Whole-genome sequencing reveals oncogenic mutations in mycosis fungoides

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Key Points

- High-throughput sequencing of MF revealed multiple mutations within epigenetic and cytokine pathways that may drive disease.
- Pharmacologically targeting the JAK3 pathway in MF results in cell death and may be an effective treatment of this disease.

Introduction

Cutaneous T-cell lymphoma (CTCL) is a skin malignancy with significant mortality in advanced disease (27% 5-year survival).1 Mycosis fungoides (MF), the most common form of CTCL, classically presents as patches and plaques on the skin with disease progression leading to skin tumors, and lymph node, blood, and visceral involvement. The etiology of MF is unknown, but is thought to include chronic antigenic stimulation through viral/bacterial exposure,2,3 environmental exposures,4 and altered microRNA (miRNA) expression.5,6

A variety of genetic aberrations have been identified in MF. Specifically, mutations in the tumor suppressor p53 (TP53) were detected in 24% of tumor-stage MF, but less often in earlier-stage MF.7 TP53 mutations commonly demonstrate an UV B signature, with C-T transitions, suggesting UV-based therapy of early-stage disease may contribute to these mutations.8 Loss of other tumor suppressor genes, including CDKN2A and CDKN2B,8,9 and increased expression of NAV3, JUNB, and c-MYC, as well as hypermethylation of mismatch repair genes, are reported.9 MF has not been associated with a specific translocation. However, MF can have chromosomal gains and losses, with the most common being gains within chromosome 7 and losses within chromosomes 1, 10, and 17.10-12 It is unclear whether any of the alterations reported have a role in the development of MF.

The pathogenesis of mycosis fungoides (MF), the most common cutaneous T-cell lymphoma (CTCL), is unknown. Although genetic alterations have been identified, none are considered consistently causative in MF. To identify potential drivers of MF, we performed whole-genome sequencing of MF tumors and matched normal skin. Targeted ultra-deep sequencing of MF samples and exome sequencing of CTCL cell lines were also performed. Multiple mutations were identified that affected the same pathways, including epigenetic, cell-fate regulation, and cytokine signaling, in MF tumors and CTCL cell lines. Specifically, interleukin-2 signaling pathway mutations, including activating Janus kinase 3 (JAK3) mutations, were detected. Treatment with a JAK3 inhibitor significantly reduced CTCL cell survival. Additionally, the mutation data identified 2 other potential contributing factors to MF, ultraviolet light, and a polymorphism in the tumor suppressor p53 (TP53). Therefore, genetic alterations in specific pathways in MF were identified that may be viable, effective new targets for treatment. (Blood. 2015;126(4):508-519)

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Methods

Tissue procurement

Five patients with active tumor-stage MF underwent informed consent for participation in this study. Punch biopsies were obtained from a single MF tumor (tumor content ≥80%) and from clinically uninvolved skin (detailed in supplemental Methods, see supplemental Data, available on the Blood Web site). For 1 patient, an additional biopsy was obtained from a separate tumor, and the tumor cells cultured (without a matrix) as previously described.31 Additionally, frozen, hanked skin biopsies from 25 MF patients were retrospectively identified. All samples were obtained under an institutional review board–approved protocol (IRB# 111143).

WGS, exome, and Sanger sequencing/analysis

Genomic DNA from tissue was isolated using standard methods (details in supplemental Methods). Paired-end sequencing (PE-75) of genomic DNA from MF tumor and matched normal skin was performed (Illumina Genome Analyzer Ix). Reads were aligned to the human reference genome (hg19) using the Burrows-Wheeler Aligner (BWA),22 followed by realignment and recalibration of the initial alignment using the Genome Analysis Toolkit (GATK).31 Somatic variants were detected using VarScan224 and Strelka25 (additional details in supplemental Methods). The detected variants by either tool were pooled for follow-up analysis. Candidate somatic variant calls were inspected using Integrated Genomics Viewer (IGV). Copy-number variations were detected by Control-FREEC copy number and allelic content caller (control-FREEC).26,27 The Clipping Reveals Structure (CREST) tool was used to detect gross structural variations (ie, translocations).28 For validation, polymerase chain reaction (PCR) amplification of selected somatic single nucleotide variants (SNVs) and indels followed by Sanger sequencing was performed. Primer sequences are available upon request.

For exome sequencing, genomic DNA from CTCL cell lines was subjected to Exome capture (NimbleGen v3, 6Gb, Illumina HiSeq platform, PE-100). Reads were mapped as described in the previous paragraph followed by local realignment, score recalibration, and marking duplicate reads. Variants were called using the function “pileup2cns” implemented in VarScan2 (additional details in supplemental Methods).29 Variant annotations were based on ANNOVAR.29

Targeted ultra-deep sequencing and analysis

For targeted ultra-deep sequencing for JAK3, NOTCH2, and TP53, the Illumina MiSeq platform was used. After samples were library-prepped and run on the MiSeq, reads were mapped to hg19 using BWA,22 and realigned using GATK.31 Variants were called using SAMtools and BCFTools (additional details in supplemental Methods).30

UV signature analysis

We calculated the proportion of samples with dinucleotide mutations (dinucleotide polymorphism [DNP]) and those specifically with CC→TT mutations in MF tumors and 8 cancer types from which we previously collected somatic mutation data from large-scale next-generation sequencing projects.31 We next combined the SNV data from our 5 MF samples with those from 121 melanoma banked skin biopsies from 25 MF patients were retrospectively identified. All samples were obtained under an institutional review board–approved protocol (IRB# 111143).

Results

WGS of MF tumors revealed a spectrum of somatic genomic variants

To identify somatic genomic variants associated with MF, we performed WGS of tumor and matched uninvolved skin from 5 patients with tumor-stage MF (patient characteristics in Table 1). The mean depth of coverage for the WGS ranged from 32 to 44 for all samples (supplemental Table 1). WGS revealed an average of 27 602 (12 106-70 110) somatic SNVs and small insertions/deletions (indels), and an average of 102 (17-297) nonsynonymous or stop-gain exonic SNVs and indels across the 5 tumor samples (supplemental Tables 2-4). Indels in protein coding regions were infrequent (supplemental Tables 2 and 4). Using CREST analysis, 1 to 3 interchromosomal translocations were identified in tumors from patients 2, 3, and 5 (supplemental Table 5, supplemental Figure 1). Tumors in patients 1 and 4 had more translocations with 60 and 14, respectively. No translocations were similar between patients and were not further investigated.

The mutations identified were distributed without notable hotspots across the chromosomes (Figure 1A), and most mutations were novel (not known single nucleotide polymorphisms [SNPs] in dbSNP or in the 1000 Genome Project; supplemental Tables 3-4; supplemental Figures 1-2). Similar profiles of somatic mutation types were detected in the tumors, with intergenic and intronic mutations accounting for the vast majority of mutations (Figure 1B). There was variability in a small percentage of splicing and stop-gain mutations between samples. The mutation frequency in the 5 MF tumors was 5 to 26 per Mb with a mean of 12 mutations per Mb per tumor (supplemental Table 6), which is similar to the mutation frequency in lung cancers and melanoma.34

Sanger sequencing was performed on 92 SNVs and 5 indels to assess the accuracy of the WGS analysis. The SNVs and indels chosen included a random sampling and those with high oncogenic potential. We validated 90% of the alterations (supplemental Figure 3), and those that validated had a range of quality score and depth of coverage.

UV signature in mutations from MF tumors

Because MF is a hematologic disease of the skin, we sought to determine whether the types of mutations identified by WGS were consistent with those caused by UV radiation. Evaluating all variants identified within the 5 MF tumors, transition (C-T or G-A) mutations were commonly observed, accounting for 40% to 65% of SNVs. Two tumors (patients 3 and 4) also had an increased frequency of A-G or T-C transitions (~20%; Figure 2A). An examination of the dinucleotide variations (DNP) in MF samples compared with other cancers showed MF had a very high proportion of DNP (Figure 2B, supplemental Table 7). A hallmark of UV-induced mutations are transition mutations at dipyrimidines that are CC. Notably, CC DNPs in MF tumors were at a similar high proportion as those in melanoma, a malignancy known to have a UV component, and significantly higher than in other cancers (Figure 2B). Similar results were obtained when the number or the proportion of CC:GG to TT:AA mutations was evaluated (Figure 2B,
supplemental Figure 4). These results suggest that UV exposure may have a role in generating the abundant CC-TT mutations.

Evaluating the mutation signatures in MF and melanoma with NMF analysis,31,32 we observed 4 distinct mutation signatures, including 3 consistent with UV exposure (signatures 2, 3, and 4; Figure 2C). The high bars in each signature indicate characteristic mutation types in their trinucleotide context (Figure 2C). Signature 1 is unique to MF. We next assessed the mutation signature profile in individual MF and melanoma tumors for each signature. The magnitude of each signature as a component of the overall mutation load in each sample shows that the UV-related signatures are a factor in all 5 MF samples (Figure 2D). Specifically, for signature 2, MF tumors from patients 1 and 5 showed high coefficients, indicating that this UV-related signature significantly contributes to their mutation profiles (Figure 2D). There were moderate coefficients for patients 1 and 4 for signature 3 and patients 2 to 4 for signature 4. Although these analyses may be limited by sample number, the data suggest UV may be contributing to MF.

**Mutations identified in common pathways**

In the subset of genes containing exonic mutations, those likely to have a role in oncogenesis were identified. Pathway analysis of the nonsynonymous, exonic SNVs and indels revealed a diverse set of biological processes affected (Figure 3A).33,34 Within the 5 patients, there were 42 genes implicated in other malignancies, through gain of function (oncogenes), loss of function (tumor suppressor), or aberrant expression (Figure 3B). Of note, 41% of these mutations were C to T transitions, further supporting a role for UV in MF. Five of the 42 genes had mutations in 2 of the MF tumor samples: NCor1, FAT3, CLUAP1, BAI3, and ZEB1 (Figure 3B and supplemental Tables 3, 4, and 8). Additionally, 10 other genes not previously linked to cancer or that have uncharacterized roles in cancer had exonic, nonsynonymous mutations in >1 tumor sample (supplemental Table 8).

To investigate similarities between tumor MF and cultured CTCL lines at the genome level, we subjected 3 CTCL cell lines (HH, Hut-78, and MyLa) to exome sequencing. We also evaluated the Cancer Cell Line Encyclopedia (CCLE), which has sequenced a panel of ~1600 genes across cell lines, including HH and Hut-78.35 Eight genes with exonic mutations in 2 or more cell lines were likely to have a role in oncogenesis were identified (Figure 3C). Among them, 1 gene was determined to have an identical mutation observed in at least 2 of the cell lines. Six of the genes that were mutated in at least 2 of the cell lines were also reported in HH and/or Hut-78 in the CCLE (Figure 3C).

Given the poorly understood therapeutic success of histone deacteylase inhibition in patients with CTCL,36 the genes involved in epigenetic regulation were of specific interest. Four of five patient tumors sequenced and all CTCL lines had mutations in genes integral to chromatin remodeling and histone modification. Mutations were identified in genes that mediate histone methylation, acetylation, ubiquitination, and chromatin remodeling (Figure 3D). Evaluation of the 10 genes mutated in patient tumors showed that they were significantly enriched compared with 24,349 known protein-coding genes in NCBI RefSeq \( (P = 2.565 \times 10^{-11}) \) Fisher exact test. Notably, mutation of Ncor1, a corepressor that recruits histone deacetylases to chromatin, was present in a third of the samples (2 patients and 1 cell line).

Because Th2-skewed cytokine expression has a role in MF,38 we assessed genes that regulate cytokine expression or cytokine signal transduction. We identified mutations within cytokine pathway genes, and specifically in the IL-2 signaling pathway, in all but 1 of the MF tumor samples (Figure 3E). Mutations in the IL-2R signaling proteins (JAK3 kinase and its downstream target STAT5), as well as proteins that regulate IL-2 and IL-2R expression were present. Evaluation of the 4 genes mutated in patient samples showed that they were significantly enriched compared with 24,349 known protein-coding genes in NCBI RefSeq \( (P = 8.684 \times 10^{-11}) \) Fisher exact test. We also identified mutations in cytokine signaling pathway genes in the HH and Hut-78 cell lines (Figure 3E).

**SCNAs in MF**

Somatic copy-number alterations (SCNAs) were assessed in each patient. We observed both broad copy-number events that span a chromosome arm or an entire chromosome and focal copy-number events that affect much smaller regions than a chromosome arm (Figure 4A). Broad SCNAs were observed in all patients except patient 5, with patient 1 displaying the highest number of such events. For example, the MF tumor in patient 1 had only 1 copy of chromosome 3 and 13, copy-number loss of 6q, 10p, and 21q, and copy-number gain of 7q across all 5 patients in both their clinical stage.

**Table 1. MF tumor samples subjected to WGS**

<table>
<thead>
<tr>
<th>Patient</th>
<th>Ethnicity</th>
<th>Sex</th>
<th>Age, y</th>
<th>Tumor site</th>
<th>Normal tissue site</th>
<th>Pathology</th>
<th>Past therapy</th>
<th>Current therapy</th>
<th>TNMB</th>
<th>Clinical stage</th>
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</thead>
<tbody>
<tr>
<td>1</td>
<td>White</td>
<td>Female</td>
<td>70</td>
<td>Knee</td>
<td>Back</td>
<td>Nontransformed</td>
<td>XRT</td>
<td>Bexarotene, vorinostat</td>
<td>T3, N0, M0, B0</td>
<td>IIB</td>
</tr>
<tr>
<td>2</td>
<td>White</td>
<td>Male</td>
<td>67</td>
<td>Upper arm</td>
<td>Abdomen</td>
<td>Granulomatous</td>
<td>NB-UVB</td>
<td>Bexarotene</td>
<td>T3, N2, M0, B0</td>
<td>IIB</td>
</tr>
<tr>
<td>3</td>
<td>White</td>
<td>Male</td>
<td>41</td>
<td>Forearm</td>
<td>Abdomen</td>
<td>Nontransformed</td>
<td>Bexarotene</td>
<td>NB-UVB, topical bexarotene</td>
<td>T3, N0, M0, B0</td>
<td>IIB</td>
</tr>
<tr>
<td>4</td>
<td>Black</td>
<td>Male</td>
<td>35</td>
<td>Scalp</td>
<td>Forearm</td>
<td>Large cell transformed</td>
<td>XRT, ECP bexarotene, romidepsin</td>
<td>Gemcitabine</td>
<td>T3, N2, M0, B2</td>
<td>IVA</td>
</tr>
<tr>
<td>5</td>
<td>White</td>
<td>Female</td>
<td>69</td>
<td>Upper arm</td>
<td>Abdomen</td>
<td>Nontransformed</td>
<td>None</td>
<td>None</td>
<td>T3, N0, M0, B0</td>
<td>IIB</td>
</tr>
</tbody>
</table>

ECP, extracorporeal photopheresis; NB-UVB, narrow-band UV B; TNMB, tumor, node, metastasis, blood; XRT, local radiation therapy.
Figure 1. Mutation profiles of patient tumors subjected to WGS. (A) Each patient is plotted with a unique color. The vertical bars indicate the number of predicted, somatic SNVs in a 5-Mb window. (B) Pie charts of functional categories of the predicted somatic SNVs. For each patient, the left pie chart depicts the proportion of somatic mutations in categories based on their genomic location, including intergenic, intronic, ncRNA, and exonic SNVs. The right chart shows the breakdown of different mutation types in coding regions, including upstream/downstream, UTR, nonsynonymous, synonymous, stop-gain SNVs, and SNVs located in splicing sites. Only SNVs not included in dbSNP (build 131) or the 1000 Genome Project (November 2010) are shown. ncRNA, noncoding RNA; UTR, untranslated region.
Figure 2. Mutations in MF appear to have a UV signature. (A) Frequency of different mutation classes in each MF patient. (B) Samples with any DNP and with mutations that specifically change CC in different cancer types (left). The number of CC:GG to TT:AA mutations in the coding regions of each cancer type (right). BRCA: breast cancer, n = 507; CRC: colon and rectal cancer, n = 224; EC: endometrial carcinoma, n = 248; GBM: glioblastoma, n = 290; LUAD: lung adenocarcinoma, n = 182; Mel: melanoma, n = 121; OVCA: ovarian carcinoma, n = 316; SQCC: squamous cell lung cancer, n = 177. (C) Four mutation signatures of MF and melanoma tumors. Each color denotes 1 of the 6 possible trinucleotide mutations. The y-axis denotes the relative coefficient of each substitution to the corresponding signature. (D) Coefficient of each MF (red) and melanoma (blue) tumor for each signature. Only SNVs not in dbSNP (build 131) or the 1000 Genome Project (November 2010) were included.
normal and tumor sample, indicating these gains were germline in origin (Figure 4C).

**TP53 genetic variation is associated with MF**

A quarter of tumor-stage MF have somatic TP53 mutations, whereas TP53 mutations are less common in earlier disease.40 Somatic TP53 mutations were not detected in any of the 5 MF tumors evaluated by WGS. Specific polymorphisms within TP53 are associated with susceptibility to malignancy, including TP53 Pro72Arg.41 Given its potential significance in MF,7 we also evaluated TP53 polymorphisms. In addition to the initial 5 tumor-stage patients, we also obtained banked tumor tissue from 25 patients with MF. Of the 30 MF samples, 8 (27%) were homozygous Pro/Pro, 6 (20%) were heterozygous Arg/Pro, and 16 (53%) were homozygous Arg/Arg (Table 2). There were 4 samples from black patients (13%) and 26 from white patients. Comparing our results to a published control white population (n = 335), there was a significant increase in the frequency of Pro/Pro in our MF patients (27% vs 6%, χ² test, P < .001) and a similar frequency of Arg/Arg (53% vs 58%, χ² test, P = .45). When comparing only our white patients (n = 26) to the control white population, as there are ethnic differences in allele frequency,43 we still obtained an increased frequency of Pro/Pro (19% vs 6%, χ² test, P = .01). Fourteen of the 30 MF patients (47%) had at least 1 Pro allele, and the overall Pro allele frequency was increased compared with control (33% vs 24%, χ² test, P = .03). These data suggest the proline residue may be contributing to susceptibility of MF.

**Alteration of Notch pathway in MF**

The Notch protein family consists of 4 transmembrane receptors and 5 ligands that are integral to embryogenesis and cell-fate decisions.44,45 Activating mutations in NOTCH2 were implicated in the development of both B-cell lymphomas and murine T-cell leukemia.45-47 In addition to the increased copy numbers of NOTCH2 and NOTCH2NL in all patients subjected to WGS (Figure 4C), we also identified other mutations affecting the Notch pathway. For example, we identified with targeted ultra-deep sequencing a patient with early plaque-stage MF who had a NOTCH2 nonsynonymous SNV (C4699T, R1567W), which is predicted to be deleterious, and is located in the negative regulatory region between 2 previously identified gain-of-function mutations found in marginal zone lymphomas.46

**Identification of JAK3-activating mutations in MF**

JAK3-, A572V-, and A573V-activating mutations (within the pseudo-kinase regulatory domain) were reported in NK/T-cell lymphomas (Figure 5A). The FERM domain also harbors activating mutations (L156P, R172Q, E183G) in ATLL.17 We identified an activating JAK3 point mutation in 1 of the 5 MF tumor samples by WGS (c.C1718T, p.A573V) with a variant allele frequency of 41.67%. Sanger sequencing validated the mutation and showed mutation of a single allele at the 1718 position (Figure 5B). A biopsy of a separate tumor from the same patient placed in short-term culture showed the identical JAK3 mutation (Figure 5B). Sanger sequencing of JAK3 in 3 CTCL cell lines revealed
the same JAK3 C1718T (A573V) activating mutation in the Hut-78 cells (Figure 5C). JAK3 was not mutated in HH or MyLa cells. These data suggest this JAK3 mutation may be a contributing factor to MF.

To assess the rate of JAK3 mutation in a larger MF population, we performed targeted ultra-deep sequencing on 25 MF samples, 17 of which were early stage (stage Ia-IIa). Although we did not detect overt evidence of JAK3 mutations in these samples, 5 of them had ~1% allele frequency of previously reported activating JAK3 mutations (L156P, E183G, A572V, and A573V). Given the very low frequency of these variations and limited DNA remaining, these data could not be validated.
by other sequencing methods. Of note, early-stage MF have a low tumor burden and are inherently heterogeneous (mixture of cell types).

**CTCL cells are sensitive to a JAK3 inhibitor**

A JAK3 inhibitor, tofacitinib, has recently been US Food and Drug Administration (FDA) approved for rheumatoid arthritis treatment, as it inhibits the proliferation of activated T cells.\(^4^8\) Given our results demonstrating the presence of activating JAK3 mutations in MF and the Hut-78 line, we postulated tofacitinib may be useful as a therapeutic agent in MF/CTCL (Figure 3E). Evaluation of the JAK3/STAT5 signaling pathway in 3 CTCL lines showed increased phospho-STAT5 in Hut-78, compared with HH cells (Figure 5D). MyLa cells had robust levels of phospho-STAT5 likely due to increased STAT5 protein levels (Figure 5D). Following exposure to 200 nM tofacitinib, there were significantly reduced cell numbers for all 3 CTCL lines (Figure 5E). Notably, treatment of the CTCL lines with increasing concentrations of tofacitinib (1-500 nM) revealed a significantly reduced half-maximal inhibitory concentration (IC\(_{50}\)) in Hut-78 compared with HH and MyLa cells (2-way analysis of variance [ANOVA], \(P\) < .001; Figure 5F). All 3 lines had an IC\(_{50}\) well below the tofacitinib concentrations achieved in humans (85-107 ng/mL) based on FDA-approved dosage.\(^4^9\) There was a dramatic reduction of phospho-STAT5 in both Hut-78 and MyLa cells at tofacitinib concentrations as low as 10 nM (Figure 5G). Therefore, JAK3 inhibition may be therapeutically useful for treating CTCL with activating mutations in JAK3 as well as those malignancies with an overactive JAK3/STAT5 signaling pathway.

**Discussion**

As the pathogenesis of MF is poorly understood, and previous focused genetic studies have failed to identify mutations common to MF tumors,\(^7^9\) we performed WGS of MF tumors. Our analysis revealed a wide breadth of mutations with overlap between tumors, and many of the mutations in shared biologic pathways. We identified mutations in known oncogenes and tumor suppressor genes as well as genes not previously linked to cancer and certainly not to MF.

Of note, we identified mutations in many genes involved in epigenetic regulation (Figure 3D). Of the genes mutated, 3 of them have a specific role in the methylation of H3K4, which is associated with open chromatin and active transcription.\(^5^0\) MLL2 is a tumor suppressor and histone methyltransferase that helps to methylate H3K4.\(^5^1\) MLL2 mutations were identified in B-cell lymphomas, and our novel mutation (C8788T, P2930S) is in a similar region as several previously identified.\(^5^2\) SETD1A, a component of a histone methyltransferase complex that can mono-, di-, or trimethylate H3K4,\(^5^3\) had a missense mutation in 1 MF tumor. RNF20, an E3 ubiquitin ligase that ubiquitinates histone 2B, which is a prerequisite for methylation of H3K4,\(^5^4\) had a mutation in a coiled region in 1 tumor. Our results are interesting given the successful treatment of some MF patients with histone deacetylase inhibitors (HDACi), which have been shown to stimulate methylation of H3K4.\(^5^5\) We hypothesize mutations in the genes that regulate H3K4 methylation lead to reduced methylation, closed chromatin, and reduced transcription of tumor suppressors, which can be reversed with HDACi. These will be important areas in the future on which to focus research efforts.

Specific TP53 polymorphisms are linked to many malignancies,\(^5^6\) and we determined that the Pro72Arg polymorphism may be associated with MF. The proline allele is linked with improved DNA repair and cell cycle arrest, whereas the arginine allele is associated with enhanced apoptosis.\(^5^7\) Within dbSNP the G nucleotide (Arg) is the minor allele, as we indicated; however, both proline and arginine alleles are reported as the minor allele, and ethnicity can influence allele variance.\(^4^3\) Although a previous MF study did not find an increased frequency of Pro/Pro (n = 24;
European population), it demonstrated a 29% proline allele frequency in MF vs 19% in controls.⁵⁷ Within our cohort (30 patients), there was a significant increase of both the Pro/Pro genotype and the presence of at least 1 proline allele. Although this is a limited cohort, this TP53 polymorphism may be associated with susceptibility to MF, and suggest that a population-based study would be beneficial.

Notch signaling is involved in a multiple cellular processes, including embryogenesis and cell-fate decisions, including lymphocyte differentiation.⁴⁴,⁴⁵ Interestingly, NOTCH2 mutations, gain of function, copy-number gains, and loss of function have been associated with epithelial to mesenchymal transition in cancers,⁶⁵ inactivating mutations in 2 MF tumors. Although increased ZEB1 is associated with increased IL-2 proliferative signal through phosphorylation of STAT proteins (Figure 3E), it has activating mutations in NK/T-cell lymphoma (35%), ATLL (11%), and a single MF tumor. In the current study, we identified the same activating JAK3 mutation in an MF tumor evaluated with WGS and a CTCL line. Ultra-deep sequencing of an additional 25 MF samples revealed very low allele frequencies of known activating JAK3 mutations in 5 samples that we were unable to validate. Thus, although interesting, these data should be taken with caution. It is possible that the heterogeneous (mixture of cell types) nature of the biopsies of patch- and plaque-stage MF and the low tumor burden of these samples (~10% in patch stage) may be masking the true incidence of JAK3 mutation, but it is also possible that this frequency is within the margin of error or indicates a subpopulation. These findings highlight the difficulty of assessing tumor genetics in low tumor burden, heterogeneous samples. Regardless, an activating JAK3 mutation was verified in 1 of 12 (8.3%) MF tumor-stage samples and in 1 of 3 CTCL cell lines. These results indicate JAK3-activating mutations may be a driver of a portion of MF.

Importantly, we determined that our CTCL cell lines were sensitive to treatment with JAK3 inhibition. Hut-78, which harbored the activating JAK3 mutation, was the most sensitive to tofacitinib. Although there were no JAK3 mutations in HH or MyLa cells, they still demonstrated sensitivity to JAK3 inhibition, indicating that they are dependent on the pathway for their growth/survival. In support of this idea, MyLa cells had increased levels of STAT5 phosphorylation and IL-2 production and MF tumor cell growth (Figure 3E). Therefore, ZEB1 mutation and loss of heterozygosity (LOH) may be an important contributor to MF.

JAK3, which participates in transducing the IL-2 proliferative signal through phosphorylation of STAT proteins (Figure 3E), has activating mutations in NK/T-cell lymphoma (35%), ATLL (11%), and a single MF tumor. In the current study, we identified the same activating JAK3 mutation in an MF tumor evaluated with WGS and a CTCL line. Ultra-deep sequencing of an additional 25 MF samples revealed very low allele frequencies of known activating JAK3 mutations in 5 samples that we were unable to validate. Thus, although interesting, these data should be taken with caution. It is possible that the heterogeneous (mixture of cell types) nature of the biopsies of patch- and plaque-stage MF and the low tumor burden of these samples (~10% in patch stage) may be masking the true incidence of JAK3 mutation, but it is also possible that this frequency is within the margin of error or indicates a subpopulation. These findings highlight the difficulty of assessing tumor genetics in low tumor burden, heterogeneous samples. Regardless, an activating JAK3 mutation was verified in 1 of 12 (8.3%) MF tumor-stage samples and in 1 of 3 CTCL cell lines. These results indicate JAK3-activating mutations may be a driver of a portion of MF.

An interesting finding from our WGS analysis was that there appears to be a link between UV exposure and tumor-stage MF. There was a high frequency of UV-induced DNP mutations and mutation signatures consistent with UV exposure in all 5 MF tumors, which is comparable to melanoma, a malignancy with a known UV component. Only 2 of the 5 patients had UV therapy, but the sun exposure history for all the patients is unknown. It was proposed previously that UV exposure of MF may hasten progressive disease with UV signature TP53 mutations in advanced disease. It is concerning that although often effective, UV treatment of early-stage MF may create mutations leading to advanced disease, as well as increase risk of secondary skin cancer. Although this is not unique to MF, as secondary malignancies are correlated with other cancer therapies, our data again raise the possibility that UV exposure, including phototherapy, may contribute to MF.

Both WGS and targeted ultra-deep sequencing of specific genes are useful tools to help better detect mutations within specific genes and pathways in malignancies, as we discovered in MF. We identified genes in specific cytokine signaling pathways, cell-fate pathways, and epigenetic regulatory processes mutated in MF,
and a possible link to UV exposure, resulting in a better understanding of the pathogenesis of this disease. Consequently, new therapeutic options for MF have been identified and others can be developed.

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Authorship

Contribution: L.Y.M. and C.M.E. designed the experiments, analyzed the data, prepared figures, and wrote the manuscript; P.J. and Z.Z. analyzed the sequencing data, prepared figures, and assisted in writing the manuscript; K.B.D. assisted in designing the sequencing of patient samples, validating the WGS data, and evaluating data; L.Y.M., D.A.B., R.J.D., and D.H. performed experiments; J.A.Z. provided samples and participated in discussions of the data; J.P.Z. did the pathological evaluation and participated in discussions of the data; U.D. assisted in Figure 5F preparation and participated in data discussions; and all authors reviewed and edited the manuscript.

Conflict-of-interest disclosure: The authors declare no competing financial interests.
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