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Inhibition of Aurora-kinases for Tailored Risk Adapted Treatment of Multiple Myeloma

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Abstract

Genetic instability and cellular proliferation have been associated with Aurora-kinase expression in several cancer entities, including multiple myeloma. Therefore, the expression of Aurora-A, -B and -C was determined by Affymetrix DNA-microarrays in 784 samples including two independent sets of 233 and 345 CD138-purified myeloma-cells from previously untreated myeloma-patients. Chromosomal aberrations were assessed by comprehensive iFISH and proliferation of primary myeloma-cells by propidium-iodine staining. The effect of the clinical Aurora-kinase inhibitor VX680 on proliferation of 20 human-myeloma-cell-lines and survival of 5 primary myeloma-cell-samples was tested. We found Aurora-A and -B to be expressed at varying frequencies in primary myeloma-cells of different patient-cohorts, Aurora-C in testis-samples only. Myeloma-cell samples with detectable vs. absent Aurora-A expression show a significantly higher proliferation rate, but neither a higher absolute number of chromosomal aberrations present (aneuploidy) nor of subclonal aberrations (chromosomal instability). VX680 induces apoptosis in all myeloma-cell-line- and primary myeloma-cell-samples tested. Presence of Aurora-A expression delineates significantly inferior event-free and overall-survival in two independent cohorts of patients undergoing high-dose chemotherapy, independent of conventional prognostic factors, i.e. serum-ß2-microglobulin or ISS-stage. In conclusion, using gene expression profiling, Aurora-kinase inhibitors as promising therapeutic option for newly-diagnosed patients can be tailoredly given to patients with adverse prognosis, expressing Aurora-A.
Introduction

Multiple myeloma is an incurable malignant disease of clonal plasma cells which accumulate in the bone marrow causing clinical signs and symptoms related to the displacement of normal hematopoiesis, formation of osteolytic bone lesions, and production of monoclonal protein \(^1\). Multiple myeloma cells (MMC) at the time of diagnosis are characterized by a low proliferation rate that increases in relapse \(^2\). Presence of proliferation correlates with adverse prognosis \(^3;4\). At the same time, myeloma cells harbor a high median number of chromosomal aberrations \(^5;6\), often associated with genetic instability \(^6\). Proliferation \(^7\) and chromosomal instability \(^8\) in turn are associated with the expression of Aurora-kinases. In several cancer entities, Aurora-kinases have been implicated in tumor formation and progression \(^9;14\).

Aurora-kinases represent a family of conserved mitotic regulators comprising three closely related members, i.e. Aurora-A, -B and -C \(^9\). Aurora-C expression is restricted to germ cells, where it regulates spermatogenesis \(^15\). Aurora-A associates with the centrosome and spindle microtubules being required for centrosome separation and spindle assembly \(^7\). Aurora-B is a member of the chromosomal passenger complex, as such sequentially recruited to centromeres, spindle midzone and midbody as mitosis progresses \(^16\), and required for chromosome biorientation, the spindle assembly checkpoint, and cytokinesis \(^7\). Inhibition of Aurora-A and -B exhibits distinct phenotypic features: loss of Aurora-A activity induces a centrosome separation defect and a monopolar spindle phenotype \(^17;18\); inhibition of Aurora-B generates polyploidy through defects in cytokinesis, which ultimately leads to a loss in cell viability \(^19;22\). Aurora-A and -B expression has been detected by quantitative real-time PCR (qRT-PCR) in myeloma cell lines \(^23;24\) and small series of myeloma patients \(^23;24\). Aurora-kinase inhibitors like VX680 have been shown to abrogate proliferation and induce apoptosis in human myeloma cells lines and primary myeloma cells \(^23;25\).

We assess here the expression of Aurora-A, -B, and -C in 784 Affymetrix gene expression profiles of malignant plasma cells from previously untreated myeloma patients compared to normal bone marrow plasma cells (BMPC), their non-malignant proliferating precursors (polyclonal
plasmablastic cells, PPC), and human myeloma cell lines (HMCL). We find that in our data set 24% of previously untreated myeloma patients express *Aurora-A*. We show myeloma cells expressing *Aurora-A* kinase to have a higher proliferation-rate, whereas the number of chromosomal aberrations (aneuploidy) is not higher compared to myeloma cells with absent *Aurora-A* expression. The same holds true for subclonal aberrations (i.e. genetic instability), which are less frequent in myeloma cell-samples expressing *Aurora-A*. *Aurora-A* kinase expression in turn is significantly associated with an inferior event-free (EFS) and overall survival (OAS) in two independent cohorts of a total of 513 myeloma-patients treated with high-dose chemotherapy (HDT) and autologous stem cell transplantation (ASCT).

Aurora-kinase inhibitors (including VX680 tested here) are very active on human myeloma cell lines and primary myeloma cells and represent a promising weapon in the therapeutic arsenal against multiple myeloma. Gene expression profiling allows an assessment of Aurora-kinase expression and thus in turn a tailoring of treatment to patients expressing these kinases.
Materials and Methods

Patients and healthy donors

Patients presenting with previously untreated MM (n=233) or monoclonal gammopathy of unknown significance (MGUS, n=12) at the University Hospitals of Heidelberg and Montpellier and 14 healthy normal donors were included in the study approved by the institutional review board of the Medical Faculty of the Ruprecht-Karls-University Heidelberg, Germany, the institutional review board of the University of Arkansas for Medical sciences, AR, USA, and the institutional review board of the CHU Montpellier, France, for the respective patients. Written informed consent was obtained in accordance with the Declaration of Helsinki. The first 65 patients comprise the training group (TG), the 168 additional the independent validation group (VG). Patients were diagnosed, staged and response to treatment was assessed according to standard criteria 26-28. 168 patients underwent frontline HDT with 200 mg/m² melphalan and ASCT according or in analogy to the GMMG-HD3-trial 29. Survival data were validated by an independent cohort of 345 patients treated within the total therapy 2 protocol 30. For clinical parameters, see Supplementary Table S1.

Samples

For an overview, see Supplementary Table S2. Plasma cells from the bone marrow were purified using CD138-microbeads (Miltenyi Biotech, Bergisch Gladbach, Germany) and purity was assessed by flow cytometry (FACSCalibur, Becton Dickinson, Heidelberg, Germany). An aliquot of unpurified (whole) bone marrow (WBM) of patients (n=57) and healthy donors (n=7) was obtained after NH₄-lysis as published 31. An alternate aliquot was subjected to FACS-sorting (FACSaria, Becton Dickinson) in CD3⁺-, CD14⁺-, CD15⁺-, and CD34⁺-cells. Peripheral CD27⁺-memory B-cells (MBC) were generated as published 32. Testis-samples (n=5) were obtained from healthy donors.

The HMCL XG-1, XG-2, XG-3, XG-4, XG-5, XG-6, XG-7, XG-10, XG-11, XG-12, XG-13, XG-14, XG-16, XG-19, XG-20 were generated at INSERM U847 as published 33-35. HG-1 was generated in the Multiple Myeloma Research Laboratory Heidelberg. U266, RPMI-8226, LP-1, OPM-2, SKMM-2, AMO, JJN-3, KMS-12-BM, L363, NCI, MOLP-8 (Deutsche Sammlung von
Mikroorganismen und Zellkulturen GmbH, Braunschweig, Germany) were cultured as recommended. PPC (n=12) \(^{36}\), osteoclasts \(^{32}\) (OC; \(n=5\)) and mesenchymal stromal cells (MSC, \(n=19\)) \(^{37}\) were generated and analyzed as previously published.

**Chemicals**

The 4,6 diaminopyrimidine VX680 (Cyclopropanecarboxylicarid[4[4(4-methyl-piperazin-1-yl)-6-(5-methyl-2H-pyrazol-3-ylamino)-pyrimidin-2-ylsulfanyl]-phenyl]-amide; ACC corporation, San Diego, CA, USA) was dissolved in dimethyl sulfoxide (DMSO; Merck, Darmstadt, Germany) and stored at a final concentration of 10 mM at -80 °C.

**Interphase-FISH (iFISH)**

Interphase-FISH-analysis was performed on CD138-purified plasma cells as described \(^{5}\) using a set of probes for the chromosomal regions 1q21, 6q21, 8p21, 9q34, 11q23, 11q13, 13q14.3, 14q32, 15q22, 17p13, 19q13, 22q11, as well as the translocations t(4;14)(p16.3;q32.3) and t(11;14)(q13;q32.3), (Poseidon-probes, Madison, WI, USA). Ploidy-status, clonal/subclonal aberrations for a single aberration (i.e. present in \(\geq 60\%\) vs. 20-59 % of assessed MMC) were defined as published \(^{5}\). A modified copy number score (CS) \(^{5,38}\) (excluding gains of 1q21) and the score of Wuilleme et al. \(^{39}\) using chromosomes 5, 15, 19 (CSW) was used to assess ploidy. For the assessment of the absolute number of chromosomal aberrations, 105 patients were assessed for the translocations t(4;14) and t(11;14) as well as numerical aberrations of the chromosomal regions 11q13, 11q23, 1q21, 17p13, 13q14.3, 14q32. For the assessment of the presence of subclonal aberrations, a MMC-sample was considered to contain a subclonal aberration, if at least one aberration was detected in \(\geq 70\%\) of the myeloma cells present, and at least one other aberration was detected in 20-59 % of assessed MMC.

**Gene expression profiling (GEP)**

RNA extraction was performed using the RNeasy kit (Qiagen, Hilden, Germany), the SV-total RNA extraction kit (Promega, Mannheim, Germany) and Trizol (Invitrogen, Karlsruhe, Germany). Labeled cRNA was generated using the small sample labeling protocol vII (Affymetrix, Santa Clara, CA, USA), and hybridized to U133 A+B GeneChip microarray (Affymetrix) for training,
and U133 2.0 plus arrays for validation group, according to the manufacturer’s instructions. When different probe sets were available for the same gene, we chose the probe set yielding the maximal variance and the highest signal. Aurora-A (Hs00233470_m1), Aurora-B (Hs00177782_m1) and Aurora-C (Hs00152930_m1, all Applied Biosystems, Darmstadt, Germany) gene expression was analyzed by qRT-PCR using the ABI Prism 7700 Sequence Detection System. The expression data are deposited in ArrayExpress under the accession numbers E-MTAB-81 and E-GEOD-2658.

Measurement of proliferation by ³H-thymidine

Proliferation of 20 HMCL was investigated according to published protocols. Per well, 10,000 cells were cultured in 96-well-plates in RPMI-1640 (Biochrom, Berlin, Germany) containing 10 % FCS (Invitrogen) with or without (medium control) graded concentrations (10, 2, 0.4, 0.08, 0.0032 µM) of VX680. DMSO at the highest concentration present in the 10 µM well served as DMSO-control. For the HG- and XG-lines, 2 ng/ml IL-6 (R&D Systems, Wiesbaden, Germany) was added. Proliferation was evaluated after 5 days of culture: cells were pulsed with 37 kBq of ³H-thymidine for 18 h, harvested and ³H-thymidine uptake measured.

Measurement of proliferation of primary myeloma cells by propidium iodine

The Plasma Cell Labeling Index, i.e. the percentage of MMC in S-phase, was determined by flow cytometry using a FACSCalibur. WBM (10⁶ cells per tube) was incubated with 20 µl of either control IgG-FITC, CD38-FITC (both Beckman Coulter, Krefeld, Germany, clone A07795 and A07778) and CD138-FITC (Diaclone, Stamford, CT, U; 954.501.010), respectively. After NH₄-lysis, cells were resuspended with propidium iodine (PI-) solution (1mg/ml PI in 1x citrate buffer containing 0.1 % Tween 1mg/ml RNase A (Sigma-Aldrich, Schnelldorf, Germany)) for 45 min at 4 °C. The percentage of CD138⁺ S-phase cells was determined using ModFit software (Verity Software House, Topsham, ME, USA) using a rectangular mathematical model for calculating the S-phase fraction in % of the selected CD138⁺ plasma cells.

Survival of primary myeloma cells

Primary MMC cultured together with their bone marrow microenvironment (negative fraction of plasma cell purification) of 5 newly-diagnosed patients were exposed to concentrations of 100, 20,
4, 0.8, 0.16, 0.032 µM VX680. Cell viability was measured by CD138-FITC (IQ products, Groningen, Netherlands, clone B-A38) / PI (Pharmingen, Heidelberg, Germany) staining after 6 days of culture and referred to the medium and DMSO-control, respectively. One µl of PI with a concentration of 50 µg/ml was used.

**Apoptosis induction**

XG-1 and XG-10 were cultured in 24-well-plates at 10^5 cells per well in RPMI-1640 containing 10 % FCS and 2 ng/ml IL-6 with or without 1 µM VX680. After 8, 24, 48 and 72 h of culture, cells were stained for annexin V-FITC and PI according to the manufacturer’s instructions (Pharmingen) and analyzed on a FACS Aria.

**Intracellular staining for Aurora-A and -B**

Intracellular Aurora-A (clone 35C1; Abcam, Cambridge, United Kingdom) and -B (Cell signaling technology, Danvers, MA, USA) expression of 10 HMCL was measured by flow cytometry using a fixation and permeabilization kit (eBioscience, San Diego, CA, USA). Overlays were established using the Infinicyt 1.1 Software (Cytognos, Salamanca, Spain).

**Western blotting**

Cells were pelleted and resuspended in lysis buffer containing 10mM Tris-HCl (pH 7.05), 50 mM sodium chloride, 30 mM sodium pyrophosphate decahydrate, 50 mM sodium fluoride, 5 µM zinc chloride, 1 % Triton-X 100, and a protease/phosphatase inhibitor cocktail (Complete mini tablets; Roche, Basel, Switzerland). After pelleting, supernatants were mixed with loading buffer (Roti, Carl Roth, Karlsruhe, Germany), heated for 5 min at 95 °C and separated on 10 % NuPAGE Bis-tris gels (Invitrogen). Immunodetection was performed using the WesternBreeze Kit (Invitrogen). Membranes were incubated with antibodies against Aurora-A, -B (see above) and β-actin (Ab5, Becton Dickinson) as loading control. HELA cells served as positive control.

**Statistical analysis**

Gene expression data were germa-normalized. To assess presence or absence of gene expression independently of Affymetrix-mismatch-probesets, the “Presence- Absence calls with Negative
Probesets (PANP)” algorithm \(^46\) was used. Differential gene expression was assessed using empirical Bayes statistics in linear models for microarray data \(^47\). \(P\)-values were adjusted for multiple testing controlling the false-discovery-rate as defined by Benjamini and Hochberg at a level of 5 % \(^48\). Expression profiles of 439 samples (233 MM, 14 BMPC, 12 PPC, 12 MGUS, 40 HMCL, 13 MBC, 64 WBM, 19 MSC, 5 CD3, 5 CD14, 5 CD15, 5 CD34, 7 OC, and 5 testis) divided in TG \((n=113, \text{MM } n=65)\) and VG \((n=257, \text{MM } n=168)\) were analyzed. As a further validation, 345 samples of newly-diagnosed myeloma patients from the Arkansas-group were analyzed.

Event-free survival \(^29\) and overall survival \(^29\) were investigated for the 168 patients (48 TG, 120 VG) undergoing HDT and ASCT using Cox’s proportional hazard model. Two groups of patients with presence and absence of Aurora-A expression were delineated. Findings were validated using the same strategy on the independent group of 345 patients from the Arkansas-group. For myeloma cells, association of chromosomal aberrations and clinical parameters with gene expression was calculated using two-sample t-statistic. Differences in clinical parameters between defined groups were investigated by analysis-of-variance (ANOVA). Correlation was measured using the Spearman correlation coefficient. Correlation with categorical variables was measured using the Kendall’s tau coefficient. For assessing the relationship between categorical variables, Fisher’s Exact Test was used. The centrosome-index was calculated as published by Chng et al. \(^49\). For the calculation on the Arkansas-group, our 7 BMPC samples were normalized together with the 345 MMC-samples.

The gene expression based proliferation index is calculated as described in Supplementary Text S1. In all statistical tests, an effect was considered as statistically significant if the \(P\)-value of its corresponding statistical test was not greater than 5 %. All statistical computations were performed using R \(^50\) version 2.7.0 and Bioconductor \(^51\), version 2.2.
Results

Expression of Aurora-A, -B and -C

First, we assessed expression (Table 1,2) and differential expression (Supplementary Table S3A,B) of Aurora-A, -B, and -C in primary myeloma cells, normal bone marrow plasma cells, their precursors, as well as normal and myelomatous bone marrow. In our data set, Aurora-A and -B are expressed in 24 % and 3 % of primary myeloma cells and all PPC as well as HMCL. In the Arkansas-data, Aurora-A, -B, and -C are expressed in 48/345 (14 %), 12/345, (3 %) and 0/345 myeloma cell-samples, respectively. The mean expression of Aurora-A and -B is significantly and by several orders of magnitude higher in proliferating plasmablastic cells and cell lines compared to non-proliferating MBC, or BMPC (each P<.001 in TG and VG; Supplementary Table S3A,B, Figure 1A). Aurora-A and -B are expressed in almost all bone marrow samples of healthy individuals and myeloma patients (Table 1). Here, the mean expression of Aurora-B is significantly different (lower) in myelomatous compared to normal bone marrow (4.85 vs. 3.81 arbitrary units, P=.009). A significant stage dependent differential gene expression could be found for Aurora-A between myeloma cells from early- (MGUS and MMI) and advanced stage (MMII and MMIII) patients (P=.01 and .01, in TG and VG, respectively). Aurora-A and -B expression correlates significantly in the VG (r_s=0.59, P<.001) and Arkansas-group (r_s=0.61, P<.001).

Validation of gene expression by qRT-PCR, western blotting and flow cytometry

To validate Aurora-kinase expression detected by gene expression profiling, we performed qRT-PCR, western blotting and flow-cytometric staining. Aurora-A expression in terms of “presence” or “absence” (C_t-value ≥ 35) by qRT-PCR is consistent with results by PANP in 10/11 primary myeloma cell-samples. One sample “absent” by qRT-PCR is judged “marginal” by PANP. Aurora-A expression by GEP strongly correlates with dC_t value by qRT-PCR (r_s= -0.87, P<.001). Aurora-B expression is consistent with results by PANP in 3/6 samples. All samples are “present” by qRT-PCR but three are judged “absent” by PANP. Aurora-B expression by GEP strongly correlates with dC_t value by qRT-PCR (r_s= -0.83, P=.06). Aurora-C expression by qRT-PCR is
consistent with absence of expression detected by PANP in 5/6 samples. One sample “present” by qRT-PCR (Ct-value 34.8) is judged “absent” by PANP. *Aurora-C* expression by GEP strongly correlates with the dCt value obtained by qRT-PCR (r= -0.94, P=.02). *Aurora-A* and -*B* expression in HMCL was further validated by western blotting (Figure 2B) and intracellular flow cytometry (exemplary data shown in Figure 2C).

**Association of Aurora-kinase expression with proliferation and chromosomal aberrations**

To investigate the biological impact of Aurora-kinase expression, we assessed the association with proliferation, chromosomal aberrations and presence of subclonal aberrations as detected by iFISH, and a published centrosome-index 49.

The expression of *Aurora-A* correlates with proliferation in terms of the plasma cell labeling index assessed by PI staining (n=66, r=0.45, P<.001) and the gene expression based proliferation index in TG (r=0.62, P<.001) and VG (r=0.87, P<.001). The same holds true for the latter for *Aurora-B* in the VG (r=0.64, P<.001).

The absolute number of chromosomal aberrations is not significantly different between myeloma cells expressing or not expressing *Aurora-A* (mean 5.1 and 5.1, n=105 patients, n=8 FISH-probes tested). Presence/absence of *Aurora-A* expression does not significantly interrelate to the presence/absence of hyperdiploidy as determined by either CS or CSW, neither does the presence/absence of *Aurora-A* expression interrelate to the presence of any of the single aberrations t(11;14), t(4;14), or numerical aberrations of 17p13, 9q34, 15q22, 19q13, 4p16, 14q32 or 22q11.

Interestingly, in patients with presence of *Aurora-A* expression, gains of 11q13 (P=.03) and 11q23 (P=.004) are significantly less frequent compared to those with absent *Aurora-A* expression. The opposite holds true for patients with gain of 1q21 (P=.002) or deletion of 13q14 (P=.03) as well as deletion of 8p21 (P=.03): MMC of patients with present *Aurora-A* expression show a significantly higher number of these respective aberrations. For a gain 1q21, for which data were also available for the Arkansas-group, the same observation was made (n=244, P<.001). However, subclonal aberrations *per se* are significantly more frequent in MMC with absent *Aurora-A* expression (67 with vs. 37 without) compared to present *Aurora-A* expression (18 vs. 23, P =.03). For single
aberrations, subclonal presence (vs. full-clonal or absence of the respective aberration) is significantly more frequent in MMC of patients with absence of Aurora-A expression for gains of 11q13 ($P<.03$), 11q23 ($P<.001$), and losses of 13q14 ($P=.009$). Losses of 17p13 marginally fail significance ($P=.06$). Losses of 8p21 are significantly ($P=.04$) more frequent in patients with presence of Aurora-A expression.

The centrosome-index correlates with Aurora A-expression in our series (TG $r=.53$, $P<.001$; VG $r=.32$, $P<.001$) and the Arkansas-data ($r=.43$, $P<.001$). The centrosome-index is significantly predictive for EFS and OAS in the Arkansas-group (EFS $P=.005$, OAS, $P<.001$) on which it has been derived, but not our series (EFS $P=.09$, OAS, $P=.99$; Supplementary Figure S1).

**Prognostic value of Aurora-kinase-expression**

Next, we investigated whether the presence of Aurora-kinase expression has a prognostic impact in newly-diagnosed myeloma patients treated with HDT and ASCT.

Presence of Aurora-A expression in MMC is an adverse prognostic factor in terms of EFS and OAS in our data ($n=168$, EFS $P=.003$, hazard ratio (HR) 2.02, confidence interval (CI) [1.26,3.25], OAS, $P=.03$, HR 2.31, CI [1.04,5.15]) and the Arkansas-group ($n=345$, $P=.004$, HR 1.74, CI [1.18,2.57], OAS, $P=.008$, HR 1.85, CI [1.16,2.95]), Figure 3. The expression-signal of Aurora-A as single continuous variable is significantly predictive for EFS in the VG ($n=120$, $P<.001$) and the Arkansas-group ($P<.001$). The same holds true for OAS in the Arkansas-group ($P<.001$) and it marginally failed significance for our VG ($n=120$, $P=.06$).

In a Cox-model tested with the international staging system (ISS), presence of Aurora-A expression appears as independent prognostic factor for EFS in our data (Aurora A-expression ($P=.003$), ISS ($P=.009$)), and the Arkansas-data (Aurora-A expression ($P<.001$), ISS ($P<.001$)). For OAS, Aurora-A expression marginally fails independence ($P=.08$, ISS ($P=.009$)) in our data-set, but is significantly independent in the Arkansas-data (Aurora-A expression ($P<.001$), ISS ($P<.001$)).

In a Cox-model tested with serum-β2-microglobulin (B2M) as continuous variable, presence of Aurora-A expression appears as independent prognostic factor for EFS in our data (Aurora-A expression ($P=.002$), B2M ($P=.006$)), and the Arkansas-data (Aurora-A expression ($P<.001$), B2M
Activity of VX680 on myeloma cell lines and primary myeloma cells

Given the expression and prognostic value of Aurora-kinases, we tested the activity of the pan-Aurora-kinase inhibitor VX680 that has previously shown activity on a small series of human myeloma cells, on a large series of 20 myeloma cell lines. VX680 significantly inhibits proliferation of all HMCL investigated (Figure 4A1). The maximum inhibition at 10 µM ranges from 64.4 % (HG-1) to 100 % (OPM-2). The concentration to reduce proliferation to half of the control value (IC50) is reached in all myeloma cell lines; it ranges from 0.003 - 2.715 µM, Figure 4A2. No significant correlation could be found between the expression of Aurora-A, -B or hyaluron mediated motility receptor (HMMR) and the IC50 of 12 myeloma cell lines tested (see discussion).

VX680 significantly inhibits the survival of primary myeloma cells cultivated within their bone marrow microenvironment from 5/5 newly-diagnosed myeloma patients at a concentration of 4 µM (50 - 100 % inhibition, Figure 4B1). At the same dose level, VX680 induces significant but lower toxicity within the bone marrow microenvironment (Figure 4B2). Four of four samples, for which sufficient RNA was available, showed an expression of Aurora-A by qRT-PCR.

Next, XG-1 and XG-10 were cultured for 3 days with or without VX680. Cell viability and apoptosis were determined by flow-cytometric analysis of annexin V-binding and PI uptake after 8, 24, 48 and 72 h. Exposure of XG-1 and XG-10 against 1 µM VX680 induces apoptosis after 8 h (XG-1: 23.5 % vs. 24.1 %, XG-10: 16.4 % vs. 13.3 %) to 72 h (XG-1: 17.9 % vs. 85.5 %, XG-10: 9.0 % vs. 49.9 %), Figure 4C, exemplary data shown for XG-10.
Discussion

Expression of Aurora-kinases

In our set of previously untreated myeloma patients (n=233), expression of Aurora-A and -B could be detected in 24% and 3% of purified myeloma cell-samples. The same percentage of Aurora A- and -B expression could be found for the subgroup of patients treated with HDT and ASCT (n=168, 23.2% and 3.5%, respectively). In the independent data set of Shaughnessy et al. (n=345), the same percentage of patients (3%) expresses Aurora-B, but Aurora-A expression could only be detected in 14% of patients. This observation could not be explained by the use of U133 A+B chips for part of our patients, as in these, the percentage of patients expressing Aurora-A is even lower (10.4% of patients in the HDT-cohort). Likewise, the use of a double amplification (our data) instead of single amplification protocol (Arkansas-group) could not be taken as explanation, as one would rather expect a higher percentage of detection in the single-amplification group. In an additional set of patients treated within the GMMG-HD4 trial, an Aurora-A expression can be detected in 43/70 (61%) of cases. Taken together, the percentage of patients expressing Aurora-A seems to be quite variable in different patient populations, indicating to assess Aurora-A expression when testing Aurora-inhibitors within clinical trials.

“Presence” of gene expression as determined by gene expression profiling based on PANP needs to be interpreted as presence above the background level (“threshold”) seen for unspecific hybridization for negative strand matching probesets. As such a background correction is not performed when analyzing qRT-PCR data, the latter might have a detection threshold within the background of gene expression. This is one possible explanation why in a previously published small patient series, Evans et al. found by qRT-PCR all CD138-purified myeloma samples to express Aurora-A (5/5) \(^{23}\) and Aurora-B (7/7) \(^{24}\). Alternate explanations are a more advanced (relapsed) patient population, or a contamination by other cell types, as in these series, purity of CD138-sorted plasma cells was only assessed by morphology, and expression of Aurora-A and -B could be detected in almost all our bone marrow samples (Table 1,2).
The lower frequency of *Aurora-B* compared to *Aurora-A* expression in the same sample as detected by GEP seems likewise to be related to the detection-threshold: (i) In normal plasma cells, the expression levels of *Aurora-A* and -B are of the same height (Supplementary Table S3A-B), but the differential expression of *Aurora-A* in proliferating plasmablastic cells and myeloma cell lines is higher compared to *Aurora-B* (Supplementary Table S3A,B). (ii) Despite of a tight correlation between qRT-PCR and gene expression profiling, qRT-PCR shows a comparable expression level for myeloma cell lines in terms of *Aurora-A* and -B expression (mean C_t 27.6 vs. 28.2). (iii) All samples expressing *Aurora-B* by PANP express likewise *Aurora-A*.

**Biological implications**

*Aurora*-kinases have been associated with proliferation 7 and genetic instability 8 in different cancer entities 9–14, including multiple myeloma 25. *Aurora-A* and -B are expressed in all myeloma cell lines and proliferating 36 (non-malignant) plasmablastic cells, and are significantly higher expressed in both of these compared to memory B-cells or normal plasma cells. At the same time, expression of *Aurora-A* and -B correlates with the plasma cell labeling index determined by PI-staining as well as the gene expression based proliferation index. Furthermore, the percentage of primary myeloma cells of untreated patients expressing *Aurora-A* is in agreement with the low proliferative rate of these. The same holds true for the significant increase in *Aurora-A* expression from early- to late-stage plasma cell dyscrasias in both our training and validation group. Thus, *Aurora*-kinase expression is obviously associated with proliferation in multiple myeloma.

As chromosomal aberrations can be detected by iFISH in almost all primary myeloma cells 5,6, e.g. in all our patients tested, but only myeloma cells from a minor fraction of myeloma patients express *Aurora-A* or -B, *Aurora*-kinase expression in CD138-positive primary myeloma cells cannot be the cause of aneuploidy or ongoing genetic instability in myeloma. Two further strong arguments are given by the fact that *Aurora-A* or -B expression height (or presence of expression) neither correlates with the median number of chromosomal aberrations in an individual sample, nor the presence of subclonal (i.e. emerging) aberrations, but to the contrary, presence of subclonal
aberrations at all is significantly associated with the absence of Aurora-expression. The same holds true if the presence of specific subclonal aberrations is considered (11q13, 11q23, 13q14) vs. clonal gain or normal copy number state with the exception of deletions of 8p21. It is interesting to denote that aberrations of 1q21 (gains), 13q14.3 (deletions) and 8p21 (deletions), the first two associated with advanced stages \(^{52,53}\), are significantly more frequent in myeloma cells expressing *Aurora-A*. At the same time, gains of 11q13 and 11q23 are less frequent in myeloma cells expressing *Aurora-A*.

It cannot, however, be ruled out by our analysis that Aurora-expression is present in a small fraction of myeloma cell precursors, *i.e.* putative proliferating “myeloma stem cells” thereby creating genomic instability, with this presence of Aurora-expression not being maintained in the differentiated non-proliferating progeny. If this is the case, one would expect the presence of Aurora-expression within the myeloma stem cells leading to a higher number of clonal and subclonal chromosomal aberrations in advanced stage or relapsed compared to early-stage myeloma patients. At the same time, structural aberrations or point mutations in the Aurora-kinase genes of myeloma cells might be present as indicator for a respective role of Aurora-kinase expression in putative myeloma stem cells. Both investigations, being beyond the scope of this paper, are currently performed by our group.

Taken together, it is unlikely that Aurora-kinase expression in CD138-purified myeloma cells drives genetic instability in myeloma, but is, as a high labeling index and the presence of chromosomal aberrations associated with disease “progression”, rather a sign of proliferative, “advanced” myeloma cells.

**Prognostic value of Aurora-kinase expression**

Presence of *Aurora-A* expression in myeloma cells is an adverse prognostic factor in terms of EFS and OAS in our data \((n=168)\) and the Arkansas-group \((n=345, \text{Figure 3})\), as is the expression-height of *Aurora-A* as single continuous variable. In a Cox-model tested with either ISS or B2M (as continuous variable), presence of *Aurora-A* expression appears as independent prognostic factor for EFS and OAS. Of note, *Aurora-A* (*STK6*) is also one of the genes within the gene expression
based high-risk score for myeloma. \textsuperscript{54} Aurora-A kinase expression in our data set correlates with the gene expression based centrosome index that has likewise shown prognostic significance on the Arkansas-dataset \textsuperscript{49} (Supplementary Figure S1). Thus, direct assessment of Aurora-A kinase expression allows identifying a poor-risk patient population independent of B2M or ISS.

**Inhibition of proliferation of HMCL and primary MMC by Aurora-kinase inhibitors**

Proliferation of all tested 20 myeloma cell lines is significantly inhibited by VX680 with an IC50 ranging from 0.003 to 2.715 µM (Figure 4A2). The observed minimal IC50 is in the range of data from Tyler et al. \textsuperscript{19} who have shown an inhibition of in vitro kinase activity by VX680 in terms of phosphorylation of histone-H3 by Aurora-A, -B, and -C (with IC50-values of 36, 18, and 25 nM, respectively); Aurora-A and -B kinase activity was completely abrogated at 1 µM \textsuperscript{19}. VX680 induced a marked delay in mitosis, presumably due to Aurora-A dependent effects on spindle bipolarity, but did override a mitotic arrest imposed by several experimental agents, indicating an effect on the spindle assembly checkpoint, likely through Aurora-B inhibition \textsuperscript{19}. Our data are in agreement with VX680 having been shown to induce a dose-dependent inhibition of proliferation on myeloma cell lines \textsuperscript{25}. Inhibition of proliferation by VX680 was accompanied in our hands by apoptosis induction in myeloma cell lines and primary myeloma cell-samples in agreement with findings from Shi et al. \textsuperscript{25}. Driven by the published observation that forced over-expression of Aurora-A reduces \textsuperscript{25}, and downregulation by siRNA increases \textsuperscript{23} the susceptibility of myeloma cell lines towards Aurora-kinase inhibitors, we investigated whether the IC50 in 12 myeloma cell lines tested might correlate with the expression level of Aurora-A. However, in our hands, it did not. The same observation was made in terms of expression of HMMR, for which a forced up-regulation was described to increase, and a down-regulation to decrease the sensitivity of the respective myeloma cell line \textsuperscript{25}, but does not in our data. This might be explained by a plethora of growth and survival-factors differentially expressed between HMCL causing a high inter-cell-line variance in terms of sensitivity against VX680 (Figure 4A2). Thus, only if Aurora-A or HMMR- expression is the single parameter varied (e.g. by siRNA) within one cell line, differences in the sensitivity of the respective HMCL might be seen.
Implications for myeloma treatment

The presence of multiple chromosomal aberrations in multiple myeloma suggests that, during the evolution of myeloma, a disruption of cell cycle checkpoints has occurred. This would normally arrest cells at the G1/S and G2/M transitions or at mitosis when DNA-damage or spindle abnormalities have occurred, thus allowing for potential damage-repair \(^{25}\). Alternatively or in addition, these checkpoints might be “overruled” by intrinsic aberrant or increased expression of D-type cyclins and myeloma growth and survival factors (“strained” checkpoints). In both cases, myeloma cells might be particularly susceptible to the induction of apoptotic death in mitosis (the so-called mitotic catastrophe \(^{55}\)) when further assaults on the mitotic machinery are induced. Experimental evidence for the latter is given by the fact that VX680 inhibits proliferation of both, CD3/CD28- or phytohaemagglutinin-stimulated peripheral blood lymphocytes and myeloma cell lines, but induces apoptosis only in human myeloma cell lines \(^{25}\). Thus, Aurora-kinase inhibitors are indicated in multiple myeloma not because myeloma cells show a genetic instability, but because Aurora-kinase inhibitors target and inhibit the proliferation of myeloma cells. They might do this especially efficient because they (additionally) induce apoptosis in the presence of strained or deranged cell-cycle checkpoints present in primary myeloma cells and human myeloma cell lines.

Conclusion

*Aurora-A* is expressed at varying frequencies in CD138-purified myeloma cells of newly-diagnosed patients. Its expression significantly interrelates with proliferation, but not with a higher number of chromosomal aberrations (aneuploidy) or subclonal aberrations (genetic instability). Using gene expression profiling, Aurora-kinase inhibitors as promising therapeutic option for newly-diagnosed patients can be tailoredly given to patients expressing *Aurora-A* with adverse prognosis.
Acknowledgments

We thank Véronique Pantesco, Katrin Heimlich, Maria Dörner and Margit Happich for technical assistance. This work was supported in part by grants from the Hopp-Foundation, Germany, the University of Heidelberg, Germany, the National Centre for Tumor Diseases, Heidelberg, Germany, the Tumorzentrum Heidelberg/Mannheim, Germany, the European Myeloma Stem Cell Network (MSCNET) funded within the 6th framework program of the European Community, the Deutsche Krebshilfe, Bonn, Germany, the Ligue Nationale Contre Le Cancer (équipe labellisée), Paris, France. It is also part of a national program called “Carte d’Identité des Tumeurs” (CIT) funded by the Ligue Nationale Contre le Cancer, France.
Authorship

Contribution: D.H. designed research, wrote the paper and participated in the microarray experiments; T. Meißner, A.B. and T.H. performed statistical analysis, A.S. performed experiments and participated in the writing of the paper; J.L., V.B., J.B. and J.Z. participated in the analysis of the data; J.F.R., U.B., J.H. and M.H. collected bone marrow samples and clinical data; B.B. and J.S. performed the total therapy 2-trial and J.S. the microarray experiments for the Arkansas-group; J.M., K.N. and A.K. participated in the writing and review of the paper; J.D.V participated in the microarray experiments; A.J. contributed in performing the interphase-FISH experiments; T.R., T. Möhler, B.K. and H.G. participated in the analyzing of the data and in the writing of the paper.

Conflict-of-interest disclosure: The authors declare no competing financial interests.

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References


Figure Legends

Figure 1. Expression of *Aurora-A* (*AURKA*), -B (*AURKB*), and -C (*AURKC*) as determined by gene expression profiling in memory B-cells (MBC), polyclonal plasmablastic cells (PPC), normal plasma cells (BMPC), myeloma cells (MMC), and myeloma cell lines (HMCL) within the (A) training and (B) validation group. An asterisk (*) indicates a significant difference of the indexed population compared to both, BMPC and MBC at a level of *P* < .001.

Figure 2. Validation of gene expression by quantitative real-time PCR, western blotting and flow cytometry. To validate gene expression data, quantitative real-time PCR for *Aurora-A*, -B, -C in (A1) 10 myeloma cell lines (HMCL) and (A2) 10 primary myeloma cells (pMMC) was performed. Shown are -dCt-values (reference gene 18S-RNA). (B) Gene expression data were further validated by western blotting. Shown are the blots of 10 cell lines for Aurora-A and -B with β-actin as loading control and HELA cells as positive control, respectively. (C) Intracytoplasmatic expression of Aurora-A and -B as determined by flow cytometry. Shown is the cell line OPM-2. Light grey line: control without primary antibody, black line: measurement with primary and secondary antibody.

Figure 3. Prognostic relevance of *Aurora-A* expression for two independent groups of patients treated with high-dose therapy and autologous stem cell transplantation. Shown are (A) event-free (EFS) and (B) overall survival (OAS) in our cohort of patients (left side) and the Arkansas-group (right side) for absence (black curve) vs. presence (grey curve) of *Aurora-A* expression in CD138-purified myeloma cells. Presence of *Aurora-A* expression in myeloma cells is an adverse prognostic factor in terms of EFS and OAS in both groups.

Figure 4. Inhibition of proliferation of myeloma cell lines as well as survival of primary myeloma cells and cells of the bone marrow microenvironment. (A1) Inhibition of proliferation of 20 myeloma cell lines by the pan-Aurora-kinase inhibitor VX680 in graded concentrations vs. medium- and DMSO-control, respectively, measured by ³H-thymidine uptake. Two independent experiments were performed in triplicates. (A2) The IC50 (in µM) and maximal inhibition at
10 µM (IMAX10 in %) are shown. (B1) Survival of primary myeloma cells (pMMC) cultured within their bone marrow microenvironment (negative fraction of plasma cell purification) is significantly inhibited compared to the medium control as determined by staining with anti-CD138-FITC antibody and propidium iodine. An asterisk (*) indicates a significant decrease between the medium control and the respective VX680 concentration. (B2) Survival of cells within the bone marrow microenvironment (BMME cells; negative fraction of plasma cell purification) was determined as described above for pMMC. An asterisk (*) indicates a significant decrease between the medium control and the respective VX680 concentration. (C) Induction of apoptosis by VX680 at 1 µM as determined by annexin V-staining after 8, 24, 48 and 72 h.
Figure 2.
**Figure 2.**

**B**

<table>
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<th>XG-1</th>
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<th>XG-11</th>
<th>XG-13</th>
<th>XG-19</th>
<th>LP-1</th>
<th>U266</th>
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</table>

**C**

![Histogram of Aurora-A and Aurora-B expression in OPM-2 cells](aurora-a_and_b_histogram.png)
Figure 3.

A  
EFS HM-group  

$P = .003$

B  
OAS HM-group  

$P = .03$

EFS Arkansas-group  

$P = .004$

OAS Arkansas-group  

$P = .008$
### Table A2

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Figure 4.

B1

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by Bernard Klein on March 20, 2009.
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From
Figure 4.

B2

**BMME 1**

**BMME 2**

**BMME 3**

**BMME 4**

**BMME 5**

Fraction of living BMME cells

VX680-concentration µM

medium DMSO  0.032  .16  .8  4  20  100

medium DMSO  0.032  .16  .8  4  20  100

medium DMSO  0.032  .16  .8  4  20  100

medium DMSO  0.032  .16  .8  4  20  100

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Figure 4.

C

<table>
<thead>
<tr>
<th>Time</th>
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<tr>
<td>8 h</td>
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<td>48 h</td>
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<td>72 h</td>
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Table 1. Presence of expression of Aurora-kinase A, -B, and -C.

<table>
<thead>
<tr>
<th>Gene Symbol</th>
<th>Probeset</th>
<th>MBC present [%] (n=13)</th>
<th>PPC present [%] (n=12)</th>
<th>BMPC present [%] (n=14)</th>
<th>MGUS present [%] (n=233)</th>
<th>MM present [%] (n=233)</th>
<th>HMCL present [%] (n=40)</th>
<th>ND-WBM present [%] (n=7)</th>
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Table 1. Presence of expression of Aurora-kinase A (AURKA), -B (AURKB), -C (AURKC) as judged by PANP in normal bone marrow plasma cells (BMPC), proliferating polyclonal plasmablastic cells (PPC), memory B-cells (MBC), multiple myeloma cells (MMC), cells from patients suffering from monoclonal gammopathy of unknown significance (MGUS), human myeloma cell lines (HMCL) and the bone marrow of normal donors (ND-WBM) as well as myeloma patients (MM-WBM).
Table 2. Presence of expression of Aurora-kinase A, -B and -C.

<table>
<thead>
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<th>Gene Symbol</th>
<th>Probeset</th>
<th>CD3 present [%] (n=5)</th>
<th>CD14 present [%] (n=5)</th>
<th>CD15 present [%] (n=5)</th>
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<th>MSC-ND present [%] (n=7)</th>
<th>MSC-MGUS present [%] (n=5)</th>
<th>MSC-MM present [%] (n=7)</th>
<th>OC present [%] (n=7)</th>
<th>Testis [%] (n=5)</th>
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Table 2. Presence of expression of *Aurora-kinase A* (*AURKA*), -B (*AURKB*), -C (*AURKC*) as judged by PANP in subfractions of the bone marrow (mesenchymal stromal cells (MSC) from normal donors (ND-MSC), patients with monoclonal gammopathy of unknown significance (MSC-MGUS) as well as myeloma patients (MSC-MM); osteoclast (OC)), and testis-samples, respectively.