Cardiotoxicity of tyrosine kinase inhibitors in chronic myelogenous leukemia therapy
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Abstract
Emerging evidence suggests that the three tyrosine kinase inhibitors currently approved for the treatment of patients with chronic myelogenous leukemia (CML) – imatinib, dasatinib, and nilotinib – have potential cardiotoxic effects. The mechanisms behind these events, and the relations between them, are largely unclear. For example, relative to dasatinib and nilotinib, severe congestive heart failure and left ventricular dysfunction are rare but prominent with imatinib treatment, particularly in patients receiving higher doses (>600 mg/day). In comparison with imatinib, prolongation of the QT interval is relatively common in patients treated with either dasatinib or nilotinib. In contrast to nilotinib, pericardial effusions are observed with both imatinib and dasatinib. It is suggested that these data, an evaluation of cardiac status, use of concomitant medications, and potential risk factors should be considered in the management of CML.

Introduction
BCR-ABL tyrosine kinase is a key molecule responsible for the pathophysiology of chronic myelogenous leukemia (CML). Tyrosine kinase inhibitors (TKIs) directed against BCR-ABL are currently the cornerstone of treatment for patients with CML. Imatinib (Gleevec®; Novartis Pharmaceuticals Corporation, New Jersey, USA) was the first TKI approved for CML. Imatinib has marked response and survival benefits over interferon-α plus low-dose cytarabine and is presently the only TKI licensed for first-line treatment.1 However, despite the remarkable success of imatinib, many patients discontinue treatment because of either resistance or intolerance to this drug.

In the pivotal phase III IRIS (International Randomized Study of Interferon and STI571) trial, primary resistance was observed in 24% of patients, and secondary resistance presented as relapse in 17% of patients and disease progression in 7% after 4.5 years.2 After six years of follow-up, 34% of patients had discontinued imatinib treatment, mostly (14%) because of an unsatisfactory therapeutic effect (defined as lack of efficacy/progression), but a number of patients (5%) also stopped receiving the drug as a result of adverse events (AEs) or abnormal laboratory values.3 Further treatment options therefore continue to be developed.

Second-line choices for treatment include increasing the dose of imatinib, or changing therapy to dasatinib [Sprycel®; Bristol-Myers Squibb Co. (BMS), New York, USA] or nilotinib (Tasigna®; Novartis Pharmaceuticals Corp., New Jersey, USA). Newer agents for imatinib failures are still under clinical development.4 Dasatinib potently inhibits BCR-ABL.5 Compared with imatinib, dasatinib has 325-fold greater activity against native BCR-ABL.6 Furthermore, dasatinib inhibits all imatinib-resistant mutations of this molecule (key mediators of imatinib resistance) except the T315I mutant, which is resistant to all currently available TKIs.6,7 It is, however, relatively insensitive to the F317L mutation.3 Dasatinib was originally approved at the dosage of 70 mg twice daily, following data from the START (the SRC/ABL Tyrosine Kinase Inhibition Activity: Research Trials of Dasatinib) program of clinical studies.8,9 In November 2007, the label for dasatinib was changed to include updated dosing information, safety information from more than 2,100 patients, and data from a randomized comparison with high-dose imatinib.10 The recommended starting dose for patients with chronic phase (CP) CML is now 100 mg once daily. The starting dose for advanced disease remains 70 mg twice daily.

Nilotinib (an analog of imatinib) is also active against all imatinib-resistant BCR-ABL mutations except T315I.11 However, nilotinib is relatively inactive against common mutations in the ATP-binding P-loop domain,4 and those at the F359 residue in the catalytic domain, of BCR-ABL.12 Patients harboring such mutations at baseline do not respond to nilotinib and progress quickly during treatment. Furthermore, patients who do not have these mutations at baseline may eventually develop them when resistance to imatinib occurs.13 Nilotinib has recently been approved by the FDA for treating imatinib-resistant and -intolerant patients with CP or accelerated phase CML, but not those suffering from blast crisis.

TKIs share a number of common adverse effects (AE), including neutropenia and thrombocytopenia. Cardiotoxicities of the TKIs have become a clinical concern. This review provides an overview of common AEs and cardiotoxicities associated with these TKIs.

Common adverse events associated with tyrosine kinase inhibitors
A number of AEs are associated with TKIs. The most common ones are hematologic. In the IRIS study, anemia, neutropenia, and thrombocytopenia (all grades) were each reported by 45-61% of patients receiving imatinib.1 Grade 3-4 anemia, neutropenia, and thrombocytopenia were observed in 3%, 14%, and 8% of patients, respectively. The updated dasatinib label includes data from the START-R trial, a comparison of dasatinib 70 mg twice daily with high-dose imatinib (800 mg/day) in patients with CP CML resistant to standard-dose imatinib.13 In this study, the incidences of grade 3-4 thrombocytopenia and neutropenia for high-dose imatinib were 14% and 39%, respectively. The START-R study also showed that dasatinib has significant response and progression-free survival benefits compared with high-dose imatinib, and that significantly more patients discontinue high-dose imatinib than dasatinib.

The 100 mg once daily dose for dasatinib in patients with CP CML was approved as a result of data from a phase III dose modification study.11 Shah and colleagues compared dasatinib 100 mg once daily, 50 mg twice daily, 140 mg once daily, and 70 mg twice daily doses in CP CML. In this study, response rates were...
similar between doses, and overall efficacy was equivalent between the 70 mg twice daily and 100 mg once daily regimens. In the 100 mg once daily dose there was significantly less grade 3-4 thrombocytopenia (22%; p=0.004) compared with the 70 mg twice daily dose (37%). Grade 3-4 anemia, leukopenia, and neutropenia were observed in 10%, 16% and 33% of patients, respectively. The 100 mg once daily dose was also associated with the lowest frequencies of treatment discontinuation (overall: 16% vs. 23%; toxicity alone: 4% vs. 11%, respectively).

For nilotinib, the incidence rates of grade 3-4 thrombocytopenia (28%), neutropenia (28%), and anemia (8%) at the recommended dose (800 mg/day) in patients with CP CML appear to be similar to those for dasatinib 100 mg once daily,13,14. The incidences of non-hematologic AEs are much lower than those for hematologic events for all TKIs, and are broadly similar between TKIs at their current recommended doses. Cutaneous toxicity is more common for TKIs against receptor tyrosine kinases.15

Cardiotoxicities observed in chronic myeloid leukemia

Current evidence suggests that TKIs have potential cardiotoxic effects. Cardiac AEs reported include palpitations, arrhythmia, QT prolongation, pericardial effusions, myocardial ischemia, myocardial infarction, and congestive heart failure (CHF). All of the clinically available BCR-ABL inhibitors report the potential for cardiotoxicity in their respective package inserts (Table 1).16,17,18

Imatinib

Although rare, severe CHF and left ventricular dysfunction have been reported during imatinib treatment, especially in patients with risk factors or comorbidities. In the IRIS trial, this was observed in 0.7% of patients.12,18 Cardiotoxicity associated with high-dose imatinib was not reported during the START-R study.19 In smaller studies of high-dose imatinib, existing CHF was exacerbated in one study (6% of patients), and was related to mortality in another (4% of patients).20,21

In a large review of patients enrolled in imatinib clinical studies (n=1,276), 1.8% of patients had symptoms suggestive of systolic heart failure. Most had risk factors for cardiac failure. In total, 0.6% of patients had cardiac events attributed to imatinib treatment. The authors concluded that CHF is rare during imatinib treatment.22 A similar incidence of CHF (1%) was reported in a nine-year retrospective review at a single institution.23 Reports of smaller studies also confirm that cardiac failure is a rare feature of imatinib therapy.24,25

The mechanism underlying imatinib-induced cardiac failure is currently unclear. In an in vitro study, physiological concentrations of imatinib significantly and adversely affect mitochondrial membrane potential, apoptosis, cell viability, and cellular ultrastructure.26 This cardiotoxic effect may be linked to inhibition of BCR-ABL. Imatinib was reported to cause stress-induced and dose-dependent mitochondrial changes in murine ventricular myocytes, which was reduced by re-engineering the imatinib molecule such that BCR-ABL inhibition was hampered.27,28 Nonetheless, the re-engineered molecule may have had altered activities besides reduced BCR-ABL inhibition.

A second cardiac AE associated with imatinib therapy is fluid retention manifesting as pericardial effusion. Grade 3-4 fluid retention reactions, which included pericardial effusions, were reported in 2% of patients in the IRIS study and in 6% of all other CML clinical studies.29 Frank periarditis has been observed in <0.1% of patients receiving imatinib (all indications).30 Other cardiac AEs include tachycardia, hypertension, hypotension, flushing, and peripheral coldness, were each reported in 0.1-1.0% of patients.31

Precautions and general guidelines for dose adjustment for cardiac AEs associated with imatinib treatment are included in the prescribing information and have been summarized in Table 2. CML patients with existing cardiac disease or cardiac risk factors should be monitored and treated accordingly.32,33 Patients should also be weighed regularly and monitored for signs and symptoms of fluid retention. Unexpected weight gain should be investigated carefully, and treated appropriately.34,35 Significant fluid retention (local or general) can usually be managed by interrupting imatinib treatment and using diuretics or other supportive care.36 In severe cases of fluid retention, imatinib should be withheld until this is resolved.

Nilotinib

Nilotinib can cause QT prolongation and sudden death and carries a black box warning for these side effects.37 Nilotinib prolongs the QT interval in a concentration-dependent manner, and the common AEs occurred in 1-10% of all patients in clinical trials.20 In a study in healthy volunteers, nilotinib was associated with a maximum mean QT interval increase of 18 ms (adjusted for placebo).38 In imatinib-resistant patients with CML, nilotinib caused a maximum mean QT change of 10 ms from baseline; QT increase >60 ms and QT >500 ms associated with nilotinib were reported in 2.1% and <1% of patients, respectively. Sudden deaths were reported in 0.6% of patients in a clinical study, and at a similar frequency in an expanded access study. The early occurrences of some of these deaths relative to the start of nilotinib treatment suggest that ventricular repolarization may have contributed to their occurrence.19

Another common cardiac AE associated with nilotinib therapy is myocardial ischemia. In the pivotal phase II study, it was reported in 7% (21/321) of patients with CP CML receiving nilotinib as second-line treatment.31 Uncommon cardiac events (0.1-1% of all patients in clinical trials) include cardiac failure, angina pectoris, atrial fibrillation, pericardial effusion, coronary artery disease, cardiomegaly, cardiac murmur, and bradycardia. Rare events (of uncertain frequency) include myocardial infarction, ventricular dysfunction, pericarditis, cardiac flutter, and extrasystoles.18

The mechanisms underlying these AEs are still unclear. In a manner similar to imatinib, nilotinib was found to decrease the cellular viability of rat cardiomyocytes cultured in vitro, although the integrity of the mitochondrial membrane potential was unaffected.32 However, nilotinib has also been found to inhibit human ether-a-go-go related gene (hERG) potassium currents with an IC50 of 0.66 μM. This concentration is approximately one-tenth the expected Cmax for this compound, well within therapeutic levels. This mechanism is likely to underlie nilotinib-induced QT prolongation. Inhibition of hERG channels is established as a cause of QT prolongation for a number of compounds, and is a significant barrier in the development of new drugs.32 Indeed, the phase II development of the aurora kinase inhibitor MK-0457 (VX-680) was recently suspended, pending a full analysis of all efficacy and safety data. The decision was based on preliminary safety data, in which QT prolongation was observed in one patient.33

The potential for QT prolongation and sudden death associated with nilotinib, although rare, necessitates vigilant monitoring. In particular, ECGs should be performed at baseline, seven days after initiation of treatment, periodically throughout therapy, and following dose adjustments. Electrolyte levels should be monitored periodically throughout therapy. Nilotinib is contraindicated for patients with hypokalemia, hypomagnesemia or long QT syndrome.33 The nilotinib prescribing information recommends dose adjustments for QT prolongation, presented in Table 2.

Dasatinib

The events of dasatinib-induced QT prolongation are rare although a warning for such a possible event is given. In single-arm studies of dasatinib, nine patients (1%) had QT prolongation reported as an AE.2 The mean QT
interval increased by 3-6 ms (Fridercia’s method); this increase was not clinically relevant.\(^3\)\(^4\) In total, <1% of patients had a QT increase to >500 ms. In contrast with nilotinib, the IC\(^{50}\) for dasatinib for the inhibition of hERG currents (14.3 \(\mu\)M) is 100 times the expected C\textsubscript{max} for this drug.\(^2\)\(^7\) This may explain why QT prolongation is more clinically prominent for nilotinib than it is for dasatinib.

Common cardiac AEs (observed in 1 – < 10% of all patients in clinical trials) include arrhythmia and palpitations. Severe pericardial effusions have been reported in 1% of all patients in all clinical studies, and the prescribing information for dasatinib includes a warning for this toxicity.\(^2\)\(^7\) Severe CHF has also been reported in 1% of all patients.\(^2\)\(^7\) In single-arm studies, CHF or ventricular dysfunction occurred in 4% (20/911) of patients.\(^3\)\(^4\) However, in the dose optimization study, dasatinib 100 mg once daily was not associated with any incidence of severe CHF or pericardial effusion in any patient; both AEs were reported in patients receiving the 70 mg twice daily dose (all grades, 4%; grade 3-4, 3%).\(^2\)\(^7\)

Both pericardial effusions and cardiac failure associated with dasatinib therapy may be caused by similar mechanisms to those associated with imatinib therapy, although, in contrast with imatinib, dasatinib has not been found to significantly affect mitochondrial membrane potential, apoptosis, cell viability, or cellular ultrastructure at physiological con-

### Table 1. Reported incidence of cardiotoxicity during tyrosine kinase inhibitors treatment.

| Toxicity                        | \(n\)  | Incidence (%) | Reference |
|--------------------------------|-------|---------------|-----------|
| **Imatinib**                    |       |               |           |
| Pericardial effusions           | NR    | 6             | Novartis\(^{23}\) |
| Systolic heart failure          | 1276  | 1.8           | Atallah \textit{et al.}\(^{27}\) |
| Congestive heart failure        | 553   | 0.7           | Novartis\(^{23}\) |
| Left ventricular dysfunction    | 553   | 0.7           | Novartis\(^{23}\) |
| Cardiac failure                 | NR    | 0.1-1.0       | Novartis\(^{23}\) |
| Flushing                        | NR    | 0.1-1.0       | Novartis\(^{23}\) |
| Hypotension                     | NR    | 0.1-1.0       | Novartis\(^{23}\) |
| Hypertension                    | NR    | 0.1-1.0       | Novartis\(^{23}\) |
| Peripherial coldness            | NR    | 0.1-1.0       | Novartis\(^{23}\) |
| Tachycardia                     | NR    | 0.1-1.0       | Novartis\(^{23}\) |
| Pericarditis                    | NR    | <0.1          | Novartis\(^{23}\) |
| **Dasatinib**                   |       |               |           |
| Severe pericardial effusions    | NR    | 1             | BMS\(^{13}\) |
| Congestive heart failure        | 911   | 4             | Brave \textit{et al.}\(^{34}\) |
| Arrhythmia                      | NR    | 1-<10         | BMS\(^{13}\) |
| Palpitations                    | NR    | 1-<10         | BMS\(^{13}\) |
| QT prolongation >500 ms         | -300  | <1            | BMS\(^{13}\) |
| Angina pectoris                 | NR    | 0.1-<1        | BMS\(^{13}\) |
| Cardiomegaly                    | NR    | 0.1-<1        | BMS\(^{13}\) |
| Myocardial infarction           | NR    | 0.1-<1        | BMS\(^{13}\) |
| Pericarditis                    | NR    | 0.1-<1        | BMS\(^{13}\) |
| Ventricular arrhythmia          | NR    | 0.1-<1        | BMS\(^{13}\) |
| Acute coronary syndrome         | NR    | <0.1          | BMS\(^{13}\) |
| Myocarditis                     | NR    | <0.1          | BMS\(^{13}\) |
| **Nilotinib**                   |       |               |           |
| Myocardial ischemia             | 321   | 7             | Kantarjian \textit{et al.}\(^{31}\) |
| Increase in QTC \(>60\) ms from BL | 232  | 2.1           | Novartis\(^{18}\) |
| Palpitations                    | NR    | 1-10          | Novartis\(^{18}\) |
| Angina pectoris                 | NR    | 0.1-1         | Novartis\(^{18}\) |
| Atrial fibrillation             | NR    | 0.1-1         | Novartis\(^{18}\) |
| Bradycardia                     | NR    | 0.1-1         | Novartis\(^{18}\) |
| Cardiac failure                 | NR    | 0.1-1         | Novartis\(^{18}\) |
| Cardiac murmur                  | NR    | 0.1-1         | Novartis\(^{18}\) |
| Cardiomegaly                    | NR    | 0.1-1         | Novartis\(^{18}\) |
| Coronary artery disease         | NR    | 0.1-1         | Novartis\(^{18}\) |
| Pericardial effusion            | NR    | 0.1-1         | Novartis\(^{18}\) |
| Death\(^{a}\)                   | 867   | 0.6           | Novartis\(^{18}\) |

\(\text{BL, baseline}; \text{NR, not reported, representing all patients within each agent’s clinical trials.} \text{Possibly resulting from ventricular repolarization.}\)
Article

Table 2. Precautions and dose modifications on the emergence of cardiac events during tyrosine kinase inhibitors treatment.

| Agent    | Dose modification                                           | Precautions                                                                 |
|----------|-------------------------------------------------------------|------------------------------------------------------------------------------|
| Imatinib | On the emergence of a severe event, withhold until the event has resolved. Resume at an appropriate dose, depending on initial severity of event. | Carefully monitor any patient with or at risk of cardiac failure. All patients with signs or symptoms of cardiac failure should be evaluated and treated. |
| Dasatinib| On the emergence of a severe event, withhold until the event has resolved or improved. Resume at an appropriate dose, depending on initial severity of event. | Administer with caution to patients with or at risk of QTc prolongation: those with hypokalemia, hypomagnesemia, or congenital long QT syndrome; or taking medicines known to prolong QT, including anti-arrhythmic drugs, and cumulative high-dose antiarrhythmic therapy. Correct hypokalemia or hypomagnesemia prior to administration. |
| Nilotinib| QTc > 480 ms: 1) withhold therapy; correct serum potassium and magnesium levels if below normal; and review concomitant medication; 2) resume at prior dose in <2 weeks if QTc returns to <450 ms and <20 ms of baseline; 3) reduce dose to 400 mg/day if QTc 450-480 ms after 2 weeks; 4) discontinue if QTc returns to >480 ms after dose reduction; 5) repeat ECG assessment approx. 7 days after any dose adjustment. | Do not administer to patients with long QT syndrome. Do not administer drugs known to prolong QT, and strong CYP3A4 inhibitors. Correct hypokalemia or hypomagnesemia prior to administration, and periodically monitor serum electrolyte levels during therapy. Perform ECGs at baseline, 7 days after treatment starts, periodically as indicated clinically, and after any dose adjustment. |

Conclusions

Although rare, severe CHF and left ventricular dysfunction have occurred with imatinib treatment, especially in patients receiving higher doses. Cardiac failure is also an uncommon feature of nilotinib and dasatinib therapy. However, CHF has not yet been reported in patients with CP CML receiving dasatinib with a starting dose of 100 mg once daily. Compared with imatinib, palpitations and prolongation of the QT interval are relatively common with both dasatinib and nilotinib. Nilotinib carries a black box warning for such AE and sudden death has been observed. These AEs may be related to BCR-ABL inhibition and therefore be genuine class effects. Further research is needed to clarify the mechanisms underlying these effects.

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