

A recurrent 11q aberration pattern characterizes a subset of *MYC*-negative high-grade B-cell lymphomas resembling Burkitt lymphoma

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Introduction

Burkitt lymphoma (BL) is an aggressive B-cell lymphoma characterized by typical morphological, immunophenotypic, and molecular features.¹ The t(8;14)(q24;q32) hallmark translocation or its variants,

which juxtapose the *MYC* oncogene to one of the 3 immunoglobulin (*IG*) loci, is detectable by cytogenetic or, currently, molecular cytogenetic techniques in almost all cases of BL.²⁻⁴

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There is an ongoing discussion about whether BL is without exception defined by the presence of an *IG-MYC* translocation as the driving mutation or whether a small subset of true BL that lacks an *IG-MYC* translocation or even a *MYC* breakpoint does exist.^{1,5} In the latter case, it could be speculated that alternative mechanisms substitute for the hallmark *MYC* activation, pathogenomic for the vast majority of BL.^{4,6} A recent study on *MYC*-negative endemic BL has suggested that downregulation of microRNA hsa-mir-34b could be such a pathogenetic mechanism substituting *MYC* translocations.⁵ Alternatively, a common cell of origin could determine the typical features of BL in *MYC*-positive and -negative cases.

Various studies have reported lymphomas with clinical, histomorphologic, immunophenotypic, or gene expression features of BL lacking a detectable *MYC* translocation by fluorescence in situ hybridization (FISH).^{5,7-9} Nevertheless, caution is warranted before concluding that these indeed represent true *MYC*-negative BL, because the scattering of breakpoints in the *MYC* and *IG* loci along with small insertions of one locus into the other can render *MYC* breaks undetectable even if an extensive set of FISH probes is applied.¹⁰ Moreover, there is no gold standard to define BL in the absence of an *MYC* break. Prototypic BL that fulfills all characteristics of the World Health Organization (WHO) classification are rare.^{1,7} This in turn makes the distinction between BL and so called “intermediate lymphomas between Burkitt and diffuse large B-cell lymphomas (DLBCL)” difficult.¹ In a recent cytogenetic study, Pienkowska-Grela et al described 4 lymphomas in young adult patients displaying typical BL morphology that bona fide lacked an *MYC* rearrangement by FISH and chromosome analyses. Remarkably, these cases shared a recurrent abnormality within chromosome 11 described as dup(11)(q23q13). This result led them to reclassify the cases as intermediate between DLBCL and BL because of the immunophenotypical and genetic features.⁸

Similarly, Poirel et al observed by cytogenetic review of childhood mature B-cell non-Hodgkin lymphomas that the 13 cases classified as BL without a detectable *MYC* translocation shared alterations with the *MYC*-positive BL except for the existence of more frequent aneuploidy and the presence of a derivative chromosome 11q.¹¹

In the present study, we investigated the incidence of *MYC*-negative BL defined by gene expression analysis using 2 different classifiers. These cases were analyzed by histologic, immunophenotypic, and various molecular means. We show that the very rare cases of these *MYC*-negative BL display a recurrent pattern of alterations on chromosome 11q. We subsequently show that this pattern was also present in a subset of high-grade B-cell lymphomas with features of, but not ultimately fulfilling, the diagnostic criteria of BL. Clinical, morphologic, and molecular characterizations of these lymphomas with the typical 11q abnormality patterns suggest that these represent a distinct subset of *MYC*-negative high-grade B-cell lymphomas with features resembling but not identical to BL, and identify candidate genes potentially involved in their pathogenesis.

Material and methods

Lymphoma materials

A global overview on the cohorts studied and the selection of cases included in the present study is given in Figure 1. This study was performed as part of the Molecular Mechanisms in Malignant Lymphoma (MMML) Network Project of the Deutsche Krebshilfe, for which approval was obtained by the Institutional Review Board of the Medical Faculty Kiel under 403/05 as well as the study central. The protocols of the Berlin-Frankfurt-Münster (BFM)

clinical trials have also been approved by central and local review boards. This study was conducted in accordance with the Declaration of Helsinki.

Cohort 1: lymphomas called by both GEP classifiers (mBL and BL-PAP classifiers) BLs (n = 59)

To identify *MYC*-negative lymphomas with a BL gene expression profiling (GEP), 2 independently developed gene expression classifiers, the molecular-BL (mBL) index⁷ and the BL-pathway activation pattern (BL-PAP classifier)¹² were applied to a cohort of 753 previously characterized lymphomas from the MMML Network. The 2 algorithms concordantly selected a total of 59 BL by GEP (cohort 1). Two of these cases lacked a *MYC* translocation and were thus considered true *MYC*-negative BL.

Cohort 2: high-grade B-cell lymphomas and cell lines with features of BL, *MYC* negative (n = 14)

Twelve *MYC*-negative lymphomas, 5 of which were previously reported,^{8,13,14} and SU-DHL-5 and HT cell lines, derived from *MYC*-negative high-grade B-cell lymphomas (www.dsmz.de), were included for cytogenetic and genetic analyses. Information on case selection is detailed in the supplemental Methods and in supplemental Table 1, available on the *Blood* Web site. All biopsy specimens were evaluated by at least 2 hematopathologists according to the WHO classification (Table 1; supplemental Table 2).¹ Case 10, although its cytology showed features of BL, was initially classified as follicular lymphoma grade 3 with simultaneous DLBCL (supplemental Figure 1).^{13,14}

Minimal regions of gain and loss in chromosome 11q were defined in this series.

Cohort 3: MMML cases with 11q gain/loss pattern determined by CGH-array (n = 6, including 2 cases from cohort 1 and 1 case from cohort 2)

A total of 514 MMML cases with available comparative genomic hybridization (CGH)-array data were screened for aberrations in the minimal regions previously described in cohort 2. Six cases showing at least a partial gain and a partial loss within these regions were selected for cohort 3. Cases with non-mBL GEP, relapsed samples, and DLBCL cases *IG/MYC*-positive, *IGH/BCL2*-positive DLBCL, and *BCL6* break-positive were excluded. Gene expression and immunohistochemical and genetic features of these cases were compared with 2 different sets of reference samples. The first set included all samples that exhibited an mBL GEP and *IG-MYC* positivity (n = 46). The second set was composed of DLBCL samples (n = 198). All cases selected for these 2 sets had no imbalances in the minimal 11q regions.

Immunohistochemistry, conventional cytogenetics, and FISH

Immunohistochemical, conventional cytogenetic, and FISH analyses were performed as previously described.¹⁵⁻¹⁷ A description of the FISH probes is summarized in the supplemental Methods. Methods on cases 4 through 7 have been previously published.⁸

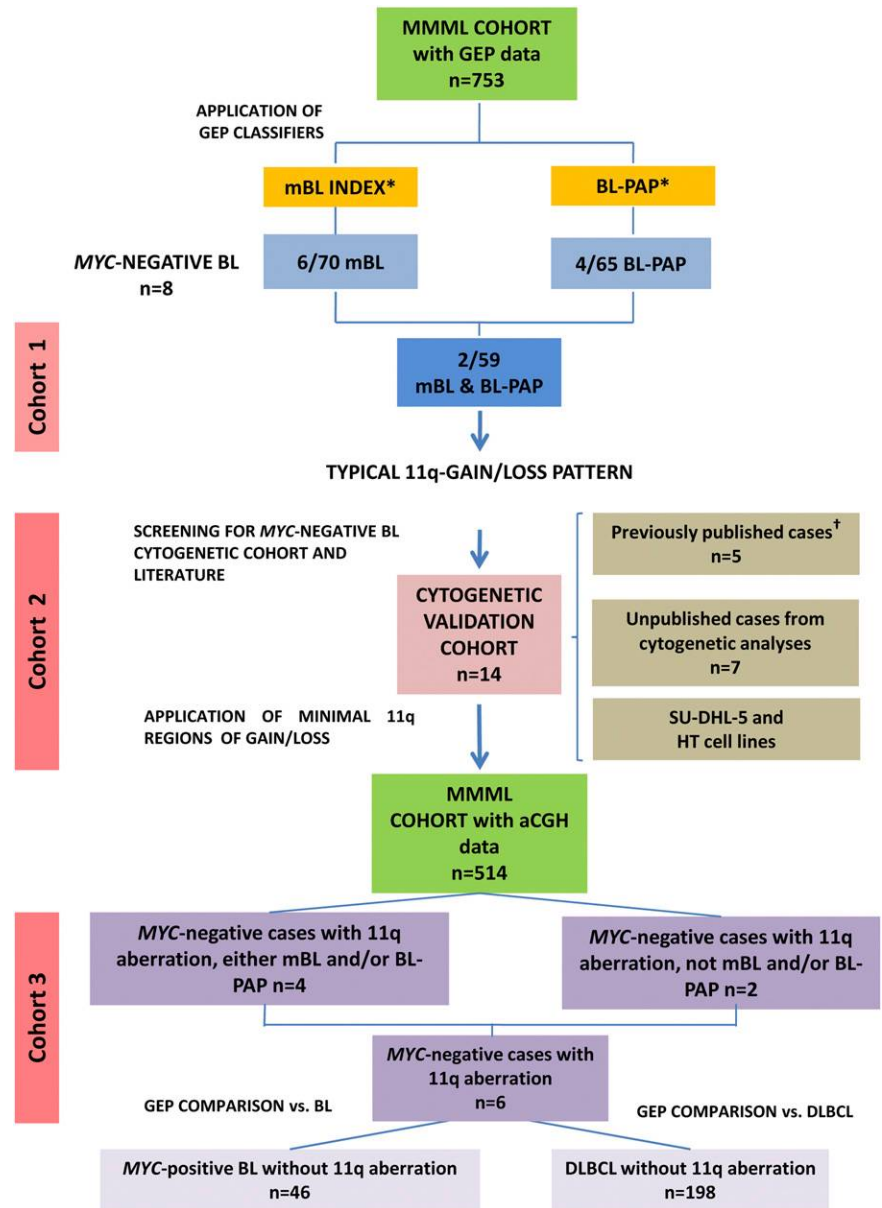
Copy number and mutational analyses

SNP 6.0 array (Affymetrix, Santa Clara, CA) was used in cases 1 through 3 and 8 and cell lines as previously described.¹⁸ Cases 4 through 7 and 9 through 12 were analyzed using Agilent Human CGH Microarray platforms owing to methodologic requirements for formalin fixed paraffin embedded tissues (GEO-Number GSE47508). Gains and losses were defined by using different software (see supplemental Methods). Mutational analysis of the *ATM* gene in case 1 was performed as previously described.^{19,20} *FLII* and *ETS1* genes' mutational analyses were carried out in selected samples and cell lines (supplemental Table 3). *ID3* mutational analysis was performed as previously described.²¹

Bisulfite pyrosequencing analyses

The hsa-mir-34b methylation pattern was determined by bisulfite pyrosequencing on cases 1 through 7, selected *IG-MYC* cell lines (CA-46, DAUDI, and RAJI), and primary *IG-MYC* BL samples (n = 6) according to standard

Figure 1. Diagram of the strategy used for the identification of potential *MYC*-negative BL. One case from cohort 2 is also part of cohort 3 (MMML series). *mBL⁷ and BL-PAP GEP classifiers.¹² †Previously published cases.^{8,13}



protocols. The results were evaluated with Pyro Q-CpG 1.0.9 software (Biotage AB, Uppsala, Sweden) (supplemental Table 3).

Custom exome sequencing analysis in cell lines

Custom exome sequencing was performed in SU-DHL-5 and HT cell lines outsourced to Aros (www.arosab.com/). Details on analysis are provided in the supplemental Methods.

Gene expression analyses

To evaluate the impact of 11q alterations in the GEP, differential gene expression analysis was performed as previously described. Details on the analysis are provided in the supplemental Methods.

Statistical methods

Comparison of immunohistochemical and genetic features between cases with aberrant 11q-gain/loss pattern and each reference set individually was performed using methods previously described.⁷ Genetic complexity was defined as the number of observed copy number (CN) aberrations per tumor sample. Complexity was compared between groups by the Kruskal-Wallis rank-sum test.

MicroRNAs target gene analysis

Test for overrepresentation of microRNA targets within a list of differentially expressed genes was performed in multiple hypergeometric tests as implemented in the HTSanalyzeR package (version 2.8.0).²²

Western blot analysis

Protein extract preparation and western blot experiments were performed as previously described using standard procedures. Details are provided in the supplemental Methods.

Results

MYC-negative lymphomas with a gene expression profile of BL are rare (cohort 1)

To identify BL, we applied 2 different GEP classifiers for BL, the mBL index⁷ and the BL-PAP classifier¹², to a total of 753 aggressive

Table 1. Immunohistochemical and clinical characteristics of the 12 *MYC*-negative B-cell lymphomas and SU-DHL-5 and HT cell lines (cohort 2)

Case no.	Age, gender	Morphologic diagnosis	Localization	Immunohistochemistry									Treatment protocol	Outcome, follow-up	Pathology panel
				MYC	CD20	CD10	CD5	BCL2	BCL6	MUM1	Ki67	EBER			
Case 1	7, male	Aggressive B-NHL, BL	LN(c)	+	+	+	-	-	+	-	90%	-	B-NHL BFM 04, treatment modified due to AT	In remission, alive, 62 mo	WK, IO, MMML panel
Case 2	7, male	BL	LN (colon)	-	+	+	-	-	+	na	95%	-	NHL-BFM-95	In remission, alive, 146 mo	WK, IO
Case 3	16, female	High-grade B-NHL, atypical BL	LN	-	+	+	na	+	+	+	95%	-	NHL-BFM-95	In remission, alive, 43 mo	WK, IO
Case 4	18, male	BL	LN(c)		+	+	-	-	+	na	>95%	-	GMALL 2008	In complete remission, alive, 48 mo	WK, GR
Case 5	22, male	BL	LN (ing)		+	+	-	-	-	na	>95%	-	GMALL 2002	In complete remission, alive, 36 mo	WK, GR
Case 6	25, male	BL	LN (ab)		+	+	-	-	+	na	>95%	-	CODOX-m 2003	Dead (progression), 12 mo	WK, GR
Case 7	37, female	BL	LN(c),T		+	+	-	-	+	na	>95%	-	GMALL 2002	In complete remission, alive, 72 mo	WK, GR
Case 8	11, male	Mature B-cell lymphoma/lymphoblastic B-lymphoma	LN(c)		+	+	-	-	+	na	>95%	na	EORTC 58951	Alive, 3 mo	WK, EJ
Case 9	12, male	BL	leg	+	+	+	-	-	+	-	>95%	-	B-NHL BFM 04	Alive, 8 mo	WK, IO
Case 10	6, female	FL	LN (c)		+	+	na	-	+	-	>95%	-	Ritux + B-NHL BFM 04	Alive, 24 mo	WK, IO
Case 11	9, male	Atypical BL	LN(ing)		+	+	-	-	+	-	>95%	-	COG-ANHL01P1	In complete remission, alive, 10 mo	WK, EJ
Case 12	76, male	BL	LN (c)	-	+	+	na	w+	na	na	>95%	na		Alive, na	WK, IO
SU-DHL-5	17, female	B-NHL (DLBCL)			+	+				+		-			
HT	70, male	B-NHL (DLBCL)			+	+				+		-			

-, negative; +, positive; ab, abdomen; c, cervical; EBER, Epstein-Barr–encoded RNA test; FL, follicular lymphoma; ing, inguinal; LN, lymph node; na, not available; NHL, non-Hodgkin lymphoma; T, tonsil; w, weak.

Cases 4, 5, 6, and 7 correspond to cases 1, 2, 3, and 4 from Pienkowska-Grela et al.⁸ Case 10 displayed a simultaneously follicular and diffuse grown pattern. Although its cytology showed features of BL, it was classified as FL grade 3 (minor component) with simultaneous DLBCL (major compartment), and it was included in a previous series of pediatric lymphomas already published; corresponds to case 23 from Oschlies et al.¹³ and pFL8 from Martin-Guerrero et al.¹⁴ Information about SU-DHL-5 and HT cell lines was obtained from DSMZ (www.DSMZ.de). Lymphoma samples were scored positive if >25% of the tumor cells stained positive except for MYC, where the cutoff was established in 40%.

lymphomas included in the cohort of the MMML network. Based on the mBL and BL-PAP indices, we classified 70 and 65 lymphomas, respectively, as molecularly defined BL. An *IG-MYC* fusion was identified in 63 of 70 (90%) mBL and 60 of 65 (92%) BL-PAP. One additional case identified by both classifiers carried a non-*IG-MYC* translocation, leaving 6 of 70 mBL and 4 of 65 BL-PAP lacking a *MYC*-translocation.

Supervised analysis comparing *IG-MYC*-positive and *MYC*-negative BL revealed 708 and 266 genes differentially expressed in the mBL and BL-PAP groups, respectively (false discovery rate = .05). The *MYC* gene itself was significantly lower expressed at the transcript level in both *MYC*-negative mBL and *MYC*-negative BL-PAPs compared with their *MYC*-positive counterparts ($P = .014$) (Figure 2).

The thorough gene expression analyses left only 2 *MYC*-negative lymphomas (MPI-086, MPI-382) from a total of 59 lymphomas,

which were called “BL” by both GEP classifiers applied. Interestingly, both cases were diagnosed in young males aged 14 and 24 years, showed a germinal center B (GCB)-like cell of origin gene expression signature, and were initially diagnosed as BL but reclassified by the pathology panel as high-grade B-cell lymphoma, not otherwise specified (NOS), and DLBCL, centroblastic, respectively. They had nodal presentation and the immunophenotype was that typical of BL. Both cases lacked the t(14;18)*IGH-BCL2* fusion, *BCL6* breaks, or *MALT1* breaks by FISH. Moreover, both shared low *MYC*-expression with 1.9-fold lower expression than in *IG-MYC*-positive cases.

Despite the considerable CN alterations found in the 2 cases (12 and 9 alterations, respectively), both shared only one aberration, which was a peculiar pattern of 11q22-q24 gain and 11q24-qter loss (11q-gain/loss) (supplemental Figure 2).

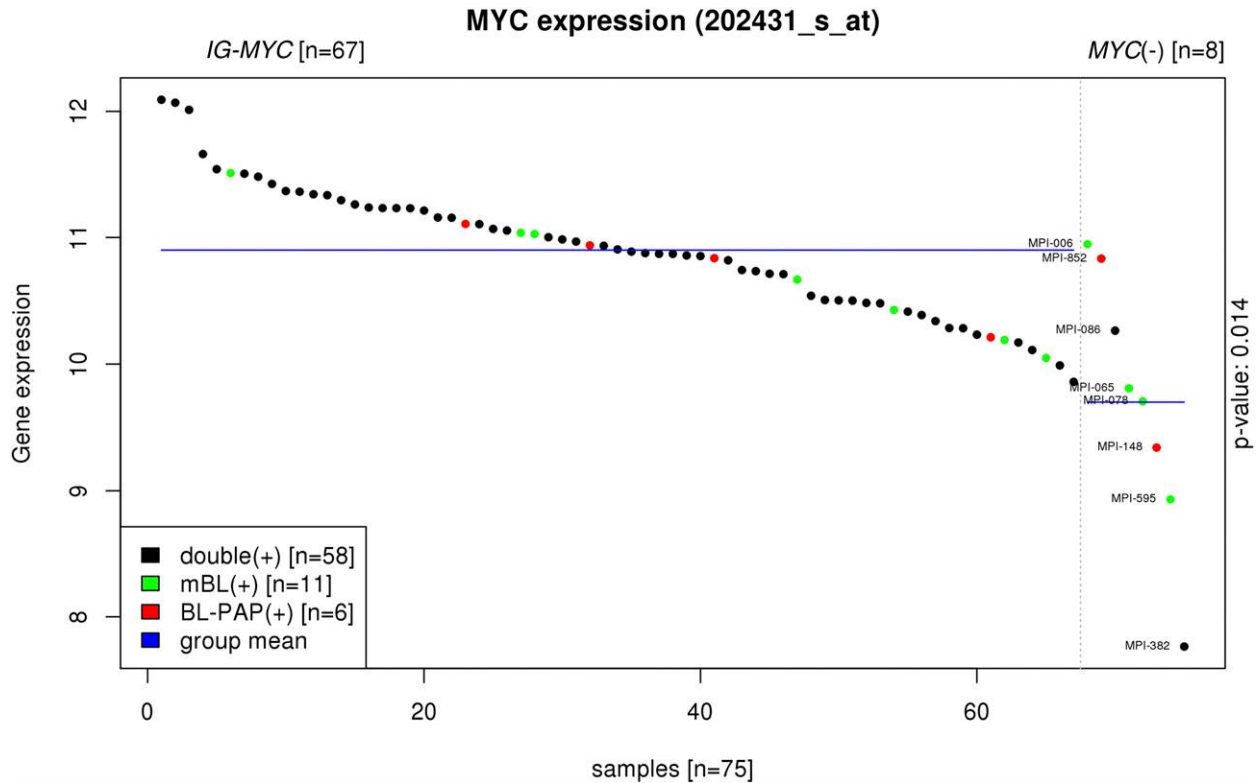


Figure 2. Supervised analysis comparing MYC expression between *IG-MYC*-positive and *MYC*-negative lymphomas from mBL and BL-PAP groups. mBL: BL cases defined by mBL index (green); BL-PAP: BL cases defined by BL-PAP index (red); double: BL cases defined by both mBL and BL-PAP indices (black) ($P = .014$).

Cytogenetic and array-based characterization of the chromosome 11 aberrations pattern in *MYC*-negative lymphomas resembling BL (cohort 2)

Cytogenetic and concomitant aberrations. Because a recent cytogenetic study suggested a similar pattern appearing as partial trisomy 11 resulting from a dup(11)(q23q13) to characterize lymphomas with BL morphology without *MYC* gene rearrangement,⁸ an extensive genetic investigation of this chromosome 11 aberration pattern was initiated. In addition to the cases described by Pienkowska-Grela et al.,⁸ we identified 8 lymphomas with features of BL predominantly in children and young adults. The mean age of the whole series was 20.5 years (range 6-76).

The median follow-up of the patients was 36 months. All patients but one (case 6) were alive at last follow-up (Table 1). The cytogenetic analyses confirmed the absence of a Burkitt translocation and other recurrent *IG* translocations such as the t(14;18)/*BCL2* breaks. Despite the lack of any detectable *MYC* translocation, these cases resembled BL in many aspects including morphologic features (“starry sky” pattern, many mitoses, and very few infiltrating reactive lymphocytes) and immunophenotype (mostly CD20⁺/CD10⁺/*BCL2*⁻ with Ki67 positivity close to 100%) (Figure 3; supplemental Table 2). Epstein-Barr virus RNA expression was not detectable in 10 cases analyzed (Table 1). Cytogenetically, the chromosome 11 aberration was mostly described as inverted duplication 11q13-q23 (5/9). In addition to the chromosome 11 aberrations, 6q14-q24 and 18q21-q22 changes were recurrent in the cases, being present in 2 and 3 of the 9 evaluable cases, respectively (Table 2).

The cell lines SU-DHL-5 and HT were selected as potential models of this subgroup of lymphoma because they showed the same

aberration pattern 11q-gain/loss (cytogenetics or arrays). None of the cases and cell lines carried *ID3* mutations, which have been recently shown to be present in >60% of BL.^{21,23,24}

To define the pattern of chromosome 11 aberrations at high resolution, we performed single nucleotide polymorphism (SNP) array analyses on DNA extracted from fresh frozen cultured cells (cases 1-3, 8, SU-DHL-5, and HT) and oligonucleotide CGH-array on DNA from formalin fixed paraffin embedded tissue tumor samples (cases 4-7 and 9-12) (Figure 4). Common regions of alteration in addition to 11q-gain/loss pattern were gains of 7q34-qter (3 cases), 12pter-p12.2 (4 cases), 18q21.2 (3 cases), and 19pter-p13.2 (2 cases), and losses of 6q14.3-q22.2 (3 cases) (supplemental Table 4). No recurrent regions of CN neutral loss of heterozygosity (CNN-LOH) were observed.

Minimal gained/amplified 11q region. With regard to chromosome 11q, interstitial gains in 11q23.2-q23.3 were detected in 13 of 14 samples analyzed in the cohort 2 (92.9%). Of those, 5 of 13 samples included focal high-level amplifications in 11q23.3 (38.5%) (cases 2, 6, 9, 10, and HT). The minimal region of gain was defined as chr11:114 530,818-117 939,359bp (hg18) and the minimal region of amplification as chr11:117 107,361-117 939,359bp (hg18) (supplemental Table 5). Remarkably, the *ATM* gene was located in the region of gain in 8 of 14 samples, but it is noteworthy that one lymphoma (case 1, MPI-626) occurred in a patient with ataxia telangiectasia associated with a homozygous germline deletion of exons 9 and 10 of the *ATM* gene (c.663_1065del403; EX9_10del) present in a stretch of CNN-LOH.

Recently, hsa-mir-34b was described to be downregulated in BL cases that were negative for *MYC* translocation.⁵ Remarkably, the hsa-mir-34b is located in 11q23.1 (chr11:110 888,873-110 888,956bp,

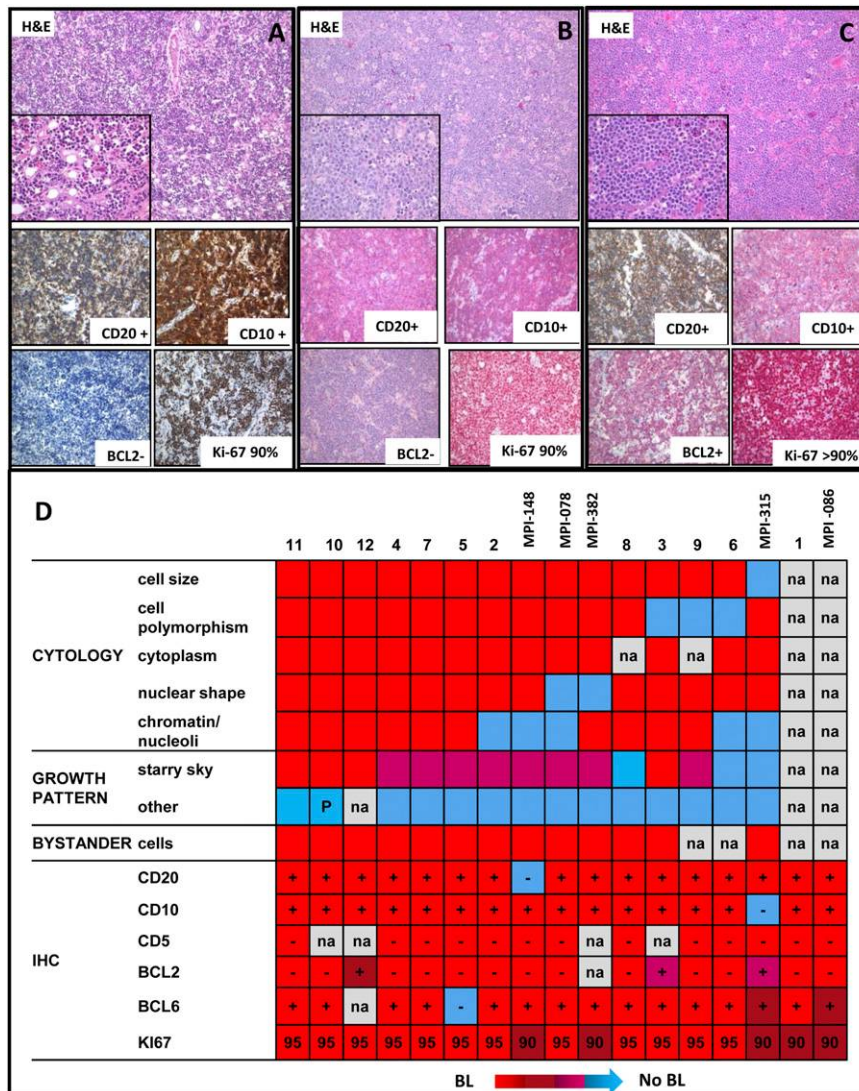


Figure 3. Morphologic features of *MYC*-negative high-grade lymphomas. (A) Hematoxylin and eosin preparation showing the morphology and CD20, CD10, BCL2, and Ki67 expression by immunohistochemistry of (A) case 1, (B) case 2, and (C) case 3 in cohort 2. (D) Heatmap showing the morphologic and the immunohistochemical features of the 17 *MYC*-negative high-grade lymphomas. The morphologic characteristics included information of the cytology, growth pattern, and bystander cells. Red, pro-Burkitt: medium cell size; no cell polymorphism; narrow cytoplasm; round nuclear shape; multiple, small, and paracentric nucleoli; "starry-sky" growth pattern; scattered bystander cells; and sparse eosinophils. Blue, no BL-like: large cell size; cell polymorphism; abundant cytoplasm; irregular nuclear shape; multiple, small, eccentric (centroblast-like) nucleoli/single, large, central (immunoblast-like) nucleolus/finely, granular (lymphoblast-like) other chromatin; absent "starry-sky" growth pattern; abundant bystander cells. Immunohistochemical analyses included data on CD20, CD10, CD5, BCL2, BCL6, and Ki67. -, negative; +, positive; IHC, immunohistochemistry; na, not available; P, partial.

hg18) and thus in the recurrently gained rather than in the lost region. Thus we analyzed the DNA methylation of hsa-mir-34b. Heterogeneous methylation patterns of hsa-mir-34b were found in cases 1 through 7 from cohort 2 (supplemental Figure 3). In addition, we found no significant enrichment of hsa-mir-34b annotated target genes (Molecular Signatures Database, MSigDB) among the genes that were differentially expressed between cases with the 11q-gain/loss pattern and *IG-MYC*-positive mBL nor between these cases and DLBCL.

Minimal loss/homozygously lost 11q region. Telomeric losses of 11q affecting 11q24.1-qter were observed in the whole cohort 2. The common region of deletion was chr11:126 977 015-134 445 937bp (hg18). Interestingly, one patient (case 9) displayed a focal homozygous deletion within this region located at chr11:127 322 011-128 846 569bp (hg18) (supplemental Figure 4A; supplemental Table 4). The minimal region of loss defined by this homozygous deletion contains, among others, the genes *FLII* and *ETSI*, which have been shown to be involved in hematologic neoplasms and to be mutated in lymphomas, respectively.^{25,26}

The 11q-breakpoints between regions of gain and loss were not conserved in the different cases, and breakpoints scattered from 117 939 359 to 126 977 015bp (hg18) from 11pter (supplemental Figure 4B).

Identification of lymphomas with the typical 11q-gain/loss pattern in the MMML series (cohort 3)

Having defined the minimal regions of gain and loss in the retrospective cohort 2, we investigated the available CGH-array data of the MMML for cases with simultaneous gain and loss of the minimal regions in 11q. This search yielded a total of 6 cases. Excluding the 2 initial *MYC*-negative BL and one case that was also in cohort 2 (case 1, MPI-626), this left us with 3 additional cases. Again, these cases resembled BL in many molecular features. Notably, 2 of these cases were classified as BL by one, but not both, of the GEP algorithms (MPI-078 and MPI-148). Table 3 provides an overview on all 6 cases identified from the MMML cohort in which the typical 11q-gain/loss pattern was observed in the absence of an *MYC*, *BCL2*, or *BCL6* translocation. A comparison of pathological and molecular features contrasting these 6 cases with mBL and DLBCL from the MMML cohort is provided as supplemental Table 6. These results once more indicate that based on CD10 positivity, BCL2 negativity, and high Ki67 expression, the identified cases resembled more BL than DLBCL. Moreover, the cases had an excellent survival rate. Specifically, global survival analysis of the 16 cases from cohort 2 and cohort 3 compared with *MYC*-positive BL patients younger than 40 years

Table 2. CN and cytogenetic aberrations of the 12 cases of MYC-negative B-cell lymphomas and SU-DHL-5 and HT cell lines (cohort 2)

Case no.	Large CN alterations (>4Mb)	Karyotype	t(8;14)	MYC BAP/ IGH BAP	MYC BAP 1/2	BCL2 MALT1 BAP	CEP11/ATM/ FDX	CEP11/11qter	CCND1 BAP
Case 1	Gains (7q34-qter, 8q23.1-q24.22, 11q12.2-q23.3, 13q22.3-q33.2, 18q12.3-q21.1); ampl (12q12-q13.12, 18q21.1-q22.1); losses (4q21.21-q22.1, 6q13-q22.2, 11q23.3-qter, 13q33.2-qter, 18q22.1-q23)	46,XY,del(6)(q14q24), add(7)(p21), dup(11)(q13q23) ,t(12;18)(q12;q23), ins(18;?)(q21;?)[18]	-	-	-	- (3 copies)	CEP11x2;ATMx3;FDXx3		
Case 2	Gains (11q12.1-q23.2); ampl (11q23.3-qter); losses (10q23.2-q23.32, 11q23.3-qter)	46,XY,? dup(11)(q12q23) , der(18)?t(13;18)(q21;q21)[15]	-	-	-/-	-	CEP11x2; ATMx2;FDXx2		
Case 3	Gains (3p26.3-q13.11, 3q22.1-q29, 10q24.32-q26.3, 18q21.1-q23); ampl(3q12.1-q13.11); losses (3q13.11-q22.1, 6q14.3-q24.2, 11q23.3-q25, Xp22.33-qter)	47,X,-X,+3,add(3)(q11)x2, t(6;10)(p22;q22),der(6)t(6;10)(p22;q22)del(6)(q12q23), add(11)(q23) ,+mar [12]/46, idem,t(9;13)q10;p10), -mar[5]	-	-	-/-	-	CEP11x2; ATMx2;FDXx2		
Case 4	Gains (11q13.4-q23.3); losses (11q23.3-qter)	46,XY, dup(11)(q23q13) [20]	-	-	-	-	CEP11x2; ATMx3;FDXx3	CEP11x2 11qterx1	
Case 5	Gains (7q34-qter, 11q13.4-q24.1,12); losses (11q24.1-qter)	46-47,XY, dup(11)(q23q13) -13,+ 1~2mar[15]/46-47,sl,+add(8)(p21),+der(14)t(4;14)(q31;q13), add(18)(q22)[cp5]	-	-	-	-			
Case 6	Gains (11q13.1-q23.3, 19p13.3-p13.2); ampl (11q23.3); losses (11q23.3-qter)	46,XY, dup(11)(q23q13) [9]	-	-/+	-	-	CEP11x2; ATMx3;FDXx3	CEP11x2 11qterx1	CCND1x3
Case 7	Gains (11q13.1-q24.1, 12); losses (11q24.1-qter)	47,XX, dup(11)(q23q13) , +12[cp21]	-	-/+	-	-		CEP11x2 11qterx1	CCND1x3
Case 8	Gain (7q31.1-qter; 11q12.1-q24.2; 12pter-p12.2; 12p12.1-qter); losses (9q21.11-q31.1; 11q24.2-qter)	47,XY,del(9)(q22q34), der(11)(11pter->11q23::?11q23->11q11:?) ,+12[7]/47,sl, add(17)(p11)[2]	-	-	-	-			
Case 9	Gain (11q13.1-q23.3); ampl (11q14.3-q23.3); losses (2q14.3-q31.1; 11q23.3-qter)	na	-	-	-/-	-	ATMx4-5; FDXx4-5		
Case 10	Gain (10; 11q12.1-q23.3; 11q23.3); losses (11q23.3-qter)	na	-	-	-/-	-			
Case 11	Gain (8q24.13-q24.23; 11q13.1-q23.3); losses(7q11.21-q11.22; 7q11.22-q11.23; 11q23.3-qter)	46, XY, add(11)(q23) [6]/ 46,XY[14]	-	-	-/-	-		CEP11X2 11qterx1	
Case 12	Gain (11q12.1-q22.3); losses (11q22.3-qter)	na	-	-	-	-		CEP11X2 11qterx1	
SU-DHL-5	Gains (11q23.2-q23.3, 12pter-q13, 19pter-p13.2); losses (6pter-p22.1, 6q13-qter, 11q23.3-qter)	47,XX,add(6)(p21), del(6)(q13), +12,del(12)(q12)	-	-	-/-	-			
HT	Gain (10q21.1-q22.2, 11q22.3-q23.2); ampl (11q22.3-q23.2; 11q23.2-q23.3); losses (2p25.3-p24.1, 2p22.3-p16.2, 2p16.1-p15, 2p15-p12, 2q13-q24.1, 4pter-p12, 11q23.3-qter)	46(42-46)<2n>XY,+2, der(2;4)(p10;q10), der(2)(del)(2)(p11?q21), dup(10)(q11q22-23), dup(11)(?q23qter)	-	-	-/-	-			

-, negative; +, positive; BAP, break apart.

Bold indicates chromosomal alterations that involve chromosome 11.

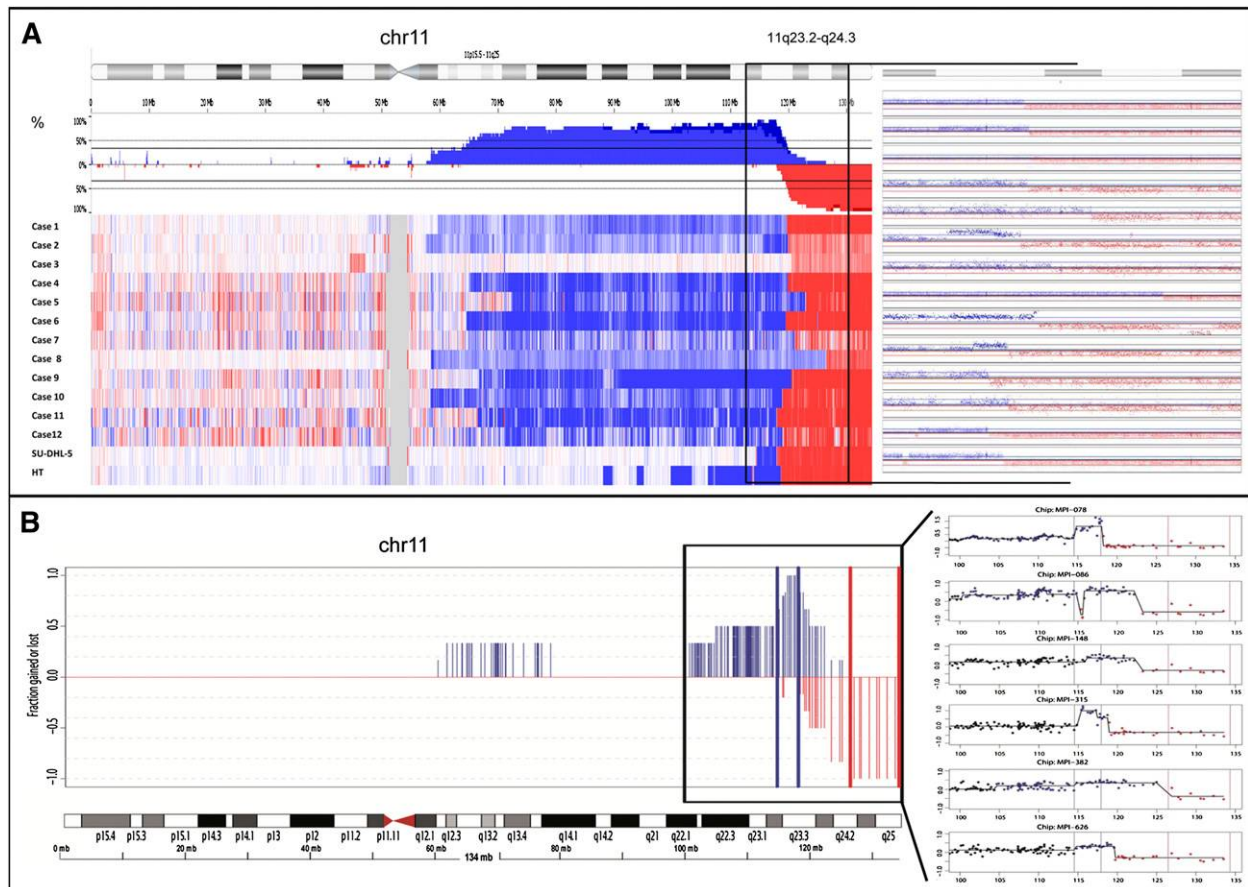


Figure 4. 11q alterations in *MYC*-negative high-grade lymphomas. (A) Chromosomal view of chromosome 11 analyzed by SNP arrays in cohort 2 ($n = 14$, including SU-DHL-5 and HT cell lines). (B) Chromosomal view of chromosome 11 analyzed by custom CGH-array in cohort 3 ($n = 6$).

($n = 51$ from the MMML series) showed that this subgroup of cases has no significantly different prognosis than typical BL of this age range (3-year overall survival: 92% vs 76%, $P = .12$) (supplemental Figure 5). In terms of genetic complexity and in line with the results of cohort 2, cohort 3 showed recurrent losses of chromosome 6q as the second most frequent genetic alteration in these cases (4/6 cases) (supplemental Table 4). Combining both data sets, the minimal region of loss was in 6q14.1-q21 and included the *EPHA7* and *PRDM1* tumor suppressor genes.^{27,28} Overall, the cases with the typical 11q-gain/loss pattern showed a significantly higher chromosomal complexity based on CGH-array compared with mBL and DLBCL (12.2 vs 4.2 and 7.2 CN alterations, respectively; $P < .0001$).

Identification of target genes of the duplicated/deleted 11q regions

To identify candidate genes potentially deregulated through the gains/amplifications in 11q23 and the telomeric losses in 11q24, we compared the GEP of the 6 lymphomas from the MMML with the typical 11q-gain/loss pattern (cohort 3) to *IG-MYC* mBL ($n = 46$) and DLBCL ($n = 198$) (supplemental Tables 7 and 8). Several genes located in the minimal region of gain at 11q23.3 (*PAFAH1B2*, *ZNF259*, *PCSK7*, *CEP164*, *UBE4A*, and *ATP5L*) were significantly overexpressed in 11q-gain/loss pattern cases compared with both mBL and DLBCL (supplemental Figure 6A-B).

To investigate whether the transcriptional upregulation also translates into changes in protein expression, we performed Western blot analyses of the candidate gene *PAFAH1B2* in cell lines with and

without 11q-gain/loss pattern as well as in BL cell lines (supplemental Figure 6C). Also on the protein level, this gene was overexpressed in HT and SU-DHL-5 compared with cell lines without an 11q-gain/loss pattern.

In the minimal region of loss at 11q24.1-qter, the *FLII*, *SNX19*, *NCAPD3*, and *ACAD8* genes showed significantly lower mRNA expression in cases with an 11q-gain/loss pattern (supplemental Figure 7A-B). Thus sequence analysis of *FLII* was initiated. In addition, the *ETS1* gene located in the focal homozygous deletion of case 9 entered into mutational screening. Sequencing of all exons of *FLII* showed no mutations in the 12 analyzed samples with sufficient material available (data not shown). In contrast, 7 point mutations were detected in *ETS1* in 4 of 16 cases studied (supplemental Table 9; supplemental Figure 7C). Two mutations were silent and 5 were nonsilent, corresponding to 3 transitions and 2 transversions. In total, 3 mutations were missense, 1 was located in the splice donor site of intron 1, and the last one was an inactivating mutation (p.Y154X).

In addition to the Y→X/del mutation that removes the DNA binding domain in case 8, no clear functional consequences of any mutation could be derived based on protein modeling, because the modified residues lie outside of regions with resolved structures.

To explore recurrence of *ATM*, *ETS1*, and *FLII* gene mutations, data from custom exome sequencing of HT and SU-DHL-5 cell lines with regard to genes located at 11q were investigated. However, the cell lines showed no mutation of the genes in the minimal regions of loss including *ETS1* and *FLII* (data not shown). *ATM* was also not mutated in these cell lines.

Table 3. Morphologic, genetic, immunohistochemical, and clinical data of the 6 MMML cases carrying 11q-gain/loss pattern selected for GEP analysis (cohort 3)

Case no.	Age, sex	Diagnosis	Location	FISH		Immunohistochemistry										Mutations					GEP	Molecular diag	PAP diag†	Outcome, follow-up
				MYC break	BCL6 break	CD20	CD10	CD5	BCL2	BCL6	MUM1	Ki67	EBER	IGHV	BCL6	MYC	ABC/GCB	MYC	BCL6	IGHV				
MPI-078	8, male	Centroblastic, DLBCL	LN(c)	-	-	+	+	+	-	+	+	+	+	95	-	mut	mut	wt	GCB	mBL	mind-L	In complete remission, alive, 119 mo		
MPI-086¶	14, male	High grade B-cell lymphoma	LN(c)	-	-	+	+	+	-	+	+	+	+	90	-	mut	mut	wt	GCB	mBL	BL-PAP	In complete remission, alive, 161 mo		
MPI-148*	26, male	Atypical BL	extranodal	-	-	-	+	-	-	+	+	+	-	90	-	mut	mut	wt	GCB	intermediate	BL-PAP	In complete remission, alive, 163 mo		
MPI-315	49, female	Centroblastic, DLBCL	intestine	-	-	+	-	+	+	+	+	+	+	90	-	mut	na	na	GCB	intermediate	PAP-1	In complete remission, alive, 106 mo		
MPI-382¶	24, male	Centroblastic, DLBCL	LN	-	-	+	+	na	na	+	+	+	na	90	-	mut	na	na	GCB	mBL	BL-PAP	In complete remission, alive, 91 mo		
MPI-626†	7, male	Malignant B-cell lymphoma	LN(c)	-	-	+	+	-	-	+	+	+	-	90	-	mut	na	wt	GCB	intermediate	mind-L	In remission, alive, 62 mo		

-, negative; +, positive; ABC, activated B cell; c, cervical; diag, diagnosis; EGER, Epstein-Barr-encoded RNA test; GCB, germinal center B cell; GEP, gene expression profiling; LN, lymph node; mind-L, molecularly individual lymphoma; mut, mutated; na, not available; PAP, pathway activation pattern algorithm; wt, wild-type.

*This case presented the following karyotype: 46,XY,t(7;18)(q32;q21),inv dup(11)(q23q13),add(15)(q22),inv(16)(q11q24).

†This case corresponds to case 1 from cohort 2 and was investigated using SNP and CGH-array platforms.

‡Molecular diagnoses described in Hummel et al and Benitink et al, respectively.^{7,12}

¶These two cases belong to cohort 1. Lymphoma samples were scored positive if >25% of the tumor cells stained positive except for MYC, where the cutoff was established in 40%.

Differential gene expression in lymphomas with the typical 11q-gain/loss pattern

Finally, we compared the global GEP of cohort 3 with those of BL and DLBCL. There were 1194 and 1826 genes differentially expressed between 11q-gain/loss cases and *IG-MYC*-positive BL (n = 46) or DLBCL (n = 198), respectively (supplemental Figures 8 and 9). Compared with DLBCL, cases with 11q-gain/loss pattern showed upregulation of genes involved in cell cycle and proliferation and downregulation of major histocompatibility complex/immune-interferon responses, cell death, and signaling, most of them also with features distinguishing BL from DLBCL. Compared with BL, 11q-gain/loss cases showed downregulation of RNA binding activity and metabolic processes and upregulation of RNA degradation.

Discussion

Here we report the clinical, morphologic, and molecular characterization of a subset of *MYC*-negative, high-grade B-cell lymphomas sharing similarities with but not being BL. The search for *MYC*-negative lymphomas with a GEP of BL showed that only 2 cases of 59 were concordantly called “BL” by 2 GEP classifiers, stating that these cases are in general rare. In other series, *MYC*-negative BL accounted for 14% to 22% of all BL. Nevertheless, many of these studies either do not apply a full set of FISH probes necessary to exclude rare variants of the *IG-MYC* translocation or are not based on gene expression.^{5,29} In addition, a recent study applying an extensive set of FISH probes also reached a similar low frequency of *MYC*-negative BL as described herein.³⁰ Of note, 4 of 6 *MYC*-negative mBL and 2 of 4 *MYC*-negative BL-PAP were differently classified by the alternative BL classifier, suggesting that in a high percentage (50-66%) of even molecularly defined *MYC*-negative BL, the diagnosis should be questioned. The significantly lower expression of *MYC* mRNA in the *MYC*-translocation-negative mBL/BL-PAPs supports the view that *MYC* indeed is not genomically activated in these cases, rendering a technical failure of FISH in detection of the translocation unlikely.

Both molecularly defined *MYC*-negative BL cases shared a recurrent pattern of chromosome 11q abnormalities that was subsequently found in a series of 12 high-grade B-cell lymphomas with features of BL and in 2 cell lines. A similar pattern was identified in a set of 6 *MYC*-negative cases from MMML series classified as mBL/intermediate by GEP. The comparison of pathological and molecular features between these 6 cases compared with DLBCL and BL showed that these tumors resembled more BL than DLBCL, reinforcing the idea that this group of samples may represent a genetic variant of BL lacking the *MYC* translocation. Nevertheless, for the sake of diagnostic clarity, it is proposed here to restrict the term BL to *IG-MYC*-positive cases and call the lymphomas described herein “high-grade B-cell lymphoma with features of BL, *MYC*-, with 11q-gain/loss pattern.”

Alterations of the 11q chromosomal region are not frequent in typical BL^{31,32} but gains/amplifications of 11q22-q24 are found in approximately 8% to 15% of DLBCL^{33,34} and also in GC-derived *IRF4*-translocation-positive lymphomas.³⁵ Nevertheless, the region of gain in these cases (minimal region at 11q23.2-q23.3) is not coincident with the minimal region observed in the present series. Indeed, in the DLBCL and *IRF4*-translocation-positive lymphomas, genes like *ETS1* or *FLII*, which are included in our minimal region of loss (11q24.1-qter) and are even homozygously deleted in one case, are considered as oncogenes in DLBCL because they cooperate in

sustaining DLBCL proliferation and viability and regulate genes involved in germinal center differentiation,^{34,36} contrasting the findings herein, which rather suggest a tumor-suppressive function.

The incidence of 11q-gain/loss pattern in the total group of 514 samples from MML with CGH-array data are the following: 0.7% in DLBCL (2/267), 1.6% in *IG-MYC* mBL (1/63), 16.7% cell lines (2/12), 0% FL (0/67), 5.2% intermediate (4/77), 50% other mBL without *IG-MYC*, (3/6), and 0% others (0/22).

When we screened the Mitelman (<http://cgap.nci.nih.gov/Chromosomes/Mitelman>) and Progenetix (www.Progenetix.org) databases (September 2012) for pediatric lymphomas, we found 10 of 31 cases (32%) diagnosed as BL/Atypical without the t(8;14) translocation or variants presented alterations on chromosome 11.

A potential candidate oncogene in the gained region in 11q in the lymphomas described herein is *PAFAH1B2*, which we showed to be selectively upregulated also on the protein level in cell lines containing the typical 11q pattern (supplemental Figure 6C). The *PAFAH1B2* gene has been previously recognized as target of *IGH* locus translocations in 2 lymphoma cases.³⁷ Such *IGH* translocation leads typically to oncogene activation and it is therefore consequent to suggest *PAFAH1B2* (previously named *PAFAH1A2*) as an oncogene in lymphomagenesis. Furthermore, it has been found to interact at protein level with *ZFP36L1*,³⁸ another partner of *IGH* translocations described in chronic lymphocytic leukemia^{39,40} and the target of hypermutation in DLBCL.⁴¹ It is also intriguing to speculate that, given the recurrent co-amplification of several genes in 11q, there is a cooperative oncogenic function, as has been proposed for 9p24 gains in some subsets of B-cell lymphomas.^{42,43}

In the minimal region of loss, *FLII* and *ETS1* were the candidate tumor suppressor genes, both located in the region of focal homozygous loss. The transcription factor *FLII* is known to be involved in lymphoma development in mice,²⁵ and we found it significantly downregulated in the cohort of *MYC*-negative lymphomas with the 11q aberrations from cohort 3. Nevertheless, this gene displayed no mutations in any of the samples analyzed. The *ETS1* gene, however, was found to be homozygously lost in one and mutated in four of the 16 lymphomas investigated. In addition, *ETS1* has been identified as an oncogenic fusion partner of the c-Myb proto-oncogene product in the E26 avian leukemia virus⁴⁴ and has been found to be recurrently mutated in non-Hodgkin lymphomas,²⁶ although the functional consequences of the detected mutations remain to be defined. Our attempts to model the effect of the mutations yielded no striking results because of the low number of mutations found in our series and the fact that none of the point mutations were within resolved 3-dimensional structures. Additional structures or biophysical studies of *ETS1* would be required to resolve the possible roles of these positions in protein function. High levels of expression associated with gains of the 11q24.3 (including *FLII* and *ETS1*) have been also seen in these genes in DLBCL patients,³⁴ although a different mechanism seems to be involved in these cases.

Recently, several studies showed that *ID3* mutations and/or *TCF3*-activating mutations in BL lead to activation of PI3K signaling.^{21,23,24} Moreover, PI3K pathway activation has been shown to cooperate with *MYC* activation in the pathogenesis of BL.⁴⁵ Our series displayed no mutations in *ID3*. Nevertheless, PI3K signaling activity in the cases described herein are distributed similarly to that of *MYC*-positive BL, applying the pathway activity quantification approach of Sander et al (supplemental Figure 10).⁴⁵ These findings could suggest that either alternative mechanisms cause similar pathway deregulations like in *MYC*-positive BL, or that the cell of origin of the lymphomas with features of BL characterized by the 11q aberration is the same as that of *MYC*-positive BL.

In contrast to Leucci et al,⁵ we failed to identify a role of hsa-mir-34b in *MYC*-negative BL. These results are in agreement with a recent study performed by Lenze et al in which the *MYC*-negative BL cases did not show downregulation of this microRNA.⁴⁶ It is remarkable that this microRNA is located in a region of gain in 6 of the 7 cases with 11q aberration studied.

In conclusion, we have identified a particular molecular subset of high-grade B-cell lymphomas that share a recurrent pattern of 11q aberration. Cases are morphologically and phenotypically similar to BL but differ with regard to the *MYC* translocation, the 11q-gain/loss pattern and the more complex karyotypes. Of note, lymphomas with the typical 11q-gain/loss pattern seem to have more frequently nodal presentation than BL from patients younger than 40 years (82% vs 55%, $P = .074$). The specific characteristics detected in this group of lymphomas raise the question of whether these lymphomas truly belong to the entity of BL or whether they could be considered as another set of high-grade, mature B-cell lymphomas sharing features of, but not being, BL. We provide evidence that the presence of the typical 11q-gain/loss pattern could be a genetic hallmark that defines this group of cases. The pattern of aberrations along with the results of our molecular analyses suggest that activation of genes included in the region of gain in 11q may cooperate with the functional inactivation of genes in the recurrent region of loss, including *FLII* and *ETS1*, most likely through transcriptional mechanisms in this newly identified subgroup of lymphomas.

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Authorship

Contribution: I.S. and I.M.-G. performed molecular analysis, analyzed data, and wrote the manuscript; R.W. performed molecular analysis and analyzed data; M.K., C.W.K., M.L., M.R., and

R. Spang performed statistical analyses; J.R., B.P.-G., H.G.D., M.H., R.K., R.A.F.M., I.N., R.B.R., D.S., M. Schlesner, R. Scholtysik, C.S., M. Szczepanowski, I.V., and S.W. performed molecular analysis; B.B., A.C., C.D.-W., E.S.J., C.L., J.L., and L.T. provided clinical data and patient samples; P.A., I.O., G.R., and W.K. provided samples and performed pathology review; and R. Siebert designed the project, analyzed data, and wrote the manuscript.

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