Advances in understanding the pathogenesis of chronic myeloid leukemia (CML) and implementation of the therapy with tyrosine kinase inhibitors (TKI) could be considered as a prototype of successful fight against cancer. However, for an optimally responding patient it is recommended to follow the TKI therapy indefinitely. The question about the possibility of safe TKI treatment discontinuation in certain clinical situations was raised and is currently under close investigation worldwide. Currently, imatinib discontinuation trials have shown that about 60% of eligible patients experienced molecular recurrence within 6 months of treatment discontinuation, while the remaining 40% remained in defined deep molecular response throughout the duration of mostly two years follow-up. Interestingly, retreatment with the same TKI or another TKI was successful in the vast majority of patients demonstrating molecular recurrence of the disease. These findings support the concept of safe TKI treatment discontinuation and its usefulness for a specific subset of CML patients. However, recent data are not sufficient for TKI discontinuation attempts outside clinical trials yet. Because of the high risk of potentially problematic molecular recurrences of the pathological clones, the key question is to find the right predictive marker of TKI discontinuation success, however it stays unsolved yet. This minireview brings a concise summary of this hot topic with a realistic view from clinical routine.

Key words: tyrosine kinase inhibitors, imatinib, discontinuation, chronic myeloid leukemia

Chronic myeloid leukemia (CML) is a clonal hematopoietic stem cell disorder consequent to the origin of the BCR-ABL1 fusion gene, which is predominantly generated by a t(9;22)(q34;q11) reciprocal translocation. The product of the oncogene, bcr-abl1 fusion protein, has a constitutive tyrosine kinase activity and is believed to play a crucial role in the development of CML. Biological therapy with tyrosine kinase inhibitors (TKIs) specifically disables this activity and effectively clears malignant clone in the vast majority of CML patients. Imatinib mesylate, first-generation TKI, has become the standard first-line treatment of chronic phase CML during the last decade. Second-generation TKIs (dasatinib, nilotinib) were shortly developed to overcome the resistance or intolerance to imatinib. Clinical trials with next generation TKIs (bosutinib, ponatinib) are in progress. Several studies of imatinib as first-line therapy have published comparable long-term results. The estimated event-free survival at 6 years was 83%, and the estimated rate of freedom from progression to accelerated phase and blast crisis was 93% in the IRIS study [1]. Practice guidelines issued by the European LeukemiaNet (ELN) and the National Comprehensive Cancer Network (NCCN) both recommend that a patient who is responding optimally to TKI treatment continues indefinitely at the standard recommended dose [2, 3]. However, some situations in clinical routine raised the question about the possibility of safe TKI treatment discontinuation. These situations involve issues like chronic toxicity, patient compliance, pregnancy or patient’s wish. Furthermore, financial issues of TKI treatment are being considered not only in economies with limited financial
sources (approximately 5,000 new cases are diagnosed with CML in Europe each year).

Rare cases with long-term stable minimal residual disease or complete molecular response after cessation of interferon alpha therapy had been reported and they laid the theoretical basis for the TKI discontinuation attempts [4]. Soon after 2000, when imatinib treatment came into clinical routine, case reports of patients who stopped the treatment were reported. Few years later, first clinical studies focusing on imatinib dose modification or discontinuation started to be published [5-7].

**Response to TKI treatment.** At present, the most important prognostic factor for CML patients is the response to TKI treatment. The response is measured based on cytogenetic (G-banding, FISH) and molecular genetic (qRT-PCR) methods. The definition of optimal response has evolved during the last few years. At present, an optimal response to any TKI as first-line treatment is defined by at least partial cytogenetic response (PCyR; 1-35% Ph+ metaphases) at 3 months, at least complete cytogenetic response (CCyR; No Ph+ metaphases) at 6 months and at least major molecular response (MMR; BCR-ABL1 transcripts ≤ 0.1% on the International Scale) at 12 months [2]. Molecular response (MR) is mostly assessed according to the International Scale (IS) as the ratio of BCR-ABL1 transcripts to ABL1 transcripts and it is reported as BCR-ABL1 % on a log scale, where 10%, 1%, 0.1%, 0.01%, 0.0032%, and 0.001% correspond to a decrease of 1, 2, 3, 4, 4.5, and 5 logs, respectively, below the standard baseline that was used in the IRIS study [2]. A BCR-ABL1 expression of 0.1% (MR0.1) corresponds to major molecular response (MMR). The proportion of patients who achieved MMR after 12 months of 400 mg imatinib daily (standard dose) ranged from 18% to 58% in the published studies [2, 3]. A high number of commercially available kits for molecular genetic analyses from different manufacturers for different platforms are available on the market. It is the responsibility of each laboratory to choose some of the kits to fit with the recommended criteria and obtain results, which are comparable with other laboratories.

**Success rate for TKI discontinuation.** The nonrandomized multicentre Stop Imatinib (STIM) study was one of the largest prospective studies of TKI treatment discontinuation organized to date [8]. Imatinib treatment was discontinued in patients who maintained MR31 (defined as undetectable BCR-ABL1 with assay sensitivity of ≥5 log below IS-standardized baseline) for at least 2 years. Of 100 patients enrolled, 61 patients had molecular recurrence after a median of 30 months of follow-up (58 patients within 7 months) and 39 patients maintained MR30 after a median of 22 months of follow-up. The overall probability of maintaining MR30 at 12 months and 24 months was 43% and 39%, respectively [8, 9]. Molecular recurrence was defined as BCR-ABL1 ≥ 0.001% IS in the STIM study.

The results of another prospective study, TWISTER, were published in 2013 [10]. Imatinib treatment was stopped in 40 patients who maintained MR32 for at least 2 years. With a median of 42 months of follow-up post-treatment cessation, 18 patients (45%) had not demonstrated molecular relapse (defined as loss of MMR or two consecutive occurrences of detectable BCR-ABL1), and the Kaplan–Meier estimate of rate of treatment-free remission at 2 years was 47.1% [9, 10]. Molecular relapse was defined as loss of MMR or two consecutive occurrences of detectable BCR-ABL1 in TWISTER study.

In the HOVON 51 study, patients received escalating doses of imatinib in combination with escalating doses of cytarabine according to the study protocol. Thirty-three patients from the HOVON 51 study with an MR45 for at least 2 years who were still on imatinib treatment were randomized between continuation of imatinib (arm A, n = 18) or discontinuation of imatinib (arm B, n = 15) [11]. After a median follow up of 36 months since randomization, 3 patients (17%) in arm A and 10 patients (67%) in arm B had a molecular relapse. All 3 relapsing patients in arm A had also stopped imatinib after randomization. All but one relapsing patient relapsed within 7 months after discontinuation of imatinib. The molecular relapse rate at 12 and 24 months after randomization was 0% and 6% (arm A) and 53% and 67% (arm B) respectively [11].

Takahashi et al. [12] conducted a nationwide retrospective survey of the clinical outcomes of CML patients after discontinuation of imatinib in Japan. Among 3,242 imatinib-treated CML patients, they identified 50 (1.5%) who had discontinued imatinib for at least six months; of these they were able to analyze 43 in full. A complete molecular response (CMR) was defined as detection of no BCR-ABL1 transcript in a real-time quantitative-polymerase chain reaction (RT-PCR) assay (n=24), nested reverse transcriptase-polymerase chain reaction (RT-PCR) assay (n=14), or a highly sensitive transcription-mediated amplification (TMA) method (n=5). These PCR methods could detect at least a 4-log reduction in the BCR-ABL1 transcript (international scale <0.01%). The reasons for which imatinib was discontinued were adverse events (n=18), patient's request due to cost (n=14), patient's desire to become pregnant (n=3), and long undetectable residual disease (n=8). The median duration of CMR before cessation was 27.4 months (range 0.9-79.6 months). The median period of cessation was 22.4 months (range 6.2-97.9 months). Molecular recurrence was detected in 19 patients (44%). The relapse free survival (RFS) rate at five years was estimated to be 47% while median RFS was determined to be 41 months using the Kaplan–Meier method [12].

The higher rates of deep molecular responses achievable with nilotinib and dasatinib suggest that use of second-generation TKIs might broaden the pool of candidates eligible for treatment discontinuation. To date, only very limited clinical data on discontinuation of nilotinib or dasatinib therapy are available.

**Summary of qualification criteria for TKI discontinuation.** Above mentioned prospective trails had differed in the qualification criteria for TKI discontinuation and there has not been published any consensus yet. Length of TKI treatment duration, level of response, and duration of response were the...
main criteria used [8, 10, 11]. At least 3 years of TKI treatment duration was demanded in the STIM and TWISTER study. At least MR<sup>45</sup> for at least 2 years were the minimal required level of response and its duration [8, 10, 11]. The ongoing EURO-SKI study (NCT01596114) has these main inclusion criteria: CML in chronic phase, duration of TKI treatment before enrolment at least 3 years, at least MR<sup>48</sup> for at least one year (personal communication).

**Predictive factors for TKI discontinuation success.** Statistically significant predictive factors differed between published studies as well. A trend for a shorter time to BCR-ABL1 negativity in nonrelapsing patients versus relapsing patients was found in the pilot study conducted by Rousselot et al. [6]. The absence of relapse was also not significantly associated with the length of interferon alpha (IFN-α) exposure prior to imatinib in this study [6]. Highest Sokal score group and female sex were predictive of worsened prognosis, whereas a long duration of imatinib was predictive of improved prognosis in the STIM study conducted by Mahon et al. [8]. Significant interaction with relapse risk was observed only for duration of IFN-α treatment and time to achieve undetectable minimal residual disease after switching from IFN-α to imatinib treatment in the TWISTER study performed by Ross et al. [10]. Three (75%) of 4 Sokal high-risk patients relapsed vs 45% of low-risk patients (log-rank P > .05) in this study [10]. The retrospective study of Takahashi et al. [12] found that based on multivariate regression analysis, imatinib dose intensity and prior IFN-α administration were independently predictive of molecular recurrence within 12 months. The depth of the molecular response should be a factor influencing long-term duration of molecular recurrence within 12 months. The risk of relapse on TKI cessation is dependent on the functional immune response and the intrinsic immunogenicity of the CML cells [14, 15].

**Retreatment after relapse.** Importantly, in patients who demonstrated molecular recurrence after TKI discontinuation, retreatment with the same TKI or another TKI resulted in high rates of response, mostly as deep as that observed during initial TKI treatment [6–12]. Loss of MMR was recently recommended as a practical and safe criterion for restarting therapy by Rousselot et al. [13]. It is too early to draw any conclusions about survival of continuing versus discontinuing patients.

**Limitations of current data.** Clinical data on TKI discontinuation obtained from published studies are difficult to compare. The cohorts varied among the clinical studies in many criteria such as how long they had been on TKI therapy, what level of response they had achieved, how long that response had been maintained before discontinuation, and how molecular recurrence was defined. Milder criteria for TKI discontinuation might expand the low pool of candidates, but their heterogeneity may lead to higher rates of relapse and hamper the ability to define the right candidate characteristics. Another limitation of the published studies is the duration of follow-ups, which could not be sufficient to draw relevant conclusions [14]. Moreover, each of the published studies suggested different predictive factors of TKI discontinuation success. Current data about TKI discontinuation don’t allow us to predict any long-term consequences of TKI discontinuation and retreatment on the clinical course of CML as well.

**Models of operational cure of CML.** Several models for the achievement of a prolonged drug-free remission in CML are discussed in the literature [9, 14, 15]. The models are not mutually exclusive, and different pathways to drug-free remission might apply in different patients. In the Model of stem cell depletion, there is a rapid initial drop in the level of BCR-ABL1, which is thought to reflect the clearance of mature CML progeny that are sensitive to TKI. There follows a second phase with a shallow gradient, which is thought to reflect the gradual depletion of the less-sensitive CML stem cells. The slow but progressive depletion of CML progenitors during TKI treatment may be explained by the apoptosis that is dependent on cell cycling [14, 15]. In the Model of immunological control, a reduction in the level of minimal residual disease by TKI therapy is sufficient to overcome T-cell anergy and enables the emergence of an autologous immunological response that suppresses, but may not eradicate, the whole CML clone. The risk of relapse on TKI cessation is dependent on the functional immune response and the intrinsic immunogenicity of the CML cells [14, 15].

**Future directions.** Many of the above-mentioned limitations should be solved by numerous ongoing clinical trials conducted or planned worldwide. In Europe these are STIM 2 (imatinib), STOP 2GTKI (nilotinib, dasatinib), EURO-SKI (all TKIs), and German CML Study V (nilotinib), in the United States study organized by the University of Michigan Cancer Center (imatinib) and ENESTgoal (nilotinib), in Japan Dasatinib Stop Trail (dasatinib), in Korea study organized by the Seoul St. Mary’s Hospital (imatinib). Other trials are organized by the pharmaceutical companies – Bristol-Myers Squibb (DASFREE, dasatinib), Novartis Pharmaceuticals (ENESTop and ENESTFreedom, both nilotinib). Furthermore, French investigators from around the STop Imatinib (STIM) trial have started to explore the feasibility of second attempt to discontinue imatinib in patients with second sustained CMR who experienced molecular recurrence after first attempt of imatinib discontinuation [16]. Their preliminary data suggest that such a second attempt could be possible and could be beneficial for a small subset of patients [16].

Leukemic stem cells are believed to play a crucial role in leukemic hematopoiesis. Strategies to eradicate both proliferating mature progeny and stem cells will continue to be a focus of much clinical research in the field of CML. Research efforts have focused mainly on combination approaches – TKIs plus other drugs.

Despite the remarkable success of TKIs against bcr-abl1, secondary resistance develops in a significant subset of patients, most often due to point mutations in the BCR-ABL1 tyrosine kinase domain or insensitivity of leukemic stem cells to TKIs [17]. One of the current approaches to overcome this
resistance is launch of new non-TKI drugs with a different mechanism of action. One of them is the natural alkaloid omacetaxine mepesuccinate that inhibits protein synthesis and induces cell death independently from the presence of bcr-abl1 mutations [18].

Conclusions

In general, imatinib discontinuation trails have shown that about 60% of eligible patients experienced molecular recurrence within 6 months of treatment discontinuation, while the remaining 40% remained in defined deep MR throughout the duration of mostly two years follow-up. It seems certain that achievement of an MMR is not enough to decide on TKI treatment discontinuation and deeper MR is needed. Interestingly, retreatment with the same TKI or another TKI was successful in the vast majority of patients demonstrating molecular recurrence of the disease. These findings support the concept of safe TKI treatment discontinuation and its usefulness for a specific subset of CML patients. Regardless of economic issues, major benefits from TKI treatment discontinuation could have patients with adverse events and pregnant women. However, recent data are not sufficient for TKI discontinuation attempts outside clinical trials yet. Furthermore, the high risk of potentially problematic molecular relapse can be only hardly acceptable for many of the optimally responding patients, who don’t experience any significant discomfort connected with the TKI therapy. The key issue is to find the right predictive marker of TKI discontinuation success and it waits for the results of the current research.

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