The soluble CD40 ligand sCD154 in systemic lupus erythematosus

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We found that the plasma of patients with active systemic lupus erythematosus (SLE) could induce a human B-cell line (Ramos) to express high levels of immune accessory molecules that are commonly found on blood B cells of patients with active SLE. The ability of SLE plasma to induce such phenotypic changes could be abrogated by neutralizing antibodies specific for the CD40 ligand (CD154) but not by antibodies to TNF-α. Immunoprecipitation studies with anti-CD154 identified a 20-kDa protein in the plasma of SLE patients with active disease, but not in plasma of normal donors, indicating that such plasma contained soluble CD154 (sCD154). Using a quantitative ELISA method, we found that the plasma of patients with active disease had levels of sCD154 that were significantly higher than those found in plasma of normal donors. Levels of CD154 transcripts in SLE blood lymphocytes correlated with the relative concentrations of sCD154 found in SLE plasma. Furthermore, plasma levels of sCD154 correlated with the titers of anti–double-stranded DNA autoantibody and with clinical disease activity. These studies indicate that sCD154 of patients with SLE may act as a functional ligand for CD40 that is associated with SLE disease activity.


Introduction
Systemic lupus erythematosus (SLE) is an autoimmune disease characterized by prominent lymphocyte activation, resulting in production of pathogenic IgG autoantibodies such as anti-double-stranded DNA antibody (anti-dsDNA Ab). Such autoantibodies may play a critical role in the progression of lupus nephritis (1). Whereas low amounts of low-affinity anti-dsDNA Ab also can be detected in the sera of healthy adults, high serum titers of anti-dsDNA Ab’s of the IgG isotype are found nearly exclusively in patients with SLE. Moreover, the levels of such IgG anti-dsDNA Ab’s appear related to disease activity.

Aberrant expression of immune costimulatory molecules may contribute to this pathophysiology. Studies indicate that the blood lymphocytes of SLE patients often express higher levels of immune accessory molecules, such as CD54, CD80, CD86, and CD95, than the blood lymphocytes of normal adults (2–5). High-level expression of CD80 or CD86 may contribute to pathologic presentation of self antigens to T cells and/or the production of pathologic anti-DNA autoantibodies. Consistent with this notion, the production of pathologic autoantibodies by lupus-prone New Zealand black (NZB) and New Zealand white (NZW) F1 mice can be ameliorated by CTLA4-Ig (6, 7), a recombinant protein that can block CD80/CD86→CD28 interactions (8). This has led to speculation that aberrant expression of these costimulatory molecules may contribute to the T-cell activation seen in patients with this disease.

Normal B cells can be induced to express immune costimulatory molecules by activated T cells. Activated CD4 T cells can express CD40 ligand (CD154), a molecule that can engage CD40 on the B-cell surface (9). This triggers a cascade of events that ultimately results in expression of a variety of heretofore nonexpressed stimulatory surface accessory molecules, such as CD80 (B7-1) (10–15). High-level expression of CD154 has also been detected on T cells from patients with active SLE, indicating that such cells may have exaggerated expression of this stimulatory molecule (16, 17). Conceivably, the exaggerated expression of CD154 could account for the high-level expression of immune accessory molecules on B cells of patients with active disease. Moreover, high-level expression of CD154 may be required for disease activity, which appears to be the case in animal models of SLE. Early et al., for example, reported that the anti-dsDNA Ab production in NZB/NZW F1 mice could be suppressed by treatment with anti-mouse CD154 antibody in vivo (18). Also, lpr/lpr mice made genetically defective in their ability to express CD154 did not develop IgG rheumatoid factor or anti-dsDNA (19).
On the other hand, soluble proteins released from activated T cells may contribute to immune activation (20–23). TNF-α, for example, is a protein that can exist as either a soluble molecule or a membrane-associated glycoprotein (24–26). Either form of the protein can augment B-cell expression of CD80 and other immune costimulatory molecules (27) and cause polyclonal B-cell activation (28). In this study, we examined whether the cell-free plasma of patients with SLE could also induce the expression of immune accessory molecules on human B cells.

**Methods**

After informed consent, blood was obtained from patients (22–69 years old) who satisfied diagnostic criteria of the American College of Rheumatology (ACR) for SLE (29) or from normal age-matched control donors. Whole blood was collected into tubes containing EDTA or heparin, and was separated immediately by centrifugation at 100 g at 4°C. The plasma was harvested and stored at –80°C until analyzed. Mononuclear cells were isolated from the cell pellet using density-gradient centrifugation in Histopaque 1077 (Sigma Chemical Co., St. Louis, Missouri, USA). The cells were analyzed immediately or were suspended in FCS containing 10% DMSO for storage in liquid nitrogen. The Burkitt’s lymphoma B-cell line Ramos was obtained from the American Type Culture Collection (Rockville, Maryland, USA) and cultured in RPMI-1640 supplemented with 10% FCS. Neutralization antibodies specific for human CD154 (TRAP-1), TNF-α (mAb1), or IFN-γ (NIB42) were obtained from PharMingen (San Diego, California, USA). Matrix metalloproteinase inhibitors BB94 and KB-R8301 were obtained from Ko Okumura (Juntendo University, Tokyo, Japan). Sodium azide was purchased from Sigma Chemical Co.

**Flow cytometry.** Isolated mononuclear cells were washed and suspended in staining medium (SM) consisting of RPMI-1640, 3% FCS, 0.05% NaN3, and 1 μg/mL propidium iodide (PI). The cells were stained for flow cytometry in SM containing saturating amounts of fluorescein-conjugated mAb’s. We used fluorescein-conjugated mAb’s specific for human CD54 or CD3 (Caltag Laboratories Inc., Burlingame, California, USA) or CD95 or CD154 (PharMingen). Alternatively, the cells were stained with fluorescein-conjugated isotype control antibody (MOPC-21; PharMingen). After 30 minutes at 4°C, the cells were washed with SM and analyzed by flow cytometry using a FACScalibur (Becton Dickenson Immunocytometry Systems, San Jose, California, USA). Dead cells that stained brightly with PI were excluded from the analyses. The relative expression of surface antigen is described as the mean fluorescence intensity ratio (MFIR). The MFIR equals the mean fluorescence intensity (MFI) of cells stained with a fluorescein-conjugated antigen-specific mAb divided by the MFI of cells stained with a fluorescein-conjugated isotype control mAb.

**Immunoprecipitation and immunoblotting of soluble CD154.** Human plasma (500 μL) from SLE patients or healthy donors were centrifuged at 10,000 g before immunoprecipitation. After preclearing with normal mouse IgG1 MOPC-21–conjugated, CNBr-activated Sepharose (20 μL; Pharmacia Biotech AB, Uppsala, Sweden), aliquots were immunoprecipitated with TRAP-1–conjugated, CNBr-activated Sepharose (20 μL) for 2 hours at 4°C. After washing with wash buffer (0.05% Tween-20 in PBS), the 1% SDS eluates were subjected to SDS-PAGE under nonreducing conditions and electroblotted onto nitrocellulose membranes (Bio-Rad Laboratories Inc., Hercules, California, USA). The blots were treated overnight at 4°C with blocking buffer (2% BSA and 0.05% Tween-20 in Tris-buffered saline). The blots were washed 2 times with wash buffer (0.05% Tween-20 in TBS) and incubated for 2 hours at room temperature with polyclonal rabbit antibody to human CD154 (C-20 or D-19; Santa Cruz Biotechnology Inc., Santa Cruz, California, USA). After 4 washings, the blots were incubated with horseradish peroxidase–conjugated (HRP-conjugated) goat anti-rabbit IgG antibody (Caltag Laboratories Inc.) followed by SuperSignal substrate (Pierce Chemical Co., Rockford, Illinois, USA). The blots were then exposed to BIOMAX x-ray film (Eastman Kodak Co., Rochester, New York, USA) for a few seconds.

**Figure 1**

Changes in surface antigen phenotype of Ramos B cells after culture in human plasma depleted of fibrinogen. Ramos B cells were cultured in media supplemented with 50% human plasma either without (a and b) or with (c and d) a neutralizing mAb specific for CD154. The cells were examined for expression of CD54 (a and c) or CD95 (b and d) using flow cytometry. The histograms depict the relative cell number and the logarithmic fluorescence intensity. Open histograms represent staining of Ramos cells with an isotype control mAb. Hatched and filled histograms depict the staining of Ramos cells with antigen-specific mAb after culture in plasma from a control donor (NOR-plasma) or patient with SLE (SLE-plasma), respectively.
ELISA for soluble CD154. We measured the levels of soluble CD154 (sCD154) by an ELISA that used 2 non-cross-blocking CD154 mAb’s (clones TRAP-1 and M90) obtained from Hideo Yagita and Ko Okumura (Juntendo University). For this, polystyrene microtiter plates (Corning-Costar Corp., Cambridge, Massachusetts, USA) were coated with TRAP-1 at 3 μg/mL in 0.1 M phosphate buffer (pH 9.0). The plates were washed with PBS and treated for 2 hours at room temperature with blocking buffer (10% FCS and 0.01% sodium azide in PBS). The plates were washed 4 times with wash buffer (0.05% Tween-20 in PBS), and then incubated overnight at 4°C with human plasma diluted in blocking buffer. For our standard curve, we used a recombinant sCD154 obtained from Bender MedSystems (Vienna, Austria). The plates were washed 4 times with wash buffer and then incubated for 1 hour with biotinylated M90 mAb at 3 μg/mL in blocking buffer. Washed plates were then treated sequentially with avidin and then biotinylated HRP (biotin-HRP) (Elite VECTASTAIN; Vector Laboratories, Burlingame, California, USA). After washing plates with blocking buffer, the plates were allowed to react with its substrate, 3,3′,5,5′-tetramethylbenzidine (TMB) peroxidase (Kirkegaard & Perry Laboratories Inc., Gaithersburg, Maryland, USA). The OD at 450 nm was determined using an ELISA microplate reader (Molecular Devices Corp., Sunnyvale, California, USA).

Quantitative CD154 RT-PCR ELISA. ELISA-based quantitative competitive RT-PCR for CD154 was as described previously (30). Total RNA from PBMCs (1-5 × 10^6 cells) was extracted with the Total RNaseasy Kit (QIAGEN Inc., Valencia, California, USA). The competitor RNA was generated from the insert-containing CD154 cDNA cloned into pBluescript (Stratagene, La Jolla, California, USA), using the Superscript Preamplification System. Varying amounts of competitor RNA were added to separate wells of isolated total RNA that subsequently were converted into cDNA using the First Strand cDNA Synthesis Kit (GIBCO BRL, Grand Island, New York, USA). PCR reactions were performed for 35 cycles, and the products were purified with a QIAQuick Spin PCR Purification Kit (QIAGEN Inc.). Standard DNA of CD154 was amplified from pBSII KS using T3 primer (5′-TTAACCCTCACTAAAG-3′) and T7 primer (5′-biotin-AATACGACTCACTATAG-3′). PCR product serial dilutions of 1:3, ranging from 1 pM to 500 aM, were used to generate a standard curve.

The biotinylated PCR products were captured onto microtiter plates (Becton Dickinson Labware, Franklin Lakes, New Jersey, USA) coated with streptavidin (Sigma Chemical Co.). PCR products were loaded into 2 wells and incubated at room temperature for 1 hour. The plates were treated with 0.1 N NaOH for 10 minutes to remove the sense-strand DNA, and then were washed with 0.1x SSC and BW buffer. The plates were incubated with oligonucleotide probes labeled with digoxigenin-11-dideoxy-UTP (Boehringer Mannheim, Indianapolis, Indiana, USA) at 65°C for 20 minutes and then at 42°C for 90 minutes. After washing with HYBE buffer (6x SSC, 0.1% N-lauroyl sarlosine) and blocking buffer (0.5% Genius blocking reagent [Boehringer Mannheim] in Tris-HCl buffer), HRP-conjugated antidigoxigenin antibody (Boehringer Mannheim) in blocking buffer was added and incubated for 30 minutes at room temperature; then plates were allowed to react with its substrate TMB peroxidase. The OD at 450 nm was determined using an ELISA microplate reader.

Figure 2
Immunoblot analyses of sCD154 isolated from culture supernates of human CD154–transfected HeLa cells (a and b) or human plasma samples (c). Cell-free supernatants (SUP), or plasma from a control donor (H1) or each of 2 SLE patients (S1 and S2), were absorbed onto anti-CD154 mAb–coupled Sepharose beads. The affinity-bound material was eluted, and then subjected to SDS-PAGE and immunoblot analysis with anti-CD154 antisera specific for the extracellular (a and c) or cytoplasmic (b) domain of CD154, respectively. The left lanes in a and b (labeled MW) are proteins of known molecular size that serve as standards. To the left of each panel are lines indicating the migration distance of each of these proteins, along with its molecular size. The arrows at the right of each panel indicate the expected migration distance of sCD154.
Standard curves plotting the moles of RNA product versus OD were made for the standard cDNA reactions. The equations describing these standard curves were then used to calculate the moles of wild-type or competitor DNA present in the unknown PCR reactions, based on the ODs obtained in the ELISA readings. The ratio of wild-type DNA to competitor DNA was then plotted against the known quantity of competitor RNA added in the initial samples. The ratio of 1 was interpreted to mean that the amount of target RNA equaled the amount of competitor RNA added to the sample. The molecules of target RNA per PBMC were then calculated based on the following formula: \[ \text{[moles target CD154 RNA]} \times (6 \times 10^{23} \text{ molecules per mole}) \times \text{(dilution factor of test RNA)}]/\text{(percent of PBMCs in total cell population)}. \]

Measurement of disease activity. The SLE disease activity index (SLEDAI) score and anti-dsDNA Ab titer were determined at the time of blood sampling. Disease activity for lupus patients was measured using the SLEDAI score (31). Plasma anti-dsDNA levels were measured using the Farr technique at Specialty Laboratories (Santa Monica, California, USA).

Statistical analyses. The mean sCD154 plasma concentrations of patients with SLE and the mean concentrations of control donors were compared using the Mann-Whitney nonpaired nonparametric test. Relationships between the level of sCD154 protein and CD154 mRNA or anti-dsDNA Ab titer in SLE patients were both analyzed with the Pearson correlation test. Because the Pearson correlation test assumes the usual bivariate Gaussian distribution, we also analyzed these relationships using the Spearman rank order correlation test, which makes less stringent assumptions.

Results

Plasma from patients with active SLE can upregulate expression of immune accessory molecules on Ramos B cells. We assessed phenotypic changes in Ramos B cells after culturing the cells in media supplemented with plasma from patients with SLE. Briefly, Ramos B cells were cultured for 48 hours in media supplemented with 50% plasma from patients with SLE or from normal control donors. We found that the plasma from patients with active SLE could induce such cells to express higher levels of CD54 and CD95, whereas plasma from healthy donors could not (Figure 1, a and b). Moreover, plasma from patients with active SLE could induce Ramos B cells to express CD80 after 72 hours in culture (data not shown). The ability of SLE plasma to induce increased expression of CD54, CD95, or CD80 could be
inhibited by anti-human CD154 mAb (Figure 1, c and d), but not by anti-human TNF-α or anti-human IFN-γ mAb’s (data not shown). The increased expression of CD54, CD80, or CD95 is similar to that noted for leukemia B cells stimulated with cells expressing membrane-bound CD154 (32). Collectively, these data imply that the plasma of patients with active SLE could induce expression of immune accessory molecules on B cells, and that this activity was mediated by sCD154.

sCD154 in plasma of patients with active SLE. To look for sCD154 in SLE plasma, we first established whether we could detect sCD154 in the culture supernatants of a HeLa cell line (designated CD154-HeLa) that was transfected with an expression vector encoding the full-length human CD154. Culture supernatants of CD154-HeLa were incubated with Sepharose beads conjugated with anti-CD154 mAb. This process captured a protein of approximately 20 kDa that could be identified with immunoblot analysis using an antibody specific for the COOH-terminal (extracellular) region of CD154 (Figure 2a), but not with an antibody specific for the NH2-terminal (cytoplasmic) region of CD154 (Figure 2b). The size of this protein was similar to the 19- to 20-kDa sCD154 glycoprotein isolated from the supernatants of activated T cells (33). We then examined whether we could immunoprecipitate sCD154 from the plasma of patients with active SLE or healthy donors. As seen in Figure 2c, sCD154 could be isolated from the plasma of patients with active SLE (S1 and S2), but not from plasma of a normal healthy donor (H1).

To measure the levels of sCD154 in the plasma of SLE patients, we developed a sensitive and specific sandwich ELISA using 2 non–cross-blocking anti-CD154 mAb’s (clones TRAP-1 and M90). Recombinant homotrimeric sCD154 was used as a positive standard control for the ELISA (Figure 3a). The minimum concentration of sCD154 that we could detect with this assay was approximately 40 pg/mL. The specificity of this ELISA was confirmed by its ability to detect human sCD154 protein, but not murine sCD154, human soluble Fas ligand, or human soluble CD27 (data not shown).

To assess whether the sCD154 detected in the plasma of patients with SLE was released from cells after phlebotomy, we examined changes in plasma levels of sCD154 from collected blood samples over time. Plasma was isolated at various times after phlebotomy from whole-blood samples kept at room temperature or at 4°C. The measured sCD154 levels in blood samples kept at 4°C were constant over the first 6 hours after sample collection (Figure 3b). Moreover, these levels were similar to those found in plasma processed immediately after phlebotomy. However, the sCD154 levels in plasma isolated from whole blood kept at room temperature increased over time, reaching a plateau value after 2 hours (Figure 3b). Addition of matrix metalloproteinase inhibitors BB94 or KB-R8301 to the blood samples immediately after phlebotomy could inhibit this rise in sCD154 over time (data not shown). However, addition of sodium azide to blood samples in order to reach a final concentration of 0.05% did not inhibit the release of sCD154 over time (data not shown). These data indicate that sCD154 may be cleaved from membrane CD154 by a matrix metalloproteinase and released into the plasma ex vivo through a process that does not require oxidative phosphorylation. Nevertheless, patients with active SLE apparently have elevated levels of sCD154 in vivo, because high levels of sCD154 could be detected in the plasma from such patients even when it was isolated immediately after phlebotomy.
We determined the levels of sCD154 in the plasma of 21 healthy subjects and 26 SLE patients who had different degrees of disease activity. Plasma samples were isolated from whole-blood samples that were chilled immediately to 4°C after phlebotomy and then processed within 30 minutes (Figure 3c). In the control group, the plasma concentration of sCD154 was 290 ± 340 pg/mL (mean ± SD). Moreover, half of the tested plasma samples from the control donors did not have detectable amounts of sCD154 (≤40 pg/mL), as determined by ELISA (Figure 3c). However, the plasma of SLE patients had significantly higher levels of sCD154 (6,800 ± 4,300 pg/mL; mean ± SD) than the control plasma samples (\( P < 0.0001 \) by Mann-Whitney unpaired nonparametric test).

**Correlation between expression levels of CD154 mRNA and sCD154.** To examine the levels of CD154 mRNA in the blood mononuclear cells of patients with SLE, we used an ELISA-based quantitative competitive RT-PCR to measure the numbers of CD154 transcript per cell, as described (30). This method titers known amounts of a CD154 competitor transcript containing an engineered mutation that can be detected with oligonucleotide hybridization.

We found that the blood mononuclear cells of 10 patients with active SLE had an average of 7,800 ± 1,100 (SD) transcripts of CD154 mRNA per cell. The number of transcripts per cell was significantly higher than that noted in the blood mononuclear cells of 10 healthy control donors (\( P < 0.001 \) by Mann-Whitney unpaired nonparametric test). Furthermore, we found that the levels of CD154 transcripts per cell for individual patients correlated with the relative levels of sCD154 detected in their respective plasma samples (Pearson \( r = 0.779, P = 0.008 \); Spearman \( r = 0.782, P < 0.02 \) (Figure 4a). However, the relative levels of sCD154 transcripts did not have such a significant correlation with the relative MFIR values for membrane-bound CD154 detected on the cells in respective samples using flow cytometry (Spearman \( r = 0.194, P = 0.584 \) (Figure 4b).

**Correlation between sCD154 plasma concentration and dsDNA titer.** We compared the SLEDAI of each patient with the level of sCD154 detected in the plasma. The levels of SCD154 were highest in the plasma of patients with highly active disease who had a SLEDAI score of 10 or greater. These 10 patients had higher mean plasma levels of SCD154 (mean ± SD of 10,680 ± 2,840 pg/mL). Patients with less-active disease (SLEDAI scores between 2 and 9) had a significantly lower mean plasma level of SCD154 (\( m = 3,880 ± 2,080 \) pg/mL, \( n = 16 \)) than patients with SLEDAI scores of 10 or greater (\( P < 0.0001 \) by Mann-Whitney unpaired nonparametric test; Figure 5a). Nevertheless, the mean sCD154 plasma level of these patients was still significantly higher than that of normal donors (\( m = 423 ± 437 \) pg/mL, \( n = 10, P < 0.0001 \) by Mann-Whitney unpaired nonparametric test; Figure 5a).

We also compared the anti-dsDNA Ab titers with sCD154 concentration in SLE plasma. A highly significant linear correlation was found between plasma anti-dsDNA Ab titers and sCD154 plasma levels (Pearson \( r = 0.837, P < 0.0001 \); Spearman \( r = 0.758, P < 0.001 \) \( n = 23 \)) (Figure 5b). We also examined serial plasma samples collected over a 17-week period from one patient who initially had a high plasma level of sCD154.
(10,194 pg/mL) and a high titer of anti-dsDNA Ab (650 IU/mL) (Figure 5b, square 1). This patient's plasma levels of sCD154 and titer of anti-dsDNA Ab rose (Figure 5b, square 2) and fell (Figure 5b, squares 3 and 4) concomitantly after treatment, to final values of 1,114 pg/mL and 34 IU/mL, respectively. However, no correlation was found between sCD154 plasma levels and proteinuria (data not shown).

Discussion

In this study, we found that cell-free plasma of patients with SLE could induce Ramos B cells to express high levels of CD54 and CD95. The ability of SLE plasma samples to induce such phenotypic changes could be abrogated by pretreating the plasma with neutralizing mAb's specific for CD154 (Figure 1). Immunoprecipitation and immunoblot analyses revealed that the plasma from patients with active disease possessed a protein of approximately 20 kDa that reacted with mAb's specific for the extracellular domain of CD154 (Figure 2). Collectively, these data indicate that the plasma of patients with active SLE contains a CD154 polypeptide that has biologic activity.

CD154 exists ordinarily as a 39-kDa type II membrane glycoprotein (34). This protein is expressed transiently by activated CD4+ T cells (30, 35–37). However, CD154 also can be cleaved from the cell surface of activated T cells, releasing a sCD154 of approximately 18–20 kDa (38). This sCD154 can apparently form homotrimers that can induce B-cell activation or differentiation (33, 38). Moreover, several in vitro studies have revealed that sCD154, or a soluble recombinant CD154 chimeric protein, can induce antigen-presenting cells (APCs) to express immune accessory molecules in a manner similar to that of plasma from patients with active SLE (33, 39, 40).

The sCD154 in plasma of patients with active SLE is most likely derived from the membrane-bound form of CD154 on activated T cells. Previous reports demonstrated that the absolute numbers of blood T cells that express surface CD154 are increased in patients with active lupus (16, 17). Moreover, T cells from SLE patients could also maintain high-level surface expression of CD154 in culture, indicating that such T cells have constitutive or exaggerated expression of CD154. Consistent with this, we found that the blood mononuclear cells of patients with active SLE expressed significantly higher levels of CD154 transcripts per cell than the blood mononuclear cells of control donors. However, the expression levels of surface CD154 on such cells did not correlate well with disease activity (data not shown). On the other hand, the plasma concentrations of sCD154 did have a significant correlation with disease activity (Figure 5). The soluble molecule does not appear to be encoded by a truncated CD154 mRNA, because only full-length transcripts could be identified in the mononuclear cells of patients with SLE (data not shown). Furthermore, the level of sCD154 found in the plasma had a significant correlation with the level of CD154 mRNA detectable in blood mononuclear cells using quantitative RT-PCR (Figure 4a). As such, it appears that sCD154 in the plasma of patients with SLE results from cleavage of the full-length form of the CD154 molecule.

Consistent with this notion, we found that the plasma concentrations of sCD154 could be inhibited by chilling the sample to 4°C or by adding matrix metalloproteinase inhibitors, such as BB94 or KB-R8301, to the collected sample. This inhibition indicates that the release of sCD154 is a temperature-dependent process that may require a matrix metalloproteinase. This process is similar to that required for release of TNF-α or soluble Fas ligand (41, 42).

Despite the ex vivo production of sCD154, we found that plasma immediately isolated from the collected blood samples of patients with SLE still contained high concentrations of sCD154. This concentration does not appear secondary to the immediate release of sCD154. One study indicated that platelets can express CD154 within seconds after activation in vitro (43). For this reason, we collected the blood samples in tubes containing heparin sulfate to prevent thrombus formation and platelet activation. Because we found that the ex vivo release of sCD154 was temperature-dependent, we chilled subsequent blood samples immediately to 4°C and processed them soon thereafter. Blood samples were collected from SLE patients and control donors in an identical fashion. If our methods stimulated an immediate release of sCD154 from platelets, then the plasma collected from control donors would also be expected to contain elevated levels of sCD154. Instead, the plasma from patients with active SLE had consistently higher levels of sCD154 than plasma from control donors. Furthermore, the significant correlation between the levels of lymphocyte CD154 mRNA and plasma concentrations of sCD154 (Figure 4a) also argues against the notion that sCD154 is released from platelets at the time of sample collection or processing. Instead, it appears that patients with active SLE have elevated levels of sCD154 in vivo.

The elevated plasma levels of sCD154 appear to be associated with autoimmune disease activity. The plasma samples from most normal donors did not have sCD154 at levels that could be detected in our assay (e.g., 240 pg/mL). Moreover, the plasma samples of patients with other diseases, such as progressive systemic sclerosis, pneumonia, Sjögren's syndrome, or multiple myeloma, also had little or no detectable sCD154 (data not shown). On the other hand, all the plasma samples from patients with SLE contained detectable amounts of sCD154. Furthermore, the sCD154 levels in plasma correlated with anti-dsDNA Ab titers and clinical disease activity, as assessed by the SLEDI (31) (Figure 5).

In view of this data, we hypothesize that the correlation defines a causal relationship. Similar to membrane-bound CD154, elevated expression of sCD154
may contribute to immune activation of APCs and stimulation of autoantibody-producing B cells in patients with SLE. In addition to its ability to activate B cells, monocytes, or dendritic cells, CD154 also may induce inflammatory changes in CD40-bearing non-hematopoietic cells. For example, endothelial cells can be induced to upregulate surface expression of CD54, CD62E, and CD106 (44) upon CD40 ligation, thereby assuming a phenotype more conducive to inflammation. Conceivably, the vasculitis that is often observed in patients with highly active SLE may be due, in part, to high levels of circulating sCD154. Consistent with this notion, we found that patients with active vasculitis in the setting of rheumatoid arthritis also have high plasma levels of sCD154 (data not shown; T. Tamura, S. Kobayashi, K. Kato, T.J. Kipps, and H. Brando, manuscript in preparation).

Similarly, it is possible that sCD154 may play a role in renal pathology that often develops in patients with SLE. Because monomeric, and even homotrimeric, sCD154 falls below the 64-kDa cutoff for renal filtration, sCD154 may pass through the renal glomerulus and become sequestered or undergo absorption in the proximal tubules. Accordingly, sCD154 may have a greater influence on proximal tubular epithelial cells than the blood lymphocytes that express this molecule on the cell surface. It is noteworthy that proximal tubular epithelial cells also express CD40. Moreover, ligation of CD40 on these cells stimulates them to express inflammatory chemokines, such as IL-8, MCP-1, and RANTES (45). Although we failed to detect a significant correlation between the plasma level of sCD154 and proteinuria in patients with SLE, further studies are necessary to determine whether the plasma levels of sCD154 correlate with progression of lupus nephropathy.

Future studies are required to determine whether the plasma level of sCD154 has value in predicting flare-ups in SLE disease activity. It is possible that the level of sCD154 may rise and fall faster than that of anti-dsDNA Ab’s during flare-ups and remissions in disease activity, respectively. If so, then assessing the plasma concentration of sCD154 may prove to be a more reliable way to monitor disease activity in patients with SLE.

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