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Alternative exon usage in TRIM21 determines the antigenicity of Ro52/TRIM21 in systemic lupus erythematosus

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Abstract

The origin and mechanisms of autoantigen generation in systemic lupus erythematosus (SLE) are poorly understood. Here, we identified SLE neutrophils activated in vivo by interferon as a prominent source of Ro52/TRIM21 (hereafter Ro52), a critical autoantigen historically thought to be primarily generated by keratinocytes in SLE. Different to mononuclear cells and keratinocytes, SLE neutrophils are enriched in several unique Ro52 species containing a core sequence encoded by exon-4 (Ro52Ex4) in TRIM21. Ro52Ex4 is the main target of anti-Ro52 antibodies and is found in two Ro52 variants (Ro52α and a novel isoform termed Ro52γ) upregulated in SLE neutrophils. Further analysis of Ro52γ revealed a new subset of autoantibodies against a unique C-terminal domain (Ro52γCT) generated from a frameshift due to the lack of exon-6 in Ro52γ. Antibodies to Ro52Ex4 and Ro52γCT distinguish SLE patient subsets characterized by distinct clinical, laboratory, treatment and transcriptional profiles, which are not discerned by the “classical” anti-Ro52 antibodies. Together, these studies uncover interferon-activated neutrophils as a key source of unique immunogenic forms of Ro52 in SLE. Moreover, the finding of Ro52Ex4 and Ro52γCT as core targets of anti-Ro52 antibodies focus interest on Ro52γ as the potential isoform toward which immunological tolerance is initially lost in SLE.
Introduction

Ro52, also known as tripartite motif-containing protein 21 (TRIM21), is an interferon (IFN)-induced E3 ubiquitin ligase that drives negative feedback regulation during inflammation (1, 2), and has also been identified as a cytosolic antibody receptor involved in the intracellular clearance of antibody-coated viruses, such as adenovirus (3). Initially described as part of the Ro antigenic particle, Ro52 is among the first autoantigens discovered in systemic lupus erythematosus (SLE) (4, 5), a multisystemic autoimmune disease characterized by sustained IFN signaling and high titer autoantibodies leading to immune-mediated tissue damage (6, 7). Antibodies to Ro52 are frequently detected before clinical onset in SLE and are found in up to 40% of patients with established disease (1, 8).

Although Ro52 is mainly expressed by immune cells under steady state conditions (2, 9), its pathogenic relevance in SLE has been centered on keratinocytes, mainly because of the initial association between antibodies against the Ro particle with photosensitivity and cutaneous lupus (10-13). In particular, the redistribution of Ro52 on the cell surface and apoptotic blebs of keratinocytes in response to ultraviolet radiation is considered the main mechanism related to the immunogenic source of Ro52 in SLE (14, 15). Unexpectedly, during the search for in vivo IFN-induced autoantigens expressed by neutrophils in patients with SLE, we discovered Ro52 as a prominent neutrophil autoantigen with multiple structural forms, of which expression is related to in vivo IFN-induced activation in SLE neutrophils. Understanding the pathologic significance of Ro52 expression in SLE neutrophils is the focus of research addressed in the present study.

Results

Neutrophils express distinct forms of Ro52 linked to IFN activation in SLE

While neutrophils have gained major interest as a source of interferogenic signals in SLE, such as the release of genomic and oxidized mitochondrial DNA that activate
plasmacytoid dendritic cells (16-20), it is noteworthy that neutrophils are also important targets of IFN-I in this disease (16), suggesting the possibility that these cells may be an important source of autoantigens induced by IFN in SLE. To search for neutrophil autoantigens linked to IFN activation in SLE, we initially identified in vivo activated cells by IFN through the study of protein expression of the IFN induced protein with tetratricopeptide repeats 3 (IFIT3) —as marker of IFN-induced activation (21)— in freshly isolated peripheral blood neutrophils and mononuclear cells (PBMCs) from consecutive patients with SLE (Figure 1A).

SLE neutrophils and PBMCs with the lowest and highest IFIT3 expression (hereafter IFN low and IFN high, respectively) were then pooled according to cell type and IFIT3 expression (Figure 1B) and used to screen SLE sera (n = 20) by immunoblotting to identify neutrophil autoantigens whose expression parallel that of IFIT3. From these studies, we initially focused our interest on five SLE sera detecting a common pattern of 3 bands of ~43, 47 and 52 kDa, which were highly expressed by IFN-high SLE neutrophils, but absent in IFN-low SLE neutrophils (Figure 1C). Among these bands, the 52kDa species was also expressed in PBMCs regardless of IFN status (Figure 1C). Using two-dimensional electrophoresis and mass spectrometry, this set of bands was identified as Ro52 (Supplemental Figure 1).

Using commercial antibodies, we further addressed whether Ro52 has unique patterns of expression in SLE neutrophils and PBMCs according to IFIT3 levels (Figure 1, D-H). Intriguingly, different patterns of Ro52 were detected depending on the commercial antibody utilized. Two mouse monoclonals and one rabbit polyclonal antibody to Ro52 —in which the target epitopes are not disclosed— (Figure 1, D-F, respectively) reproduced identical patterns of Ro52 detected in neutrophils and PBMCs by SLE serum (Figure 1C), including some additional bands of ~27-33 kDa found in neutrophils. In contrast, two commercial antibodies against the N-terminal and C-terminal regions (amino acid residues
51-100 and 421-470, respectively) showed detection of Ro52—albeit modest—in PBCMs and poor detection of the set of Ro52 bands enriched in IFN-high neutrophils (Figure 1, G and H). Considering that the antibodies are directed against different regions in Ro52, we hypothesized that these discrepancies may be explained by the presence of distinct structural forms of Ro52, either transcriptional variants, degradation products or both.

**SLE neutrophils express novel splicing variants of Ro52**

To gain insights into the source of the different Ro52 species found in IFN-high SLE neutrophils, we focused on the search for Ro52 splicing variants. Ro52 is encoded by the TRIM21 gene, which is split into 7 exons (Figure 2A). The 5’ untranslated sequence is divided between exon 1 and 2, and the initiation codon is located in exon 2 (22). In order to identify novel Ro52 isoforms, we analyzed publicly available RNAseq data-sets (GSE149050 and GSE123949) of three circulating immune cell types [including neutrophils, classical monocytes (cMo) and T cells] (23) and keratinocytes from patients with SLE (24). Using the ‘new Tuxedo’ pipeline (25), we identified three TRIM21 transcripts in circulating cells: one corresponding to the sequence encoding the full-length protein (termed TRIM21α/Ro52α), a second transcript corresponding to TRIM21β/Ro52β (resulting from the splicing of exon 3 to exon 5, skipping exon 4) (22), and a novel variant termed TRIM21γ/Ro52γ that results from the alternative splicing of exon 5 to exon 7, skipping exon 6 (Figure 2A).

This analysis also demonstrated that the expression of Ro52 variants varies depending on the cell type. Thus, compared to controls, SLE T cells and cMo showed a significant upregulation of the Ro52α and Ro52β transcripts, respectively (Figure 2, B and C), while SLE neutrophils were distinguished by a significant upregulation of both Ro52α and Ro52γ transcripts (Figure 2, B and D, respectively). In contrast to circulating cells in SLE, keratinocytes only express TRIM21α/Ro52α and a new variant termed
TRIM21δ/Ro52δ that results from the alternative splicing of exon 6 to an internal splicing site in exon 7 (Supplemental Figure 2). Interestingly, even though SLE keratinocytes are hyperresponsive to IFNα2 (24), the expression of TRIM21α/Ro52α and TRIM21δ/Ro52δ was not different in SLE compared to healthy control keratinocytes either at baseline or after IFN stimulation (Supplemental Figure 2).

**Autoantibodies against Ro52 target a sequence encoded by exon 4 in TRIM21.**

To investigate whether the distinct patterns of Ro52 detection by SLE sera and commercial antibodies in SLE leukocytes is explained by targeting unique Ro52 isoforms, we focused on splicing variants upregulated in circulating SLE cells. Ro52α, which corresponds to full length Ro52, is a molecule of 52 kDa consisting of 475 amino acid residues (22). The deduced structure contains three distinct domains. The amino-terminal region includes two zinc-finger motifs (a RING-finger and a B-box), followed by a coiled-coil stretch including a putative leucine zipper and a B30.2/PRYSPRY domain in the C-terminal end (Figure 3A) (1). Ro52β is a ~45 kDa protein that lacks the sequence encoded by exon 4 (amino acid residues 167-245), which includes the leucine zipper and part of the coiled-coil domain (Figure 3B) (22). Ro52γ is a new variant consisting of a C-terminally truncated protein lacking the B30.2/PRYSPRY domain. The predicted product is a protein of 287 amino acids (~33 kDa) containing the RING, B-box, and coiled-coil domains, followed by a novel C-terminal sequence of 35 amino acid residues generated from a frameshift at the junction of exon 5 and 7 (Figure 3C).

To address the detection of Ro52 isoforms by SLE sera and commercial anti-Ro52 antibodies, cell lysates from HEK293 cells transfected to express the different Ro52 variants were analyzed by immunoblotting. Strikingly, all SLE sera and commercial antibodies recognizing the set of Ro52 bands in IFN-high SLE neutrophils (Figure 1, C-F) were specific for Ro52α and Ro52γ (Figure 3, D-I), demonstrating that the main epitope
targeted by antibodies against the Ro52 neutrophil variants is located within a sequence encoded by exon 4 (hereafter Ro52Ex4), which is missing in Ro52β. Since Ro52β contains both the N-terminal and C-terminal domains (Figure 3B), it is likely that the ~45 kDa band detected in PBMCs by antibodies to the N-terminal and C-terminal regions of Ro52 (Figure 1, G and H, respectively) corresponds to Ro52β, which is consistent with the upregulation of its transcript in SLE monocytes (Figure 2C). These data also suggest that Ro52β protein is not detected in neutrophils.

Using the monoclonal antibody D-12 (Figure 1D), we further demonstrated that neutrophils from consecutive patients with SLE are highly enriched with Ro52 species containing the Ro52Ex4 epitope, which exhibited a strong correlation with IFIT3 protein expression ($R^2 = 0.728, P <0.001$) (Figure 4, A and C). In contrast, a single band of Ro52 kDa was detected in SLE PBMCs, which also correlated —albeit less strongly— with IFIT3 levels ($R^2 = 0.271, P = 0.044$) (Figure 4, B and D). In cells from healthy donors, however, although the Ro52 kDa species of Ro52 was prominent in PBMCs, it was absent in neutrophils (Figure 4E). Moreover, the 43 kDa and 47 kDa species showed variable expression in control neutrophils, which was minimal compared to SLE neutrophils (Figure 4E). The set of ~27-33 kDa bands were present both in healthy and SLE neutrophils (Figure 4E). Together, these data demonstrate that compared to PBMCs and healthy control neutrophils, SLE neutrophils are highly enriched with Ro52 species containing the Ro52Ex4 sequence, which expression correlates with the IFN-induced marker IFIT3.

**Splicing variation of exons 4 and 6 in TRIM21 determines the antigenic targets of autoantibodies to Ro52 in SLE.**

Although Ro52Ex4 is well detected by SLE sera when found in the context of Ro52α and Ro52γ (Figure 3D), we further addressed whether the isolated Ro52Ex4 sequence is sufficient for efficient antibody recognition. Thus, we generated recombinant
proteins containing the Ro52Ex4 amino acid sequence either alone or in combination with the sequences encoded by flanking exons 3 and/or 5, i.e. exon 4, exons 4-5, exons 3-4, and exons 3-4-5 (amino acid residues 167-245, 167-253, 137-245, and 137-253, respectively) (Figure 5A), and tested their recognition by SLE sera using immunoblotting (Figure 5B). Although Ro52Ex4 is the target of SLE sera in Ro52α and Ro52γ (Figure 3D), the presence of exon 3 importantly enhanced or was necessary for the efficient detection of the isolated Ro52Ex4 protein sequence (Figure 5B). Because SLE sera lack recognition of Ro52β (Figure 3D), this excludes the possibility that the region encoded by exon 3 is independently immunogenic in SLE. Rather, it is likely that the exon 3 encoded sequence facilitates antibody binding by stabilizing the epitope encoded by exon 4. Alternatively, it is possible that the sequence encoded at the junction between exons 3 and 4 may play some role in autoantibody recognition.

Since the region encoded by exon 3 enables efficient detection of antibodies to Ro52Ex4 (i.e. anti-Ro52Ex4 antibodies), we used the recombinant protein containing amino acid residues 137-245 to screen for antibodies in SLE sera from the SPARE lupus cohort, for which extensive clinical and serologic variables are available (26, 27), and 80 healthy controls. Demographic, clinical and laboratory features of the SLE cohort are summarized in Supplemental Table 1. SLE patients had significantly elevated serum levels of anti-Ro52Ex4 antibodies compared to healthy controls (p<.0001). Using a cut-off value determined by ROC curve analysis, 50% of SLE patients (96/191) were positive for anti-Ro52Ex4 antibodies and 5% of controls (4/80) (Figure 5C).

Since Ro52Ex4 is found in variants Ro52α and Ro52γ, which are both transcriptionally upregulated in SLE neutrophils (Figure 2B and D), either isoform could serve as the antigen driving the production of anti-Ro52Ex4 antibodies in SLE. Nevertheless, it is intriguing that autoantibodies against the C-terminal half of Ro52α (amino acid residues 268-475, also shared by Ro52β) are rare (i.e. 0%) among anti-Ro52
antibodies in SLE (28). While it is possible that the canonical C-terminal region of Ro52α is not a self-immunogen in SLE, it is also possible that the Ro52 variant responsible for the production of antibodies against Ro52Ex4 does not contain the classic C-terminal sequence. This focused our attention on Ro52γ, which differs from Ro52α in that it possesses a unique C-terminal sequence caused by a frameshift at the junction of exon 5 and 7. Following the same principle that other regions in self-immunogenic Ro52 variants should be targeted by autoantibodies in SLE, we looked for antibodies against the unique C-terminal sequence found in Ro52γ (hereafter Ro52γCT). Importantly, a BLAST search for this novel sequence against the catalogue of annotated eukaryotic and prokaryotic proteins showed no similarity to any other known protein. Antibodies to Ro52γCT were significantly increased in SLE compared to healthy controls (p <0.001) and found in 22.5% (43/191) of patients with SLE and 3.8% (3/80) healthy controls (Figure 5D).

In the SPARE cohort, 39% (74/191) of the patients have antibodies to Ro52 (hereafter anti-Ro52'classic' antibodies) as detected by the clinically available assay (Supplemental Table 1). Analysis of the intersection of the different anti-Ro52 antibodies revealed that 88% (65/74) of SLE patients positive for anti-Ro52'classic' antibodies are also positive for anti-Ro52Ex4 antibodies (Figure 5E), demonstrating that positivity to anti-Ro52Ex4 antibodies identifies the majority of anti-Ro52'classic' antibodies in SLE. Further analysis of the small subset of SLE sera positive for anti-Ro52'classic' antibodies, but negative for antibodies to Ro52Ex4 (n=9), revealed that these sera target the N-terminal sequence (i.e. anti-Ro52Nt) shared by Ro52α, β, and γ (Figure 5F). This pattern of detection importantly contrasts with anti-Ro52Ex4 serum, which only recognizes Ro52α, Ro52γ and the RoEx3-4 protein sequence (Figure 5G). Thus, antibodies to Ro52Nt represent 4.7% (9/191) of anti-Ro52 antibodies in the SLE cohort, which is consistent with a previous study showing that 4% of anti-Ro52 positive SLE sera recognize the N-terminal sequence in Ro52 (amino acid residues 1–127) (28).
Antibodies targeting unique regions in Ro52 distinguish distinct clinical subsets and disease activity in SLE

From the analysis of Ro52 expression in SLE neutrophils and further mapping of antibody subsets based on recognition of Ro52 isoforms, antibodies to Ro52 in SLE are classified in three subsets: anti-Ro52Nt, anti-Ro52Ex4 and anti-Ro52yCt antibodies. Interestingly, the distinct antibodies to Ro52 identify clinically relevant endotypes within SLE (Figure 6). Anti-Ro52Nt antibody positivity was associated with lymphadenopathy and intriguingly, were less likely to be associated with lower levels of C4 (Figure 6A and Supplemental Table 2). Antibodies to Ro52Ex4 were significantly associated with history of sepsis, renal failure, digital gangrene, anemia, and antibodies to La and RNP (Figure 6A and Supplemental Table 2). In contrast, patients with anti-Ro52yCT antibodies showed an increased frequency of stroke, features of secondary Cushing’s syndrome (i.e., moon facies and buffalo hump), and were more likely treated with Mycophenolate (Figure 6A and Supplemental Table 2).

At time of visit, the presence of anti-Ro52Nt antibodies showed limited value for SLE disease activity. Rather, these antibodies showed negative associations with laboratory and clinical features linked to disease activity. For instance, this group of patients showed lower SLEDAI, lupus activity index (LAI), renal and hematologic activity, higher C3 and C4, and low use of prednisone (Figure 6B and Supplemental Table 3). In contrast, anti-Ro52Ex4 antibodies were associated with lower lymphocyte and higher neutrophil counts, lower C3, higher SLEDAI and LAI, increased rash and renal activity, and higher prednisone doses (Figure 6B and Supplemental Table 3). Moreover, anti-Ro52yCT antibodies were associated with elevated ESR, lower hsCRP and C3, and a tendency for higher SLEDAI score (Figure 6B and Supplemental Table 3).
Despite the prominent overlap with antibodies to Ro52Ex4, anti-Ro52‘classic’ antibodies were only associated with anemia, anti-La and sepsis, but not with features of disease activity as anti-Ro52Ex4 antibodies (Supplemental Tables 2 and 3). These differences are likely explained because anti-Ro52Nt antibodies, which negatively associate with disease activity, are within the pool of anti-Ro52‘classic’ antibodies. Moreover, anti-Ro52Ex4 antibody positivity was found in 26 additional patients negative for anti-Ro52‘classic’ antibodies. Thus, autoantibodies targeting specific domains in Ro52 variants appear to be more informative than anti-Ro52‘classic’ antibodies to identify clinical subsets in SLE.

Anti-Ro52 antibody subsets exhibit distinct transcriptional profiles in SLE

Patients with SLE display unique blood transcriptional signatures associated with immune pathways activated during active disease (29). In particular, it is interesting that the IFN signature has been linked with antibodies to the Ro particle (30). Since anti-Ro52 antibody subsets correlated with distinct clinical features in SLE, we further addressed the relationship of these antibodies with transcriptional fingerprints activated in SLE. Using whole blood gene expression data collected in parallel with the samples used to measure anti-Ro52 antibodies, we performed a three-way comparison between SLE patients positive for antibodies to Ro52Nt, Ro52Ex4 and Ro52γCt (Figure 7A).

We identified 926 differentially expressed transcripts (DETs) between the three anti-Ro52 antibodies (Figure 7A and Supplemental file 1). DETs associated with anti-Ro52Ex4 (n=138) were predominantly enriched in genes involved in immune mediated pathways including IFN, intracellular DNA sensing, B-cell receptor signaling, necroptosis and degranulation (Figure 7B, Supplemental file 1, and Supplemental figure 3). Interestingly, a group of common DETs between anti-Ro52γCT and anti-Ro52Ex4 (n=174) was mainly enriched in pathways related to IFN signaling (including increased expression
of TNFSF13B, TLR7, IFIT3 and IRF5) and antigen presentation. In contrast, DETs exclusive to anti-Ro52γCT (n=385) contained genes related to p53 signaling, RNA metabolism, ribosome biogenesis, mitochondrial function, and negative-regulation of IFN-I (Figure 7B, Supplemental file 1 and Supplemental figure 3). DETs associated with anti-Ro52Nt antibodies lacked enrichment of IFN-stimulated genes (ISGs), which is consistent with the low disease activity in this small group of patients. Instead, these antibodies were associated with increased vascular endothelial growth factor, TNF signaling and apoptosis (Figure 7B, Supplemental file 1, and Supplemental figure 3). Using gene set variation analysis (GSVA) (31) to quantify the activity of selected pathways on individual SLE patients, we further confirmed that antibodies to Ro52Ex4 have the most significant association with the IFN signature, followed by anti-Ro52γCT antibodies and no association of anti-Ro52Nt antibodies with the IFN signature was found (Figure 7C).

Discussion

The origin and mechanisms underlying the production of autoantigens in SLE remain unclear. Since core autoantigens in SLE — such as Ro52, Ro60, La, histones, dsDNA and RNP — are normally expressed under steady state conditions, it has been hypothesized that self-immunogenic forms of these molecules are generated under unique inflammatory environments amplified in SLE. When we examined for autoantigens targeted by SLE serum in in vivo IFN-activated neutrophils from patients with SLE, we were struck by the finding that Ro52 is a prominent IFN-induced autoantigen in SLE neutrophils, which is absent or minimally detected at the protein level in both healthy control and “steady-state” SLE neutrophils. Moreover, IFN-activated SLE neutrophils contained multiple Ro52 species recognized by autoantibodies, which were absent in PBMCs. By combining the analysis of an RNAseq dataset of peripheral blood SLE leukocytes, epitope mapping with commercial anti-Ro52 antibodies, and the study of a
large cohort of patients with SLE, we further concluded that the large majority of antibodies to Ro52 in SLE are directed against an epitope encoded by exon 4 in TRIM21 —found in Ro52α and Ro52γ— which is the major target of anti-Ro52 antibodies in in vivo IFN-activated SLE neutrophils. The significant association of anti-Ro52Ex4 and anti-Ro52γCT antibodies with the IFN signature provides additional evidence that this set of autoantibodies is mechanistically related to the IFN-induced activation in SLE.

Although the expression of Ro52α and Ro52γ can explain the binding of anti-Ro52Ex4 antibodies to SLE neutrophils, the sole expression of these isoforms (molecular weights 52 and ~33 kDa, respectively) is insufficient to elucidate the origin of the broad range of bands containing the Ro52Ex4 epitope, which are detected in IFN-activated SLE neutrophils. While we cannot discard the existence of additional Ro52 isoforms, an alternative hypothesis is that Ro52α and Ro52γ suffer additional modifications, creating complex patterns of Ro52 detection. In the case of Ro52α, cleavage and/or trimming of the N-terminal and C-terminal regions, leaving fragments containing the core Ro52Ex4 sequence intact, could explain the detection of multiple Ro52 species below 52 kDa. Regarding Ro52γ, this isoform would require modifications that increase the molecular weight of the protein. It is noteworthy that both SLE and healthy control neutrophils contain molecular weight bands below 33 kDa, which were detected by the three commercial antibodies against Ro52Ex4, supporting that these bands are likely generated from Ro52α/Ro52γ. As an additional hypothesis, it is possible that Ro52 is normally degraded in neutrophils under steady state conditions, but instead accumulates during SLE disease activity as result of increase expression and likely less degradation. Although neither of these possibilities are exclusive of each other, the most striking finding is that the protein species containing Ro52Ex4 are both enriched in in vivo IFN-activated neutrophils and are the main target of anti-Ro52 antibodies in SLE. This finding is unlikely to be coincidental, but rather mechanistically related.
Ro52α is an abundant isoform constitutively expressed by several cell types, such as keratinocytes and immune cells (2, 9, 14). If Ro52α is the main immunogen in SLE, it is intriguing that the major target of anti-Ro52 antibodies is limited to Ro52Ex4, sparing the C-terminal half of the molecule (28). In contrast, however, Ro52γ is targeted both at Ro52Ex4 and the C-terminal domain, offering it an alternative self-antigen to explain the subsets of anti-Ro52 antibodies found in SLE—i.e. anti-Ro52Nt, anti-Ro52Ex4, and anti-Ro52γCT—, as well as the limited humoral response against C-terminal Ro52α. The proposal that the N-terminal and C-terminal regions of Ro52α may be cleaved/trimmed, enriching for immunogenic fragments containing Ro52Ex4, is also a potential mechanism to explain the antibody specificity to Ro52Ex4 without targeting the C-terminal domain. Indeed, both models are not exclusive, but complementary. Importantly, whereas it is possible that other cell types in different tissues may generate similar Ro52 patterns as IFN-activated neutrophils in SLE, our work demonstrates that neutrophils are the main cellular source of multiple Ro52 protein species containing Ro52Ex4 in peripheral blood in SLE. In this scenario, large amounts of immunogenic Ro52 may become accessible to the immune system as result of neutrophil death in SLE, which may initiate an immune response to Ro52.

Interestingly, the striking difference in the phenotype of two Ro52/Trim21 knockout mice, in which one develops lupus-like disease and the other has a normal life span (2, 9), has been attributed to the potential production of a truncated Ro52 protein that is overexpressed in the mouse that develops lupus (32, 33). Like Ro52γ, except for the lack of the unique Ro52γ-CT domain, the predicted truncated protein carries the RING domain, B-box and the coiled-coil domain containing the homologous sequence of Ro52Ex4, which is instead encoded by exon 5 in mouse Trim21 (32). In the context of our findings, it is possible that lupus-like disease in these mice is driven by the truncated Ro52γ-like protein, which may work either as an autoantigen or through a unique pro-inflammatory
function resulting from the lack of the B30.2/PRYSPRY domain, which mediates protein–protein interactions and Fc receptor function (34, 35).

Our finding that SLE neutrophils activated by IFN are highly enriched in Ro52 species targeted by the majority of anti-Ro52 antibodies in SLE, underscores neutrophils as the main cellular source of self-immunogenic Ro52 in peripheral blood in SLE. Moreover, the study uncovers the isoform-specific domains Ro52Ex4 and Ro52γCT as the core targets of anti-Ro52 antibodies in SLE, which could be used as potential biomarkers of disease state and to unravel disease mechanisms associated with SLE.

Methods

Study design and participants.

The objective of the study was to identify neutrophil autoantigens with unique patterns of expression linked to in vivo activation by IFN in SLE. Peripheral blood mononuclear cells (PBMCs) and neutrophils were purified from 19 consecutive SLE patients. Briefly, after Ficoll-Hypaque isolation of mononuclear cells, neutrophils were isolated by 2 cycles of red blood cell lysis using ACK lysing buffer (Quality Biological). Cells were lysed and boiled immediately after purification in SDS-sample buffer. Sera from 80 healthy controls and 191 SLE patients from the “Study of biological Pathways, Disease Activity and Response markers in patients with Systemic Lupus Erythematosus” (SPARE) (26, 27) cohort were screened for the presence of anti-Ro52 antibodies. SPARE is a prospective observational cohort that has been extensively described previously (26). Briefly, adult patients (age 18 to 75 years-old) who met the definition of SLE per the revised American College of Rheumatology classification criteria were eligible into the study (36). At baseline, the patient's medical history was reviewed, and information on current medications was recorded. Patients were followed-up over a 2-year period. Patients were treated according to standard clinical practice. Disease activity was
assessed using the Safety of Estrogens in Lupus Erythematosus: National Assessment (SELENA) version of the Systemic Lupus Erythematosus Disease Activity Index (SLEDAI) (37) and physician global assessment (PGA) (38). C3, C4, anti-dsDNA (Crithidia), complete blood cell count and urinalysis were performed at every visit. Study participants also underwent whole blood gene expression analysis at baseline using the Affymetrix GeneChip HT HG-U133+ (26, 27). All samples were obtained under informed written consent approved by the Johns Hopkins Institutional Review Board.

**Autoantigen discovery.**

PBMC and neutrophil lysates from patients with SLE were used to detect IFIT3 (IFN-induced protein with tetratricopeptide repeats 3) by immunoblotting as a marker of IFN activation (21). To identify neutrophil-specific SLE autoantigens linked to IFN activation, cell lysates with low and high IFIT3 expression were pooled according to cell type and IFIT3 expression and used to screened by immunoblotting sera from 20 consecutive patients with SLE. The identity of autoantigens of interest was further determined by two-dimensional electrophoresis and mass spectrometry (MS), as previously described (39).

**Antibodies.**

Mouse monoclonal anti-human IFIT3 (H00003437-B01) was purchased from Abnova, mouse monoclonal anti-human Ro52 (clone D-12) from Santa Cruz Biotechnology, rabbit anti-N-terminal TRIM21 polyclonal antibody from Origene (TA335782), rabbit anti-human TRIM21 polyclonal antibody from Proteintech (121081-1-AP), mouse anti-human TRIM21 monoclonal antibody from Proteintech (671361-1-Ig), rabbit anti-C-terminal TRIM21 polyclonal antibody from Sigma (AV381248), mouse anti-
human β-actin from Sigma (A5316), and mouse anti-human histone H3 from Cell Signaling (96C10).

**Cloning, production and expression of recombinant Ro52 isoforms and domains.**

The coding sequence of Ro52α was amplified from SLE neutrophil cDNA and used as template to generate Ro52β and Ro52γ by deleting exons 4 and 6, respectively. Exon deletion was performed using the Q5 Site-Directed Mutagenesis Kit (New England Biolabs). All Ro52 isoforms were cloned into pcDNA3.1 and pET28a(+). The sequences spanning exons 4, 4-5, 3-4 and 3-4-5 were amplified by PCR using Ro52α as template and cloned into pET28(+). Recombinant proteins were expressed in *E. coli* BL21 (DE3) and purified by Ni-NTA affinity chromatography. All recombinant proteins are N-terminal His-tagged. The complete C-terminal domain of Ro52γ (Ro52γCT) was generated as a 35mer synthetic peptide (SPHHSGRHSQSVADTFRRSETSEAWRHPAEHTWK). HEK-293T cells were transiently transfected with plasmids to express Ro52 isoforms using lipofectamine 2000 (Invitrogen). HEK-293 cells were lysed after 48 hours post-transfection and used for immunoblotting analyses.

**Detection of antibodies to Ro52, Ro52Ex4 and Ro52γCT.**

“Classical” anti-Ro52 antibodies were detected by ELISA using a commercial kit from INOVA (QUANTA Lite SS-A 52, Part number 704505). Anti-Ro52Ex4 and anti-Ro52γCT antibodies were measured in serum/plasma by homemade ELISA. Briefly, Nunc Maxisorp plates were coated with 4 μg/mL recombinant Ro52 exon 3-4 to detect anti-Ro52Ex4 antibodies or 5 μg/mL of Ro52γCT was covalently attached to Nunc CovaLink NH plates to detect anti-Ro52γCT antibodies. The plates were blocked for one hour with phosphate buffered saline plus 0.1% Tween-20 (PBST) with 3% non-fat milk. Serum/plasma was diluted 1:1000 in PBST 1% non-fat milk and assayed in duplicate using
antigen-conjugated plates and plates without antigen for background subtraction. HRP conjugated goat anti-human IgG was used as a secondary antibody (Diluted at 1:10,000 in PBST 1% nonfat milk). Anti-Ro52Ex4 and anti-Ro52γCT antibody arbitrary units were calculated using a standard curve made of a serial diluted serum from a high-titer SLE patient. Anti-Ro52Nt antibodies were detected by immunoblotting using recombinant Ro52α, Ro52β, Ro52γ and the Ro52 exon 3-4 encoded sequence.

**Differential Gene expression and enrichment analyses.**

Gene expression analysis from the SPARE cohort was previously described.(27) CEL files were subjected to RMA background correction, and quantile normalization, using the Oligo package (29). To select only expressed genes in whole blood, we filtered out transcripts that had a raw signal < 100 in less than 10% of samples with the genefilter R package. All calculations and analyses were performed using R (ver 4.2.1) and Bioconductor (ver 3.15.2) (40). Differentially expressed transcripts (DETs) were analyzed using the R package ‘limma’ using a multivariate linear model adjusted of anti-dsDNA positivity and SLEDAI (41). Gene set enrichment analyses were done using the online platform Metascape.org (42). Three-way differential gene expression analyses were done using the package ‘volcano3D’ (43) by combining the results from the pairwise comparisons between anti-Ro52Ex4, anti-Ro52γCT, and anti-Ro52Nt antibodies, and the F test calculated with the R package ‘limma’. To annotate the transcriptomes associated to each anti-Ro52 antibody type, we performed gene set enrichment analyses using the online platform Metascape.org (42). Briefly, in order to do the enrichment analyses, we split the differentially expressed transcripts (DET) into lists according to the results of three-way analysis. Then, the lists of genes were uploaded into the Metascape.org platform. An adjusted p-value <0.05 was considered as significant. Activity of the interferon
pathways was calculated using gene-set variation analysis (GSVA) with the R package GSVA (31).

**Isoform analysis in RNAseq data**

To discover new isoforms of the Ro52 antigen, we reanalyzed RNAseq data from neutrophils, classical monocytes (cMo), and T-cells from 24 SLE patients and 12 healthy controls from the publicly available data set GSE149050 (23). In addition, we analyzed RNAseq data from keratinocytes from 7 HC and 7 SLE subjects deposited at GSE123949 (24). Briefly, the fastq files were aligned to the human genome reference build 38 (GRCh38/hg38) using the splicing-aware aligner HISAT2 (44). Sample’s BAM files were further processed using StringTie to quantify de novo assembled transcripts (45). Visualization of the de novo assembled Ro52 (TRIM21) transcripts was done using Ballgown (46).

**Statistical analysis.**

Comparisons of continuous variables between two groups were done using Student’s T test. Fisher’s exact test was used for univariate analysis on SPARE cohort variables, the exact 2x2 package in R was used to calculate the p-value, OR, and 95% CI. Effect size (Cohen’s D) between anti-Ro52 positive and negative SLE subjects was calculated using the ‘psych’ R package (47). Pairwise comparisons between of pathway activity were done using Wilcoxon’s pairwise test. Statistical significance was set at p < 0.05. The statistical analyses were carried with the R software version 4.2.1.

**Author contributions**

All authors were involved in drafting the article or revising it critically for important intellectual content, and all authors approved the final version to be published.
Study conception and design
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Experimental work
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Acquisition of data
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Analysis and interpretation of data
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References


33. Espinosa A, Brauner S, Ambrosi A, Kuchroo V, and Wahren-Herlenius M. Comment on "Gene Disruption Study Reveals a Nonredundant Role for
TRIM21/Ro52 in NF-kappa B-Dependent Cytokine Expression in Fibroblasts Response. *Journal of Immunology.* 2009;183(12):7620-1.


Figure 1. SLE neutrophils overexpress distinct forms of Ro52 in relation to interferon (IFN) pathway activation. (A) Neutrophils (PMN) and PBMCs from 19 consecutive patients with SLE were analyzed by immunoblotting using antibodies to IFIT3, histone H3 (H3) and β-actin (loading controls). Representative samples from 9 patients are shown. (B-H) Lysates from SLE neutrophils and PBMCs with low and high activation by IFN based on IFIT3 expression (B) were used to screen 20 SLE sera (C) and five commercial antibodies to Ro52 (D-H). Histone H3 is shown as loading control in B. In (C), data from five SLE sera recognizing a set of autoantigens overexpressed in SLE neutrophils with high IFIT3 expression are shown.
Figure 2. Neutrophils express splicing variants of TRIM21/Ro52. (A) Schematic representation of the transcripts corresponding to TRIM21 isoforms found in a publicly available RNAseq dataset (GSE1491050) from classical monocytes (cMo), neutrophils (PMN), and T cells from SLE patients (n=24) and healthy controls (HC, n=12) using the ‘new tuxedo’ pipeline. Each solid block represents an exon. (A-C) Differential expression analyses of TRIM21α (B), TRIM21β (C), and TRIM21γ (D) between HC and SLE according to cell type. Pairwise comparisons between HC and SLE were done using Wilcoxon’s Test. *p<0.05, **p<0.01.
Figure 3. Autoantibodies against Ro52 target a sequence encoded by exon 4 in TRIM21. (A-C) Schematic representation of exon usage and structural domains in Ro52 isoforms. (D-I) SLE sera (D) and commercial anti-Ro52 antibodies (E-I) from Figure 1C-H were used to immunoblot cell lysates from HEK293 cells transfected with mock (empty vector, EV) or plasmids expressing Ro52α, Ro52β or Ro52γ. The arrows denote the Ro52 isoforms. Detection of Ro52α, Ro52β and Ro52γ by the N-terminal anti-Ro52 antibody in H is also shown as loading control. RING, “Really interesting new gene”; BB, B-box domain; CC, coiled-coil; LZ, Leucine-zipper.
Figure 4. Relationship between IFIT3 and Ro52 expression in neutrophils and PBMCs from SLE patients and healthy controls. (A) Neutrophils (PMN) and (B) PBMCs from 19 consecutive patients with SLE were analyzed by immunoblotting using antibodies to IFIT3, Ro52 (D-12 mouse monoclonal antibody), histone H3 (H3) and β-actin (loading controls). Representative data from 10 patients are shown. (C and D) Correlation between the expression of IFIT3 and Ro52 in PMN and PBMCs (C and D, respectively). The
expression of Ro52 and IFIT3 was quantified by densitometry from the corresponding bands in A and B, and the values were fitted to a linear regression model. (E) PBMCs and PMN from 12 healthy controls (Ctrl) and one patient with SLE were analyzed by immunoblotting using antibodies to IFIT3, Ro52 (D-12 mouse monoclonal antibody) and H3 (loading control). Representative data from 6 healthy controls are shown. The SLE samples were included for comparison.
**Figure 5. Patients with SLE have autoantibodies targeting distinct Ro52 isoforms.**

(A) Schematic representation showing the regions encoded by TRIM21 exon 4, exons 4-5, exons 3-4, and exons 3-4-5 in Ro52.  
(B) Recombinant proteins containing the sequence encoded by TRIM21 exon 4, exons 4-5, exons 3-4, and exons 3-4-5 were detected by immunoblotting using SLE sera positive for anti-Ro52Ex4 antibodies. Representative data from 6 sera are shown.  
(C and D) Levels of antibodies to Ro52Ex4 (C) and Ro52γCT (D) in sera from the SPARE cohort (SLE) and healthy controls (HC). Comparisons were done using Student’s T test.  
(E) Venn diagram depicting the anti-Ro52 antibody intersections (overlap) in 128/191 SLE patients positive for anti-Ro52Ex4, anti-Ro52γCT, and/or anti-Ro52‘classic’ antibodies.  
(F-G) Recombinant Ro52α, Ro52β, Ro52γ and the Ro52 exon 3-4 encoded sequence were used to analyze by immunoblotting SLE sera from the none
overlapping anti-Ro52 ‘classic’ antibodies (n=9) (representative data from 5 sera are shown) (F), and anti-Ro52Ex4 serum (G). ****p<0.0001.
Figure 6. Anti-Ro52 antibody subsets are associated with distinct clinical endotypes in SLE. (A) Clinical and laboratory associations present during the clinical course of SLE according to anti-Ro52 antibody type. Odds ratios (OR) were calculated against SLE patients Negative for the corresponding anti-Ro52 autoantibody type using a 2x2 Table. Error bars correspond to 95% confidence interval (CI). Associations were tested using chi-square test or Fisher’s exact test. (B) Left panel, effect size (Cohen’s D) between positive vs. negative SLE patients for each anti-Ro52 antibody type over clinical variables and disease activity evaluated at time of visit. Right panel, heatmap of -log10 (p values)
showing the significant associations among anti-Ro52 autoantibody types and variables obtained at time of visit. P values were calculated with the Student’s t test. DBP: Diastolic blood pressure, UrPr/Cr: Urinary protein/creatinine ratio, hsCRP: high-sensitivity c-reactive protein, ESR: erythrocyte sedimentation rate, ACL: anti-cardiolipin antibodies, RVVT: Dilute Russell Viper Venom Time, SLEDAI: Systemic Lupus Erythematosus Disease Activity Index, LAI: Lupus Activity Index. *p<0.05, **p<0.01, *** p<0.001.
Figure 7. Anti-Ro52 antibody subsets are associated with distinct transcriptional immune-mediated pathways in SLE. (A and B) Three-way differentially expressed transcript (DET) analysis between SLE patients positive for anti-Ro52Ex4 (n=95), anti-Ro52Nt (n=9), and anti-Ro52γCT (n=21) antibodies. DET between the three anti-Ro52 antibody types (n=926) are shown in a 3D volcano plot (A) and in a radial plot (B). Significance was calculated using the volcano3D package by combining the results of the F test and pairwise comparisons between anti-Ro52Ex4, anti-Ro52Nt, and anti-Ro52γCT, using a multivariate linear model adjusted by anti-DNA positivity and SLEDAI. Color code denotes significant DET in anti-Ro52Ex4 (red), anti-Ro52γCT (green), anti-Ro52Nt (blue),
and overlapping genes between anti-Ro52Ex4 and anti-Ro52γCT (yellow), anti-Ro52Nt and anti-Ro52Ex4 (purple), anti-Ro52γCT and anti-Ro52Nt (light blue). Representative genes from the enriched pathways on each DET subset are labeled. (C) Activity of the IFN pathway in anti-Ro52 negative (n=63), and SLE patients positive for anti-Ro52Ex4 (n=95), anti-Ro52Nt (n=9), and anti-Ro52γCT (n=21) antibodies. Pathway activity was calculated using gene set variation (GSVA) score. Comparisons between groups were done using the pairwise Wilcoxon test. *p<0.05, ****p<0.0001.