Single cell immune profiling by mass cytometry of newly diagnosed chronic phase chronic myeloid leukemia treated with nilotinib

Stein-Erik Gullaksen,1 Jørn Skavløn,1 Sonia Gavasso,2 Vinko Tosevski,4 Krzysztof Warzocha,1 Claudia Dumrese,6 Augustin Ferrant,7 Tobias Gedde-Dahl,1 Andrzej Hellmann,9 Jeroen Janssen,10 Boris Labar,11 Alois Lang,12 Valeed Majeed,13 Georgi Mihaylov,14 Jesper Stenfort,15 Lief Stenke,1 Andreas Hochhaus,20 Leif Stenke,16 Josef Thaler,17 Noortje Thielen,10 Andreas Hochhaus,20 Henrik Hjorth-Hansen,21,22 Satu Mustjoki,23 Sieghart Sopper,26 Francis Giles,28 Kimmo Porkka,25 Dominik Wolf25,27 and Bjørn Tore Gjertsen28

1Centre of Cancer Biomarkers CCBIO, Department of Clinical Science, Precision Oncology Research Group, University of Bergen, Norway; 2Department of Clinical Medicine, University of Bergen, Norway; 3Neuroimmunology Lab, Haukeland University Hospital, Bergen, Norway; 4Mass Cytometry Facility, University of Zurich, Switzerland; 5Department of Hematology, Institute of Hematology and Transfusion Medicine, Warsaw, Poland; 6Flow Cytometry Facility, University of Zurich, Switzerland; 7Hematology Department, Cliniques Universitaires St Luc, Brussels, Belgium; 8Department of Medicine, Oslo University Hospital, Norway; 9Department of Hematology, Medical University of Gdańsk, Poland; 10Department of Hematology, VU University Medical Center, Amsterdam, the Netherlands; 11Department of Hematology, University Hospital Center Rebro, Zagreb, Croatia; 12Internal Medicine, Hospital Feldkirch, Austria; 13Department of Hemato-Oncology, Stavanger University Hospital, Norway; 14Clinic for Hematology, University Hospital Sofia, Bulgaria; 15Hematology Unit, Aarhus University Hospital, Denmark; 16Department of Medicine, Karolinska University Hospital, Stockholm, Sweden; 17Department of Internal Medicine IV, Wels-Grieskirchen Hospital, Wels, Austria; 18Department of Hematology, University Hospital Leuven, Belgium; 19Department of Internal Medicine – Hematology, University Hospital Hradec Kralove, Czech Republic; 20Department of Hematology and Medical Oncology, Universitätsklinikum Jena, Germany; 21Department of Hematology, St Olavs Hospital, Trondheim, Norway; 22IKM, NTNU, Trondheim, Norway; 23Hematology Research Unit Helsinki, University of Helsinki and Helsinki University Hospital Comprehensive Cancer Center, Department of Hematology, Finland; 24Department of Clinical Chemistry, University of Helsinki, Finland; 25Department of Hematology and Oncology, Innsbruck Medical University and Tyrolean Cancer Research Institute, Innsbruck, Austria; 26NMDTI, Robert H. Lurie Comprehensive Cancer Center of Northwestern University, Chicago, IL, USA; 27Medical Clinic 3, Oncology, Hematology and Rheumatology, University Hospital Bonn (UKB), Germany and 28Department of Internal Medicine, Haukeland University Hospital, Bergen, Norway

ABSTRACT

Monitoring of single cell signal transduction in leukemic cellular subsets has been proposed to provide deeper understanding of disease biology and prognosis, but has so far not been tested in a clinical trial of targeted therapy. We developed a complete mass cytometry analysis pipeline for characterization of intracellular signal transduction patterns in the major leukocyte subsets of chronic phase chronic myeloid leukemia. Changes in phosphorylated Bcr-Abl1 and the signaling pathways involved were readily identifiable in peripheral blood single cells already within three hours of the patient receiving oral nilotinib. The signal transduction profiles of healthy donors were clearly distinct from those of the patients at diagnosis. Furthermore, using principal component analysis, we could show that phosphorylated transcription factors STAT3 (Y705) and CREB (S133) within seven days reflected BCR-ABL1 at three and six months. Analyses of peripheral blood cells longitudinally collected from patients in the ENEST1st clinical trial showed that single cell mass cytometry appears to be highly suitable for future investigations addressing tyrosine kinase inhibitor dosing and effect. (clinicaltrials.gov identifier: NCT061177)
Introduction

Chronic myeloid leukemia (CML) is a hematologic stem cell disorder driven by transcription of the fusion protein Bcr-Abl1, a constitutively active tyrosine kinase. The deregulated kinase activity is sufficient to maintain the CML phenotype and leads to activation of several pathways, including JAK/STAT, PI3K/Akt and Ras/MAPK. Accordingly, specific tyrosine kinase inhibitors (TKIs) targeting Bcr-Abl1 have revolutionized CML treatment. The majority of patients display excellent cytogenetic and molecular responses and remain free of disease progression. However, for many CML patients, a cure will probably not be achieved by TKIs alone because of the persistence of leukemic stem cells that may cause relapse. Furthermore, a significant number of patients are switched from one TKI to another because of insufficient efficacy or adverse effects. Early identification of suboptimal responders to TKI could facilitate optimal dosing and choice of TKI thereby limiting adverse events and securing optimal molecular responses. This could be accomplished by the direct assessment of Bcr-Abl1 kinase activity in the leukemic cells early after start of TKI therapy.

Pharmacodynamic analysis of TKI effect on Bcr-Abl1 tyrosine kinase activity and subsequent downstream signal transduction in CML patients has been technically challenging and thus rarely exploited in clinical trials. However, recent advances in single cell analysis of intracellular signal transduction show promise both in diagnostics and in therapeutic leukemia monitoring. Cytometry by Time of Flight (CyTOF) allows single cell characterization of immunophenotypes and intracellular signaling status in healthy individuals and patients with hematologic malignancies. The technique enables measurement of more than 40 parameters per cell, thereby dramatically increasing the dimensionality of acquired data compared to conventional flow cytometry. This is achieved by exploiting the resolution of mass spectrometry and stable rare-earth isotope-conjugated antibodies to perform highly multiplexed assays on single cells. This has permitted the dynamic signal transduction events within the hematologic hierarchy in acute myeloid leukemia to be evaluated.

A panel of antibodies targeting more than 30 cell-surface and intracellular phospho-specific epitopes covering the major signaling molecules in Bcr-Abl1 oncogenic signaling (pBCR Y177, pAb1 Y245, pCRKL Y207, pSTAT1 Y701, pSTAT3 Y705, pSTAT5 Y694, pCREB S133, pERK 1/2 T202/Y204, pS6 ribosomal protein S235/36 and S240/44) were developed. By analyzing longitudinally collected samples we show that both TKI-mediated changes in signal transduction and immunophenotypes is measurable, and propose that a 1-tube-based analysis by mass cytometry is a feasible biomarker strategy in clinical trials of leukemia.

Methods

Patients

Patients were eligible for the present study if they were enrolled in the international phase IIIb ENESt1tst study. Patients were enrolled when they had newly diagnosed chronic phase CML, were 18 years of age or older, and had a World Health Organization (WHO) performance status no higher than 2. The demographic and clinical parameters of all patients included in this study (n=17) are shown in Online Supplementary Tables S1 and S2. Treatment consisted of nilotinib 300 mg BID. A peripheral blood (PB) sample was collected before the first nilotinib dose, and three hours, seven days and 28 days after the start of treatment. The study was performed in accordance with the Declaration of Helsinki, and all patients provided written informed consent. The study was approved by the local institutional review boards of all participating centers and is registered at clinicaltrials.gov identifier: 01061177. PB and bone marrow (BM) were collected from healthy individuals after written informed consent (University of Bergen, Norway, local ethical committee approval 2012/1045).

Materials

The primary material was fixed and erythrocytes lysed by Lyse/Fix buffer (BD Phosflow) in local hospital laboratories and immediately frozen in saline for long-term storage and shipping at -80°C. The longitudinal PB samples from each patient (before first dose of nilotinib, and after 8 hours, 7 days, and 28 days) were barcoded for identification, pooled and stained with the antibody panels (Online Supplementary Table S3) according to the manufacturer’s recommendation (Maxpar Phospho-Protein Staining Protocol). In the preliminary dataset #1, a protocol for barcoding was adapted from Zunder et al. (Online Supplementary Figure S1) and barcoded reagents were a kind gift from Prof. Bernd Bodenmuller, University of Zurich, Switzerland. The acquisition of samples was performed on the CyTOF2 mass cytomter (Fluidigm) at the Mass Cytometry Facility, University of Zurich, Switzerland. In dataset #2, a commercially available barcoding kit was used (Fluidigm). Acquisition of this dataset was performed using the Helios mass cytomter (Fluidigm) at the Flow Cytometry Core Facility, University of Bergen, Norway.

Statistical analysis

Friedman non-parametric test with Dunn’s multiple comparison was used to find statistically significant changes in phosphorylation of intracellular targets (before, and after 3 hours and 7 days of treatment). Wilcoxon matched-pairs signed rank test was used to identify statistically significant changes in cell abundances before and after initiation of nilotinib therapy. Prism software (GraphPad Software Inc.) was used to calculate P-values and plot the graphs. P<0.05 was considered statistically significant.

Results

The immunophenotype and signal transduction in CD34+ cells of CML patients

It has been proposed that both number of hematopoietic stem and progenitor cells (CD34+CD38low and CD34+CD38high, respectively), and their signaling response patterns to TKIs are important for response prediction in CML. Therefore, we performed an initial analysis of the PB CD34+ compartment before and after receiving the first dose (3 hours and 7 days) of nilotinib in 17 patients enrolled in the ENESt1st study. The 3 samples from each patient were barcoded, pooled and stained with a validated panel of antibodies (Online Supplementary Figures S2 and S3). The high dimensional single-cell data for each patient were clustered using the SPADE algorithm, where manual annotation identified several major hematologic cell subsets, including the CD34+ cells. This population was further gated manually into CD34+CD38low and CD34+CD38high cells. Due to inter-patient variation in CD38 expression range, the gating was tailored to each
longitudinal patient sample set. Only patients for whom a minimum of 100 cells could be identified as CD34⁺CD38⁻low and CD34⁺CD38⁻high for all samples were included in the final analysis (n=10). To characterize the debulking of the tumor load at the single cell level and over time, all CD34⁺ cells were analyzed using viSNE algorithm.²¹ This allowed single cells from different samples to be graphed in a unifying 2-dimensional plot, where its (x,y) position contains information about cell surface marker expression (see Figure 1A for 2 representative patients’ sample sets). CD34⁺CD38⁻low cells resided in the lower part of the viSNE plot, while CD34⁺CD38⁻high cells resided in the middle/top right. It has been suggested that CD25 can be used to identify leukemic stem cells in CML.²⁹ In this pool of CD34⁺ cells collected from PB, CD34⁺CD25⁺ cells are strongly aggregated towards the very bottom of the viSNE plot, in the CD34⁺CD38⁻low region. During seven days of TKI treatment, we saw a statistically significant (P≤0.01) depletion of CD34⁺ cells from PB leukocytes (Figure 1B). Furthermore, after gating of CD34⁺CD25⁺ double positive cells, we saw a statistically significant depletion of CD34⁺CD25⁺ cells compared to CD34⁺ cells at seven days of therapy (Figure 1C). The 85th percentile metal intensity of pAbl Y245, pCRKL Y207, pSTAT3 Y705 and pSTAT5 Y694 was calculated for the manually gated CD34⁺CD38⁻low and CD34⁺CD38⁻high single cell populations. Comparing the changes of phosphorylation level across patients over time, we found a statistically significant increase in pSTAT3 Y705 in the CD34⁺CD38⁻high population (P≤0.05) between samples collected before and after seven days of TKI therapy (Figure 1D). Based on these observations in the CML progenitor compartment, we decided to examine more highly differentiated CML cells based on a more extensive panel of antibodies characterizing the myeloid and lymphoid lineages in greater detail (Online Supplementary Figures S4 and S5).

Mass cytometry identified hematologic remission in clinical trial samples

To validate our methodological approach, we analyzed a second cohort of patients (n=6) enrolled in the ENEST1st clinical trial. From each patient, 3 longitudinally collected samples (before, and after 3 hours and day 7 of TKI treat-
ment) were barcoded, along with 4 healthy PB and 4 healthy BM samples, using the commercially available 20-plex metal barcoding kit (Fluidigm). Each longitudinal set of 3 samples from a patient, and healthy PB and BM, was analyzed using the SPADE clustering algorithm, and the major leukocyte subsets were manually identified and annotated (Figure 2A). The relative abundance of a selection of cellular subsets quantified in longitudinal samples collected before and after seven days of TKI therapy is shown in Figure 2B and C. The abundance of the corresponding healthy cell subset is shown in a lighter shade, identified either from the healthy PB or BM samples (n=4 each). Similar to previous results (Online Supplementary Figure S3), already at day 7 of TKI therapy we observed a statistically significant change in the relative abundance of several cellular subsets. We measured a statistically significant reduction of myelocytes and progenitor cells (CD34+CD38−), and an expansion of mature neutrophils, CD8+ T cells, and activated T cells, (Wilcoxon matched pairs signed rank test P<0.05 for all).

The direct measurement of phosphorylated Bcr-Abl1 in single cells from patients treated with nilotinib

The phosphorylation level of several key signaling molecules in the Bcr-Abl1 signal transduction network was characterized in each patient, as a function of time and nilotinib treatment (before, and after 3 hours and 7 days of TKI therapy; n=8), and in the healthy PB and BM (n=4 each) (Figure 3A). In general, these data support our initial findings (Online Supplementary Figure 3C), reproducing the clearly distinct signaling status observed in healthy PB and CML at diagnosis, including the attenuated level of pSTAT3 Y694 phosphorylation in the CML neutrophils. In the larger patient cohort (n=8), we are able to measure a statistically significant decrease in pCRKL Y207 as a function of Bcr-Abl1 inhibition in the basophils (between before and after 7 days on TKI; P≤0.05), and an increase in pSTAT5 Y694 in the CD4 T helper cells (between diagnosis and after 3 hours of TKI; P≤0.01). We also measured a clear trend of pAbl Y245 downregulation in the CD34−CD38+ population as a function of TKI-therapy (simple Wilcoxon test after 7 days of TKI P=0.0313). The seemingly ubiquitous regulation of pCREB S133, initial decrease and subsequent increase, reached statistical significance in the neutrophils (between before and after 7 days, and between diagnosis and after 3 hours of TKI; P≤0.05 for both) but was observed in several cellular subsets. In the hematopoietic stem and progenitor compart-

**Figure 2. High-resolution single cell immune profiles of healthy and patient leukocytes.** Longitudinally collected samples (before, after 3 hours and day 7 on nilotinib) from 8 patients in the ENEST1st trial, together with 4 healthy peripheral blood (PB) and bone marrow (BM) samples, were barcoded using the 20-plex metal barcoding kit (Fluidigm). (A) Data from the longitudinal samples from each patient, and the 4 healthy PB and BM samples, were pooled and clustered using the SPADE algorithm and manually annotated to identify cellular subsets. The SPADE tree analysis of patient 4702_0004 is shown. The size of each node represents the number of cells clustered and the expression of pSTAT3 Y705 is color-coded. The red bubble highlights the mature neutrophil population. (B and C) The relative abundance of the major PB populations identified in samples collected at diagnosis and after seven days of tyrosine kinase inhibitor (TKI) therapy is shown for the patient cohort (n=8, error bars showing standard error of mean, SEM), with the appropriate subpopulation in the healthy samples shown in a lighter shade. CD34+ cells could only be identified in 7 out of 8 patients. Wilcoxon matched-pairs rank test was used to identify statistically significant changes from before and after seven days of TKI therapy, where P≤0.05 was considered statistically significant.
Figure 3. Single cell signaling profiles correlate to BCR-ABL1* molecular response. The phosphorylation level (75th percentile of metal intensity) of the intracellular Bcr-Abl1 signaling network (n=10) was measured in the longitudinal samples (before, after 3 hours and after 7 days of nilotinib) in the patient cohort (n=8), together with 4 healthy peripheral blood (PB) and bone marrow (BM) samples. (A) We observed distinctly different states of signal transduction in the patient samples compared to the healthy controls. (B) The Lin-CD34+CD38lo and Lin+CD34+CD38hi cell populations were manually gated from the CD34+ population identified by SPADE, and the intracellular signal transduction measured. CD34+ cells could only be identified in 7 out of 8 patients. The same strategy was followed for healthy CD34+ cells. For both (A and B) statistical significance was determined using Friedman non-parametric with Dunn’s multiple comparison test, where $P \leq 0.05$ was considered statistically significant ($*P \leq 0.05$, **$P \leq 0.01$). (C) The BCR-ABL1* of each patient during tyrosine kinase inhibitor (TKI) therapy. The patient cohort was divided into two groups based on the BCR-ABL1* at three and six months (red and green). (D) An unsupervised PCA analysis of the signal transduction arcsinh fold change compared samples before and after seven days of nilotinib treatment in the mature neutrophils (Neutro) and myelocytes (Myelo) was performed using Unscramble software (CAMO Software). The categorical color-coding from (C) was superimposed on the resulting PCA plot, and the dashed line was drawn manually.

TKI-mediated changes in single cell signal transduction reflects BCR-ABL1* at 3 and 6 months

Finally, we examined whether the observed TKI-induced changes in signal transduction networks could provide prognostic information. We split the patients according to their BCR-ABL1* at three and six months into two response groups (Figure 3C, low in green and high in red). The arcsinh fold change of each signal transduction epitope (n=10) between diagnosis and after seven days of TKI treatment was calculated for the neutrophils and myelocytes for all patients (n=8). We performed an unsupervised multivariate principle component analysis (PCA) of the fold change data using Unscrambler X (CAMO Software), and then manually overlaid the response group affiliation (Figure 3D). The PCA analysis revealed underlying differences in signaling status between the response groups, driven mainly by the rates of downregulation of pCREB S133 and upregulation of pSTAT3 Y705.

Discussion

Mass cytometry enabled the measurement of more than 30 intracellular and cell surface markers on each single cell, with an additional six channels reserved for metal barcoding for sample multiplexing. The convenient single tube labeling of multiplexed samples combined with semi-automated analysis makes the technique highly efficient. We were able to recapitulate the expected hematologic response, and could also demonstrate the debulking of leukemic CD34+ cells in the PB by immunophenotyping as early as one week after start of therapy. Importantly, by simultaneously probing key intracellular phosphorylation targets of the Bcr-Abl1 signaling network, we monitored changes in signal transduction of individual cell types for each patient undergoing TKI therapy. Unsupervised principle component analysis of these early changes in signal transduction allowed patients to be identified according to their BCR-ABL1*, indicating a possible future prognostic impact of this approach.

The proportion of leukemic stem cells (as determined by FISH) in the BM of chronic phase CML have been shown to have a prognostic significance. In the PB of our patients, we were able to monitor the therapy-dependent debulking of CD34+ cells (Figure 1B), and both CD34+CD38low and CD34+CD38hi cells (data not shown), at day 7 of therapy. Interestingly, at day 7 we saw a statistically significant depletion of CD34+CD25+, a putative CML stem cell subset compared to all CD34+ cells (Figure 1C). This could indicate an increased sensitivity to TKI or alternatively an increased bone marrow homing of this cell subset. Early relative increase of lymphocytes in the PB of CML patients treated with TKI has also previously been shown to have prognostic importance. Our immune profiling of patient leukocytes allowed a high-resolution picture of PB subpopulations, enabling the
simultaneous monitoring of the size of all major hematopoietic lineages. Here, the analysis of healthy PB and BM side-by-side with primary material clearly delineates cell populations dominating in CML compared to healthy individuals (Figure 2). Interestingly, we were able to measure statistically significant changes in abundances for several cellular subpopulations, including an increase in PB CD8+ cytotoxic T cells already after seven days of TKI therapy.

We believe that a particular strength of our single cell analysis is the direct monitoring of nilotinib-induced changes in signal transduction. This includes the primary target kinase for nilotinib, ABL1, as well as its immediate substrates like pCRKL Y2073 and pivotal ABL1-affected signal transduction pathways in myeloid neoplasias, e.g. the STAT3/5 pathway.34,35 In the STAT3/5 pathway,16,17,33 a decrease of pCREB S133 as early as three hours after first TKI dosing in several cellular subsets followed by an increase after seven days. Interestingly, we see that, compared to non-CML cell lines, K562 cells express high levels of pCREB S133 at baseline that was reduced by TKI treatment (Online Supplementary Figure S3C). Using a principle component analysis to analyze changes in signal transduction between diagnosis and after seven days of TKI treatment, we could discriminate patient groups according to their BCR-ABL134 at three and six months (Figure 3D). The PCA analysis was driven mainly by mutual regulation of pCREB S133 and pSTAT3 Y705, and underscores a potential prognostic application for this assay.

In summary, here we demonstrate the first human study of signal transduction in cancer cells from nilotinib-treated CML patients in a prospective clinical trial, demonstrating significant phosphorylation of CREB and STAT3 in chronic phase CML patients early after start on therapy. We propose that mass cytometry, enabling simultaneous measurement of more than 30 parameters on single cells, may provide a highly detailed characterization of CML that allows comprehensive 1-tube-based monitoring of the disease in patients during TKI therapy. The method outlined here will allow early changes in individual patient’s immunophenotype and signal transduction as a function of TKI therapy to be monitored in clinical trials. Furthermore, this approach may shed light on the relation between the kinetics of TKI-modulated signal transduction and clinical response. We anticipate this technology to have high clinical impact by providing cancer cell signal transduction profiles in patients close to real time after start of TKI therapy. Additional work on standardization of antibody panels is a prerequisite if mass cytometry is to be established as a future tool to guide signal transduction-targeted therapy.

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