (24) |
| PD | 43 (22) | 8 (9) | 51 (18) |
| Death | 12 (6) | 12 (13) | 24 (8) |
| Investigator request | 19 (10) | 4 (4) | 23 (8) |
| Patient request | 5 (3) | 3 (3) | 8 (3) |
| Unsatisfactory response (efficacy)† | 7 (4) | 1 (1) | 8 (3) |
| Lost to follow up | 0 (0) | 0 (0) | 0 (0) |
| Discontinuation of study by sponsor | 0 (0) | 0 (0) | 0 (0) |
| CCyR were similar among patients aged under 65 years and 65 years or over (Online Supplementary Table S3). Among responders, the Kaplan-Meier estimated probability of maintaining a response was similar at years 5 and 2: 71% and 76%, respectively, for an MCyR; 69% and 75% for a CCyR; and 68% and 70% for MMR (Table 2 and Figure 1). Overall, 41 of 156 (26%) responders lost MCyR, 37 of 150 (25%) lost CCyR, and 25 of 82 (30%) lost MMR. Few patients lost response after year 2 (7 lost MCyR, 10 lost CCyR, and 2 lost MMR). At the last assessment prior to discontinuation, 81% (n=127) of all 156 responders had an MCyR and 68% (n=106) had a CCyR of 141 responders without a baseline CCyR, 111 attained an MCyR and 98 attained a CCyR.

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reduction to 300 mg/day, and one (2%) patient lost MCyR. Among patients who achieved an MCyR after a dose reduction, median duration of response (non-Kaplan-Meier) was longer for patients receiving 400 mg/day versus 300 mg/day (167 days vs. 17 days); median MCyR duration (non-Kaplan-Meier) were similar in patients who had a response before and after dose reduction (283 days vs. 260 days) (Online Supplementary Table S4).

Of the 224 (79%) patients with a known baseline mutation status, 79 (35%) had at least one mutation in the BCR-ABL1 kinase domain, most of whom were in the IM-R cohort [IM-R, n=75 of 156 (47%); IM-I, n=6 of 61 (9%)] (Online Supplementary Table S5). Thirteen patients had multiple mutations, all of whom were IM-R. A total of 43 unique BCR-ABL1 mutations were evident, most commonly F359V (n=9), M551T (n=8), M244V (n=6), G250E (n=6), and T315I (n=9). Among evaluable patients with a mutation other than T315I, most (44 of 69, 64%) attained/maintained an MCyR; response rates were similar among patients without a mutation (75 of 130, 58%) and appeared lower among those with multiple mutations (6 of 12, 50%). Among evaluable patients with mutations that are sensitive (n=50), moderately resistant (n=12), and highly resistant (n=12) to bosutinib (see Figure 2 legend for definitions), the cumulative MCyR rates were 67%, 50%, and 53%, respectively, with corresponding CCyR rates of 57%, 42%, and 33%. Among evaluable patients with mutations of unknown sensitivity (n=24) and patients for whom mutation status was unknown (n=54), the MCyR rates were 63% and 65%, respectively, with corresponding CCyR rates of 50% and 57%.

Of 104 (37%) patients evaluated for BCR-ABL1 kinase domain mutations before and during bosutinib therapy, 26 had at least one newly-emerging mutation; this was most commonly T315I (n=9), V299L (n=5), and M244V (n=2). Of 104 (37%) patients evaluated for BCR-ABL1 kinase domain mutations before and during bosutinib therapy, 26 had at least one newly-emerging mutation; this was most commonly T315I (n=9), V299L (n=5), and M244V (n=2). Of 104 (37%) patients evaluated for BCR-ABL1 kinase domain mutations before and during bosutinib therapy, 26 had at least one newly-emerging mutation; this was most commonly T315I (n=9), V299L (n=5), and M244V (n=2). Of 104 (37%) patients evaluated for BCR-ABL1 kinase domain mutations before and during bosutinib therapy, 26 had at least one newly-emerging mutation; this was most commonly T315I (n=9), V299L (n=5), and M244V (n=2). Of 104 (37%) patients evaluated for BCR-ABL1 kinase domain mutations before and during bosutinib therapy, 26 had at least one newly-emerging mutation; this was most commonly T315I (n=9), V299L (n=5), and M244V (n=2). Of 104 (37%) patients evaluated for BCR-ABL1 kinase domain mutations before and during bosutinib therapy, 26 had at least one newly-emerging mutation; this was most commonly T315I (n=9), V299L (n=5), and M244V (n=2). Of 104 (37%) patients evaluated for BCR-ABL1 kinase domain mutations before and during bosutinib therapy, 26 had at least one newly-emerging mutation; this was most commonly T315I (n=9), V299L (n=5), and M244V (n=2). Of 104 (37%) patients evaluated for BCR-ABL1 kinase domain mutations before and during bosutinib therapy, 26 had at least one newly-emerging mutation; this was most commonly T315I (n=9), V299L (n=5), and M244V (n=2). Of 104 (37%) patients evaluated for BCR-ABL1 kinase domain mutations before and during bosutinib therapy, 26 had at least one newly-emerging mutation; this was most commonly T315I (n=9), V299L (n=5), and M244V (n=2). Of 104 (37%) patients evaluated for BCR-ABL1 kinase domain mutations before and during bosutinib therapy, 26 had at least one newly-emerging mutation; this was most commonly T315I (n=9), V299L (n=5), and M244V (n=2). Of 104 (37%) patients evaluated for BCR-ABL1 kinase domain mutations before and during bosutinib therapy, 26 had at least one newly-emerging mutation; this was most commonly T315I (n=9), V299L (n=5), and M244V (n=2). Of 104 (37%) patients evaluated for BCR-ABL1 kinase domain mutations before and during bosutinib therapy, 26 had at least one newly-emerging mutation; this was most commonly T315I (n=9), V299L (n=5), and M244V (n=2). Of 104 (37%) patients evaluated for BCR-ABL1 kinase domain mutations before and during bosutinib therapy, 26 had at least one newly-emerging mutation; this was most commonly T315I (n=9), V299L (n=5), and M244V (n=2). Of 104 (37%) patients evaluated for BCR-ABL1 kinase domain mutations before and during bosutinib therapy, 26 had at least one newly-emerging mutation; this was most commonly T315I (n=9), V299L (n=5), and M244V (n=2). Of 104 (37%) patients evaluated for BCR-ABL1 kinase domain mutations before and during bosutinib therapy, 26 had at least one newly-emerging mutation; this was most commonly T315I (n=9), V299L (n=5), and M244V (n=2). Of 104 (37%) patients evaluated for BCR-ABL1 kinase domain mutations before and during bosutinib therapy, 26 had at least one newly-emerging mutation; this was most commonly T315I (n=9), V299L (n=5), and M244V (n=2). Of 104 (37%) patients evaluated for BCR-ABL1 kinase domain mutations before and during bosutinib therapy, 26 had at least one newly-emerging mutation; this was most commonly T315I (n=9), V299L (n=5), and M244V (n=2). Of 104 (37%) patients evaluated for BCR-ABL1 kinase domain mutations before and during bosutinib therapy, 26 had at least one newly-emerging mutation; this was most commonly T315I (n=9), V299L (n=5), and M244V (n=2). Of 104 (37%) patients evaluated for BCR-ABL1 kinase domain mutations before and during bosutinib therapy, 26 had at least one newly-emerging mutation; this was most commonly T315I (n=9), V299L (n=5), and M244V (n=2). Of 104 (37%) patients evaluated for BCR-ABL1 kinase domain mutations before and during bosutinib therapy, 26 had at least one newly-emerging mutation; this was most commonly T315I (n=9), V299L (n=5), and M244V (n=2). Of 104 (37%) patients evaluated for BCR-ABL1 kinase domain mutations before and during bosutinib therapy, 26 had at least one newly-emerging mutation; this was most commonly T315I (n=9), V299L (n=5), and M244V (n=2).
or more after transformation: one patient who trans-
formed on day 14 continued bosutinib treatment for
another six years and subsequently continued treatment
in the extension study; another patient who transformed
on day 246 discontinued treatment two years later for PD.
Among 153 patients remaining on treatment after year 2,
only 2 (both IM-R) had on-treatment transformation to AP
after this time (on days 734 and 2165). Eleven of the 15
patients with on-treatment transformation had responses
to bosutinib, including 4 with best responses of MCyR, 3
with CCyR, and 4 with CHR. The cumulative incidence
of on-treatment PD/death was higher by 4% at year 5
[19% (IM-R, 23%; IM-I, 10%)] versus year 2 [15% (IM-R,
19%; IM-I, 7%)]; 42% of patients discontinued treatment

Figure 1. Duration of response. Duration of major cytogenetic
response (MCyR) (A) and complete cytogenetic response
(CCyR) (B) among responders. Open circles indicate censored
observations. IM-I: imatinib-intolerant; IM-R: imatinib-resis-
tant; n: number; d: day.
without on-treatment PD/death before year 5. Long-term outcomes are reported according to age in Online Supplementary Table S3.

Kaplan-Meier probability of OS at year 5 was 84% (IM-R, 81%; IM-I, 90%) versus 91% (IM-R, 88%; IM-I, 98%) at year 2; 40% of patients were censored prior to year 5, and 31% enrolled in the extension study for continued treatment or follow up for longer-term survival (Figure 3). A total of 45 (16%) deaths occurred on study; 24 through year 2, 5 after year 5, and 10 within 30 days of the last bosutinib dose. Patients aged under 65 years had a higher OS rate compared with those aged 65 years or over (85% vs. 82%); P = 0.0108.

**Tables and Figures**

**Figure 2. Predictors of response loss, disease progression, and death.** Closed circles represent major cytogenetic response (MCyR) duration and open circles represent complete cytogenetic response. Based on multivariate Cox regression models. Parameters failing to meet elimination criteria (0.20) not shown. Hazard ratios > 1 indicate worse outcome. P-values were not adjusted for multiple comparisons; significant P-values are in bold. Definitions of covariates can be found in the Online Supplementary Methods. On-treatment characteristics are Cox time-dependent covariates. *Baseline factor for durable response model. BOS: bosutinib; IM: imatinib; LFT: liver function test; Ph+: Philadelphia chromosome positive; y: years; CI: confidence interval.

**Figure 3. Kaplan-Meier estimated overall survival.** Open circles indicate censored observations. Overall survival was calculated as the first date of study dosing until the date of death; patients without events were censored at the last contact. Per protocol, patients were followed for overall survival for two years after treatment discontinuation. Analysis includes data from a long-term extension study. IM-I: imatinib-intolerant; IM-R: imatinib-resistant; n: number.
vs. 77%) (Online Supplementary Table S3). Causes of death were PD [n=26 (58%); IM-R: n=23; IM-I: n=3], AE unrelated to bosutinib [n=16 (36%); IM-R: n=14; IM-I: n=2], and unknown cause [n=3 (7%); all IM-I]. None of the 45 deaths were assessed as treatment-related. Four deaths occurred within 30 days of the last bosutinib dose through year 2 (all IM-R) and 4 occurred during years 3-5 (2 IM-R and 2 IM-I patients).

Predictors of response duration, PFS, and OS
Significant (P<0.05) baseline factors predictive of MCyR or CCyR loss were Ph⁺ ratio ≥95% versus ≤35% and late versus early disease stage (Figure 2). No on-treatment factors were predictive of MCyR duration; however, experiencing treatment-emergent thrombocytopenia was predictive of loss of CCyR (P=0.0471). Several baseline factors predictive of decreased OS were identified: age ≥65 years versus <65 years, Ph⁺ ratio ≥95% versus ≤35%, lack of an MCyR by week 12, higher baseline peripheral blood blasts, and having a BCR-ABL1 mutation at baseline that is either sensitive or highly resistant to bosutinib. Among on-treatment factors examined, experiencing an abnormal liver function test (LFT) was predictive of increased OS. Notably, prior response or resistance to imatinib did not predict duration of cytogentic response or long-term survival outcomes.

Factors predictive of decreased PFS were Ph⁺ ratio ≥95% versus ≤35%, lack of MCyR by week 12, and higher baseline peripheral blood blasts (Figure 2). The on-treatment factor of receiving a bosutinib dose reduction to 400 mg due to AEs was predictive of increased PFS.

Safety and tolerability
The most common any grade hematologic treatment-emergent AEs (TEAEs) were thrombocytopenia [42% (Grade 3/4, 25%)], anemia [29% (Grade 3/4, 13%)], and neutropenia [16% (Grade 3/4, 10%)] (Online Supplementary Table S6). The most common non-hematologic TEAEs were diarrhea [86% (Grade 3/4, 10%)], nausea [46% (Grade 3/4, 2%)], vomiting [37% (Grade 3/4, 4%)], and rash [36% (Grade 3/4, 9%)]. Most newly-occurring AEs (MedDRA preferred terms not reported for the same patient previously for those on treatment during a given year (1 year = 365.25 days). *Includes the high-level group terms (HLGTs) cardiac arrhythmias, pericardial disorders, and heart failures under the cardiac disorders system organ class (SOC); relevant preferred terms (PTs) (cardiac death, sudden cardiac death, sudden death) under the general disorders and administration site SOC conditions; relevant PTs (decreased ejection fraction, abnormal electrocardiogram QT interval, prolonged electrocardiogram QT, long QT syndrome, congenital long QT syndrome, torsade de points, ventricular tachycardia) under the SOC investigations.

Figure 4. Incidence of newly-occurring adverse events (AEs) over time. Denominators are the number of patients on treatment during the indicated years (NB: incidences of certain AEs appear higher in later years compared with previous years due to a lower number of patients on treatment). Newly-occurring AEs were those not experienced by the same patient previously among patients on treatment during a given year (1 year = 365.25 days). *Includes the high-level group terms (HLGTs) cardiac arrhythmias, pericardial disorders, and heart failures under the cardiac disorders system organ class (SOC); relevant preferred terms (PTs) (cardiac death, sudden cardiac death, sudden death) under the general disorders and administration site SOC conditions; relevant PTs (decreased ejection fraction, abnormal electrocardiogram QT interval, prolonged electrocardiogram QT, long QT syndrome, congenital long QT syndrome, torsade de points, ventricular tachycardia) under the SOC investigations. **HLGTs included: coronary artery disorders, atherosclerosis, stenosis, vascular insufficiency and necrosis, embolism and thrombosis; high-level terms (HLTs) included arterial therapeutic procedures (excluding aortic), central nervous system hemorrhages and cerebrovascular accidents, central nervous system vascular disorders not elsewhere classified (NEC), non-site specific vascular disorders NEC, peripheral vascular disorders NEC, and all subordinated terms. ***HLGTs included: vascular hypertensive disorders and cardiac and vascular investigations (excluding enzyme tests), the HLT vascular tests NEC (including blood pressure); PTs included: abnormal blood pressure, abnormal ambulatory blood pressure, increased systolic blood pressure, increased diastolic blood pressure, increased blood pressure, abnormal systolic blood pressure, and increased blood pressure. **HLT included: renal failure and impairment; PTs included: blood creatinine abnormal, blood creatinine increased, creatinine renal clearance abnormal, creatinine renal clearance decreased, glomerular filtration rate abnormal, glomerular filtration rate decreased. ALT: alanine aminotransferase; AST: aspartate aminotransferase; URTI: upper respiratory tract infection; UTI: urinary tract infection; n: number.
year) were experienced by patients during year 1 (99.6%) of treatment, with rates somewhat lower in years 2 (74%), 3 (68%), 4 (52%), and 5 (57%) (Figure 4). Common AEs (in ≥5 patients) newly-occurring in year 3 were cough [5% (n=8)], increased blood creatinine [5% (n=7)], and pyrexia [4% (n=6)]; most events were grade 1/2. Common newly-occurring AEs in year 4 were increased blood creatinine [5% (n=6)] and pleural effusion [5% (n=7)]; 2 events (both grade 4 pleural effusion considered probably not related to bosutinib) resulted in hospitalization. No newly-occurring AEs were reported in more than 5 patients in year 5.

Adverse events led to treatment discontinuation in 69 (24%) patients throughout the study, including one who also discontinued due to PD and another who discontinued due to subject request as the primary reason. AEs resulting in treatment discontinuation in 2% or more of patients overall were thrombocytopenia [6% (n=17)], neutropenia [2% (n=6)], and alanine aminotransferase increased [2% (n=6)]. Of these 69 patients, 28 (41%) discontinued treatment without attempting a dose reduction to less than 500 mg/day. The majority (86%) of discontinuations due to AEs occurred during the first two years of treatment (Online Supplementary Table S2). AEs led to treatment discontinuation in 7 patients in years 3-5: coronary artery disease, scleroderma, and renal failure in year 3; ascites and serositis (same patient), increased blood creatinine, and pulmonary hypertension in year 4; and thrombocytopenia in year 5.

Although diarrhea was the most common AE [IM-R, 86% (n=167); IM-I, 85% (n=76)], in most instances this was grade 1 or 2 [IM-R, 76% (n=149); IM-I, 75% (n=67)] (Online Supplementary Table S6). Only 12 (3%) patients had maximum grade 3/4 events and 14 (all grade; grade 3/4, n=1) had events considered by the investigator to be treatment related. The median (range) time to first renal AE was 673 (8-2695) days. Renal AEs led to treatment discontinuation in 3 patients (1 each in years 1, 2, and 3) and death in one patient (acute kidney injury in year 1 related to PD and unrelated to bosutinib).

Cross-intolerance

Eighty-nine patients were intolerant to prior imatinib (Online Supplementary Table S7). Of 85 patients with a specific AE reported as the reason for discontinuation of imatinib, 52 (61%) experienced the same AE with bosutinib that led to imatinib discontinuation, most commonly hematologic AEs (thrombocytopenia, n=12; neutropenia, n=5; anemia, n=5) or gastrointestinal AEs (diarrhea, n=6; nausea, n=4); 14 (16%) had cross-intolerance (defined as having discontinued bosutinib due to the same AE that led to prior imatinib discontinuation). Twenty-five (29%) patients experienced the same grade 3/4 AE while on bosutinib. No patient receiving bosutinib died due to the same AE that led to intolerance to prior imatinib.

Discussion

After five years of follow up, the final results of this phase I/II study demonstrated durable efficacy and acceptable long-term safety for second-line bosutinib in patients with CP CML resistant or intolerant to imatinib. The estimated probabilities of responders maintaining an MCyR or CCyR at year 5 (71%, 69%) decreased modestly from the estimated probabilities at year 2 (76%, 78%). Resistance and intolerance to prior imatinib did not appear to result in differences in response durability, as rates observed at years 2 and 5 were similar for both IM-R and IM-I patients. Additionally, late disease progression was uncommon, supporting the observed response durability [although 36 (13%) patients discontinued after year 5, potentially biasing the interpretation of subsequent outcomes]. Cumulative response rates at years 5 and 2 were similar (year 5: MCyR, 60% and CCyR, 50%; year 2: MCyR, 58% and CCyR, 46%). However, it should be noted that results reported here are based on a finalized database resulting in slight differences from previously published data.10

The response rates achieved in this study are compara-
ble to those observed in studies of second-line nilotinib and dasatinib. With similar follow-up durations, CCyR rates of 57% and 49% were reported with nilotinib and dasatinib, respectively, compared with 47% with bosutinib in the present study.4,8,13 Estimated rates of on-treatment PD/death (19%) and transformation to AP/BP CML (5%) remain low with bosutinib; only 2 IM-R patients had on-treatment transformation to AP after year 2, although there is a potential bias from patients lost to follow up. Similar rates of transformation were observed with second-line dasatinib (5%).4,11 The estimated OS rate at 5 years is high, with a modest decrease from the 2-year OS rate (84% vs. 91%). This 5-year rate is also comparable to those reported for dasatinib (91%), nilotinib (87%), and ponatinib (81%) in CP CML patients after prior TKI failure.8

Responses were observed in all but 2 (T315I and M244V) of the 26 patients with newly-emerging BCR-ABL1 mutations. All but 14 of 14 patients with newly-emerging mutations that are highly resistant to bosutinib13 had a best response of at least CHR; 5 (36%) had a best response of at least PCyR. Effects of dose reductions on response were limited as most patients who dose reduced dose attained/maintained an MCyR. Only 4% and 2% of patients who reduced dose to 400 mg/day and 300 mg/day, respectively, lost their previously achieved MCyR.

Gastrointestinal toxicities remained the most commonly reported AEs overall at the 5-year follow up (diarrhea, 86%; nausea, 46%; vomiting, 37%). Initial events occurred early, with incidences through year 2 of 84% for diarrhea, 45% for nausea, and 37% for vomiting.11 Although diarrhea was common, grade 3 events occurred in only 10% of patients (no grade 4), and only 4 patients discontinued because of this AE, all within two years of initiating bosutinib. Grade 3/4 hematologic AEs, such as thrombocytopenia (25%) and neutropenia (10%), occurred at rates similar to or lower than those observed with second-line dasatinib (24% and 36%), nilotinib (30% and 51%), and ponatinib (55% and 23%).4,5,8 Rates of cross-intolerance between bosutinib and prior imatinib were low, suggesting that most patients intolerant to imatinib therapy may be successfully treated with bosutinib.

Given the long-term nature of TKI therapy, late-emerging toxicities are of concern, particularly cardiac and vascular events. In a study of bosutinib versus imatinib as first-line treatment for CP CML, the incidence of cardiac and vascular AEs with bosutinib was low and similar to that of imatinib.1,3 In the present study, the incidence of newly-occurring cardiac and vascular AEs with second-line bosutinib remained low after year 2. However, most (85%) discontinuations due to AEs as the primary reason occurred within the first two years; thus, patients remaining on treatment after year 2 may have a lower risk of experiencing these events. The incidence of renal AEs, while low, remained similar in years 3-5. Bosutinib has been associated with a decrease in glomerular filtration rate that is typically modest and potentially reversible (similar to what has been reported with imatinib).19,20 Dose adjustments are recommended in patients with baseline and treatment-emergent renal impairment.6,19 Careful monitoring, supportive care, and prompt management of toxicities may allow patients to continue treatment long term.

Most baseline and on-treatment factors examined appeared not to be predictive of response duration, OS, or PFS. Baseline Ph’ ratio ≤35% (vs. ≥55) was associated with all 3 types of long-term outcomes (MCyR duration but not CCyR duration). Lower percentage of peripheral blood blasts at baseline and MCyR by week 12 were associated with both improved OS and PFS. Having a baseline BCR-ABL1 mutation, regardless of sensitivity to bosutinib, was predictive of decreased OS and, interestingly, having an abnormal LFT on-treatment was predictive of increased OS. This unexpected result may be due to increased bosutinib exposure levels resulting from the underlying cause of the abnormal LFT, leading to an increase in efficacy; however, population pharmacokinetics modeling from this study has found no relationship between baseline LFTs and bosutinib pharmacokinetics. Notably, prior response or resistance to IM did not predict any long-term outcomes. Because P-values were not adjusted for multiple comparisons, marginally significant P-values should be interpreted with caution.

The potent and durable activity and distinct toxicity profile of bosutinib confirm it is an important option for treating CML patients in the second-line setting, as demonstrated by its long-term efficacy and safety in these patients; a 10-year follow up is planned for patients enrolled in an ongoing extension study.

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