Impact of BCR-ABL <10% at 6 Months on Outcome of Patients with CML with a Poor Response at 3 Months
CME INFORMATION

OVERVIEW OF ACTIVITY

Each year, thousands of clinicians, basic scientists and other industry professionals sojourn to major international oncology conferences, like the American Society of Hematology (ASH) annual meeting, to hone their skills, network with colleagues and learn about recent advances altering state-of-the-art management in hematologic oncology. As such, these events have become global stages where exciting science, cutting-edge concepts and practice-changing data emerge on a truly grand scale. This massive outpouring of information has enormous benefits for the hematologic oncology community, but the truth is it also creates a major challenge for practicing oncologists and hematologists.

Although original data are consistently being presented and published, the flood of information unveiled during a major academic conference is unprecedented and leaves in its wake an enormous volume of new knowledge that practicing oncologists must try to sift through, evaluate and consider applying. Unfortunately and quite commonly, time constraints and an inability to access these data sets leave many oncologists struggling to ensure that they’re aware of crucial practice-altering findings. This creates an almost insurmountable obstacle for clinicians in community practice because they are not only confronted almost overnight with thousands of new presentations and data sets to consider but they are also severely restricted in their ability to review and interrogate the raw findings.

To bridge the gap between research and patient care, this CME activity will deliver a serial review of the most important emerging data sets on the management of chronic myeloid leukemia (CML) from the latest ASH meeting, including expert perspectives on how these new evidence-based concepts may be applied to routine clinical care. This activity will assist medical oncologists, hematologists, hematology-oncology fellows and other healthcare professionals in the formulation of optimal clinical management strategies and the timely application of new research findings to best-practice patient care.

LEARNING OBJECTIVES

- Evaluate the impact of early molecular response or dose interruption of tyrosine kinase inhibitors (TKIs) on the prognosis of patients with CML.
- Compare and contrast the benefits and risks of nilotinib versus imatinib therapy in patients with newly diagnosed chronic-phase CML.
- Appraise recent clinical data on the effect of switching to nilotinib in patients with a suboptimal response to imatinib therapy versus continuation of imatinib at a higher dose.
- Analyze the outcomes of the STIM1 and STIM2 studies of discontinuation of imatinib in patients with a deep molecular response, and consider these results in the management of CML.
- Assess the efficacy and safety of ponatinib as initial therapy and in patients with TKI-resistant CML.

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FACULTY — The following faculty (and their spouses/partners) reported real or apparent conflicts of interest, which have been resolved through a conflict of interest resolution process:

- Jorge E Cortes, MD
  - D B Lane Cancer Research
  - Distinguished Professor for Leukemia Research
  - Deputy Chairman, Section Chief of AML and CML
  - Department of Leukemia
  - The University of Texas MD Anderson Cancer Center
  - Houston, Texas
  - Consulting Agreements: Bristol-Myers Squibb Company, Genentech BioOncology, Lilly, Novartis Pharmaceuticals Corporation, Pfizer Inc, Sanofi; Contracted Research: Bristol-Myers Squibb Company, Celgene Corporation, Novartis Pharmaceuticals Corporation, Pfizer Inc, Sanofi.
- Hagop M Kantarjian, MD
  - Chairman and Professor, Leukemia Department
  - The University of Texas MD Anderson Cancer Center
  - Houston, Texas

No real or apparent conflicts of interest to disclose.

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Hardware/Software Requirements:
A high-speed Internet connection
A monitor set to 1280 x 1024 pixels or more
Internet Explorer 7 or later, Firefox 3.0 or later, Chrome, Safari 3.0 or later
Adobe Flash Player 10.2 plug-in or later
Adobe Acrobat Reader
(Optional) Sound card and speakers for audio

Last review date: February 2014
Expiration date: February 2015
Sometimes I have to pinch myself to see if this is a dream or if I really have a job listening to and learning from the great minds in our chosen field. Last week was a perfect reminder of just how cool “work” can be when within the space of a few days my calendar included extensive interviews with Drs Jorge Cortes and then Hagop Kantarjian. As deputy chair and chair of MD Anderson’s Department of Leukemia, respectively, these 2 investigators lead a unique clinical and research powerhouse that has contributed perhaps as much to the care of patients with these and other related hematologic disorders as any other institution in the world.

To get a sense of just how prolific they are, peruse the 2013 ASH abstracts and you will find that Drs Cortes and Kantarjian helped author 103 oral presentations and posters, including 30 on chronic myelogenous leukemia (CML) alone. As such, and not surprisingly, each of these conversations focused heavily on that disease — which has become the poster child for targeted oncologic treatment — and below find the bottom line on their thoughts about how the data sets from New Orleans helped address the following important questions in CML.

1. **What are the key early markers of response, and when should consideration be given to switching to another tyrosine kinase inhibitor (TKI)?**

Another MD Anderson leukemia maven and chair of the NCCN CML guidelines committee, Dr Susan O’Brien frequently reinforces the important concept that although there are many reasons to seek deep molecular responses (DMR), the classic and most important endpoint is complete cytogenetic response (CCyR) — a milestone that is achieved faster and more frequently with the second-generation agents, nilotinib and dasatinib. The question of whether suboptimal molecular response should trigger a switch to another TKI ties directly into the issue of selection of up-front therapy and whether long-term outcomes are compromised when residual disease is present.
Equally relevant and looming in the background is a fascinating question of “quality” and cost associated with oncology care. Specifically, imatinib is due to go off patent in January 2015, and it is expected that this will dramatically lower the annual tab (about $90,000 with imatinib, and with nilotinib and dasatinib closer to $100,000). With a current prevalence of about 100,000 CML cases in the United States alone — a number that will likely double in the next 3 decades before plateauing — researchers, clinicians and policy makers will almost certainly continue the debate about the value of starting with imatinib (the soon-to-be less costly and perhaps slightly less effective agent) and reserving second-generation treatment for patients with higher-risk disease and those with suboptimal initial responses to imatinib. How these potential resource savings stack up against others in oncology related to, for example, futile care and unnecessary imaging will be discussed extensively, and more globally Dr Kantarjian has taken a leadership role in organizing a group of “CML experts” (including Dr Cortes) who have been on a dedicated and major offensive attacking the current CML cost structure.

At ASH we witnessed a number of related papers that tie in to the issue of imatinib versus the rest, including the 36-month update of the ENESTcmr study. This landmark Phase III effort demonstrated that among patients in CCyR but with detectable BCR-ABL transcripts, those randomly assigned to switch to nilotinib achieved more DMRs compared to those continuing on imatinib (47% with nilotinib versus 33% with imatinib at 36 months). This benefit came with greater toxicity, which may in part be attributable to the trial design in that patients who transitioned to nilotinib were already tolerating imatinib well.

On a similar note, an ASH data set presented by Dr Cortes from the Phase III LAsoR trial revealed that switching to nilotinib versus escalating the dose of imatinib in patients who experienced suboptimal response resulted in a better rate of CCyR at 6 months (49% versus 42%, respectively), although the findings were not statistically significant ($p = 0.3844$).

Finally, a retrospective analysis of 3- and 6-month responses in early trials of imatinib demonstrated that some patients who achieve an optimal response by 6 instead of 3 months have long-term outcomes comparable to those who achieved an optimal response at 3 months, suggesting that waiting a few additional months before considering a change in treatment is a rational approach.

Proponents of using imatinib as initial treatment in standard-risk situations often point out that so far, no survival benefit has been demonstrated using the second-generation agents — possibly because these drugs also effectively rescue patients experiencing disease progression on imatinib. Thus, although DMR is an intuitively appealing goal, until further research identifies more accurately who can cease TKI treatment (now there’s a cost saving!), there will be debate and controversy about what to start with and when and if to make a switch. This is particularly true as more follow-up occurs
with the landmark second-generation trials, some of which are documenting more long-term complications, such as the 5-year update of the ENESTnd trial presented at ASH that now shows not only deeper molecular responses with nilotinib but also an increasing number of cardiovascular events.

2. Are there situations in which it is safe to discontinue TKI treatment?

At ASH we saw more data from 2 French studies (STIM 1 and 2) attempting to define the outcomes of patients with prolonged (more than 2 years) DMRs who discontinued treatment. These studies and others have documented that when taken off therapy more than half the patients experience relapse — usually quickly — and the remainder fare well off treatment. Importantly, although most patients experiencing relapse can be effectively salvaged with the same or a different TKI, at this point there is no way to pick who will do well without treatment and therefore neither professor employs this approach outside a trial setting, although Dr Kantarjian notes that if ongoing research shows how to identify these patients, both long-term toxicity and financial costs can be avoided.

Interestingly, Dr Cortes commented on one situation in which a variation of this stopping strategy is often a consideration — specifically, in women with CML who wish to become pregnant — and so far he has managed about 2 dozen carefully selected patients, most of whom have not required retreatment until after childbirth.

Another fascinating and somewhat related ASH report documented that in a major Phase III trial of dasatinib versus imatinib patients starting treatment who missed doses due to toxicities like cytopenias had significantly worse 3-month outcomes. Importantly, this effect appears to occur when missing even 1 dose (in the case of imatinib) and increases with the number of doses missed.

3. What is the current role of ponatinib?

In December 2012 this pan-BCR-ABL “super TKI” was approved by the FDA, but last October it was pulled off the market due to toxicity concerns, mainly arteriothrombotic events. By December ponatinib was once again available, accompanied by a new black box warning and a Risk Evaluation and Mitigation Strategy program designed to help clinicians more effectively evaluate the risks and benefits of using the agent.

In discussing ponatinib, Dr Kantarjian noted that the approved daily dose of 45 mg not uncommonly leads to toxicities such as hypertension, vasospastic reactions, pancreatitis and skin rashes that are not acceptable in the up-front setting, where safer effective choices exist. In this regard an MD Anderson single-arm pilot study of 51 patients presented at ASH was amended to include a starting dose of 30 mg daily. Regardless, accrual was suspended in October, as in another major Phase III up-front study comparing ponatinib to imatinib.
However, in discussing the updated ASH results from the pivotal PACE trial in relapsed disease, Dr Kantarjian reiterated that ponatinib, when used in that indicated setting, can be a life-altering therapy, particularly for those with BCR-ABL T315I mutations. He also pointed out that the vaso-occlusive reactions that have been observed with this drug occur infrequently with the other TKIs.

Next on this series, we provide an update on ASH reports in lymphoma, including encouraging data sets on the nonchemotherapy combination of lenalidomide and rituximab, the antibody-drug conjugate brentuximab vedotin and a fascinating paper on crizotinib in ALK-positive lymphoma.

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Impact of BCR-ABL <10% at 6 Months on Outcome of Patients with CML with a Poor Response at 3 Months

Presentation discussed in this issue

Branford S et al. Any BCR-ABL reduction below 10% at 6 months of therapy significantly improves outcome for CML patients with a poor response at 3 months. Proc ASH 2013;Abstract 254.

Slides from a presentation at ASH 2013 and transcribed comments from a recent interview with Jorge E Cortes, MD (1/24/14)

Any BCR-ABL Reduction Below 10% at 6 Months of Therapy Significantly Improves Outcome for CML Patients with a Poor Response at 3 Months

Branford S et al.

Proc ASH 2013;Abstract 254.
Background

- The molecular response at 3 months after commencement of tyrosine kinase inhibitor (TKI) therapy for patients with CML has prognostic significance.
- Analyses by Neelakantan et al suggest that additional measurement of BCR-ABL1 transcript levels at 6 months adds little prognostic value to the 3-month result (Blood 2013;121:2739).
- However, another recent study based on cytogenetic response concluded that for patients with poor response at 3 months, assessing the response at 6 months may provide a better predictor of long-term outcome (Haematologica 2013;98:1686).
- **Study objective:** To evaluate the prognostic importance of assessing both the 3- and 6-month molecular response for patients with chronic-phase CML (CML-CP).


Study Design

- The study included patients with CML-CP enrolled in consecutive clinical trials of first-line imatinib from 2000 to 2011 (n = 528).
  - Many patients were treated before alternative TKIs were available, but 89 switched therapy.*
- The utility of BCR-ABL as a predictor of death (overall survival), progression (AP/BC: progression-free survival), treatment failure (failure-free survival) and major molecular response (MMR) was assessed.
- Patients were divided according to the 2013 European LeukemiaNet (ELN) definitions of 3- and 6-month molecular response:
  - 3 mo, optimal ≤10% or warning >10%
  - 6 mo, optimal <1%, warning 1-10% or failure >10%

*Study was not powered to assess the effect of treatment intervention

Outcomes at 4 Years for Patients in the Optimal (≤10%) versus Warning (>10%) Category at 3 Months

<table>
<thead>
<tr>
<th>Outcome at 4 y</th>
<th>Optimal (n = 406)</th>
<th>Warning (n = 100)</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Overall survival</td>
<td>97%</td>
<td>89%</td>
<td>0.0003</td>
</tr>
<tr>
<td>Progression-free survival</td>
<td>99%</td>
<td>86%</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Failure-free survival</td>
<td>83%</td>
<td>46%</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>MMR</td>
<td>89%</td>
<td>42%</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>


Survival of Patients in the 3-Month Warning Category Grouped by Category at 6 Months

Overall Survival (n = 89)

- Optimal 100%
- Warning 100%
- Failure 71%

Progression-Free Survival (n = 89)

- Optimal 100%
- Warning 94%
- Failure 72%

With permission from Branford S et al. Proc ASH 2013;Abstract 254.
MMR for Patients in the 3-Month Warning Category Grouped by Category at 6 Months

- Patients in the warning category at 3 months who have BCR-ABL1 <10% at 6 months have improved outcomes
- No significant difference in any outcome assessment after 6 months between those who were in the optimal category at 3 months and 6 months versus those in the warning category at 3 months who moved to the optimal category at 6 months

With permission from Branford S et al. Proc ASH 2013;Abstract 254.

Patients at High Ongoing Risk of Poor Response

Change of BCR-ABL1 level from baseline to 3 months was important for outcome

With permission from Branford S et al. Proc ASH 2013;Abstract 254.
Use of Halving Time to Predict Outcome for Patients at High Ongoing Risk of Poor Response

When BCR-ABL1 was measured as a continuous covariate, patients with the same value at 3 months had better outcomes if their baseline value was higher.


Outcomes for Patients in the Warning Category at 3 Months by Halving Time Responses

<table>
<thead>
<tr>
<th>Outcome at 4 y</th>
<th>Overall (n = 100)</th>
<th>Halving time response</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Overall (n = 100)</td>
<td>≤90 d (n = 79)</td>
<td>&gt;90 d (n = 19)</td>
</tr>
<tr>
<td>Overall survival</td>
<td>89%</td>
<td>93%</td>
<td>69%</td>
</tr>
<tr>
<td>Progression-free survival</td>
<td>86%</td>
<td>90%</td>
<td>69%</td>
</tr>
<tr>
<td>Failure-free survival</td>
<td>46%</td>
<td>56%</td>
<td>7%</td>
</tr>
<tr>
<td>MMR</td>
<td>42%</td>
<td>53%</td>
<td>5%</td>
</tr>
</tbody>
</table>

The halving time at 3 months may also be predictive of overall and progression-free survival for the 35 patients who subsequently met the ELN failure criteria at 6 months.

Author Conclusions

- BCR-ABL1 >10% at 3 months is a poor risk category.
- Not all patients with a BCR-ABL1 value >10% at 3 months have a high ongoing risk of treatment failure.
  - Any reduction below 10% by 6 months may improve outcome.
  - The rate of reduction over the first 3 months is an important factor for outcome and could be considered when making therapeutic decisions.


Investigator Commentary: BCR-ABL Levels <10% at 6 Months Significantly Improve Outcome for Patients with CML-CP with a Poor Response at 3 Months

This study showed that some patients who do not have a good molecular response at 3 months may be able to catch up at 6 months, whereas others continue to have a poor response. The patients who catch up at 6 months have the same good prognosis as those who achieve the response at 3 months. Those who do not catch up even by 6 months will have a poor outcome. This has important implications for how we care for patients who have BCR-ABL levels >10% at 3 months.

My recommendation is not to change treatment for any patient at 3 months but to ensure that the patients are monitored at 6 months. I would consider changing the treatment for those who continue to respond poorly.

With imatinib, about a third of patients don’t achieve a good response at 3 months and about half of these patients will continue to fare poorly at 6 months. However, with dasatinib or nilotinib, only 10% to 15% of patients will not have a good response at 3 months and half of those will continue to respond poorly at 6 months. That is a rationale for using dasatinib or nilotinib as up-front therapy.

Interview with Jorge E Cortes, MD, January 24, 2014