The Clinical Impact of Low Doses of Dasatinib in Patients with Chronic Myeloid Leukemia

Salih AKSU1, Fahri SAHIN2, Burak UZ1, Selim A. YAVUZ3, Hilmi ATAY4, Engin KELKITLI4, Mehmet TURGUT4, Mustafa PEHLIVAN5, Meltem O. AKAY6, Emel GURKAN7, Muzaffer DEMIR8, Selda KAHRAMAN9, Fatih DEMIRKAN9, Senay PAYDAS7, Güray SAYDAM9, Ibrahim C. HAZNEDAROGLU6

1 Hacettepe University Faculty of Medicine, Department of Hematology, Ankara
2 Ege University, Faculty of Medicine, Department of Hematology, Izmir
3 Istanbul University, Faculty of Medicine, Department of Hematology, Istanbul
4 Ondokuz Mayıs University, Faculty of Medicine, Department of Hematology, Samsun
5 Gaziantep University Faculty of Medicine, Department of Hematology, Gaziantep
6 Osmangazi University, Faculty of Medicine, Department of Hematology, Eskisehir
7 Cukurova University, Faculty of Medicine, Department of Hematology, Adana
8 Trakya University, Faculty of Medicine, Department of Hematology, Edirne
9 Dokuz Eylül University, Faculty of Medicine, Department of Hematology, Izmir, TURKEY

ABSTRACT

We report our experience in 41 patients with chronic phase (CF)-chronic myeloid leukemia (CML) who had discontinued imatinib switched to dasatinib, retrospectively. The CF-CML patients received dasatinib at starting dose of 100 once daily. Dose adjustment were observed in 11 patients, respectively. In case of other circumstances, treatment has been continued with a lower dose if needed. The median dose of dasatinib was 100 mg daily (range: 50 to 140 mg). We conclude that even low-dose dasatinib therapy is an effective and safe in second line treatment of CML patients.

Keywords: Chronic myeloid leukemia, Dasatinib, Treatment dosage

ÖZET

Kronik Myelositer Lösemi Tedavisinde Düşük Doz Dasatinib’in Klinik Etkinliği

İmatinib tedavisinde aylan faz kronik miyeloid lösemi tanısı alan ve dasatinib tedavisine geçen 41 tam yanıt- li olduğu bilinen ve son 1 yılın aylık olarak aldığı dozlan bilinen hasta restrospektif olarak bir yıl süresince hastaların ortalaması dasatinib dozlan araştırma. Bu hastaların hepsinde dasatinib tedavi başlangıc dozu 100 mg/gün idi. Onbir hastada doz ayarlanmas yapıldığını gözlemdi. Çeşitli nedenlerden dolayı, ihtilaç olunduğunda tedaviye daha düşük bir dasatinib dozu ile devam edildiği gözlenmiştir. Medyan dasatinib dozu 100 mg idi (aralık: 50-140 mg). Sonuç olarak, düşük doz dasatinib bile KML hastalarının ikiçini basamak tedavisinde hastalarda etkin ve güvenilir bir tedavi seçeneğidir.

Anahtar Kelimeler: Kronik miyeloid lösemi, Dasatinib, Tedavi dozlarını
INTRODUCTION

Chronic myeloid leukemia (CML), a clonal myeloproliferative disorder of blood stem cells, has been caused by a genetic abnormality which is the breakpoint cluster gene between chromosomes 9 and 22. Tyrosine kinase inhibitors (TKIs) including imatinib, dasatinib and nilotinib which have created the opportunity for tailored CML treatment and if eligible stem cell transplantation (SCT) are current treatment options for CML. In Turkey, the current standard first-line therapy for patients with newly-diagnosed chronic phase-CML is imatinib. Dasatinib and nilotinib have still been using in clinical practice to manage the treatment of CML patients who had resistance and intolerant to imatinib. Thus, TKIs are essential for the treatment of CML disease and it’s continuity is also crucial.

The patients with CML frequently need transient treatment interruptions and dose reductions during their treatment with 2nd-generation TKIs due to adverse events, patient’s compliance problems. Dasatinib (Sprycel; N-(2-Chloro-6-methylphenyl)-2-[[6-[[4-(2-hydroxyethyl)-1-piperazinyl]-2-methyl-4-pyrimidinyl]amino]-5-thiazolcarboxamide monohydrate, Bristol-Myers Squibb) is a potent, orally ABL kinase inhibitor and binds to both the active and also inactive con-formation of the ABL kinase domain. Nilotinib (Tasigna; N-[3-[3-(1H-imidazolyl)propoxy] phenyl]-4-methyl-3-[[4-(3-pyridinyl)-2-pyrimidinyl]amino] benzamide; Novartis Pharmaceuticals), a novel orally bioavailable derivative of imatinib, is a tyrosine kinase inhibitor with improved target specificity. Like imatinib, nilotinib inhibits Bcr-Abl by binding to only an inactive conformation of the ABL kinase domain.

Low adherence to long-term maintenance therapy is a known challenge in a number of chronic diseases. Adherence among patients with chronic diseases averages only 50%. In CML, one study indicated that 51% of the patients take < 85% of their prescribed dose within the first year of their treatment. The correlation between poor adherence to imatinib and clinical outcomes including poor response rates, high loss of response rates has been previously demonstrated among CML patients in clinical study settings. Yood et al showed that in the patients who received nilotinib as a 2nd-line treatment, adherence problems have been reported almost two times more compared with dasatinib ≤ 100 mg/day. 10 Age, sex, number of concomitant medications, number and type of comortibities, dosing schedule (e.g. once vs twice/ day), dosing restrictions (e.g. taking with / without food), and/or number intensity of adverse events could influence patients adherence.

Treatment complexity is another factor that may affect adherence to BCR-ABL inhibitors. To decrease the risk of gastrointestinal irritation, imatinib doses should be taken with a meal and a large glass of water and the dose of imatinib 400 mg/day. The recommended daily dose of nilotinib (800 mg/ day) requires twice-daily dosing and doses should be taken ~12 hours apart. Because of the following, no food should be consumed 2 hours before and 1 hour after each nilotinib dose. However, the recommended daily dose of dasatinib (100 mg/day) can be taken in a single dose without fasting and with or without a meal. In case of severe AEs, reduced dose of imatinib, nilotinib and dasatinib are 300, 400 and 80 mg/day. However, dasatinib has a rich diversity of tablet formulation like 20, 50, 70, 80, 100 mg. Thus, it is the only tyrosine kinase inhibitor which has adjustable dose application.

The use of lower dose of TKIs may be resulted in reduced efficacy of therapy. There are several studies in order to clarify this correlation when using imatinib for patients with CML. Sneed et al. has stated that the patients who have received lower dose of imatinib have a worse cytogenetic response rate. In addition, the higher initial imatinib doses may cause the better response rates, and more adverse events. The incidences of myelosuppression and pleural effusion were significantly lower with dasatinib 100 mg daily compared with dasatinib at 70 mg twice daily. Close monitoring and timely intervention are necessary for patients at risk for developing pleural effusions. One study showed that in patients with CML who are resistant or intolerant to imatinib, lymphocytosis has been reported during dasatinib treatment and has been associated with increased incidence of pleural effusion. Cytophenic response rates to dasatinib were presented as higher in the group of patients with pleural effusion. DASISION study has also indicated the similar results.
Visani et al. demonstrated that dasatinib, even at low doses, induces and maintains responses (cytogenetic, hematologic and molecular) in real-life CML patients who were resistant and/or intolerant to imatinib treatment. In addition, the “weekend drug holiday” in patients who developed pleural effusion during dasatinib treatment, a reduced schedule (3-5 days on and 4/2 days off), showed the same efficacy. In this study, we thus analysed the significance of dose reductions and treatment interruptions among patients treated with dasatinib.

MATERIALS, METHODS AND RESULTS
Forthy-one patients who have not entered blastic or accelerated phase in the last year and using dasatinib for at least one year, registered to the CML enrollment study between the dates 24.07.2007 and 13.03.2012 with known dosages of their medication on a monthly basis and who are CML patients with "Complete" response and in chronic phase for the last year meet these criteria among the 132 CML patients treated by dasatinib at nine universities which are located in Turkey. Dasatinib doses that were taken in the last 12 months starting from the last records, patient age and gender, reason of the change in dose if exists, dasatinib initiation dose, reason to switch to the second level TKI, last dose of the imatinib, and if the mutation analysis exists or not were identified in the database. Mean dasatinib dose of these patients, monthly and in the last year and mean drug costs monthly and yearly were calculated from the refunding foundation point of view. Effect of the clinical and demographic factors found in this database to the decrease in the dosage was assessed. Mean drug costs were calculated by calculating the mean of the monthly used medications that correspond to each patient's 12 months medication dose. Monthly collected daily dosage data were calculated for each patient one by one and mean cost was calculated with respect to the Social Security Institution point of view including a month corresponds to a day. SPSS for Windows version 17 was used in all of the analysis. Demographic information of the patients are presented in Table 1. Average age is 50 and 46.3 % of the patients are women.

Mean dosage of the patients regarding the months are presented in Table 2. While the mean dosage was 92 in the first month, this mean was observed as 90 in the last month.

The line graph regarding the mean dasatinib dosages patients were using was presented below (Figure 1). It can be observed from the graph that the used dasatinib dosage is presenting a decreasing trend.

The frequencies of the reasons of the changes in doses are reported as 11 (26.8%) patients with change in doses. Mean dasatinib is 92 mg that the patients were initially using. Patients were observed to use minimum 40 mg, maximum 140 mg dasatinib at the study initiation (Table 3).

The reason of the patients’ switching to the second line TKI are presented in Table 4. 80.4% of the patients were observed to switch to the second line TKI because of inefficacy, 7.3% of them switched
because of no response, 9.8% of them switched because of intolerance problem and adverse effects.

Descriptive statistics regarding the last imatinib dosages that were used by the patients were presented in Table 5. Mean dose of imatinib that the patients last used is 486 mg. The imatinib dosage that were lastly used by the patients were minimum 300 mg and maximum 800 mg.

It is observed that mutation analysis was performed in 43.9% of the patients and was not performed in 56.1% of the patients that were enrolled to the study (Table 6).

Average dosage cost per month is presented in Table 7. While the dosage cost of the patients were 153 TL initially, it was observed to be approximately 148 TL in the last month of study. On this respect, we can claim that monthly costs decrease with the dose reduction and daily costs are reduced down to 148 TL in patients that are stable or complete response. Annual average cost is 152 ± 3 TL. This explains providing the cheapest treatment choice in the 2nd level in chronic CML treatment.

The reduction in average dose cost of patients are presented in Figure 2.

**Table 3. Dasatinib initiation dose, mg/day**

<table>
<thead>
<tr>
<th>Statistics</th>
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<tbody>
<tr>
<td>Mean ± Std. Deviation, mg/day</td>
<td>92 ± 23</td>
</tr>
<tr>
<td>Minimum</td>
<td>40</td>
</tr>
<tr>
<td>Maximum</td>
<td>140</td>
</tr>
</tbody>
</table>

**Table 4. Reasons to switch to the second line TKI**

<table>
<thead>
<tr>
<th>Reason</th>
<th>f (%)</th>
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</thead>
<tbody>
<tr>
<td>Inefficacy, secondary resistance</td>
<td>33 (80.4%)</td>
</tr>
<tr>
<td>No response, primary resistance</td>
<td>3 (7.3%)</td>
</tr>
<tr>
<td>Tolerance problem regarding AEs</td>
<td>4 (9.8%)</td>
</tr>
<tr>
<td>Other</td>
<td>1 (2.4%)</td>
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</tbody>
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**DISCUSSION**

The introduction of the imatinib has surprisingly improved the survival time for patients with CML. However, upon resistance or intolerance to imatinib, appropriate therapeutic modifications are required. Use of second-line therapy with dasatinib or nilotinib may overcome imatinib resistance or intolerance.8 Dasatinib has allowed a shift in the treatment of patients with CML, across all phase of disease, who develop resistance or intolerance to imatinib in Turkey. A retrospective study has been designed to understand CML patients treated with dasatinib as second-line therapy reflecting the real-world clinical practice. A total of 41 patients from different universities in Turkey satisfied the selection criteria and were included in the analysis. The main reason of the switch from imatinib (80.4%) has been reported as secondary resistance. It is very well-known that the patients with CML require do-
Table 7. Average Dosage Costs Of Patients Per Months

<table>
<thead>
<tr>
<th>Month</th>
<th>Average dosage cost (TL)</th>
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<tbody>
<tr>
<td>1</td>
<td>153</td>
</tr>
<tr>
<td>2</td>
<td>152</td>
</tr>
<tr>
<td>3</td>
<td>152</td>
</tr>
<tr>
<td>4</td>
<td>152</td>
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</tr>
<tr>
<td>11</td>
<td>147</td>
</tr>
<tr>
<td>12</td>
<td>148</td>
</tr>
</tbody>
</table>

pleural effusion, it is needed to restart with lower doses of dasatinib in certain cases and then to increase to targeted dose of dasatinib. This article provided the rationale that in these cases, the patients have been treated appropriately even with lower doses of dasatinib for relatively longer periods without any doubt for progression risk.

In recently published reviews by Buyukan et al., it was summarized the last decade history of CML regarding the amazing developments of therapeutic side of the disease including TKI’s and understanding the molecular background for this development. In this review articles, it was briefly mentioned about the potential dose variations of dasatinib in different type of patients and in different clinical situations with acceptable efficacy and safety.

There have been many reports showing the potential effectiveness of different dosage regimens for dasatinib which could potentiate the individualized therapy based on the patient’s characteristics and comorbidities even facilitating the use of desired drug without changing the drug for the physicians.

This study underlined the real-world clinical and economic outcomes associated with the use of dasatinib. Dasatinib has a wide tablet formulations like 20, 50, 70 mg in Turkey and thus can provides an adjustable dose application. Even at low doses of dasatinib are associated with higher response rates in Turkish patients with CML. While further studies are warranted, this research provides clinical and economic evidences to help clinicians and healthcare administrators in selecting second-line treatment for CML patients resistant or intolerant to imatinib.
REFERENCES
19. Gleevec® (imatinib) SmPC. Novartis.
20. Tasigna® (nilotinib) SmPC. Novartis.
21. SPRYCEL® (dasatinib) SmPC. Bristol-Myers Squibb.


Correspondence
Dr. Salih AKSU
Hacettepe Üniversitesi Tıp Fakültesi
İç Hastalıklar ABD, Hematoloji Ünitesi
Sihhiye, Ankara / TURKEY

Tel: (+90.312) 305 15 43
Fax: (+90.312) 305 16 14
e-mail: saksu@hacettepe.edu.tr