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# BIOLOGICAL AND CLINICAL IMPLICATIONS OF BIRC3 MUTATIONS IN CHRONIC LYMPHOCYTIC LEUKEMIA 

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## SUMMARY

The current shift of therapy of chronic lymphocytic leukemia (CLL) towards novel targeted agents mandates the identification of molecular predictors to inform on who can still benefit from chemoimmunotherapy and who can be instead early considered for novel targeted agents. Fludarabine, cyclophosphamide, and rituximab (FCR) is the most effective chemoimmunotherapy regimen for the management of CLL and represents the current standard of care for young and fit patients devoid of TP53 disruption. A retrospective multicenter cohort of 287 untreated patients receiving first-line FCR was analyzed by targeted next generation sequencing of 24 recurrently mutated genes in CLL. By univariate analysis adjusted for multiple comparisons BIRC3 mutations identify a poor prognostic subgroup of patients failing FCR (median progression free survival: 2.2 years, $\mathrm{p}<0.001$ ) similar to cases harboring TP53 mutations (median progression free survival: 2.6 years, $\mathrm{p}<0.0001$ ). BIRC3 mutations maintained an independent association with an increased risk of progression with a hazard ratio of 2.8 ( $95 \%$ confidence interval 1.45.6, $p=0.004$ ) in multivariate analysis adjusted for TP53 mutation, 17p deletion and IGHV mutation status. The functional implications of BIRC3 mutations are largely unexplored and little is known about the prognostic impact of BIRC3 mutations in CLL cohorts homogeneously treated with first line FCR. By immunoblotting analysis, we showed that the non-canonical NF-кB pathway is active in BIRC3 mutated cell lines and in primary CLL samples, as documented by the stabilization of MAP3K14 and by the nuclear localization of p52. In addition, BIRC3 mutated primary CLL cells are less sensitive to fludarabine. If validated, BIRC3 mutations may be used as a new molecular predictor to select high-risk patients for novel frontline therapeutic approaches.

## SOMMARIO

Le terapie innovative per la leucemia linfatica cronica (CLL) includono nuovi agenti i quali richiedono l'identificazione di predittori molecolari per determinare i pazienti che possono ancora beneficiare della chemio-immunoterapia e coloro che, invece, necessitano di trattamento con nuovi farmaci. La terapia di prima linea per pazienti giovani, in buone condizioni cliniche e privi di aberrazioni del gene TP53, prevede I'utilizzo dello schema immunochemioterapico FCR, la combinazione dei farmaci Fludarabina, Ciclofosfamide e Rituximab. II DNA genomico di 287 pazienti, trattati con terapia secondo lo schema FCR, è stato raccolto alla diagnosi e sottoposto al sequenziamento di 24 geni ricorrentemente mutati nella CLL, mediante tecnica di Next Generation Sequencing (NGS). In analisi univariata, le mutazioni di BIRC3 identificano un sottogruppo di pazienti con prognosi sfavorevole (sopravvivenza mediana libera da progressione: 2,2 anni, $\mathrm{p}<0,001$ ) simile ai casi che presentano mutazioni di TP53 (sopravvivenza mediana libera da progressione: 2,6 anni, $p<0,0001$ ). Le mutazioni di $B I R C 3$ rimangono associate ad un maggior rischio di progressione (HR) in analisi multivariata corretta per mutazione di TP53, delezione 17p e stato mutazionale di IGHV: HR 2,8 ( $95 \%$ I.C. $1,4-5,6, p=0,004$ ). Le implicazioni funzionali delle mutazioni di BIRC3 sono in gran parte inesplorate e poco si conosce circa il loro impatto prognostico in coorti di pazienti trattati omogeneamente con FCR. Mediante analisi di immunoblotting è stato dimostrato che la via non canonica di NF-кB è attiva sia nelle linee cellulari BIRC3 mutate che nelle cellule primarie, come documentato dalla stabilizzazione di MAP3K14 e dalla localizzazione nucleare di p52. Inoltre, le cellule primarie BIRC3 mutate sono meno sensibili al trattamento con Fludarabina. Se validate, le mutazioni di BIRC3 potrebbero essere utilizzate come nuovo predittore molecolare per selezionare pazienti ad alto rischio per nuovi approcci terapeutici in prima linea.

## 1. INTRODUCTION

Chronic lymphocytic leukemia (CLL) is the most common type of adult leukemia in the western world with marked genetic and clinical variability. ${ }^{1}$ The clinical course of CLL ranges from a very indolent condition, with a nearly normal life expectancy, to rapidly progressive leading to early death. ${ }^{2}$ Fludarabine, cyclophosphamide, and rituximab (FCR) is the most effective chemoimmunotherapy regimen for the management of CLL, and represents the current standard for untreated patients who are young and in good physical condition ${ }^{3,4}$ except for patients with TP53 alterations. ${ }^{5}$ Though the majority of CLL patients receiving FCR as frontline therapy are destined to relapse, a subgroup of cases may experience a durable first remission. The current shift of therapy of CLL towards novel targeted agents mandates the recognition of molecular predictors to identify patients who can still benefit from chemoimmunotherapy and those who should instead be considered for novel targeted agents upfront. In the case of FCR, the immunoglobulin heavy chain genes (IGHV) mutational status and Fluorescence In Situ Hybridization (FISH) karyotype stratify: i) low-risk patients carrying mutated IGHV genes and devoid of both del11q and del17p who maximally benefit from such treatment; ii) intermediate-risk patients harboring unmutated IGHV genes and/or del11q in the absence of del17p who are a case mix of good and poor responders to FCR; iii) high-risk patients harboring del17p who are unsuitable for chemoimmunotherapy. ${ }^{3}$ Deletion of 17 p and TP53 mutations capture most, routinely analyzed in clinical practice, but not all patients who are refractory to chemo-immunotherapy, which prompts the identification of additional biomarkers associated with early failure of FCR. ${ }^{3-5}$ B-cell neoplasia often pirates signaling pathways by molecular lesions to promote survival and proliferation. Though according to bioinformatics criteria BIRC3 (also known as cIAP2) is one of the candidate driver genes of CLL, the functional implications of BIRC3 mutations are partially unexplored. ${ }^{7-9}$

Nuclear factor-кB (NF-кB) signaling is a key component CLL development and evolution. ${ }^{10}$ Two NF-кB pathways exist namely, canonical and noncanonical. ${ }^{11}$ The former is triggered by the B -cell receptor (BCR) signaling via the Bruton's tyrosine kinase (BTK), while the latter is activated by members of the tumor necrosis factor (TNF) cytokine family. ${ }^{12}$ Upon receptor binding, the TRAF3/MAP3K14-TRAF2/BIRC3 negative regulatory complex of non-canonical NF-KB signaling is disrupted, MAP3K14 (also known as NIK), the central activating kinase of the pathway, is released and activated to induce the phosphorylation and proteasomal processing of p100, thereby leading to the formation of p52-containing NF-кB dimers. The p52 protein dimerizes with RelB to translocate into the nucleus, where it regulates gene transcription. BIRC3 is a negative regulator of non-canonical NF-кB. Physiologically, BIRC3 catalyzes MAP3K14 protein ubiquitination in a manner that is dependent on the E3 ubiquitinine ligase activity of its C-terminal RING domain. MAP3K14 ubiquitination results into its proteasomal degradation. ${ }^{13}$ Also, little is known about the prognostic impact of BIRC3 mutations in CLL cohorts homogeneously treated with first line FCR.

## 2. AIM OF THE STUDY

We aimed at refining the genetic-based stratification of FCR-treated CLL patients.

The aims of this study are:

- To identify molecular predictors in FCR treated patients;
- To assess the biological features underlying chemo-refractoriness to FCR.


## 3. MATERIALS AND METHODS

### 3.1. Patients

The study was designed as a retrospective observational analysis from a multicenter cohort of 287 (275 with complete clinical and molecular data) untreated CLL receiving first-line therapy with FCR in 17 different hematological centers. The following biological material was collected: i) 280 tumor genomic DNA (gDNA) and 7 tumor RNA isolated from peripheral blood (PB) before treatment start; and ii) paired germline gDNA from saliva from 14 cases. Normal gDNA from 22 healthy donors was also used to set the experimental background of the deep next generation sequencing (NGS) approach. Tumor and normal gDNA was extracted according to standard procedures. ${ }^{14}$ Tumor RNA was extracted according to the TRIzol Reagent protocol (Life Technologies). The clinical database was updated in April 2018. Patients provided informed consent in accordance with local Institutional Review Board requirements and the Declaration of Helsinki. The study was approved by the Ethical Committee of the Ospedale Maggiore della Carità di Novara associated with the Amedeo Avogadro University of Eastern Piedmont (study number CE 67/14)

### 3.2. Cancer personalized profiling by deep sequencing (CAPP-seq)

A targeted resequencing gene panel ${ }^{15}$ was designed to include: i) coding exons plus splice site of 24 CLL genes known to be implicated in CLL pathogenesis and/or prognosis; ii) 3'UTR of NOTCH1; and iii) enhancer and promoter region of PAX5 (size of the target region: 66627bp) (Table S1). ${ }^{8,9}$ Tumor and germline gDNA were quantified using the Quant-iT ${ }^{T M}$ PicoGreen dsDNA Assay kit (ThermoFisher Scientific) and 400 ng were sheared through sonication (Covaris M 220 focused-ultrasonicator) before library construction to obtain 200-bp fragments. The size of the DNA fragments was checked using the Bioanalyzer
(Agilent Technologies). The NGS libraries for gDNA were constructed using the KAPA Library Preparation Kit (Kapa Biosystems) and NGS libraries for RNA were constructed using RNA Hyper Kit (Roche) following the manufacturer's instructions. Hybrid selection was performed with the custom SeqCap EZ Choice Library (Roche NimbleGen). Multiplexed libraries ( $\mathrm{n}=10$ per run) were sequenced using 300-bp paired-end runs on a MiSeq sequencer (Illumina) to obtain a coverage of at least 2000 x in $>90 \%$ of the target region (66627bp) in $80 \%$ of cases (Table S2).

### 3.3. Bioinformatic pipeline for variant calling after CAPP-seq

Initially, FASTQ sequencing reads were deduped. We deduped FASTQ sequencing reads from gDNA by utilizing the FastUniq v1.1 software, that collapses as duplicate reads only those fragments (read pairs) with $100 \%$ sequence identity that also share genomic coordinates. The same approach was also used to dedupe germline gDNA and normal gDNA from 22 healthy donors, to avoid the introduction of biases in variant calling due to the application of different deduplication protocols. Then, the deduped FASTQ sequencing reads were locally aligned to the hg19 version of the human genome assembly using the BWA v.0.6.2 software with the default setting, and sorted, indexed and assembled into a mpileup file using SAMtools v.1. The aligned read families were processed with mpileup using the parameters -A -d 10000000. For cases provided with paired germline gDNA, single nucleotide variations and indels were called in tumor gDNA vs germline gDNA, respectively, with the somatic function of VarScan2 using the parameters min-coverage 1 --min-coverage-normal 1 --min-coverage-tumor 1 --min-var-freq 0--min-freq-for-hom 0.75 --somatic-pvalue 0.05 --min-avg-qual 20 --strand-filter 1 --validation 1 . For cases lacking paired germline gDNA, single nucleotide variations and indels were called in tumor gDNA using the CNS function of VarScan2 using the
parameters --min-coverage 0 --min-readge 2 --min-avg-qual 20 --min-var-freq 0 --min-freq-for-hom 0.75 --p-value 0.05 --strand-filter 1 --output-vcf 1 --variants 0 . The variants called by VarScan 2 were annotated using the SeattleSeq Annotation 138 tool by using the default setting. Variants annotated as SNPs according to dbSNP 138 (with the exception of TP53 variants that were manually curated and scored as SNPs according to the IARC TP53 database), intronic variants mapping > 2 bp before the start or after the end of coding exons, and synonymous variants were then filtered out. The following strict post-processing filters were then applied to the remaining variants to further improve variant call confidence. To filter out variants below the base-pair resolution background frequencies in gDNA across the selector, for cases provided with paired germline gDNA, the Fisher's exact test was used to test whether the frequency of the variant called by VarScan 2 was significantly higher from that called in the corresponding paired germline gDNA, after adjusting for multiple comparisons by Bonferroni test [multiple comparisons corrected p threshold $=$ 0.00000018761163 , corresponding to alpha of $0.05 /(66627 \times 4$ alleles per position]. Accordingly, variants represented in $>10$ reads of the paired germline and/or variants with a somatic $p$ value from VarScan2 > 0.00000018761163 were no further considered. To filter out systemic sequencing errors, a database containing all germline and normal gDNA background allele frequencies was assembled. Based on the assumption that all background allele fractions follow a normal distribution, for both cases provided with paired germline gDNA and cases lacking paired gDNA, a Z-test was employed to test whether a given variant in the tumor gDNA differed significantly in its frequency from typical germline or normal gDNA background at the same position in all the other germline and normal gDNA samples, after adjusting for multiple comparisons by Bonferroni test [multiple comparisons corrected p threshold $=0.00000018761163$, corresponding to alpha of $0.05 /(66627 \times 4$ alleles per position]. Variants that did not pass this filter were no
further considered. Variant allele frequencies for the resulting candidate mutations and the background error rate were visualized using IGV.

### 3.4. Statistical analysis

Progression free survival (PFS) was the primary endpoint and was measured from date of treatment start to date of progression according to IWCLL-NCI guidelines (event), death (event) or last follow-up (censoring). Overall survival (OS) was measured from date of initial presentation to date of death from any cause (event) or last follow-up (censoring). Survival analysis was performed by Kaplan-Meier method and compared between strata using the Log-rank test. A false discovery rate approach was used to account for multiple testing, and adjusted p-values were calculated using the Bonferroni correction. A maximally selected rank statistic was used to determine the optimal cut-off for variant allele frequency (VAF) based on the Log-rank statistics. A cut-off of $3 \%$ of VAF was set for TP53 mutations and of $10 \%$ for all the other genes. The adjusted association between exposure variables and PFS was estimated by Cox regression. Internal validation of the multivariate analysis was performed using a bootstrap approach to estimate means and confidence intervals of hazard ratios (HR), and percentage of selection for each variable in the model. The number of bootstrap samples used was 1000. Statistical significance was defined as p value < 0.05. The analysis was performed with the Statistical Package for the Social Sciences (SPSS) software v.24.0 (Chicago, IL), with R statistical package 3.1.2 and with GraphPad version 7 (GraphPad Software Inc).

### 3.5. Cell studies

The human CLL cell line MEC1, the SMZL cell lines SSK41, VL51, and the MCL cell lines MAVER-1, Z-138 and JEKO-1 were cultured under standard conditions in RPMI-1640 medium with L-glutamine supplemented with $10 \%$ fetal calf serum (FCS), Penicillin (100 U/ml) and Streptomycin (100 U/ml) (Sigma Aldrich). Human HEK-293T cells were maintained in Iscove's Modified Dulbecco Medium (IMDM) supplemented with $10 \%$ fetal calf serum, $100 \mathrm{U} / \mathrm{ml}$ penicillin, $100 \mathrm{U} / \mathrm{ml}$ streptomycin and 2 mM L-glutamine (Sigma Aldrich) under identical conditions.

Three primary cells samples known to harbor heterozygous inactivating mutations of BIRC3 were included in the experiments. Two BIRC3 wild type cases were used as controls.

### 3.6. Western blot analysis

The entire non-canonical NF-кB pathway was assessed using the following specific primary antibodies: anti-BIRC3 (Cell Signaling, \#3130), anti-TRAF2 (Cell Signaling, \#4712), anti-TRAF3 (Cell Signaling, \#4729), anti-MAP3K14 (Cell Signaling, \#4994), anti-Phospho-NF-кB2 p100 (Cell Signaling, \#4810), anti-NF-kB2 p100/p52 (Cell Signaling, \#4882). Anti- $\beta$-actin (Sigma Aldrich, \#A2066) was used as loading control. The Qproteome Nuclear Protein Kit (Qiagen) was used according to the manufacturer's instructions to isolate nuclear proteins from cells. Anti- $\beta$-tubulin (Sigma Aldrich, \#T5201) and anti-BRG1 (G-7) (Santa Cruz Biotechnology, \#17796) were used as controls for the purity of the cytoplasmic and nuclear fractions, respectively. Horseradish peroxidase-conjugated goat anti-mouse (LI-COR, \#926-80010) or anti-rabbit (LICOR, \#926-80011) antibodies were used to highlight binding by enhanced chemiluminescence with the

Clarity Western ECL Substrate (Biorad). Image acquisition and densitometric analyses were performed using the Molecular Imager Gel Doc XR System and the Quantity One software (Biorad).

### 3.7. RNA extraction and gene expression profiling

Total RNA was extracted from exponentially growing cell lines by TRIzol reagent (Life Technologies), and retro-transcribed using the Reverse Transcription Kit (Applied Biosystems). Quantitative real-time PCR (qRT-PCR) was conducted with the Step One Plus apparatus (Step One software 2.0; Applied Biosystems) using commercially available TaqMan Gene expression assays (TNFAIP3: Hs00234713_m1; NFKB2: Hs00174517_m1; NFKBIA: Hs00153283_m1; NFKBIE: Hs00234431_m1; PLEK: Hs00950975_m1; WNT10: Hs00228741_m1; IL2RG: Hs00953624_m1; RELB: Hs00232389_m1; MALT1: Hs01120052_m1) (BIRC3: Hs00985031_g1) (Life Technologies). Reactions were done in triplicate from the same cDNA (technical replicates). The comparative CT method ( $\Delta \Delta C T$ ) was used to calculate relative expression levels of the gene under analysis, using GAPDH (Hs03929097_g1) as internal references.

### 3.8. Knockdown of MAP3K14 by RNA interference

Lentiviruses expressing 3 short hairpin RNAs (shRNAs) targeting MAP3K14, as well as the scrambled shRNA, were produced and cloned into the BamHI/HindIII cloning sites of the pGFP-C-shLenti vectors (OriGene Technologies). Within the $5^{\prime}$-LTR and $3^{\prime}$-LTR regions, each pGFP-C-shLenti vector contains an shRNA expression cassette driven by an U6 promoter, a puromycin resistance marker driven by a SV40 promoter and a GFP driven by a CMV promoter. The shRNA expression cassette consists of 29 bp target-gene-specific sequence, a 7 bp loop, and another 29 bp reverse complementary sequence, followed by a

TTTTTT termination sequence. The HEK293T cell line was co-transfected with expression (3 different pGFP-

C-MAP3K14-shLenti or pGFP-C-non-effective-shLenti) vectors and adjuvant vectors (pMDL, REV and VSVG). Fluorescence microscope was utilized to check the expression of the GFP in the transfected HEK293T cell line. After virus titration, the VL51 cell line was infected with lentiviruses harboring the shRNAs against MAP3K14 and the scrambled through a spinoculation protocol. After four days, infected cells were monitored by flow cytometry for the expression of the GFP and were selected by puromycin ( $1.5 \mu \mathrm{~g} / \mathrm{mL}$ ). Cell viability was monitored by Tripan blue counting.

### 3.9. Inhibitor studies

Cells were put under starvation in RPMI 0.1\% Fetal Bovine Serum (FBS) 24 h before treatment. Then they were seeded at 8000 cells per well in a 96 -well U-bottom plate and treated with $1 \mu \mathrm{M}, 5 \mu \mathrm{M}$ and 10 $\mu \mathrm{M}$ of Ibrutinib (PCI-32765, Selleckchem) or vehicle (DMSO). Relative growth was determined by a CellTiter Glo (CTG) Luminescent Cell Viability Assay (Promega) 72h and 96h after treatment, according to the manufacturer's instructions and luminescence was quantified using a Victor X (PerkinElmer) multilabel reader. Treatments were done in triplicate (biological replicates).

### 3.10. In vitro drug responses in primary CLL cells

Leukemic cells were purified using Ficoll-Hypaque (Sigma Aldrich) from PB of CLL patients. Staining with CD19 and CD5 confirmed that in all samples leukemic cells were $>90 \%$. Patients were then divided into BIRC3 mutated (MUT) or wild-type (WT). TP53 mutated samples (and BIRC3 WT) were selected as positive control (i.e., cells intrinsically resistant to therapies). Cells were cultured in RPMI 10\% FCS (200 $\mu \mathrm{l}$, all
reagents from Sigma) at a density of $5 x 106 / \mathrm{ml}$ and both dose- and time-dependent responses were analyzed. Specifically, CLL cells were exposed to fludarabine) for 24-48 hours. Fludarabine was used at 1-5-$10-25 \mu \mathrm{M}$ and venetoclax at 5-10-50-100-500-2000 nM .

### 3.11. Apoptosis assay

Drug-induced apoptosis was measured using the eBioscience ${ }^{\text {TM }}$ Annexin V Apoptosis Detection Kit APC (ThermoFisher) following the manufacturer's instruction. Data were acquired using a FACSCanto II cytofluorimeter (BD Biosciences) and processed with DIVA v6.1.3 and FlowJo Version 9.01 (TreeStar). Apoptosis assays were analyzed using the two-way ANOVA test.

## 4. RESULTS

### 4.1 Patients harboring BIRC3 mutations are at risk of failing FCR

Mutational profiling was performed in 287 patients who received first line FCR. The baseline features of the study cohort were consistent with progressive, previously untreated CLL (Table 1). The median follow-up was 6.8 years, with a median PFS and OS of 4.6 and 11.7 years, respectively (Table 1) consistent with clinical trial cohorts. ${ }^{16}$ As expected, SF3B1 and NOTCH1 were the most frequently mutated genes identified in $13.9 \%$ and in $13.6 \%$ of patients respectively, followed by TP53 in $9.4 \%$ and ATM in $6.9 \%$ of patients, reflecting the data reported in previous studies. ${ }^{8,9,17}$ Overall, 154/287 (53.6\%) cases harbored at least one non-synonymous somatic mutation in one of the 24 CLL genes included in our panel (range: 1-5 mutation per patient), which is consistent with the typical mutational spectrum of the coding genome of CLL requiring first line treatment. (Figure 1; Table S3). ${ }^{8,9,18}$ Outside of the coding genome, we identified one single mutation in the $3^{\prime}$ region of NOTCH1 (c.*378A>G) already reported. ${ }^{8}$

By univariate analysis adjusted for multiple comparisons, among the genes analyzed in our panel, only TP53 mutations (median PFS of 2.6 years; $\mathrm{p}<0.0001$ ) and BIRC3 mutations (median PFS of 2.2 years; p < 0.001) (Figure 2 A) associated with significantly shorter PFS (Table 2). The PFS after FCR of BIRC3 mutated patients was similar to that of cases harboring TP53 disruption (Figure 2 B). Consistently, BIRC3 mutated patients had a lower likelihood of achieving complete response (22.2\%) at the end of FCR compared to BIRC3 wild type cases ( $76.7 \%$; $\mathrm{p}=0.001$ ). Well known molecular prognostic biomarkers of CLL, such as unmutated IGHV gene status and 17p deletion also associated with a significantly shorter PFS, supporting the representativeness of the study cohort (Table 2). By multivariate analysis including variables showing a
multiplicity adjusted significant association with PFS, BIRC3 mutations maintained an independent association with PFS, with a HR of 2.8 (95\% C.I. 1.4-5.6, $\mathrm{p}=0.004$ ) (Table 2).

### 4.2 BIRC3 mutations associate with activation of non-canonical NF-кB signaling

In order to comprehensively map unique BIRC3 mutations in CLL, we compiled somatically confirmed variants identified in the current CLL study cohort with those identified in previous studies ${ }^{17}$ or listed in public CLL mutation catalogues (Figure 3 A ). Virtually all BIRC3 mutations were represented by frameshift or stop codons clustering in two hotspot regions comprised between amino acid 367-438 and amino acid 537-564. BIRC3 variants were predicted to generate aberrant truncated transcripts causing the elimination or truncation of the C-terminal RING domain of the BIRC3 protein. The RING domain of BIRC3 harbors the E3 ubiquitin ligase activity that is essential for proteasomal degradation of MAP3K14, the central activating kinase of the noncanonical NF-кB signaling. This observation points to non-canonical NF-кB activation through MAP3K14 stabilization as the predicted functional consequence of BIRC3 mutations in CLL. The non-canonical NF-кB signaling was profiled by immunoblotting in B-cell tumor cell lines and primary CLL cells with different genetic make-up in the non-canonical NF-кB pathway to verify whether BIRC3 mutations lead to constitutive non-canonical NF-кВ activation. Additional genetic features of the above mentioned cell lines and primary CLL cells are shown in Table S4. In the VL51 SMZL cell line and in the MEC1 CLL cell lines, both harboring endogenous truncating mutations of the BIRC3 gene, non-canonical NF-KB signaling was constitutively active, as documented by the stabilization of MAP3K14, phosphorylation of NF-KB2, its processing from p100 to p52, as well as nuclear localization of p52 (Figure 3 B-D). Consistent with the biochemical clues of non-canonical NF-кB activation, the gene expression signature of the VL51 and MEC1
cell lines was significantly enriched of non-canonical NF-кB target genes (Figure 3 E-F). Non-canonical NFкB signaling in BIRC3 mutated cells was consistent with that of MCL cell lines known to harbor a disrupted TRAF3/MAP3K14-TRAF2/BIRC3 negative regulatory complex by loss of TRAF3 or TRAF2. ${ }^{19}$ As BIRC3 mutated cell lines, also primary CLL samples harboring inactivating mutations of BIRC3 showed stabilization of MAP3K14 and NF-кB2 processing from p100 to p52 (Figure 3 C), thus confirming that non-canonical NF-кB activation is also a feature of primary cells harboring BIRC3 variants.

MAP3K14 was genetically targeted by shRNA to test whether BIRC3 mutated cells are addicted of its stabilization. Compared to non-targeting shRNA, the most efficient anti MAP3K14 shRNA-D resulted in a partial silencing of MAP3K14 and in a decreased NF- $\mathrm{KB}_{2}$ processing from p 100 to p 52 . This translated into a reduced cell viability of the BIRC3 mutated VL51 cell line transduced with shRNA-D. This observation indicates that MAP3K14 stabilization is a vulnerability of BIRC3 mutated cells (Figure 4).

In order to test the contribution of BTK to noncanonical NF-кB signaling when it is activated through BIRC3 mutations, BIRC3 mutated cell lines, as well as cell lines harboring a disrupted or competent TRAF3/MAP3K14-TRAF2/BIRC3 negative regulatory complex were treated with ibrutinib at different dosage and non-canonical NF-кB signaling activation probed by immunoblotting of the NF-кB2 processing from p100 to p52. Processing from p100 to p52 was unaffected by ibrutinib treatment in cell lines harboring BIRC3 mutations (Figure 5) or a disrupted TRAF3/MAP3K14-TRAF2/BIRC3 negative regulatory complex consistent with the notion that BIRC3 mutations activate non-canonical NF-кB by bypassing BTK blockade by ibrutinib ${ }^{19}$.

### 4.3 BIRC3 mutations confer resistance to fludarabine in primary CLL cells

We performed in vitro pharmacological studies on primary CLL cells to verify the vulnerabilities of BIRC3 mutated cells. CLL cells purified from patients carrying BIRC3 mutations were treated with increasing doses of fludarabine. Drug-induced apoptosis was compared to samples harboring TP53 mutations, which represent a control for fludarabine resistance. CLL cells devoid of genetic lesions on either BIRC3 or TP53 were adopted as a control cohort for fludarabine sensitivity. Molecular characteristic of the ex-vivo CLL cells are listed in Table S5. BIRC3 mutated cells showed a delayed fludarabine-induced cell death, as no response was observed after 24-hour treatment, at variance with TP53 and BIRC3 wild type samples. At this time point, cell viability curves of BIRC3 mutated samples were almost completely overlapping with that of TP53 disrupted samples, which are known to be fludarabine resistant (Figure 6 A). At 48 hours, BIRC3 mutated cells had viability that was lower than that of TP53 mutated samples, but higher than that of TP53 and BIRC3 wild type samples (Figure 6 B).

In order to assess whether BIRC3 mutations interfere with apoptosis, primary CLL cells were treated with venetoclax. Venetoclax treatment resulted in similar reduction of cell viability in BIRC3 mutated cells, TP53 mutated cells and BIRC3/TP53 wild type cells (Figure 6 C, D). Such divergent sensitivity to fludarabine and venetoclax of BIRC3 mutated CLL cells indirectly suggests that BIRC3 mutations likely affect the upstream DNA damage response pathway rather than the downstream apoptosis among mechanisms of cell death induction.

## 5. DISCUSSION

The results of this study provide the evidence that: i) BIRC3 mutated patients fail FCR chemoimmunotherapy analogous to cases harboring TP53 disruption; and that ii) BIRC3 mutations associate with activation of the non-canonical NF-кB pathway and with resistance to fludarabine in vitro.

The mere presence of somatic mutations is insufficient to implicate a gene in cancer. Cancer geneticists and bioinformaticians differentiate "passengers" events, likely being randomly acquired, to distinguish them from mutations targeting candidate "cancer driver" genes, likely implicated in the tumor biology, according to a statistical definition. Any given gene is labeled as candidate "cancer driver" if it harbors somatic point mutations at a statistically significant rate or pattern in cancer samples. In CLL, more than 40 genes fulfill the statistical definition of candidate "cancer driver", including BIRC3, but few of them are biologically validated (i.e. SF3B1, NOTCH1, TP53, ATM, FBXW7). ${ }^{8,9,20-23}$ The BIRC3 (Baculoviral IAP Repeat Containing 3) gene codes for a protein that ubiquitinates and negatively regulates the central activating kinase of the non-canonical NF-KB pathway, namely MAP3K14. ${ }^{24,25}$ In lymphoid malignancies, the NF-кB pathway is a pivotal and positive mediator of cell proliferation and survival. ${ }^{7,26,27}$ In CLL, BIRC3 mutations are absent in monoclonal B-cell lymphocytosis (MBL) patients, are rare at the time of diagnosis (3-4\%), but are detectable in approximately $25 \%$ of fludarabine refractory patients. ${ }^{17}$ In this study, we verified the biological consequences of BIRC3 mutations by showing that they associate with activation of the noncanonical NF-кB pathway, that BIRC3 mutated lymphoid cells are addicted of non-canonical NF-кB pathway, and that BIRC3 mutated CLL are resistant to fludarabine both in vitro and in patients. It still remains to be clarified whether NF-кB activation is the only molecular pathway that causes chemo-refractoriness in BIRC3 mutated CLL or whether other mechanisms are also involved. ${ }^{27-31}$

The introduction of FCR has represented a breakthrough in the management of young and fit CLL patients with an improvement in both PFS and OS compared to previous regiments. In both clinical trials and real life cohorts, ${ }^{3-5}$ IGHV mutation status and TP53 disruption sorted out as strong predictors of poor response to FCR. However, these molecular biomarkers do not fully capture all high-risk patients destined to relapse. We propose BIRC3 mutations as a new biomarker for the identification of high-risk patients failing FCR similarly to cases harboring TP53 disruption. If validated in independent series, BIRC3 mutations may turn out as a new molecular predictor of FCR resistance to be use for selecting patients to be treated with novel targeted agents.

Non-canonical NF-кB activation by BIRC3 mutations by-pass the block of BTK by ibrutinib. Consistently, NF-кB activation and cell survival is unaffected by ibrutinib in both CLL cells (our study) and mantle cell lymphoma cells. ${ }^{19}$ If this pre-clinical evidence will be validated in ibrutinib-treated patients, BIRC3 mutations may translate in a biomarker also for informing selection of novel agents.

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## 7. TABLES

Table 1. Clinical data of FCR-treated CLL patients

| Characteristics | Number of patients (\%) | Total |
| :---: | :---: | :---: |
| Male | $198(69.0 \%)$ | $\mathrm{N}=287$ |
| Female | $89(31.0 \%)$ |  |
| Binet A | $33(11.5 \%)$ | $\mathrm{N}=287$ |
| Binet B-C | $254(88.5 \%)$ | $\mathrm{N}=280$ |
| IGHV mutated | $100(35.7 \%)$ | $\mathrm{N}=274$ |
| IGHV unmutated | $180(64.3 \%)$ | $\mathrm{N}=273$ |
| 17p deletion | $13(4.7 \%)$ | $\mathrm{N}=273$ |
| No 17p deletion | $261(95.3 \%)$ | $\mathrm{N}=272$ |
| 11q deletion | $47(17.2 \%)$ |  |
| No 11q deletion | $226(82.8 \%)$ |  |
| 13q deletion | $111(40.7 \%)$ |  |
| No 13q deletion | $162(50.3 \%)$ |  |
| Trisomy 12 | $50(18.4 \%)$ |  |
| No Trisomy 12 | $222(81.6 \%)$ |  |
| Median Follow-up (years) |  |  |
| Median PFS | 6.8 |  |
| PFS \% (7-years) | 4.6 |  |
| Median OS (years) | $11.7 \%$ |  |
| OS \% (7-years) | $75.5 \%$ |  |

PFS, progression free survival; OS, overall survival; IGHV, immunoglobulin heavy variable gene.

Table 2. Univariate and multivariate analysis of PFS

|  | Univariate analysis |  |  |  |  | Multivariate analysis |  |  |  | Internal bootstrapping validation |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Characteristics | 7-y PFS <br> (\%) | Median PFS (y) | 95\% CI | P | P* | HR | LCI | UCI | P | HR | LCI | UCI | Bootstrapping selection (\%) |
| Binet A | 40.3\% | 4.5 | 2.4-6.6 | 0.356 | - | - | - | - | - | - | - | - | - |
| Binet B-C | 30.0\% | 4.6 | 3.8-5.4 |  |  | - | - | - |  | - | - | - |  |
| IGHV mutated | 49.3\% | 6.5 | 3.8-9.2 | <0.001 | 0.003 |  |  |  | 0.001 |  |  |  | 98.8\% |
| IGHV unmutated | 23.0\% | 3.9 | 3.5-4.4 |  |  | 1.8 | 1.3 | 2.6 |  | 1.9 | 1.3 | 2.7 |  |
| No 11q deletion | 33.4\% | 5.0 | 4.2-5.9 | 0.025 | 0.700 | - | - | - | - | - | - | - | - |
| 11q deletion | 13.9\% | 3.6 | 2.4-4.9 |  |  | - | - | - |  | - | - | - |  |
| No 17p deletion | 33.0\% | 4.8 | 4.1-5.6 | <0.0001 | <0.0001 | - | - | 5 | <0.0001 |  | - | - | 99.5\% |
| 17p deletion | nr | 1.1 | 0-2.6 |  |  | 4.0 | 2.2 | 7.5 |  | 4.9 | 2.5 | 9.8 |  |
| TP53 Wild type | 33.8\% | 5.4 | 4.3-5.8 | <0.0001 | <0.001 | - |  |  | 0.030 | - |  | - | 73.3\% |
| TP53 Mutated | nr | 2.8 | 2.0-3.5 |  |  | 1.7 | 1.1 | 2.8 |  | 1.8 | 1.1 | 3 |  |
| BIRC3 Wild type | 32.2\% | 4.8 | 4.1-5.6 | <0.001 | 0.005 | - | - | - | 0.004 | - | - | - | 91.1\% |
| BIRC3 Mutated | nr | 2.2 | 0.9-3.5 |  |  | 2.8 | 1.4 | 5.6 |  | 3.4 | 1.6 | 7.3 |  |
| EGR2Wild type | 31.5\% | 4.7 | 3.9-5.4 | 0.015 | 0.420 | - | - | - | - | - | - | - | - |
| EGR2Mutated | nr | 1.5 | 0-3.8 |  |  | - | - | - |  | - | - | - |  |
| ATM Wild type | 32.5\% | 4.8 | 4.1-5.6 | 0.029 | 0.812 | - | - | - | - | - | - | - | - |
| ATM Mutated | nr | 3.2 | 2.4-4.1 |  |  | - | - | - |  | - | - | - |  |

$P$, P-value; $P^{*}$, Bonferroni correction; PFS, progression free survival; Cl , confidence interval; HR , hazard ratio; LCI, lower confidence interval; UCI, upper confidence interval; IGHV, immunoglobulin heavy variable gene; nr, not reached

## 8. FIGURES LEGENDS

Figure 1. Mutational profile of the FCR-treated cohort. Case-level mutational profiles of 287 patients FCRtreated patients. Each column represents one tumor sample, each row represents one gene. The fraction of tumors with mutations in each gene is plotted on the right. The number and type of mutations in each patient is plotted above the heat map. Mutations are highlighted in red. IGHV mutational status, 17p deletion and 11q deletion are plotted in the bottom of the heatmap.

Figure 2. Kaplan-Meier estimates of progression free survival in BIRC3 mutated patients. (A) Cases harboring BIRC3 mutations are represented by the red line. Cases wild type for this gene are represented by the blue line. (B) Cases harboring BIRC3 mutations are represented by the red line. Cases harboring TP53 disruption (including TP53 mutation and/or 17p deletion) are represented by the yellow line. Patients devoid of BIRC3 mutation and TP53 disruption are represented by the blue line. The Log-rank statistics p values are indicated adjacent curves.

Figure 3: Non-canonical NF-кВ pathway is active in BIRC3 mutated CLL cell lines and primary samples.
(A) Disposition of BIRC3 mutations across the protein. The mutations identified by Landau et al. ${ }^{9}$, Puente et al. ${ }^{8}$ and from public CLL mutation catalogue (COSMIC v85) are plotted in grey. Individual BIRC3 mutations identified in the current studied cohort and in our previous study ${ }^{17}$ are plotted in red. (B) Western blot analysis of BIRC3 protein expression and NF-кB2 activation and processing in the SMZL cell lines SSK41, VL51 and in the CLL cell line MEC1, carrying wild-type or disrupted BIRC3. The MAVER-1 and

Z-138 cell lines were used as positive controls of non-canonical NF-кB activation, harboring genetic activation of non-canonical NF-kB signaling. The JEKO-1 and HEK 293T cell lines were used as negative controls for non-canonical NF-кB signalling. $\alpha$-actin was used as a loading control. Color codes indicate the gene status in each cell lines. The aberrant BIRC3 band expressed in MEC1 and VL51 cell lines correspond in size to the predicted BIRC3-truncated protein, encoded by the mutant allele. (C) Western blot analysis showing BIRC3 expression and NF-kB2 processing in purified primary tumor cells from 5 CLL and SMZL patients carrying wild-type or disrupted BIRC3. Color codes indicate the gene status in each cell lines. The aberrant BIRC3 bands in patients 09321, 14462 and 12603 correspond in size to the predicted BIRC3-truncated protein encoded by the mutant allele. $\alpha$-actin was used as a loading control. (D) Western blot of whole cell extract, cytoplasmic or nuclear fractions of the SMZL and CLL cell lines probed for the NF-кB2 subunits p100 and p52. The MAVER-1 and Z-138 cell lines served as positive controls while the JEKO-1 and HEK 293T cell lines were used as negative controls. $ß$-tubulin and BRG1 served as controls for the purity of the cytoplasmic and nuclear fractionations, respectively. (E) GSEA enrichment score and distribution of non-canonical NF-кB target genes along the rank of transcripts differentially expressed in the SMZL cell lines SSK41, VL51 and in the CLL cell line MEC1. The JEKO-1 cell line was used as negative control. (F) Validation of non-canonical NF-кB target genes expression in the same SMZL and CLL cell lines as determined by quantitative real-time RT-PCR. Changes of genes expression were normalized to GAPDH expression; relative quantities were $\log _{2}$ normalized to control samples (MCL cell line JEKO-1).

Figure 4: Knockdown of MAP3K14 by RNA interference in VL51 cells. (A) Western blot analysis for MAP3K14 expression and for NF-кB2 processing of p100 to p52. (B) VL51 cells viability assessed by trypan
blue after transduction with lentiviral vectors expressing the shRNAD_MAP3K14 (in red), a scrambled shRNA (in blu), and in non transfected cells (in green).

Figure 5: Non-canonical NF-кB pathway is not switched off by ibrutinib in BIRC3 mutated cell lines. Western blot showing p100/p52 expression in (A) MEC1 and (B) VL51 cell lines that harbors BIRC3 mutations. (C) MAVER-1 and (D) Z-138 cell lines, known to be affected by noncanonical NF-кB pathway gene mutations and resistant to ibrutinib were used as positive controls. (E) JEKO-1 cell line, known to be devoid of NF-кB pathway gene mutations and sensitive to ibrutinib was used as negative control. All cell lines were treated with different concentrations of ibrutinib for 72 and 96 hours.

Figure 6: Responses of primary cells lines to fludarabine and venetoclax. Viability of BIRC3 mutated ( $\mathrm{n}=$ 6 patients, red line), TP53 mutated ( $n=8$ patients, black line) and wild type ( $n=7$ patients, blue line) primary CLL cells treated with different concentrations of fludarabine for (A) 24 hours and (B) 48 hours and of venetoclax for (C) 24 hours and (D) 48 hours The pairwise $p$ values have been listed in the tables below the respective figures. $M$, mutated; WT , wild type; $N T$, not treated.

## 8. FIGURES

Figure 1. Mutational profile of the FCR-treated cohort.


Figure 2. Kaplan-Meier estimates of progression free survival in BIRC3 mutated patients.


Figure 3: Non-canonical NF-кB pathway is active in BIRC3 mutated CLL cell lines and primary samples.

A


B


D


E


F


Figure 4: Knockdown of MAP3K14 by RNA interference in VL51 cells.


Figure 5: Non-canonical NF-KB pathway is not switched off by ibrutinib in BIRC3 mutated cell lines.


Figure 6: Responses of primary cells lines to fludarabine and venetoclax.

A $24 \mathrm{~h}{ }^{125}$

|  | NT | Fludarabine <br> $\mathbf{1 m M}$ | Fludarabine <br> 5 mM | Fludarabine <br> $\mathbf{1 0 m M}$ | Fludarabine <br> $\mathbf{2 5 m M}$ |
| :---: | :---: | :---: | :---: | :---: | :---: |
| TP53-M vs. <br> BIRC3-M | $\mathrm{p}>0.9999$ | $\mathrm{p}=0.9447$ | $\mathrm{p}=0.8923$ | $\mathrm{p}=0.9997$ | $\mathrm{p}=0.4311$ |
| TP53-M vs. <br> WT | $\mathrm{p}>0.9999$ | $\mathrm{p}=0.0567$ | $\mathrm{p}=0.0031$ | $\mathrm{p}=0.0078$ | $\mathrm{p}=0.0005$ |
| BIRC3-M <br> vs. WT | $\mathrm{p}>0.9999$ | $\mathrm{p}=0.1117$ | $\mathrm{p}=0.0112$ | $\mathrm{p}=0.0115$ | $\mathrm{p}=0.0184$ |

$\pm$ TP53-M; BIRC3-WT

- TP53-WT; BIRC3-M
- TP53-WT; BIRC3-WT


|  | Fludarabine ( $\mu \mathrm{M}$ ) |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  | NT | $\begin{array}{\|c\|} \hline \begin{array}{c} \text { Fludarabine } \\ 1 \mathrm{mM} \end{array} \\ \hline \end{array}$ | Fludarabine 5 mM | $\begin{array}{\|c\|} \hline \begin{array}{c} \text { Fludarabine } \\ 10 \mathrm{mM} \end{array} \\ \hline \end{array}$ | $\begin{array}{\|c\|} \hline \begin{array}{c} \text { Fludarabine } \\ 25 \mathrm{mM} \end{array} \\ \hline \end{array}$ |
| TP53-M vs. BIRC3-M | $\mathrm{p}>0.9999$ | $\mathrm{p}=0.0837$ | $\mathrm{p}=0.0309$ | $\mathrm{p}=0.0320$ | $\mathrm{p}=0.0053$ |
| $\begin{gathered} \hline T P 53-\mathrm{M} \text { vs. } \\ \text { WT } \end{gathered}$ | p>0.9999 | $\mathrm{p}=0.0737$ | $\mathrm{p}=0.0001$ | $\mathrm{p}=0.0004$ | p<0.0001 |
| $\begin{gathered} \hline \text { BIRC3-M } \\ \text { vs. WT } \\ \hline \end{gathered}$ | $\mathrm{p}>0.9999$ | $\mathrm{p}=0.9982$ | $\mathrm{p}=0.2043$ | $\mathrm{p}=0.4756$ | $\mathrm{p}=0.2742$ |



D 48 h


|  | NT | $\begin{array}{\|c\|} \hline \text { Venetoclax } \\ 5 \mathrm{nM} \end{array}$ | Venetoclax 10nM | Venetoclax 50 nM | $\begin{array}{\|c\|} \hline \text { Venetoclax } \\ 100 \mathrm{nM} \end{array}$ | $\begin{array}{\|c\|} \hline \text { Venetoclax } \\ \mathbf{5 0 0 n M} \end{array}$ | $\begin{array}{\|c\|} \hline \text { Venetoclax } \\ 2000 \mathrm{nM} \end{array}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\begin{array}{\|c} \text { TP53-M vs. } \\ \text { BIRC } 3-\mathrm{M} \end{array}$ | $\mathrm{p}>0.9999$ | $\mathrm{p}=0.9512$ | $\mathrm{p}=0.5282$ | $\mathrm{p}=0.9119$ | $\mathrm{p}=0.4524$ | $\mathrm{p}=0.7524$ | $\mathrm{p}=0.7524$ |
| $\underset{\text { WT }}{T P 53-\mathrm{M}} \mathrm{vs} .$ | $\mathrm{p}>0.9999$ | $\mathrm{p}=0.9800$ | $\mathrm{p}=0.6789$ | $\mathrm{p}=0.3778$ | $\mathrm{p}=0.7438$ | $\mathrm{p}=0.8597$ | $\mathrm{p}=0.8597$ |
| $\begin{array}{\|c} \hline \begin{array}{c} \text { BIRC3-M } \\ \text { vs. WT } \end{array} \\ \hline \end{array}$ | $\mathrm{p}>0.9999$ | $\mathrm{p}=0.8844$ | $\mathrm{p}=0.9547$ | $\mathrm{p}=0.2959$ | $\mathrm{p}=0.8518$ | $\mathrm{p}=0.9722$ | $\mathrm{p}=0.9722$ |

## 9. SUPPLEMENTARY TABLES

Table S1. Target region

| Gene | hg19 chromosome | hg19 coding exon start plus splice site (2bp) | hg19 coding exon stop plus splice site (2bp) |
| :---: | :---: | :---: | :---: |
| NRAS | chr1 | 115251156 | 115251277 |
|  | chr1 | 115252188 | 115252351 |
|  | chr1 | 115256419 | 115256601 |
|  | chr1 | 115258669 | 115258781 |
| XPO1 | chr2 | 61705955 | 61706103 |
|  | chr2 | 61708318 | 61708418 |
|  | chr2 | 61709513 | 61709676 |
|  | chr2 | 61710090 | 61710228 |
|  | chr2 | 61711070 | 61711242 |
|  | chr2 | 61712901 | 61713099 |
|  | chr2 | 61715298 | 61715408 |
|  | chr2 | 61715721 | 61715908 |
|  | chr2 | 61717775 | 61717913 |
|  | chr2 | 61719168 | 61719335 |
|  | chr2 | 61719458 | 61719618 |
|  | chr2 | 61719700 | 61719885 |
|  | chr2 | 61720048 | 61720190 |
|  | chr2 | 61721027 | 61721228 |
|  | chr2 | 61722588 | 61722750 |
|  | chr2 | 61724012 | 61724144 |
|  | chr2 | 61725806 | 61725929 |
|  | chr2 | 61725998 | 61726050 |
|  | chr2 | 61726846 | 61727031 |
|  | chr2 | 61729129 | 61729177 |
|  | chr2 | 61729382 | 61729447 |
|  | chr2 | 61749744 | 61749820 |
|  | chr2 | 61753553 | 61753658 |
|  | chr2 | 61760905 | 61761032 |
| SF3B1 | chr2 | 198257027 | 198257187 |
|  | chr2 | 198257694 | 198257914 |
|  | chr2 | 198260778 | 198261054 |
|  | chr2 | 198262707 | 198262842 |
|  | chr2 | 198263183 | 198263307 |
|  | chr2 | 198264777 | 198264892 |
|  | chr2 | 198264974 | 198265160 |
|  | chr2 | 198265437 | 198265662 |
|  | chr2 | 198266122 | 198266251 |
|  | chr2 | 198266464 | 198266614 |
|  | chr2 | 198266707 | 198266856 |
|  | chr2 | 198267278 | 198267552 |
|  | chr2 | 198267671 | 198267761 |
|  | chr2 | 198268307 | 198268490 |
|  | chr2 | 198269798 | 198269903 |
|  | chr2 | 198269997 | 198270198 |
|  | chr2 | 198272720 | 198272845 |
|  | chr2 | 198273091 | 198273307 |
|  | chr2 | 198274492 | 198274733 |


|  | chr2 | 198281463 | 198281637 |
| :---: | :---: | :---: | :---: |
|  | chr2 | 198283231 | 198283314 |
|  | chr2 | 198285150 | 198285268 |
|  | chr2 | 198285751 | 198285859 |
|  | chr2 | 198288530 | 198288700 |
|  | chr2 | 198299694 | 198299723 |
| MYD88 | chr3 | 38180153 | 38180521 |
|  | chr3 | 38181353 | 38181491 |
|  | chr3 | 38181877 | 38182061 |
|  | chr3 | 38182246 | 38182341 |
|  | chr3 | 38182621 | 38182777 |
| FBXW7 | chr4 | 153244035 | 153244303 |
|  | chr4 | 153245334 | 153245548 |
|  | chr4 | 153247156 | 153247385 |
|  | chr4 | 153249358 | 153249543 |
|  | chr4 | 153250822 | 153250939 |
|  | chr4 | 153251882 | 153252022 |
|  | chr4 | 153253746 | 153253873 |
|  | chr4 | 153258952 | 153259090 |
|  | chr4 | 153268080 | 153268225 |
|  | chr4 | 153271192 | 153271278 |
|  | chr4 | 153332453 | 153332955 |
| IRF4 | chr6 | 393150 | 393370 |
|  | chr6 | 394818 | 395009 |
|  | chr6 | 395844 | 395937 |
|  | chr6 | 397105 | 397254 |
|  | chr6 | 398825 | 398937 |
|  | chr6 | 401421 | 401779 |
|  | chr6 | 405015 | 405132 |
|  | chr6 | 407452 | 407600 |
| NFKBIE | chr6 | 44226953 | 44227023 |
|  | chr6 | 44227777 | 44228021 |
|  | chr6 | 44228185 | 44228278 |
|  | chr6 | 44229360 | 44229587 |
|  | chr6 | 44230294 | 44230401 |
|  | chr6 | 44232716 | 44233502 |
| POT1 | chr7 | 124464016 | 124464130 |
|  | chr7 | 124465304 | 124465413 |
|  | chr7 | 124467266 | 124467361 |
|  | chr7 | 124469306 | 124469398 |
|  | chr7 | 124475331 | 124475470 |
|  | chr7 | 124481025 | 124481234 |
|  | chr7 | 124482859 | 124483019 |
|  | chr7 | 124486994 | 124487054 |
|  | chr7 | 124491924 | 124492007 |
|  | chr7 | 124493024 | 124493194 |
|  | chr7 | 124499009 | 124499168 |
|  | chr7 | 124503402 | 124503696 |
|  | chr7 | 124510963 | 124511097 |
|  | chr7 | 124532318 | 124532436 |
|  | chr7 | 124537217 | 124537227 |
| BRAF | chr7 | 140434397 | 140434572 |
|  | chr7 | 140439610 | 140439748 |
|  | chr7 | 140449085 | 140449220 |
|  | chr7 | 140453073 | 140453195 |


|  | chr7 | 140453985 | 140454035 |
| :---: | :---: | :---: | :---: |
|  | chr7 | 140476710 | 140476890 |
|  | chr7 | 140477789 | 140477877 |
|  | chr7 | 140481374 | 140481495 |
|  | chr7 | 140482819 | 140482959 |
|  | chr7 | 140487346 | 140487386 |
|  | chr7 | 140494106 | 140494269 |
|  | chr7 | 140500160 | 140500283 |
|  | chr7 | 140501210 | 140501362 |
|  | chr7 | 140507758 | 140507864 |
|  | chr7 | 140508690 | 140508797 |
|  | chr7 | 140534407 | 140534674 |
|  | chr7 | 140549909 | 140550014 |
|  | chr7 | 140624364 | 140624503 |
| NOTCH1 | chr9 | 139390512 | 139392020 |
| NOTCH1_3'UTR | chr9 | 139388893 | 139390524 |
| PAX5 | chr9 | 37370906 | 37371645 |
|  | chr9 | 37033543 | 37034202 |
| EGR2 | chr10 | 64572967 | 64574230 |
|  | chr10 | 64575619 | 64575789 |
| NFKB2 | chr10 | 104155676 | 104155777 |
|  | chr10 | 104155999 | 104156101 |
|  | chr10 | 104156155 | 104156296 |
|  | chr10 | 104156471 | 104156590 |
|  | chr10 | 104156650 | 104156822 |
|  | chr10 | 104157048 | 104157175 |
|  | chr10 | 104157273 | 104157452 |
|  | chr10 | 104157727 | 104157852 |
|  | chr10 | 104157958 | 104158064 |
|  | chr10 | 104158131 | 104158290 |
|  | chr10 | 104158485 | 104158631 |
|  | chr10 | 104159034 | 104159264 |
|  | chr10 | 104159323 | 104159485 |
|  | chr10 | 104159826 | 104159961 |
|  | chr10 | 104160024 | 104160258 |
|  | chr10 | 104160401 | 104160591 |
|  | chr10 | 104160693 | 104160816 |
|  | chr10 | 104160926 | 104161098 |
|  | chr10 | 104161225 | 104161325 |
|  | chr10 | 104161491 | 104161684 |
|  | chr10 | 104161794 | 104161926 |
|  | chr10 | 104161998 | 104162143 |
| ATM | chr11 | 108098352 | 108098425 |
|  | chr11 | 108098501 | 108098617 |
|  | chr11 | 108099903 | 108100052 |
|  | chr11 | 108106395 | 108106563 |
|  | chr11 | 108114678 | 108114847 |
|  | chr11 | 108115513 | 108115755 |
|  | chr11 | 108117689 | 108117856 |
|  | chr11 | 108119658 | 108119831 |
|  | chr11 | 108121426 | 108121801 |
|  | chr11 | 108122562 | 108122760 |
|  | chr11 | 108123542 | 108123641 |
|  | chr11 | 108124539 | 108124768 |
|  | chr11 | 108126940 | 108127069 |


|  | chr11 | 108128206 | 108128335 |
| :---: | :---: | :---: | :---: |
|  | chr11 | 108129711 | 108129804 |
|  | chr11 | 108137896 | 108138071 |
|  | chr11 | 108139135 | 108139338 |
|  | chr11 | 108141789 | 108141875 |
|  | chr11 | 108141976 | 108142135 |
|  | chr11 | 108143257 | 108143336 |
|  | chr11 | 108143447 | 108143581 |
|  | chr11 | 108150216 | 108150337 |
|  | chr11 | 108151720 | 108151897 |
|  | chr11 | 108153435 | 108153608 |
|  | chr11 | 108154952 | 108155202 |
|  | chr11 | 108158325 | 108158444 |
|  | chr11 | 108159702 | 108159832 |
|  | chr11 | 108160327 | 108160530 |
|  | chr11 | 108163344 | 108163522 |
|  | chr11 | 108164038 | 108164206 |
|  | chr11 | 108165652 | 108165788 |
|  | chr11 | 108168012 | 108168111 |
|  | chr11 | 108170439 | 108170614 |
|  | chr11 | 108172373 | 108172518 |
|  | chr11 | 108173578 | 108173758 |
|  | chr11 | 108175400 | 108175581 |
|  | chr11 | 108178622 | 108178713 |
|  | chr11 | 108180885 | 108181044 |
|  | chr11 | 108183136 | 108183227 |
|  | chr11 | 108186548 | 108186640 |
|  | chr11 | 108186736 | 108186842 |
|  | chr11 | 108188098 | 108188250 |
|  | chr11 | 108190679 | 108190787 |
|  | chr11 | 108192026 | 108192149 |
|  | chr11 | 108196035 | 108196273 |
|  | chr11 | 108196783 | 108196954 |
|  | chr11 | 108198370 | 108198487 |
|  | chr11 | 108199746 | 108199967 |
|  | chr11 | 108200939 | 108201150 |
|  | chr11 | 108202169 | 108202286 |
|  | chr11 | 108202604 | 108202766 |
|  | chr11 | 108203487 | 108203629 |
|  | chr11 | 108204611 | 108204697 |
|  | chr11 | 108205694 | 108205838 |
|  | chr11 | 108206570 | 108206690 |
|  | chr11 | 108213947 | 108214100 |
|  | chr11 | 108216468 | 108216637 |
|  | chr11 | 108218004 | 108218094 |
|  | chr11 | 108224491 | 108224609 |
|  | chr11 | 108225536 | 108225603 |
|  | chr11 | 108235807 | 108235947 |
|  | chr11 | 108236050 | 108236235 |
| BIRC3 | chr11 | 102195241 | 102196095 |
|  | chr11 | 102196195 | 102196298 |
|  | chr11 | 102198781 | 102198863 |
|  | chr11 | 102199626 | 102199678 |
|  | chr11 | 102201728 | 102201974 |
|  | chr11 | 102206695 | 102206953 |


|  | chr11 | 102207489 | 102207534 |
| :---: | :---: | :---: | :---: |
|  | chr11 | 102207638 | 102207833 |
| KRAS | chr12 | 25368375 | 25368496 |
|  | chr12 | 25378546 | 25378709 |
|  | chr12 | 25380166 | 25380348 |
|  | chr12 | 25398206 | 25398318 |
| MAP2K1 | chr15 | 66679683 | 66679767 |
|  | chr15 | 66727362 | 66727577 |
|  | chr15 | 66729081 | 66729232 |
|  | chr15 | 66735615 | 66735697 |
|  | chr15 | 66736991 | 66737047 |
|  | chr15 | 66774090 | 66774219 |
|  | chr15 | 66777325 | 66777531 |
|  | chr15 | 66779563 | 66779632 |
|  | chr15 | 66781550 | 66781616 |
|  | chr15 | 66782053 | 66782103 |
|  | chr15 | 66782837 | 66782955 |
| MGA | chr15 | 41961082 | 41962166 |
|  | chr15 | 41988262 | 41989231 |
|  | chr15 | 41991040 | 41991149 |
|  | chr15 | 41991251 | 41991367 |
|  | chr15 | 41999915 | 42000067 |
|  | chr15 | 42000291 | 42000416 |
|  | chr15 | 42002878 | 42003557 |
|  | chr15 | 42005338 | 42005704 |
|  | chr15 | 42019367 | 42019614 |
|  | chr15 | 42021351 | 42021557 |
|  | chr15 | 42026699 | 42026802 |
|  | chr15 | 42028368 | 42028906 |
|  | chr15 | 42032240 | 42032411 |
|  | chr15 | 42034733 | 42035380 |
|  | chr15 | 42040824 | 42041135 |
|  | chr15 | 42041298 | 42042823 |
|  | chr15 | 42046624 | 42046775 |
|  | chr15 | 42049955 | 42050057 |
|  | chr15 | 42052510 | 42052737 |
|  | chr15 | 42053926 | 42054058 |
|  | chr15 | 42054316 | 42054570 |
|  | chr15 | 42057073 | 42057270 |
|  | chr15 | 42058191 | 42059488 |
| TP53 | chr17 | 7572927 | 7573010 |
|  | chr17 | 7573925 | 7574035 |
|  | chr17 | 7576851 | 7576928 |
|  | chr17 | 7577017 | 7577157 |
|  | chr17 | 7577497 | 7577610 |
|  | chr17 | 7578175 | 7578291 |
|  | chr17 | 7578369 | 7578556 |
|  | chr17 | 7579310 | 7579592 |
|  | chr17 | 7579698 | 7579723 |
|  | chr17 | 7579837 | 7579912 |
| IKZF3 | chr17 | 37922032 | 37922756 |
|  | chr17 | 37933893 | 37934030 |
|  | chr17 | 37944500 | 37944637 |
|  | chr17 | 37947658 | 37947846 |
|  | chr17 | 37948915 | 37949196 |


|  | chr17 | 37985629 | 37985751 |
| :---: | :---: | :---: | :---: |
|  | chr17 | 37988320 | 37988434 |
|  | chr17 | 38020322 | 38020429 |
| RPS15 | chr19 | 1438374 | 1438477 |
|  | chr19 | 1438785 | 1438911 |
|  | chr19 | 1440007 | 1440262 |
|  | chr19 | 1440337 | 1440471 |
| SAMHD1 | chr20 | 35521335 | 35521471 |
|  | chr20 | 35526223 | 35526364 |
|  | chr20 | 35526841 | 35526949 |
|  | chr20 | 35532558 | 35532654 |
|  | chr20 | 35533765 | 35533908 |
|  | chr20 | 35539619 | 35539738 |
|  | chr20 | 35540862 | 35540957 |
|  | chr20 | 35545123 | 35545235 |
|  | chr20 | 35545350 | 35545454 |
|  | chr20 | 35547765 | 35547924 |
|  | chr20 | 35555583 | 35555657 |
|  | chr20 | 35559161 | 35559280 |
|  | chr20 | 35563430 | 35563594 |
|  | chr20 | 35569440 | 35569516 |
|  | chr20 | 35575139 | 35575209 |
|  | chr20 | 35579837 | 35580046 |
| ASXL1 | chr20 | 30946548 | 30946665 |
|  | chr20 | 30954176 | 30954279 |
|  | chr20 | 30955479 | 30955582 |
|  | chr20 | 30956807 | 30956936 |
|  | chr20 | 31015920 | 31016061 |
|  | chr20 | 31016117 | 31016235 |
|  | chr20 | 31017130 | 31017244 |
|  | chr20 | 31017693 | 31017866 |
|  | chr20 | 31019113 | 31019297 |
|  | chr20 | 31019375 | 31019492 |
|  | chr20 | 31020672 | 31020798 |
|  | chr20 | 31021076 | 31021730 |
|  | chr20 | 31022224 | 31025151 |
| ZMYM3 | chrX | 70460766 | 70460960 |
|  | chrX | 70461075 | 70461196 |
|  | chrX | 70462018 | 70462276 |
|  | chrX | 70462818 | 70462936 |
|  | chrX | 70463677 | 70463832 |
|  | chrX | 70464150 | 70464322 |
|  | chrX | 70464638 | 70464745 |
|  | chrX | 70465187 | 70465337 |
|  | chrX | 70465516 | 70465694 |
|  | chrX | 70465834 | 70465950 |
|  | chrX | 70466201 | 70466364 |
|  | chrX | 70466443 | 70466544 |
|  | chrX | 70467193 | 70467362 |
|  | chrX | 70467582 | 70467758 |
|  | chrX | 70468010 | 70468164 |
|  | chrX | 70468286 | 70468376 |
|  | chrX | 70468534 | 70468653 |
|  | chrX | 70468867 | 70469021 |
|  | chrX | 70469309 | 70469531 |


|  | chrX | 70469874 | 70470055 |
| :--- | :--- | ---: | ---: |
|  | $\operatorname{chrX}$ | 70470280 | 70470578 |
|  | $\operatorname{chrX}$ | 70471026 | 70471096 |
|  | $\operatorname{chrX}$ | 70471406 | 70471453 |
|  | $\operatorname{chrX}$ | 70472437 | 70473105 |

Table S2. Target region with $\geq 1000 X$ and $\geq 2000 X$ coverage

| Material | Sample ID | Target Region Coverage (\%) |  |
| :---: | :---: | :---: | :---: |
|  |  | $\geq$ 1000X | $\geq$ 2000X |
| RNA | 3389 | 29.6 | 19.9 |
| RNA | 3390 | 49.8 | 28 |
| RNA | 3392 | 86.8 | 31.1 |
| RNA | 3393 | 55.2 | 33.6 |
| RNA | 3394 | 58.6 | 37.5 |
| RNA | 3395 | 91.7 | 38.8 |
| RNA | 3396 | 62 | 39.5 |
| gDNA | 4822 | 66.5 | 44.5 |
| gDNA | 4937 | 88.2 | 46.7 |
| gDNA | 4997 | 92.3 | 88.2 |
| gDNA | 5564 | 67.6 | 49 |
| gDNA | 5868 | 86 | 58 |
| gDNA | 6415 | 92.6 | 64 |
| gDNA | 7274 | 91 | 67.4 |
| gDNA | 7302 | 86.5 | 73.8 |
| gDNA | 8461 | 98.3 | 74.5 |
| gDNA | 9248 | 98.4 | 74.5 |
| gDNA | 9289 | 95.9 | 75.1 |
| gDNA | 9291 | 89.7 | 75.2 |
| gDNA | 10676 | 100 | 99.9 |
| gDNA | 10851 | 89.2 | 75.4 |
| gDNA | 12162 | 98.3 | 76.3 |
| gDNA | 12640 | 98.9 | 90.2 |
| gDNA | 12955 | 99.8 | 97.4 |
| gDNA | 13318 | 99.9 | 99.2 |
| gDNA | 13426 | 66.4 | 46.5 |
| gDNA | 13443 | 72.2 | 40.7 |
| gDNA | 14261 | 90.2 | 78 |
| gDNA | 14293 | 95.8 | 78.9 |
| gDNA | 14419 | 94.2 | 79.3 |
| gDNA | 14462 | 99.9 | 96.3 |
| gDNA | 14687 | 100 | 99.8 |
| gDNA | 14821 | 69.6 | 40.3 |
| gDNA | 14830 | 75.4 | 31.6 |
| gDNA | 15065 | 99.7 | 79.5 |
| gDNA | 15257 | 97.8 | 79.6 |
| gDNA | 15280 | 98.3 | 80 |
| gDNA | 15361 | 95.7 | 80.7 |
| gDNA | 15367 | 95.6 | 80.9 |
| gDNA | 15426 | 87.8 | 80.9 |
| gDNA | 15448 | 68.7 | 39.2 |
| gDNA | 15467 | 96.2 | 82.5 |
| gDNA | 15505 | 96.3 | 82.8 |
| gDNA | 15522 | 96.7 | 83.6 |
| gDNA | 15537 | 97.8 | 83.7 |
| gDNA | 15562 | 96.5 | 83.7 |
| gDNA | 15597 | 93.8 | 84.6 |
| gDNA | 15871 | 98.5 | 84.8 |
| gDNA | 15891 | 97.5 | 85 |
| gDNA | 15997 | 91.3 | 85.6 |


| gDNA | 16243 | 99.9 | 98.8 |
| :---: | :---: | :---: | :---: |
| gDNA | 17548 | 99.6 | 97.4 |
| gDNA | 17698 | 86.6 | 68 |
| gDNA | 18723 | 97.7 | 86 |
| gDNA | 18736 | 98.1 | 86.2 |
| gDNA | 19184 | 97.5 | 86.5 |
| gDNA | 19189 | 97 | 87 |
| gDNA | 19363 | 86.9 | 74.3 |
| gDNA | 19402 | 99.7 | 97.9 |
| gDNA | 19405 | 100 | 99.6 |
| gDNA | 19533 | 98.2 | 87 |
| gDNA | 20010 | 98.6 | 87.3 |
| gDNA | 20196 | 90.1 | 87.6 |
| gDNA | 20802 | 98.1 | 87.8 |
| gDNA | 20803 | 98.2 | 88 |
| gDNA | 20804 | 99.1 | 91.2 |
| gDNA | 20805 | 98.6 | 90.3 |
| gDNA | 20806 | 98.2 | 89.1 |
| gDNA | 20807 | 98.4 | 90.1 |
| gDNA | 20808 | 96.2 | 89.4 |
| gDNA | 20809 | 96.3 | 89.7 |
| gDNA | 20810 | 99.8 | 91.1 |
| gDNA | 20811 | 98.4 | 90.5 |
| gDNA | 20812 | 95.0 | 90.1 |
| gDNA | 20813 | 98.6 | 90.4 |
| gDNA | 20814 | 98.4 | 90.4 |
| gDNA | 20815 | 99.1 | 90.7 |
| gDNA | 20816 | 99.2 | 90.9 |
| gDNA | 20817 | 95.6 | 91.2 |
| gDNA | 20818 | 99.4 | 91.7 |
| gDNA | 20819 | 98.4 | 91.7 |
| gDNA | 20820 | 98.8 | 92.3 |
| gDNA | 20821 | 98.8 | 92.3 |
| gDNA | 20822 | 98.2 | 92.5 |
| gDNA | 20823 | 99.3 | 92.8 |
| gDNA | 20824 | 98.9 | 92.9 |
| gDNA | 20825 | 99.3 | 93.1 |
| gDNA | 20826 | 98.7 | 93.3 |
| gDNA | 20827 | 98.1 | 93.4 |
| gDNA | 20828 | 99.8 | 93.4 |
| gDNA | 20829 | 99 | 93.5 |
| gDNA | 20830 | 98.2 | 93.7 |
| gDNA | 20831 | 98.9 | 93.7 |
| gDNA | 20832 | 99.2 | 93.8 |
| gDNA | 20833 | 99 | 93.9 |
| gDNA | 20834 | 99.5 | 94.5 |
| gDNA | 20835 | 99.3 | 94.6 |
| gDNA | 20844 | 99.3 | 94.7 |
| gDNA | 20845 | 98.4 | 94.9 |
| gDNA | 20846 | 99.4 | 95.3 |
| gDNA | 20847 | 99.9 | 95.4 |
| gDNA | 20848 | 99.1 | 95.4 |
| gDNA | 20849 | 99.4 | 95.4 |
| gDNA | 20850 | 98.8 | 95.4 |
| gDNA | 20851 | 99.2 | 95.6 |


| gDNA | 20852 | 99.2 | 95.6 |
| :---: | :---: | :---: | :---: |
| gDNA | 20853 | 99.7 | 95.9 |
| gDNA | 20854 | 99.7 | 95.9 |
| gDNA | 20855 | 99 | 96.1 |
| gDNA | 20856 | 99.2 | 96.1 |
| gDNA | 20857 | 99.4 | 96.2 |
| gDNA | 20858 | 99.4 | 96.3 |
| gDNA | 20859 | 99.9 | 96.5 |
| gDNA | 20860 | 99.8 | 96.5 |
| gDNA | 20861 | 99.4 | 96.6 |
| gDNA | 20864 | 99.4 | 96.6 |
| gDNA | 20865 | 99.1 | 96.7 |
| gDNA | 20866 | 99.7 | 96.7 |
| gDNA | 20867 | 99.3 | 96.8 |
| gDNA | 20868 | 99.4 | 96.8 |
| gDNA | 20869 | 99.8 | 96.8 |
| gDNA | 20870 | 99.6 | 96.8 |
| gDNA | 20871 | 99.1 | 96.9 |
| gDNA | 20872 | 99.3 | 96.9 |
| gDNA | 20873 | 99.1 | 96.9 |
| gDNA | 20874 | 99.9 | 97 |
| gDNA | 20875 | 99.4 | 97.1 |
| gDNA | 20876 | 99.2 | 97.1 |
| gDNA | 20896 | 99.4 | 97.1 |
| gDNA | 20897 | 99.1 | 97.2 |
| gDNA | 21006 | 99.4 | 97.2 |
| gDNA | 21007 | 99.6 | 97.2 |
| gDNA | 21008 | 99.3 | 97.2 |
| gDNA | 21009 | 99.4 | 97.3 |
| gDNA | 21010 | 99.4 | 97.3 |
| gDNA | 21011 | 99.1 | 97.3 |
| gDNA | 21012 | 99.2 | 97.3 |
| gDNA | 21013 | 99.5 | 97.3 |
| gDNA | 21014 | 99.4 | 97.4 |
| gDNA | 21015 | 99.6 | 97.4 |
| gDNA | 21016 | 99.3 | 97.4 |
| gDNA | 21017 | 99.2 | 97.4 |
| gDNA | 21018 | 99.3 | 97.4 |
| gDNA | 21019 | 99.3 | 97.5 |
| gDNA | 21020 | 99.5 | 97.5 |
| gDNA | 21021 | 99.4 | 97.5 |
| gDNA | 21022 | 99.2 | 97.5 |
| gDNA | 21045 | 99.2 | 97.6 |
| gDNA | 21046 | 99.6 | 97.6 |
| gDNA | 21047 | 99.4 | 97.6 |
| gDNA | 21048 | 99.5 | 97.6 |
| gDNA | 21049 | 99.4 | 97.7 |
| gDNA | 21050 | 99.3 | 97.7 |
| gDNA | 21051 | 99.5 | 97.7 |
| gDNA | 21052 | 99.6 | 97.7 |
| gDNA | 21053 | 99.6 | 97.7 |
| gDNA | 21054 | 99.5 | 97.8 |
| gDNA | 21095 | 99.9 | 97.8 |
| gDNA | 21096 | 99.6 | 97.8 |
| gDNA | 21097 | 99.4 | 97.8 |


| gDNA | 21098 | 99.6 | 97.8 |
| :---: | :---: | :---: | :---: |
| gDNA | 21099 | 99.4 | 97.8 |
| gDNA | 21189 | 99.5 | 97.8 |
| gDNA | 21190 | 99.5 | 97.8 |
| gDNA | 21191 | 99.4 | 98 |
| gDNA | 21193 | 99.8 | 98 |
| gDNA | 21194 | 99.7 | 98 |
| gDNA | 21195 | 99.7 | 98 |
| gDNA | 21196 | 99.5 | 98 |
| gDNA | 21197 | 99.4 | 98.1 |
| gDNA | 21198 | 99.6 | 98.1 |
| gDNA | 21199 | 99.5 | 98.1 |
| gDNA | 21291 | 99.7 | 98.1 |
| gDNA | 21292 | 99.5 | 98.1 |
| gDNA | 21293 | 99.6 | 98.1 |
| gDNA | 21294 | 99.8 | 98.2 |
| gDNA | 21295 | 99.7 | 98.2 |
| gDNA | 21296 | 99.6 | 98.2 |
| gDNA | 21297 | 99.8 | 98.2 |
| gDNA | 21299 | 99.8 | 98.2 |
| gDNA | 21300 | 99.5 | 98.3 |
| gDNA | 21301 | 99.6 | 98.3 |
| gDNA | 21302 | 99.8 | 98.3 |
| gDNA | 21303 | 99.5 | 98.3 |
| gDNA | 21304 | 99.5 | 98.3 |
| gDNA | 21305 | 99.9 | 98.3 |
| gDNA | 21306 | 99.6 | 98.4 |
| gDNA | 21307 | 99.7 | 98.4 |
| gDNA | 21308 | 99.6 | 98.4 |
| gDNA | 21309 | 99.8 | 98.4 |
| gDNA | 23050 | 99.7 | 98.4 |
| gDNA | 23051 | 99.6 | 98.5 |
| gDNA | 23052 | 99.7 | 98.5 |
| gDNA | 23053 | 99.9 | 98.5 |
| gDNA | 23054 | 99.6 | 98.5 |
| gDNA | 23055 | 99.8 | 98.5 |
| gDNA | 23056 | 99.9 | 98.5 |
| gDNA | 23057 | 99.6 | 98.5 |
| gDNA | 23058 | 99.8 | 98.5 |
| gDNA | 23059 | 99.8 | 98.5 |
| gDNA | 23060 | 99.9 | 98.6 |
| gDNA | 23061 | 99.8 | 98.6 |
| gDNA | 23062 | 99.7 | 98.6 |
| gDNA | 23063 | 99.6 | 98.6 |
| gDNA | 23064 | 99.7 | 98.6 |
| gDNA | 23065 | 99.7 | 98.7 |
| gDNA | 23066 | 99.7 | 98.7 |
| gDNA | 23067 | 99.6 | 98.7 |
| gDNA | 23068 | 99.9 | 98.7 |
| gDNA | 23069 | 99.8 | 98.7 |
| gDNA | 23070 | 99.9 | 98.7 |
| gDNA | 23071 | 99.6 | 98.7 |
| gDNA | 23072 | 100 | 98.7 |
| gDNA | 23073 | 99.9 | 98.7 |
| gDNA | 23074 | 99.7 | 98.7 |


| gDNA | 23075 | 99.9 | 98.7 |
| :---: | :---: | :---: | :---: |
| gDNA | 23076 | 99.9 | 98.8 |
| gDNA | 23077 | 99.7 | 98.8 |
| gDNA | 23078 | 99.7 | 98.8 |
| gDNA | 23079 | 99.6 | 98.8 |
| gDNA | 23080 | 99.9 | 98.8 |
| gDNA | 23081 | 99.9 | 98.8 |
| gDNA | 23082 | 99.8 | 98.8 |
| gDNA | 23083 | 99.9 | 98.8 |
| gDNA | 23084 | 99.9 | 98.8 |
| gDNA | 23085 | 99.9 | 98.9 |
| gDNA | 23086 | 99.9 | 98.9 |
| gDNA | 23087 | 99.9 | 98.9 |
| gDNA | 23088 | 99.9 | 98.9 |
| gDNA | 23089 | 99.9 | 98.9 |
| gDNA | 23090 | 99.9 | 98.9 |
| gDNA | 23091 | 99.7 | 98.9 |
| gDNA | 23092 | 99.5 | 98.9 |
| gDNA | 23093 | 99.8 | 98.9 |
| gDNA | 23094 | 99.7 | 98.9 |
| gDNA | 23095 | 99.9 | 98.9 |
| gDNA | 23096 | 99.9 | 98.9 |
| gDNA | 23097 | 99.9 | 98.9 |
| gDNA | 23098 | 99.9 | 98.9 |
| gDNA | 23099 | 99 | 99 |
| gDNA | 23100 | 99.9 | 99 |
| gDNA | 23101 | 99.7 | 99 |
| gDNA | 23102 | 99.9 | 99 |
| gDNA | 23103 | 99.9 | 99 |
| gDNA | 23104 | 99.9 | 99 |
| gDNA | 23105 | 99.9 | 99 |
| gDNA | 23106 | 99.7 | 99.1 |
| gDNA | 23107 | 99.9 | 99.1 |
| gDNA | 23108 | 99.9 | 99.1 |
| gDNA | 23109 | 99.9 | 99.1 |
| gDNA | 23110 | 99.9 | 99.1 |
| gDNA | 23111 | 99.9 | 99.1 |
| gDNA | 23112 | 99.9 | 99.1 |
| gDNA | 23113 | 99.9 | 99.1 |
| gDNA | 23114 | 99.9 | 99.1 |
| gDNA | 23115 | 99.8 | 99.1 |
| gDNA | 23116 | 99.9 | 99.1 |
| gDNA | 23117 | 99.9 | 99.1 |
| gDNA | 23118 | 99.9 | 99.2 |
| gDNA | 23119 | 99.9 | 99.2 |
| gDNA | 23120 | 99 | 99.2 |
| gDNA | 23121 | 99.9 | 99.2 |
| gDNA | 23122 | 99.8 | 99.2 |
| gDNA | 23123 | 99.9 | 99.2 |
| gDNA | 23124 | 99.9 | 99.3 |
| gDNA | 23125 | 99.9 | 99.3 |
| gDNA | 23126 | 99.9 | 99.3 |
| gDNA | 23127 | 99.9 | 99.4 |
| gDNA | 23128 | 99.9 | 99.4 |
| gDNA | 23129 | 99.9 | 99.4 |


| gDNA | 23130 | 99.9 | 99.4 |
| :--- | :---: | :---: | :---: |
| gDNA | 23131 | 99.9 | 99.4 |
| gDNA | 23132 | 99.9 | 99.4 |
| gDNA | 23133 | 99.9 | 99.4 |
| gDNA | 23134 | 99.9 | 99.5 |
| gDNA | 23135 | 99.9 | 99.5 |
| gDNA | 23136 | 99.8 | 99.5 |
| gDNA | 23137 | 99.9 | 99.5 |
| gDNA | 23138 | 100 | 99.5 |
| gDNA | 23139 | 99.9 | 99.5 |
| gDNA | 23140 | 100 | 99.6 |
| gDNA | 23141 | 100 | 99.6 |
| gDNA | 23142 | 99.9 | 99.6 |
| gDNA | 23143 | 99.9 | 99.6 |
| gDNA | 23144 | 100 | 99.8 |
| gDNA | 23145 | 100 | 99.9 |
| gDNA | 23146 | 100 | 99.9 |

Table S3. Somatic non-synonymous mutations discovered in tumor samples

| ID Sample | Gene | RefSeq | \#CHROM | POS | REF | VAR | cDNA | AA | VAF in tumor samples |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 3392 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 34.4 (386/1119) |
| 3396 | TP53 | NM_000546.5 | chr17 | 7577610 | T | TC | c.673-2A>G | - | 5.13 (44/857) |
| 4822 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 36 (581/1614) |
| 4937 | SF3B1 | NM_012433.2 | chr2 | 198267371 | G | T | c.1986C>A | p.H662Q | 31.15 (3202/10278) |
| 4997 | TP53 | NM_000546.5 | chr17 | 7578394 | T | A | c.536A>T | p.H179L | 89.57 (2233/2491) |
| 5564 | ATM | NM_000051.3 | chr11 | 108206593 | G | A | c. $8173 \mathrm{G}>\mathrm{A}$ | p.D2725N | 18.78 (496/2641) |
| 5564 | ATM | NM_000051.3 | chr11 | 108224528 | C | T | c.8707C>T | p.P2903S | 23.06 (751/3257) |
| 5564 | POT1 | NM_015450.2 | chr7 | 124493080 | C | A | c.422G>T | p.G272V | 15.23 (545/3579) |
| 7302 | NOTCH1 | NM_017617.3 | chr9 | 139390759 | CG | C | c.7431_7431delC | p.A2478fs*5 | 43.4 (447/1030) |
| 10676 | BRAF | NM_004333.4 | chr7 | 140534588 | A | T | c. $325 \mathrm{~T}>\mathrm{A}$ | p.F109I | 40.4 (5204/12861) |
| 10676 | SF3B1 | NM_012433.2 | chr2 | 198266834 | T | C | c. 2098 A>G | p.K700E | 20.3 (2415/11876) |
| 12162 | MGA | NM_001164273.1 | chr15 | 41989130 | A | AT | c.1922_1923insT | p.T642fs*15 | 11.59 (361/3115) |
| 12162 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 41.64 (513/1232) |
| 12955 | FBXW7 | NM_033632 | chr4 | 153258983 | G | A | c.832C>T | p.R278* | 15.86 (550/3467) |
| 13443 | BIRC3 | NM_001165.4 | chr11 | 102207656 | AC | A | c.1639_1639delC | p.Q547fs*21 | 10.5 (55/523) |
| 14293 | TP53 | NM_000546.5 | chr17 | 7578553 | T | C | c.377A>G | p.Y126C | 13.65 (731/5353) |
| 14419 | ATM | NM_000051.3 | chr11 | 108170513 | AT | A | c.5079_5079delT | p.D1693fs*21 | 86.18 (2014/2332) |
| 14419 | MGA | NM_001164273.1 | chr15 | 42054328 | TAA | T | c.7513_7514delAA | p.K2504fs*31 | 36.5 (803/2199) |
| 14419 | NFKBIE | NM_004556.2 | chr6 | 44232738 | TGTAA | T | c.759_762delTTAC | p.Y254fs*13 | 21 (285/1355) |
| 14419 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 12.79 (169/1320) |
| 14462 | BIRC3 | NM_001165.4 | chr11 | 102201928 | TA | T | c.1281_1281delA | p.R428fs*19 | 31.17 (944/3028) |
| 14462 | BIRC3 | NM_001165.4 | chr11 | 102207519 | AG | A | c.1609_1609delG | p.E537fs*31 | 27 (678/2511) |
| 14462 | XPO1 | NM_003400.3 | chr2 | 61719472 | C | T | c.1711G>A | p.E571K | 36.9 (1803/4877) |
| 14687 | ATM | NM_000051.3 | chr11 | 108201101 | C | T | c.7468C>T | p.L2490F | 33.8 (5139/15183) |
| 15257 | SF3B1 | NM_012433.2 | chr2 | 198266611 | C | T | c. $2225 \mathrm{G}>\mathrm{A}$ | p.G742D | 30.16 (2557/8475) |
| 15257 | XPO1 | NM_003400.3 | chr2 | 61719472 | C | T | c.1711G>A | p.E571K | 30.4 (2882/9475) |
| 15280 | SF3B1 | NM_012433.2 | chr2 | 198266834 | T | C | c.2098A>G | p.K700E | 18.28 (647/3535) |
| 15361 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 38.84 (3004/7720) |
| 15367 | SF3B1 | NM_012433.2 | chr2 | 198266834 | T | C | c. $2098 \mathrm{~A}>\mathrm{G}$ | p.K700E | 10.13 (220/2171) |
| 15367 | TP53 | NM_000546.5 | chr17 | 7577539 | G | A | c.742C>T | p.R248W | 38.49 (1411/3662) |
| 15448 | POT1 | NM_015450.2 | chr7 | 124537225 | C | CA | c.2_3insT | initiating Methionine lost | 27.2 (61/224) |
| 15448 | TP53 | NM_000546.5 | chr17 | 7578190 | T | G | c.659A>C | p.Y220S | 23.5 (2002/8499) |
| 15505 | KRAS | NM_033360.2 | chr12 | 25398255 | G | T | c.64C>A | p.Q22K | 21.95 (1222/5559) |
| 15505 | SAMHD1 | NM_015474 | chr20 | 35526905 | CA | C | c.1545_1545delT | p.l515fs*35 | 43.16 (3089/5442) |


| 15505 | SAMHD1 | NM_015474 | chr20 | 35526278 | C | T | c.1693G>A | p.A565T | 50.02 (3232/6457) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 15505 | TP53 | NM_000546.5 | chr17 | 7577548 | C | T | c.733G>A | p.G245S | 19.95 (979/4903) |
| 15537 | BIRC3 | NM_001165.4 | chr11 | 102201966 | G | GA | c.1318_1319insA | p.S441fs*3 | 10.34 (156/1505) |
| 15537 | BIRC3 | NM_001165.4 | chr11 | 102201960 | G | T | c.1312G>T | p.E438* | 11.79 (186/1586) |
| 15537 | BIRC3 | NM_001165.4 | chr11 | 102207656 | AC | A | c.1639_1639delC | p.Q547fs*21 | 20.99 (390/1853) |
| 15597 | SF3B1 | NM_012433.2 | chr2 | 198267484 | G | A | c.1873C>T | p.R625C | 10.8 (677/6265) |
| 15597 | TP53 | NM_000546.5 | chr17 | 7577610 | T | C | c. $673-2 A>G$ | - | 22.44 (724/3227) |
| 15891 | TP53 | NM_000546.5 | chr17 | 7579372 | GC | G | c.314_314delG | p.G105fs*18 | 18.41 (648/3505) |
| 15891 | TP53 | NM_000546.5 | chr17 | 7577610 | T | A | c. $673-2 A>G$ | - | 33.55 (787/2208) |
| 16243 | ATM | NM_000051.3 | chr11 | 108160350 | C | T | c.4258C>T | p.L1420F | 31.7 (2184/6880) |
| 16243 | SF3B1 | NM_012433.2 | chr2 | 198267489 | T | C | c.1868A>G | p.Y623C | 16.9 (1066/6299) |
| 17698 | NOTCH1 | NM_017617.3 | chr9 | 139390800 | AG | A | c.7390_7390delC | p.L2464fs*13 | 44.2 (1955/4422) |
| 18723 | TP53 | NM_000546.5 | chr17 | 7578208 | T | C | c. $641 \mathrm{~A}>\mathrm{G}$ | p.H214R | 7.01 (502/7158) |
| 18736 | FBXW7 | NM_033632.3 | chr4 | 153253770 | C | T | c. $963 \mathrm{G}>\mathrm{A}$ | p.W321* | 23.23 (1103/4742) |
| 18736 | IRF4 | NM_002460.3 | chr6 | 394951 | T | G | c.347T>G | p.L116R | 14.58 (1202/8245) |
| 19184 | ZMYM3 | NM_201599 | chrX | 70472961 | CA | C | c.144_144delT | p.G49fs*64 | 18.86 (458/2423) |
| 19363 | SF3B1 | NM_012433.2 | chr2 | 198267359 | C | G | c.1998G>C | p.K666N | 26.67 (1767/6625) |
| 19363 | XPO1 | NM_003400.3 | chr2 | 61719186 | T | C | c.1871A>G | p.D624G | 28.31 (1129/3987) |
| 19405 | NOTCH1 | NM_017617.3 | chr9 | 139390585 | C | T | c.7606G>A | p.V25361 | 32.98 (6297/19088) |
| 19533 | XPO1 | NM_003400.3 | chr2 | 61719472 | C | T | c. $1711 \mathrm{G} \times \mathrm{A}$ | p.E571K | 44.82 (3115/6948) |
| 20010 | SF3B1 | NM_012433.2 | chr2 | 198266822 | T | A | c.2110A>T | p.1704F | 12.14 (753/6200) |
| 20010 | SF3B1 | NM_012433.2 | chr2 | 198267371 | G | T | c.1986C>A | p.H662Q | 15.4 (686/4454) |
| 20010 | TP53 | NM_000546.5 | chr17 | 7578413 | C | G | c.517G>C | p.V173L | 11.54 (261/2260) |
| 20196 | IKZF3 | NM_012481.4 | chr17 | 37985732 | G | A | c.71C>T | p.A24V | 51.19 (2296/4484) |
| 20802 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 37.13 (822/2209) |
| 20803 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 23.3 (916/3923) |
| 20804 | IKZF3 | NM_012481.4 | chr17 | 37947776 | A | C | c.485T>G | p.L162R | 27.22 (1316/4834) |
| 20804 | ZMYM3 | NM_201599 | chrX | 70472962 | AG | A | c.143_143delC | p.P48fs*65 | 24.43 (579/2369) |
| 20807 | MYD88 | NM_002468.4 | chr3 | 38182641 | T | C | c. $794 \mathrm{~T}>\mathrm{C}$ | p.L265P | 21.99 (1087/4940) |
| 20808 | SF3B1 | NM_012433.2 | chr2 | 198267360 | T | G | c.1997A>C | p.K666T | 22.11 (1782/8056) |
| 20810 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 34.54 (1636/4731) |
| 20811 | SF3B1 | NM_012433.2 | chr2 | 198266713 | C | T | c.2219G>A | p.G740E | 10.96 (424/3865) |
| 20812 | IKZF3 | NM_012481.4 | chr17 | 37947776 | A | C | c.485T>G | p.L162R | 42.39 (3787/8922) |
| 20812 | NOTCH1 | NM_017617.3 | chr9 | 139390975 | G | A | c. $7216 \mathrm{C}>\mathrm{T}$ | p.Q2406* | 41.51 (2956/7119) |
| 20812 | RPS15 | NM_001018.3 | chr19 | 1440439 | C | T | c. $413 \mathrm{C}>$ T | p.S138F | 14.08 (309/2187) |
| 20813 | MGA | NM_001164273.1 | chr15 | 42028793 | TA | T | c.4332_4332delA | p.L1444fs*36 | 32.37 (1714/5295) |
| 20818 | ASXL1 | NM_015338.5 | chr20 | 31024704 | G | A | c.4189G>A | p.G1397S | 48.02 (2053/4270) |
| 20820 | XPO1 | NM_003400.3 | chr2 | 61719472 | C | A | c.1711G>T | p.E571* | 10.68 (240/2247) |
| 20821 | BRAF | NM_004333.4 | chr7 | 140481402 | C | G | c.1406G>C | p.G469A | 35.17 (1666/4732) |


| 20821 | EGR2 | NM_000399.3 | chr10 | 64573167 | C | G | c.1231G>C | p.D411H | 34.43 (2156/6253) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 20822 | ATM | NM_000051.3 | chr11 | 108126954 | GA | G | c.2138_2138delA | p.T713fs*20 | 10.87 (332/3055) |
| 20822 | BIRC3 | NM_001165.4 | chr11 | 102201945 | GAA | G | c.1298_1299delAA | p.R434fs*3 | 10.88 (420/3857) |
| 20822 | IRF4 | NM_002460.3 | chr6 | 394951 | T | G | c. $347 \mathrm{~T}>\mathrm{G}$ | p.L116R | 29.37 (1664/5661) |
| 20823 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 25.53 (1971/5742) |
| 20827 | IKZF3 | NM_012481.4 | chr17 | 37947776 | A | C | c.485T>G | p.L162R | 41.37 (1814/4377) |
| 20831 | ATM | NM_000051.3 | chr11 | 108178641 | CG | C | c.5693_5693delG | p.R1898fs*19 | 11.94 (151/1265) |
| 20832 | ATM | NM_000051.3 | chr11 | 108175463 | A | T | c. $5558 \mathrm{~A}>\mathrm{T}$ | p.D1853V | 50.28 (2634/5237) |
| 20832 | XPO1 | NM_003400.3 | chr2 | 61719472 | C | T | c.1711G>A | p.E571K | 44.23 (2407/5441) |
| 20833 | BIRC3 | NM_001165.4 | chr11 | 102207704 | CA | C | c.1687_1687delA | p.E564fs*4 | 17.15 (996/5794) |
| 20833 | ZMYM3 | NM_201599 | chrX | 70472444 | A | T | c.662T>A | p.L221* | 85.83 (1950/2268) |
| 20834 | KRAS | NM_033360.2 | chr12 | 25398262 | C | G | c.57G>C | p.L19F | 28.85 (1123/3893) |
| 20844 | TP53 | NM_000546.5 | chr17 | 7579419 | AG | A | c.267_267delC | p.S90fs*33 | 3.16 (86/2723) |
| 20845 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 10.23 (424/4132) |
| 20845 | RPS15 | NM_001018.3 | chr19 | 1440436 | C | T | c.413C>T | p.S138F | 22.71 (422/1857) |
| 20845 | XPO1 | NM_003400.3 | chr2 | 61719186 | T | C | c.1871A>G | p.D624G | 38.73 (2087/5385) |
| 20846 | TP53 | NM_000546.5 | chr17 | 7578176 | C | T | c. $672+1 \mathrm{G}>\mathrm{A}$ | - | 28.1 (726/2584) |
| 20847 | BRAF | NM_004333.4 | chr7 | 140453136 | A | T | c.1799T>A | p.V600E | 17.69 (820/4633) |
| 20849 | SF3B1 | NM_012433.2 | chr2 | 198267359 | C | A | c.1998G>T | p.K666N | 20.2 (1033/5110) |
| 20850 | SF3B1 | NM_012433.2 | chr2 | 198266834 | T | C | c.2098A>G | p.K700E | 12.16 (460/3779) |
| 20852 | ATM | NM_000051.3 | chr11 | 108143456 | C | G | c.3161C>G | p.P1054R | 48.39 (4558/9414) |
| 20852 | NFKBIE | NM_004556.2 | chr6 | 44232738 | TGTAA | T | c.759_762delTTAC | p.Y254fs*13 | 81.93 (3260/3973) |
| 20852 | SF3B1 | NM_012433.2 | chr2 | 198266834 | T | C | c.2098A>G | p.K700E | 13.94 (718/5150) |
| 20853 | ATM | NM_000051.3 | chr11 | 108143456 | C | G | c.3161C>G | p.P1054R | 51.51 (2525/4894) |
| 20853 | BIRC3 | NM_001165.4 | chr11 | 102207684 | AC | A | c.1667_1667delC | p.T556fs*12 | 49.27 (2360/4789) |
| 20853 | SF3B1 | NM_012433.2 | chr2 | 198266611 | C | T | c. $2225 \mathrm{G}>\mathrm{A}$ | p.G742D | 43.96 (1797/4083) |
| 20855 | NFKBIE | NM_004556.2 | chr6 | 44232738 | TGTAA | T | c.759_762delTTAC | p.Y254fs*13 | 13.6 (507/3712) |
| 20857 | EGR2 | NM_000399.3 | chr10 | 64573248 | G | T | c.1150C>A | p.H384N | 27.07 (1313/4848) |
| 20857 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 41.62 (1463/3505) |
| 20860 | MGA | NM_001164273.1 | chr15 | 42028430 | AT | A | c.3969_3969delT | p.Y1323* | 22.38 (1126/5032) |
| 20860 | TP53 | NM_000546.5 | chr17 | 7577123 | A | C | c. $815 \mathrm{~T}>\mathrm{G}$ | p.V272G | 5.11 (248/4847) |
| 20861 | EGR2 | NM_000399.3 | chr10 | 64573248 | G | T | c.1150C>A | p.H384N | 10.1 (720/7121) |
| 20861 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 23.64 (1042/4396) |
| 20868 | SF3B1 | NM_012433.2 | chr2 | 198267360 | T | G | c.1997A>C | p.K666T | 19.46 (1184/6077) |
| 20868 | SF3B1 | NM_012433.2 | chr2 | 198267361 | T | C | c.1996A>G | p.K666E | 20.3 (1249/6152) |
| 20870 | ATM | NM_000051.3 | chr11 | 108160480 | T | G | c.4388T>G | p.F1463C | 49.82 (3142/6305) |
| 20872 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 14.06 (532/3781) |
| 20872 | TP53 | NM_000546.5 | chr17 | 7577108 | C | A | c.830G>T | p.C277F | 5.21 (344/6605) |
| 20875 | XPO1 | NM_003400.3 | chr2 | 61719471 | T | C | c.1712A>G | p.E571G | 20.02 (1086/5420) |


| 20876 | SF3B1 | NM_012433.2 | chr2 | 198267361 | T | C | c.1996A>G | p.K666E | 41.44 (2390/5766) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 20897 | BIRC3 | NM_001165.4 | chr11 | 102207656 | AC | A | c.1639_1639delC | p.Q547fs*21 | 75.16 (838/1115) |
| 21007 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 30.48 (1704/5585) |
| 21007 | TP53 | NM_000546.5 | chr17 | 7578460 | A | T | c.470T>A | p.V157D | 6.84 (511/7466) |
| 21008 | SF3B1 | NM_012433.2 | chr2 | 198266709 | C | A | c. $2223 \mathrm{G}>\mathrm{T}$ | p.K741N | 10.72 (452/4217) |
| 21010 | IRF4 | NM_002460.3 | chr6 | 394951 | T | G | c. $347 \mathrm{~T}>\mathrm{G}$ | p.L116R | 54.95 (2315/4210) |
| 21010 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 33.41 (449/1340) |
| 21011 | BRAF | NM_004333.4 | chr7 | 140481402 | C | G | c.1406G>C | p.G469A | 41.92 (1971/4693) |
| 21012 | FBXW7 | NM_033632.3 | chr4 | 153247289 | G | A | c.1513C>T | p.R505C | 42.2 (1834/4342) |
| 21012 | NOTCH1 | NM_017617.3 | chr9 | 139391200 | C | CCACA | c.6990_6991insTGTG | p.A2331fs*24 | 30.69 (421/1353) |
| 21012 | TP53 | NM_000546.5 | chr17 | 7578535 | T | A | c.395A>T | p.K132M | 7.85 (244/3109) |
| 21013 | SF3B1 | NM_012433.2 | chr2 | 198266611 | C | T | c. $2225 \mathrm{G}>\mathrm{A}$ | p.G742D | 13.18 (653/4951) |
| 21014 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 20.25 (873/4302) |
| 21015 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 46.48 (1517/3261) |
| 21015 | SF3B1 | NM_012433.2 | chr2 | 198266821 | A | T | c. $2111 \mathrm{~T}>\mathrm{A}$ | p.I704N | 47.02 (2466/5244) |
| 21015 | TP53 | NM_000546.5 | chr17 | 7577117 | A | T | c. $821 \mathrm{~T}>\mathrm{A}$ | p.V274D | 16.17 (876/5410) |
| 21015 | XPO1 | NM_003400.3 | chr2 | 61719472 | C | T | c.1711G>A | p.E571K | 13.72 (811/5912) |
| 21016 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 14.06 (532/3781) |
| 21017 | ATM | NM_000051.3 | chr11 | 108143456 | C | G | c.3161C>G | p.P1054R | 78.34 (3292/4200) |
| 21017 | IKZF3 | NM_012481.4 | chr17 | 37985732 | G | A | c.71C>T | p.A24V | 47.79 (3051/6380) |
| 21018 | TP53 | NM_000546.5 | chr17 | 7578400 | G | A | c. 530 C > $T$ | p.P177L | 5.38 (288/5353) |
| 21018 | XPO1 | NM_003400.3 | chr2 | 61719472 | C | T | c.1711G>A | p.E571K | 22.72 (2210/9718) |
| 21019 | FBXW7 | NM_033632.3 | chr4 | 153245446 | G | A | c.1745C>T | p.S582L | 17.19 (1314/7640) |
| 21019 | TP53 | NM_000546.5 | chr17 | 7577548 | C | A | c.733G>T | p.G245C | 3.19 (199/6226) |
| 21019 | TP53 | NM_000546.5 | chr17 | 7577538 | C | T | c. $743 \mathrm{G}>\mathrm{A}$ | p.R248Q | 4.28 (273/6383) |
| 21019 | TP53 | NM_000546.5 | chr17 | 7577106 | G | A | c.832C>T | p.P278S | 5.56 (440/7903) |
| 21019 | TP53 | NM_000546.5 | chr17 | 7577120 | C | T | c. $818 \mathrm{G}>\mathrm{A}$ | p.R273H | 15.34 (1286/8382) |
| 21021 | TP53 | NM_000546.5 | chr17 | 7577121 | G | A | c. $817 \mathrm{C}>$ T | p.R273C | 26.49 (656/2473) |
| 21022 | NOTCH1 | NM_017617.3 | chr9 | 139390861 | G | A | c.7330C>T | p.Q2444* | 18.12 (1219/6726) |
| 21045 | SF3B1 | NM_012433.2 | chr2 | 198266834 | T | C | c. $2098 \mathrm{~A}>\mathrm{G}$ | p.K700E | 11.07 (350/3163) |
| 21046 | XPO1 | NM_003400.3 | chr2 | 61719471 | T | A | c. $1712 \mathrm{~A}>\mathrm{T}$ | p.E571V | 14.34 (663/4625) |
| 21047 | MGA | NM_001164273.1 | chr15 | 42042450 | TC | T | c.6646_6646delC | p.Q2216fs*22 | 29.7 (1549/5215) |
| 21047 | SF3B1 | NM_012433.2 | chr2 | 198267361 | T | C | c.1996A>G | p.K666E | 40.33 (1914/4745) |
| 21048 | MAP2K1 | NM_002755 | chr15 | 66727453 | A | G | c.169A>G | p.K57E | 13.53 (710/5245) |
| 21048 | RPS15 | NM_001018.3 | chr19 | 1440423 | G | A | c.400G>A | p.G134R | 20.62 (632/3064) |
| 21050 | ATM | NM_000051.3 | chr11 | 108143456 | C | G | c.3161C>G | p.P1054R | 46.61 (2718/5829) |
| 21050 | NOTCH1 | NM_017617.3 | chr9 | 139390765 | CG | C | c.7425_7425delC | p.V2475* | 19.53 (739/3781) |
| 21050 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 20.79 (794/3815) |
| 21051 | SF3B1 | NM_012433.2 | chr2 | 198267491 | C | A | c.1866G>T | p.E622D | 14.5 (744/5129) |


| 21095 | SF3B1 | NM_012433.2 | chr2 | 198266834 | T | C | c. 2098 A>G | p.K700E | 10.71 (596/5558) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 21095 | TP53 | NM_000546.5 | chr17 | 7578375 | GCTAT | G | c.551_554delATAG | p.D184fs*62 | 36.15 (2539/7008) |
| 21098 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 77.44 (3453/4448) |
| 21099 | TP53 | NM_000546.5 | chr17 | 7578461 | C | A | c.469G>T | p.V157F | 17.9 (1408/7866) |
| 21189 | RPS15 | NM_001018.3 | chr19 | 1440414 | C | T | c.391C>T | p.P131S | 30.92 (983/3178) |
| 21190 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 17.07 (320/1873) |
| 21190 | ZMYM3 | NM_201599 | chrX | 70461153 | TA | T | c.3807_3807delT | p.l1270fs*2 | 12.8 (210/1563) |
| 21191 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 22.05 (484/2190) |
| 21193 | RPS15 | NM_001018.3 | chr19 | 1440436 | C | T | c.413C>T | p.S138F | 21.18 (270/1275) |
| 21193 | TP53 | NM_000546.5 | chr17 | 7577570 | C | G | c.711G>C | p.M2371 | 23.01 (421/1829) |
| 21194 | TP53 | NM_000546.5 | chr17 | 7578460 | A | T | c.470T>A | p.V157D | 33.24 (995/2990) |
| 21196 | EGR2 | NM_000399.3 | chr10 | 64573332 | C | T | c.1066G>A | p.E356K | 46.87 (2378/5070) |
| 21196 | NFKBIE | NM_004556.2 | chr6 | 44232738 | TGTAA | T | c.759_762delTTAC | p.Y254fs*13 | 29.99 (702/2340) |
| 21296 | NFKBIE | NM_004556.2 | chr6 | 44232738 | TGTAA | T | c.759_762delTTAC | p.Y254fs*13 | 23.85 (899/3770) |
| 21300 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 13.51 (302/2235) |
| 21300 | ZMYM3 | NM_201599 | chrX | 70471427 | CT | C | c.691_691delA | p.S231fs*26 | 55.71 (1093/1962) |
| 21301 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 28.54 (764/2673) |
| 21301 | SAMHD1 | NM_015474 | chr20 | 35526278 | C | T | c.1693G>A | p.A565T | 52.67 (2193/4164) |
| 21302 | ATM | NM_000051.3 | chr11 | 108200961 | G | A | c.7328G>A | p.R2443Q | 13.56 (480/3539) |
| 21304 | IKZF3 | NM_012481.4 | chr17 | 37947776 | A | C | c. $485 \mathrm{~T}>\mathrm{G}$ | p.L162R | 14.63 (936/6389) |
| 21304 | MGA | NM_001164273.1 | chr15 | 42021500 | CA | C | c.3797_3797delA | p.Q1266fs*25 | 24.72 (1448/5854) |
| 21304 | MGA | NM_001164273.1 | chr15 | 41988277 | A | AT | c.1069_1070insT | p.A358fs*5 | 26.82 (1201/4466) |
| 21307 | RPS15 | NM_001018.3 | chr19 | 1440421 | T | G | c. $398 \mathrm{~T}>\mathrm{G}$ | p.I133S | 39.94 (1203/3010) |
| 21308 | SF3B1 | NM_012433.2 | chr2 | 198266834 | T | C | c.2098A>G | p.K700E | 13.08 (289/2206) |
| 23050 | BIRC3 | NM_001165.4 | chr11 | 102207656 | AC | A | c.1639_1639delC | p.Q547fs*21 | 28.5 (2711/9510) |
| 23050 | TP53 | NM_000546.5 | chr17 | 7578542 | G | C | c. $388 \mathrm{C}>\mathrm{G}$ | p.L130V | 41.1 (1803/4386) |
| 23054 | ATM | NM_000051.3 | chr11 | 108155201 | G | A | c. $3993+1 \mathrm{G}>\mathrm{A}$ | - | 30 (1106/3695) |
| 23054 | ATM | NM_000051.3 | chr11 | 108143456 | C | G | c.3161C>G | p.P1054R | 52.2 (2399/4595) |
| 23054 | RPS15 | NM_001018.3 | chr19 | 1440432 | C | T | c.409C>T | p.H137Y | 35.4 (955/2699) |
| 23057 | ATM | NM_000051.3 | chr11 | 108142006 | C | T | c.2950C>T | p.Q984* | 94.5 (1768/1870) |
| 23057 | XPO1 | NM_003400.3 | chr2 | 61719472 | C | T | c.1711G>A | p.E571K | 53.4 (1504/2814) |
| 23057 | ZMYM3 | NM_201599 | chrX | 70469355 | G | A | c. $1426 \mathrm{C}>\mathrm{T}$ | p.Q476* | 82.8 (3236/3906) |
| 23058 | NFKBIE | NM_004556.2 | chr6 | 44232738 | TGTAA | T | c.759_762delTTAC | p.Y254fs*13 | 16.57 (643/3879) |
| 23058 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 40.75 (2095/5133) |
| 23063 | ATM | NM_000051.3 | chr11 | 108160350 | C | T | c.4258C>T | p.L1420F | 47.8 (3023/6316) |
| 23063 | SF3B1 | NM_012433.2 | chr2 | 198266834 | T | C | c. $2098 \mathrm{~A}>\mathrm{G}$ | p.K700E | 12.2 (786/6441) |
| 23065 | MYD88 | NM_002468.4 | chr3 | 38182641 | T | C | c.794T>C | p.L265P | 17.3 (184/1063) |
| 23068 | NFKBIE | NM_004556.2 | chr6 | 44232738 | TGTAA | T | c.759_762delTTAC | p.Y254fs*13 | 49.6 (3380/6806) |
| 23070 | BIRC3 | NM_001165.4 | chr11 | 102207672 | C | T | c.1654C>T | p.Q552* | 17.82 (784/4399) |


| 23070 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 27.71 (908/3276) |
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| 23074 | SF3B1 | NM_012433.2 | chr2 | 198266611 | C | T | c. $2225 \mathrm{G}>\mathrm{A}$ | p.G742D | 46.2 (3284/7103) |
| 23075 | ATM | NM_000051.3 | chr11 | 108114679 | G | A | c.497-1G>A | - | 39.35 (1672/4249) |
| 23075 | ATM | NM_000051.3 | chr11 | 108172375 | T | A | c.5178T>A | p.C1726* | 39.99 (2785/6963) |
| 23075 | ATM | NM_000051.3 | chr11 | 108160350 | C | T | c.4258C>T | p.L1420F | 49.89 (3499/7013) |
| 23075 | RPS15 | NM_001018.3 | chr19 | 1440432 | C | T | c.409C>T | p.H137Y | 16.41 (462/2814) |
| 23076 | XPO1 | NM_003400.3 | chr2 | 61719303 | C | A | c.1754G>T | p.C585F | 48.6 (3396/6977) |
| 23077 | ATM | NM_000051.3 | chr11 | 108119720 | G | GA | c.1126_1127insA | p.S377fs*2 | 27.7 (1263/4552) |
| 23077 | SF3B1 | NM_012433.2 | chr2 | 198266834 | T | C | c.2098A>G | p.K700E | 49 (3488/7113) |
| 23078 | EGR2 | NM_000399.3 | chr10 | 64573248 | G | T | c.1150C>A | p.H384N | 48.7 (3886/7965) |
| 23079 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 10.8 (530/4866) |
| 23079 | NRAS | NM_002524.4 | chr1 | 115258744 | C | T | c.38G>A | p.G13D | 49.2 (1884/3822) |
| 23080 | SF3B1 | NM_012433.2 | chr2 | 198266494 | T | C | c. $2342 \mathrm{~A}>\mathrm{G}$ | p.D781G | 18.87 (783/4148) |
| 23082 | NFKBIE | NM_004556.2 | chr6 | 44232738 | TGTAA | T | c.759_762delTTAC | p.Y254fs*13 | 29.36 (1477/5030) |
| 23082 | SAMHD1 | NM_015474 | chr20 | 35555633 | CAT | C | c.646_647delAT | p.M216fs*2 | 39.39 (1530/3884) |
| 23082 | SF3B1 | NM_012433.2 | chr2 | 198266834 | T | C | c.2098A>G | p.K700E | 35.09 (2582/7357) |
| 23085 | SF3B1 | NM_012433.2 | chr2 | 198266834 | T | C | c.2098A>G | p.K700E | 26.24 (415/1581) |
| 23086 | FBXW7 | NM_033632.3 | chr4 | 153247366 | C | T | c.1436G>A | p.R479Q | 44.08 (1792/4067) |
| 23086 | MGA | NM_001164273.1 | chr15 | 42042600 | AG | A | c.6796_6796delG | p.N2267fs*68 | 72.5 (1002/1382) |
| 23088 | SF3B1 | NM_012433.2 | chr2 | 198266834 | T | C | c. 2098 A>G | p.K700E | 18.17 (714/3929) |
| 23089 | NOTCH1 | NM_017617.3 | chr9 | 139390145 | T | C | c. ${ }^{*} 378 \mathrm{~A}>\mathrm{G}$ | - | 10 (507/5041) |
| 23089 | SF3B1 | NM_012433.2 | chr2 | 198266834 | T | C | c. $2098 \mathrm{~A}>\mathrm{G}$ | p.K700E | 29.7 (1230/4141) |
| 23091 | TP53 | NM_000546.5 | chr17 | 7577538 | C | T | c.743G>A | p.R248Q | 44.2 (2624/5938) |
| 23093 | MYD88 | NM_002468.4 | chr3 | 38182641 | T | C | c.794T>C | p.L265P | 50.22 (4631/9220) |
| 23094 | ASXL1 | NM_015338.5 | chr20 | 31023328 | CT | C | c.2814_2814delT | p.A939fs*6 | 10.3 (750/7309) |
| 23094 | ASXL1 | NM_015338.5 | chr20 | 31022628 | G | T | c. $2113 \mathrm{G}>\mathrm{T}$ | p.E705* | 32.6 (2222/6818) |
| 23094 | XPO1 | NM_003400.3 | chr2 | 61719472 | C | T | c.1711G>A | p.E571K | 11.6 (794/6847) |
| 23096 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 15.47 (442/2857) |
| 23097 | KRAS | NM_033360.2 | chr12 | 25378562 | C | T | c.436G>A | p.A146T | 14.8 (970/6527) |
| 23106 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 17.4 (611/3508) |
| 23106 | POT1 | NM_015450.2 | chr7 | 124499045 | T | C | c. $668 \mathrm{~A}>\mathrm{G}$ | p.Y223C | 38.3 (1714/4469) |
| 23106 | RPS15 | NM_001018.3 | chr19 | 1440417 | G | A | c. $394 \mathrm{G}>\mathrm{A}$ | p.G132S | 39.1 (1066/2723) |
| 23107 | NFKBIE | NM_004556.2 | chr6 | 44232738 | TGTAA | T | c.759_762delTTAC | p.Y254fs*13 | 14.2 (746/5269) |
| 23108 | SF3B1 | NM_012433.2 | chr2 | 198265476 | T | C | c. $2681 \mathrm{~A}>\mathrm{G}$ | p.D894G | 23.7 (1147/4840) |
| 23112 | SF3B1 | NM_012433.2 | chr2 | 198266834 | T | C | c. $2098 \mathrm{~A}>\mathrm{G}$ | p.K700E | 38.5 (2460/6375) |
| 23113 | TP53 | NM_000546.5 | chr17 | 7577551 | C | T | c.730G>A | p.G244S | 55.6 (1837/3302) |
| 23115 | EGR2 | NM_000399.3 | chr10 | 64573332 | C | T | c.1066G>A | p.E356K | 16.35 (1084/6629) |
| 23118 | NOTCH1 | NM_017617.3 | chr9 | 139390793 | CGTGGGCA | C | c.7391_7397delTGCCCAC | p.L2464fs*11 | 29.7 (603/2028) |
| 23118 | POT1 | NM_015450.2 | chr7 | 124537227 | T | A | c.1A>T | p.M1L | 23.9 (579/2414) |


| 23118 | ZMYM3 | NM_201599 | chrX | 70469448 | C | A | c.1333G>T | p.G445* | 15 (332/2200) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 23118 | ZMYM3 | NM_201599 | chrX | 70469002 | G | T | c.1488C>A | p.Y496* | 49.6 (1038/2089) |
| 23120 | NFKBIE | NM_004556.2 | chr6 | 44232738 | TGTAA | T | c.759_762delTTAC | p.Y254fs*13 | 25.22 (915/3627) |
| 23121 | NFKBIE | NM_004556.2 | chr6 | 44232738 | TGTAA | T | c.759_762delTTAC | p.Y254fs*13 | 20.3 (645/3175) |
| 23124 | IKZF3 | NM_183229.2 | chr17 | 37947776 | A | C | c. $485 \mathrm{~T}>\mathrm{G}$ | p.L162R | 14.2 (764/5348) |
| 23127 | ATM | NM_000051.3 | chr11 | 108143456 | C | G | c.3161C>G | p.P1054R | 49.66 (5187/10444) |
| 23128 | NFKBIE | NM_004556.2 | chr6 | 44232738 | TGTAA | T | c.759_762delTTAC | p.Y254fs*13 | 19.5 (786/4029) |
| 23129 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 15.34 (393/2561) |
| 23129 | SF3B1 | NM_012433.2 | chr2 | 198266711 | T | C | c. $2221 \mathrm{~A}>\mathrm{G}$ | p.K741E | 45.5 (3228/7081) |
| 23129 | XPO1 | NM_003400.3 | chr2 | 61719472 | C | T | c.1711G>A | p.E571K | 45.4 (4136/9097) |
| 23131 | SF3B1 | NM_012433.2 | chr2 | 198267491 | C | A | c.1866G>T | p.E622D | 39 (3585/9180) |
| 23132 | MGA | NM_001164273.1 | chr15 | 41961741 | C | T | c.649C>T | p.Q217* | 29.8 (2153/7211) |
| 23132 | XPO1 | NM_003400.3 | chr2 | 61719472 | C | T | c.1711G>A | p.E571K | 38 (2909/7641) |
| 23133 | IKZF3 | NM_183229.2 | chr17 | 37947776 | A | C | c.485T>G | p.L162R | 21.04 (1759/8358) |
| 23133 | NOTCH1 | NM_017617.3 | chr9 | 139390648 | CAG | C | c.7541_7542delCT | p.P2514fs*4 | 11.85 (574/4841) |
| 23133 | ZMYM3 | NM_201599 | chrX | 70469358 | G | A | c.1423C>T | p.Q475* | 21.18 (805/3800) |
| 23135 | MYD88 | NM_002468.4 | chr3 | 38182259 | T | C | c.695T>C | p.M232T | 29.1 (1890/6486) |
| 23137 | SF3B1 | NM_012433.2 | chr2 | 198266611 | C | T | c. $2225 \mathrm{G}>\mathrm{A}$ | p.G742D | 31.9 (2402/7525) |
| 23140 | SF3B1 | NM_012433.2 | chr2 | 198267369 | G | A | c.1988C>T | p.T663I | 39.18 (2933/7485) |
| 23141 | IKZF3 | NM_183229.2 | chr17 | 37947776 | A | C | c. $485 \mathrm{~T}>\mathrm{G}$ | p.L162R | 44.6 (2695/6031) |
| 23143 | SF3B1 | NM_012433.2 | chr2 | 198266834 | T | C | c. $2098 \mathrm{~A}>\mathrm{G}$ | p.K700E | 23.88 (1942/8129) |
| 23145 | ATM | NM_000051.3 | chr11 | 108155085 | A | G | c. $3878 \mathrm{~A}>\mathrm{G}$ | p.N1293S | 36.6 (2670/7285) |
| 23145 | IKZF3 | NM_183229.2 | chr17 | 37947776 | A | C | c.485T>G | p.L162R | 38.14 (2282/5983) |

Table S4. 11q deletion, 17p deletion and TP53 mutational status of tumor cell lines and primary CLL cells

| Sample ID | 11q deleted | 17p deleted | TP53 mutated |
| :--- | :---: | :---: | :---: |
| MEC1 | no | yes | no |
| SSK41 | no | no | no |
| VL51 | no | yes | no |
| JEKO1 | no | yes | yes |
| MAVER1 | yes | yes | yes |
| Z138 | no | no | no |
| 9321 | no | no | no |
| 14462 | yes | no | no |
| 12603 | no | no | no |
| 11731 | no | no | no |
| 12600 |  |  | no |

Table S5. Primary cell lines divided into BIRC3/TP53 mutated or wild-type (WT).

| Sample ID | IGHV | TP53 | BIRC3 |
| :--- | :---: | :---: | :---: |
| 13443 | Unmutated | WT | Mutated |
| 9696 | Unmutated | WT | Mutated |
| 9482 | Unmutated | WT | Mutated |
| 14324 | Unmutated | WT | Mutated |
| PMN019 | Unmutated | WT | Mutated |
| PMN012 | Unmutated | WT | Mutated |
| 11480 | Unmutated | Mutated | WT |
| 13344 | Unmutated | Mutated | WT |
| 7916 | Unmutated | Mutated | WT |
| 13730 | Unmutated | Mutated | WT |
| 9311 | Mutated | Mutated | WT |
| PMN081 | Unmutated | Mutated | WT |
| 14326 | Unmutated | Mutated | WT |
| 11214 | Unmutated | WT | WT |
| 10666 | Unmutated | WT | WT |
| 11815 | Unmutated | WT | WT |
| 19361 | Unmutated | Unmutated | WT |
| 10872 | Unmutated | WT | WT |
| 10650 |  | WT | WT |
| PMN059 |  |  | WT |
| 10320 |  |  |  |

