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KMT2D mutations and *TP53* disruptions are poor prognostic biomarkers in mantle cell lymphoma receiving high-dose therapy: a FIL study

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Authors contribution

DR, SC, FB, ML and GG designed and performed research; SF, DR, ADR, ViSt, PG, MGdS, AS, AM, AF, AP, SC, ML and GG enrolled patients; SF, DB, DN, MP, ML and AZ provided biological samples SF, DR, AR, AB, VaSp, IK, FD, CF and DB performed research; SF, DR, AR, AB, VaSp, AE, RM and IK analysed data; CWE, CD, AK, MJ and KG provided data for the validation; SF, DR, AE, RM, FB, ML and GG wrote the paper.

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ABSTRACT

In recent years, the outcome of mantle cell lymphoma has improved, especially in younger receiving cytarabine-containing chemoimmunotherapy and autologous patients, stem cell transplantation. Nevertheless, a proportion of mantle cell lymphoma patients still experience early failure. To identify biomarkers anticipating failure of intensive chemotherapy in mantle cell lymphoma, we performed target resequencing and DNA profiling of purified tumor samples collected from patients enrolled in the prospective FIL-MCL0208 phase III trial (high-dose chemoimmunotherapy followed by autologous transplantation and randomized lenalidomide maintenance). Mutations of KMT2D and disruption of TP53 by deletion or mutation associated with an increased risk of progression and death, both in univariate and multivariate analysis. By adding KMT2D mutations and TP53 disruption to the MIPI-c backbone, we derived a new prognostic index, the "MIPI-genetic". The "MIPI-g" improved the model discrimination ability compared to the MIPI-c alone, defining three risk groups: i) low-risk patients (4-years progression free survival and overall survival of 72.0% and 94.5%); ii) intermediaterisk patients (4-years progression free survival and overall survival of 42.2% and 65.8%) and iii) highrisk patients (4-years progression free survival and overall survival of 11.5% and 44.9%). Our results: i) confirm that TP53 disruption identifies a high-risk population characterized by poor sensitivity to conventional or intensified chemotherapy; *ii*) provide the pivotal evidence that patients harboring *KMT2D* mutations share the same poor outcome as patients harboring *TP53* disruption; and *iii*) allow to develop a tool for the identification of high-risk mantle cell lymphoma patients for whom novel therapeutic strategies need to be investigated. (Trial registered at clinicaltrials.gov identifier: NCT02354313)

INTRODUCTION

The introduction of high dose cytarabine-containing chemoimmunotherapeutic regimens and autologous transplantation (ASCT) have considerably improved the outcome of young fit mantle cell lymphoma (MCL) patients. Nonetheless, approximately 20 to 25% of MCL patients demonstrate inadequate efficacy of intensified chemoimmunotherapy as they are either primary refractory or relapse within two years from autologous stem cell transplantation.¹⁻⁵

Clinical and pathological scores, including the MCL international prognostic index (MIPI),⁶ the Ki-67 proliferative index,⁷ and their combination in the MIPI-c score, stratify MCL patients in groups at different risk of relapse.⁸ However, none of these tools has sufficient positive predictive value to trigger the development of tailored schedules specifically designed for high risk patients.⁹

Several recurrent mutations have been described in MCL, affecting DNA repair genes and cell cycle regulators (*TP53*, *ATM*, *CCND1*), epigenetic regulation genes (*KMT2D*, *WHSC1*) and cell-signaling pathways genes (*NOTCH1-2*, *BIRC3*, *TRAF2*).¹⁰⁻¹² The proof of principle that MCL genetics can impact on disease outcome stems from studies that have focused on the *TP53* tumor suppressor gene, including both mutations and 17p deletions.¹³⁻¹⁷

We prospectively assessed the clinical impact of a panel of genomic alterations in a cohort of young MCL patients treated with high dose chemoimmunotherapy and ASCT from the Fondazione Italiana Linfomi (FIL) "MCL0208" phase III trial.¹⁸ The results document that *KMT2D* mutations associate with poor outcome in MCL and, along with *TP53* mutations and 17p deletions, might be integrated in a new prognostic score to segregate a subgroup of patients who obtain minimal or no benefit from intensive chemoimmunotherapy. The prognostic score was validated in an independent series of cases.

METHODS

Patients series

The FIL-MCL0208 (NCT02354313) is a phase III, multicenter, open-label, randomized, controlled study, designed to determine the efficacy of lenalidomide as maintenance versus observation in young (18-65 years old), fit, advanced stage (Ann arbor II-IV) MCL patients after first line intensified and high-dose chemo-immunotherapy followed by ASCT. Cases of non-nodal MCL were excluded.¹⁹ The clinical trial, as well as the ancillary mutational study, were approved by the Ethical Committees of all the enrolling Centers. All patients provided written informed consent for the use of their biological samples for research purposes, in accordance with Institutional Review Boards requirements and the Helsinki's declaration. Clinical results of the fist interim analysis of the trial were already presented.¹⁸ Further information are supplied in the Supplementary appendix.

Biological samples

Tumor cells were sorted from the baseline BM samples by immunomagnetic beads (CD19 MicroBeads,human-Miltenyi Biotec GmbH, Bergisch Gladbach, Germany) and stocked as dry pellets. Tumor DNA was extracted according to DNAzol protocol (Life Technologies). Germline DNA was obtained from PB mononuclear cells collected under treatment and proven to be tumor free by MRD analysis. Further information are supplied in the Supplementary appendix.

Next generation sequencing

A targeted resequencing panel (target region: 37'821 bp) (Table S1) including the coding exons and splice sites of 7 genes (*ATM*, *TP53*, *CCND1*, *WHSC1*, *KMT2D*, *NOTCH1* exon 34, *BIRC3*) that are recurrently mutated in \geq 5% of MCL tumors was specifically designed.¹⁰⁻¹² We also included in the panel *TRAF2*²⁰ and *CXCR4*.²¹ NGS libraries preparation was performed using TruSeq Custom Amplicon sequencing assay according to manufacturer's protocol (Illumina, Inc., San Diego, CA). Multiplexed libraries (n=48 per run) were sequenced using 300-bp paired-end runs on an Illumina MiSeq sequencer, (median depth of coverage 2356x). A robust and previously validated bioinformatics pipeline was used for variant calling (Supplementary Appendix). Copy number variation analysis methods^{22,23} are supplied in the Supplementary Appendix.

Minimal residual disease analysis

For MRD purposes, MCL diagnostic BM and PB samples were investigated for IGH gene rearrangements and BCL1/IGH MTC by qualitative PCR.²⁴⁻²⁶ Both BM and PB samples were analyzed for MRD at specific time points during and after treatment. Further information are supplied in the Supplementary appendix.

Statistical analysis

The primary outcome of the clinical study was progression-free survival (PFS) and secondary outcomes included overall survival (OS).²⁷ The adjusted effects of mutations and exposure variables (MIPI-c and blastoid variant) on PFS and OS were estimated by Cox regression. To compare clinical baseline features between patients enrolled in the molecular study and patients not included in the analysis, we used Mann-Whitney test for continuous variables and Pearson's chi-squared test for categorical variables. Statistical analyses were performed using Stata 13.0 and R 3.4.1. Further information are supplied in the Supplementary appendix.

Validation set

The Nordic Lymphoma Group "MCL2" and "MCL3", phase 2, prospective trials¹⁷ were used for independent validation of our findings. In particular, the raw sequencing data of the study by *Eskelund* were reanalyzed according to our bioinformatics pipeline (detailed before), to get a uniform mutation calling.

RESULTS

Patients characteristics

Out of the 300 patients enrolled in the FIL-MCL0208 clinical trial, 186 (62%) were provided with CD19⁺ sorted tumor cells from the BM and evaluable for both mutations and copy number abnormalities. Moreover, four more patients were provided with the copy number abnormalities data only. Baseline features of cases included in the molecular study overlapped with those of cases not included in the molecular analysis because of lack of tumor material in the BM aspirates. As expected, tumor cells were obtained more frequently in cases with BM infiltration documented by morphological or flow-cytometry analysis (Table 1). Overall, this observation did not introduce a selection bias, since cases evaluable for genomic studies showed a similar outcome to that of cases not analyzed, both in terms of PFS and OS (Figure S1).

Description of genomic alterations

At least one somatic non-synonymous mutation affecting genes of the target region was observed in 69.8% of patients (130/186) (Figure 1, Figure S2; Table S2). Mutated genes were *ATM* (41.9%), followed by *WHSC1* (15.6%), *KMT2D* (12.4%), *CCND1* (11.8%), *TP53* (8.1%), *NOTCH1* (7.5%), *BIRC3* (5.9%) and *TRAF2* (1.1%). *KMT2D* deletion occurred in 1.6% of patients (3/190) and *TP53* deletion in 13.2% patients (25/190). *TP53* was inactivated by mutations or deletions in 31/186 (16.6%) cases, including 8/186 (4.3%) mutated/deleted cases, 16/186 (8.6%) deleted but not mutated cases, and 7/186 (3.7%) mutated but not deleted cases. *KMT2D* was inactivated by mutations or deleted but not mutated but not mutated cases, and 22/186 (11.8%) mutated but not deleted cases.

KMT2D mutations and TP53 disruption associate with poor outcome in MCL

By univariate analysis, mutations of *KMT2D* were associated with poor clinical outcome in terms of both PFS and OS. At 4 years, the PFS of *KMT2D* mutated patients was 33.2% vs 63.7% (p<0.001) in wild type cases (Figure 2 A). The OS of *KMT2D* mutated patients was 62.3% vs 86.8% (p=0.002) in wild type patients (Figure 2 B). Consistent with previous reports, both *TP53* mutations and deletion associated with shorter PFS and OS at 4 years (Figure 2 C,D and Figure 3). In detail, the negative prognostic impact for *TP53* disruption was equal for all the three inactivation modalities, which were then considered as a single group for further analyses (Figure S3). No further survival analysis was performed on *KMT2D* deletions, given the low frequency of this genetic lesion. All the other investigated mutations did not show strong association with PFS or OS (Figures S4-6 and Table S3).

Patients harboring *TP53* disruption were significantly enriched in known high-risk features of MCL. Indeed, 48.3% of the *TP53* disrupted patients had Ki-67 \geq 30%, 37.9% scored in the higher MIPI-c risk classes (i.e. "intermediate-high" and "high"), and 22.6% showed blastoid morphology. Conversely, 45.5% of cases harboring *KMT2D* mutations scored in the higher MIPI-c risk classes but did not associate with Ki-67 expression or blastoid morphology (Table S4). Moreover, *KMT2D* mutated patients showed slightly higher beta-2 microglobulin (B2M) median values, as well as higher prevalence of B symptoms and bulky disease (>5 cm) than wild type patients (all p<0.05). Interestingly, also *TP53* disrupted patients showed slightly higher B2M median values (p<0.05) than wild type patients (Table S4) and were associated with a high rate of disease progression during treatment (9 out of 31 patients, 29%). Moreover, *TP53* disrupted patients reached lower levels of MRD negativity after ASCT, if compared with wild type ones: 35% vs 58% in BM (p=0.06) and 58% vs 80% in PB (p=0.04), respectively. Similar trends were seen for *KMT2D* mutated patients (46% vs 55% in BM and 58% vs 79%), albeit not statistically significant (Table S5). Analogous to the Nordic

Lymphoma Group MCL2 and MCL3 trials¹⁷, also in our study morphological BM involvement was significantly associated with the presence of mutations in any of the genes analysed (p < 0.05). However, both *TP53* disruptions and *KMT2D* mutations were equally distributed in patients with and without BM involvement (p = 0.26 and p = 0.32, respectively).

By multivariate analysis adjusted for the validated risk factors MIPI-c and blastoid variant, both *KMT2D* mutations and *TP53* disruptions maintained an independent increased hazard of progression and death (Table 2 and S6). Patients carrying at least one of these genetic lesions, namely *KMTD2* mutations, *TP53* mutations or deletion (n=49/186, 26.3%), had a 4-year PFS of 32.0% vs 69.9% of wild type patients (p<0.0001) and a 4-year OS of 65.1% vs 90.3% (p<0.0001), respectively (Figure 4).

Integration of a genetic score into the MIPI-c: the "MIPI-g" model

In order to integrate the clinical impact of *KMT2D* mutations and *TP53* disruptions into the MIPI-c prognostic index (complete data available for 172 patients), we assigned a score to each of the single variables, based on the multivariate Cox regression analysis. MIPI-c low, low-intermediate and intermediate-high risk classes scored 0 points, MIPI-c high-risk class scored 1 point, while *KMT2D* mutations as well as *TP53* disruption scored 2 points (Table 3). Patients were then grouped into three risk classes, according to their total score, in the "MIPI-genetic" index ("MIPI-g"), namely: *i*) 0 points, low risk group (LR 121 patients, 70.3%); *ii*) 1-2 points, intermediate risk group (IR 38 patients, 22.1%); *iii*) \geq 3 points, high risk group (HR 13 patients, 7.6%). PFS and OS at 4-years for low-, intermediate-, and high-risk groups were 72.0%, 42.2%, 11.5% (p<0.0001) and 94.5%, 65.8%, 44.9% (p<0.0001), respectively (Figure 5). The MIPI-g index improved the model discrimination ability, with a C-statistics of 0.675 for PFS (bootstrapping corrected 0.654) and 0.776 for OS (bootstrapping corrected 0.747), as compared to MIPI-c alone (C-statistics 0.592 and 0.7, respectively).

Validation set

Most *KMT2D* variants considered in the Nordic study have been removed by our mutational calling, since these were missense variants not reported in COSMIC. At the end of the re-analysis, from the original 28 mutations, 21 were excluded. Two previously unrecognized frameshift mutations have been identified by our bioinformatics pipeline, overall accounting for a total of 9 *KMT2D* mutations (all disrupting, as expected for *KMT2D*) in the Nordic validation series. In the Nordic validation series, *KMT2D* mutated patients showed a similar increased risk for OS, with a median OS 12.7 years (95% C.I. not evaluable) for wild type vs. 8.4 (95% C.I 0-17.6) for mutated cases. The Nordic validation series also replicated the MIPI-g score. The re-analysis of *TP53* mutations confirmed the original data of Eskelund *et al.*, with median OS of 12.7 (95% C.I. not evaluable) for wild type cases and 2.0 years (95% C.I, 1.2-2-8) for mutated cases. Consistently, also the MIPI-g validation on the Nordic series showed similar results: 4-year OS for LR (N=103), IR (N=36) and HR (N=13) MIPI-g groups were 91.3%, 72.2%, 15.4%.

DISCUSSION

To identify new molecular predictors in MCL, we performed targeted resequencing and DNA profiling of purified tumor samples collected from younger patients enrolled in the ASCT-based prospective FIL-MCL0208 phase III trial (NCT02354313). Our study documents that: *i) KMT2D* mutations are a novel, independent, adverse genetic biomarker in MCL, impacting both on PFS and OS (Figure 2 A,B); *ii) TP53* aberrations (both mutations and deletion) prospectively confirm their adverse prognostic value in younger MCL patients receiving high-dose chemo-immunotherapy followed by ASCT, both in terms of PFS and OS (Figures 2 C,D and Figure 3); *iii)* identification of either *KMT2D* mutations or *TP53* disruption (or both) defines a high-risk group of young MCL patients whose outcome is still not satisfactory despite intensive immunochemotherapy and ASCT (Figure 4); *iv*) these

biomarkers may be incorporated into a "genetic" MIPI-c ("MIPI-g") model, accounting for three risk classes (Figure 5), that improve the C-statistics discrimination ability on survival, if compared to MIPI-c alone.

The adverse prognostic value of *TP53* mutations in MCL has been already observed in some retrospective series,¹³⁻¹⁷ and has been recently confirmed in a combined series from two, ASCT-based, phase II trials of the Nordic Lymphoma Group.¹⁷ *TP53* deletions impacted on both PFS and OS in the randomized, phase III European MCL Network "Younger" trial,¹⁶ while these data were not confirmed by multivariate analysis in the Nordic study, due to the high association with *TP53* mutations.¹⁷ Our prospective study performed in a similar patient population of young MCL patients demonstrates that the presence of either *TP53* mutations or deletions or both associates with poor prognosis. Importantly, although *TP53* aberrations associated with elevated Ki-67, higher MIPI-c classes and blastoid morphology, their impact on survival was independent of these known risk factors. Moreover, *TP53* disrupted patients show higher levels of MRD positivity after ASCT, as described in Table S5. Finally, some previous studies reported also a negative impact of *NOTCH1* mutations in univariate analysis,^{10,17} however in our cohort these mutations were not an independent predictor of survival, as most of them co-occurred with *TP53* mutations.

In the FIL-MCL0208 trial, *KMT2D* mutations emerged as a novel biomarker heralding chemoimmunotherapy failure, with a predictive value similar to that of *TP53* aberrations. *KMT2D* (Lysine Methyltransferase 2D), also known as *MLL2*, acts as a tumor suppressor gene mutated in several B-cell lymphoma types, including 10-15% of MCL.²⁸⁻³¹ Even though *KMT2D* mutated patients of the FIL-MCL0208 trial scored in the higher MIPI-c risk classes, they did not show either elevated Ki-67 or blastoid morphology, suggesting that *KMT2D* mutations capture high-risk patients not otherwise identifiable through conventional pathologic parameters.

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To the best of our knowledge, the adverse impact on cancer survival of *KMT2D* mutations has not been documented to date. No impact on survival was found for *KMT2D* mutations in the Nordic study.¹⁷ The lack of impact on survival of *KMT2D* mutations in the Nordic MCL series might be related to two main reasons. First, in the Nordic series, most *KMT2D* mutations were missense sequence variants (15/28) not reported as somatic variants in the COSMIC database, and therefore not fulfilling the criteria of "true" mutations. Conversely, in our series 74% of *KMT2D* mutations were protein truncating events, as expected²⁸⁻³¹. Second, since Eskelund et al. performed mutational analysis in unsorted bone marrow samples, the low or absent tumor content of many cases might lead to underestimate *KMT2D* "true" mutations. By applying our bioinformatics pipeline to the raw sequencing data of the MCL2 and MCL3 Nordic Lymphoma Group trials, we validated the poor prognostic role of *KMT2D* mutations in an independent prospective cohort.

The independent adverse prognostic value of *TP53* and *KMT2D* aberrations prompted us to integrate the molecular results into the MIPI-c,⁸ aiming at further improving its ability to discriminate high-risk patients. The "MIPI-genetic" ("MIPI-g") was able to divide patients into three risk classes, on the basis of a simple score given to each variable (namely: MIPI-c class, *TP53* disruption and *KMT2D* mutations). Patients in the high "MIPI-g" risk groups may deserve new treatments, and a simple tool like the MIPI-g might be proposed in a future, "tailored" trial to select high-risk MCL patients for targeted experimental strategies.

Our study suffers from some limitations. The analyses were performed only on CD19+ sorted BM cells and no tissue control is available at the moment; this issue might represent a limit for the extrapolation of the results to lymph-node samples, as across-compartment heterogeneity of the mutational landscape is described in MCL.¹⁰ However, the CD19+ selection approach we used, increasing the purity of tumor cells and, consequently, the sensitivity of our mutational approach, ensured that all the analyzed samples are representative of MCL. Therefore, we set a VAF threshold of

10% to call a mutation, accordingly to ERIC guidelines for the mutational analysis of the *TP53* gene in chronic lymphocytic leukemia.³² Although we acknowledge that the present validation relies on a limited number of *KMT2D* mutated patients, we note that the Nordic trials are currently the only prospective studies with prompt available mutational data, adequate clinical follow-up and similar characteristics (i.e. patients age and treatment schedule), to validate our original findings from the "FIL-MCL0208" trial.

The impact of lenalidomide maintenance in the FIL-MCL0208 trial on the described genetic aberrations has not been addressed, as complete data on randomization are not available yet. However, it should be noted that due to the high number of progressive diseases in the aberrant *TP53/KMT2D* group, 27 patients have been finally randomized but only 9 actually started lenalidomide maintenance. Therefore, it is unlikely that lenalidomide might play a clear role in driving the outcome of these patients and the trial will probably not be able to fully address this issue even with longer follow-up.

In conclusion, our results show that *KMT2D* mutated and/or *TP53* disrupted younger MCL patients are a high-risk population, characterized by poor sensitivity even to intensified chemo-immunotherapy. Given the negative prognostic impact of these genetic lesions, they might be used to select high-risk patients for novel therapeutic approaches that can circumvent these detrimental genetic lesions. As in other lymphoid disorders, novel non-chemotherapeutic strategies specifically designed for high-risk patients need to be investigated in MCL. Besides the approved drugs lenalidomide and ibrutinib, new molecules such as the Bcl-2 inhibitor venetoclax might be very promising for these chemorefractory patients, especially for *TP53* disrupted cases.^{33,34} Moreover, as the majority of *KMT2D* mutated and/or *TP53* disrupted patients of our series actually achieve a response, though shortlasting after ASCT, an alternative consolidation with allogeneic transplantation deserves investigation.

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Characteristics	Patients analysed for mutations and/or CNV (N=190)	Patients not analysed for mutations and CNV (N=110)	P-value
Median age	57	58	0.987
Gender			
Female	47 (24.7%)	18 (16.4%)	0.090
Male	143 (75.3%)	92 (83.6%)	
Ki-67			
<30%	126 (71.6%)	61(64.2%)	0.210
≥30%	50 (28.4%)	34 (35.8%)	
Median WBC	74500/ul	75000/ul	0.567
ECOG			
0	144 (75.8%)	87 (79.2%)	0 700
1	40 (21.1%)	19 (17.3%)	0.722
2	6 (3.2%)	4 (3.6%)	
Median LDH	275.5 UI/L	298	0.848
Risk class MIPI			
Low	114 (60.0%)	66 (60.0%)	0 5 6 0
Intermediate	49 (25.8%)	24 (21.8%)	0.562
High	27 (14.2%)	20 (18.2%)	
Risk class MIPI-c	, , , , , , , , , , , , , , , , , , ,	· · ·	
Low	88 (50.0%)	45 (47.4%)	
Low-Intermediate	49 (27.8%)	30 (31.6%)	0.685
Intermediate/High	25 (14.2%)	10 (10.5%)	
High	14 (8.0%)	10 (10.5%)	
BM Invasion		. ,	
No	26 (13.9%)	37 (33.9%)	<0.001
Yes	161 (86.1%)	72 (66.1%)	
Median BM invasion by flow (%)	10%	0.8%	<0.0001
Histology			
MCL Classic	174 (91.6%)	100 (90.9%)	0.842
MCL blastoid variant	16 (8.4%)	10 (9.1%)	
Bulky mass			
No	124 (65.3%)	78 (70.9%)	0.315
Yes	66 (34.7%)	32 (29.1%)	

Table 1. Clinical and biological baseline characteristics of the patients included and not included in the molecular analysis

CNV, Copy Numer Variation Analysis; WBC, White Blood Cells; ECOG, Eastern Cooperative Oncology Group; LDH, Lactate dehydrogenase; MIPI, MCL International Prognostic Index; MIPI-c, Combined MIPI; BM, Bone Marrow

	Progression free survival					Overall survival						
		Univariat	e	Multivariate (MIPI-c and blastoid variant adjusted)		Univariate			Multivariate (MIPI-c and blastoid variant adjusted)			
Genes	HR	95% CI	P-value	HR	95% CI	P-value	HR	95% CI	P-value	HR	95% CI	P-value
ATM mut	1.29	0.84-1.97	0.245	1.19	0.77-1.83	0.432	1.52	0.62-2.51	0.527	1.05	0.52-2.12	0.887
WHSC1 mut	1.53	0.90-2.60	0.119	1.51	0.87-2.61	0.140	0.85	0.30-2.41	0.755	0.741	0.25-2.15	0.581
CCND1 mut	0.83	0.41-1.66	0.595	0.94	0.46-1.92	0.860	0.75	0.23-2.48	0.643	1.01	0.29-3.53	0.980
KMT2D mut	2.59	1.50-4.48	0.001	2.74	1.55-4.84	0.001	3.20	1.48-6.92	0.003	2.48	1.12-5.46	0.024
TP53 mut	2.84	1.57-5.13	0.001	2.55	1.36-4.78	0.003	5.28	2.44-11.45	<0.0001	2.78	1.09-7.06	0.032
NOTCH1 mut	1.86	0.93-3.72	0.078	1.57	0.76-3.24	0.226	1.34	0.41-4.40	0.629	0.61	0.17-2.12	0.609
BIRC3 mut	0.88	0.32-2.41	0.807	0.70	0.25-1.96	0.500	1.84	0.56-6.08	0.315	1.15	0.33-3.98	0.822
TP53 del	3.51	2.09-5.88	< 0.0001	3.13	1.73-5.68	<0.001	4.46	2.14-9.29	< 0.0001	2.65	1.06-6.59	0.036
TP53 dis	3.39	2.10-5.45	<0.0001	3.17	1.87-5.38	<0.0001	4.26	2.09-8.67	<0.0001	2.65	1.10-6.37	0.030

Table 2. Uniavariate and multivariate Cox-regression analysis in terms of PFS an OS

Del, deleted; dis, disrupted; PFS, Progression Free Survival; OS, overall survival; HR, Hazard Ratio; CI, Confidence Interval.

Table 3. The MIPI-g score

Variables	Beta- coefficients	Points
KMT2D mutations	1,035,607	2
TP53 disruption	1,113,875	2
MIPI-c		
Low	-	0
Low-Intermediate	-	0
Intermediate-High	-	0
High	0.6847757	1

MIPI-c, Combined Mantle Cell International Prognostic Index

FIGURE LEGENDS

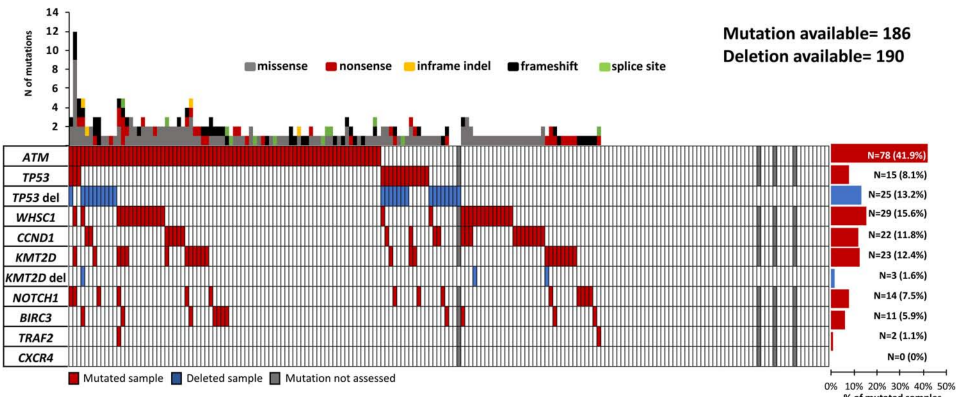
Figure 1. Overview on prevalence and molecular spectrum of non-synonymous somatic mutations discovered in patients' tumor DNA. Heatmap representing the mutational profiles of 186 MCL cases, genotyped on tumor DNA (and four additional patients with copy number abnormalities data only). Each column represents one patient, each row represents one gene. The fraction of patients with mutations in each gene is plotted on the right. The number of aberrations in a given patient is plotted above the heatmap.

Figure 2. Prognostic impact of *KMT2D* **and** *TP53* **mutations.** Kaplan-Meier estimates of progression free survival and overall survival of *KMT2D* (A, B), and *TP53* (C, D) mutated versus wild type patients. Cases harboring mutations (mut) in these genes are represented by the yellow line. Cases wild type (wt) for these genes are represented by the blue line. The Log-rank statistics *p* values are indicated adjacent curves.

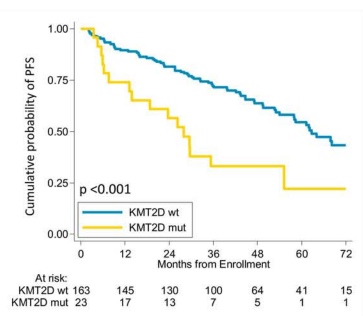
Figure 3. Prognostic impact of *TP53* deletion. Kaplan-Meier estimates of progression free survival (A) and overall survival (B) of *TP53* deleted versus wild type patients. Cases with *TP53* deletion are represented by the yellow line. Cases without *TP53* deletion are represented by the blue line. The Logrank statistics p values are indicated adjacent curves.

Figure 4. Prognostic impact of combined *KMT2D* mutations and *TP53* disruption. Kaplan-Meier estimates of (A) progression free survival and (B) overall survival of patients harboring *KMT2D* mutations and/or *TP53* disruption (mutations and/or deletions). Cases harboring at least one of these 3 genetic lesions are represented by the yellow line. Cases without these genes are represented by the blue line. The Log-rank statistics p values are indicated adjacent curves.

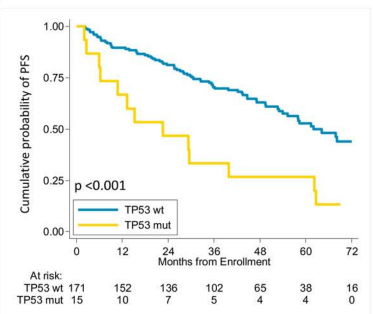
Figure 5. The "MIPI-g" model. Kaplan-Meier estimates of (A) progression free survival and (B) overall survival of patients harboring *KMT2D* mutations and/or *TP53* disruption (mutations and/or deletions) integrated into the MIPI-c. Low MIPI-g risk cases are represented by the blue line, intermediate MIPI-g cases by the yellow line and high MIPI-g cases by the red line. The Log-rank statistics p values are indicated adjacent curves.

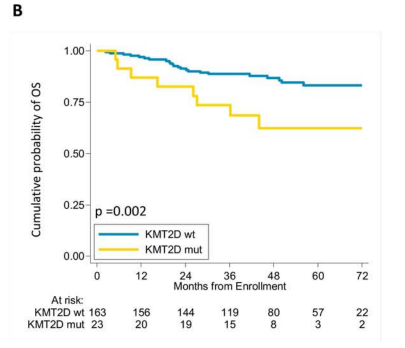


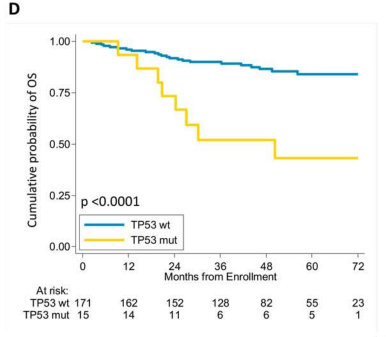
% of mutated samples



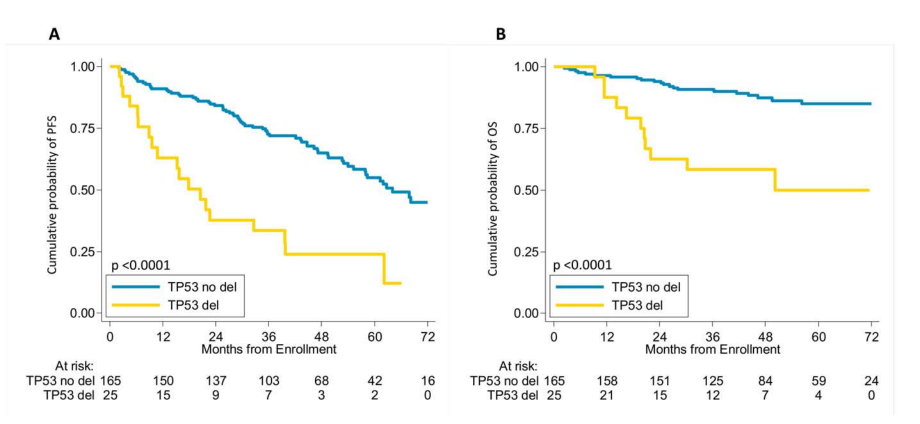
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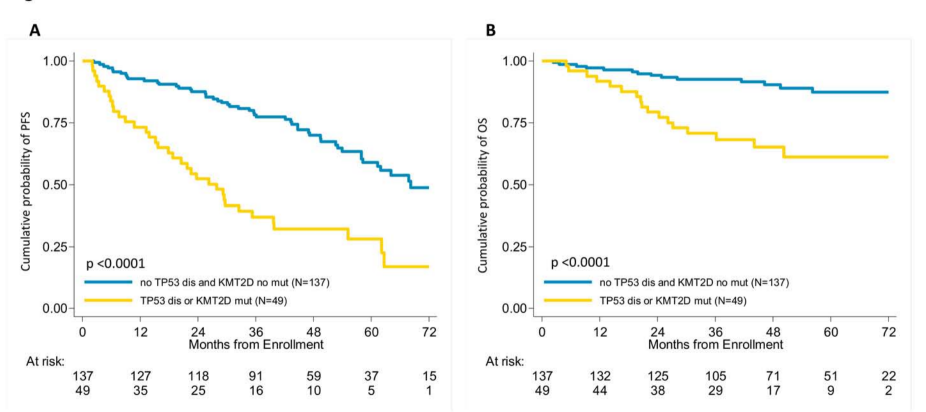


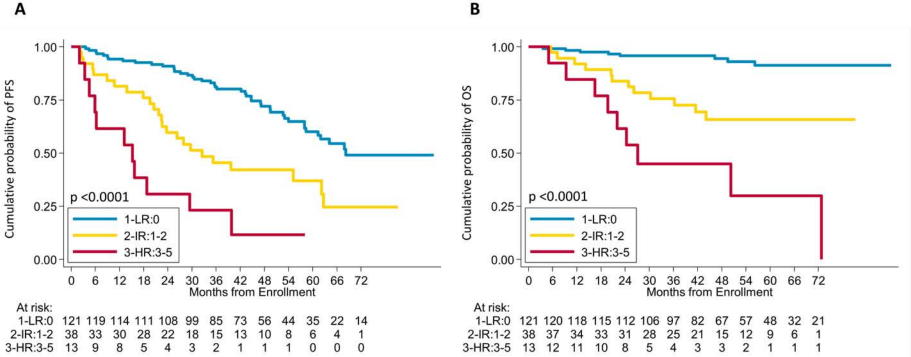




Α







SUPPLEMENTARY APPENDIX

KMT2D mutations and *TP53* disruptions are poor prognostic biomarkers in MCL receiving high-dose therapy: a FIL study

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SUPPLEMENTARY METHODS

Patients series

The FIL-MCL0208 (NCT02354313) is a phase III, multicenter, open-label, randomized, controlled study, designed to determine the efficacy of lenalidomide as maintenance versus observation in young (18-65 years old), fit, advanced stage (Ann arbor II-IV) MCL patients who achieved complete or partial remission after first line intensified and high-dose chemotherapy plus rituximab followed by ASCT. Briefly, patients received 3 R-CHOP-21, followed by R-high-dose cyclophosphamide (4g/m2), 2 cycles of R-high-dose Ara-C (2g/m2 q12x3 d) and ASCT conditioned by using the BEAM or FEAM regimen. From May 2010 to August 2015, a total of 300 patients were enrolled in the study. Cases of non-nodal MCL were excluded.¹ All patients required to have a biopsy proving MCL, including evidence of cyclin D1 overexpression or t(11;14)(q13;q32) translocation. MCL diagnosis was confirmed by centralized pathological revision according to WHO criteria.¹ The clinical trial, as well as the ancillary mutational study, were approved by the Ethical Committees of all the enrolling Centers. All patients provided written informed consent for the use of their biological samples for research purposes, in accordance with Institutional Review Boards requirements and the Helsinki's declaration. Clinical results of the fist interim analysis of the trial were already presented.² Final, unblinded results are not available at the moment and have not been yet presented anywhere.

Biological samples

Bone marrow (BM) and peripheral blood (PB) samples were collected, as per protocol, at baseline and at several time-points during follow-up, corresponding to the pre-planned time-points for minimal residual disease (MRD) analysis. To identify and quantify the rate of tumor invasion, flow cytometry (FC) was performed both on BM and PB with the following antibodies: anti-CD19 APC, anti-CD23 PE, anti-CD5 FITC, and anti-CD20. Tumor cells were sorted from the baseline BM samples by immunomagnetic beads (CD19 MicroBeads,human-Miltenyi Biotec GmbH, Bergisch Gladbach, Germany) and stocked as dry pellets.

Tumor DNA was extracted according to DNAzol protocol (Life Technologies). Germline DNA was obtained from PB mononuclear cells collected under treatment and proven to be tumor free by MRD analysis.

Next generation sequencing

A targeted resequencing panel (target region: 37'821 bp) (Table S1) including the coding exons and splice sites of 7 genes (*ATM*, *TP53*, *CCND1*, *WHSC1*, *KMT2D*, *NOTCH1* exon 34, *BIRC3*) that are recurrently mutated in \geq 5% of MCL tumors was specifically designed.³⁻⁵ We also included in the panel *TRAF2* ⁶ and *CXCR4*.⁷ The gene panel was analyzed in tumor DNA from baseline BM CD19⁺ purified MCL cells (186 cases) and, for comparative purposes to filter out polymorphisms, in the paired normal genomic DNA (105 cases). NGS libraries preparation was performed using TruSeq Custom Amplicon sequencing assay according to manufacturer's protocol (Illumina, Inc., San Diego, CA). Multiplexed libraries (n=48 per run) were sequenced using 300-bp paired-end runs on an Illumina MiSeq sequencer, (median depth of coverage 2356x). To avoid the loss of *NOTCH1* ex 34 c.7544_7545delCT mutation, that is included in a region poorly covered by the target design, all MCL cases were also analyzed by amplification refractory mutation system (ARMS) polymerase chain reaction (PCR).

Bioinformatic analysis

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. A cut-off of 10% of variant allele frequency (VAF) was set for variant calling. Among cases provided with both tumor and paired normal gDNA, single nucleotide variations and indels were called using

the somatic function of VarScan2. The variants called by VarScan2 were annotated by 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 postprocessing filters were then applied to the remaining variants to further improve variant call confidence. Accordingly, variants represented in >10 reads of the paired germline and/or variants with a somatic p value from VarScan2 >3.305e⁻⁷ [multiple comparisons corrected p threshold=3.305e⁻ ⁷, corresponding to alpha of 0.05/(37'821 x 4 alleles per position)] were no further considered. Variant allele frequencies for the resulting candidate mutations and the background error rate were visualized using IGV. Among patients lacking the paired normal gDNA, single nucleotide variations and indels were called in tumor gDNA with the cns function of VarScan2. The variants called by VarScan2 were annotated by 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. Only protein truncating variants (i.e. indels, stop codons and splice site mutations), as well as missense variants not included in the dbSNP 138 and annotated as somatic in the COSMIC v78 database, were retained.

Copy number variation analysis

DNA profiling was performed on DNA samples derived from baseline BM CD19+ purified tumor cells using the HumanOmni2.5 arrays (Illumina, San Diego, CA, USA). Copy number status of the genes included in the targeted resequencing panel (*KMT2D* and *TP53*) was assessed after genomic profiles segmentation with the Fast First-derivative Segmentation Algorithm, as previously described.^{8,9}

Minimal residual disease analysis

For MRD purposes, MCL diagnostic BM and PB samples were investigated for IGH gene rearrangements and BCL1/IGH MTC by qualitative PCR. Briefly, IGH were screened using forward consensus primers annealing the IGH-V-regions and a reverse primer complementary to the JH region. BCL1/IGH MTC translocation was investigated by nested-PCR approach, as described.¹⁰⁻¹² After direct sequencing, FASTA files alignment was performed by IMGT/V-QUEST (http://imgt.org) and BlastN tool (NCBI, https://blast.ncbi.nlm.nih.gov/Blast.cgi), in order to define rearranged loci nomenclature, chromosomic breakpoints, and to assess patient specific nucleotide insertions (N insertions), then used to design allele specific oligonucleotides primers for nested-PCR MRD monitoring. Therefore, both BM and PB samples were analyzed for MRD at specific time points during and after treatment.

Statistical analysis

The primary outcome of the clinical study was progression-free survival (PFS). PFS was calculated from the date of enrolment into the clinical study to the date of disease progression (event), death from any causes (event), or last follow up (censoring).¹³ Secondary outcomes included overall survival (OS), measured from the date of enrolment into the clinical study to the date of death from any causes (event), or last follow up (censoring). Time-to-event outcomes (PFS and OS) were estimated using the Kaplan-Meier method and compared between groups using the Log-rank test. The adjusted effects of mutations and exposure variables (MIPI-c and blastoid variant) on PFS and OS were estimated by Cox regression. To compare clinical baseline features between patients enrolled in the molecular study and patients not included in the analysis, we used Mann-Whitney test for continuous variables and Pearson's chi-squared test for categorical variables. A Cox model for PFS was estimated including MIPI-c and clinically impacting genetic alterations, and an additive score was computed according to the proportion between each predictor coefficient and the lowest one. The Cox model was internally validated using 1000 bootstrap samples and the C-statistic correct

for optimism was also provided. Patients were then grouped in classes of risk according to their total score using the nonparametric tree modelling technique of classification and regression tree (CART) analysis. Statistical analyses were performed using Stata 13.0 and R 3.4.1. The outcome data for the present analysis were updated as of December, 2017 the randomization arms being still blinded.

SUPPLEMENTARY FIGURE LEGENDS

Figure S1. Survival analysis for patients enrolled in the MCL0208 clinical trial included and not included in the present molecular study. Kaplan-Meier estimates of progression free survival (A) and overall survival (B) of patients with available DNA included in the present molecular study (in blue) and of patients without available DNA not included in the present molecular study (in yellow). The Log-rank statistics *p* values are indicated adjacent curves.

Figure S2. Disposition of identified gene mutations across the protein. Mutations identified in the studied cohort are plotted above the protein divided into the main domains. Missense mutations are plotted in green, stop codon mutations in red, splicing mutations in black, frameshift mutations in red, in-frame mutations in yellow.

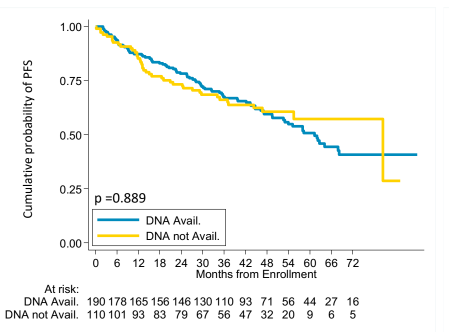
Figure S3. Prognostic impact of *TP53* mutation and *TP53* deletion. Kaplan-Meier estimates of progression free survival (A) and overall survival (B) of *TP53* mutated patients, *TP53* deleted patients, *TP53* mutated and deleted patients, versus wild type patients. Cases with *TP53* mutation are represented by the yellow line, cases with *TP53* deletion are represented by the red line, cases with *TP53* mutation and deletion are represented by the black line, cases with *TP53* mutation and deletion are represented by the blue line. The Log-rank statistics p values are indicated adjacent curves.

Figure S4. Prognostic impact of *NOTCH1* **mutation and** *ATM* **mutation.** Kaplan-Meier estimates of (A) progression free survival and (B) overall survival of patients harboring *NOTCH1* mutation and Kaplan-Meier estimates of (C) progression free survival and (D) overall survival of patients harboring *ATM* mutation. Cases harboring *NOTCH1* or *ATM* mutation are represented by the yellow line. Wild

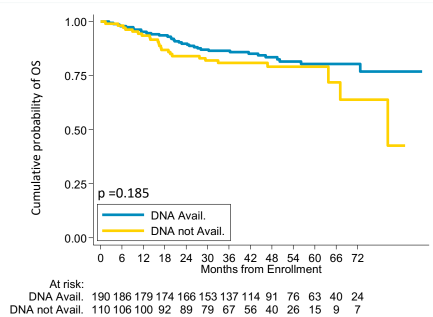
type cases are represented by the blue line. The Log-rank statistics p values are indicated adjacent curves.

Figure S5. Prognostic impact of WHSC1 mutation and CCND1 mutation. Kaplan-Meier estimates of (A) progression free survival and (B) overall survival of patients harboring *WHSC1* mutations and Kaplan-Meier estimates of (C) progression free survival and (D) overall survival of patients harboring *CCND1* mutation. Cases harboring *WHSC1* or *CCND1* mutation are represented by the yellow line. Wild type cases are represented by the blue line. The Log-rank statistics *p* values are indicated adjacent curves.

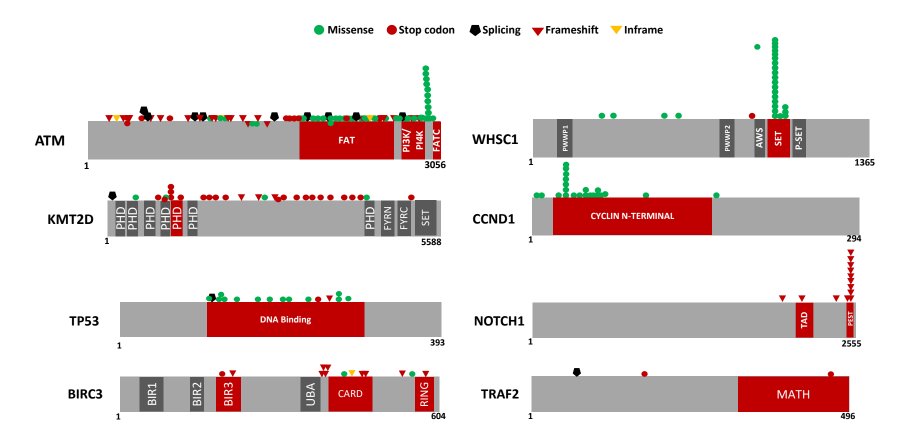
Figure S6. Prognostic impact of *BIRC3* mutation and *TRAF2* mutation. Kaplan-Meier estimates of (A) progression free survival and (B) overall survival of patients harboring *BIRC3* mutation and Kaplan-Meier estimates of (C) progression free survival and (D) overall survival of patients harboring *TRAF2* mutation. Cases harboring *BIRC3* or *TRAF2* mutation are represented by the yellow line. Wild type cases are represented by the blue line. The Log-rank statistics p values are indicated adjacent curves.

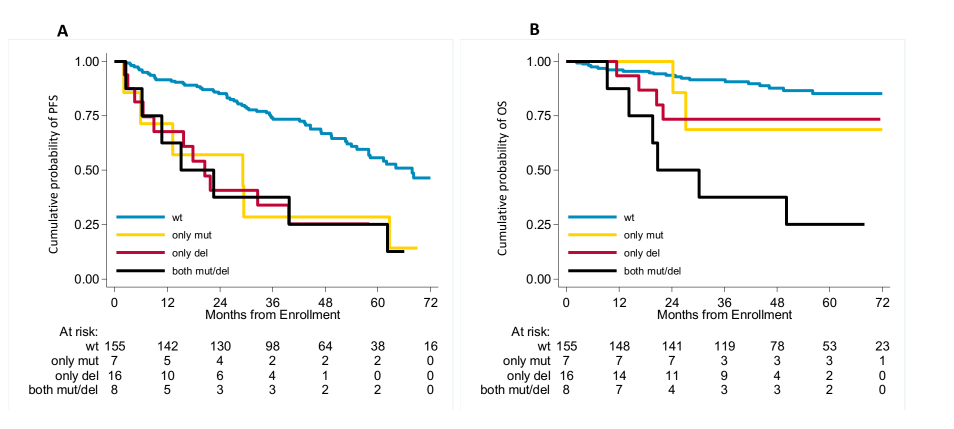


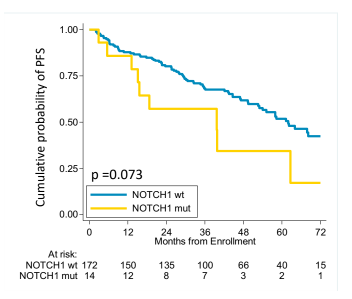
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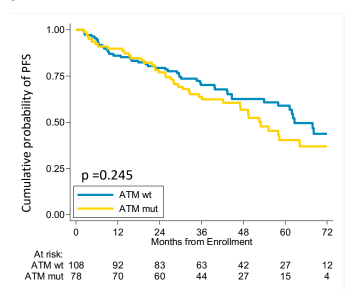
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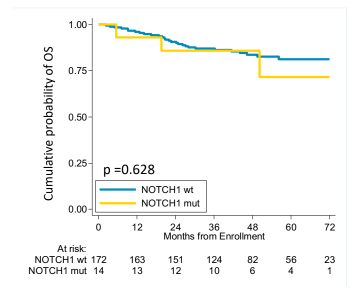




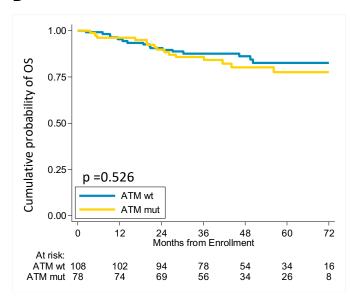


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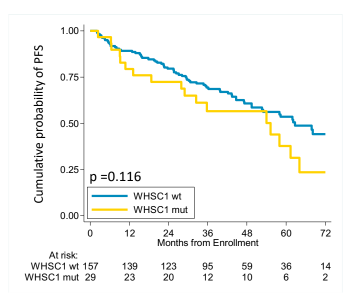


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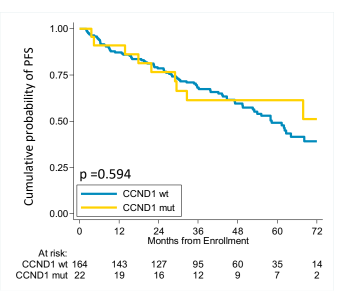


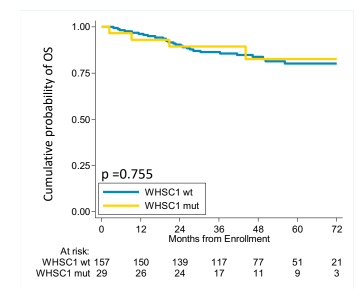
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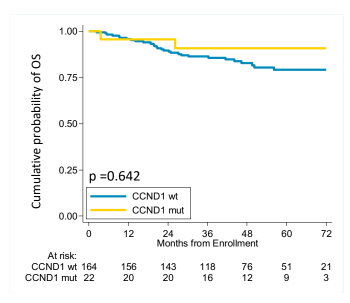


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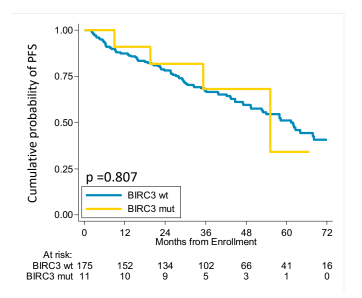




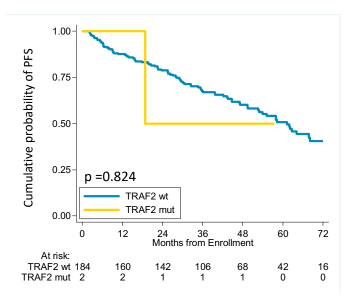
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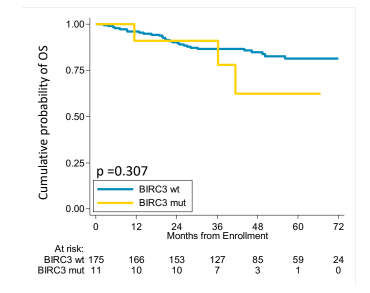


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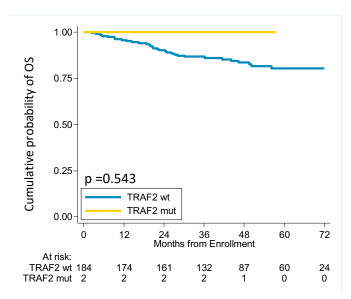


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SUPPLEMENTARY TABLES EXCEL FILE ONLY

Table S1. Target region

Table S2. Somatic non-synonymous mutations discovered in the 186 cases of the molecular study

Table S3. Log-rank univariate analysis in terms of PFS and OS according to mutational status

Table S4. Clinical and biological baseline features of KMT2D and TP53 disrupted patients

Table S5. MRD assessment after ASCT in TP53 disrupted and in KMT2D mutated patients

Table S6. Multivariate analysis of TP53 disruption and KMT2D mutations

SUPPLEMENTARY REFERENCES

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