



# Making Treatment-Free Remission (TFR) Easier in Chronic Myeloid Leukemia: Fact-Checking and Practical Management Tools

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## Abstract

In chronic-phase chronic myeloid leukemia (CML), tyrosine kinase inhibitors (TKIs) are the standard of care, and treatment-free remission (TFR) following the achievement of a stable deep molecular response (DMR) has become, alongside survival, a primary goal for virtually all patients. The GIMEMA CML working party recently suggested that the possibility of achieving TFR cannot be denied to any patient, and proposed specific treatment policies according to the patient's age and risk. However, other international recommendations (including 2020 ELN recommendations) are more focused on survival and provide less detailed suggestions on how to choose first and subsequent lines of treatment. Consequently, some grey areas remain. After literature review, a panel of Italian experts discussed the following controversial issues: (1) *early prediction of DMR and TFR*: female sex, non-high disease risk score, e14a2 transcript and early MR achievement have been associated with stable DMR, but the lack of these criteria is not sufficient to exclude any patient from TFR; (2) *criteria for first and subsequent line therapy choice*: a number of patient and drug characteristics have been proposed to make a personalized decision; (3) *monitoring of residual disease after discontinuation*: after the first 6 months, the frequency of molecular tests can be reduced based on MR4.5 persistence and short turnaround time; (4) *prognosis of TFR*: therapy and DMR duration are important to predict TFR; although immunological control of CML plays a role, no immunological predictive phenotype is currently available. This guidance is intended as a practical tool to support physicians in decision making.

## Key Points

In chronic-phase chronic myeloid leukemia, treatment-free remission (TFR) after tyrosine kinase inhibitor therapy has become, with survival, a primary goal for virtually all patients.

The possibility of achieving TFR can no longer be denied to any patient, but some issues regarding the optimal selection of candidates remain unsolved.

We provide practical tools intended to facilitate the decision-making process and maximize the rate of TFR.

## 1 Introduction

Tyrosine kinase inhibitors (TKIs) have drastically improved the outcome of patients with chronic-phase (CP) chronic myeloid leukemia (CML). Normal survival and good quality of life without life-long treatment are now the goals of therapy [1]. Treatment-free remission (TFR) after TKI therapy has become a feasible and safe option in selected CP-CML patients treated with imatinib mesylate (IM) or second-generation (2G) TKIs, such as nilotinib (the only TKI that can be discontinued by label [2]) and dasatinib. Beginning with the STIM1 (Stop Imatinib) study, a number of randomized controlled trials (RCTs) and real-world studies have demonstrated that 30–70% of patients achieving a sustained, stable deep molecular response (DMR) can successfully discontinue therapy and obtain TFR [2–28]. Albeit the inclusion criteria were heterogeneous, most studies relied on a minimum duration of TKI therapy of 3 years and a sustained DMR of  $\geq 1$  year [1, 29]. Long-term data indicate a TFR rate of

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45% at 60 months [5, 30] and 39% at 77 months [2, 3] for patients treated with IM as first-line therapy or second-line after interferon (IFN), and of up to 56% at 60 months for patients receiving nilotinib or dasatinib as first- or second- line therapy [31].

All 2GTKIs have been tested as first-line treatment against IM in company-sponsored RCTs [32–34], but no data on direct comparisons are available yet [35]. No statistically significant survival benefit has been demonstrated for any specific TKI [1, 36, 37]. In the DASISION [33] and ENESTnd [32] trials, superiority of dasatinib and nilotinib was observed only in age-selected populations of young and adult individuals and, therefore, the results cannot be generalized.

The mean age of patients included in trials of TKI discontinuation was similar, between 49 and 63 years. Only in three studies age was shown to have a prognostic role in the maintenance of response, with an advantage for older patients [7, 25, 38]. Age is relevant, because younger patients more frequently express the desire for TFR. However, no age restriction can be set since older patients are just as interested in TFR as younger patients and have a higher probability of achieving a stable DMR [1, 37].

According to the Gruppo Italiano Malattie Ematologiche dell'Adulto (GIMEMA) CML Working Party (WP) recommendations, both survival and TFR are therapeutic goals for patients with CP-CML, and the possibility of achieving TFR can no longer be denied to any patient. TFR should be considered as the primary goal for virtually all patients; treatment policies should be aimed at maximizing the TFR rate while limiting toxicity and costs [37]. However, other international recommendations are more conservative [1, 36] and, consequently, some grey areas remain.

To shed light on some controversial issues and provide practical tools to support physicians in their practice, four Italian experts selected the early prediction of DMR and TFR, the choice of first and later lines of therapy, the optimal frequency of molecular monitoring after the first 6 months of discontinuation, and TFR prognosis as topics that may have a remarkable impact on clinical practice. In collaboration with 17 more hematologists, four working groups were established. They discussed the available data in detail and shared their experience during four online meetings (one per topic) held in April/May 2020. During a plenary meeting in July 2020, the panelists agreed on the final contents to include in the present paper and all of them approved this final version.

## 2 Early Prediction of Deep Molecular Response (DMR) and Treatment-Free Remission (TFR): Is It Possible?

Achievement of stable DMR (defined as at least a 4-log decrease in BCR-ABL1 transcript based on the International Scale [IS] [MR4] [39] for > 2 years [40]) is a prerequisite

to attempting TFR and, therefore, the early identification of DMR predictors is important. Thus far, gender, disease risk, transcript type, and achievement of an early molecular response (EMR) have been attributed a predictive role for attaining DMR in some, but not all, studies; still, their weight cannot be quantified.

### 2.1 Gender

In multivariate analysis, two studies reported a higher probability of achieving a stable DMR in female patients even though the reasons remain unclear [41, 42]. Moreover, in STIM1, female sex was associated with a higher rate of TFR at 12 months (70% vs. 46% in men) [2].

The panel agreed to consider female sex as predictive of achieving stable DMR.

### 2.2 Disease Risk

The Sokal score has long been used to assess the baseline risk of CML in patients treated with conventional chemotherapy,  $\alpha$ -IFN and TKIs. However, according to the most recent recommendations [1, 37], the European Treatment and Outcome Study (EUTOS) long-term survival (ELTS) score should be preferred to assess the risk of progression at baseline and decide the first-line treatment. Despite the fact that the Sokal score and the ELTS score share the same parameters (i.e., hematologic data, spleen size and age), age has a lower negative prognostic impact on the Sokal score. Both scores predict the probability of achieving a major molecular response (MMR), DMR or deeper response [43, 44]. Interestingly, unlike the Sokal score, the ELTS score can predict the achievement of MMR (99%, 87% and 75% in low-, intermediate- and high-risk patients, respectively;  $P = 0.001$ ) and DMR (82%, 61% and 50% in patients with a low, intermediate and high ELTS score, respectively;  $P = 0.005$ ) in CML patients older than 65 years treated with frontline IM or nilotinib therapy [43]. A recent retrospective study investigated the predictive value of the ELTS score with regard to treatment response in patients with CP-CML. Of the Sokal, Hasford, EUTOS and ELTS scores, EUTOS and ELTS predicted MMR at 12 months but ELTS was the only scoring system that predicted DMR at any time (65.0%, 43.7% and 23.5% in low- intermediate- and high-risk groups, respectively;  $P = 0.00006$ ).

Table 1 presents the studies assessing the role of the Sokal and ELTS score at diagnosis as predictors of TFR achievement and maintainance. The largest prospective trial conducted so far (EURO-SKI,  $N = 758$ ) failed to find a predictive role for the Sokal score or the ELTS score in maintaining an MMR 6 months after the discontinuation of TKI therapy [13]. Despite some experiences suggesting

a relevant role of disease risk in predicting TFR [3, 21, 45], the GIMEMA and ELN recommendations, based on the EURO-SKI data [13], stated that the possibility of TFR cannot be denied to intermediate- or high-risk patients.

Low-risk patients have a higher probability of achieving a deeper response than intermediate- and high-risk patients, but it is still unclear whether low-risk patients have a higher probability of maintaining TFR after discontinuation.

The panel agreed that no patient, if eligible, should be excluded from a TFR based on the disease risk.

### 2.3 Transcript Type

The presence of a typical BCR-ABL1 transcript (i.e., e13a2 [b2a2] or e14a2 [b3a2]) is a prerequisite for proper molecular monitoring before and after cessation of TKI therapy [1, 45]. In contrast, in the presence of an atypical transcript (found in nearly 2% of newly diagnosed CML patients [46]), TFR should not be attempted because of the lack of standardized monitoring methods [1, 36, 45]. Thus, although a recent report on seven patients harboring atypical transcripts has shown that treatment discontinuation may be attempted [47, 48], it cannot be recommended yet.

Most studies support the association of the presence of the e14a2 transcript with a faster achievement of a sustained DMR and with a higher rate and duration of TFR after both IM and 2GTKI therapy [41, 49–56] (Table 2). Moreover, in some studies, a better outcome has been reported for patients with the e14a2 transcript if compared to patients with the e13a2 transcript [49, 50], but data were not conclusive.

Interestingly, the EUTOS study on 67 patients with newly diagnosed CML showed that the level of the e14a2 transcript was generally lower than that of the e13a2 transcript (median 33.6%, range 2.9–58.3% vs. 41.6%, range 17.5–101.5) and that the amplification of e14a2 was on average 2.8% less efficient than that of e13a2 ( $P = 0.0026$ ) [57]. This difference in the efficiency of amplification by real-time quantitative polymerase chain reaction (RT-qPCR) may contribute to explaining the differences in the time to achieve an MMR observed among patients with CML harboring the e13a2 and the e14a2 transcripts. Recent studies comparing the performance of RT-qPCR and droplet digital PCR (ddPCR) in the quantification of both BCR-ABL1 transcripts reported higher levels of the e13a2 variant by RT-qPCR and no difference by ddPCR [58, 59], questioning the reliability of BCR-ABL1 measurement by RT-qPCR especially in patients with the e13a2 transcript. This issue is critical for the routine care of CML patients and the results of the ongoing multicenter study are eagerly awaited [57].

The panel agreed that the presence of a typical e14a2 transcript should be considered highly predictive of achieving stable DMR.

### 2.4 Early Molecular Response

EMR is defined as  $\text{BCR-ABL}^{\text{IS}} \leq 10\%$  at 3 months [60]. The rate of EMR at 3 months was between 60% and 80% after treatment with IM, 84% with dasatinib (vs. 64% with IM) [8] and 97% with nilotinib [61]. Notably, patients attaining an EMR at 3 months were shown to reach a complete cytogenetic response (CCyR), an MMR and an MR4.5 by 5 years more frequently than those who did not [8], and had a higher probability of achieving and maintaining the TFR [42, 62–64].

The panel agreed to consider EMR as a predictor of a stable DMR.

Some studies have found that the decrease in BCR-ABL1 transcript levels – defined as BCR-ABL1 halving time (HT)—predicts clinical outcomes and MR regardless of the TKI [65–69]. A recent retrospective study [68] investigated whether the HT was able to discriminate which CML patients were likely to achieve DMR and could be considered for TKI discontinuation. HT was defined as a greater than twofold (if  $\text{BCR-ABL1/ABL1}^{\text{IS}}$  levels were  $\geq 0.01\%$ ) or fivefold (if  $\text{BCR-ABL1/ABL1}^{\text{IS}}$  levels were  $< 0.01\%$ ) decrease in two consecutive analyses. Results showed that the early (at 6 months) achievement of an MMR followed by a further BCR-ABL1 decrease to a DMR ( $\geq \text{MR4.0}$ ) may be considered a selective parameter favoring the rapid identification of individuals potentially eligible for discontinuation of TKI therapy.

In the retrospective study by Shanmuganathan et al. [69], the HT at month 3 was significantly shorter in patients with sustained TFR than in those with molecular relapse, and was a significant predictor of sustained TFR. Based on the HT, patients were divided into quartiles (Q1:  $< 9.35$  days; Q2: 9.35–13.95 days; Q3: 13.96–21.85 days; Q4:  $> 21.85$  days), with the probability of sustained TFR decreasing from 80% in Q1 to 4% in Q4 ( $P < 0.01$ ). Notably, the late relapses had HT in Q3 and Q4, and all patients with HT in Q4 experienced molecular relapse during the follow-up [69].

The panel recognized the relevance of HT as a predictor of DMR but did not endorse its implementation in clinical practice because of the complexity of its calculation.

Table 3 reports the variables discussed above and their impact on TFR planning. In practice, the calculation of

**Table 1** Studies assessing the predictive value of the Sokal and ELTS score for TFR

Ref	N	Outcome measure	Disease risk			P
			Low	Intermediate	High	
TWISTER [6] (Sokal)	40	Rate of TFR 3 years after stopping IM	51.1%	36.5%	25.0%	
STIM1 long term [4] (Sokal)	100	Cumulative incidence of molecular recurrence (SHR [95%CI])—Univariate analysis	Low + intermediate		2.01 (0.98–4.11)	0.05
		Cumulative incidence of molecular recurrence (SHR [95%CI])—Multivariate analysis	Low + intermediate		2.22 (1.11–4.442)	<b>0.024</b>
		Cumulative incidence of molecular recurrence over time	≈ 5.8%	≈ 82%	≈ 82%	<b>0.048</b>
ENESTfreedom 96 weeks [22] (Sokal)		Rate of TFR (95%CI)	61.3% (48.1–73.4)	50.0% (35.5–64.5)	28.6% (13.2–48.7)	
EURO-SKI [14] (Sokal and ELTS)	758	Probability of maintaining MMR 6 months after TKI discontinuation (OR [95%CI])				
		Sokal	Ref.	0.91 (0.60–1.37)	0.90 (0.52–1.55)	0.88
		ELTS	Ref.	0.86 (0.54–1.37)	0.79 (0.36–1.77)	0.72
Castagnetti et al. (2018) [43] (Sokal and ELTS)		All patients				
		Rate of MMR				
		Sokal	89%	86%	70%	< <b>0.001</b>
		ELTS	89%	83%	65%	< <b>0.001</b>
		Rate of DMR				
		Sokal	68%	63%	49%	< <b>0.001</b>
		ELTS	73%	53%	42%	< <b>0.001</b>
		Elderly (> 65 years)				
		Rate of MMR				
		Sokal	94%	88%	83%	0.471
		ELTS	100%	87%	75%	<b>0.001</b>
		Rate of DMR				
Sokal	67%	60%	74%	0.245		
ELTS	82%	61%	50%	<b>0.005</b>		
Italian observational study [26] (Sokal)	293	Predictive value for molecular relapse HR (95% CI)—Univariate analysis	Ref.	0.74 (0.47–1.17)		0.19
		Predictive value for molecular relapse HR (95% CI)—Multivariate analysis	Ref.	0.92 (0.54–1.97)		0.76
					2.07 (1.16–3.71)	<b>0.01</b>
Sato et al. 2020 [44] (Sokal and ELTS)		Rate of MMR by 12 months				
		Sokal	53.3%	56.2%	45.5%	0.409
		ELTS	56.3%	52.1%	11.8%	<b>0.00137</b>
		Rate of DMR at any time				
Sokal	63.3%	56.2%	50.9%	0.223		
ELTS	65.0%	43.7%	23.5%	<b>0.000057</b>		

Bold indicates statistically significant *P*-values (< 0.05)

*N* number of patients, *TFR* treatment-free remission, *IM* imatinib mesylate, *SHR* sub distribution hazard ratio, *95%CI* 95% confidence interval, *MMR* major molecular response, *ELTS* European Treatment and Outcome Study long-term survival, *OR* odds ratio, *pts* patients, *HR* hazard ratio

**Table 2** Selection of studies reporting the association between the e14a2 transcript and molecular response and/or TFR as clinical outcome

Ref	N	Outcome measure	Results	P
Bonifacio et al. (2015) [48]	320	Rate of stable MR4 (among pts with optimal response to IM)		
		At 3 months	e13a2: 23.2% e14a2: 47.8% both: 21%	<b>0.003<sup>a</sup></b>
		At 6 months	e13a2: 28.8% e14a2: 48.6% both: 33.3%	<b>0.027<sup>a</sup></b>
		At 12 months	e13a2: 25.9% e14a2: 53.7% both: 40%	<b>0.037<sup>a</sup></b>
Jain et al. (2016) [49]	e13a2: 200 e14a2: 196 both: 85	Cumulative rate of MMR	e13a2: 79% e14a2: 91% both: 95%	<b>0.0001</b>
		Cumulative rate of MR4.5	e13a2: 57% e14a2: 79% both: 80%	<b>0.00001</b>
		MMR at 12 months: OR for e14a2 (95%CI)	e14a2: 5.85 (3.01–11.37) both: 3.29 (1.61–6.75)	< <b>0.001</b>
Castagnetti et al. (2017) [50]	e13a2: 203 e14a2: 290	Median time to MMR	e13a2: 12 months e14a2: 6 months	<b>0.001</b>
		Overall estimated probability of MMR	e13a2: 83% e14a2: 88%	< <b>0.001</b>
		Median time to MR4.0	e13a2: 61 months e14a2: 41 months	
		Overall estimated probability of MR4.0	e13a2: 52% e14a2: 67%	<b>0.001</b>
Claudiani et al. (2017) [51]	e13a2: 27 e14a2: 37	Cumulative incidence of loosing MR3 (95%CI)	e13a2: 64% (50–77) e14a2: 35% (15–56)	-
		Cumulative incidence of MR3 loss over time for e14a2: HR (95%CI)		
		Univariate analysis	0.4 (0.18–0.85)	<b>0.019</b>
		Multivariate analysis	0.38 (0.18–0.84)	<b>0.016</b>
Pagnano KBB et al. (2017) [52]	e13a2: 56 e14a2: 94 both: 20	Rate of optimal MR (BCR-ABL levels <10%) at 3 months	e13a2: 60% e14a2: 84% both: 75%	<b>0.02</b>
Breccia M et al. 2018 [41]	e13a2: 97 e14a2: 108 both: 3	Rate of MR4.5	e13a2: 31% e14a2: 43% both: 42%	<b>0.02</b>
		Probability to achieve MR4.5—Multivariate analysis: HR (95%CI) of e14a2 vs e13a2	1.6 (1.3–2.0)	<b>0.03</b>
Shanmuganathan et al. (2018) [53]	e13a2: 35 e14a2: 42 both: 18	Rate of TFR at 12 months	e13a2: 34% e14a2: 65%	<b>0.008</b>
		Probability to remain in TFR at 12 months—Multivariate analysis, HR (95%CI) of e14a2/both vs e13a2	2.24 (1.07–4.67)	<b>0.032</b>
D’Adda et al. (2019) [54]	e13a2: 67 e14a2: 106	Rate of MR4 at 60 months	e13a2: 52.4% e14a2: 82.2%	<b>0.008</b>
		Rate of sustained DMR at 60 months	e13a2: 26.9% e14a2: 47.2%	<b>0.003</b>
		Rate of TFR after 12 months	e13a2: 22±14% e14a2: 61±8%	<b>0.005</b>
Genthon A et al. (2020) [55]	e13a2: 51 e14a2: 63	Rate of MMR at 12 months	e13a2: 50.1% e14a2: 66.7%	<b>0.048</b>
		OR (95%CI)	3.25 (1.30–8.18)	<b>0.01</b>

Bold indicates statistically significant P-values (< 0.05)

<sup>a</sup>e14a2 vs e13a2 or e13a2/e14a2

N number of patients, MR4 a 4-log decrease in BCR-ABL1 transcript based on the International Scale, IM imatinib mesylate, MMR major molecular response, OR odds ratio, 95%CI 95% confidence interval, HR hazard ratio, MR molecular response, TFR treatment-free remission, DMR deep molecular response

BCR/ABL1 HT is complex and not yet standardized, therefore it was excluded from the tool.

### 3 Criteria for the Choice of First and Subsequent Lines of Tyrosine Kinase Inhibitor (TKI) Therapy

#### 3.1 Importance of Physician-Patient Communication

Effective communication represents the basis in establishing and strengthening the therapeutic alliance between physicians and patients, allowing the patient's expectations, objectives and attitudes to be explored [70]. In the

**Table 4** Choice of the first-line therapy according to the GIMEMA CML WP proposal [38]

	18–40 years	41–65 years	66–80 years	> 80 year
Low ELTS score	2GTKIs	<b>IM</b> –2GTKIs	IM	IM
Int ELTS score	2GTKIs	2GTKIs	<b>IM–2GTKIs</b>	IM
High ELTS score	2GTKIs	2GTKIs	<b>IM–2GTKIs</b>	IM

Bold indicates the grey zone in which therapy choice is more challenging

GIMEMA Gruppo Italiano Malattie EMatologiche dell'Adulto, ELTS European Treatment and Outcome Study for CML (EUTOS) Long-Term Survival, 2GTKIs second-generation tyrosine kinase inhibitors, IM imatinib mesylate

**Table 3** Tool summarising the factors that permit to plan a TFR: in orange, the conditions where a TFR is either not recommended or hardly achievable; in yellow and green, the intermediate and best possible scenarios, respectively

Factor	ORANGE	YELLOW	GREEN	Notes
<b>AT DIAGNOSIS</b>				
Gender		Male	Female	Female sex is predictive of a higher DMR rate vs male. Few studies showed a benefit in terms of TFR duration
Disease risk		High	Non-high	A high baseline risk allows to predict an inferior DMR rate vs non-high risk. The duration of TFR is not clearly risk-related. The ELTS score should be preferred
Type of transcript	Atypical transcript	e13a2 (b2a2)	e14a2 (b3a2)	The e14a2 transcript is associated with higher DMR rates and more durable TFR vs e13a2
<b>DURING TREATMENT</b>				
MR at 3 months	Level of transcript >10%	Level of transcript 1-10%	Level of transcript <1%	Failure of EMR at 3 and 6 months is associated to very low chances of achieving a DMR
MR at 6 months	Level of transcript >1%	Level of transcript 0.1-1%	Level of transcript <0.1%	
TKI, overall treatment duration	<3 years	3-5 years (IM) 3-4 years (2GTKIs)	>5 years (IM) >4 years (2GTKIs)	"Green milestones" predict optimal TFR duration
DMR duration	MR4.5: <1.0 year MR4.0: <2 years	MR4.5: 1-2 years MR4.0: 2-3 years	MR4.5: ≥ 2years MR4.0: ≥ 3 years	

DMR deep molecular response, TFR treatment-free remission, ELTS European Treatment and Outcome Study long-term survival, MR molecular response, EMR early molecular response, TKI tyrosine-kinase inhibitor, IM imatinib mesylate, 2GTKI second-generation TKI

context of TFR, despite the clear advantages, treatment discontinuation may raise concerns in patients because of fear of the unknown (individual probability of disease relapse) [71–76] and has been associated with anxiety and depression [77]. Thus, the physician’s ability to effectively communicate with patients, providing adequate information to reassure and motivate them, plays a central role [2]. This is particularly important when the treatment choice is controversial (see below). In the case of older patients, the involvement of caregivers is essential.

The panel endorsed the ELN 2020 recommendations [1] advising consideration of TFR after careful discussion with the patient, highlighting the importance of shared decision making.

### 3.2 Choosing the First-Line Therapy

TKIs are the standard of care for the first-line treatment of CP-CML, but current international recommendations, including the ELN recommendations, do not provide simple criteria to choose a specific inhibitor for each individual patient [1, 36, 45]. In view of treatment discontinuation, the first practical suggestion on how to choose the first-line therapy comes from the GIMEMA CML WP, who suggest that the patient’s age (i.e., 18–40 years; 41–65 years; 66–80 years; > 80 years) and the ELTS score should be considered [37] (Table 4).

The panel endorsed this proposal acknowledging that the choice is more challenging in some grey zones (i.e., low-risk patients aged 41–65 years and intermediate-/high-risk individuals aged 66–80 years), where the choice of first-line treatment needs a careful evaluation of each individual patient [37].

### 3.3 The Choice in Controversial Cases

The panel proposed to consider a number of patients’ and drugs’ characteristics to steer the choice of therapy in controversial cases (Table 5).

The panel acknowledged the importance of considering the type and severity of concomitant diseases [37, 78]; the patient’s comorbidities should be carefully evaluated and managed to minimize the risk of TKI toxicity, also in the long term [79]. Moreover, the panel included the patient’s expectations, individual attitudes and personal objectives among the factors to be taken into account when deciding whether TFR is a priority. A minor role was attributed to drug costs and physicians’ personal experience.

The panel acknowledged that, when the treatment goal is TFR, a longer therapy duration increases the chance of achieving it, even with the 2GTKIs [13].

Consequently, before TKI discontinuation, a dose reduction could be considered to reduce the side effects while

**Table 5** Tool to help steer the choice of therapy in first and second line (and beyond)

	Factors to be considered for the choice of therapy	
	First line	Second line (and beyond)
Patient	Disease risk (ELTS score, CCA in Ph+) Age Comorbidities (type, severity) Personal expectations and objectives	Reason of switch (Int vs. Fail/Warn) BCR-ABL1 mutations <sup>a</sup> Type of first-line TKI (IM vs. 2GTKIs) Comorbidities (severity, type) Age Personal expectations and objectives
Drug	Potency (efficacy; time to response) Side Effects (long-term safety)	Potency (efficacy; time to response) Side Effects (long-term safety)
Endpoints	Overall survival Treatment-free remission Quality of life	Overall survival Quality of life Treatment-free remission

ELTS European Treatment and Outcome Study for CML (EUTOS) Long-Term Survival, CCA clonal chromosomal abnormalities, Ph+ Philadelphia-positive, Int intolerance, Fail failure (i.e., loss of MMR), Warn warning (i.e., carefully consider the current treatment for continuation or change based on the patient’ characteristics comorbidities and tolerance), TKI tyrosine-kinase inhibitor, IM imatinib mesylate, 2GTKI second-generation TKIs

<sup>a</sup>As reported in the GIMEMA recommendations [38], of the 35 most frequent BCR-ABL1 kinase domain mutations associated with TKI resistance: all but five confer resistance to imatinib: M244V, L248V, G250E, Q252H, Y253H, E255V/K, L273M, D276G, T277A, E279K, F311L, T315I, F317L/V/I/C, M351T, E355G, F359C/I/V, E379K, L384M, L387M/F, H398R/P, E459K and F486S; Y253H, E255V/K, T315I and F359C/I/V to nilotinib; V299L, T315I/A, and F317L/V/I/C to dasatinib; E255V/K, V299L and T315I to bosutinib; T315M/L to ponatinib

continuing treatment [80–84]. The non-randomized, phase 2 De-Escalation and Stopping Treatment with Imatinib, Nilotinib, or Dasatinib (DESTINY) study evaluated the de-escalation to half the standard dose for 12 months followed by discontinuation for 24 months with frequent RT-qPCR monitoring [83]. In patients with stable MR4, 1 year of de-escalation before stopping resulted in a 2-year relapse-free survival of 72%, with improvement of common treatment-related symptoms [83]. Further insights, not only on TFR success but also on the incidence of the TKI withdrawal syndrome, will be provided by the ongoing Italian phase 2, prospective multicentre De-escAlation and discontinuation of Nilotinib ThErapy (DANTE) trial, evaluating the rate of TFR in selected CP-CML patients treated with nilotinib at half the standard dose during a consolidation period of 12 months, followed by complete cessation of therapy [84].

### 3.4 When to Switch the TKI

Approximately 40–50% of the patients on TKI therapy undergo a switch from first- to second-line therapy. The decision to change depends on the treatment response at 3, 6, 12 and 24 months. As shown in Table 6, the criteria for response definition (i.e., optimal, failure or warning) are clear and are similar across the ELN 2020 recommendations [1] and the GIMEMA proposal [37]. The only difference relies on the fact that the ELN definitions refer to treatment changes aimed at achieving optimal survival, whereas the GIMEMA response criteria aim at the achievement of a TFR. In particular, for the 24-month milestone the GIMEMA WP has given a more stringent definition of ‘optimal’ response, warning and failure [1, 37].

The panel endorsed the GIMEMA WP criteria [37].

### 3.5 Choosing the Second-Line Therapy

In keeping with other international recommendations, the GIMEMA WP did not provide clear indications on which TKI should be preferred for second-line therapy.

The panel acknowledged that the choice of second-line treatment (and beyond) is influenced by the reasons for switching, the presence and the type of BCR-ABL mutations, and the type of first-line TKI, considering that, in the case of resistance or warning to first-line treatment, the subsequent chances of reaching a TFR are very low or low, respectively [23, 31].

The variables to be considered in this setting are summarized in Table 5.

**Table 6** Comparison of response criteria [1, 36, 38]

	RESPONSE	GIMEMA 2019	NCCN 2019	ELN 2020
3 months	Optimal	≤ 10	≤ 10	≤ 10
	Warning	-	> 10	> 10
	Failure	> 10 (confirmed)	-	-
6 months	Optimal	≤ 1	≤ 10	≤ 1
	Warning	1-10	-	1-10
	Failure	> 10	> 10	> 10
12 months	Optimal	≤ 0.1	≤ 1	≤ 0.1
	Warning	0.1-1	1-10	0.1-1
	Failure	> 1	> 10	> 1
24 months	Optimal	≤ 0.01	≤ 1	≤ 0.1
	Warning	0.1-0.01	-	0.1-1
	Failure	> 0.1	> 1	> 1

*GIMEMA* Gruppo Italiano Malattie EMatologiche dell'Adulto, *ELN* European LeukemiaNet, *NCCN* National Comprehensive Cancer Network

The panel endorsed the 2020 ELN recommendations [1]: In the case of intolerance, any TKI can be used; in the case of resistance to IM, the choice is a 2GTKI, whereas in the case of resistance to a 2GTKI, the choice is ponatinib.

## 4 The Frequency of Monitoring During TFR

### 4.1 The Frequency of Molecular Monitoring After Discontinuation: Focus on the Second Semester

After cessation of therapy, the frequency of monitoring must ensure patient safety, promptly detecting possible loss of MMR and allowing resumption of therapy before the loss of cytogenetic or hematological responses. Albeit relapse occurs predominantly within the first 6 months of TFR [3, 13, 47], the response may be lost even 2 years after discontinuation [85]: data from 128 patients included in the prospective discontinuation trial A-STIM indicate that very late recurrence (i.e., > 2 years after discontinuation) occurs in nearly 14% of patients. Moreover, patients with a stable MR4.5 had a 100% probability of remaining in TFR for more than 2 years versus 65.4% in those with an unstable remission [85]. Notably, the kinetics of relapse differs between patients relapsing very early or those who do not [86].

The first indications on the frequency of monitoring were, and are, rather stringent [87]. In clinical practice, it is expected that the frequency of monitoring will be prolonged. Current guidelines consistently recommend monitoring patients more frequently during TFR than during therapy



(i.e., monthly by RT-qPCR for at least 6 months [1, 36, 37]), but there is no agreement about the frequency of monitoring after the first 6 months [1, 36], and neither did the GIMEMA WP reach a consensus [37].

Shanmuganathan et al. [88] evaluated different monitoring algorithms. They found that performing molecular tests every 2 months for the first 6 months and every 3 months thereafter in patients maintaining MR4.5 provided the best balance between reduced testing and shorter delays in detecting relapse and resuming TKI [88]. A lower frequency (i.e., every 3 months) resulted in many patients losing cytogenetic and hematological responses, which is not acceptable [88]. The need for a stringent follow-up for early diagnosis of MMR loss has been confirmed in a sub-analysis of an Italian observational study, which retrospectively investigated the modalities of molecular monitoring performed in 227 patients with CML in a real-life setting [25, 47]. By month 6, 46% of patients had three or fewer molecular assessments and 77.5% of those off treatment (N = 71) who relapsed, did so during the first 6 months. The authors calculated that if a 3-month monitoring schedule was adopted, > 50% of patients would have experienced a delay in the detection of MR loss up to month 6, very few patients between month 6 and 12, and even fewer afterwards [47].

The panel agreed that, given the absence of factors reliably predicting the stability of TFR during the first 6 months, the monthly monitoring is mandatory. In difficult situations (i.e., advanced age, comorbidities, difficulty accessing the lab), it is advisable to avoid offering the TFR. After the first 6 months, the monitoring frequency depends on the careful evaluation of what happened before, and can be adjusted based on the stability of MR4.5 (Table 7).

At present, it is not possible to establish whether any baseline factor can influence the frequency of monitoring.

## 4.2 Turnaround Time and Possible Role of Digital Droplet Polymerase Chain Reaction (PCR)

Current indications on the turnaround time (TAT) of the RT-qPCR test results are different: within 15 days for the National Comprehensive Cancer Network (NCCN) guidelines [36], within 4 weeks for Hughes et al. [89], and “rapidly” for the ELN 2020 recommendations [1]. During the first 6 months off treatment, it is mandatory to receive the results within 4 weeks, that is before the next blood withdrawal. In Italian real-life, the TAT is about 3 weeks, which is adequate in the first 6 months. Afterwards, the mean TAT of the reference lab must be considered to ensure prompt intervention in the case of MMR loss.

**Table 7** Frequency of molecular monitoring in case of stable and unstable MR4.5 maintained during the first 6 months after discontinuation. After month 6, patients with unstable MR4.5 must be monitored at closer intervals than those with a stable MR4.5. When personalizing the frequency of monitoring, it is important to consider the turnaround time (TAT) of the reference lab

Months after TKI cessation	MR4.5 up to month 6	
	Stable	Unstable
0–6	Monthly	Monthly
7–12	Every 2–3 months	Every 1.5–2 months
> 12	Every 4 months	Every 3 months

TKI tyrosine-kinase inhibitor, MR4.5 molecular response defined as a 4.5-log decrease in BCR-ABL1 transcript based on the International Scale, Stable MR4.5 up to month 6 confirmed MR4.5 in all QPCR tests performed in the first 6 months after discontinuation

The panel agreed that if a TAT of 4 weeks cannot be assured, the interval between the molecular tests can be extended up to the higher limit of the intervals reported in Table 7.

Novel monitoring techniques with improved sensitivity for detection of residual disease, such as ddPCR, could help to refine the selection of candidates for TFR and provide useful information during TFR. Experiences in this field are discordant. In a cohort of 142 CML patients monitored for a median time of 24 months, ddPCR was more sensitive and accurate than standard RT-qPCR in measuring minimal residual disease [90]; among the 111 patients who discontinued treatment, a threshold value of 0.468 BCR-ABL1 copies/ $\mu$ L at TKI stop was able to predict the maintenance of TFR [90]. In contrast, RT-qPCR was unable to discriminate patients with a higher risk of MMR loss after discontinuation [90]. Similarly, the evaluation of BCR-ABL1 by a combination of DNA and RNA measurement during DMR maintenance before TKI cessation was more effective than standard RT-qPCR performed on cDNA in predicting the success of TFR [91]. Interestingly, a recent update of the Italian discontinuation ISAV study showed that ddPCR was able to detect increasing BCR-ABL1 values in most non-relapsing patients after 36 months after discontinuation, reinforcing the need for regular long-term monitoring of TFR patients in order to detect late progressions [92].

However, other studies failed to identify differences in the sensitivity for BCR-ABL1 detection between qPCR and ddPCR [93, 94].

The panel agreed that standard RT-qPCR is the technique of choice during TFR. ddPCR is a potential alternative to RT-qPCR [95, 96] but not yet standard-

ized [97]. Therefore, its implementation in clinical practice is not yet advisable.

## 5 The Prognosis of TFR

### 5.1 Duration of Therapy

Therapy duration is one of the most relevant prognostic factors for the success of TFR. The EURO-SKI trial demonstrated that in patients receiving first-line IM, longer treatment duration was associated with a continuous, per-year increase in the probability of maintaining MMR at 6 months (odds ratio [OR] per year: 1.14, 95% confidence interval [CI] 1.05–1.23;  $P = 0.0010$ ) [13]. Recently, duration of nilotinib therapy before the start of TFR was identified as the only significant predictor of a durable TFR in a pooled analysis of data from ENESTfreedom and ENESTop [98]. The authors found that each 1-month increase in therapy duration was associated with a 3.6% (95% CI 1.5–5.8) increase in the odds of durable TFR at week 24 [98].

The GIMEMA CML WP agreed that, to ensure safety and avoid disease progression, the minimum duration of treatment should be 3 years with a 2GTKI and 5 years with IM, but no consensus was reached on the definition of ‘optimal’ duration [37]. The ELN 2020 recommendations, published soon after the GIMEMA proposal, acknowledged a duration of TKI therapy > 5 years with IM and > 4 years for 2GTKI among the ‘minimal’ requirements (i.e., stop allowed) for therapy cessation. A duration > 5 years was defined as ‘optimal’ (i.e., stop recommended for consideration) [1]. These differences in the number of years considered by the GIMEMA CML WP and ELN are due to the lack of definitive, prospectively validated cut-off values; still, the longer the duration of therapy, the higher the likelihood of maintaining MMR at 6 months [13].

The panel endorsed the minimum and optimal criteria reported in the ELN recommendations [1] (Table 3).

### 5.2 Duration of the DMR

Similar to therapy duration, no unequivocal cut-off values are available for DMR duration; indeed, those provided by the GIMEMA CML WP and ELN differ from each other [1, 37]. In patients treated with IM as first-line therapy, longer DMR duration (OR per year: 1.13, 95% CI 1.04–1.23;  $P = 0.0032$ ) was associated with increasing probability of MMR maintenance at 6 months [13].

In clinical practice, a score to predict the success of TFR would be very useful, but despite many attempts, at present none is available. Claudiani et al. proposed a score including

duration of MR4, previous TKI resistance, age at diagnosis and transcript type [99]. Nonetheless, the score has not been independently validated and it does not include any indication on the minimal duration of DMR.

The panel agreed with the minimal and optimal criteria of DMR duration reported in the 2020 ELN recommendations [1] (Table 3).

### 5.3 Immunological Factors: Is It Possible to Develop an Immunological Score?

The mechanisms underpinning the lack of overt relapse in about 50% of patients off treatment are still unclear. A possible role has been attributed to host factors, such as the effector-mediated immune surveillance (impaired in CML patients before TKI therapy and restored by the use of TKIs) and/or to the leukemic stem cells (LSCs) [81, 100, 101]. However, although the peripheral blood cell populations involved in TFR are well characterized, the LSC compartment and target antigens are not well known [102], and to date there is no immunological predictive phenotype that may be used in clinical practice.

The immunological control of CML changes throughout the different phases of the disease (pre-TKI therapy, during TKI therapy and during TFR), with immunoescape failing progressively and immunosurveillance prevailing upon TKI therapy [100]. Immunoescape is unanimously defined as the result of the activity of programmed cell death protein 1-positive T cells, regulatory T cells (Tregs) and myeloid-derived suppressor cells belonging to the BCR/ABL1 clone. Conversely, natural killer (NK) cells and cytotoxic T lymphocytes are responsible for immunosurveillance. Over time, during TKI therapy, the system first reaches equilibrium and then, in the TFR phase, immunosurveillance prevails over immunoescape [100].

Regarding the control of disease at the LSC level the panel discussed the possible existence of a particular antigen able to stimulate the immune system, leading to disease control and culminating in the achievement and maintenance of TFR. The findings from Bocchia et al. [103] do not support the hypothesis that CD26 might be the actual target of immunosurveillance: indeed, the authors found circulating CD26+ LSCs in a relevant proportion of CP-CML patients during first-line TKI therapy and in TFR, with similar numbers (on therapy: median 0.014 cells/ $\mu$ L; range 0.0012–0.66 cells/ $\mu$ L; in TFR: median 0.015 cells/ $\mu$ L; range 0.006–0.76 cells/ $\mu$ L) [103]. Moreover, the number of residual LSCs did not correlate with the BCR-ABL1/ABL1<sup>IS</sup> ratio [103].

The panel agreed that there are no convincing data supporting a key role of any particular antigen in the immune control of the minimal residual disease.

Other published scores were analyzed but the data were not conclusive: (1) the ‘effector-suppressor’ score [104], based on the absolute count of NK cells, Tregs and MSDCs at the time of discontinuation; (2) a score based on individual immunologic configurations (i.e., quiescent LSCs, proliferating LSCs and effector cells, with a complex mathematical model) [105]; (3) a score based on proteomics and (inflammatory) cytokines [106].

Finally, a recent study conducted on 19 patients reported detection of BCR/ABL1 on genomic DNA from sorted B lymphocytes, suggesting that the lineage-specific assessment of minimal residual disease in individual patients may contribute to predicting the success of TFR [107].

The panel acknowledged the lack of tools based on immunological parameters in clinical practice for the prediction of TFR and that more research is needed. The panel agreed that the total duration of TKI treatment and DMR duration are the pivotal factors to predict the stability of TFR. Other parameters, namely the transcript type, the rapidity of the MR, the Sokal or ELTS risk score, gender and age, may significantly influence the success of TFR, but data are not conclusive.

## 6 Conclusions and Future Perspectives

TFR is a complex matter and, despite the large number of clinical trials conducted thus far, several issues remain unsolved. Of the current recommendations and proposals, only those from the GIMEMA CML WP focused on TFR as a primary goal of therapy.

In this paper, we addressed the topics that we deemed as the most urgent based on our clinical practice experience (early prediction of DMR and TFR; criteria for first and subsequent line therapy choice; monitoring of residual disease after discontinuation; prognosis of TFR). The originality (and main strength) of our work, compared with similar recent papers reviewing the literature and discussing controversial points, resides in the practical tools provided for each topic, which are intended to facilitate the decision-making process in daily practice.

However, a few points remained unsolved. In the future, it will be important to: (1) clarify whether the amplification efficiency of the e13a2 and e14a2 transcripts is really different, because this would have an impact on their predictive value of DMR; (2) assess the role of first or subsequent-line of treatment with 2G-3GTKIs and new drugs in maximizing the probability of TFR without increasing toxicity; (3) implement novel sensitive techniques for the standardized monitoring of residual disease, such as ddPCR; (4) elucidate the mechanisms behind the immunological control of CML,

in order to identify possible markers to predict the risk of relapse after TKI cessation through an immunological score.

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## Declarations

**Conflicts of interest** *Fausto Castagnetti* has received consultancy fees and honoraria from Novartis, Incyte, Pfizer and Bristol Myers Squibb; *Gianni Binotto* has received honoraria and speaker fees from Novartis, Pfizer, Incyte, BMS-Celgene; *Isabella Capodanno* has received honoraria and speaker fees from Novartis, Pfizer and Incyte; *Atto Billio* declares no conflicts of interest; *Elisabetta Calistri* declares no conflicts of interest; *Francesco Cavazzini* has received honoraria and speaker fees from Novartis, Pfizer and Incyte; *Monica Crugnola* has received honoraria from Novartis and Incyte; *Antonella Gozzini* has received honoraria from Novartis, Incyte and Bristol Myers Squibb; *Gabriele Gugliotta* has received honoraria/consultancy fees from Novartis, Pfizer and Incyte; *Mauro Krampera* has received honoraria/consultancy fees from Novartis; *Alessandro Lucchesi* has received honoraria from Novartis, Incyte and Pfizer; *Anna Merli* declares no conflicts of interest; *Maria Cristina Miggiano* declares no conflicts of interest; *Claudia Minotto* has received honoraria for serving on the advisory board from Novartis; *Monica Poggiaspalla* declares no conflicts of interest; *Marzia Salvucci* declares no conflicts of interest; *Barbara Scappini* has received honoraria and speaker fees from Novartis, Amgen Sanofi, Jazz and Incyte; *Mario Tiribelli* has received honoraria and speaker fees from Novartis, Pfizer and Incyte; *Elena Trabacchi* declares no conflicts of interest; *Gianantonio Rosti* has received consultancy fees and honoraria from Novartis, Incyte, Pfizer and Bristol Myers Squibb; *Sara Galimberti* has received honoraria from Novartis, Incyte and Pfizer; *Massimiliano Bonifacio* has received consultancy fees from Novartis, Pfizer, Incyte and Amgen.

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