Genomic arrays identify high-risk chronic lymphocytic leukemia with genomic complexity: a multicenter study

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ABSTRACT

Complex karyotype identified by chromosome-banding analysis has been shown to have prognostic value in chronic lymphocytic leukemia (CLL). Genomic arrays offer high-resolution genome-wide detection of copy-number alterations (CNA) and could therefore be well equipped to detect the presence of a complex karyotype. Current knowledge on genomic arrays in CLL is based on outcomes of single-cen-
ter studies, in which different cutoffs for CNA calling were used. To further determine the clinical utility of genomic arrays for CNA assessment in CLL diagnostics, we retrospectively analyzed 2,293 arrays from 13 diagnostic laboratories according to established standards. CNA were found outside regions captured by CLL fluorescence in situ hybridization probes in 34% of patients, and several of them, including gains of 8q, deletions of 9p and 18p (P<0.01), were linked to poor outcome after correction for multiple testing. Patients (n=972) could be divided into three distinct prognostic subgroups based on the number of CNA. In multivariable analysis only high genomic complexity, defined as ≥5 CNA, emerged as an independent adverse prognosticator for time to first treatment (hazard ratio: 2.15; 95% confidence interval: 1.36-3.41; P=0.001) and overall survival (hazard ratio: 2.54, 95% confidence interval: 1.54-4.17; P<0.001; n=528). Lowering the size cutoff to 1 Mb in 647 patients did not significantly improve risk assessment. Genomic arrays detected more chromosomal abnormalities and, in terms of risk stratification, performed at least as well as simultaneous chromosome banding analysis as carried out in 122 patients. Our findings indicate that genomic array is an accurate tool for CLL risk stratification.

**Introduction**

Chronic lymphocytic leukemia (CLL) is both clinically and genetically highly heterogeneous. The advent of efficacious, albeit expensive, targeted treatment regimens for CLL has highlighted the need for more accurate risk stratification. Diverse CLL-specific biomarkers are used to predict clinical course and survival in CLL including Rai1 and Binet staging, immunoglobulin heavy variable (IGHV) gene somatic hypermutation status,3,4 fluorescence in situ hybridization (FISH)-detected specific chromosomal alterations,5,6 and gene somatic hypermutation status.3,4 These observations demonstrate that genomic analysis of genetic aberrations provides valuable additional prognostic information compared to FISH analysis alone, hence refining risk-adapted stratification of patients.

Chromosome banding analysis (CBA) of in vitro mitogen-stimulated peripheral blood samples provides a low-resolution whole-genome view of the cytogenetic landscape of CLL, thereby overcoming the targeted nature of FISH which precludes a comprehensive assessment of the genomic landscape of CLL.7-9 Relatively small studies employing CBA in CLL identified the presence of a complex karyotype (CK; defined as ≥3 cytogenetically-visible structural and/or numerical aberrations) as a prognostic marker for refractoriness not only to chemoimmunotherapy,10,11 but also to novel targeted agents such as ibritinib12 and venetoclax.13 Within the European Research Initiative on CLL (ERIC), we recently performed a retrospective study of 5,290 patients with available CBA data and found that cytogenetic complexity with ≥5 CK patients, urged for a reappraisal of the performance of genomic arrays in 2,293 CLL patients from 13 European CLL laboratories.

Patients and established biomarkers

Overall, 2,293 CLL patients, diagnosed according to International Workshop on CLL criteria,26 were included in the present multicenter, retrospective study. Thirteen different CLL diagnostic laboratories connected to ERIC were involved (Online Supplementary Figure S1). In total, 572 of the 2,293 patients have been included in previous publications.19,27 Cases included for survival analyses were mostly recruited within 2 years (69.1%) or 1 year (59.1%) after diagnosis. An overview of the patients’ characteristics (demographics and biological features) is provided in Online Supplementary Table S1.

TP53 mutation analysis (Online Supplementary Table S2) and determination of the somatic hypermutation status of the clonotypic rearranged IGHV genes were performed and interpreted according to ERIC guidelines (Online Supplementary Methods).19,12 In a subgroup of patients (260/2,293) interphase FISH analysis...
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using probes for the regions 13q14 (BCL2 and/or DLEU1), 11q22.3 (ATM), 17p13.1 (TP53) and trisomy 12 was performed concurrently with genomic array analysis (i.e., same date of sampling). Simultaneous CBA (i.e., same date of sampling) was conducted in 122/2,293 patients. Metaphases for CBA were induced with stimulation protocols based on phorbol-12-myristate-13-acetate or immunostimulatory cytosine guanine dinucleotide-oligonucleotide DSP30 plus interleukin as previously described. The study was performed according to national and international ethical and legal recommendations with approval from the local Ethics Review Committees of the participating institutes.

Genomic array analysis

Genomic arrays were performed using several commercially available high-density (array-based comparative genomic hybridization/single nucleotide polymorphism array) platforms applied for diagnostics in the participating centers (Online Supplementary Table S3 and Online Supplementary Methods). CNA routinely assessed in CLL by targeted FISH revealed by genomic arrays in this study were included in all analyses irrespective of size, while for other CNA a size cutoff of ≥5 Mb was used according to Schoumans et al. Copy-number neutral loss of heterozygosity events, detected by single nucleotide polymorphism arrays, were not included in the counting of CNA. CNA with a distance ≥5 Mb between the two CNA, and putative chromothripsis events were counted as one event.

Statistical analysis

Statistical analysis was performed using SPSS version 24 and R. A Pearson χ2 test was used to evaluate the independence between categorical factors, and a Kruskal-Wallis test was applied to compare the distributions of continuous variables between groups of patients. Time-dependent receiver operating characteristic (ROC) curve analysis was performed evaluating different time-points from the date of array analysis (Online Supplementary Methods).

Overall survival (OS) and time to first treatment (TTFT) were calculated from the date of sampling for array analysis to the last date of follow-up. Only patients untreated at the date of sampling (n=972) were included in the survival analyses unless otherwise stated. The median follow-up was 44 months with a total of 353 survival events. Kaplan-Meier survival curves were compared by log-rank testing. Univariable Cox regression analysis was used to evaluate the effect of CNA, for which there were at least ten events and established clinico-biological features, on OS. Survival of cases with chromothripsis was also evaluated than that of cases with TP53abn or del(11q) without chromothripsis (P<0.03) (Online Supplementary Figure S3B).

We focused on the ten most frequent CNA other than del(13q), del(11q), trisomy 12 and del(17p) in our cohort (indicated by asterisks in Online Supplementary Figure S4) and investigated whether these CNA correlated with IGHV gene somatic hypermutation status and CNA captured by FISH probes applied in CLL. This resulted in several significant associations, such as (i) U-CLL with gain of 2p and 8q and loss of 6q (Online Supplementary Figure S5); (ii) del(11q) with gain of 2p and 8q and loss of 4p; (iii) trisomy 12 with loss of 14q; (iv) del(13q) with loss of 18p and 14q and, (v) del(17p) with gain of 3q and 8q, and loss of 4p, 8p, 15q and 18p (Online Supplementary Figure S6A-D). A summary of all significant correlations of CNA (corrected P<0.01) detected in the entire cohort with predefined CLL prognostic subgroups is presented in Table 1 and for all correlations in Online Supplementary Table S4.

In a subgroup of 260 patients with FISH and genomic array results available from the same sample, we observed 92.2% concordance (90/983 of regions analyzed by FISH) between the two approaches. Compared to genomic arrays, FISH detected statistically more cases of del(13q) (58.6% vs. 49.8%, P<0.001) and del(11q) (51.3% vs. 25.5%, P=0.003); while the detection rates of trisomy 12 (12.9% vs. 10.4%, P=0.07) and del(17p) (9.3 vs. 8.9%, P=1.00) were not statistically different (Online Supplementary Table S5). FISH-identified CNA not captured by genomic arrays (60/983) were mostly subclonal (median FISH clone size of 19%). *Vice versa,* 17/983 CNA were detected by genomic arrays only, including six cases of del(17p) (median size 19.11 Mb) and four of del(11q) (median size 17.67 Mb). To investigate the discriminatory power of FISH versus genomic arrays, a concordance index was calculated for genomic arrays versus FISH for del(11q) and del(17p) for both TTFT and OS. The higher number of del(11q) detected by FISH did result in a higher concordance index for TTFT (57% vs. 53%) while the concordance index for OS was 54% for both techniques. The concordance index for del(17p) was similar for genomic arrays and FISH for both TTFT (52% vs. 51%) and OS (57% vs. 55%). These data confirm that genomic
Figure 1. Overview of copy-number alterations detected by genomic arrays. (A) Percentages of patients with del(13)(q14), del(11)(q22.3) (ATM), trisomy 12 (+12) or del(17)(p13.1) (TP53) detected by genomic arrays irrespective of size. (B-E) Percentages of patients with different chromosomal losses (B), gains (C), monosomies (D) and trisomies (E). (F) Percentages of patients with putative chromothripsis events containing TP53abn in black and cases without TP53abn but del(11q)-positive in white. In four del(11q) cases, TP53 mutation status was not determined.
arrays show high concordance with FISH with the advantage that (other) potentially clinically relevant abnormalities can be elucidated on a genome-wide scale.

**Recurrent copy-number alterations detected by genomic arrays are associated with adverse outcome**

The effect of CNA detected by genomic arrays on OS was tested in untreated patients at the date of sampling (n=972), mostly within 2 years from diagnosis (69.1%; median time after diagnosis, 7 months). A significant impact of well-established risk factors for CLL, including Binet B/C, U-CLL, TP53abn and del(11q) (Figure 2 and Online Supplementary Figure S7) on OS was confirmed. Next, we analyzed the effects of CNA outside the established CLL regions which were detected in at least ten cases (Figure 2). Minimal common regions of deletion or amplification are presented in Online Supplementary Table S6. Gains of 8q (P=0.01; encompassing MYC in >95% of cases), loss of 9p (P<0.001; involving SMARCA2 [46%] and CDKN2A [85%]) and 18p (P<0.001; including USP14 [100%]) were significantly associated with a shorter OS in univariable Cox regression analysis after correction for multiple testing.

**Patients with chronic lymphocytic leukemia can be grouped into three prognostic subgroups based on copy-number alteration profile**

ROC curve analysis revealed that based on the total number of CNA above the 5 Mb cutoff, including aneuploidy events (see the Online Supplementary Methods for details of this analysis), the cohort could be subdivided into three prognostic subgroups: low-genomic complexity (GC) (0-2 CNA; n=793), intermediate-GC (3-4 CNA; n=122) and high-GC (≥5 CNA; n=57). The results of the ROC curve analysis were replicated by a different statistical approach, using maximally selected rank statistics (see the Online Supplementary Methods for details of this analysis).32

Results were supported with bootstrapping based on 100 random bootstrap samples from the original sample (with replacement) and showed relatively high percentages for the Youden index and the maximally selected estimated cut-point (Online Supplementary Methods). Low-GC cases were associated with indolent disease: Binet A, IGHV mutated-CLL, and low incidences of TP53abn and del(11q) (P<0.01). Intermediate-GC and high-GC cases were associated with a more advanced clinical stage compared to low-GC cases (P=0.003). High-GC cases were enriched for TP53abn and U-CLL (P<0.001) compared to intermediate-GC and low-GC cases (Table 2). Calculation of associations of established biomarkers and CNA with prognostic subgroups identified differences between the prognostic GC subgroups (Figure 3). Interestingly, loss of 9p and loss of 15q were strongly associated with high-GC (P<0.001), not significantly associated with intermediate-GC and negatively associated with low-GC (P<0.001). A positive correlation with high-GC was observed for all CNA except for trisomies 12,18 and 19.

Low-GC was associated with a more favorable clinical outcome (median OS: 10.17 years, 95% CI: 8.92-11.42 years) compared to the outcomes of cases with intermediate-GC (median OS: 7.02 years, 95% CI: 4.79-9.25 years, P=0.001) and high-GC, who had the shortest OS (median OS: 4.79 years, 95% CI: 1.14-4.96 years, P<0.001) (Figure 4A). Similar results were observed for TTFT (Figure 4B) between the three different prognostic subgroups (P=0.001).

**High genomic complexity is an independent prognostic risk factor in chronic lymphocytic leukemia**

Univariable Cox regression analysis revealed that the GC subgroups intermediate-GC and high-GC (P<0.001) as well as established CLL prognostic factors – male sex, age

### Table 1. Correlations of copy-number alterations with predefined prognostic subgroups of chronic lymphocytic leukemia.

<table>
<thead>
<tr>
<th>CLL subgroup</th>
<th>losses: 13q14*, 11q22.3, 17p13.1, 14q, 6q, 8p, 13q,other, 4p</th>
<th>gains: 2p, 8q</th>
<th>‘trisomies: none</th>
<th>‘chromothripsis: none</th>
</tr>
</thead>
<tbody>
<tr>
<td>del(11)(q22.3)</td>
<td>losses: 8p, 18p, 4p, 3q, 11p, 4q, 12p, 6q</td>
<td>gains: 2p, 8q, 22q, 21q, 7p, 6q</td>
<td>trisomies: 12*, 22</td>
<td>chromothripsis: none</td>
</tr>
<tr>
<td>trisomy 12</td>
<td>losses: 13q14*, 11q22.3*, 14q, 8p*</td>
<td>gains: 2p*</td>
<td>trisomies: 18, 18</td>
<td>chromothripsis: none</td>
</tr>
<tr>
<td>del(13)(q14)</td>
<td>losses: 14q*, 13q,other, 18p</td>
<td>gains: 17q*, 13q</td>
<td>trisomies: 12*</td>
<td>chromothripsis: none</td>
</tr>
<tr>
<td>del(17)(p13.1)</td>
<td>losses: 8p, 18p, 4p, 15q, 9p, 3p, 4q, 13q,other, 6p, 2q, 20p, 10q, 9q, 18q, 2p, 19p, 10p, 5p, 9q, 17q, Xp, 9, 21q, 13, Yq, 7q, 1p, 11p, 14q</td>
<td>gains: 8q, 3q, 17q, 15q, 11q, 3p, 5q, 1p, 13q, 11p</td>
<td>trisomies: none</td>
<td>chromothripsis: 8, 5, 6, 17</td>
</tr>
</tbody>
</table>

Significantly correlated copy-number abnormalities (corrected for multiple testing; P<0.01). Negative correlations are indicated by an asterisk (*). CLL: chronic lymphocytic leukemia; U-CLL: with unmutated immunoglobulin heavy chain variable gene (IGHV).
>70, Binet stage B/C disease, U-CLL, TP53abn and del(11q) – were associated with adverse outcome (P<0.05). (Online Supplementary Tables S7 and S8). High-GC cases had significantly shorter TTFT and OS in the CLL subsets U-CLL (P<0.001), IGHV gene-mutated CLL (P=0.001) and TP53abn/del(11q) (P=0.01) compared to low-GC cases (Online Supplementary Figures S8-S10).

Furthermore, including the established factors – male gender, age >70, Binet stage B/C, U-CLL, TP53abn and del(11q) – in the multivariable Cox regression model, high-GC retained statistical significance as a binary predictor (with the categories ≥5 CNA and <5 CNA) including the above-mentioned factors in the multivariable Cox regression model for TTFT and OS (P=0.002) (Online Supplementary Tables S9 and S10).

Lower copy-number alteration size cutoff does not significantly improve risk stratification

In routine clinical practice, a 5 Mb threshold for reporting CNA other than del(13q), del(11q) and del(17p) is used, as described in the publication of Schoumans et al. In order to identify whether the recommended 5 Mb threshold or a 1 Mb threshold may be a better discriminator for prognostication, a subgroup of genomic arrays (n=647; with both 1 Mb and 5 Mb cutoff results available) was analyzed applying both cutoffs. Lowering the size cutoff to 1 Mb resulted in the detection of 290 additional

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**Subgroup** | **nr. of patients analyzed** | **Positive cases analyzed** | **Hazard ratio for OS (95% CI)**
--- | --- | --- | ---
Established Biomarkers | | | |
U-CLL | 628 | 286 | 4.04 (3.16-5.17)
TP53abn | 749 | 82 | 2.73 (2.01-3.70)
Binet B/C | 913 | 328 | 2.17 (1.74-2.69)
Established regions | | | |
Loss.17p13.1 | 961 | 56 | 3.68 (2.56-5.29)
Loss.11q22.3 | 961 | 158 | 2.04 (1.60-2.61)
TRIS.12 | 961 | 115 | 1.22 (0.88-1.71)
Loss.13q14 | 961 | 519 | 0.88 (0.71-1.08)
Other | | | |
Loss.9p | 961 | 17 | 3.24 (1.82-5.77)
Loss.18p | 961 | 31 | 3.18 (1.97-5.12)
Gain.17q | 961 | 13 | 2.50 (1.11-5.62)
Gain.8q | 961 | 34 | 2.46 (1.49-4.07)
Loss.4p | 961 | 21 | 2.23 (1.22-4.07)
Loss.20p | 961 | 14 | 2.22 (1.10-4.48)
Gain.2p | 961 | 58 | 1.84 (1.23-2.75)
Loss.6q | 961 | 31 | 1.81 (0.99-3.30)
Loss.14q | 961 | 38 | 1.79 (1.00-3.20)
Loss.15q | 961 | 11 | 1.68 (0.75-3.78)
Gain.3q | 961 | 14 | 1.50 (0.67-3.37)
Loss.8p | 961 | 29 | 1.45 (0.77-2.71)
Gain.13q | 961 | 12 | 1.32 (0.42-4.12)
Loss.1q | 961 | 16 | 1.26 (0.56-2.81)
Loss.13q.other | 961 | 13 | 0.74 (0.24-2.31)
TRIS.19 | 961 | 11 | 0.46 (0.12-1.86)
TRIS.18 | 961 | 10 | 0.43 (0.11-1.73)

Figure 2. Effect of copy-number alterations on clinical outcome in chronic lymphocytic leukemia. Forest plot representing the results of univariable Cox regression analysis of copy-number alterations detected by array (with at least 10 events) and the effect of chronic lymphocytic leukemia (CLL) prognostic factors. A hazard ratio of less than 1.00 indicates a lower risk of overall survival. The size of each square is proportional to the amount of data available. U-CLL denotes patients with an unmutilated immunoglobulin heavy chain variable gene and TP53abn denotes patients with del(17p) and/or a TP53 mutation. Loss.13q,other are patients with a del(13q) not containing the established 13q14 minimally deleted region detectable by the diagnostic fluorescence in situ hybridization probe, and TRIS denotes patients with a trisomy. Only patients untreated at the date of DNA sampling were included for survival analysis. 95% CI: 95% confidence interval.
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Table 2. Correlations of the three genomic complexity subgroups (based on the complexity of the array profile) with other chronic lymphocytic leukemia risk factors

<table>
<thead>
<tr>
<th>N=972</th>
<th>Low-GC</th>
<th>Intermediate-GC</th>
<th>High-GC</th>
<th>P-values</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male</td>
<td>500/761, 66%</td>
<td>81/115, 70%</td>
<td>35/53, 66%</td>
<td>0.606</td>
</tr>
<tr>
<td>&gt;70 years</td>
<td>179/793, 23%</td>
<td>31/122, 25%</td>
<td>20/57, 35%</td>
<td>0.089</td>
</tr>
<tr>
<td>Binet B/C</td>
<td>256/752, 33%</td>
<td>55/118, 47%</td>
<td>26/54, 48%</td>
<td>0.003</td>
</tr>
<tr>
<td>U-CLL</td>
<td>212/515, 41%</td>
<td>50/81, 62%</td>
<td>23/33, 76%</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>TP53abn†</td>
<td>29/602, 5%</td>
<td>26/102, 25%</td>
<td>27/54, 50%</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>del(11)q(11q23)</td>
<td>84/793, 11%</td>
<td>52/122, 43%</td>
<td>26/57, 46%</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

Table 2: Correlations of the three genomic complexity subgroups (based on the complexity of the array profile) with other chronic lymphocytic leukemia risk factors.

-abn: del(17)q(p11.1) and/or TP53 mutation. GC: genomic complexity; U-CLL: chronic lymphocytic leukemia with unmutated IGHV genes. GC categories: low-GC=0-2 copy-number alterations (CNA); intermediate-GC=3-4 CNA; high-GC=≥5 CNA detected by array. A Pearson χ² test was used to calculate the P-values with a combination of subgroups.

Discussion

In our study CLL patients with ≥5 CNA emerged as a separate subgroup with aggressive disease and an adverse outcome, which was independent of several well-established factors in CLL, such as IGHV and TP53 status. A disadvantage of a collection of retrospective multicenter data is the risk of variation introduced by combining results from different genomic array platforms. Nevertheless, marked variation in this study is less likely since the data were methodically accepted and uniformly interpreted using commonly accepted guidelines which provide tools for filtering and calling CNA and single nucleotide polymorphism data. Another clear disadvantage of such real-world retrospective data is that clinical correlations, although indicative, must be considered with great caution as treatments were not uniform and we were limited to the date of sample collection (instead of diagnosis) for the TTFT analysis. In current International Working Group CLL guidelines, targeted FISH is recommended for the detection of genomic aberrations. In our multicenter study overall concordance between array-based comparative genomic hybridization data and targeted FISH results (subgroup, n=260) was high (>90%).

Nevertheless, genomic arrays resulted in the detection of fewer CNA for regions captured by FISH, which could be explained by the superior detection limit of FISH. However, the increased detection of CNA by FISH was not directly associated with a better risk assessment, in agreement with the findings in other genomic array stud-
ies, providing a rationale that genomic arrays can be used, besides FISH, in CLL diagnostics. Direct comparison of genomic arrays with FISH in prospective clinical trials is necessary before genomic arrays can be accepted to replace FISH in routine clinical practice for CLL prognostics.

Since our data were collected retrospectively from patients during follow-up after various chemoimmunotherapy regimens, we could not correlate the effects of CNA on OS with specific treatments. CNA such as del(4p), del(9p) and del(18p) co-occurred with other CNA and were associated with other CLL biomarkers. For this reason, the prognostic impact of these individual CNA must be considered with caution. Furthermore, the applicability of our results needs to be established for novel agents, as the prognostic value following chemoimmunotherapy might be different. This is exemplified by a recent update of the RESONATE-2 study in which

Figure 4. Effect of genomic complexity on clinical outcome in chronic lymphocytic leukemia. (A, B) Kaplan-Meier plots presenting the overall survival (A) and time to first treatment (B) of patients with chronic lymphocytic leukemia divided into three categories of genomic complexity (GC) based on the total number of copy-number alterations (CNA): low-GC (0-2 CNA), intermediate-GC (3-4 CNA) and high-GC (≥5 CNA) and defined by receiver operating characteristic curve analysis. 95% CI: 95% confidence interval.
del(11q) was prognostically favorable in ibrutinib-treated patients.9,10 Additionally, del(9p) was recently correlated with resistance to ibrutinib.47 The inferior outcome of patients with loss of 9p is in line with the findings of a recent study on the effect of loss of 9p24.3 containing SMARCA2.48 Loss of 9p24.3 and/or mutations in the SWI-SNF chromatin remodeling complex were identified in all nonresponders to ibrutinib and venetoclax. Deletion of SMARCA2 results in upregulation of the prosurvival protein BCL-XL which could protect cells from apoptosis.49

The differences in OS and TTFT in the distinct GC subcategories based on ROC analysis in this study are in agreement with recent findings regarding CBA-defined CK and the independent effect of high-GC (with ≥5 numerical and/or structural abnormalities),12 which were characterized by a distinct distribution of aberrations affecting a broad spectrum of chromosomes.12 Similarly, high-GC detected by genomic arrays was negatively associated with trisomy 12 and, in contrast to intermediate-GC, not positively associated with trisomy 18 and 19 (Figure 3). High-CК cases detected by CBA and lacking del(17p) or a Sanger-detected (‘clonal’) TP53 mutation were all negative for subclonal TP53 mutations when sequenced by next-generation sequencing.50 It is conceivable that recently identified potential drivers for high-GC in CK, such as SAMHD1 and SETD2, attributed to high-GC in a subset of patients.9,42-44 The resolutions of the genomic array platforms used in this study are higher than the resolution that can be achieved by CBA. Differences in the detection of chromosomal aberrations between genomic arrays and CBA in our study could be related to the lower resolution of CBA and/or the use of phosphor-12-myristate-13-acetate in most CBA cases instead of the more sensitive method involving cytosine guanine dinucleotides plus interleukin 2.12 In contrast to genomic arrays, CBA selects for cells with proliferative potential as a prognostic marker and is able to detect balanced translocations. Although balanced rearrangements (such as balanced translocations) cannot be detected by genomic arrays, chromosome regions involved in unbalanced translocations can be recognized more precisely by arrays, possibly revealing new recurrent abnormalities. Our finding that, in terms of risk stratification, genomic arrays do not underperform compared with CBA implies that the inability of genomic arrays to detect balanced rearrangements does not impede adequate risk assessment. This could be explained by the fact that, in contrast to other mature B-cell tumors, CLL is defined by the presence of CNA with a paucity of balanced rearrangements.3,46

In CLL diagnostics genomic arrays are cost-effective as a stand-alone method when compared to FISH plus CBA. Only recently, the integration of genomic arrays has been recommended by the Cancer Genomics Consortium working group as having evidence-based clinical utility for a broad spectrum of hematologic malignancies.3,46-49

In conclusion, our study highlights the strengths of genome-wide approaches for risk stratification in CLL and we identified CLL patients with high-GC, defined as ≥5 CNA detected by genomic arrays, as an independent subgroup with dismal clinical outcome. We, therefore, recommend implementation of comprehensive genomic profiling in future clinical trials.

**Disclosures**

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**Contributions**


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**Table 3.** Multivariable analysis for time to first treatment.

<table>
<thead>
<tr>
<th></th>
<th>HR</th>
<th>95% CI</th>
<th>P-values</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male</td>
<td>1.05</td>
<td>0.84-1.32</td>
<td>0.674</td>
</tr>
<tr>
<td>&gt;70 years</td>
<td>1.10</td>
<td>0.83-1.46</td>
<td>0.490</td>
</tr>
<tr>
<td>Binet/BC</td>
<td>3.37</td>
<td>3.04-4.94</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>U-CLL*</td>
<td>3.25</td>
<td>2.51-4.21</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>7p53ab1</td>
<td>1.12</td>
<td>0.76-1.65</td>
<td>0.567</td>
</tr>
<tr>
<td>del(11q)(q22.3)</td>
<td>1.22</td>
<td>0.92-1.63</td>
<td>0.166</td>
</tr>
<tr>
<td>GC (3 categories)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intermediate-GC vs. low-GC</td>
<td>1.00</td>
<td>0.73-1.38</td>
<td>0.984</td>
</tr>
<tr>
<td>High-GC vs. low-GC</td>
<td>2.15</td>
<td>1.36-3.41</td>
<td>0.001</td>
</tr>
</tbody>
</table>

1 TP53ab1 = del(17)(p13.1) and/or TP53 mutation. HR: hazard ratio; 95% CI: 95% confidence interval of the hazard ratio; U-CLL: chronic lymphocytic leukemia with unmutated IGHV genes; GC: genomic complexity; GC categories: low-GC=0-2 copy-number alterations (CNA); intermediate-GC=3-4 CNA; high-GC=≥5 CNA detected by array.

**Table 4.** Multivariable analysis for overall survival.

<table>
<thead>
<tr>
<th></th>
<th>HR</th>
<th>95% CI</th>
<th>P-values</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male</td>
<td>1.26</td>
<td>0.96-1.64</td>
<td>0.094</td>
</tr>
<tr>
<td>&gt;70 years</td>
<td>2.48</td>
<td>1.86-3.31</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Binet/BC</td>
<td>1.47</td>
<td>1.13-1.91</td>
<td>0.004</td>
</tr>
<tr>
<td>U-CLL*</td>
<td>3.84</td>
<td>2.85-5.17</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>7p53ab1</td>
<td>1.56</td>
<td>1.03-2.36</td>
<td>0.037</td>
</tr>
<tr>
<td>del(11q)(q22.3)</td>
<td>0.92</td>
<td>0.67-1.26</td>
<td>0.605</td>
</tr>
<tr>
<td>GC (3 categories)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intermediate-GC vs. low-GC</td>
<td>1.19</td>
<td>0.81-1.73</td>
<td>0.373</td>
</tr>
<tr>
<td>High-GC vs. low-GC</td>
<td>2.54</td>
<td>1.54-4.17</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

1 TP53ab1 = del(17)(p13.1) and/or TP53 mutation. HR: hazard ratio; 95% CI: 95% confidence interval of the hazard ratio; U-CLL: chronic lymphocytic leukemia with unmutated IGHV genes; GC: genomic complexity; GC categories: low-GC=0-2 copy-number alterations (CNA); intermediate-GC=3-4 CNA; high-GC=≥5 CNA detected by array.
A.C.L., PB, KS, DO, JCS, RR, and APK wrote the first draft; and all authors were involved in revising the manuscript and approved the final version.

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