Ristocetin-induced platelet aggregation for monitoring of bleeding tendency in CLL treated with ibrutinib

L Kazianka1,2, C Drucker3, C Skrabs1,2, W Thomas4, T Melchardt5, S Struve6, M Bergmann6, PB Staber1,2, E Porpacz1,2, C Einberger1,2, M Heinz1,2, A Hauswirth1,2, M Raderer2,7, I Pabinger1,2, R Thalhammer8, A Egle5, C-M Wendtner6, G Follows4, G Hoermann8, P Quehenberger8, B Jilma3 and U Jaeger1,2

Bleeding because of impaired platelet function is a major side effect of the Bruton’s tyrosine kinase (BTK) inhibitor ibrutinib. We quantitatively assessed ristocetin-induced platelet aggregation (RIPA) in 64 patients with chronic lymphocytic leukemia (CLL) under ibrutinib at 287 time points. Eighty-seven bleeding episodes in 39 patients were registered (85 Common Toxicity Criteria (CTC) grade 1 or 2, 2 CTC grade 3) during a median observation period of 10.9 months. At times of bleeding, RIPA values were significantly lower (14 vs 28 U; \( P < 0.0001 \)). RIPA was impaired in patients receiving concomitant antiplatelet therapy or anticoagulation (14 vs 25 U, \( P = 0.005 \)). A gradual decline of median RIPA values was observed with increasing bleeding severity. Importantly, no CTC grade 2 or 3 bleeding were observed with RIPA values of > 36 U. Sequential monitoring indicated a decrease of RIPA values from a median of 17 to 9 U within 2 weeks after initiation of treatment as well as an increase above the critical threshold of 36 U within 7 days when ibrutinib was paused. Low RIPA values were similar during treatment with another BTK inhibitor, CC292. Quantitative assessment of platelet function is a practical tool to monitor bleeding tendency under BTK-inhibitor therapy.

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INTRODUCTION

Targeting Bruton’s tyrosine kinase (BTK) with the small-molecule ibrutinib has significantly improved progression-free and overall survival of patients with chronic lymphocytic leukemia (CLL) and mantle cell lymphoma and is currently investigated in other B-cell lymphomas.1–5 The drug is now widely used in clinical routine. Its mechanism of action is based on the inhibition of BTK as part of the B-cell receptor signaling cascade.6–10 BTK is also expressed in other hematopoietic lineages, including platelets.11–13 Inhibition of this kinase may therefore result in relevant hematologic side effects. Indeed, bleeding episodes were observed in 44% of patients in the CLL registration trial1 and in up to 61% of patients after a longer observation period.14

Bleeding is usually mild (Common Toxicity Criteria (CTC) grades 1–2, corresponding to spontaneous bruising or petechiae), but more severe bleeding is observed in 8% of patients.14 CTC grade 3 or 4 bleeding occurs in ~5% of patients after trauma.1,4,15

BTK is a Tec family kinase and acts as a signaling molecule in von Willebrand factor (vWF)- and collagen-induced platelet activation.12 The corresponding receptors glycoprotein VI (GPVI, collagen receptor) and the glycoprotein Ib-IX-V complex (GPIb, vWF receptor) have previously been shown to be affected by BTK inhibition: in vitro binding of platelets from ibrutinib-treated patients to vWF matrix was significantly impaired16 as well as platelet response to collagen using light transmission aggregometry.11

Of note, vWF antigen and activity in plasma remain within normal range.17 The dependency of GPIb-IX-V signaling on BTK has also been shown in vivo using a BTK-deficient mouse model.13 However, there is a possibility that TEC, another member of the Tec kinase family, compensates for the loss of BTK in platelets as it is indicated by the partially preserved response to collagen in human X-linked agammaglobulinemia patients.18 Nevertheless, the cysteine residue that is required for the covalent binding of ibrutinib to an adenosine triphosphate-binding site can also be found in TEC.13 Taken together, there is strong evidence that pharmacologic inhibition of BTK and other Tec kinases by ibrutinib results in impaired platelet function.

Proper management of bleeding complications is important because they may affect the length and intensity of ibrutinib treatment. In addition, up to 50% of CLL patients with cardiovascular comorbidities are under concomitant treatment with anticoagulants or platelet function inhibitors.19,20

We used an easy to perform method for quantitative assessment of vWF/ristocetin-induced platelet aggregation (RIPA) in ibrutinib-treated CLL patients in order to (1) correlate the degree of impairment of platelet function with bleeding tendency and (2) monitor platelet aggregation during drug therapy and before interventions.

SUBJECTS AND METHODS

Subjects

This observational study included 64 CLL patients from 4 different centers in 3 countries (Cambridge, Munich, Salzburg and Vienna). Patients received...
Ibrutinib at a target dose of 420 mg daily and were either included in the RESONATE study (28 patients), in the ibritunib named patient program (12 patients) or received ibritunib in routine clinical use according to the licensed indication in Europe (24 patients). Of the patients, 18% were previously untreated but had a 17p deletion or TP53 mutation, whereas 82% had relapsed or refractory CLL with or without 17p/TP53 aberrations. The study was approved by the ethics committee of the Medical University of Vienna (ethics committee Nr 1631/2012 and 11/2005) and informed consent was obtained. Eleven patients had multiple measurements before start and during ibritunib therapy and 53 patients were investigated under stable ibritunib intake.

Patients were seen regularly (at least monthly) in their corresponding centers. Bleeding tendency was recorded at each time point by standardized questions including bruising, petechiae and epistaxis, and a physical examination was performed. Clinical information included clinical stage, genetics, lines of therapy and concomitant treatment, particularly therapy with antiplatelet agents or anticoagulants. Patient characteristics are shown in Table 1. Severity of bleeding was scored according to the CTC grading scale (version 4.03: 14 June 2010) used in the RESONATE protocol.

Controls included 13 CLL patients currently not on treatment, 11 patients receiving other treatments for CLL (4 with immunochemotherapy (rituximab plus fludarabine/cyclophosphamide or bendamustine) and 7 with phosphatidylinositol-3-kinase inhibitors (idelalisib, duvelisib)) as well as 1 patient with X-linked agammaglobulinemia and proven BTK protein deficiency.

Platelet function assessments

Fresh blood specimens (5 ml) for analysis of platelet function were drawn into hirudine containing tubes (Vacuette, Krefelsanuender, Austria) and analyzed within 3 h in a Multiplate Analyzer (Roche, Vienna, Austria) software. Statistical methods comprised χ² analysis or Fisher’s exact test to compare baseline characteristics, and t-test or analysis of variance where applicable for RIPA analysis. Correlations were determined by Pearson’s r. The P-values of < 0.05 (two sided) were considered statistically significant.

RESULTS

Patient characteristics and bleeding events

The study cohort consisted of a typical CLL cohort treated with ibritunib with a median age of 71 (range: 40–92) years, 2 (range: 0–6) prior treatment lines and aberrations of TP53 (17p deletion and/or TP53 mutation) in 47.6%. The median observation period under ibritunib was 10.9 months. Importantly, 28.1% were treated with concomitant antiplatelet (acetyl salicylic acid, clopidogrel, 18.8%) or anticoagulation therapy (coumarin, heparin, novel direct oral anticoagulants, 10.9%) or both (in 1 case).

At least one bleeding event was recorded in 39 of the 64 patients (60.9%; Table 1). Bleeding was mild in most patients (CTC grade 1 or 2) and only two CTC grade 3 bleedings were observed. The total number of bleeding episodes was 87, again predominantly grade 1 or 2. None of the patients had a grade 4 or 5 bleeding event. Of note, 60 of the 64 patients (93.8%) were still responding and on ibritunib after 10.9 months. Reasons for discontinuation or pause are shown in Table 1.

Bleeding tendency, platelet number and anticoagulation

The occurrence of bleeding events under ibritunib was significantly associated with lower platelet counts (116.5 G/l, range: 38–303 vs 137 G/l, range: 51–328; P = 0.0012; Supplementary Figure 1). Patients receiving antiplatelet therapy or anticoagulation (13/18, 72%) were more frequently affected than patients without concomitant treatment (26/46; 57%, P = 0.25). Patients with anticoagulation had the highest frequency of bleeding (6 of 7 patients; Table 2). However, this was not statistically significant.

Impairment of platelet aggregation during ibritunib treatment

In total, 287 quantitative assessments were performed by RIPA (median: 3 per patient). CLL patients without current treatment (median 57 U, range: 8–111) or during immunochemotherapy (median 68.5 U, range: 27–94) had RIPA values close to normal, whereas patients under ibritunib had considerably impaired RIPA values, although with a wide range (median 19.5 U, range: 0–199; Supplementary Figure 2).
The major goal of this study was to establish an association between bleeding and platelet function during ibrutinib treatment. The median RIPA measured at times of bleeding events under ibrutinib was significantly impaired (median 14 U, range: 0–76) compared with time points when no bleeding was observed (median 28 U, range: 0–199; P < 0.0001; Figure 1a). Collagen-induced platelet aggregation testing confirmed the findings obtained by RIPA measurements. Platelet function showed significantly lower values at time points when bleeding occurred (P = 0.0015; Supplementary Figure 3). We confirmed that RIPA values with this assay correlate with platelet numbers (P < 0.0001; Supplementary Figure 4). However, low RIPA values were associated with bleeding regardless of platelet count (Figure 1b).

**Bleeding tendency and platelet function**

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**Effect of concomitant anticoagulation**

RIPA was significantly impaired in patients receiving concomitant antiplatelet and/or anticoagulant medication. Ibrutinib-treated CLL patients under concomitant antiplatelet and/or anticoagulant therapy showed reduced RIPA values (median 14 U, range: 0–118 U) when compared with patients without concomitant medication (median 25 U, range: 0–199 U; P = 0.005).

**Bleeding severity and platelet function**

We next analyzed platelet function in relation to the severity of the event expressed by CTC grade: a steady decline of median RIPA values was observed from time points without bleeding (28 U) to CTC grade 1 (14.5 U), grade 2 (12 U) and grade 3 (8.5 U) events (Figure 3). Importantly, no CTC grade 2 or 3 bleeding was observed with RIPA values of >36 U. These data indicate that impairment of platelet function is associated with severity of bleeding and that significant bleeding does not occur beyond a defined threshold.

**Monitoring of platelet function during ibrutinib therapy**

Consecutive samples before and after start of ibrutinib were available in 11 patients. A decline of platelet aggregation from 17 to

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**Table 2. Characteristics of bleeding and nonbleeding patients**

| Characteristic                  | Bleeding | No bleeding | P-value |
|---------------------------------|----------|-------------|---------|
| Total number (n = 64)           | 39/64 (60.9%) | 25/64 (39.1%) | NA |
| Median body weight, kg (range)  | 74 (48–108) | 69.5 (44–114) | 0.49 |
| Male, %                         | 64.1%    | 64%         | 0.99 |
| Antiplatelet or anticoagulant Tx| 13/39 (33.3%) | 5/25 (20%) | 0.25 |
| Antiplatelet therapy            | 8/39 (20.5%) | 4/25 (16%) | 0.65 |
| Oral AC/heparin                 | 6/39 (15.4%) | 1/25 (4%) | 0.15 |
| Median platelet count, G/l (range) | 116.5 (38–303) | 137 (51–328) | 0.0012 |
| Median hemoglobin, mg/dl (range) | 12.6 (9.2–16.7) | 12.1 (8.3–17.1) | 0.16 |
| Median WBC, G/l (range)         | 21.8 (4.3–347.8) | 16.0 (2.2–397) | 0.41 |

Abbreviations: AC, anticoagulant; NA, not available; Tx, treatment; WBC, white blood cell count. Bold value indicates significant P-value.

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**Figure 1.** RIPA measurements in CLL patients during stable ibrutinib therapy. (a) RIPA measurements were significantly lower at the time when bleeding-related adverse events occurred (median 14 vs 28 U, P < 0.0001). (b) When patients were grouped by platelet count, RIPA results remained impaired at the time of bleeding-related adverse events (P = 0.003 and P < 0.0001, respectively).

**Figure 2.** RIPA in ibrutinib-treated CLL patients with concomitant antiplatelet and/or anticoagulant medication. Ibrutinib-treated CLL patients under concomitant antiplatelet and/or anticoagulant therapy showed reduced RIPA values (median 14 U, range: 0–118 U) when compared with patients without concomitant medication (median 25 U, range: 0–199 U; P = 0.005).

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9 U was observed when therapy was initiated \((P = 0.019; \text{Figure 4})\). The median interval between measurements was 13 days. When ibrutinib was paused or stopped (9 instances) the median RIPA values increased from 12 to 54 U \((P = 0.004; \text{median interval 18 days; \text{Figure 5a}})\). Short-term RIPA kinetics were available in 5 patients \((\text{Figure 5b})\): some variability of platelet function recovery was observed with 4 of 5 patients crossing the imaginary threshold of 36 U within 7 days, whereas recovery was slow in 1 patient. These data indicate that quantitative assessment of RIPA may be used to monitor platelet function, for example, in the context of planned surgical interventions.

Platelet function with higher doses of ibrutinib and other BTK inhibitors

We also assessed RIPA in patients receiving higher doses of ibrutinib (560 mg daily) in mantle cell lymphoma patients \((n = 7)\). RIPA values were comparable to those in CLL patients \((\text{Figure 6})\). Ibrutinib also inhibits other kinases including TEC kinases that are also involved in the regulation of platelet function. Interestingly, patients under treatment with a different BTK inhibitor, CC292 \((n = 6)\), showed almost identical RIPA values \((19.5 \text{ vs 23 U})\). Two different measurements in a patient with a hereditary BTK deficiency \((\text{X-linked agammaglobulinemia})\) showed less impairment of platelet function.

DISCUSSION

Ibrutinib has become an integral part of the CLL treatment algorithm. The drug is well tolerated, but bleeding is a frequent side effect.\(^{14}\) Here we show that quantitative assessment of RIPA is...
a practical tool to monitor and manage bleeding tendency, and may thereby increase the safety of ibrutinib.

The theoretical basis for the impairment of platelet function has been well described by others13,16,18 (Supplementary Figure 6). BTK is an important component of the signaling cascades downstream of the GPIb and GPVI platelet receptors.3,18 Therefore, ristocetin- as well as collagen-induced platelet aggregation are impaired and could both be used as diagnostic tools.16 Unfortunately, assessment of platelet function by light transmission aggregometry has turned out to be tedious, variable and operator dependent. Here we used an easy to perform whole blood aggregometry method that is frequently used in clinical routine. We chose the RIPA assay for our investigations because of its slightly better discriminatory power in preliminary experiments. However, similar results were obtained with collagen-induced platelet aggregation (Supplementary Figure 3). There are some uncertainties as to the contribution of BTK in comparison with other Tec kinases in impairing platelet function as ibrutinib inhibits several other kinases. However, the data obtained with a different BTK inhibitor, CC292, argue for a strong involvement of BTK in platelet function. It remains to be established whether this will translate into clinical bleeding tendency when other selective BTK inhibitors enter the clinic.25,26

The major clinical finding of this study is that low RIPA values are strongly associated with bleeding tendency. This is supported by the following observations: (1) a decline in platelet aggregation was observed when ibrutinib is initiated; (2) platelet function was significantly impaired in CLL patients under stable ibrutinib therapy compared with CLL patients under immunochemotherapy or without current intervention; and (3) at times when bleeding events occurred, RIPA values were significantly lower (14 vs 28 U). This also is in line with previous observations made by other groups.11,16,17 Bleeding under ibrutinib is usually mild, but can be severe in 5 to 8% of patients.14,16 A practical clinical implication of our study is that RIPA values decrease with the degree of bleeding. Thus, it seems possible to define a threshold (36 U in this study) above which no major bleeding events (CTC grade ≥ 3) will occur. This will allow monitoring of bleeding tendency with RIPA tests before elective surgery or in cases of emergency, where platelet transfusions may be indicated.16 We note that the recommended washout period of 7 days before surgical interventions correlated well with the increase in RIPA above the threshold level of 36 U in most patients. However, this may be different in some patients in whom monitoring will add important information for clinical decisions.

We observed an association of platelet number with bleeding tendency confirming previous observations (Supplementary Figure 1), indicating that bleeding events were more frequent during the initial treatment phase of CLL patients, when platelets are still low.14 In addition, there was a strong correlation between RIPA and platelet number in the assay used for this study, as reported during the evaluation of the test (Supplementary Figure 4).23 However, the association of low RIPA values and bleeding under ibrutinib was still significant regardless of platelet counts (Figure 1b). In addition, the median platelet values were quite high with a median of 116.5 G/l in bleeding patients. Increased bleeding tendency with concomitant antiplatelet or anticoagulation therapy has been reported and has led to the recommendation that warfarin should not be used together with ibrutinib.10 There was a trend for, but no significant association of, concomitant antiplatelet or anticoagulation treatment with bleeding in our study. This is probably because of low patient numbers. However, in our study low RIPA values were associated with both antiplatelet and anticoagulation therapy indicating a higher propensity for bleeding.

There are some limitations to this study. The RIPA impedance test has to be performed on site within a limited time frame with fresh samples. Frozen or shipped samples cannot be used. On the other hand, the analyzer used is now widely available in larger hospitals. In addition, similar and reliable results could be obtained in 4 different centers in 3 countries (median RIPA values were 26 U (range: 0–199; Vienna), 14 U (range: 1–148; Cambridge), 16 U (range: 2–76; Munich) and 25 U (range: 3–68; Salzburg)). Measurements under stable ibrutinib intake on consecutive visits showed little variability (Supplementary Figure 7). There was no significant statistical difference between centers. The test has a reasonably low intrasubject variation (coefficient of variation < 10% in healthy donors and < 16% in patients under ibrutinib) confirming its clinical reliability. The threshold for major bleeding events is currently only based on 64 CLL patients and 287 measurements. A more detailed and formal evaluation such as for other CLL diagnostic tests will be required for robust generalized diagnostics.27,28 Reporting of bleeding events is dependent on thorough clinical evaluation of patients at any visit. We have therefore developed a questionnaire that can be used in the future. Last but not the least, further confirmation in larger studies is still needed.

It is important to note that 93.8% of our CLL patients were still responding to ibrutinib after 10.9 months and many will remain on the drug for much longer. Monitoring of bleeding tendency with platelet function tests may therefore contribute to enhance safety, adherence and the successful long-term use of ibrutinib.

CONFLICT OF INTEREST

MB, PS, AE, CW-M and UJ received honoraria from Janssen, LC, CD, CS, WT, TM, SS, EP, CE, MH, AH, MR, IP, RT, GH, PQ and BJ declare no conflict of interest.

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