Switching to second-generation tyrosine kinase inhibitor improves the response and outcome of frontline imatinib-treated patients with chronic myeloid leukemia with more than 10% of BCR-ABL/ABL ratio at 3 months

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Abstract
Chronic myeloid leukemia patients display heterogeneous responses to imatinib. Survival depends on baseline clinical characteristics (including prognostic scoring systems) and on early response (such as >10% BCR-ABL/ABL ratio at 3 months of therapy). The results of switching to second-generation tyrosine kinase inhibitors (2GTKIs) may contain a bias since, in the majority of these studies, patients who switch treatment due to intolerance or failure are censored or excluded. We analyzed the Spanish Registry data on switching in an intention-to-treat analysis of patients in standard clinical practice. Switching to 2GTKIs improves responses from 45% to 75% of complete cytogenetic response (CCyR) and from 15% to 45% of major molecular response (MMR) in the group without molecular response 1 (MR1) at 3 months and from 70% to 87% in CCyR and from 52% to 87% in MMR in the group with MR1. The final response rate is poorer in the group with no MR1 at 3 months. Nevertheless, the differences in the rates of response were not translated into differences in major events (transformations or deaths), and the final progression-free survival and overall survival were similar.

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**Introduction**

The introduction of tyrosine kinase inhibitors (TKIs) has proved to be a major advance in the management of patients with chronic myeloid leukemia in chronic phase (CML-CP). Imatinib therapy has radically changed the prognosis for patients with CML, and most patients enjoy a near-normal life expectancy [1]. Recently, efforts have been made to define molecular markers, or clinical milestones, that predict the course of patient outcomes more reliably than cytogenetics but, to-date, such findings are not fully reproducible. CML patients display heterogeneous response to therapy, with survival often depending on baseline clinical characteristics. Until recently, the stratification was made according to two scores developed in CML patients treated with chemotherapy [2], and interferon (Euro) [3]. In 2011, the European Leukaemia Net (ELN) evaluated more than 2000 patients with early chronic phase (ECP) CML treated with front-line imatinib; the objective was to develop a new prognostic scoring system: the European Treatment and Outcome Study (Eutos) score [4]. Since the publication of the system, however, there have been conflicting reports on the predictive value of the Eutos score. The Spanish RELMC (Registro Español de Investigación y Tratamiento de Leucemia Mieloide Crónica) is a hospital-based registry, which includes only out-of-study patients. Monitoring and treatment are at the physician’s discretion, and do not reflect any clinical pathway. There is no common protocol and no common recommendations since the aim of the Registry is to monitor the current practice of CML management in Spain.

Recently, a cut-off value of BCR-ABL/ABL ratio of >10% at 3 months has been proposed to discriminate patients with a poorer outcome when treated with imatinib [5], dasatinib [6], nilotinib [7], and bosutinib [8]. The aims of the present study were to discriminate the influence of this response variable on outcomes in CML patients, and to assess the value of preclinical scores (Sokal, Euro, and Eutos) in this model. In addition, data are limited on how this outcome is affected by the change to second-generation TKI (2GTKI). The switching to 2GTKIs introduces a bias in the majority of studies because patients who switch treatment due to intolerance or failure are censored and, as such, “increasing” the probability of response in their group [9]. To avoid this bias, the variable “time-to-imatinib failure” was introduced into our outcomes analysis.

**Methods**

**Patients and therapy**

We retrospectively analyzed 374 patients with CML in first CP (treated with imatinib as first choice, and outside of clinical trials). Table 1 summarizes patients’ characteristics. Complete hematologic response (CP), complete cytogenetic response (CCyR), and major molecular response (MMR) were defined using conventional criteria. Follow-up and treatment decisions were decided by hematologists according to their own clinical judgment, that is, no preset protocol. Cytogenetic or molecular data were included until the time that imatinib treatment was discontinued, for whatever reason, or at last follow-up.

**Detection of BCR-ABL1 transcripts**

BCR-ABL1 transcripts were measured in venous blood samples taken at 12-week intervals, using real-time quantitative polymerase chain reaction (RTQ-PCR) as described previously [10]. Results were expressed as percentage ratios relative to an ABL1 internal control. BCR-ABL1 transcript samples were not centralized but all RTQ-PCR analyses were performed in the same laboratory for each patient during follow-up. In our registry of patients treated with frontline imatinib, 156 (41.7%) of 374 had a molecular ratio of BCR-ABL1/ABL1 determined at 3 months. During follow-up, 43 of 156 patients discontinued imatinib and had switched to receive dasatinib (n = 30), or nilotinib (n = 13), while five patients were treated with two TKI’s; dasatinib–nilotinib in sequence (n = 4) and nilotinib–dasatinib in sequence (n = 1).

**Table 1. Patient characteristics.**

|                          | All patients | Patients with RTQ-PCR at 3 months |
|--------------------------|--------------|----------------------------------|
| N                        | 374          | 156/374 (42%)                    |
| Duration imatinib (days) | 19 (1–180)   | 19 (1–180)                       |
| Male/Female              | 229 (61%)/145 (39%) | 93 (60%)/63 (40%)               |
| Age, years (range)       | 52 (15–88)   | 54.2 (15–86)                     |
| Sokal index              |              |                                  |
| Low                      | 138 (39%)    | 62 (40%)                         |
| Intermediate             | 172 (48%)    | 78 (50%)                         |
| High                     | 44 (12%)     | 15 (10%)                         |
| Euro index               |              |                                  |
| Low                      | 170 (48%)    | 82 (53%)                         |
| Intermediate             | 165 (47%)    | 65 (42%)                         |
| High                     | 19 (5%)      | 6 (5%)                           |
| Eutos index              |              |                                  |
| Low                      | 349 (91%)    | 138 (91%)                        |
| High                     | 30 (9%)      | 14 (9%)                          |

RTQ-PCR, real-time quantitative polymerase chain reaction.
Those patients with treatment failure at 3 months, as defined by ELN06, were excluded from the statistical analyses. Probabilities of overall survival (OS), progression-free survival (PFS), and time-to-imatinib failure (TIF) were calculated using the Kaplan–Meier method. In TIF, an event was defined as a progression to advanced phase, death, or imatinib discontinuation (for whatever reason). For PFS, the corresponding events were death, progression to accelerated phase (AP), or blast crisis (BC) [11]. The probabilities of cytogenetic and molecular responses were calculated using the cumulative incidence procedure. The log-rank test or a Cox regression model was used to compare OS, PFS, and TIF. Univariate and multivariate analyses were performed in accordance with standard methods; variables with \( P < 0.10 \) in univariate analyses were entered in the multivariate analysis. Unless stated in the text, all the analyses were performed on an intention-to-treat basis. Differences between groups were evaluated by the chi-square test and Mann–Whitney \( U \) test for categorical and continuous variables, respectively. Survival probabilities were estimated by the Kaplan–Meier method and compared with the log-rank test. Univariate and multivariate analyses were performed to identify potential factors predicting early cytogenetic responses. Multivariate analysis of response used the logistic regression model and, for time-to-event outcomes, we used the Cox proportional hazard regression.

**Results**

The median follow-up in the current series of patients was 54 months (range: 1–174 months). The distributions of BCR-ABL/ABL ratios at 3 months according to Sokal, Euro, and Eutos are summarized in Table 2. Sokal, Euro, or Eutos risk scores were not significantly associated with cut-off points, but the proportion of patients with ratio >10% was higher in Sokal and Euro high-risk groups.

**Response to imatinib**

The rate of best CCyR and MMR reached with imatinib therapy were 81% and 75.4%, respectively. The data are summarized in Table 3. The percentages of subsequent best CCyR and MMR in the group with MR1 at 3 months were significantly higher than those of patients not achieving MR1 (95% confidence interval [CI]: 92–88% vs. 77–58%; \( P < 0.001 \)).

**Time-to-imatinib failure**

During follow-up (median 80 months), 43 patients had their therapy changed to a 2GTKI. Of these, 30 were assigned to dasatinib, 13 to nilotinib, while five patients were treated with two TKIs: dasatinib–nilotinib in sequence (\( n = 4 \)) and nilotinib–dasatinib in sequence (\( n = 1 \)). The outcome was an 8-year probability of imatinib failure-free survival of 60%. In the group with MR1 at 3 months, 23 (22%) of 106 patients switched to 2GTKIs because of lack of efficacy (\( n = 4 \)), intolerance (\( n = 10 \)), and loss of response (\( n = 9 \)). The distribution was different in the group that did not achieve MR1, that is, intolerance (\( n = 8 \)), loss of response (\( n = 1 \)), and lack of efficacy (\( n = 11 \)) for a total of 20 (40%) of 50 patients (Table 4).

In univariate analysis, only having MR1 at 3 months was significantly associated with higher probability of imatinib treatment failure (long-rank test \( P = 0.026 \)) (Fig. 1) and this variable was the only one associated with poorer outcome in multivariate analysis (hazard ratio: 1.91; 95% [CI]: 1.61–2.20; \( P = 0.027 \)) (Table 5).

**Response rate to 2GTKIs after switching**

Switching to 2GTKIs improves the mean rates of response in both groups (from 45% to 75% of CCyR and from 15% to 45% of MMR in the group without MR1 at 3 months and from 70% to 87% in CCyR and from 52% to 87% in MMR in the group with MR1) (Table 3). This improvement in mean response rate with switching was reflected in the best response rates obtained in the groups; the CCyR and MMR were 92% and 88% in the group with MR1 at 3 months and the corresponding values were 77% and 58% in the non-MR1 group (\( P < 0.001 \)). Nevertheless, the rates of response were not translated into differences in major events (transformations or deaths).

### Table 2. Index score distributions according to BCR-ABL/ABL ratio at 3 months.

| Risk score | Sokal index | Euro index | Eutos index |
|------------|-------------|------------|-------------|
|            | Low | Intermediate | High | Low | Intermediate | High | Low | High |
| \( \leq 10% \) | 45/62 (73%) | 53/78 (68%) | 8/15 (53%) | 59/82 (72%) | 42/65 (65%) | 3/6 (50%) | 93/138 (67%) | 10/14 (71%) |
| 106/156 (68%) | 17/62 (27%) | 25/78 (32%) | 7/15 (47%) | 23/82 (28%) | 23/65 (35%) | 3/6 (50%) | 45/138 (33%) | 4/14 (29%) |
| \( > 10% \) | 50/156 (32%) | 0.353 | 0.402 | 0.758 |
Table 6. The final PFS and OS were similar in both groups (Fig. 2).

**Discussion**

The observation of prognostic significance of response to TKI after 3 months of the start of therapy has triggered a debate on the optimal management of these patients. Currently, there are no data available from prospective studies suggesting that molecular response-driven change in therapy among these patients alters the final outcome. The NCCN has recommended that these patients should be offered a change in therapy [12], but ELN2013 considered this status as a need for caution [13].

Despite the prognostic importance of MR1 at 3 months, the actual incidence of MR1 has not been clearly established in real practice. When we analyzed prospective studies comparing imatinib with 2GTKIs (ENESTND, DASISION, and BELA trials) the proportion of patients with no MR1 at 3 months in the imatinib group was similar (33.3%, 36%, and 34.5% in the three trials, respectively) [6–8]. We have seen a similar rate (32%) in our study, confirming that at least one third of patients will fail to respond to the treatment (assuming that the proportion of patients at high-risk is comparable with our study).

Our population sample had a lower percentage of patients with a high-risk index (Sokal, Eutos, or Euro score) at diagnosis than that reported in other series. These scores seem to be good predictors of response at 3 months. Patients with high-risk score at baseline had a higher probability of failing to achieve MR1 at 3 months. Moreover, despite the negative influence of the diagnostic score, the prognostic weight does not seem to be maintained if patients achieved MR1 at 3 months; and this response is the only variable to remain significant in the multivariate analysis.

All changes in treatment decisions were made according to the ELN recommendations; most patients changed at different times and for different reasons [14, 15]. However, no patient in this study switched before 3 months (we excluded patients from the statistical analysis who were considered failures according to the ELN06 recommendations at 3 months), and many of them had high-risk scores. To the best of our knowledge, our study is the first to describe the actual treatment given to the patients when BCR-ABL/ABL ratio at 3 months is the variable of interest. Other studies have analyzed the outcomes with respect to this ratio, but they have not reported the influence of the change in therapy, and they do not provide information about the specific treatment given as second line. The only exception is the TIDEL II study, the intervention in which depends on specific
molecular markers [16]. We can conclude from our results that these changes in therapy (at different times depending on the different recommendations) have enabled a significant number of patients to improve their response rates. However, the final outcomes in terms of PFS and the OS are similar in all patients (despite the poor initial forecast in patients with no MR1 at 3 months).

Our results show that the 10% threshold at 3 months discriminates, in a statistically significant manner, the probability of obtaining CCyR and MMR subsequently, not only when first-line imatinib is considered but also

| BCR-ABL/ABL ratio at 3 months | Switch 2GTKIs | Causes | Time to switch (years) |
|-------------------------------|--------------|--------|-----------------------|
|                               |              | Intolerance | No optimal response | Loss of response | 1   | 2   | 3   |
| ≤10%                          | Yes: 23/106 (22%) | 10/23 (43%) | 4/23 (17%) | 9/23 (39%) | 10/23 (43%) | 5/23 (22%) | 8/23 (35%) |
| 106/156 (68%)                 | No: 83/106 (88%) | 10/23 (43%) | 5/23 (22%) | 5/23 (22%) | 8/23 (35%) |
| More than 10%                 | Yes: 20/50 (40%) | 8/20 (40%) | 11/20 (55%) | 1/20 (5%) | 8/20 (40%) | 7/20 (35%) | 5/20 (25%) |
| 50/156 (32%)                  | No: 30/50 (60%) | <0.017 | 0.008 |

2GTKI, second generation tyrosine kinase inhibitor.
when all lines of TKI treatment are taken into account. However, this did not translate in to differences in OS or in PFS. This lack of difference in survival outcomes between the two groups has not been concordant in all recent studies. Hanfstein et al. [17] reported significantly better PFS and OS probabilities in patients who achieved MCyR or BCR-ABL/ABL ratio ≤10% with 3 months of i-matinib therapy, while Marin et al. [5] reported that achieving BCR-ABL-ABL ratio ≤9.84% at 3 months was linked to higher probabilities of CCyR, MMR, MR, PFS, and OS in patients treated with imatinib. In all arms of the ENESTND trial, early molecular response failure (>10% BCR-ABL ratio at 3 months) was associated with lower rates of molecular response, an increased risk of progression, and lower OS compared to those achieving early molecular response [7]. In the DASISION study [6], early molecular response failure was associated with lower rates of molecular response, an increased risk of progression, and lower PFS and OS.

None of these studies have described the treatment of patients after imatinib failure. The influence of switching treatment on the response and on the survival outcomes were not explored (data were not reported and patients censored for response). Furthermore, the influence of early treatment change was not analyzed. As such, it is important to highlight that, in the ENESTND trial, early

| Covariates                | β    | SE   | Wald | df  | P    | HR  |
|---------------------------|------|------|------|-----|------|-----|
| Gender                    | −0.060 | 0.295 | 0.041 | 1   | 0.839 | 0.942 |
| Sokal risk group          | 0.116 | 0.277 | 0.175 | 1   | 0.675 | 1.123 |
| Euro risk group           | 0.306 | 0.312 | 0.964 | 1   | 0.326 | 1.358 |
| Eutos risk group          | −0.172 | 0.470 | 0.134 | 1   | 0.714 | 0.842 |
| Ratio BCR-ABL/ABL at 3 months (>10% vs. ≤10%) | 0.649 | 0.295 | 4.859 | 1   | 0.027 | 1.914 |

SE, standard error; df, degrees of freedom; HR, hazard ratio.

| BCR-ABL/ABL ratio at 3 months | AP | BC | Deaths | Deaths or transformations |
|-------------------------------|----|----|--------|--------------------------|
| ≤10%                          | 2/106 (2%) | 1/106 (1%) | 4/106 (4%) | 7/106 (7%) |
| >10%                          | 0/47 (0%)  | 1/50 (2%)  | 3/50 (6%)  | 4/50 (8%)  |

P-value 0.084 0.584 0.531 0.751

progressions were common in the imatinib arm (7 of the 15 patients progressed between 3 and 6 months, representing a rate of 2.4% in 3 months). In the IRIS trial, the first annual rate of progression to AP and BC [11] was 1.5% [18].

Conversely, in a recent analysis of all patients from clinical trials, the MD Anderson group reported the same lack of difference we found in our study. They analyzed early cytogenetic response (at 3 and 6 months) and observed differences in time-to-failure, but not in OS [19]. In their study, only one patient progressed to BC at 3 months and no transformations occurred between 3 and 6 months. None of the patients had switched treatment in that period while, in our study, up to 40% switched within the first year.

In our series, switching to 2GTKI, although associated with significant improvement in response rates, does not
appear to overcome the poor outcome prediction of the 10% cut-off at 3 months; the response rate is poorer in the group with no MR1 at 3 months. The reason for lack of translation of the difference in responses into probabilities of better PFS or OS is not clear. It could reflect the efficacy of the treatment in third line and beyond. The explanation is hampered by the reason(s) for switching being different in the two groups, that is, more secondary resistance in the MR1 group, more primary resistance in the group without MR1 at 3 months.

In summary, our results confirm (in real-life practice within the setting of an intention-to-treat analysis and with a long follow-up) that not obtaining a BCR-ABL/ABL ratio of ≤10% at 3 months is a warning sign because it compromises the potential response to treatment even when switching to a 2GTKI. This emphasizes the importance of evaluating new treatment approaches when this threshold is not met.

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**Conflict of Interest**

Casado—Novartis, BMS, and Pfizer: Consultancy, Speakers Bureau. García-Gutiérrez—Novartis, BMS, and Pfizer: Membership on the entity’s Board of Directors or advisory committees. Martínez-López—Celgene: Honoraria; Steegmann—Novartis, BMS: Consultancy, Membership on the entity’s Board of Directors or advisory committees, Speakers Bureau; Pfizer: Consultancy, Speakers Bureau.

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