# Imatinib in the treatment of chronic myeloid leukemia in children and adolescents is effective and well tolerated: Report of the Polish Pediatric Study Group for the Treatment of Leukemias and Lymphomas

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A – research concept and design; B – collection and/or assembly of data; C – data analysis and interpretation; D – writing the article; E – critical revision of the article; F – final approval of the article

## Abstract

**Background.** Chronic myeloid leukemia (CML) constitutes only 2–3% of all leukemias in pediatric patients. Philadelpha chromosome and BCR-ABL fusion are genetic hallmarks of CML, and their presence is crucial for targeted molecular therapy with tyrosine kinase inhibitors (TKIs), which replaced hematopoietic stem cell transplantation (HSCT) as a standard first-line therapy. The disease in pediatric population is rare, and despite molecular and clinical similarities to CML in adults, different approach is needed, due to the long lifetime expectancy and distinct developmental characteristics of affected children.

**Objectives.** The objective of this study is to evaluate treatment with imatinib in Polish pediatric patients with CML.

**Material and methods.** We analyzed the results of treatment with imatinib in 57 pediatric patients (June 2006 – January 2016) from 14 Polish pediatric hematology and oncology centers.

**Results.** In the study group, 40 patients continued imatinib (median follow-up: 23.4 months), while in 17 the treatment was terminated (median follow-up: 15.1 months) due to therapy failure. In the latter group, 13 patients underwent HSCT, while 4 switched to second-generation TKIs. The 5-year overall survival rate (OS) in the study group was 96%, and the 5-year event-free survival (EFS) was 81%.

**Conclusions.** Our results confirm that the introduction of TKI therapy has revolutionized the treatment of CML in the pediatric population by replacing the previous method of treatment with HSCT and allowing a high percentage of OS and EFS.

**Key words:** children, imatinib, adolescents, chronic myeloid leukemia

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**Funding sources**

None declared

**Conflict of interest**

None declared

**Acknowledgements**

The authors thank Jennifer Grbevski and Nicole Grbevski for their careful English language editing.

**Received** on May 28, 2016

**Reviewed** on June 7, 2016

**Accepted** on October 27, 2016

**DOI**

10.17219/acem/66462

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Introduction

Chronic myeloid leukemia (CML) is a myeloproliferative neoplasm typically diagnosed in the adult population and is relatively rare in children with the incidence of 0.6–1.2 million per year. CML is a clonal disorder of hematopoietic progenitor cells resulting from the balanced translocation (9;22) called the Philadelphia chromosome (Ph) at the molecular level, resulting in the formation of a fusion gene BCR-ABL. The BCR-ABL protein encoded by the fusion gene exhibits tyrosine kinase activity and promotes uncontrolled proliferation of pluripotent stem cells in bone marrow (BM). CML has a 3-phase course: chronic phase (CML-CP); accelerated phase (CML-AP); and blast crisis phase (CML-BC). CML is most commonly diagnosed in the CML-CP and only in about 10% of cases in advanced phases: CML-AP or CML-BC.²

Before the implementation of imatinib, hydroxyurea +/- interferon alpha remained the first-line treatment of CML, followed by HSCT after achieving hematologic remission.³ The identification of tyrosine kinase inhibitors (TKI) with BCR-ABL blocking ability revolutionized the CML therapy due to pharmacological control of leukemic clone. TKI-based therapy has proved to be very effective and quickly led to the withdrawal of HSCT as the first-line treatment. A first-generation TKI – imatinib was approved for the treatment of adult patients in 2001. Then, in 2003, it was approved, for the therapy in children. It should be emphasized that treatment with TKI is not a way to cure CML, as in most patients leukemia cells are still present. However, a unique feature of these drugs is a significant reduction of the risk of CML progression.² In the era of TKIs, the CML-CP can last beyond 20 years.

The recommended starting dose of imatinib for children is 260–300 mg/m² (max daily dose: 400 mg) in CP, 400 mg/m² (max daily dose: 600 mg) in AP, and 500 mg/m² (max daily dose: 800 mg) in the blastic phase.³ In recent years, due to the development of the targeted therapy and the implementation of TKIs in pediatric patients, there has been significant progress in the treatment of CML, but the data is relatively limited due to low incidence of CML. One should remember that long-term side effects of TKI therapy may occur, because the drug has only been in use for approx. 15 years. Despite the excellent results of imatinib therapy, one should not forget about HSCT. It is the only method for obtaining definite cure of CML. HSCT is the first-line treatment in patients with CML who have become resistant to the first- and second-generation TKIs, or when serious side effects of the therapy occur and there is a matched donor available. In some cases, the preference of HSCT by the patient or his/her parents is also very important. It is worth noting that HSCT should be considered more frequently in patients diagnosed with CML before puberty due to growth impairment after TKI therapy. The transplant is in fact the only method that can lead to the cure of this disease.

Material and methods

We conducted a retrospective, nation-wide analysis of the imatinib therapy results in children and adolescents with CML in Poland. The study group consisted of 57 patients (M = 35, F = 22) from 14 Polish pediatric hematology and oncology centers treated in 2006–2016. The majority of patients (n = 54) were diagnosed in the chronic phase and only 3 patients in the accelerated phase. The diagnosis of CML was made according to the 4th edition of WHO classification of hematopoietic and lymphoid tissues, with mandatory molecular confirmation of BCR-ABL fusion. The first-line treatment with imatinib was performed accordingly to previous I-BFM recommendations. Patients and/or their legal guardians signed the appropriate consents.

The effectiveness of TKI therapy was evaluated on the basis of “milestones” of the therapy: complete hematologic remission (CHR); complete cytogenetic response (CCyR); and major molecular response (MMR). CHR was defined as leukocyte count <10 x 10³/µL, <5% basophils and <450 x 10³/µL platelets in peripheral blood, absence of myelocytes, promyelocytes and blasts in the peripheral blood, and non-palpable spleen during physical examination. Starting from the moment of CML diagnosis blood counts with blood smear should be performed every 15 days until achieving CHR, and then at least once every 3 months. Complete cytogenetic response (CCyR) was defined as the absence of Ph (+) cells in the bone marrow in classical cytogenetics or the FISH method, and evaluated within 12 months of the treatment.

Partial cytogenetic response (PCyR) was the presence of 1–35% Ph (+) cells in the bone marrow. Minor and minimal cytogenetic response was the presence of 36–65% or 66–95% Ph (+) cells in the bone marrow, respectively. Cytogenetic evaluation was performed after 3 and 6 months after the implementation of imatinib, then every 6 months until CCyR achievement, and every 12 months thereafter or in the case of treatment failure.

Molecular response was evaluated after 18 months of the treatment. Major molecular response (MMR) was defined as the BCR-ABL transcript level below 0.1% Molecular tests were performed every 3 months until MMR confirmation, and then not less than every 6 months. If there was an unsatisfactory response, testing for BCR-ABL kinase domain mutations was performed.

Statistical analysis

The statistical analysis was performed using STATISTICA v. 10.0. As events in the analysis of event-free survival (EFS) we considered the following: death of the patient; switch to second-generation tyrosine kinase inhibitors (2G TKIs) due to therapy failure or imatinib intolerance; proceeding with HSCT due to intolerance of imatinib and/or loss of cytogenetic or molecular response during imatinib therapy.
Results

Study group

The median age at CML diagnosis was 13.6 years (the youngest child was 1.2 year, the oldest 17.9 years). The most commonly reported symptoms at the diagnosis were: asthenia (n = 22); weight loss (n = 18); abdominal pain (n = 17); fever (n = 16); limb pain (n = 10); and hemorrhage (n = 7). Other reported symptoms included ecchymoses (n = 4); headache (n = 4); diplopia (n = 2); pallor (n = 2); cough (n = 2); priapism (n = 2); breast pain (n = 1); night sweats (n = 1); arthritis (n = 1); dyspnoea (n = 1); polydipsia (n = 1); hair loss (n = 1); and intramuscular hematoma (n = 1). The most frequent signs were splenomegaly (n = 43; median size of the spleen was 5.5 cm below the costal margin), and hepatomegaly (n = 34; median size of the liver was 3 cm below the costal margin). In 8 cases, the diagnosis was set on the basis of routine blood tests without other accompanying symptoms. Laboratory tests revealed significant hyperleukocytosis in the majority of patients (median 226.28 10^3/µL, range 7.17–810 10^3/µL) and thrombocytosis (median 471 10^3/µL, range 27.9–3444.7 10^3/µL). During the follow-up period, 2 patients died, both after transplantation. One of them died on day +307 after HSCT from central nervous system aspergillosis and multiorgan failure, while the other died on day +76 after HSCT from grade IV acute Graft vs Host Disease (aGvHD) and pulmonary hemorrhage. The 5-year OS in the study group was 96% (Fig. 1) and the 5-year EFS was 79% (Fig. 2).

Treatment with imatinib

All pediatric patients were qualified for first-line treatment with imatinib, according to CML-PAED II and 1-BFM-CML protocol. Hydroxyurea was administered as pre-imatinib cytoreductive phase in 21 patients (median duration: 16 days). In 1 case, anagrelide was administered alternatively for the treatment of CML-associated thrombocytosis. The median time of imatinib implementation from the moment of diagnosis was 7 days (range 0–202 days). The median initial dose was 300 mg/m² (range 220–468 mg/m²). The maximum dose was implemented in a single patient in CP, who had become resistant to the standard doses and experienced lack of CCyR and MMR in 12 and 18 months after the initiation of therapy, respectively. The dose of imatinib during therapy was modified in 23 cases. In 13 patients, the dose
had to be increased from 300 mg/m² to 400 mg/m² due to resistance to imatinib therapy. In 10 patients, the imatinib dose was reduced due to toxicity (n = 8) or satisfactory response to a higher dose (n = 2) (median dose after reduction: 170 mg/m²). Toxicity included myelotoxicity (WHO grade 4, n = 1); thrombocytopenia (n = 1); leukopenia (n = 5); and headache (n = 1). In 4 patients, 2G TKIs were implemented after imatinib: dasatinib in 3 cases, and nilotinib in 1 case. In 19 patients, due to unsatisfactory response to imatinib, testing for BCR-ABL kinase domain mutations was performed. In 3 cases, the T315I mutation was confirmed, and all these patients underwent HSCT afterwards. In 1 case, ponatinib achieved molecular remission.

**Complete hematologic remission**

Within 3 months complete hematologic remission (CHR) was achieved in 57 patients. In 2 patients, we observed hematologic relapse. One patient underwent HSCT afterwards, and the other was switched to 2G TKI-nilotinib and achieved another CHR.

**Complete cytogenetic response**

Within 12 months, complete cytogenetic response (CCyR) was evaluated in 45 out of 57 patients. In 4 cases, the follow-up period did not exceed 1 year, 4 patients underwent HSCT in less than 12 months since the start of the treatment, 2 patients were transferred to an adult ward before the evaluation, while in 2 patients cytogenetic response was not assessed. Within 12 months from the onset of therapy, CCyR was achieved in 31/45 patients (i.e., 69%). After 12 months, CCyR was observed in 9/45 cases (i.e., 20%). CCyR was not achieved in 5 cases (i.e., 11%). Out of the 5 patients who failed to achieve CCyR, 3 patients were switched to dasatinib, and 2 patients underwent HSCT afterwards.

**Major molecular response**

After 18 months, major molecular response (MMR) was evaluated in 46 out of 57 patients. In 5 cases, the follow-up period did not reach 1.5 year. Four patients underwent HSCT in less than 18 months since the start of the treatment,
and 2 patients were transferred to an adult ward. MMR was achieved in 28/46 patients (i.e., 61%) after 18 months since the beginning of the treatment. MMR later than 18 months from therapy start was seen in 8/46 patients (i.e., 17%) with median time of 24 months. MMR was not achieved in 10/46 cases (i.e., 22%). Five of these patients underwent HSCT afterwards, and 2 were switched to 2G TKI-dasatinib. In 3 cases, the dose of imatinib was increased, with continuous decrease in the BCR-ABL level. Characteristics of the study group and the results of treatment are summarized in Table 1.

Table 1. Characteristics of the study group. Results of treatment with imatinib in 57 pediatric patients (June 2006 – January 2016) from 14 Polish pediatric hematology and oncology centres

<table>
<thead>
<tr>
<th>Study group</th>
<th>Number of patients</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patients total</td>
<td>57</td>
</tr>
<tr>
<td>male</td>
<td>35</td>
</tr>
<tr>
<td>female</td>
<td>22</td>
</tr>
<tr>
<td>Phase at the diagnosis</td>
<td></td>
</tr>
<tr>
<td>chronic phase</td>
<td>54</td>
</tr>
<tr>
<td>accelerated phase</td>
<td>3</td>
</tr>
<tr>
<td>Hematologic remission</td>
<td></td>
</tr>
<tr>
<td>complete hematologic remission (CHR)</td>
<td>55</td>
</tr>
<tr>
<td>hematologic relapse</td>
<td>2</td>
</tr>
<tr>
<td>Cyto genetic response evaluable</td>
<td></td>
</tr>
<tr>
<td>CCyR within 12 months</td>
<td>45/57</td>
</tr>
<tr>
<td>CCyR after 12 months</td>
<td>31 (69%)</td>
</tr>
<tr>
<td>failed to achieve CCyR</td>
<td>9 (20%)</td>
</tr>
<tr>
<td>switch to dasatinib</td>
<td>5 (11%)</td>
</tr>
<tr>
<td>HSCT</td>
<td>2</td>
</tr>
<tr>
<td>Molecular response evaluable</td>
<td></td>
</tr>
<tr>
<td>MMR within 18 months</td>
<td>46/57</td>
</tr>
<tr>
<td>MMR after 18 months</td>
<td>28 (61%)</td>
</tr>
<tr>
<td>failed to achieve</td>
<td>8 (17%)</td>
</tr>
<tr>
<td>switch to dasatinib</td>
<td>10 (22%)</td>
</tr>
<tr>
<td>HSCT</td>
<td>5</td>
</tr>
<tr>
<td>switch to dasatinib</td>
<td>3</td>
</tr>
<tr>
<td>increase of the imatinib dose</td>
<td>5</td>
</tr>
</tbody>
</table>

**Hematopoietic stem cell transplantation**

Hematopoietic stem cell transplantation (HSCT) was performed in 13 children (M = 9, F = 4). In 4 patients, the transplant was performed in the 1st CML-CP due to the local preferences of the center. In 3 patients, the reason for HSCT was the advanced phase of CML (CML-BC, n = 1; CML-AP, n = 2), and these children were transplanted in the 2nd CP (CML-CPII), after TKI treatment. Hematologic toxicity of imatinib was the reason for a transplant in 1 case. In 5 other cases, the loss of molecular and/or cytogenetic response was observed. The donors were either matched unrelated (MUD: n = 9) or sibling (MSD: n = 4). Median time from the diagnosis of CML to HSCT was 14.8 months (range 5.7–49 months). In 9 children, HSCT was performed in the 1st CP, and in 3 cases in the 2nd CP. In 1 case, a reduced-intensity conditioning regimen according to CML-SCT I-BFM study was introduced. So far, tests have confirmed 100% allogenic chimerism and non-detectable BCR-ABL (MR 4.5) in the patients’ peripheral blood, but the follow-up period is too short (2 months) to confirm the complete success of transplantation.

**Follow-up**

Out of 57 patients enrolled in the treatment, 40 patients continue therapy with imatinib, while 17 completed the treatment. Thirty-five patients continue the therapy in pediatric centers (median follow-up: 33 months), and 3 patients in adult centers, with which we are in constant contact (median follow-up: 93 months), while 2 patients were lost to follow-up treatment after being transferred to adult centers (status at the last contact: continuation of the TKI therapy). Among patients who completed the treatment, 13 underwent HSCT (2 patients died due to complications of the post-transplant period), and 4 patients were switched to second-generation TKIs (dasatinib, n = 3; nilotinib, n = 1).

**Discussion**

Clinical data concerning the treatment of CML in pediatric population are limited because of its low incidence in children. It should be noted that for the purpose of our analysis we have gathered a relatively large group of patients (n = 57) compared with the available literature. The entire observation period amounted for 33 months in patients who continue the treatment in pediatric centers, and 93 months in patients who continue the treatment in adult centers.

**Hematopoietic stem cell transplantation**

Before the era of imatinib, HSCT was a standard first-line therapy. Among pediatric patients who underwent transplantation from MSD in the 1st chronic phase (CML-CP) in the years 1982–2004, EFS during the observation period of 3–5 years after HSCT ranged from 61 to 63%. OS ranged from 66 to 87%. In transplant patients, MUD results were inferior, with EFS ranging from 27 to 55%, and OS from 45 to 65%. The main cause of death in both groups was acute and chronic GvHD, more common in children after MUD-HSCT. In patients transplanted in the advanced phase (CML-AP or CML-BC) or the 2nd chronic phase (CML-CPII), the results were worse with EFS from 34 to 35%, and OS from 39 to 46%. In our study group, only 13 patients underwent HSCT, which was mainly due to the good response to TKI therapy in most cases. OS among these children was 86%. We should remember that despite giving up the use of HSCT as a first-line therapy, it is still the only method by which we can completely eliminate leukemia cells and cure the disease.
Treatment with imatinib

One of the first trials with TKI in children was published in 2004 by the Children's Oncology Group and it presented promising results. For the 2nd phase trial of this study, 31 pediatric patients who experienced failure of interferon therapy were qualified. In this group, all the children achieved CHR, and in 83% of them CCyR was observed. Another study covered 8 European countries and 30 patients were enrolled in it. CHR was achieved in 80% cases, and CCyR in 60% of the patients enrolled in the trial in the CP and 29% of the children enrolled in the AP. In half of the children, MMR was reported. Similar results were published in France, in the 4th phase study conducted on a group of 40 patients. The average duration of the follow-up period in this study was 16 months. CHR within 3 months was achieved in 86% of the patients, and within 6 months in 98% of the patients. CCyR was observed in 62% of the children within 1 year after the inclusion of imatinib therapy, in 34% of them MMR within 18 months after initiation of the treatment was confirmed. Similar results were achieved in a German clinical trial CML-PAED II published in year 2009. Out of 42 enrolled patients, 40 achieved CHR within 3 months. In 26 out of 28 patients, CCyR was observed within 1 year after the start of the treatment, and in 17 out of 19, MMR after 18 months from the onset of the treatment. It should be emphasized that all patients from our study group are simultaneously registered in an international database I-CML-Ped Study, so far consisting of 351 children diagnosed with CML. This study, aimed at optimizing the treatment of CML in children, is still ongoing. Preliminary results presented at the 56th American Society of Hematology (ASH) Annual Meeting are promising and comparable with the results of treatment achieved in the Polish population. Our study group consisted of 57 pediatric patients diagnosed in Poland with a median overall follow-up period of 31 months. CHR within 3 months after the implementation of the treatment with imatinib was documented in all patients. CCyR after 12 months of therapy was observed in 69% of patients, while MMR in 61% of patients after 18 months of treatment. These results correspond with the quoted literature, in particular with the results of Suttorp et al.'s analysis from 2009, and the results of treatment in the context of clinical trial I-CML-Ped Study. Moreover, when comparing the proportion of patients achieving CCyR and MMR, our results are above promising. However, despite the excellent results of imatinib therapy, one should not forget about HSCT. It is worth noting that HSCT should be considered more frequently in patients diagnosed with CML before puberty due to the fact that long-term TKI intake may in the future be the cause of short stature among these patients.

Side effects of imatinib

TKI are usually well tolerated; however, some side effects may occur. In most cases, they are classified as mild to moderate and occur mainly in patients in whom TKI therapy was introduced in advanced phases of CML. Non-hematological, relatively common side effects include nausea; vomiting; diarrhea; skin rash; swelling; limb pain; muscle spasms; bone and joint pain; headaches; weight gain; and an increase in liver enzymes. In one of our patients, we documented severe headaches, which resolved after a temporary decrease in the dose of imatinib. Milot et al. described neutropenia grade 3 or 4 in 27% of the children receiving imatinib; thrombocytopenia grade 3 or 4 in 5%; and anemia grade 3 or 4 in 2.5% of the patients. However, these cytopenias were treatable by temporary discontinuation of the therapy or the administration of granulocyte-colony stimulating factor (G-CSF) in some children with neutropenia. In 5 of our patients, we observed leukopenia, and in 1 thrombocytopenia, which also resolved after a temporary dose reduction of imatinib. Only in 1 case, myelosuppression as a side effect after imatinib implementation was a reason for HSCT. Despite the potential cardiotoxicity, hepatotoxicity, immune disorders and thyroid gland dysfunction observed in adults treated with imatinib, they have not been documented so far in children. In the group of pediatric patients with CML, the aspect of TKI therapy impact on bone metabolism is very important. Imatinib impairs the differentiation and reduces the activity of osteoblasts and osteoclasts. This can result in growth retardation in children, particularly those who are starting the treatment in prepubertal age. In our study group, we observed no impact of imatinib on calcium and phosphate metabolism. We also noticed no abnormalities in serum phosphate, calcium, parathyroid hormone (PTH) and vitamin D levels or tubular function disorders (phosphate absorption). However, we should keep in mind that particular attention should be paid in the group of the youngest patients chronically receiving TKI. They require regular and detailed clinical evaluation, and performing the panel of basic laboratory tests during each visit. Our results confirmed that the recommended daily dose of imatinib is well tolerated in pediatric patients, and severe side effects are relatively rare.

Second-generation TKIs

2G TKIs – dasatinib and nilotinib – were registered for the treatment of adult patients with CML in 2006, and dasatinib only for the therapy of children in 2007. Nilotinib is not currently recommended for the treatment of pediatric patients. 2G TKIs are recommended when intolerance or resistance to imatinib occur. They are more effective in the treatment of CML due to the linking of both active and inactive conformations of the BCR-ABL protein. In addition, they show greater activity in the case of mutations of BCR-ABL gene, associated with resistance to TKI therapy. Unfortunately, there are an even more limited number of studies on the use of 2G TKIs in children. The first reports are from 2011 as the results of the 1st
phase clinical trial conducted by the Children’s Oncology Group. Thirty-nine patients were enrolled for the treatment with dasatinib, including 9 with CML who were resistant to imatinib or who had had an adverse event after using it. In 8 patients, cytogenetic response was observed, in 3 CCyR, in 3 PCyR, in 1 minor, and in 1 minimal cytogenetic response. In 1 patient, the cytogenetic response was not possible to assess. The 1st phase study CA 180–018 from 2013 was conducted on a group of 63 pediatric patients, 17 of whom were enrolled in the chronic phase of CML, and only 3 in the advanced phase. Among patients with CML-CP, CHR was achieved in 94%, CCyR in 82%, and MMR in 47%. Patients enrolled in the study in the advanced phase of CML achieved slightly worse results. In our study group, dasatinib was administered in 3 patients, while nilotinib in 1 case. In all 4 cases, these drugs were implemented due to the failure of imatinib treatment. Patients continue the treatment with 2G TKIs with a satisfactory outcome, and no side effects have been documented. Clinical trials on the use of 2G TKIs are being carried out and are raising great hopes, especially in patients in whom imatinib has proven to be ineffective.

**Discontinuation of the treatment**

Observations in adult patients show that during the TKI therapy and after documenting the undetectable level of BCR-ABL over a period of 24 months or more, one can try to discontinue the treatment with imatinib. In approx. 40% of these patients, one can confirm continuous MMR despite cessation of the therapy. In our study group, the treatment was discontinued in 1 patient after continuous 26 months of the undetectable level of BCR-ABL. The patient remained under constant control and, after 11 months of continuous MMR, molecular relapse was confirmed (level of BCR-ABL transcript 3%). Imatinib was reintroduced and the patient achieved another MMR despite cessation of the therapy. In approx. 40% of these patients, one can confirm continuous MMR despite cessation of the therapy.

Our results confirm that the introduction of TKI therapy has revolutionized the treatment of CML in the pediatric population by replacing the previous method of treatment with HSCT and allowing a high percentage of OS (96%) and EFS (81%). Although the use of TKIs and 2G TKIs is not a cure for CML, and only reduces the risk of disease progression significantly, the results of ongoing clinical trial evaluating the safety of the treatment withdrawal after confirming a continuous 24-month undetectable level of BCR-ABL are promising. Despite the initial enthusiasm due to the excellent results of TKI therapy, there are more reports confirming that the use of imatinib is not devoid of serious side effects. However, in our study group, we observed in only 1 case myelotoxicity WHO grade 4, which was the reason for HSCT. We should keep in mind the fact that the goal of the therapy in pediatric patients should be rather to cure the disease than to suppress it, which can be achieved only by performing HSCT. In the context of very promising results of HSCT in pediatric patients with CML after reduced-intensity conditioning regimen, HSCT should be taken into consideration, especially in prepubertal children.

**Conclusions**

Our results confirm that the introduction of TKI therapy has revolutionized the treatment of CML in the pediatric population by replacing the previous method of treatment with HSCT and allowing a high percentage of OS (96%) and EFS (81%). Although the use of TKIs and 2G TKIs is not a cure for CML, and only reduces the risk of disease progression significantly, the results of ongoing clinical trial evaluating the safety of the treatment withdrawal after confirming a continuous 24-month undetectable level of BCR-ABL are promising. Despite the initial enthusiasm due to the excellent results of TKI therapy, there are more reports confirming that the use of imatinib is not devoid of serious side effects. However, in our study group, we observed in only 1 case myelotoxicity WHO grade 4, which was the reason for HSCT. We should keep in mind the fact that the goal of the therapy in pediatric patients should be rather to cure the disease than to suppress it, which can be achieved only by performing HSCT. In the context of very promising results of HSCT in pediatric patients with CML after reduced-intensity conditioning regimen, HSCT should be taken into consideration, especially in prepubertal children.

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