Targeting of Inflammatory Pathways with R2CHOP in High-Risk DLBCL

Keenan T. Hartert¹, Kerstin Wenzl¹, Jordan E. Krull¹, Michelle Manske¹, Vivekananda Sarangi², Yan Asmann³, Melissa C. Larson², Matthew J. Maurer², Susan Slager², William R. Macon⁴, Rebecca L. King⁴, Andrew L. Feldman⁴, Anita K. Gandhi⁵, Brian K. Link⁶, Thomas M. Habermann¹, Zhi-Zhang Yang¹, Stephen M. Ansell¹, James R. Cerhan², Thomas E. Witzig¹,*, Grzegorz S. Nowakowski¹,*, Anne J. Novak¹,*

¹Division of Hematology, Mayo Clinic, Rochester, MN
²Department of Health Sciences Research, Mayo Clinic, Rochester, MN
³Department of Health Sciences Research, Mayo Clinic, Jacksonville, FL
⁴Division of Hematopathology, Mayo Clinic, Rochester, MN
⁵Celgene, Summit, NJ
⁶Division of Hematology, Oncology, and Bone & Marrow Transplantation, University of Iowa, Iowa City, IA

Abstract

Diffuse large B-cell lymphoma (DLBCL) is the most common lymphoma, and front line therapies have not improved overall outcomes since the advent of immunochemotherapy. By pairing DNA and gene expression data with clinical response data, we identified a high-risk subset of non-GCB DLBCL patients characterized by genomic alterations and expression signatures capable of sustaining an inflammatory environment. These mutational alterations (PIM1, SPEN, and MYD88[L265P]) and expression signatures (NF-κB, IRF4, and JAK-STAT engagement) were associated with proliferative signaling and were found to be enriched in patients treated with RCHOP that experienced unfavorable outcomes. However, patients with these high-risk mutations had more favorable outcomes when the immunomodulatory agent lenalidomide was added to RCHOP.
We are the first to report the genomic validation of a high-risk phenotype with a preferential response towards R2CHOP therapy in non-GCB DLBCL patients. These conclusions could be translated to a clinical setting to identify the approximately 38% of non-GCB patients that could be considered high-risk and would benefit from alternative therapies to standard RCHOP based on personalized genomic data.

Introduction

As the most common form of aggressive non-Hodgkin lymphoma (NHL) afflicting nearly 30,000 patients in the USA each year, diffuse large B-cell lymphoma (DLBCL) represents a significant challenge in hematology/oncology. New DLBCL treatments remain a clinical need despite the success of rituximab combined with CHOP chemotherapy (cyclophosphamide, doxorubicin, vincristine, and prednisone), RCHOP, which results in durable responses in 60-70% of patients. Those refractory to, or who relapse following, first-line therapy have a very poor outcome, with only 20% surviving beyond 5 years despite second-line therapies. Rationally-targeted frontline strategies are needed, especially for those with high-risk disease or early clinical failure. Herein, we present the case for precision targeting of a genetically-distinct population of RCHOP-insensitive DLBCL patients with targeted therapy.

Since 2002, DLBCL has traditionally been divided into two cell of origin (COO) subcategories based on tumor gene expression profiles (GEP): Activated B-cell (ABC) and Germinal Center B-cell (GCB). Patients with ABC tumors are characterized by a more aggressive profile and active NF-κB and BCR signaling pathways while GCB cases are associated with alterations that drive aberrant chromatin-modification, PI3K signaling, and the upregulation of MYC and BCL2 through translocations or copy number gains. Recently, using tumor samples from patients treated with RCHOP, new classification models have focused on DNA alterations, including previously-identified drivers of aggressive disease such as MYD88 (L265P), CARD11, and TNFAIP3. MYD88 L265 mutations have specifically been shown to be enriched in aggressive ABC DLBCL cases. Key downstream effects of oncogenic MYD88 include activation of the NF-κB, JAK/STAT, and upregulation of inflammatory cytokines, often augmented by the loss of the inhibitory TNFAIP3 gene. The NOTCH signaling pathway has also been characterized in aggressive cases of DLBCL. To date, none of these studies have impacted the design of phase III trials for untreated DLBCL patients; rather, these trials have focused on selecting and randomizing patients based on the International Prognostic Index (IPI) and tumor GEP. However, this strategy has failed to show benefit of adding novel agents to RCHOP based on these classifiers, and higher intensity therapies such as DA-EPOCH, ibrutinib, idelalisib, obinutuzumab, and bortezomib have all failed to improve outcomes in clinical trials. The field is now focused on designing trials based on personal tumor signatures.

Building on new genetic profiling studies to personalize clinical treatment could allow clinicians to add targeted therapies to the RCHOP backbone based on individual tumor signatures. Wilson et al. showcased this methodology of precision medicine by identifying that the driver mutations of a BCR-driven, non-GCB tumor (CD79A, CD79B) can be
successfully treated with ibrutinib as a single agent and RCHOP + Ibrutinib (in patients younger than 60) when they would have likely failed standard RCHOP treatment.\textsuperscript{21, 27, 28}

Aiolos/Ikaros-degrading immunomodulatory drugs (IMiDs\textsuperscript{®}) such as thalidomide, lenalidomide, and pomalidomide play a pivotal role in the treatment of multiple myeloma.\textsuperscript{29-31} More recently, a role for lenalidomide has been reported in phase II studies for the treatment of aggressive DLBCL.\textsuperscript{32-34} The results of two clinical trials comparing RCHOP combined with lenalidomide (R2CHOP) versus RCHOP alone were recently reported.\textsuperscript{35, 36} The phase III ROBUST trial was restricted to ABC-type DLBCL and failed to show significant clinical benefit of R2CHOP, but the phase II ECOG-ACRIN1412 trial was open to patients of all COO subtypes and showed significantly superior event free and overall survival benefits for those treated with R2CHOP. Early research highlighted the ability of lenalidomide to exploit synthetic lethality in ABC cell lines by deregulating oncogenic programs.\textsuperscript{37} Lenalidomide binds Cereblon resulting in rapid degradation in transcriptional repressors Aiolos and Ikaros which leads to upregulation of interferon-stimulated genes such as IRF7 and apoptosis in B-cells, particularly those associated with the ABC-DLBCL subtype.\textsuperscript{38} In T-cells, the result is activation and IL-2 production.\textsuperscript{39} These immunomodulatory effects made lenalidomide a prime candidate for treating aggressive cases of DLBCL, especially those with an inflammatory microenvironment. Herein, we report the profile of a high-risk ABC/non-GCB subset of DLBCL driven by genomic alterations in inflammatory genes that are susceptible to front-line treatment R2CHOP but continue to experience poor outcome with RCHOP alone.\textsuperscript{33} These results showcase the success of personalized RCHOP + X therapy application based on the genetic signature and biological profile of patient tumors.

**Materials and Methods**

**Study Population**

A total of 196 patients with DLBCL were studied. Forty-seven patients with Ann Arbor stages II-IV were treated with R2CHOP from an investigator-initiated, open-label, single-arm phase II study (NCT00670358) were included in the study.\textsuperscript{33, 40} 149 newly diagnosed DLBCL cases treated with RCHOP, or R-immunochemotherapy (herein called RCHOP), and followed prospectively through the Molecular Epidemiology Resource (MER) of the University of Iowa/Mayo Clinic Lymphoma Specialized Program of Research Excellence (SPORE) were used as a matched contemporary cohort. Full details of this prospective cohort study of lymphoma outcomes have been previously published.\textsuperscript{41} All patients provided written consent at enrollment into the clinical trial or MER for use of their clinical samples. Disease progression, relapse, unplanned re-treatment after initial immunochemotherapy, and death from any cause were verified through medical record review. Cell of origin (COO) was determined in the available R2CHOP samples by the Lymph2Cx assay (nanoString, N=45).\textsuperscript{42} For the MER RCHOP cases, COO was determined by GEP (N=36), nanoString (N=68), or Hans (N=35).\textsuperscript{43} For this study, DLBCL were categorized into GCB and non-GCB (ABC and Unclassified) groups. Baseline clinical characteristics of all patients in this study are shown in Table 1 and detailed clinical information for each cohort is provided in Supplemental Table 1.
Whole Exome Sequencing

For the identification of DNA alterations in the RCHOP and R2CHOP cases, whole exome sequencing (WES) was performed. RCHOP-treated cases (N=149) were sequenced as previously described\(^\text{19}\). R2CHOP-treated cases (N=47) were sequenced as follows: DNA was extracted from formalin-fixed, paraffin-embedded (FFPE) DLBCL tumors using the QiAamp DNA FFPE Tissue Kit (Qiagen GmbH, Hilden, Germany) in the Mayo Biospecimens Accessioning and Processing Core. Prior to isolation, tumor blocks were reviewed by a Mayo Clinic hematopathologist, tumor areas were circled, and four 1 mm cores were used for DNA isolation. The minimum tumor purity for study was 30%. WES was performed at the Mayo Clinic Genome Analysis Core. Sequencing was carried out on an Illumina HiSeq 2000 at a depth of ~100 million 100 bp paired-end reads per sample. Data from all cases were mapped to human genome reference build 38 using BWA-MEM. Quality control was performed by FASTQC (v0.11.3). After realignment and recalibration by GATK (v3.4-46), SNV and INDELs from individual germline and tumor samples were called by GATK haplotype caller (v3.4-46). Variants were annotated using the Mayo Clinic in-house annotation tool BioR. Sample inclusion required a minimum of 20x coverage. Mutation data was visualized as a waterfall plot using the “complex heatmaps” R program tool\(^\text{44}\). Copy number alterations were determined and annotated for available cases as performed in Wenzl et. al. but not included in the overall analysis as no significant differences were seen between the RCHOP and R2CHOP samples. All WES data is available upon request.

Gene Expression Analysis

RNA from non-GCB RCHOP (N=104) and R2CHOP (N=42) treated tumors were assayed with the nCounter® PanCancer Pathways Panel with 730 cancer pathways genes (nanoString, Seattle WA), and analyzed with GenePattern tools from the Broad institute, including GENE-E, Comparative Marker Analysis, and Gene Neighbor analysis\(^\text{45, 46}\). GEP was normalized using standardized NanoString protocols (Positive Control Normalization and CodeSet Content Normalization). RNA from non-GCB RCHOP (N=45) and R2CHOP (N=14) treated tumors was analyzed in a second analysis.

Statistical Analysis

Graphpad Prism software and GenePattern tools were used to plot and format figures, analyze data, and calculate statistical significance\(^\text{46}\). Comparison of quantitative data between groups was done by Student’s t-test or one-way ANOVA test. Gene ontology analyses and protein-protein interactomes were generated using ToppGene Suite tools\(^\text{47}\). EFS was defined as time from diagnosis to progression or relapse, unplanned re-treatment after initial immunochemotherapy, or death from any cause. The primary clinical outcome metric used was event-free survival at 24 months (EFS24). Since most relapses in DLBCL patients occur in the first 24 months, EFS24 is an early predictor of long-term outcome\(^\text{7}\). RCHOP and R2CHOP populations were divided on the basis of EFS24 status to calculate favorability enrichment percentages for DNA alterations and T statistic metrics for gene expression among populations. Specifically, favorability enrichment percentages for DNA alterations were calculated as follows: (number of EFS24 achieved patients with a mutation/total number of EFS24 achieved patients) – (number of EFS24 failed patients with a
mutation/total number of EFS24 failed patients. A positive value indicates mutation is enriched in EFS24 achieved patients and a negative value indicates mutation is enriched in EFS24 failed patients. EFS between groups were compared using the Kaplan-Meier method and the log-rank test. Multiple clinical factors were analyzed by a Chi-square test in Graphpad Prism. All reported P values were two-sided. P values less than 0.05 were considered statistically significant.

Results

Specific DNA Mutations Predict EFS24 and R2CHOP Response in DLBCL

The high prevalence of MYD88 mutations combined with the promising clinical results of R2CHOP in ABC DLBCL suggest that there are underlying biologic and genomic differences that may correlate with tumor-specific responses. To identify a genomic signature for high-risk DLBCL that was lenalidomide-responsive, we first used WES data to determine the mutation profile of both R2CHOP and RCHOP cases and then looked for enrichment of variants in the R2CHOP responsive tumors. 47 patients with DLBCL that met eligibility criteria were enrolled in the R2CHOP cohort and were evaluated against 149 RCHOP comparison patients (Table 1). The outcome data are summarized for each treatment cohort in Table 2. The treatment cohorts did not significantly differ on the basis of either overall EFS or OS (overall survival) (Supplemental Figure 1). IPI and COO were significantly associated with outcome for the RCHOP population, but not for the R2CHOP population (Supplemental Figure 2). Mutations in known driver genes in both the R2CHOP and RCHOP cohorts, based on designation by one of the major DLBCL analyses are documented in Supplemental Figure 3 (N=211). We first identified genes that were mutated in both cohorts (N=186) and then calculated the presence of the alteration in populations that achieved or failed EFS24. Taking the difference between these values revealed whether R2CHOP treatment resulted in an increased association of a gene towards achieving EFS24. These results were plotted against each other in an XY scatter plot for all patients (Figure 1A; left panel). This analysis in all GCB and non-GCB patients together did not reveal any genes that met the criteria for R2CHOP outcome improvement (10% enrichment in RCHOP EFS24 failure and 10% R2CHOP achievement), although SPEN, PIM, and MYD88 trended towards association. In a secondary analysis of non-GCB cases only, mutations in genes (N=60) were explored due to the specific benefits of R2CHOP seen in these patients (Figure 1A; right panel) (Supplemental Table 2). Three genes were enriched in the RCHOP cases that failed EFS24 but achieved EFS24 with R2CHOP: PIM1, SPEN, and MYD88 (L265P) (Supplemental Figure 4; Table 3). PIM1 and MYD88 mutations have previously been observed to occur together, but SPEN mutations were observed to be almost entirely independent from both. A separate analysis of GCB DLBCL was also performed and no genes were associated with and EFS24 R2CHOP response. A heatmap showing differential enrichment of genes associated with non-GCB R2CHOP EFS24 response is shown in Figure 1B. Together, PIM1, SPEN, or MYD88 (L265P) mutations were present in 38.0% of all non-GCB cases (N=30/79) and are herein referred to as R2CHOP EFS24 responder alterations (RA).
We next analyzed the overall EFS of R2CHOP and RCHOP treated non-GCB cases with or without RA. In the R2CHOP cases, patients with a RA had a better overall EFS (P = 0.051) compared to wild type patients (no RA) (Figure 1C, left panel). In contrast, RCHOP treated non-GCB patients with a RA had a significantly worse overall EFS (P = 0.0004) compared to patients without a RA (Figure 1C right panel). Patients incurring a single RA were not significantly different than those incurring multiple RA, suggesting that a mutation in any one of the RA genes may predict response to R2CHOP (Supplemental Figure 5). In a secondary analysis, R2CHOP and RCHOP samples were clustered based on criteria from Chapuy et. al (Supplemental Table 2). While the sample size was small, these data suggest that the C1 and C5 subtypes were responsive to R2CHOP (Supplemental Figure 6). This is most likely due to the fact that C1 is enriched for SPEN mutations and C5 is enriched for MYD88 and PIM1 mutations.

Gene Ontology Analysis Reveals Unique Signature of R2CHOP Response Program

To explore the genetic programs and pathways that were susceptible to R2CHOP, gene ontology analysis was performed. Of the 60 driver genes with mutations in both RCHOP and R2CHOP non-GCB patient cohorts, 46 were more associated with achieving EFS24 in patients treated with R2CHOP. Gene ontology analysis revealed the top 20 cellular pathways associated with these genes (Figure 2A)\cite{12,18}. Specific genetic programs associated with the highlighted pathways are shown in Figure 2B. The cases achieving EFS24 were enriched for mutations in genes associated with chromatin modification, cytokine production, IRF4, TLR signaling, IFNG signaling, and the NOTCH and NF-κB pathways (Supplemental Table 3). These results provide insights towards high-risk pathways activated in DLBCL that are vulnerable to the addition of IMiDs.

Genetic Expression Programs with Specific Favorable Response to R2CHOP Identified

We next analyzed gene expression data from the PanCan panel of 730 B-cell-related genes to determine gene expression profiles characteristic of the high-risk/R2CHOP-profile. Gene expression data was available on 59 non-GCB DLBCL cases (45 RCHOP; 14 R2CHOP). A two-sided comparative marker analysis T statistic test was applied to assess what genes displayed differential expression based on achieving EFS24 in both populations (Supplemental Table 4)\cite{45}. Each gene was assigned a T statistic value based on this analysis. Positive values indicated greater RNA expression in the patient population that achieved EFS24, and negative values indicated greater RNA expression in the patient population that failed EFS24. The collective R2CHOP and RCHOP EFS24 T values were significantly different for 18 previously-defined gene expression signatures (Figure 3A). Non-GCB patients treated with R2CHOP had superior EFS24 when expressing genes associated with high MYD88, NF-κB, STAT3, JAK, IRF4, and OCT2 induction. The ABC and proliferation signatures were also associated with achieving EFS24 when treated with R2CHOP instead of RCHOP. IRF4 dysregulation has been previously highlighted as a mechanism of lenalidomide response\cite{37}. In contrast, cases that were associated with EFS24 failure when treated with R2CHOP expressed genes associated with the Stromal-2, lenalidomide-repressed, CNS, HRAS, TGFB, E2F3, JAK-downregulated, and MYC upregulation signatures.
To visualize gene expression relationships, normalized T statistics for R2CHOP and RCHOP were plotted against one another in an XY scatter plot. The difference between the two was used to assess whether a gene was more associated with achieving EFS24 if treated with R2CHOP. In total, 113 genes were identified; the top three were MAP3K14, IL2RB, and STAT3 on the basis of T statistic differential (Figure 3B). Other notable genes include JAK1, SOS1, and IL6R. The 113 gene set was next used to assess gene-gene interaction enrichment. The top 50 partners of this network are shown in Figure 3C. The top and bottom 5% (37/730) of genes were isolated for gene ontology analysis. The R2CHOP cases that achieved EFS24 were enriched for genes involved in cell cycle, IL6, JAK-STAT, IL2, and STAT3 pathways (Figure 3D). Of note, patients who failed R2CHOP were enriched for genes involved in PI3K-AKT, RAS, MAPK, WNT, and EGFR signaling, pathways recently highlighted as mechanisms of lenalidomide resistance in multiple myeloma. IL2RB and STAT3 transcript levels were significantly negatively correlated with the RAS signaling components RASGRP2 and HRAS (Supplemental Figure 7). Analysis of tumor microenvironment components through xCell RNA analysis revealed that a low CD8+ T-cell signature was associated with R2CHOP patients that failed EFS24 (Supplemental Figure 8). This aligns with lenalidomide’s mechanism of action involving T cell activation resulting in tumor cell cytotoxicity.

**Gene Expression Neighbor Analysis of R2CHOP Responders Reveal Mechanisms of Lenalidomide**

From the 113 genes that predicted differential EFS24 response between RCHOP and R2CHOP, 3 genes displayed the greatest associations with R2CHOP EFS24 achievement and RCHOP EFS24 failure: MAP3K14, IL2RB, and STAT3. To identify genes that were closely associated with these targets, Genetic Pearson Distance was calculated. Genes with a smaller (closer) distance to a target gene most closely match the expression patterns of that gene. Genetic distance was calculated and plotted against all 730 genes for MAP3K14, IL2RB, and STAT3 (Figure 4A). The MAP3K14 plot highlights its key role as a nonconical NF-κB activator, sharing matching expression profiles with genes such as SOS1, GSK3B, PIM1, and MYD88. Next, a protein-protein interaction network was generated using ToppGene suite tools. The top 50/359 protein-protein interaction partners between MAP3K14, IL2RB, and STAT3 based on K-step Markov prioritization are displayed (Figure 4B). However, the most specific correlative relationship emerged between IL2RB and STAT3. These two genes are so closely linked that IL2RB is the top expression neighbor to STAT3 (Figure 4C, left panel). This close relationship highlights several gene expression neighbors that are shared between IL2RB and STAT3. Genes within the top 10% of genetic distance of both are highlighted (N = 30) in the zoomed figure (Figure 4C, right panel). The MYD88 and JAK3 genes were also within the top 10% of MAP3K14 genetic distance. Other notable genes that shared expression patterns with both IL2RB and STAT3 included NOTCH2, IRAK3, JAK2, and IFNG.

**Responder Alterations are Associated with Specific Patterns of Gene Expression**

To integrate the genomic and transcriptomic data described above and identify lenalidomide sensitive mechanisms, the relationships between the RA (PIM1, SPEN, and MYD88) and the RNA pathway genes associated with R2CHOP EFS24 were explored. 15 of the 44 non-
GCB RCHOP patients with paired DNA and gene expression data had a RA. T statistic differential expression analyses were performed for all 730 PanCan genes in those cases with or without RA. The top 5% genes with overall increases or decreases in patients with individual RA are noted (Figure 5A). The top 5% genes with overall increases in patients with any of the RA are listed in Figure 5B. A graphic summarizes the signaling pathways associated with the RA and highlight a potential mechanism of lenalidomide action, including inhibition of the NOTCH, NF-κB, and JAK-STAT pathways.

Discussion

Non-GCB DLBCL cases have been linked with inferior rates of overall survival as a result of aggressive activation of survival and proliferation pathways. This includes constitutive activation of the NF-κB transcription factor family via dysregulation of genes such as MYD88 or NOTCH and results in a chronic inflammatory cytokine milieu. Past work has highlighted the potential benefit of the immunomodulatory drug IMiD® lenalidomide in this patient population, however much of this work has focused on either in vitro or clinical studies, with lack of insight on the impact of lenalidomide on patient derived samples. We have extended upon these observations by aligning clinical data with DNA and gene expression analysis from patients treated on trial with R2CHOP, and we are the first to identify a high-risk profile of non-GCB patients that benefit from the addition of lenalidomide to RCHOP. These results highlight the potential of precision medicine strategies in DLBCL to identify a vulnerable patient population, validate a genetic phenotype, and apply a personalized therapy that elicits a favorable response when standard therapy likely would not.

We report three gene alterations to be predictive of high-risk DLBCL that fail standard RCHOP yet are susceptible to R2CHOP: PIM1, SPEN, and MYD88 (L265P). Each have well-documented roles in non-GCB DLBCL, with MYD88 being supported by numerous studies. The high incidence of MYD88 mutations combined with the clinical results of R2CHOP in ABC DLBCL suggest that there are underlying biologic and genetic differences that may account for ABC-specific responses. Mutations in MYD88 have also been specifically linked with a supportive cytokine milieu that sustains an inflammatory JAK-STAT phenotype. Other work has highlighted the self-sustaining capabilities of active STAT3 signaling and its reliance on NF-κB-triggering mechanisms. PIM1 alterations and those that activate NOTCH signaling (SPEN) have also been documented to induce an inflammatory cytokine environment. In support of our finding, MYD88 and PIM1 alterations were clustered into the unfavorable C5 and MCD genomic subtypes and are associated with aggressive NF-κB signaling. SPEN has also been identified as an unfavorable marker in the non-GCB phenotype and is present in the C1 and BN2 clusters, which resemble a novel non-GCB phenotype more reliant on NOTCH and immune escape. Together these studies suggest that PIM1, SPEN, and MYD88 (L265P) are associated with high-risk non-GCB DLBCL and drive activation of inflammatory TLR signaling, NOTCH, and IRF4. While these are not known direct targets of lenalidomide, they are expressed in tumors that have vulnerabilities to lenalidomide.
Building on the genomic studies, our PanCan analysis was able to identify gene expression patterns that reflected the high-risk profile and R2CHOP susceptibility. Inflammatory pathways such as JAK-STAT, NF-κB, IRF4, MYD88, and OCT2 displayed significantly greater association with achieving EFS24 when treated with R2CHOP compared to RCHOP. High-risk patients are likely receiving benefit from lenalidomide’s interference with continuous loops of IRF4, NF-κB, and STAT transcription factor signaling. Conversely, signatures associated with R2CHOP failure included the Stromal-2 survival, lenalidomide-repressed genes, CNS lymphoma, E2F3, HRAS, TGFB, and MYC upregulation signatures. These pathways likely identify a subset of non-GCB patients that rely on a supportive tumor microenvironment (TME) and proliferative signaling.

Upon further analysis of the gene expression data we identified MAP3K14, IL2RB, and STAT3 as predictors of high-risk disease on the basis of EFS24. MAP3K14, also known as NIK, emerged as a key component of sustained noncanonical NF-κB signaling associated with MYD88, and equally interesting was the highly-correlated pair of IL2RB and STAT3 expression20, 58-60. This partnership has been previously observed, as IL2RB, IL6, and STAT3 displayed significantly responsive expression and were significantly associated with STAT3 ChIP-seq peaks54. JAK1, JAK2, JAK3, IL6R, SOS1, MYD88, NOTCH2, and IFNG were also expressed at high levels in unfavorable RCHOP cases22, 54-57, 61. Based on these results, IL2RB and STAT3 could prove to be useful predictive markers for high-risk disease and R2CHOP response. The JAK-STAT pathway and resulting cytokine signaling profile was specifically highlighted by the rrDLBCL study by Morin and colleagues, making this observation all the more fitting as a high-risk pathway20. Integrative analysis of the DNA and gene expression data revealed how each RA (PIM1, SPEN, and MYD88) uniquely influences an inflammatory phenotype. Several genes associated with NF-κB engagement displayed greater expression in patients with any of the alterations (CARD11, PLCG2, and JAK1), but the SPEN profile was uniquely associated with microenvironment and NOTCH genes, highlighting its presence in the novel C1 and BN2 genetic classifications.

In conclusion, our combined analysis of DNA and RNA across R2CHOP and RCHOP treatment cohorts identifies a high-risk non-GCB phenotype that is capable of sustaining JAK-STAT and NF-κB signaling, and is sensitive to R2CHOP. This phenotype encompasses approximately 38% of non-GCB patients, and the positive results of the ECOG-ACRIN1412 highlight the clinical success of R2CHOP. Our data supports the hypothesis that R2CHOP has activity in tumors reliant on IRF4, NF-κB, and STAT transcription factors, leading to a loss of proliferative feedback systems. Although promising and highly relevant due to the use of early phase clinical trial samples and a large comparison cohort, these conclusions require additional validation that can be done as R2CHOP trial samples with existing tissue and long term follow up become available. The RA signature generated in this work could retrospectively identify patients in larger studies (such as the ROBUST trial) that would most likely benefit from the combination. Combined with prior studies on RCHOP + ibrutinib21, 28, 62, the groundwork for a precision therapy approach in DLBCL in which DNA or RNA profiles can be used to identify patients early in treatment who may not benefit from the current standard of care, RCHOP, and who would benefit from the addition of lenalidomide or other targeted agents.
Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1: Identification of DNA Alterations that Predict High-risk non-GCB DLBCL that Respond to R2CHOP.

(A) Association of individual genes with EFS24 response to RCHOP (x-axis) and R2CHOP (y-axis) are compared on an XY scatter plot. The data from both GCB and non-GCB patients are plotted in the left panel; data from non-GCB patients only are plotted in the right panel. (B) Heatmap showing differential enrichment of alterations in non-GCB patients who fail EFS24 with RCHOP but achieve with R2CHOP. Scale shown represents percentage enriched in EFS24 population. Genes with 10% or lesser favorability in RCHOP and 10% or greater favorability in R2CHOP were designated as EFS24 responder alterations (RA). (C) Kaplan-Meier curves for event free survival of R2CHOP (N=18) and RCHOP (N=61) treated cases. 95% CI ranges are shown as dotted lines.
Figure 2: Pathway Enrichment of DLBCL Cases that Respond to R2CHOP.
(A) The top 20 pathway enrichments for genes associated with an improved R2CHOP EFS24 response are represented in a bar graph through $-\log_{10}$ false discovery rate (FDR) significance values. (B) An orbital diagram is divided into four quadrants representing ontology characteristics of R2CHOP favorable gene sets. Ontologies are labelled with their title and their $-\log_{10}$ FDR value. More significant FDR values approach the center of the orbital. The following thresholds designate $-\log_{10}$ FDR orbitals: outer orbit = 1.33 to 5, intermediate orbit = 5 to 10, and inner orbit = 10+. Size of the ontology is determined by a normalized threshold of relative genes in the hit list compared to the total number of input genes.
Figure 3: RNA Expression Analysis Reveals R2CHOP Response Pathways.
(A) Mean values of signature gene EFS24 association T statistics for each treatment are plotted as bar graphs for R2CHOP (red) and RCHOP (white) favorability. Positive T values represent greater gene expression in patients that achieved EFS24. Negative T values represent greater gene expression in patients that failed EFS24. Error bars represent the 95% confidence interval. Difference in means was observed with a student’s t test with significance achieved at $\alpha = 0.05$. (B) Individual genes are plotted in an XY scatter plot based on their EFS24 T value associations with RCHOP and R2CHOP. T statistics have been normalized to facilitate presentation. R2CHOP responder expressers are highlighted in red (N=113). (C) Top interacts of the 113 responder expressers are displayed by their significance. (D) Pathway enrichment analyses were performed for the top and bottom 5% of genes that displayed a shift in expression from poor EFS24 RCHOP cases to favorable EFS24 R2CHOP cases and vice versa. Ontologies composed of genes enriched for R2CHOP success (red) and failure (black) are documented on the basis of $-\log_{10}$ FDR value.
Figure 4: Nearest Neighbor Analysis of Top Responder Genes Reflect NF-κB, JAK/STAT, and Cytokine Signaling Programs.

(A) Plots illustrate ranked Pearson distance between responder genes and the other 729 PanCan genes. For each plot, genes to the left of the X-axis and designated in red represent genes with similar expression profiles as the target gene. Those with greater Pearson distance are plotted to the right and exhibit dissimilar expression patterns to the target gene.

(B) A bar graph highlights the top 50 protein-protein interaction partners between MAP3K14, IL2RB, and STAT3 based on k-step Markov distance. (C) A scatter plot documents the correlative genetic partners of IL2RB and STAT3 for all genes. Dotted lines designate limits for top 10% closest genes to each. 23/730 genes meet these criteria and are shaded in red. A zoomed view of these gene neighbors is displayed in the bottom plot. MYD88 and JAK3 are also top 10% gene partners with MAP3K14 and highlighted with a dark border.
Figure 5: Responder Alterations Correspond to Distinct RNA Expression Profiles.
Combined data from 44 cases that had paired WES and PanCan data was used for analysis, R2CHOP (N=13) RCHOP (N=31). 15 patients had at least one RA. (A) The top 5% and bottom 5% (N=37 each) of genes associated with the presence of each RA are visualized through dot plots. Genes with greater differential expression in the presence of an RA are closer to the top (greater T value; green) and genes with lesser expression in the presence of an RA are closer to the top (lower T value; red). Each RA is individually documented. (B) The table documents genes associated with greater differential expression in the presence of any RA. (C) A graphic summarizes the DNA alterations and the hypothesized high-risk phenotype. RA are designated green, genes associated with greater expression in yellow, and hypothesized cytokines in red.
| Characteristic                        | RCHOP (N=149) | R2CHOP (N=47) | P-Value |
|--------------------------------------|---------------|---------------|---------|
| Age median (range)                   | 64 (26-89)    | 61 (19-87)    | 0.15‡   |
| IQR                                  | 56-72         | 56-71         |         |
| Age ≥60                              | 96 (64.4%)    | 27 (57.4%)    | 0.39†   |
| Male                                 | 91 (61.1%)    | 29 (61.7%)    | 0.99†   |
| PS 2+                                | 21 (14.1%)    | 5 (10.6%)     | 0.63‡   |
| Ann arbor stage III-IV               | 93 (62.4%)    | 42 (89.4%)    | <0.0001‡|
| 2+ extranodal sites                  | 29 (19.5%)    | 14 (29.8%)    | 0.16‡   |
| LDH > ULN                            | 76 (58.0%)‡   | 29 (61.7%)    | 0.73‡   |
| IPI                                  |               |               |         |
| 0 - 1                                | 46 (30.9%)    | 16 (34.0%)    | 0.52‡   |
| 2                                    | 45 (30.2%)    | 12 (25.5%)    |         |
| 3                                    | 43 (28.9%)    | 11 (23.4%)    |         |
| 4 - 5                                | 15 (10.1%)    | 8 (17.0%)     |         |
| COO                                  |               |               |         |
| ABC                                  | 50 (33.6%)    | 14 (29.8%)    | 0.86‡   |
| GCB                                  | 78 (52.3%)    | 27 (57.4%)    |         |
| Unclassified                         | 11 (7.4%)     | 4 (8.5%)      |         |
| N-miss                               | 10 (6.7%)     | 2 (4.3%)      |         |
| EFS24 Achieved                       | 100 (67.1%)   | 34 (78.7%)    | 0.15‡   |

Abbreviations: R2CHOP, lenalidomide added to RCHOP; IQR, interquartile range; PS, performance score; IPI, international prognostic index; COO, cell of origin; EFS24, event-free survival over 24 months

* Unpaired t test
† Fischer’s exact test
‡ Chi square test
§ 18 cases without data
## Table 2

### Treatment Summary by COO

|        | RCHOP GCB | RCHOP non-GCB | R2CHOP GCB | R2CHOP non-GCB |
|--------|-----------|---------------|------------|----------------|
| N      | 78        | 61            | 27         | 18             |
| EFS24  | 74.4%     | 57.4%         | 81.5%      | 77.8%          |
| OS24   | 82.1%     | 72.1%         | 96.3%      | 88.9%          |
| EFS HR | Reference | 1.67*         | Reference  | 0.709          |
| 95% CI |           | 1.03 to 2.72  |            | 0.237 to 2.12  |
| EFS OS | Reference | 2.18**        | Reference  | 0.738          |
| 95% CI |           | 1.27 to 3.74  |            | 0.152 to 3.58  |

Abbreviations: EFS, event-free survival; OS, overall survival; HR, hazard ratio; CI, confidence interval
### Table 3

EFS24 Responder Alteration Details

| Pathway               | RCHOP Achieved | RCHOP Failed | R2CHOP Achieved | R2CHOP Failed | Proportional Difference | Result          |
|-----------------------|----------------|--------------|-----------------|---------------|--------------------------|-----------------|
| **PIM1**              | 5/35           | 8/26         | 5/14            | 0/4           | 0.522                    | Cell Survival/Proliferation, Somatic Hypermutation |
| **SPEN**              | 2/35           | 5/26         | 3/14            | 0/4           | 0.349                    | NOTCH           |
| **MYD88 (L265P)**     | 3/35           | 5/26         | 3/14            | 0/4           | 0.321                    | NFKB            |

Results are reported in decimal proportions on the basis of EFS24.