Autoantibody-mediated impairment of DNASE1L3 activity in sporadic systemic lupus erythematosus

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Antibodies to double-stranded DNA (dsDNA) are prevalent in systemic lupus erythematosus (SLE), particularly in patients with lupus nephritis, yet the nature and regulation of antigenic cell-free DNA (cfDNA) are poorly understood. Null mutations in the secreted DNase DNASE1L3 cause human monogenic SLE with anti-dsDNA autoreactivity. We report that >50% of sporadic SLE patients with nephritis manifested reduced DNASE1L3 activity in circulation, which was associated with neutralizing autoantibodies to DNASE1L3. These patients had normal total plasma cfDNA levels but showed accumulation of cfDNA in circulating microparticles. Microparticle-associated cfDNA contained a higher fraction of longer polynucleosomal cfDNA fragments, which bound autoantibodies with higher affinity than mononucleosomal fragments. Autoantibodies to DNASE1L3-sensitive antigens on microparticles were prevalent in SLE nephritis patients and correlated with the accumulation of cfDNA in microparticles and with disease severity. DNASE1L3-sensitive antigens included DNA-associated proteins such as HMGBl. Our results reveal autoantibody-mediated impairment of DNASE1L3 activity as a common nongenetic mechanism facilitating anti-dsDNA autoreactivity in patients with severe sporadic SLE.

Introduction
The serological hallmark of systemic lupus erythematosus (SLE) is the production of antibodies (Abs) to nuclear antigens (Ags), with immune complex formation leading to systemic immune activation and tissue inflammation. Anti–double-stranded DNA (anti-dsDNA) IgG Abs can be found in up to 70–80% of SLE patients, of which ∼40% have high titers rarely seen in other autoimmune disorders (Rekvig, 2015; Tan et al., 1966). High titers of anti-dsDNA IgG correlate with disease severity (Banchereau et al., 2016) and often accompany disease flares, becoming especially evident when involving the kidney (Pisetsky, 2016; Yung and Chan, 2015). Despite the clinical relevance of anti-dsDNA Abs and numerous studies supporting the pathogenicity of this reactivity, the nature of antigenic DNA and the mechanisms that prevent its recognition by autoreactive B cells still remain largely elusive.

The development of anti-dsDNA Abs involves T cell help, making it unlikely that the “naked” protein-free dsDNA is the initiator of the response. In fact, Abs to chromatin often precede the onset of anti-dsDNA Abs, implicating protein-bound DNA in the initial breakdown of tolerance, followed by the spreading of autoreactivity toward naked dsDNA (Munroe et al., 2016). In turn, the antigenic chromatin may exist as small soluble fragments or comprise larger extracellular structures such as neutrophil extracellular traps (Garcia-Romo et al., 2011; Hakkim et al., 2010; Lodol et al., 2016) and/or microparticles (MPs). Indeed, 1–2-µm-sized MPs that are derived from the cell membranes of apoptotic cells are found in the plasma of healthy subjects and SLE patients (Dieker et al., 2016; Nielsen et al., 2012). Moreover, MPs can carry apoptotic cell DNA and expose it on their surface (Casciola-Rosen et al., 1994; Radic et al., 2004;...
Ullal et al., 2011). This would effectively create an “ideal” source of antigenic DNA since it is accessible to autoreactive B cells, relatively resistant to degradation, and associated with proteins that can provide potential T cell epitopes (Beyer and Pisetsky, 2010). In addition to the physical form of DNA, the length of DNA is likely to influence its antigenicity. Anti-dsDNA Abs were proposed to bind longer fragments of naked dsDNA with higher avidity (Ali et al., 1985; Papapanteli et al., 1980), although this was shown to depend on assay conditions (Pisetsky and Reich, 1994). Overall, the precise physical form of cell-free DNA (cfDNA) that serves as an Ag in anti-dsDNA responses, and its relationship with DNA size, are not well understood.

The availability of self-DNA to autoreactive B cells can be limited by extracellular DNases that digest it (Elkon, 2018; Soni and Reizis, 2018). DNASEI3L and DNASE1 are homologous extracellular DNases that are responsible for the majority of DNase activity in the circulation (Napirei et al., 2009). DNASEI3 contains a unique positively charged C-terminal peptide that facilitates digestion of membrane- and/or protein-associated DNA, including intact chromatin (Napirei et al., 2005; Sisirak et al., 2016; Wilber et al., 2002). Homozygous null mutations in DNASEI3 cause pediatric-onset familial SLE characterized by prominent anti-dsDNA Abs and renal involvement (Al-Mayouf et al., 2011; Batu et al., 2018; Belot, 2020; Carbonella et al., 2017; Ozçakar et al., 2013). In some of these cases, the disease initially manifested as hypocomplementemic urticarial vasculitis syndrome (HUUVS; Jara et al., 2009) but almost invariably progressed to severe SLE with glomerulonephritis (Carbonella et al., 2017; Ozçakar et al., 2013). A coding polymorphism of DNASEI3 that creates a hypofunctional protein DNASEI3 R206C (Ueki et al., 2009) is associated with several seropositive autoimmune diseases, including SLE (Acosta-Herrera et al., 2019). Furthermore, DNASEI3-deficient mice rapidly develop anti-chromatin and anti-dsDNA Abs followed by SLE-like disease with renal involvement (Sisirak et al., 2016; Soni et al., 2020; Weisenburger et al., 2018). Thus, DNASEI3 represents a conserved extracellular DNase that maintains tolerance to self-DNA in a nonredundant manner. DNASEI3 was shown to digest genomic self-DNA in apoptotic MPs (Sisirak et al., 2016), as well as reduce the level of polynucleosomal DNA fragments in the circulation (Chan et al., 2020; Serpas et al., 2019). However, it is unclear how these two activities are related to one another or relevant for the role of DNASEI3 in preventing SLE. Moreover, the relevance of DNASEI3 or its DNA substrates to sporadic SLE beyond the rare genetic DNASEI3 deficiency has never been established.

This study addresses the relevance of DNASEI3 and its DNA substrates to sporadic SLE, with an emphasis on lupus nephritis (LN). We show that patients with LN often manifest reduced activity of DNASEI3 in the circulation, which is associated with auto-Abs to DNASEI3 itself. Furthermore, reduced DNASEI3 activity in LN patients is associated with cfDNA accumulation in MPs, as well as with autoreactivity toward DNA and DNA-associated proteins on the surface of MPs. In aggregate, these results establish DNASEI3 as a novel self-Ag in patients with spontaneous SLE with renal involvement and the resulting deficiency of functional DNASEI3 as a likely cause of cfDNA abnormalities that in turn facilitate anti-DNA responses.

### Results

#### The activity of DNASEI3 is reduced in SLE patients with renal involvement

Initial experiments were done to establish a reliable assay for DNASEI3 activity in the circulation. Unlike its homologue, DNASE1, DNASEI3 efficiently digests native chromatin in intact nuclei (Napirei et al., 2005). Using purified recombinant DNASEI3, we confirmed its superior ability, relative to DNASE1, to digest DNA in intact nuclei isolated from Jurkat T cells (Fig. 1A). High concentrations of DNASEI3 produced a smear pattern of nuclear DNA digest, consistent with inefficient random cleavage (Fig. 1B). In contrast, even low concentrations of DNASEI3 produced a characteristic nucleosomal ladder, suggesting efficient internucleosomal cleavage of chromatin (Fig. 1B). The distinctive ability of DNASEI3 to digest DNA in nuclei was used to measure its specific enzymatic activity in human plasma. Whereas control human plasma efficiently digested DNA in nuclei, plasma from DNASEI3-deficient patients failed to do so in the same dilution range (Fig. 1C). A lower dilution of plasma from DNASEI3-deficient patients was able to digest nuclei in our assay, likely reflecting the activity of DNASEI1 and/or nonspecific inhibition of detection (e.g., by plasma IgG; Sidstedt et al., 2018). Plasma from an SLE patient who was heterozygous for the hypomorphic R206C variant of DNASEI3 also showed a noticeable reduction (Fig. 1D), confirming the specificity and sensitivity of this assay. To measure the relative activity of DNASEI3 in patient plasma, we set the 50% inhibitory concentration (IC50) value for pooled healthy control plasma (n = 15) at 100%, the value for DNASEI3-deficient patient plasma at 0% and considered any activity below two standard deviations of the control (81%) as reduced.

We then evaluated DNASEI3 activity in several groups of SLE patients (patient demographics and average values for key readouts are provided in Table 1). Note that “renal SLE” indicates past or present renal disease and “LN” indicates renal SLE with persistent proteinuria (urine protein/creatinine ratio [UPCR] >0.5) accompanied by positive serologies (hypocomplementemia and/or elevated anti-dsDNA Abs) at the time of sampling. Average DNASEI3 activity was not significantly decreased in 14 SLE patients without current or previous renal involvement compared with healthy controls (median activity 92%; Fig. 1E). In contrast, 57% of patients with a history of renal disease (n = 37) showed a decrease in DNASEI3 activity (median activity 69%) with a significant reduction compared with both healthy controls and SLE patients who never had renal disease (Fig. 1E). Within the renal SLE group, DNASEI3 activity was lower in those with active proteinuria than in those with resolved proteinuria (i.e., inactive renal disease) defined as UPCR <0.5 at the time of sampling (Fig. 1F). Notably, plasma activity of DNASEI3 did not correlate with the degree of proteinuria (Fig. S1), arguing against protein loss through the kidney as a cause of reduced DNASEI3 activity. To further evaluate the clinical significance of reduced DNASEI3 activity, we evaluated 15 mothers of...
children with heart block and high anti-Ro Abs, most of whom were clinically asymptomatic or had only minimal symptoms of SLE but did not meet formal classification criteria (Table 1). Only three showed a reduction in DNASE1L3 activity, two of whom were the only patients who progressed to SLE with renal disease (Fig. 1E, arrows). Collectively, these data reveal a significant reduction of circulating DNASE1L3 activity in sporadic SLE patients with renal disease.

Reduced DNASE1L3 activity is associated with Abs to DNASE1L3
The known hypomorphic allelic variant of DNASE1L3, DNASE1L3 (R206C), is predominantly found in subjects of European ancestry, and indeed, only two patients in our SLE cohort carried this variant. However, reduced DNASE1L3 activity was observed in >50% of our renal SLE patients across all ethnic groups, prompting a search for a nongenetic cause. To test whether auto-Abs to DNASE1L3 might contribute to its decreased activity, we examined patient plasma by ELISA using purified DNASE1L3 as the Ag. No reactivity to DNASE1L3 was observed in plasma samples from healthy controls (Fig. 2, A–C) or DNASE1L3-deficient patients (data not shown). In contrast, 17 out of 40 (43%) evaluated patients with renal SLE showed a mean anti-DNASE1L3 Ab level greater than the threshold OD value of 0.44 (mean OD +2 SD of healthy controls; Fig. 2A). Only 3 out of 17 (18%) plasma samples from nonrenal SLE patients and one
Table 1. Demographics and readouts in different study cohorts

| Demographics                                      | Healthy control (n = 15) | DNASE1L3 deficient (n = 4) | SLE renal (n = 87) | SLE nonrenal (n = 33) | Anti-Ro* (n = 45) |
|--------------------------------------------------|-------------------------|---------------------------|-------------------|----------------------|------------------|
| Demographics                                     |                         |                           |                   |                      |                  |
| Female, n (%)                                     | 12 (80)                 | 2 (50)                    | 75 (87)           | 31 (94)              | 45 (100)         |
| Age (yr), median (IQR)                           | 36 (23–59)              | 9 (7–14)                  | 3–7 (20–78)       | 39 (27–80)           | 35 (27–64)       |
| Race/ethnicity, n (%)                            |                         |                           |                   |                      |                  |
| Hispanic/Latino                                   | 0                       | 0                         | 25 (29)           | 16 (48)              | 0                |
| Non-Hispanic/Latino                              | 0                       | 0                         | 53 (61)           | 17 (52)              | 0                |
| White                                             | 5 (33)                  | 4 (100)                   | 38 (44)           | 23 (70)              | 38 (84)          |
| Black                                             | 5 (33)                  | 0                         | 27 (31)           | 7 (21)               | 1 (2)            |
| Asian                                             | 1 (7)                   | 0                         | 21 (24)           | 2 (6)                | 3 (7)            |
| Other                                             | 1 (7)                   | 0                         | 2 (2)             | 1 (3)                | 2 (4)            |
| Not reported                                      | 3 (20)                  | —                         | —                 | —                    | 1 (2)            |
| Readouts*                                        |                         |                           |                   |                      |                  |
| DNASE1L3 activity, %                             | 100 (89–100.5, n = 15) | 0 (n = 4)                 | 68 (62–100, n = 37)| 92 (85–100, n = 14) | 100 (85–100, n = 15) |
| Anti-DNASE1L3 ELISA, OD                          | 0.15 (0.13–0.26, n = 15)| —                         | 0.40 (0.26–0.80, n = 40) | 0.30 (0.23–0.36, n = 18) | 0.19 (0.13–0.25, n = 18) |
| cfDNA total, ng/ml                               | 5.6 (3.2–8.4, n = 15)  | 10.0 (9.0–17.6, n = 4)   | 6.3 (3.4–11.8, n = 44) | 5.4 (2.6–10.2, n = 15) | 17.2 (6.5–26.5, n = 15) |
| MP cfDNA fraction, %                             | 23 (19–35, n = 15)     | 70 (40–83, n = 4)        | 62 (39–73, n = 44) | 33 (26–46, n = 15)   | 49 (43–60, n = 15) |
| DNASE1L3-sensitive binding to MPs*                | 0 (0%, n = 15)          | 3 (75%, n = 4)           | 50 (57%, n = 87)  | 5 (15%, n = 33)      | 3 (7%, n = 45)   |

Continuous variables are given as median (interquartile range [IQR]), and categorical variables are given as number as cases (%).
*The age of one control subject was unknown and was not included in the calculation.
*Not all patients in every group were tested for all readouts. The n in each of the readouts refers to the number of patients from that group who were tested for this readout.
*Table 1 shows DNASE1L3-sensitive binding at the time cfDNA and plasma activity of DNASE1L3. Fig. 6 G shows the number of patients with DNASE1L3-sensitive binding at any time point.

anti-Ro* mother with renal disease were positive in this assay (Fig. 2, A and B). The same plasma samples that showed strong reactivity to DNASE1L3 (Fig. 2 C) had minimal binding to the FLAG epitope present in this protein or immobilized DNASE1 (Fig. S2 A). To rule out any differences in immobilization between DNASE1 and DNASE1L3, we incubated plasma from SLE patients with high anti-DNASE1L3 titers with a concentration range of DNASE1 or DNASE1L3 and then tested the reactivity to plate-bound DNASE1L3 by ELISA. Representative examples in Fig. 2 D show that IgG binding was competed out by DNASE1L3, but not by DNASE1. Taken together, these data suggest a specific auto-Ab response predominantly directed toward DNASE1L3.

The binding of plasma IgG to recombinant DNASE1L3 was further confirmed by reducing SDS-PAGE followed by Western blot, in which the reactivity was observed only in plasma samples that showed anti-DNASE1L3 binding by ELISA (Fig. S2 B). In addition, DNASE1L3 was incorporated into a bead array that allows simultaneous analysis of reactivity to multiple Ags. Anti-DNASE1L3 Ab levels measured by bead array strongly correlated with those determined by ELISA (r = 0.80, P < 0.0001, n = 25; Fig. S2 C). Clq and dsDNA, Abs to which are closely linked to LN (Yung and Chan, 2015), were also included in the bead array to assess autoreactivity to these Ags. However, neither anti-Clq nor anti-dsDNA Ab levels showed a significant correlation with anti-DNASE1L3 reactivity (Fig. S2, D and E). Thus, Ab reactivity to DNASE1L3 in sporadic SLE could be demonstrated by three different methods and appears to be distinct from other autoreactivities.

Importantly, the levels of anti-DNASE1L3 Abs strongly correlated with reduced plasma activity of DNASE1L3 (Fig. 2 E). Moreover, positive anti-DNASE1L3 binding in ELISA reliably identified patients with DNASE1L3 activity below the threshold (Fig. 2 E, horizontal black dashed line; sensitivity, 71%; specificity, 89%; positive predictive value, 83%; negative predictive value, 81%). To verify that Abs to DNASE1L3 can inhibit its activity, we purified IgG from the plasma of three LN patients with positive anti-DNASE1L3 binding by ELISA (similar to LN1 and LN3 in Fig. 2 C). Preincubation of DNASE1L3 with IgG from these patients, but not with IgG from three healthy controls, decreased enzymatic activity by an average of 30% in our assay (Fig. 2 F and Fig. 2 G, left panel). Similar inhibitory effects were also observed when nuclear DNA and naked genomic DNA were used as DNA substrates (Fig. 2 F), suggesting that the Abs primarily target the core catalytic activity of DNASE1L3. In contrast, no inhibition of DNASE1 activity was observed with the same IgG concentration (Fig. S2 F and Fig. 2 G), and only minor inhibition was achieved at a 10-fold higher concentration (Fig. S2 F). Overall, these data suggest that the reduced DNASE1L3 activity...
Figure 2. Reduced DNASE1L3 activity is associated with neutralizing anti-DNASE1L3 Abs. (A-C) Anti-DNASE1L3 (1L3) Ab levels in the plasma of sporadic SLE patients as measured by ELISA. (A) Anti-DNASE1L3 (1L3) Ab OD values in individual patients at a single plasma dilution (1:40); bars represent mean ± SD. Patients included healthy controls (n = 15), renal (n = 40) and nonrenal SLE patients (n = 17) by ACR criteria, and anti-Ro Ab–positive mothers of children with neonatal lupus (Anti-Ro+, n = 18). Dashed line indicates the threshold for a positive anti-DNASE1L3 Ab (mean OD +2 SD of control). Statistical significance was determined by Kruskal–Wallis test followed by the Dunn’s multiple-comparison test; *, P < 0.05; ****, P < 0.0001 (two tailed). (B) Average OD values ± SD for plasma dilutions (1:40–1:640) in the indicated groups of SLE patients and controls. Significance levels are given for the comparison between renal and nonrenal. Statistical significance was determined by Mann–Whitney test; *, P < 0.05; **, P < 0.01; ****, P < 0.0001 (two tailed). (C) Dilution curves for healthy control plasma (Ctrl) and plasma from five SLE patients with LN (LN1–LN5). Representative of two independent experiments. (D) Cross-reactivity of anti-DNASE1L3 Abs to DNASE1 as measured in a competition assay. Recombinant DNASE1 (D1, squares) or DNASE1L3 (D1L3, circles) was incubated with plasma at a single dilution, and the binding of plasma IgG to DNASE1L3 was measured by ELISA. Shown are two representative patient samples (LN3 and LN5 from C). Dotted lines represent OD values in the absence of any competitor. Representative of two independent experiments. (E) Correlation between DNASE1L3 activity and anti-DNASE1L3 (1L3) Ab levels. Symbols represent individual patients with sporadic renal (red) or nonrenal (black) SLE. Dashed lines indicate thresholds for reduced DNASE1L3 activity (black) and positive anti-DNASE1L3 Ab (green). Results of Spearman correlation with a two-tailed P value are shown. (F and G) Inhibition of DNase activity by IgG from patients with anti-DNASE1L3 Abs. (F) Recombinant DNASE1L3 (1L3) was preincubated with purified IgG from plasma of a healthy control (Ctrl) or SLE patient with LN and used to digest DNA in native nuclei (left panel) or naked genomic DNA (right panel). Shown is the amount of remaining DNA after digestion with different concentrations of DNASE1L3. Dashed lines indicate 100% (black) and 50% (gray) of input DNA. Representative of two independent experiments. (G) Recombinant DNASE1L3 (1L3, left panel) or DNASE1 (right panel) was preincubated with purified IgG and
used to digest naked DNA (as in Fig. 1A). Shown is the relative enzymatic activity after preincubation with IgG from three controls or LN patients relative to the respective enzymes alone. Symbols represent IgG from individual patients; bars represent median ± interquartile range. Representative of two independent experiments.

in sporadic human SLE is caused primarily by specific Abs that develop against this protein.

DNASE1L3 controls the amount of DNA in MPs
We next sought to determine the impact of reduced DNASE1L3 activity on cfdNA in sporadic SLE patients. Given that DNA in circulating MPs was proposed as a physiological substrate of DNASE1L3 (Sisirak et al., 2016), we sought to characterize MP-bound and soluble cfdNA in normal human plasma. Plasma from healthy donors was subjected to several centrifugation steps to isolate a MP fraction free of cell debris and platelets. Resulting MPs were then stained for plasma membrane and DNA, using a membrane-permeable dye, and examined by confocal microscopy. Fig. 3A shows representative examples of 1–2-µm-sized particles composed of plasma membrane and containing dsDNA, suggesting that our centrifugation method indeed isolates human MPs with dsDNA. Similarly, a fraction of AnnexinV+ MP isolated from human plasma stained positive for DNA by flow cytometry (data not shown).

The fraction of MP-associated cfdNA was estimated by purifying DNA from the MP and MP-depleted fractions of 15 healthy controls and quantifying genomic DNA by quantitative PCR (qPCR) for Alu repeats in the genome. The concentration of MP-associated cfdNA ranged from 0.4–3.0 ng/ml plasma (Fig. 3B), corresponding to 17–48% (median 23%) of total cfdNA in the circulation (Fig. 3D). The three control samples with DNASE1L3 activity on or slightly below the threshold (Fig. 1E) showed the fraction of MP-associated DNA (28–37%) that was slightly above the median. In contrast, the distribution of cfdNA was strongly skewed toward the MP fraction in three out of four
DNASE1L3-deficient patients (Fig. 3 C), representing 31–86% (median 69%) of total cfDNA (Fig. 3 D). Of note, three of these four DNASE1L3-deficient patients were children diagnosed with HUVS before progression to SLE, suggesting that the observed cfDNA redistribution is not a consequence of severe SLE and/or anti-dsDNA reactivity. In our cohort of sporadic SLE patients, only those with renal SLE showed a significant increase in the MP cfDNA fraction (renal median, 62%; nonrenal median, 33%; Fig. 3 D), mirroring the changes observed with genetic DNASE1L3 deficiency. In contrast to the observed increase in MP-associated cfDNA, the total concentration of cfDNA was not significantly increased in any group, highlighting the specific effect of DNASE1L3 on cfDNA accumulation in MPs (Fig. 3 E).

The concentration of soluble cfDNA was slightly decreased in renal SLE compared with controls (2.2 versus 5.0 ng/ml), but the difference was not significant (P = 0.08). Importantly, the fraction of MP-associated cfDNA showed a significant inverse correlation with DNASE1L3 activity (Fig. 3 F); thus, patients with reduced DNASE1L3 activity harbored on average 67% of cfDNA in the MP fraction. The fraction of MP cfDNA showed a significant correlation with anti-dsDNA Ab titers (Fig. S3); nevertheless, multiple patients with a high fraction of MP-associated cfDNA had low or absent anti-dsDNA Ab titers, ruling out the coincidental isolation of DNA-containing immune complexes with MPs. In summary, while a substantial fraction of total cfDNA in healthy humans is associated with MPs, the representation of this fraction is regulated by DNASE1L3 and is significantly increased in sporadic SLE patients with reduced plasma DNASE1L3 activity.

The mouse model of DNASE1L3 deficiency was used to directly test the role of DNASE1L3 in the regulation of MP-associated DNA. The dual membrane/DNA staining protocol used for human MPs (Fig. 3 A) was used to confirm the presence of DNA-containing MPs in murine plasma (Fig. 4 A). DNA-carrying MPs in mice appeared slightly smaller than in human MPs (Fig. 4 D). Conversely, DNA-carrying MPs in mice harbored a normal amount of cfDNA in MPs but significantly more cfDNA in the soluble fraction compared with age- and sex-matched WT mice (Fig. 4 D). Conversely, Dnase1l3−/− mice harbored a normal amount of cfDNA in MPs but significantly more cfDNA in the soluble fraction compared with age- and sex-matched WT controls (Fig. 4 E). Thus, the loss of DNASE1L3, but not of its homologue, DNASE1, causes the accumulation of cfDNA in the MP fraction, implicating reduced DNASE1L3 activity in SLE patients as a likely cause of increased DNA load in MPs.

**MPs carry longer DNA fragments that are more immunogenic**

Given that a substantial fraction of human cfDNA is contained within the MP fraction, we tested whether this MP-associated DNA is qualitatively different from that in the soluble fraction. First, we estimated the length of cfDNA fragments in both fractions using the ratio of long to short amplicons recovered by qPCR (Jiang and Lo, 2016). This approach detected an increase in DNA length in the MP fraction compared with the soluble fraction of plasma from healthy controls (Fig. 5 A). Consistent with findings in the total cfDNA of DNASE1L3-deficient human patients (Chan et al., 2020), both MP-associated and soluble cfDNA from DNASE1L3-deficient patients was longer than in healthy controls (Fig. 5 A). To directly test the size distribution of MP-associated cfDNA in healthy controls, we sequenced it and computed the length of resulting fragments aligned to the genome (Jiang and Lo, 2016). No differences in genome coverage or in the end motif frequency were observed between MP-associated and soluble cfDNA (data not shown). However, the predominant DNA peak of mononucleosomal size (~180 bp) was overrepresented in the soluble fraction, whereas larger DNA fragments in the di- to trinucleosomal range (250–600 bp) were significantly overrepresented in the MP fraction (Fig. 5, B and C). Thus, MP-associated cfDNA is enriched in longer polynucleosomal fragments compared with soluble cfDNA, and MPs from DNASE1L3-deficient patients show lengthening of their cfDNA cargo compared with healthy subjects.

Given the observed enrichment of longer polynucleosomal fragments in MPs and the control of MP-associated DNA by DNASE1L3, we tested whether longer DNA fragments might represent more robust Ags. We incubated plasma from SLE patients with high anti-dsDNA titers with mononucleosomes or polynucleosomes and then tested the binding of plasma IgG to plate-bound DNA by ELISA. A representative example in Fig. 5 D shows that IgG binding to DNA was competed out by polynucleosomes, but not by mononucleosomes. This was observed using plasma samples from five patients (Fig. 5 E) and with purified IgG fractions from two of these patients (data not shown). In contrast, protein-free DNA fragments (95–831 bp) efficiently blocked IgG binding to DNA irrespective of their size (Fig. S4 A and Fig. 5 F). Moreover, IgG binding to the same fragments immobilized on a plate was slightly better for smaller fragments (Fig. S4 B) as observed previously (Pisetsky and Reich, 1994). The differences were not as prominent as observed by Pisetsky and Reich, likely because we used poly-lysine precoating to facilitate DNA binding to the plate. Thus, the impact of DNA length on auto-Ab binding is only apparent in solution and when the DNA is in its physiological chromatin state. In addition to serving as a self-Ag, cfDNA may induce innate immune responses such as the production of type I IFN (IFN-I) by plasmacytoid dendritic cells (pDCs), especially when in complexes with neutrophil-derived cationic peptides such as LL-37 (Garcia-Romo et al., 2011; Lande et al., 2011). We therefore incubated different concentrations of polynucleosomes or mononucleosomes with LL-37 and tested the ability of resulting...
complexes to activate pDCs from healthy donors. We found that polynucleosomes elicited IFN-I production at lower concentration than mononucleosomes (Fig. 5 G). Thus, longer chromatin fragments, such as those enriched in MPs, show better auto-Ab binding as well as better ability to elicit innate responses.

**DNASE1L3-sensitive MP Ags are targeted in SLE with renal involvement**

In view of reduced DNASE1L3 activity and the corresponding DNA accumulation in MPs of patients with renal SLE, we tested the functional consequences of these findings for autoreactivity in these patients. We have previously reported that DNASE1L3-deficient mice and ∼30% of human sporadic SLE patients harbored IgG binding to the surface of in vitro-generated MPs. The binding could be abolished by pretreatment of MPs with DNASE1L3, but not with DNASE1, suggesting Ab reactivity against DNASE1L3-sensitive Ags on the surface of MPs (Sisirak et al., 2016). We confirmed these observations (Fig. 6 A) and standardized the assay for an unbiased determination of DNASE1L3-sensitive binding of plasma IgG to MP Ags (referred to hereafter as "DNASE1L3-sensitive binding"). MPs are known to be heterogeneous (Dalli et al., 2013; Nielsen et al., 2011), thus the gating of MPs was deliberately broad to include all fractions below 1 µm in size; indeed, only a fraction of MP showed positive staining with IgG (Fig. 6, A–D). Among four patients with genetic DNASE1L3 deficiency, three manifested DNASE1L3-sensitive binding (Fig. 6 B). One DNASE1L3-deficient patient for whom longitudinal samples were available developed DNASE1L3-sensitive...
Figure 5. **MPs carry longer polynucleosomal fragments with better antigenic properties.** (A) Average length of cfDNA in the MP and MP-depleted (soluble) fractions of plasma from healthy controls (Ctrl; gray, n = 7) and DNASE1L3-deficient patients (IL3−/−; blue, n = 4) as determined by long to short amplicon ratio. Symbols represent individual subjects; bars represent mean. Unpaired samples (Ctrl versus IL3−/−) were compared using unpaired Student’s t test, and paired samples (MP versus soluble) were compared using paired Student’s t test; *, P < 0.05; **, P < 0.01; ****, P < 0.0001 (two tailed). (B) Size distribution of cfDNA fragments in the MP (gray) and MP-depleted (soluble, gold) fractions of plasma from healthy controls as determined by sequencing. Lines represent averages of four healthy controls. (C) Fraction of cfDNA fragments of the indicated length ranges in the MP and MP-depleted (soluble) fractions of plasma from healthy controls as determined by sequencing. Symbols represent individual healthy subjects; bars represent mean. Samples were compared using paired Student’s t test; *, P < 0.05 (two tailed). (D and E) The effect of chromatin size on binding to anti-dsDNA Abs in a competition assay using plasma of an SLE patient. D shows a representative titration of polynucleosomes (purple) and mononucleosomes (white); dashed line shows binding in the absence of competitor DNA. E shows the inhibition of binding by anti-dsDNA IgG from different SLE patients (n = 5) at the maximum inhibitor concentration (10 µg/ml). Symbols represent individual plasma samples, and bars represent median ± interquartile range. Statistical significance was determined by Mann–Whitney test; **, P < 0.01 (two tailed). (F) Inhibitory capacity of protein-free DNA fragments of the indicated lengths in bp on the binding by anti-dsDNA IgG at the maximum inhibitor concentration (10 µg/ml). Symbols represent individual SLE patients (n = 5), and bars represent median ± interquartile range. (G) The effect of chromatin size on interferon production by pDCs. Mononucleosomes (white) or polynucleosomes (purple) were incubated with antimicrobial peptide LL-37 and used to stimulate pDCs from healthy donors, and IFN-α was measured by ELISA after 24 h. Each point represents an average of two healthy donors, and bars represent SD of the mean. Data are representative of two independent experiments. Statistical significance was determined by Mann–Whitney test; *, P < 0.05 (two tailed).
binding shortly before a severe flare of LN that required temporary dialysis. This was also accompanied by a marked increase of cfDNA in the MP fraction, but not in soluble cfDNA (Fig. 6 C).

To test the prevalence of reactivity to DNAEL3-sensitive Ags on MPs in sporadic SLE, we analyzed a well-characterized cohort of SLE patients (n = 120), many of whom were used in the initial description of Ab binding to DNAEL3 described above. Of these patients, 87 (73%) had renal SLE and 62 (52%) had undergone longitudinal sampling. In total, 221 plasma samples were analyzed, revealing that 61 patients (51%) harbored DNAEL3-sensitive binding during at least one sample collection. The clinical characteristics of sporadic SLE patients organized by the presence of DNAEL3-sensitive binding are summarized in Table S1. As expected, no DNAEL3-sensitive binding was observed in any of the healthy controls analyzed (n = 15). Binding of plasma IgG to MP Ags could be prevented or reversed by the treatment of MPs with DNAEL3 before or after the incubation with patient plasma, respectively (Fig. 6 D). Importantly, patients with DNAEL3-sensitive binding showed a significant decrease of DNAEL3 activity in the plasma (Fig. 6 E) and a near-uniform increase in cfDNA in the MP fraction (Fig. 6 F). Furthermore, DNAEL3-sensitive binding was strongly associated with renal involvement: 55 of 87 (63%) renal patients compared with 5 of 33 (15%) nonrenal SLE patients (P < 0.0001; Fig. 6 G). Moreover, patients with DNAEL3-sensitive binding showed a significant enrichment in proteinuria, low complement levels (i.e., C3 and C4) levels, positive anti-dsDNA Abs (Fig. 6 H), and showed a higher median SLE disease activity index (SLEDAI) at the time of blood sampling (Fig. 6 I). In anti-Ro+ mothers, while 8 of 45 (18%) demonstrated IgG binding to MP Ags, this binding was sensitive to DNAEL3 in only three patients (Fig. 6 G). Two of these were the only subjects with renal SLE, with one patient harboring Abs to DNAEL3 (Fig. 2 A and Table 1).

To address whether DNAEL3-sensitive binding tracks with clinical response in patients with active LN, samples were obtained from 30 patients at the time of renal biopsy and longitudinally over a 52-wk period. Clinical response was defined at 52 wk as complete (n = 8, UPCR <0.5), partial (n = 3, UPCR reduced by 50% from baseline but >0.5), or none (n = 19). Detailed information on patient characteristics organized by treatment response is given in Table S2 and characteristics for each individual patient included in this analysis are shown in Table S3. At baseline, there was no significant difference in the prevalence of DNAEL3-sensitive binding between combined partial and complete responders (7/11, 63%) and nonresponders (13/19, 68%). However, at 52 wk, only 2/11 (18%) combined treatment responders still manifested DNAEL3-sensitive binding, while this reactivity remained persistent in all nonresponders (13/19, 68%; Fig. 6 J). Additionally, at baseline, there was no difference in the fraction of cfDNA located in MPs, but combined treatment responders demonstrated a shift of cfDNA toward the MP-depleted plasma fraction, while the distribution of cfDNA remained unchanged in nonresponders (Fig. 6 K). Collectively, these data suggest that Abs to DNAEL3-sensitive Ags on MPs inversely correlate with clinical response to treatment in patients with active LN.

Reactivity toward DNAEL3-sensitive Ags includes DNA-associated proteins

It was noted in evaluating the reactivity to DNAEL3-sensitive MP Ags that this did not fully mirror anti-DNA Abs. Specifically, anti-dsDNA IgG was present in >80% of patients with this reactivity but also in >50% of patients without it (Fig. 6 H). Moreover, only one of the three DNAEL3-deficient patients with DNAEL3-sensitive binding (Fig. 6 B) had anti-dsDNA Abs as assessed by ELISA (data not shown). To further explore the relationship between Abs to dsDNA and MPs, we examined a panel of 9G4+ mAbs derived from SLE patients. Abs marked by the 9G4 idiotype are prominent among SLE-associated auto-Abs and bind multiple Abs, including DNA and apoptotic cell surface.
determinants (Pugh-Bernard et al., 2001; Richardson et al., 2013; Tipton et al., 2015). Out of 20 examined 9G4+ mAbs, three showed DNASEIL3-sensitive binding (Fig. S5 A). Two of those also bound dsDNA and chromatin as expected; however, one (74C2) did not. Conversely, other clones (e.g., 88A1) reacted with dsDNA and chromatin by ELISA but did not bind MPs (Fig. S5 A). Collectively, these data suggest that reactivity to DNASEIL3-sensitive MP Ags partially overlaps with, but is distinct from, anti-dsDNA Abs and likely includes additional antigenic targets associated with DNA.

To further characterize reactivity to MP Ags, we explored the recognition of MP proteins by IgG from SLE patients using lysates of MPs harvested from apoptotic Jurkat T cells. SLE plasma samples that had shown DNASEIL3-sensitive IgG binding to MPs by flow cytomtery also demonstrated reactivity to multiple MP proteins by Western blot. Most prominently, reactivity to proteins of ∼10–15, 29, 35, 45, and 75 kD could be observed across multiple plasma samples (Fig. 7, A and B). We then took a candidate approach and explored the reactivity to histones and high-mobility group B1 (HMGB1) protein. High-mobility group proteins are structural transcription factors that can be incorporated into chromatin or released from activated or dying cells; HMGB1 in particular is an important mediator of inflammation and a common target of auto-Abs in SLE (Pisetsky, 2014a; Urbonaviciute et al., 2008; Wirestam et al., 2015). Both H2A/H2B and HMGB1 could be identified by Western blot in lysates of Jurkat cell MP, and bands of similar size (∼29 kD for HMGB1 and ∼13–15 kD for histones) were detected by SLE plasma samples with DNASEIL3-sensitive reactivity to MP Ags (Fig. 7 B). We next analyzed plasma IgG from Dnaseil3-deficient mice and observed IgG binding to proteins of similar size as those seen with human SLE plasma (Fig. 7 C). This reactivity was age dependent (Fig. 7 D), consistent with progressive autoreactivity and epitope spreading in older Dnaseil3-deficient mice (Sisirak et al., 2016).

Although Western blot analysis confirmed the reactivity of patient auto-Abs with multiple protein Ags in MPs, it did not establish whether this reactivity is DNASEIL3 sensitive. To this end, we used flow cytometry to show that HMGB1 was exposed on the surface of Jurkat cell-derived MPs and that this exposure was sensitive to DNASEIL3, but not DNASE1 (Fig. 7 E). Staining for H2A and H2B did not yield a distinct positive MP population (Fig. S5 B), likely due to the blockade of histone epitopes by the associated nucleosomal DNA on the surface of MPs. Taken together, these data demonstrate that reactivity to DNASEIL3-sensitive Ags on MPs targets not only DNA but also multiple proteins, potentially including known SLE-associated self-Ags such as HMGB1 and histones.

**Discussion**

In this study, we explored the role of DNase activity in SLE, focusing specifically on DNASEIL3, the extracellular DNase implicated in monogenic SLE. It was reported that total DNase activity may be reduced in SLE patients (Martinez-Valle et al., 2009; Sallai et al., 2005), but no correlation with disease severity was observed, and the specific DNase was not identified. More recently, a decrease in total DNase activity was observed specifically in nephritis patients (Bruschi et al., 2020), although the protein levels of both DNASE1 and DNASEIL3 appeared normal. Using a novel functional assay that is specific for DNASEIL3, we found that more than half of SLE patients with renal involvement show reduced activity of DNASEIL3 in the circulation. Given that DNASEIL3 nonredundantly protects from SLE and that a partial reduction of its activity in DNASEIL3 (R206C) carriers is associated with autoimmunity, its specific reduction in severe sporadic SLE likely contributes to the pathogenesis.

Reduced DNASEIL3 activity strongly correlated with the appearance of activity-neutralizing Abs to DNASEIL3, suggesting that DNASEIL3 is blocked by this newly described auto-Ab. Indeed, the measurement of anti-DNASEIL3 Abs by ELISA provides a robust high-throughput assay to estimate DNASEIL3 activity in sporadic SLE patients in future studies. This mechanism does not rule out other possible scenarios such as reduced DNASEIL3 protein levels, a possibility that could not be explored here due to the lack of sufficiently specific protein detection reagents. Ab-mediated blockade is consistent with the results of Bruschi et al. (2020), who implicated inhibitory circulating proteins in the reduction of total DNase activity in renal SLE. Notably, auto-Abs preferentially bound DNASEIL3 compared with DNASE1 and thus appear distinct from previously described DNASE1 inhibitory substances (Hakkim et al., 2010) or polyreactive anti-DNASE1 Abs that also bind DNA (Puccetti et al., 1995; Yeh et al., 2003). The observed Ab-mediated inhibition of DNASEIL3 resembles a similar inhibition of the complement component C1Q. Like DNASEIL3, C1Q is required for self-tolerance, as null C1Q mutations in humans cause SLE with a near-complete penetrance (Leffler et al., 2014). In addition, C1Q can itself be targeted by a specific auto-Ab that reduces C1Q levels in HUVS (Jara et al., 2009) and is also found in sporadic SLE (Orbai et al., 2015). The inhibition of DNASEIL3 by specific auto-Abs appears to recapitulate its genetic deficiency by nongenetic means, emphasizing the critical role of DNASEIL3 in tolerance and its relevance for sporadic SLE.

Self-DNA that may activate innate or adaptive immunity in SLE may comprise inflammation-associated forms such as neutrophil extracellular traps and oxidized mitochondrial DNA (Caielli et al., 2016; Garcia-Romo et al., 2011; Lood et al., 2016) or cfDNA that exists in the steady state. Given the observed effects of DNASEIL3 on the latter (Serpas et al., 2019), we explored its physical nature in circulation, including the association with MPs. We confirmed the presence of DNA-carrying MPs in normal plasma and established that ∼25% and >90% of total cfDNA is MP associated in humans and mice, respectively. The reasons for this difference between the species are unknown but may involve the shorter time and route of blood circulation in mice, reducing the lifespan of soluble cfDNA. Importantly, MP-associated cfDNA was enriched in longer polynucleosomal DNA fragments, consistent with the detection of such fragments in MPs generated in vitro (Reich and Pisetsky, 2009). Fragments of naked DNA within the 100–4,000 bp size range were shown to bind anti-DNA Abs equally well in solution (Pisetsky and Reich, 1994). We confirmed these observations but showed that for nucleosome-bound DNA, longer polynucleosomal fragments bind auto-Abs with higher affinity. We also demonstrated...
the superior capacity of polynucleosomal fragments as inducers of IFN-I production. These data are consistent with the critical role of IFN-I in SLE (Elkon and Wiedeman, 2012) and with its contribution to autoreactivity and disease in DNASE1L3-deficient mice (Soni et al., 2020). Overall, our results highlight the abundance of MP-associated cfDNA and its potential as a self-Ag that requires tight control in vivo.

The MP-associated fraction of cfDNA was increased in DNASE1L3-deficient mice and humans and correlated with reduced DNASE1L3 activity in sporadic SLE patients with nephritis. This was in striking contrast to comparable levels of total plasma cfDNA, which did not show significant differences between patient groups in our study. Although increased total cfDNA has been reported in SLE patients compared with controls (Duvvuri and Lood, 2019), this difference showed no clear association with disease activity or with LN specifically (Atamaniuk et al., 2011; Xu et al., 2018). Sequencing of total cfDNA in SLE revealed certain abnormalities such as aberrant...
genome representation and shortening due to the presence of anti-dsDNA Abs, but no consistent features serving as biomarkers of the disease (Chan et al., 2014). Our results highlight the relevance of MP-associated cfDNA for SLE pathogenesis, given its striking increase in LN relative to healthy and non-LN patients. Moreover, they confirm DNASE1L3 as an essential and specific regulator of MP-associated cfDNA, which therefore represents the physiological substrate of this enzyme. They also help explain the lengthening of total cfDNA in the plasma of Dnase1l3-deficient mice (Serpas et al., 2019) and humans (Chan et al., 2020), which apparently reflects the accumulation of MP-associated polynucleosomal fragments. Thus, auto-Ab-mediated impairment of DNASE1L3 activity in renal SLE patients results in the accumulation of a self-Ag (i.e., MP-associated cfDNA and affiliated proteins).

Auto-Ab from SLE patients were shown to bind endogenous MPs (Nielsen et al., 2012) or in vitro–generated MPs (Ullal et al., 2011), yet the nature of bound Ag and clinical significance of this reactivity remained unclear. We report that Abs to MPs are prevalent in patients with SLE and target DNASE1L3-sensitive Ag on the MP surface. We also show that this Ab reactivity is associated with reduced DNASE1L3 activity and cfDNA accumulation into MPs, likely representing a consequence of these events. Unlike ELISA-based assays for Abs to DNA and other Ags, the binding to DNASE1L3-sensitive Ag on MPs is a physiological interaction that is unbiased with respect to a specific Ag. As such, it represents an integral readout that correlates but does not directly overlap with any single reactivity, including the canonical anti-dsDNA ELISA. This likely explains why reduced DNASE1L3 activity and/or DNASE1L3-targeting Abs did not correlate with anti-dsDNA (Fig. S2 D and data not shown) but were associated with reactivity to MPs (Fig. 6 E). Indeed, we demonstrate that the reactivity to MPs involves a set of protein Ag carried by MPs, with reactivity patterns shared among different patients and even with Dnase1l3-deficient mice. One established DNASE1L3-sensitive Ag is the nucleosome (Siirak et al., 2016), consistent with its exposure on the MP surface (Ullal et al., 2011) and the presence of histones in MPs (this study); however, pure histones may not be targeted independently of the DNA. Another candidate DNASE1L3-sensitive Ag is HMGB1, a pleiotropic DNA-binding protein that is a common target of auto-Ab in SLE (Harris et al., 2012; Urbanoviciute et al., 2007). The exposure of HMGB1 on apoptotic MPs has been described (Pisetsky, 2014b), and here, we demonstrate that this phenomenon is controlled by DNASE1L3, presumably by digesting HMGB1–DNA complexes. Thus, by digesting DNA in MPs, DNASE1L3 controls the exposure of not only DNA but also multiple DNA-associated proteins, which are targeted by auto-Ab in SLE patients with reduced DNASE1L3 activity.

In conclusion, our data delineate the molecular basis of tolerogenic DNASE1L3 enzymatic activity and describe its impairment in some patients with SLE, most notably those with kidney involvement. Conceptually, our data reveal a novel mechanism of SLE pathogenesis based on auto-Ab targeting a critical “gatekeeper” of tolerance to self-DNA. The inhibition of DNASE1L3 activity by such auto-Ab recapitulates the genetic lesion of DNASE1L3-deficient patients by nongenetic means, emphasizing the key role of DNASE1L3 in sporadic SLE. Technically, our work introduces four novel readouts (DNASE1L3 activity, Abs to DNASE1L3, fraction of MP-associated cfDNA, and Abs to DNASE1L3-sensitive Ag) that correlate with each other and with active LN. Of these assays, IgG binding to DNASE1L3-sensitive Ag on MPs is a relatively simple, robust, high-throughput assay whose readout showed correlation with LN in both cross-sectional and longitudinal samples. Finally, the reduction of DNASE1L3 activity in SLE patients with nephritis warrants further exploration of DNASE1L3 as a potential therapeutic in this patient group. Our results provide an important caveat to such attempts by describing neutralizing Abs to DNASE1L3; nevertheless, the rationale for the therapeutic use of this secreted enzyme that is fundamentally linked to SLE pathogenesis appears compelling.

Materials and methods

Human subjects

The study population was composed of four subgroups: DNASE1L3-deficient patients (n = 4), the New York University (NYU) SLE cohort (n = 120), anti-Ro* mothers of children with neonatal lupus (n = 45), and healthy controls (n = 15). The NYU SLE cohort was further stratified into SLE patients with (n = 87) and without (n = 33) renal involvement according to the Systemic Lupus Erythematosus International Cooperating Clinics criteria (Petri et al., 2012).

Sporadic SLE and healthy controls

Blood specimens were obtained from several sources. Patients with SLE and healthy controls older than 18 yr of age were enrolled in the NYU SAMPLE (Specimen and Matched Phenotype Linked Evaluation) Biorepository after signing informed consent approved by the NYU School of Medicine institutional review board. SLE was assigned using classification criteria as defined by one or more of the following: (1) 1997 Revised American College of Rheumatology (ACR; Hochberg, 1997; Tan et al., 1982), (2) Systemic Lupus Erythematosus International Cooperating Clinics (Petri et al., 2012), or (3) the most recent ACR/European League Against Rheumatism (Aringer et al., 2019). Plasma for the study of SLE patients was selected randomly to represent the full spectrum of clinical disease. All SLE patients were evaluated for the status of disease activity using a modification of the original Safety of Estrogens in Lupus National Assessment (SELENA)–SLEDAI (Petri et al., 2005) in which proteinuria was scored 4 points only when the UPCR increased by more than 0.5 from the previous visit. The modification applied in this study used the SLEDAI 2K definition of proteinuria in which a UPCR of >0.5 is always scored 4 points (Gladerman et al., 2002). All other domains of the SELENA-SLEDAI were scored as originally defined. For the longitudinal follow up, a subset of SLE patients undergoing a kidney biopsy to evaluate a UPCR ratio of >0.5 (considered abnormal) was included in the study. A renal response at 52 wk was considered complete if the UPCR was <0.5 and a partial response if the baseline UPCR was reduced by 50% but did not resolve to <0.5.
Anti-Ro+ mothers of children with neonatal lupus

A second disease cohort evaluated were women enrolled in the Research Registry for Neonatal Lupus (RRNL; Buyon et al., 1998) after signing informed consent for participation approved by the NYU School of Medicine institutional review board. All participating mothers in the RRNL included in this study have a child with congenital heart block and/or characteristic rash and have Abs to at least one component of the Ro/La ribonucleoprotein complex, including 52-kD Ro, 60-kD Ro, or 48B/La, as confirmed by ELISA in the laboratory of R.M. Clancy and J.P. Buyon (Reed et al., 2012). Many of the women in the RRNL do not have any systemic disease and are only known to have anti-Ro Abs because of the workup for their child’s illness. Some mothers have an undifferentiated autoimmune disease, as they have insufficient criteria for SLE or Sjögren’s syndrome (Shiboski et al., 2017; Vitali et al., 2002) and less than half ever progress to SLE and/or Sjögren’s syndrome (Rivera et al., 2009). In this group of RRNL women, three had developed LN.

DNASE1L3-deficient patients

One patient with pediatric HUVS with renal involvement progressing to end-stage renal disease was diagnosed and treated at the Istituto Gaslini. Collection of patient samples was approved by the ethics committee of the Istituto Giannina Gaslini (approval BIOL 6/5/04). Molecular analysis by next-generation sequencing was performed in the Laboratory of Neurogenetics and Neuroinflammation at the Imagine Institute for Genetic Diseases, Paris. The patient was homozygous for the c.289-291delCA DNASE1L3 mutation, which was confirmed previously (Batu et al., 2018; Carbonella et al., 2017; Ozcakar et al., 2013) and is predicted to cause premature termination (p.Thr97Ilefs*2). Three patients with pediatric HUVS (age of onset, 9–14 yr) were diagnosed and treated at the Istituto di Ricovero e Cura a Carattere Scientifico Ospedale Pediatrico Bambino Gesù. Sample collection from patients of Ospedale Pediatrico Bambino Gesù was approved by the Institutional Ethical Committee number 1666_OPBG_2018. All patients had chronic urticarial vasculitis, hypocomplementemia, positive anti-CIQ Ab, negative antinuclear Abs, and negative anti-dsDNA by laboratory testing; one of the three showed positive anti-dsDNA by in-house ELISA. The patients were subjected to targeted resequencing on the Illumina NextSeq platform for genes associated with interferonopathies. All patients were homozygous for the c.289-291delCA DNASE1L3 mutation, which was reported previously (Belot, 2020) and is predicted to cause the same premature termination as the mutation above. All patients and/or their families signed informed consent for participation approved by the ethics committees at their respective institutions.

Animals

All animal studies were performed according to the investigator’s protocol approved by the Institutional Animal Care and Use Committee of NYU School of Medicine. Dnase1l3−/− mice on C57BL/6 (B6) background (Dnase1l3<sup>3lacZ/3lacZ</sup>) were described previously (Sisirak et al., 2016). Mice carrying a null allele of Dnase1 (Dnase1<sup>−/−</sup>) on B6 background were obtained from the Knockout Mouse Project, crossed with WT B6 mice and intercrossed to obtain Dnase1<sup>−/−</sup> mice. Mice with doxycycline-inducible ubiquitous expression of H2B-GFP fusion protein (B6; 129S4-Gt(ROSA)26Sor<sup>tm1.1(KOMP)Vlcg</sup>) were obtained from The Jackson Laboratory and intercrossed. The resulting mice heterozygous for both alleles were given oral doxycycline in water for at least 4–6 wk to induce the expression of H2B-GFP as described (Foudi et al., 2009). WT control mice of B6 background were bred in the same animal facility or purchased from Taconic and maintained in the same facility. All mice were age and sex matched when comparing groups. Male and female WT mice (57% female and 43% male) between 30 and 450 d of age (median 80 d) were analyzed (Fig. 4 C); no difference in the concentration or distribution of cfDNA was observed between mice of different age or sex. Dnase1l3-deficient males younger than 60 d were used in order to avoid any confounding effect of SLE-like disease that develops starting at 3 mo (Fig. 4 D). 1-yr-old Dnase1l3-deficient females were used in Fig. 4 E. These mice do not develop any autoreactivity at any age, and thus the age is not expected to impact our findings.

DNASE1L3 protein purification

Preliminary studies showed that the introduction of a 6-aa FLAG epitope into a predicted internal loop of the human full-length DNASE1L3 did not affect the activity of DNASE1L3, including its specific ability to digest chromatin. The resulting FLAG-DNASE1L3 open reading frame was cloned into a lentiviral vector, which was then used to transduce 293T cells followed by batch selection of stable transfectants. FLAG-DNASE1L3-expressing 293T cells were cultured in 150-mm dishes for 7 d, and the supernatants were incubated with magnetic anti-FLAG M2 beads (Sigma-Aldrich) overnight at 4°C. The beads were collected, washed, and incubated with 0.25 mg/ml FLAG peptide, and the eluted protein was quantified by SDS-PAGE with BSA as a standard.

Isolation of MPs

**Human plasma**

To obtain the MP fraction as well as the MP-depleted (soluble) fraction of human plasma, blood was collected in tubes containing EDTA (Becton Dickinson; 366643), and blood cells were removed by centrifugation at 1,400 g for 12 min at 4°C. A second centrifugation step (1,400 g for 12 min) was used to remove platelets. The resultant plasma (either fresh or stored at ~80°C) was centrifuged at 22,000 g for 60 min to pellet the MP. The supernatant and MP pellet were used to measure cfDNA in the MP-depleted (soluble) and MP plasma fractions, respectively.

**Murine plasma**

Mice were anaesthetized by i.p. injection of 100 mg/kg ketamine/10 mg/kg xylazine in sterile PBS. Plasma was isolated by cardiac puncture using a 26G syringe and transferred into EDTA-coated tubes (Kent Scientific; 41.1395.105). Blood was centrifuged at 1,000 g for 10 min, and the supernatant was kept on ice. For DNA quantification, the plasma was spun a second time at 1,000 g for 10 min. For confocal imaging studies, the...
plasma was spun a second time at 10,000 g for 10 min to remove platelets. After the second spin, the supernatant was centrifuged at 21,130 g for 60 min at 4°C to pellet MPs. The supernatant and MP pellet were used to measure cfDNA in the MP-depleted (soluble) and MP plasma fractions, respectively.

**Quantification of cfDNA**

To extract cfDNA from the MP fraction and MP-depleted fraction, samples were centrifuged as described above. Circulating cfDNA was extracted from 50–400 µl human plasma or 150–200 µl murine plasma using the QIAamp DNA Blood Mini Kit (QIAGEN). The amount of DNA in the respective samples was measured by qPCR using intra-Alu human primers or *inter-B1* mouse primers. All qPCRs were performed in 20 µl using Fast-Start Universal SYBR Green Master mix (Sigma-Aldrich 4913914001). Briefly, the method for intra-Alu human primers was as follows: initial denaturation of 12 min at 95°C followed by 45 amplification cycles of denaturation at 95°C for 20 s, annealing at 56°C for 1 min, and elongation at 72°C for 1 min (Walker et al., 2003). The mouse *inter-B1* qPCR methods have been reported previously (Zhang et al., 2010). Briefly, each reaction was subjected to an initial denaturation of 8 min at 95°C followed by 40 amplification cycles of denaturation for 30 s at 95°C, annealing for 40 s at 55°C, and extension for 1 min at 72°C.

**Estimation of cfDNA lengths by qPCR**

Two sets of primers and the respective qPCR programs described by Lou et al. were used to amplify 76 bp (short) or 200 bp (long) of the human Alu repeat sequences (Lou et al., 2015). The concentration of cfDNA as determined by the 76-bp amplicons and 200-bp amplicons were calculated, and subsequently the ratio of long to short amplicon products was determined as a measure of cfDNA length in the MP fraction and MP-depleted (soluble) plasma fraction.

**cfDNA sequencing in human patients**

Plasma from four healthy controls (two males and two females) was isolated and centrifuged to separate MP and MP-depleted fractions as described above. DNA was purified from all samples using the QIAamp DSP DNA Blood Mini Kit (QIAGEN; 61104) and quantified by qPCR. DNA libraries were prepared using the KAPA HT Library Preparation Kit (Roche) and purified using AMPure XP purification beads (Beckman Coulter) according to the manufacturer’s instructions. Adaptor-ligated libraries were analyzed on an Agilent 4200 TapeStation (Agilent Technologies) using the High Sensitivity D1000 ScreenTape System (Agilent Technologies) for quality control and quantified by Qubit (Invitrogen). The libraries were sequenced for 150 bp for each end in a paired-end format on a Hiseq 4000 or the equivalent Illumina sp300 flow cell.

**DNA sequencing analysis**

The adaptor sequences were trimmed using cutadapt (v1.18), and all of the reads from the sequencing experiment were mapped to the reference genome (hg19/GRCh37.75) using Bowtie2 (v2.2.4). Duplicate reads were removed using Picard tools (v.1.126), and the low-quality mapped reads (mapping quality <20) were removed from the analysis. The read per million normalized BigWig files were generated using BEDTools (v.2.17.0) and the bedGraphToBigWig tool (v.4). The insert sizes and their SDs were calculated using Picard tools and used to generate cfDNA length traces shown. Frequencies were calculated by dividing the number of reads from each insert size by the total number of reads acquired per sample. All downstream statistical analysis was performed in the R environment (v3.1.1). Sequence data have been deposited in the Genotypes and Phenotypes database hosted by the National Institutes of Health (BioProject accession no. PRJNA695527).

**Assessment of DNASE1L3 activity**

**Digestion of DNA substrates with DNases**

DNA substrates were incubated with different concentrations of human DNASE1 (Abcam; 73430) or human recombinant DNA-SEIL3 in the presence of 2 mM CaCl2/MnCl2 and a total volume of 10 µl for 15 min at 37°C. All dilutions of enzymes were prepared in HBSS (without magnesium and calcium) with 100 ng/ml molecular-grade BSA. Following digestion, the reactions were immediately kept on ice. The amount of remaining DNA was measured by qPCR using intra-Alu human primers as follows: initial denaturation of 10 min at 98°C followed by 45 amplification cycles of denaturation at 95°C for 5 s, annealing at 60°C for 5 s, and elongation at 72°C for 15 s. The following DNA substrates were used: 200 ng naked Jurkat DNA (Thermo Fisher) or nuclei purified from 15,000 Jurkat T cells as described previously (Nabbi and Riabowol, 2015). The digestion of nuclei was used to assess specific activity of plasma DNASE1L3, while the digestion of naked DNA was used to assess total plasma DNase activity. To assess whether plasma IgG could bind and inhibit enzymatic DNase activity, IgG was purified from healthy controls and three SLE patients with Abs to DNASE1L3 using Protein G beads. Different concentrations of DNases were preincubated with 0.1 mg/ml purified IgG at 37°C for 30 min. Preincubated DNase and untreated DNase were used to digest DNA substrates as described and the amount of remaining DNA was measured by qPCR. The DNase concentration at which 50% of the input DNA was digested (IC50) was established, and the percentage of DNase activity was compared with untreated DNase was calculated.

**Quantitative DNASE1L3 activity assay**

Several dilutions of human plasma were incubated with 15,000 Jurkat T cell-derived nuclei and 2 mM CaCl2/MnCl2 in a total volume of 10 µl for 10 min at 37°C. All plasma samples were diluted in HBSS (without magnesium and calcium) with 100 ng/ml molecular grade BSA. The remaining DNA was measured by qPCR using intra-Alu human primers as described above. The DNase concentration at which 50% of the input DNA was digested (IC50) was established, and the percentage of DNase activity was determined, considering the IC50 value for pooled healthy control plasma as 100% and the value for DNASE1L3-deficient patient plasma as 0%.

We have found that total reaction volume, digestion time, digestion solution, batch of nuclei, CaCl2/MnCl2 concentration, plasma diluent, and polymerase used for qPCR-based DNA
quantification can affect the results of this assay. Detailed protocols and troubleshooting tips will be shared with all interested investigators upon request.

ELISA

**Anti-DNase ELISA**

All plate incubations were done in a hydration chamber unless otherwise specified. Nunc MaxiSorp plates (Invitrogen; 44–2404–21) were coated with 50 µl/well of 2 µg/ml recombinant human DNASE1 (Abcam; 73430), purified recombinant DNASEI13, or FLAG peptide (Sigma-Aldrich) in PBS overnight at 4°C. Ag-coated plates were washed with PBS and blocked with 250 µl/well of PBS with 4% nonfat dry milk (NFDM) for 2–3 h at room temperature. Plates were then washed again three times with PBS and incubated with plasma diluted in PBS overnight at 4°C. Plates were then washed three times with PBS containing 1% NFDM, and bound IgG was detected using goat anti-human IgG–alkaline phosphatase conjugate (Sigma-Aldrich; A1543-1ML) diluted 1:5,000 in PBS containing 1% NFDM. The plate was then washed off with PBS+1% NFDM, and the plate was developed as described above.

**Anti-dsDNA ELISA**

Nunc MaxiSorp plates were coated with 0.01% poly-L lysine (Sigma-Aldrich) for 1 h at room temperature, washed with PBS, and coated with 50 µl/well of 10 µg/ml calf thymus DNA (Calbiochem) in PBS overnight at 4°C. Ag-coated plates were washed and then blocked with 200 µl/well of PBS containing 4% NFDM for 3 h at 4°C. Blocked plates were washed once with PBS + 1% NFDM, coated with diluted serum in PBS, and incubated overnight at 4°C. Unbound serum Abs were washed off with three washes of PBS + 1% NFDM. The bound IgG was detected using a goat anti-human IgG–alkaline phosphatase conjugate (Sigma-Aldrich; A1543-1ML) diluted 1:5,000 in PBS containing 1% NFDM. The plate was then washed off with PBS+1% NFDM, and the plate was developed as described above.

**Anti-dsDNA ELISA with DNA fragments of different length**

Nunc MaxiSorp plates were coated with 0.01% poly-L lysine (Sigma-Aldrich) for 1 h at room temperature, washed with PBS, and coated with 50 µl/well of 10 µg/ml calf thymus DNA (Calbiochem) in PBS overnight at 4°C. DNA fragments of 95 bp, 186 bp, 382 bp, and 831 bp were prepared by PCR and purified using QIAquick Gel Extraction Kit (QIAGEN). Different concentrations of PCR-amplified DNA fragments were prepared separately and mixed with an equal volume of diluted plasma. This solution was resuspended several times to mix, and 50 µl was immediately added to a freshly blocked dsDNA-coated plate. The plate was then processed as described above.

**Competition ELISA with nucleosomes**

Anti-dsDNA plates were prepared as described above. During the plate blocking, plasma and inhibitor solutions were prepared separately in PBS. Each plasma sample was prepared at a dilution that yielded an OD405nm of 0.5–0.6 so that we could assess both increases and decreases in the OD. Different concentrations of purified HeLa mononucleosomes (EpiCypher; 16–0002) and purified HeLa polynucleosomes (EpiCypher; 16–0003) were prepared separately and mixed with an equal volume of diluted plasma. This solution was resuspended several times to mix, and 50 µl was immediately added to a freshly blocked dsDNA-coated plate. The plate was then processed as described above.

**Competition ELISA with DNases**

Anti-dsDNA plates were prepared as described above. During the plate blocking, plasma and inhibitor solutions were prepared separately in PBS. Each plasma sample was prepared at a dilution that yielded a DNASEI3 ELISA OD405nm of 0.5–0.6 so that we could assess both increases and decreases in the OD. DNASEI3 plates were prepared as described above. During the plate blocking, plasma and inhibitor solutions were prepared separately in PBS. Each plasma sample was prepared at a dilution that yielded an OD405nm of 0.5–0.6 so that we could assess both increases and decreases in the OD. Different concentrations of recombinant human DNASEI (Abcam; 73430) or purified recombinant DNASEI13 were prepared separately and mixed with an equal volume of diluted plasma. This solution was resuspended several times to mix, and 50 µl was immediately added to a freshly blocked plate coated with 2 µg/ml DNASEI3. The plate was then processed as described above.

**Competition ELISA with DNA fragments of different length**

Anti-dsDNA plates were prepared as described above. During the plate blocking, plasma and inhibitor solutions were prepared separately in PBS. Each plasma sample was prepared at a dilution that yielded an OD405nm of 0.5–0.6 so that we could assess both increases and decreases in the OD. DNA fragments of 95 bp, 186 bp, 382 bp, and 831 bp were prepared by PCR and purified using QIAquick Gel Extraction Kit (QIAGEN). Different concentrations of PCR-amplified DNA fragments were prepared separately and mixed with an equal volume of diluted plasma. This solution was resuspended several times to mix, and 50 µl was immediately added to a freshly blocked dsDNA-coated plate. The plate was then processed as described above.

**Western blot**

**IgG binding against Ags on MPs**

MP proteins from 2–5 × 10⁷ Jurkat-derived MPs were separated on a 4–12% Bis-tris Western blot gel (Invitrogen; NP0326BOX) in reducing conditions and transferred to a polyvinylidene fluoride microporous membrane (Sigma-Aldrich; PVH00005). Afterwards, the membrane was blocked in 5% BSA for 60 min at room temperature and incubated overnight with plasma diluted in blocking buffer. In some instances, incubation with plasma samples was performed using a multiplex blotting system (LiCOR Biosciences). After washing with TBS with 0.1% Tween-20, membranes were incubated with HRP-conjugated anti-human IgG Ab (Jackson Immunoresearch; 1:5,000) or anti-mouse IgG Ab (eBioscience; 1:1,000) diluted in blocking buffer at room temperature for 60 min. The immunoreactive bands were visualized using an Amersham Imager 680 blot and gel imager.
IgG binding to DNases

To assess IgG binding to DNases by Western blot, 2 µg recombinant DNASEI3 (purified in our laboratory) or 2–40 µg recombinant DNASEI (Abcam; 73430) was run on a 12.5% separating gel in reducing conditions, and Western blot was performed as above.

IgG binding to HMGB1 on MPs

MP proteins from Jurkat-derived MPs were separated on a 4–12% Bis-tris Western blot gel. A polyclonal anti-HMGB1 Ab (Invitrogen; PA5–96160) was used as primary Ab (1:1,000 dilution) and detected with an HRP-conjugated anti-rabbit IgG Ab (Thermo Fisher; G-21234; 1:2,000 dilution).

IgG binding to histones on MPs

MP proteins from Jurkat-derived MPs were separated on a 4–12% Bis-tris Western blot gel. A polyclonal anti-H2A Ab (Cell Signaling Technology; 2578S) or a polyclonal anti-H2B Ab (Cell Signaling Technology; 8135S) was used as primary Ab (1:1,000 dilution) and detected with an HRP-conjugated anti-rabbit IgG Ab (Thermo Fisher; G-21234; 1:2,000 dilution).

Flow cytometry

Analysis of DNASEI3-sensitive IgG binding to MPs

MPs from Jurkat cells were generated as previously described (Sisirak et al., 2016; Ullal et al., 2011). Briefly, Jurkat cells were cultured in the presence of 1 µM staurosporine (Sigma-Aldrich; S440) overnight, harvested, and collected by centrifugation for 5 min at 400 x g. The supernatants were collected and centrifuged at 22,000 g for 30 min at 4°C to pellet MPs. Surface staining of MPs was performed as follows: 10⁵ MPs were mock-treated or treated with recombinant human DNASE1 (Abcam; 73430) or recombinant DNASEI3. Where specified, MPs were treated with DNASEI and DNASEI3-containing supernatants, as described previously (Sisirak et al., 2016), rather than recombinant proteins. DNase-treated MPs were then stained with human plasma at 1:20 dilution in a total volume of 30 µl at 4°C for 20 min. The supernatants were collected and centrifuged at 22,000 g for 30 min at 4°C, and incubated with a secondary Alexa Fluor® 488-conjugated anti-human IgG Ab (Millipore; A11001; 1:1,000). Samples were then diluted to stop staining and acquired on an Attune Nxt (Thermo Fisher) flow cytometer. Representative histograms were generated using FlowJo software (Tree Star).

To determine whether plasma IgG binds to the surface of native MPs, the respective histograms were subtracted from a control histogram. The control histogram was generated by merging histograms derived from staining native MPs with plasma from healthy controls (n = 10). To determine whether IgG binding to native MPs occurred in a DNASEI3-sensitive manner, the histogram derived from staining DNASEI3-treated MPs was subtracted from the histogram derived from staining native MPs for each plasma sample using the Overton subtraction method (Overton, 1988) as implemented in FSC Express (De Novo Software).

Analysis of MP Ags

3–4 x 10⁵ MPs were mock-treated or treated with recombinant human DNASE1 (Abcam; 73430) or recombinant DNASEI3. DNase-treated MPs were then diluted in HBSS without magnesium, calcium, or phenol red (Thermo Fisher; 14–175–095). They were centrifuged at 22,000 g for 30 min at 4°C to pelleted MPs and resuspended in HBSS (for Vybrant Green staining) or PBS for other staining.

Vybrant Green. MPs were stained in 400 µl HBSS + Vybrant Green (Invitrogen; V35004, 1:10,000) for 15 min at 37°C. The sample was then kept on ice in the dark and run on an Attune Nxt (Thermo Fisher) flow cytometer within 60 min.

HMGB1 or histones. MPs were stained in 50 µl PBS for 10 min at 4°C with HMGB1 (Invitrogen; PA5–96160, 1:50), H2A monoclonal (Cell Signaling Technology; 12349S, 1:50), H2A polyclonal (Cell Signaling Technology; 2578S, 1:50), or H2B (Cell Signaling Technology; 8135S, 1:50). After staining, samples were diluted with PBS and centrifuged at 22,000 g for 30 min at 4°C to pellet MPs. They were then stained with AFS647-conjugated anti-rabbit IgG (Invitrogen; A21245, 1:500) for 45 min at 4°C. Samples were then diluted to 400 µl with PBS and run on an Attune Nxt (Thermo Fisher) flow cytometer. MP gating was calibrated based on 1-µm sizing beads (Invitrogen).

Stimulation of human pDCs

Approximately 80 ml of blood was drawn from healthy controls into EDTA-coated tubes (BD Biosciences; 366643). The blood was then split equally between three 50-ml conical tubes and diluted to 40 ml with sterile PBS. 10 ml Ficol-Faque (GE Healthcare; 17–1440–03) was added to the bottom of the tube using a glass Pasteur pipette. The cells were spun at 400 g for 22 min with low acceleration and no breaks at room temperature. Approximately 20 ml supernatant was aspirated, and the remaining top two layers were transferred into new conical tubes. The cells were spun at 200 g for 15 min at room temperature and washed three times with sterile wash buffer (PBS, 2% FCS, and 1 mM EDTA). The cells were then counted using an Accuri C6 flow cytometer (BD Biosciences) and diluted to a density of 50,000 cells/µl.

pDCs were enriched from total peripheral blood mononuclear cells using EasySep Human Plasmacytoid DC Enrichment Kit (Stemcell; 19062A). After enrichment, the cells were immediately resuspended in full RPMI and transferred into a 96-well plate. 5,000 pDCs were added to each well in 180 µl of full RPMI, and Ag (e.g., mononucleosomes and polynucleosomes) was added in a volume of 20 µl for a total reaction volume of 200 µl. The cells were activated for 24 h at 37°C and 5% CO₂. Following activation, the plate was spun at 1,300 rpm for 10 min. The supernatant was used immediately or stored at −80°C.

Ag preparation

Purified HeLa mononucleosomes (EpiCypher; 16–0002) and purified HeLa polynucleosomes (EpiCypher; 16–0003) were incubated with LL-37 (InvivoGen; tlr-l37) for 30–40 min at room temperature in a 20-µl volume and then immediately added to purified pDCs. The pDC/Ag mixture was then resuspended several times without vortexing. LL-37 was incubated with DNA substrates at a concentration of 500 µg/ml, and this mixture was added to pDCs for a final concentration of 50 µg/ml DNA. Mononucleosomes and polynucleosomes were incubated with LL-37 at concentrations between 60 and 20 µg/ml DNA, and the final
concentrations after adding this mixture to pDCs were between
6 and 2 µg/ml. Full RPMI contained 10% FCS, 100 U/ml penicillin
and 100 µg/ml streptomycin (Gibco; 15140122), 1× MEM non-
essential amino acids solution (Gibco; 11140050), 2 mM
L-glutamine (Gibco; 25030081), 1 mM sodium pyruvate (Gibco;
11360070).

Confocal microscopy of MPs
For human MPs, blood was drawn into EDTA-coated tubes (BD Biosciences; 366643) and centrifuged at 1,000 g for 10 min. The
supernatant was then centrifuged at 10,000 g for 10 min to
isolate platelet-free plasma. Platelet-free plasma was centrifuged
at 21,130 g for 60 min at 4°C to pellet MPs. Murine MPs were
isolated as described in the Isolation of MPs section of Materials
and methods. MP pellets were resuspended in HBSS without
magnesium, calcium, or phenol red (Thermo Fisher; 14–175-095),
and 1–2 million MPs were used for staining. Samples were
stained with Vybrant DiD Cell Labeling Solution (Invitrogen;
V22887) and Vybrant Green (Invitrogen; V35004) in the dark
at 37°C for 20 min. They were diluted to 1.5 ml with HBSS to stop
the staining. The MPs were filtered through a 70-µm filter and
spun at 21,130 g for 60 min at 4°C. The pellet was washed again
with HBSS, filtered through a 70-µm filter, and spun again
21,130 g for 60 min at 4°C. The pellet was resuspended in 1 µl
HBSS and transferred to a 1.5 coverslip (Corning; 2850–22).
The sample was kept in a hydration chamber for 40 min and in-
verted onto a Poly-Prep slide (Sigma-Aldrich; P0425–72EA)
containing one drop of Diamond Antifade Mountant (Thermo
Fisher; P36961). Slides were dried for 60 min at room temper-
ature and stored at 4°C until imaged. The imaging of H2B-GFP/
Fisher; P36961). Slides were dried for 60 min at room temper-
ature and stored at 4°C until imaged. The imaging of H2B-GFP/
Fisher; P36961). Slides were dried for 60 min at room temper-
ure and stored at 4°C until imaged. The imaging of H2B-GFP/
Fisher; P36961). Slides were dried for 60 min at room temper-

Online supplemental material
Fig. S1 shows the correlation of DNASE1L3 activity with renal
function in SLE patients. Fig. S2 presents further characteri-
zation of auto-Abs to DNASE1L3 in SLE patients. Fig. S3 shows
the correlation between the MP fraction of plasma cfDNA and
anti-dsDNA Ab levels in SLE patients. Fig. S4 shows the im-
 pact of length of protein-free DNA on its binding to anti-
dsDNA IgG. Fig. S5 further characterizes auto-Ab reactivity
to MP Ags. Table S1 describes clinical characteristics of spo-
radic SLE patients with or without DNASE1L3-sensitive
binding. Table S2 summarizes clinical characteristics of SLE
patients organized by treatment response. Table S3 describes
characteristics for each individual patient included in the
analysis from Table S2.

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Y. Wang, O.A. Perez, B. Sally, V. Sisirak, C. Soni, and
G.J. Silverman performed experiments and analyzed and in-
terpreted results. A. Khodadadi-Jamayari, A. Tsirigos, and M.Y.
Kim analyzed results. A.S. Chida and I. Sanz performed experi-
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Statistical analysis
Statistical analyses were performed using Prism software ver-
sion 8 (GraphPad). Normal distribution of data were not as-
sumed unless otherwise specified. Statistical significance
between two unpaired experimental groups was determined by
nonparametric Mann-Whitney test unless otherwise specified.
For paired groups, normality was assessed using the Shapiro-
Wilk test. If the data were normally distributed, a parametric
paired t test was used. If the data were not normally distributed,
a nonparametric Wilcoxon matched-pairs signed rank test was
used. The nonparametric Kruskal–Wallis test followed by the
Dunn’s multiple-comparison test was used for statistical analysis
of more than two groups. Contingency tables were analyzed
using Fisher’s exact test or χ² test as appropriate. Correlation
between two continuous variables was explored by Spearman
correlation. All P values were two tailed, and differences were
considered significant for P values < 0.05 (*), < 0.01 (**), < 0.001
(***), and < 0.0001 (****).

Primer sequences
The primer sequences were intra-alu human: (forward), 5’-TCA
CGCCTGGAATCCAGCA-3’, and (reverse) 5’-AGCTGGAGTAC
AGGGGCC-3’; for intra-Alu human 76 bp: (forward) 5’-AGA
CCATCCTGCTAACACG-3’, and (reverse) 5’-GTTCAACGAC
CTCCTGC-3’; intra-Alu human 200 bp: (forward) 5’-AAATT
AGCCGGCCGTG-3’, and (reverse) 5’-AGACGGAGTCTCGCTG
TC-3’, intra-Alu human 831 bp: (forward) 5’-AATGCCCAG
TTCTGAGGATTGTG-3’, and (reverse) 5’-CCCTGGCCAGCAATA
GTAAATG-3’; intra-Alu human 382 bp: (forward) 5’-AAAGAT
GAAGACCGGTTGTC-3’, and (reverse) 5’-CCCCGTCTTCAAAGA
ACATT-3’; intra-alu human 186 bp: (forward) 5’-GAACCA
GCTATTACCCCTTATCC-3’, and (reverse) 5’-CCCTGGCCA
GCAATATTGGA-3’; intra-Alu human 95 bp: (forward) 5’-
AAGTGAAGACCGGTTGTC-3’, and (reverse) 5’-CCAGGTTCA
CTCAGAGTCA-3’, and (reverse) 5’-CCAAGGCTTC
TGAGATTAAG-3’.

Hartl et al.
Impaired DNASE1L3 activity in sporadic SLE
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19 of 22
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**Figure S1.**  **DNASE1L3 activity does not correlate with renal function in patients with renal SLE.** DNASE1L3 activity in plasma was measured as in Fig. 1, C–E, and is expressed as percentage of activity in healthy controls. Symbols represent DNASE1L3 activity and UPCR in individual patients with renal SLE. Dashed line indicates the threshold for proteinuria (UPCR >0.5). Results of Spearman correlation with a two-tailed P value are shown.
Figure S2. Further characterization of auto-Abs to DNASE1L3 in SLE patients. (A) Abs to DNASE1, DNASE1L3, and FLAG peptide in the plasma of renal SLE patients (LN1, LN3, and LN5 from Fig. 2C) as measured by ELISA. All Ags were plated at a concentration of 2 µg/ml. Representative of two independent experiments. (B) Ab binding to recombinant DNASE1L3 assessed by Western blot. Purified recombinant DNASE1L3 was analyzed by denaturing SDS-PAGE and probed by Western blot using plasma samples from a DNASE1L3-deficient patient (1L3 null), healthy control (Ctrl), or from SLE patients (SLE). The presence (+) or absence (−) of IgG binding to DNASE1L3 as assessed by ELISA is indicated. For sporadic SLE patients with a positive ELISA (SLE, +), both plasma and the purified IgG fraction (SLE IgG, +) were analyzed. Representative of three independent experiments. (C) Correlation between anti-DNASE1L3 (1L3) Ab levels assessed by plate-based ELISA (OD) and bead-based Ag array (mean fluorescence intensity [MFI]). Data points represent individual renal SLE patients. Results of Spearman correlation with a two-tailed P value are shown. (D) Correlation between anti-DNASE1L3 (1L3) Ab levels and anti-dsDNA Ab levels as assessed by ELISA. Symbols represent individual patients with SLE (combined renal and nonrenal). Results of Spearman correlation with a two-tailed P value are shown. (E) Correlation between anti-DNASE1L3 (1L3) Ab levels and anti-C1q Ab levels as assessed by respective bead array assays. Symbols represent individual patients with renal SLE. Results of Spearman correlation with a two-tailed P value are shown. (F) Inhibition of DNASE1 activity by IgG from patients with anti-DNASE1L3 Abs. IgG was purified from the plasma of a healthy control (Ctrl, gray) or LN patient with anti-DNASE1L3 Abs (LN, gold). Recombinant DNASE1 was preincubated with purified IgG at the same concentration as DNASE1L3 in Fig. 2F (left panel) or at a 10-fold higher IgG concentration (right panel) and used to digest naked genomic Jurkat DNA. DNASE1 without preincubation with IgG (pink) was used as a control. Shown is the amount of remaining DNA after digestion with different concentrations of DNASE1. Dashed lines indicate 100% (black) and 50% (gray) of input DNA.
Figure S3. Correlation between plasma cfDNA and anti-dsDNA Ab levels in sporadic SLE patients. Symbols represent individual patients with sporadic SLE; all patients with and without LN were included. Anti-dsDNA Ab levels were assessed by ELISA, and DNA concentrations were assessed by qPCR. Results of Spearman correlation with a two-tailed P value are shown.

Figure S4. The length of protein-free DNA does not impact binding of anti-dsDNA IgG from SLE patients. (A) The effect of DNA length on binding to anti-dsDNA Abs as measured in a competition assay. Plasma from SLE patients was incubated with increasing concentrations of PCR-amplified DNA products of different lengths (95 bp, 186 bp, 382 bp, and 831 bp), and the binding of IgG to calf thymus dsDNA was assessed by ELISA. Dashed line shows binding in the absence of competitor DNA. Shown is a representative example of the data quantified for five plasma samples (also used in Fig. 5 F). (B) The effect of DNA length on binding to anti-dsDNA Abs as measured by ELISA. Plate-bound PCR-amplified DNA products of different lengths (95 bp, 186 bp, 382 bp, and 831 bp) were used as Ag to assess anti-dsDNA Abs from plasma of SLE patients. Calf thymus dsDNA, used in standard anti-dsDNA measurements, was used as a control. Shown is a representative example of five patients tested. In both approaches, data are representative of two independent experiments.
Three tables are provided online. Table S1 shows a comparison of clinical characteristics between sporadic SLE patients with DNASE1L3-sensitive IgG binding to MP Ags versus patients with no binding. Shown are demographics, ACR criteria, SLEDAI score, and immunosuppressive treatment in these patients. Table S2 summarizes clinical characteristics of sporadic SLE patients with LN organized by treatment response. Shown are demographics, renal biopsy characteristics, UPCR, as well as the change of DNASE1L3 readouts in patients with partial/complete treatment response compared to nonresponders. DNASE1L3 readouts were assessed at baseline (time of renal biopsy) and after 52 wk of immunosuppressive treatment. Table S3 shows characteristics for each individual patient included in the analysis from Table S2. Shown are the characteristics of each LN patient who had completed 52 wk of follow-up. Given are demographics, renal biopsy characteristics, treatment dosing, and changes in IgG binding to MP Ags as well as changes to cfDNA with complete, partial, or no-treatment response.