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Genetic and Phenotypic Attributes of Splenic Marginal Zone Lymphoma

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Abstract:

Splenic marginal zone B-cell lymphoma (SMZL) is a heterogeneous clinico-biological entity. The clinical course is variable, multiple genes are mutated with no unifying mechanism, essential regulatory pathways and surrounding microenvironments are diverse. We sought to clarify the heterogeneity of SMZL by resolving different subgroups and their underlying genomic abnormalities, pathway signatures and microenvironment compositions to uncover biomarkers and therapeutic vulnerabilities. We studied 303 SMZL spleen samples collected through the IELSG46 multicenter, international study (NCT02945319) by using a multiplatform approach. We carried out genetic and phenotypic analyses, defined self-organized signatures, validated the findings in independent primary tumor meta-data and in genetically modified mouse models, and determined correlations with outcome data. We identified two prominent genetic clusters in SMZL, termed NNK (58% of cases, harboring NF-kB, NOTCH and KLF2 modules) and DMT (32% of cases, with DNA-damage response, MAPK and TLR modules). Genetic aberrations in multiple genes as well as cytogenetic and immunogenetic features distinguished NNK- from DMT-SMZLs. These genetic clusters not only have distinct underpinning biology, as judged by differences in gene-expression signatures, but also different outcome, with inferior survival in NNK-SMZLs. Digital cytometry and in situ profiling segregated two basic types of SMZL immune microenvironments termed immune-suppressive SMZL (50% of cases, associated with inflammatory cells and immune checkpoint activation) and immune-silent SMZL (50% of cases, associated with an immune-excluded phenotype) with distinct mutational and clinical connotations. In summary, we propose a nosology of SMZL that can implement its classification and also aid in the development of rationally targeted treatments.

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Genetic and Phenotypic Attributes of Splenic Marginal Zone Lymphoma

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Key Points

- SMZL comprises four distinct genetically defined molecular clusters, and two distinct phenotypically defined immune-microenvironment classes
- The molecular-based nosology of SMZL can improve disease classification and the discovery of novel biomarkers and therapeutic vulnerabilities

ABSTRACT

Splenic marginal zone B-cell lymphoma (SMZL) is a heterogeneous clinico-biological entity. The clinical course is variable, multiple genes are mutated with no unifying mechanism, essential regulatory pathways and surrounding microenvironments are diverse. We sought to clarify the heterogeneity of SMZL by resolving different subgroups and their underlying genomic abnormalities, pathway signatures and microenvironment compositions uncover biomarkers to and therapeutic vulnerabilities. We studied 303 SMZL spleen samples collected through the IELSG46 multicenter, international study (NCT02945319) by using a multiplatform approach. We carried out genetic and phenotypic analyses, defined self-organized signatures, validated the findings in independent primary tumor meta-data and in genetically modified mouse models, and determined correlations with outcome data. We identified two prominent genetic clusters in SMZL, termed NNK (58% of cases, harboring NF-kB, NOTCH and KLF2 modules) and DMT (32% of cases, with DNAdamage response, MAPK and TLR modules). Genetic aberrations in multiple genes as well as cytogenetic and immunogenetic features distinguished NNK- from DMT-SMZLs. These genetic clusters not only have distinct underpinning biology, as judged by differences in gene-expression signatures, but also different outcome, with inferior survival in NNK-SMZLs. Digital cytometry and in situ profiling segregated two basic types of SMZL immune microenvironments termed immune-suppressive SMZL (50% of cases, associated with inflammatory cells and immune checkpoint activation) and immune-silent SMZL (50% of cases, associated with an immuneexcluded phenotype) with distinct mutational and clinical connotations. In summary, we propose a nosology of SMZL that can implement its classification and also aid in the development of rationally targeted treatments.

Keywords: marginal zone, lymphoma, spleen, classification, genetics, microenvironment

INTRODUCTION

Splenic marginal zone lymphoma (SMZL) is an indolent small B-cell neoplasm deriving from lymphocytes of the splenic marginal zone and affecting the spleen, bone marrow and peripheral blood.

The incidence of SMZL is increasing, mainly because of improved diagnostic techniques resulting in more patients being diagnosed every year. However, in parallel, life expectancy of patients with SMZL is not improving. Compared to other indolent B-cell neoplasms, the survival of patients with SMZL is unsatisfactory (5-year relative survival ~79%)¹⁻³, and no breakthrough treatment advances have been seen³.

SMZL is heterogeneous at multiple levels. The clinical course is variable, with some patients having prolonged survival and a proportion (~20%) experiencing rapidly progressive disease and survival less than 5 years⁴. SMZL lacks a unifying genetic lesion. Multiple mutated genes have been identified, which are restricted to a fraction of cases.⁵⁻¹¹ Inflammatory cells are expanded in a subset of SMZL¹², suggesting the existence of different microenvironments. Clinical trials evaluating novel agents provide glimpse into signaling pathways that are essential for SMZL, but sensitivity to these agents is not always observed¹³⁻¹⁵.

The IELSG46 study (NCT02945319) is a multicenter, international, retrospective, observational study that aims at resolving the heterogeneity of SMZL into subgroups by using a multiplatform approach, with the belief that it might yield a nosology of SMZL that could be implemented for disease classification, and result in the discovery of novel biomarkers and therapeutic vulnerabilities.

METHODS

Patients

Inclusion criteria of the IELSG46 study were: i) age ≥18 years; ii) SMZL diagnosis by spleen histopathologic examination; iii) availability of tumor material from spleen collected before initiation of medical therapy; iv) availability of baseline and follow-up annotations. Patients who received any anti-tumor medical therapy before splenectomy were excluded. The study was conducted in accordance with the Declaration of Helsinki and principles of Good Clinical Practice. The study was approved by the Ethic Committee (ID: 2016-01978) and patients provided written informed consent. Coded health-related patient data and tumor biological samples were collected from eligible patients (Supplementary Methods).

Pathology review and tissue microarray

A pathology expert panel reviewed the cases (G.Bh., E.C, L.D.L., S.Dirn, L.M., S.A.P., M.A.P., M.Pa, M.Po., A.T.G.). Diagnoses were based on the WHO classification¹⁶. Cases were assembled on tissue microarrays (A.Tz) as described with slight modifications¹⁷. IGHV analysis was performed in two different ways according to quality/quantity of the starting material (Supplementary Methods).

LyV3.0 CAncer Personalized Profiling by deep Sequencing Assay

A CAPP-seq protocol was used for mutation and copy number abnormality (CNA) analysis. Libraries derived from tumor genomic DNA of FFPE (n=246) or

frozen tissues (n=57). No patient-matched normal specimens were available. The assay limit of quantification was calculated as the mean allele frequency of non-SNP variants from normal FFPE spleen "blank" samples. The assay analytical sensitivity was established by generating limiting dilutions of FFPE gDNA from one patient with active lymphoma into normal gDNA of FFPE spleen samples, resulting in expected tumor fractions between 100% and 0.2%. The number of the libraries loaded in the sequencer was tailored to obtain a coverage >2000x in >80% of the region of interest. A tumor-only, background error-suppressed approach was used for variant and copy number calling (Supplementary Methods).

RNA-seq

RNA-seq (TruSeq Stranded mRNA kit, Illumina Technology) of 45 fresh biopsy samples was used to identify gene fusions. Gene expression was assessed in FFPE biopsy samples by using HTG EdgeSeq Precision Immuno-Oncology (HTG-PIO) and HTG EdgeSeq Oncology Biomarker (HTG-OBP) Panels (HTG Molecular Diagnostics) (Supplementary Methods).

Bioinformatics and medical statistics

The following bioinformatics approaches are described in Supplementary Methods: mutation calling, CNA detection, pathway-driven clustering of mutation data, gene fusion detection, gene expression analysis, signature enrichment testing, microenvironment signatures definition and clustering, deconvolution of cell percentages from bulk transcriptomes. 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. Relative survival, defined as the ratio between actuarial survival observed in patients and expected survival of the general population matched to by geographical origin, sex, age, and calendar year of diagnosis, was calculated using the Ederer II method. Categorical variables were compared by χ^2 test and Fisher's exact test when appropriate. Continuous variables were compared by ANOVA and t-student tests when appropriate. Statistical tests were 2-sided and significance was defined as p <0.05, after correction for multiplicity where appropriate. Analyses were performed with R (http://www.r-project.org).

RESULTS

Patient characteristics

Patients diagnosed with SMZL on splenic resections were registered in the IELSG46 study by 28 centers in Europe and the US. A total of 373 patients were initially identified. Seventy cases were excluded due to alternative diagnoses on central pathology review or insufficient material (Supplementary Figure 1A). Table 1 lists the clinical characteristics of the 303 patients with confirmed SMZLs. Deletion 7q and use of IGHV1-2*04 allele, which are recurrent in SMZL^{18,19}, were detected in 26.4% and 33.9% of cases, respectively. Based on the percentage of IGHV gene identity to the germline, 11.1% of cases were 'truly unmutated' (100% homology) (Supplementary Table 1). Median follow-up after splenectomy was 10.6 years, with 86 deaths. At ten years, overall survival was 68.5% and relative survival compared to the matched general population was 82.1% (Supplementary Figure 1B). After splenectomy, which counted as first line therapy, 10.6% of patients received

additional treatment, including chemotherapy (5%) or rituximab +/- chemotherapy (5.6%).

These data confirmed the representativeness of the study cohort and the lack of biases due to the inclusion of splenectomized patients.^{3,20,21}

A genetic classifier for SMZL

We investigated mutations using the LyV3.0 CAPP-seg assay, which targeted ~280 kb of genomic space by deep sequencing and that has been specifically designed to cover the majority of coding regions known to be recurrently mutated in mature B-cell neoplasms (Supplementary Table 2). To validate the LyV3.0 CAPPseg assay, we compared in silico the enrichment of somatic mutations by LvV3.0 CAPP-seg with whole exome sequencing (WES) after simulating mutation detection using data from 32 individual SMZLs reported in the literature²². LyV3.0 CAPP-sea yielded a 23-fold increase of variant detection per sample per sequenced base pair compared to WES (Supplementary Table 3). For coding mutations, minimal increase in the mutation recovery from SMZL samples was expected by enlarging the probed genomic space over that included in LyV3.0 CAPP-seq. When applied to DNA samples from 8 FFPE tissues of normal spleen ("blank" samples) the limit of quantification of LyV3.0 CAPP-seg was 0.3%, which represented the analytical background noise threshold (Supplementary Figure 2A). When gDNA samples from 3 FFPE tissues of SMZL were diluted with control gDNA from FFPE tissues of normal spleen, the analytical sensitivity of LyV3.0 CAPP-seq was 2%, representing the lowest detectable allele frequency (Supplementary Figure 2B).

A stringent computational method designed for tumor-only samples was used to filter out artifacts, putative germline variants, variants of unknown significance and variants known to be benign or likely benign. Individual non-synonymous mutations discovered by LyV3.0 CAPP-seq (Supplementary Table 4) and recurring in >1% of SMZLs are shown in Figure 1A. The number of mutations in each individual sample was not affected by the age of the biopsy (Supplementary Figure 2C). The most frequently mutated genes were *NOTCH2*, *KLF2*, *KMT2D*, *TNFAIP3*, *NOTCH1* and *TP53*. Consistent with the diagnosis of SMZL, the rare *MYD88* mutations mapped outside the p.265 hotspot. No fusions were detected by RNA-seq. Mutations in *MYC*, *BCL2* and *BCL6*, which are surrogates for translocation of these genes adjacent to immunoglobulin genes²³, were extremely rare (Supplementary Table 4).

To segregate SMZLs into discrete genetic classes supported by coordinated mutational profiles, we started with a set of pathway-driven seed modules comprising components of several B-cell programs (Supplementary Table 5). Genes were assigned to a module based on published literature and database annotations²³⁻²⁷. Genes that were attributed to multiple modules (eg, *KLF2*) were not assigned and seeded individually. Unsupervised analysis of mutational co-occurrence between all lesion pairs revealed overall significantly stronger exclusivity between intra-pathway lesions (Figure 1B). Mutual exclusivity of mutations within a pathway could reflect functional redundancy and supported their aggregation within a seed (Figure 1C). We then applied hierarchical clustering on principal components (HCPC) and discovered four groups of tumors (clusters) with discrete genetic signatures overall accounting for 86.4% of cases, and an additional subset without detectable mutations in the interrogated genomic space (13.6% of cases) (Figure 2A). The algorithm converged on genetic clusters that for simplicity were termed NNK (58.2%

of cases, for NF-κB, NOTCH and *KLF2* modules), DMT (32.8% of cases, for DNA-damage response, MAPK and TLR modules), CBS (4.8% of cases, for cytokine, B-cell receptor signaling, and spliceosome modules) and PA (4.2% of cases, for PI3K/AKT module). Application of the same approach to an independent meta-dataset of SMZLs profiled by WES²² produced overlapping genetic clusters (Supplementary Figure 3A), thus supporting the robustness of the classification and excluding biases related to the hypothesis driven design of the genomic space we used for sequencing.

The minimal set that allowed to classify patients with SMZL into NNK and DMT clusters included 14 genes (*TP53*, *ATM*, *KLF2*, *TNFAIP3*, *NOTCH2*, *BRAF*, *MYD88*, *SPEN*, *CARD11*, *NOTCH1*, *PTPN11*, *CHD2*, *SAMHD1* and *NFKBIE*) and performed with an accuracy of 90% in the training set and of 89% in the validation set. The model was further validated in a completely independent metadataset, ²² vielding a comparable accuracy (73%) (Supplementary Figure 4).

Overall, 99.4% mutations discovered in the spleen were concordantly detected in synchronous peripheral/marrow blood, including 100% of mutations affecting the minimal set of genes. This observation indicates that molecular cluster assignment can leverage on "liquid biopsies" (Supplementary Figure 5).

NNK and DMT clusters together accounted for the vast majority (91.0%) of SMZLs with detectable mutations. NNK-SMZLs were dominated by mutations affecting NF-κB (eg, *TNFAIP3*), including non-canonical pathway genes (eg, *TRAF3*, *BIRC3*), NOTCH (eg, *NOTCH2*, *NOTCH1*, *SPEN*) and *KLF2*. *KLF2* mutations cooccurred with NOTCH mutations in 21.8% of NNK-SMZL, and with NF-kB mutations in 18.6% of cases (Figure 2B). These observations, along with the fact that NOTCH, NF-κB and KLF2 regulate MZ B-cell differentiation²⁸, and that KLF2 is a master

regulator of both NOTCH and NF-κB²⁹ signaling, suggested a pervasive oncogenic cooperation between these genetic lesions in NNK-SMZLs to hijack the MZ B-cell differentiation program. DMT-SMZLs were characterized by mutations in DNA damage response genes (eg, *TP53*, *ATM*). Mutations in MAPK (eg, *BRAF*) and TLR genes (eg, *MYD88*) were also enriched in DMT-SMZLs (Figure 2B). CBS-SMZLs harbored mutations in cytokine signaling (eg, *STAT6*, *STAT3*, *PTPN1*, *PTPRD*), B-cell receptor signaling (eg, *CD79A*, *BTK*) and spliceosome (eg, *SF3B1*, *RBMX*) genes. PA-SMZLs harbored mutations in *PIK3CA*, *ITPKB*, a non-canonical inhibitor of AKT³⁰, and *RRAGC*, an mTORC1-regulator³¹ (Figure 2B).

Molecular clusters were correlated with immunogenetic features and CNA to find specific associations (Figure 2C Supplementary Table 1 and 6). NNK-SMZLs were enriched in IGHV1-2*04 usage and 7q deletion, while conversely DMT-SMZLs showed depletion of both (Figure 2C, Supplementary Table 7).

Phenotypic differences among SMZL clusters

We applied gene expression profiling to FFPE tissue samples to explore phenotypic differences among SMZL clusters. The concordance between expression profiling of FFPE tissues and gold standard RNA-seq was validated in 57 cases with available paired frozen and FFPE SMZL specimens (Supplementary Figure 2D and E). We analyzed the entire study cohort for the expression of 2559 genes focusing on oncogenic signaling, and used 18 signatures, selected from public databases and the literature according to the genes and pathways related to the mutation signatures (Supplementary Table 8), and reflecting distinct biological processes and differentiation programs of B-cells (Supplementary Table 8 and 9). For each

signature, we calculated an activity score that is directly associated with the magnitude of a particular effect among populations of cells³². We then analyzed the correlation between the NNK- and DMT-SMZL clusters and the signatures (Figure 3A). NNK-SMZLs expressed significantly higher levels of genes belonging to the *NOTCH2* pathway and of genes that are activated by non-canonical NF-κB transcription factors³³. A proliferation signature also characterized NNK-SMZLs. Conversely, DMT-SMZLs had a signature of impaired TP53 and apoptosis functions.

We used immunohistochemistry to comparatively screen NNK-SMZLs vs DMT-SMZLs for evidence of NOTCH and NF-κB biochemical activation (Figure 3B and C; Supplementary Table 10). Consistent with the high incidence of NOTCH and NF-κB pathway mutations in NNK-SMZLs, they frequently showed nuclear staining for NOTCH, canonical- and non-canonical NF-κB proteins, while nuclear TP53 expression strongly correlated with DMT-SMZLs (Figure 3B and C).

Paired bone marrow biopsies of 96 patients were revised by the local pathologists. The bone marrow morphology was typical (i.e. nodular or nodular-interstitial pattern, with or without intrasinusoidal infiltration) in 85 cases (88.5%) and atypical (i.e. paratrabecular or diffuse patterns without intrasinusoidal infiltration) in the remaining 11 cases (11.5%). With the limitations imposed by the sample size, the majority of cases showing an atypical bone marrow morphology belonged to the DMT molecular cluster, while cases belonging to the NNK, CBS and PA molecular clusters preferentially or exclusively showed a typical bone marrow morphology (Supplementary Table 11).

Immune microenvironment of SMZL

The immune-microenvironment of SMZL is poorly understood. We thus profiled the expression of 1402 genes focusing on tumor/immune interactions, and used 16 signatures (Supplementary Table 12), reflecting distinct immune cell subtypes and immune cell programs, to virtually reconstruct the spleen microenvironment of SMZL³⁴. We analyzed the relationship of each signature among samples by applying an unsupervised clustering algorithm and obtained two major classes of predicted spleen microenvironment in SMZL (Figure 4A, Supplementary Table 13). One class, accounting for 50% SMZLs, was dominated by macrophage (eq. CD163), cytokine (eq. BAFF, IL6, IL10), chemokine (eq. CXCR335, CXCL936,37, CCL5, CX3CR138), T-cell (eg, CD3), and regulatory T-cell (eg, FOXP3) signatures, along with an expected immune-suppressive checkpoint milieu (eg, inhibitory receptors CTLA4, TIM3, PD-1, BTLA, TIGIT, LAG3 associated with T-cell exhaustion, and low expression of MHC-I genes) indicative of an immune-evasion profile (immune-suppressive class). A second class, accounting for the other 50% of SMZLs, had a microenvironment where the B-cell signatures of obvious tumor origin dominated (eg, CD19, CD20, CD79A/B, PAX5) similar to observations in immuneexcluded lymphomas (immune-silent class)³⁹. Application of the same approach to an independent meta-dataset of spleen samples of SMZL⁴⁰ produced overlapping phenotypic clusters even with a smaller sample size (Supplementary Figure 3B), thus supporting the robustness of the classification.

To confirm the different composition of the "immune-suppressive" and 'immune-silent" SMZL classes, we applied digital cytometry deconvolution⁴¹, which inferred a heterogeneous and microenvironment-rich composition in the "immune-suppressive" class of SMZL, and a high ratio of neoplastic-to-microenvironmental cells in the 'immune-silent' class (Figure 5A, Supplementary Table 14).

Orthogonal validation by immuno-phenotypical analysis confirmed the microenvironment tissue contexture derived from gene expression and digital cytometry deconvolution (Figure 5 B and C; Supplementary Tale 10). The "immune-suppressive" class of SMZL had higher intra-tumoral CD3+ T-cells, CD4+ T-cells, CD8+ T-cells, FOXP3+ T-cells, CD163+ (M2) macrophages, and CD68+ macrophages than the "immune-silent" class. The "immune-suppressive" class of SMZL also had higher intra-tumoral PD1+ cells, and PD-L1+ cells, than "immune-silent" SMZL. PD1 expression co-localized with T-follicular helper cells in tumors containing germinal centers and with tumor-infiltrating T-cells in areas devoid of germinal centers. PD-L1 expression co-localized with tumor-infiltrating macrophages (Figure 5D). The immune-microenvironment classes were evenly distributed between cases with typical and atypical bone marrow morphology (Supplementary Table 11).

The immune microenvironment classes of SMZL were equally distributed across the molecular clusters (Figure 4B). Taken together these data indicated that in SMZL, such as in DLBCL, microenvironment signatures provide additional information that is not fully captured by a multi-gene mutational signature²³. In DLBCL, individual gene mutations rather than molecular signatures provided mechanistic insights into the interaction between lymphoma cells and the immune microenvironment⁴²⁻⁴⁷. To understand at a more granular level the mechanisms by which SMZL might induce specific microenvironment patterning, we comparatively assessed the spectra of the most recurrent mutations between the "immune-suppressive" vs the "immune-silent" classes of SMZL. *KLF2* mutations were enriched in the "immune-suppressive" class of SMZL and depleted in the 'immune-silent' class (Figure 4C). The tumor mutational burden, that is an approximation for neoantigen load and corresponds to the number of non-synonymous mutations per

coding area of the target region, was also higher in the "immune-suppressive" vs the "immune-silent" classes of SMZL (1.012 mutation/Mb vs 0.826mutation/Mb, p= 0.019).

Clinical course of the SMZL clusters

Baseline clinical features distributed evenly across molecular clusters and immune microenvironment classes of SMZL (Supplementary Table 15). To understand whether any molecular cluster or immune microenvironment class of SMZL was more aggressive than the others, we correlated these variables with relative survival. Lymphoma-specific deaths accounts for less than one-half of total deaths in SMZL3. Accordingly, relative survival provides a more accurate measure of excess mortality experienced by patients than overall survival, without requiring cause of death information. When the demographic effects of age, sex, and year of diagnosis were compensated, the 10-year life expectancies of patients with NNK-, DMT- and "immune-suppressive"-SMZLs were 79.0% 85.5% and 79.6% of those expected in the matched general population (Figure 6 and Supplementary Figure 4). Conversely, the 10-year life expectancy of patients with CBS-, PA- or 'immunesilent"-SMZL was not significantly lower than that expected in the matched general population. The combination of molecular and phenotypic profiling allowed us to sort out a high-risk clinical subset whose lymphoma was characterized by having both the NNK genotype and "immune-suppressive" microenvironment, and associated with an excess mortality (10 years relative survival: 70.8%) compared to the matched general population (Figure 6 and Supplementary Figure 6).

Among baseline clinical biomarkers that are known to be associated with SMZL patients's outcome, only low Hb and albumin significantly impacted relative survival (Supplementary Figure 7). Of note, neither low Hb, nor low albumin associated with any molecular or microenvironmental SMZL subtypes, making it unlikely that they can act as confounders in the inferior relative survival of NNK and immune-suppressive SMZL (Supplementary Table 15).

DISCUSSION

Our study highlights the complexity of SMZL, which comprises four distinct genetically defined molecular clusters, and two distinct phenotypically defined immune-microenvironment classes. The molecular framework for SMZL that we present here provides an evolving understanding of its pathogenesis, and can be regarded as a building block for further refining the classification of small B-cell lymphoproliferative diseases involving the spleen.

NNK and DMT clusters together contain the vast majority of SMZLs. NNK-SMZLs differ from DMT-SMZLs genetically, phenotypically, and clinically. The NNK cluster accounts for ~60% of SMZLs and is characterized by genetic changes that have been previously associated with the disease, such as *NOTCH2* mutations, *KLF2* mutations, 7q deletion and IGHV1-2*4 usage^{5,6,18,19}. The DMT cluster accounts for ~30% of SMZLs and is characterized by mutations in genes that are broadly affected in different B-cell lymphoproliferative disorders, such as *TP53*, *ATM*, *BRAF* and *MYD88*. The enrichment of nuclear expression of NOTCH and NF-kB transcription factors and of their gene expression signatures in NKK-SMZLs, and the complementary enrichment of signatures marking dysfunction of TP53 and apoptosis in DMT-SMZLs support the biological validity of these molecular clusters. From a clinical perspective, NNK- and DMT-SMZLs show different outcomes.

The existence of NNK- and DMT-SMZLs was externally validated in independent cohorts and by genetically modified mouse models of SMZL. The robustness of the NNK and DMT molecular clusters is supported by their identification in the Oquendo et al.²² meta-dataset of SMZL genotypes, despite its heterogeneous source of tumor DNA (eg, blood, marrow, spleen), sequencing strategy, and variant calling pipeline. *In vivo* murine models mimicking SMZL confirm that *Notch2/Nf-κB* activation and *Tp53* disruption in committed/mature B-cells are distinct and sufficient mechanisms of neoplastic transformation^{48,49}. Therefore, our results strongly support the conclusion that SMZLs exist as at least two molecularly distinct subtypes that are resolvable by genomic analysis. The CBS and PA molecular clusters account for a minority of SMZL (~10%) and warrant further validations.

Splenectomy was the first line therapy in all patients of the IELSG46 cohort. Currently, splenectomy is less frequently used as first-line therapy in SMZL, and has progressively been replaced by rituximab³. Evaluation of the relationship between SMZL genetic subtypes and outcome in additional cohorts and in the setting of rituximab-based treatment will be important to confirm and extend these findings. Lymphoma treatment is an evolving field and a number of pathway inhibitors have recently been approved^{50,51}. The results of our study nominate genetic subtypes of SMZL where pathway inhibitors could be tested. For example, drugs that target B-cell receptor-dependent NF-kB activation (eg, BTK inhibitors) could be investigated in NNK-SMZLs, drugs target the PI3K/AKT/mTOR pathway (eg, PI3K inhibitors) could be investigated in PA-SMZLs, while drugs that activate the apoptotic machinery when the DNA damage response pathway is disrupted (eg, BCL2 inhibitors) could be investigated in DMT-SMZLs.

Gene expression signatures and in situ profiling segregate two basic types of SMZLs immune microenvironments with distinct mutational and clinical connotations. The "immune-suppressive" class of SMZL accounts for ~50% of cases, strongly correlates with tumor-infiltrating lymphocytes and macrophage recruitment, as well as with immune checkpoint activation, ultimately suggesting the potential role of checkpoint-inhibitors in this subset of SMZLs¹⁴. The "immune-suppressive" class of SMZLs mimics the signatures of the "host response" class of diffuse large B-cell lymphoma (DLBCL), which notably expresses high levels of molecules also observed in SMZL⁵², and frequently involves the spleen. This observation establishes a new link between SMZL and DLBCL in addition to the recently discovered BN2/C1 genotype of DLBCL^{23,53}, which shares *NOTCH2* mutations with SMZL.

We highlight the enrichment of *KLF2* inactivating mutations within the "immune-suppressive" class of SMZL. As *KLF2* is a transcription factor involved in regulating several immune responses, it is likely that a constitutive dysregulated gene-expression/epigenetic program drives immune-evasion in *KLF2* mutated SMZLs. Deregulated transcription of lymphoma cells has already been associated with modifications of the tumor- microenvironment interactions resulting in immune cell recruitment, immune cell reprogramming, and immune evasion in other lymphoma types^{42,45,46}. The precise mechanisms linking *KLF2* mutations with reshaping of the immune microenvironment remain to be established.

Gene expression and immune cell topography indicate that "immune-silent" SMZLs can be categorized as a "cold" tumor. In this setting, therapeutic strategies harnessing the killing power of T-cells in a TCR independent manner and redirecting T-cells into close proximity of target cells to form a cytolytic synapse (eg, bispecific T-cell engagers or CAR T-cells) could be investigated.

Our study has some limitations. It lacks data on gene translocations and the extent of assessable CNA is restricted to the target regions of the sequencing assay. Such biases are however largely mitigated. First, our gene panel covers all the genomic regions implicated in recurrent CNA that were previously reported in SNP array studies of SMZL, including 1q, 3q, 6q, 7q, 8p, 12q, 17p, and 18q⁵⁴. Second, contrary to DLBCL, SMZL has low genomic complexity and lacks recurrent translocations^{6,55,56}. Third, no novel recurrent fusions were detected by RNA-seq. Fourth, mutations in *MYC*, *BCL2* and *BCL6*, which are a surrogate of their translocations with an immunoglobulin gene partner,²³ were extremely rare in the cohort, consistent with the knowledge that such structural variants are virtually absent in SMZL^{55,56}.

Our study was entirely based on spleen tissue samples. Currently, SMZL can be diagnosed without the need for splenectomy by integrating bone marrow histology with cell morphology and immunophenotype in the blood and bone marrow⁵⁷. This notion prompts the question on as to how can we identify the molecular clusters and microenvironmental classes of SMZL with a minimally invasive approach (i.e. without splenectomy). The almost universal dissemination of lymphoma cells in the blood allows the use of "liquid biopsy" approaches for characterizing the genetics of SMZL. The validity of this approach is confirmed by the evidence that the molecular profile of circulating SMZL cells matches that of cells residing in the spleen⁶. Efforts to profile tumor-infiltrating immune cells in SMZL have inherent limitations if splenectomy is not performed for diagnostic or therapeutic purposes. The comparison of paired bone marrow and spleen histology may inform whether bone marrow biopsy can be used to classify SMZL according to the microenvironmental classes. Multi-parametric phenotyping of immune cells led to new lines of inquiry and

assessment of the tumor microenvironment from peripheral blood samples^{58,59}. The abundance of tumor cells in SMZL in both blood and marrow also provides an opportunity of profiling at the same time both sides of the tumor-immune cell interactome.

In summary, our multiplatform genomic analysis elucidates SMZL pathogenesis, and provides a conceptual edifice to advance the classification and development of precision therapies for SMZL.

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Authorship

Contribution: F.B. and L.T.D.B performed bioinformatics analysis, interpreted data, contributed to data interpretation and manuscript preparation; A.B. performed molecular studies and bioinformatics analysis and contributed to data interpretation and manuscript preparation; F.G. performed molecular studies and contributed to data interpretation and manuscript preparation; M.F., R.K., A.R.E. and A. Calcinotto performed animal experiments, interpreted data and contributed to manuscript revision; V.S., G.F., D.P., K.P. and A.A. performed molecular studies, contributed to data interpretation and manuscript revision; A.C., L. Ceriani, F. Bertoni and F.C. contributed to data interpretation and manuscript revision; L.B. contributed to study management and manuscript revision; M.C.P., M.G.C., G.G. and W.W. contributed to medical data management, medical statistics, data interpretation and manuscript revision; M.L., G. Bhagat, E.C., L.D.L., S. Dirnhofer, S.A.P., M.A.P., A.T.G., A. Tzankov, M.P., M. Ponzoni and L.M., provided study material, performed pathological revision and contributed to manuscript revision; L.A., M.J.B., G.B., S.B., R.B., A. Broccoli, M.M.B., V.C., S.C., P.C., E.D., L.D., S.D., F.F., G. Gaidano, J.F.G., B.G., P.G., M.G.D.S., G. Gritti, A.G., F.H., G.I., M. Ladetto, A.L.G., E.L., A.M., R.M., E.M., V.M., M.M., A. Moccia, M. Mollejo, C.M., U.N., D.G.O., F.P., F. Piazza, S.P.,

A.R., E.S., G.S., E. Santambrogio, L.S., A.S., G. Stüssi, J.T.G., G.T., C.T., C. Thieblemont, T.T., A.T., G.V., C.V., U.V., R.W., F.Z., T.Z. and P.L.Z provided study material and clinical data and contributed to manuscript revision; L.C. and H.K. performed bioinformatics analysis, contributed to data interpretation and manuscript preparation; E.Z. provided key scientific insights and contributed to data interpretation and manuscript revision; D.R. designed the study, interpreted data, and wrote the manuscript.

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FIGURE LEGENDS

Figure 1. Mutation landscape of SMZL. (A) The heatmap shows genes mutated in ≥1% of SMZL cases. In the heatmap, each row represents a gene and each column represents a SMZL case. The heatmap is manually clustered to emphasize mutational co-occurrence. The number and type of somatic mutations in any given gene is plotted in the histogram on the right of the heatmap. On the top of the heatmap are shown the cytogenetic and immunogenetic features for each SMZL sample. (B) The heatmap shows pairwise Pearson correlation coefficients between each pair of genes mutated in SMZL. Red cells indicate positive correlation, blue cells indicate negative correlation. (C) Pathways that are recurrently mutated in newly diagnosed SMZL. In the heatmap, rows correspond to genes and columns represent individual patients. Color coding is based on gene alteration status (gray, wild type; red, mutated). The heat map was manually clustered to emphasize mutational co-occurrence.

Figure 2. Pathway-driven clustering of SMZL mutations. (A) Heatmap showing SMZL mutations collapsed into pathways and clustered using hierarchical clustering on principal components (HCPC) in the SMZL cohort. (B) Heatmap showing genes entering the clustering analysis. The genes are color coded according to the assigned cluster and ordered according to the relative enrichment in the specific cluster represented as a bar on the right. Only genes with mutation rate > 0.01 are shown). (C) The heatmap shows the enrichment of IGHV mutational status and recurrent CNA in the molecular clusters. Size and color intensities of the circles are

proportional to the prevalence of each variable in the molecular cluster. The asterisk shows statistical significant differences (Fisher's exact test p <0.05)

Figure 3. Biological attributes of the molecular clusters. (A) Barplot showing differential gene expression signature enrichment between NNK and DMT cluster. Statistical significance was determined using pairwise Student's t-test on scaled activity score values and was <0.05 for all the genesets. (NF-κB noncanonical activated, p=0.004; NOTCH2 signaling, p=0.043; Apoptosis, p=0.009; p53 signaling, p=0.028; B-cell proliferation, p=0.014; IL6/JAK/STAT3 signaling, p=0.003. (B) The heatmap shows the distribution of TMA markers in NNK and DMT molecular clusters. A red color indicates the positivity of the marker, a white color indicates negativity of the markers, and grey color indicates missing data. The barplot on the left shows the -log10 p-value of a one-sided Fisher's Exact test between the two clusters for each marker. (C) TP53, NOTCH2 and p50 expression in TMA cores from SMZL spleen sections.

Figure 4. Resolution of the tumor microenvironment by gene expression analysis. (A) Heatmap of scaled activity score values related to microenvironment signatures. HCPC clustering reveals two major microenvironment classes ("immune-suppressive" and "immune-silent") in the SMZL cohort. The differential expression of the microenvironment signatures is also represented as density plots on the right.

(B) Alluvial plot showing samples distribution among microenvironment classes and mutational clusters. (C) The heatmap shows the percentages patients mutated for the top mutated genes in the two microenvironment classes. Size and color intensities of the circles are proportional to the prevalence of each variable in the

microenvironment class. The asterisk shows statistical significant differences (Fisher's exact test p < 0.05)

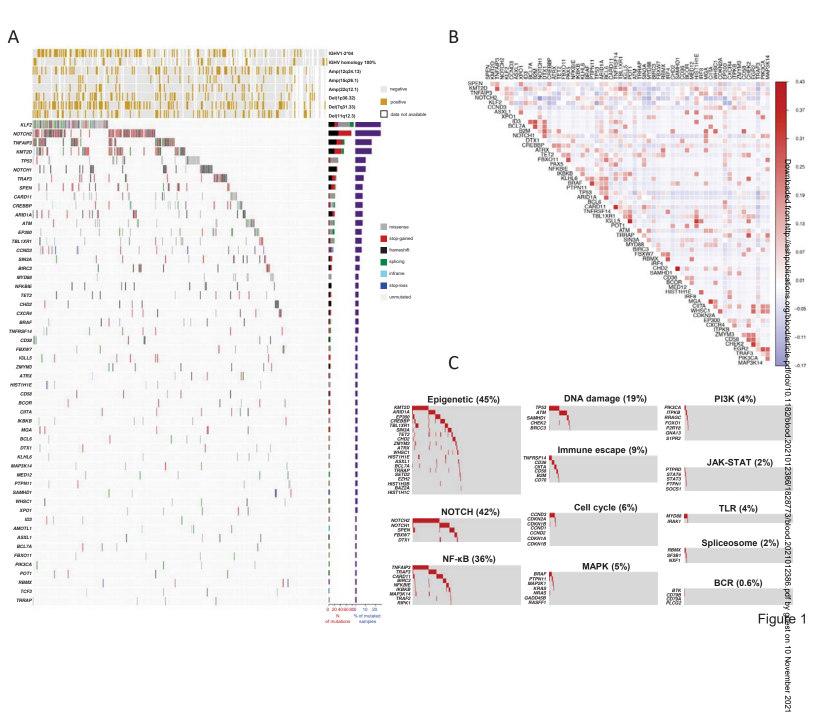
Figure 5. Orthogonal validations of the microenvironment composition. (A) Digital cytometry experiments. Gene expression data was used to infer the cell composition (covering 22 hematopoietic cellular populations) of spleen tissues. Staked barplots represent the individual cell compositions (scaled to a total of 100%). Samples (on the x axis) are ordered according to the two major microenvironment classes. (B) The heatmap shows the distribution of selected TMA markers across the two microenvironment classes. A red color indicates the positivity of the marker, a white color indicates negativity of the markers, and grey color indicates missing data. The barplot on the left shows the -log10 p-value of a one-sided Fisher's Exact test between the two classes for each marker. (C) CD3 (T-cell), CD163 (M2 macrophages), CD21 (FDC), PD1, PDL1 expression in TMA cores from SMZL spleen section. (D) The heatmap shows pairwise Pearson correlation coefficients between TMA markers. Red cells indicate negative correlation, blue cells indicate positive correlation.

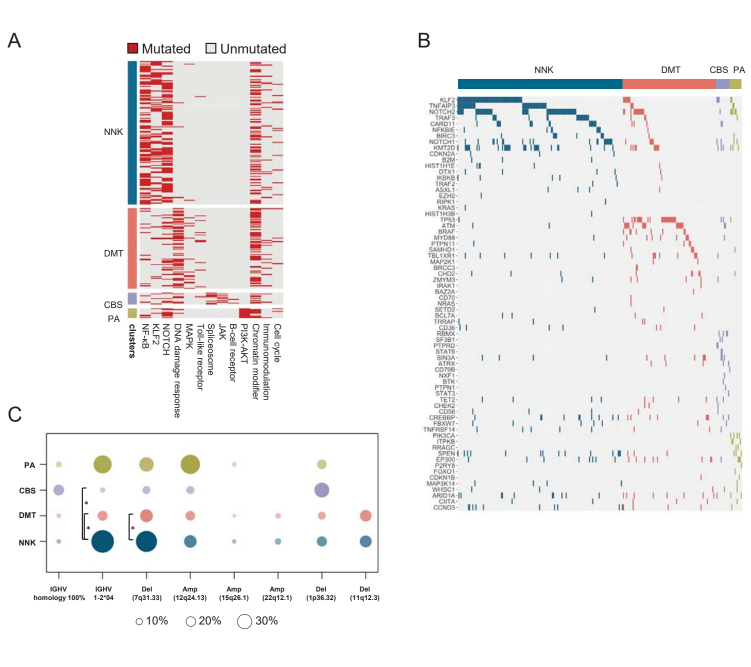
Figure 6. Relative survival of SMZL patients. Relative survival rates are presented for the two major mutational clusters (A), for the two major microenvironment classes (B) and for all their combinations (C).

| Table 1. Patients' characteristics | |
|---|--|
| Female:male ratio | 55.1:44.9 |
| Median Age | 64 (IQR=17) |
| Period of diagnosis | |
| - < 1990 | 1 (0.3%) |
| - 1990-1994 | 11 (3.6%) |
| - 1995-1999 | 32 (10.6%) |
| - 2000-2004 | 75 (24.8%) |
| - 2005-2009 | 142 (46.9%) |
| - 2010-2014 | 32 (10.6%) |
| - >2015 | 10 (3.3%) |
| ECOG scale | (5.573) |
| - 0-1 | 191 (63.0%) |
| - >1 | 16 (5.3%) |
| - missing | 96 (31.7%) |
| Ann Arbor Stage | 30 (01.170) |
| - | 0 |
| · | |
| - | 0 |
| - III | 0 |
| - IV | 272 (89.8%) |
| - missing | 31 (10.2%) |
| B symptoms | |
| - Yes | 54 (17.8%) |
| - No | 181 (59.7%) |
| - Missing | 68 (22.5%) |
| Extra hilar lymph node | |
| - Yes | 61 (20.1%) |
| - No | 156 (51.5%) |
| - Missing | 86 (28.4%) |
| Median hemoglobin | 11.6 g/dl (IQR=2.9) |
| - HPLL (< 9.5 g/dl) | 41 (13.5%) |
| - ILL (< 12 g/dl) | 142 (46.9%) |
| - Low (< 10 g/dl) | 56 (18.5%) |
| | |
| - Missing | 54 (17.8%) |
| Median platelets | 125 ×10 ⁹ /I (IQR=70) |
| - Low ($< 80 \times 10^9 / I$) | 33 (10.9%) |
| - Missing | 62 (20.5%) |
| Median lymphocytes | $3.3 \times 10^9 / I (IQR = 7.65)$ |
| - High (> 5.0 ×10 ⁹ /l) | 84 (27.7%) |
| - Normal | 135 (44.6) |
| - Missing | 84 (27.7%) |
| Median LDH | 387.5 U/I (IQR=249) |
| - High (> ULN) | 87 (28.7%) |
| - Normal | 113 (37.3%) |
| - Missing | 103 (34.0%) |
| Median albumin | 40 g/l (IQR=6.0) |
| - Low (< 35 g/l) | 23 (7.6%) |
| - Normal | 131 (43.2%) |
| - Missing | 149 (49.2%) |
| Median β2-microglobulin | 3.6 mg/l (IQR=2.07) |
| | |
| - High (> 2.5 mg/l) | 96 (31.7%) 34 (41.3%) |
| - Normal | 34 (11.2%) |
| - Missing | 173 (57.1%) |
| HCV | 10 (1 00() |
| - Positive | 12 (4.0%) |
| - Negative | 175 (57.7%) |
| - Missing | 116 (38.3%) |
| Serum monoclonal component | |
| - Yes | 61 (20.1%) |
| - IgG | <mark>21(34.5%)</mark> |
| - Ig <mark>M</mark> | <mark>39 (63.9%)</mark> |
| - Missing | 1 (1.6%) |
| - No | 138 (45.6%) |
| - Missing | 104 (34.3%) |
| LIDIT The threehold by the Colonia Marriagi Zana Lymanhama Chydy Cray | (LIDLL) access III. Like three held by the later room a Italiana Linforni (III.) |

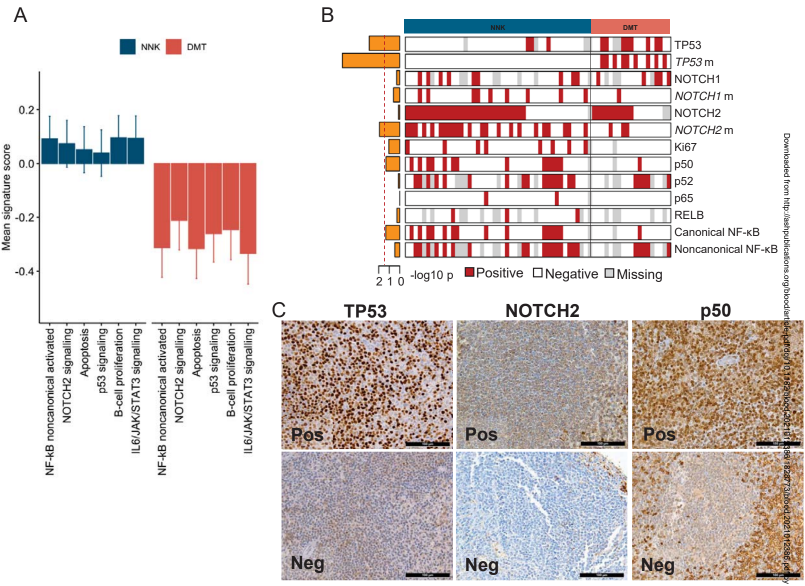
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HPLL, Hb threshold by the Splenic Marginal Zone Lymphoma Study Group (HPLL) score; IIL, Hb threshold by the Interguppo Italiano Linfomi (IIL) score; Extra hilar lymph nodes were regional in all cases and affected the celiac region.





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Figures on 10 November 2021

