Combined Immunodeficiency Due to MALT1 Mutations, Treated by Hematopoietic Cell Transplantation

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Abstract
Purpose A male infant developed generalized rash, intestinal inflammation and severe infections including persistent cytomegalovirus. Family history was negative, T cell receptor excision circles were normal, and engraftment of maternal cells was absent. No defects were found in multiple genes associated with severe combined immunodeficiency. A 9/10 HLA matched unrelated hematopoietic cell transplant (HCT) led to mixed chimerism with clinical resolution. We sought an underlying cause for this patient’s immune deficiency and dysregulation.

Methods Clinical and laboratory features were reviewed. Whole exome sequencing and analysis of genomic DNA from the patient, parents and 2 unaffected siblings was performed, revealing 2 MALT1 variants. With a host-specific HLA-C antibody, we assessed MALT1 expression and function in the patient’s post-HCT autologous and donor lymphocytes. Wild type MALT1 cDNA was added to transformed autologous patient B cells to assess functional correction.

Results The patient had compound heterozygous DNA variants affecting exon 10 of MALT1 (isoform a, NM_006785.3), a maternally inherited splice acceptor c.1019-2A>G, and a de novo deletion of c.1059C leading to a frameshift and premature termination. Autologous lymphocytes failed to express MALT1 and lacked NF-κB signaling dependent upon the CARMA1, BCL-10 and MALT1 signalosome. Transduction with wild type MALT1 cDNA corrected the observed defects.

Conclusions Our nonconsanguineous patient with early onset profound combined immunodeficiency and immune dysregulation due to compound heterozygous MALT1 mutations extends the clinical and immunologic phenotype reported in 2 prior families. Clinical cure was achieved with mixed chimerism after nonmyeloablative conditioning and HCT.

Keywords BCL10 · bone marrow transplant/hematopoietic cell transplant · CARD11 · CARMA1 · combined immunodeficiency (CID) · erythroderma · immune dysregulation

Introduction
Identification of the genetic causes of human immunodeficiencies has revealed the roles of many factors critical for
human lymphocyte development and function. Combined immunodeficiencies (CIDs) listed by the IUfS Expert Committee on Primary Immunodeficiencies [1], include a wide spectrum of gene defects underlying susceptibility to bacterial, viral and fungal infections. The most profound of these, collectively termed severe combined immunodeficiency (SCID), are disorders with few to absent autologous T cells and absent cellular and humoral immune function [1–4]. In contrast, many CID gene defects do not abrogate development or release into the periphery of T and B cells, but instead disrupt pathways critical for their effector and regulatory roles; examples are ORAI-I, STIM-1 [5], and MHC class II deficiency [6]. While over 14 different SCID genes are known [7], many patients with CID without T cell lymphopenia have as yet unidentified genetic defects. Whole exome sequencing (WES) may identify molecular causes of CID.

Studies in knockout mice and human malignancies and immunodeficiencies have delineated the intracellular signaling pathways activated by engagement of lymphocyte antigen receptors and G-protein coupled receptors [8]. NF-κB, a central mediator of activation signals, translocates from the cytoplasm into the nucleus to initiate transcription of genes that bring about lymphocyte maturation, activation and proliferation [9]. NF-κB activation and signaling is in turn controlled by multiple mechanisms, one of which is the signalosome formed from assembly of CARMA1 (also called CARD11), BCL-10 and MALT1 into the “CBM” signaling complex [10, 11]. While the precise molecular mechanisms are still not completely clear, stimulation through the T cell and B cell receptors causes phosphorylation of CARMA1, recruitment of MALT1 and BCL-10, and oligomerization of components of the CBM complex [12–14]. This in turn activates the IkB kinase complex through TNF receptor-associated factor 6 (TRAF6)-mediated ubiquitination of NF-κB essential modulator (NEMO) [15–17], leading to phosphorylation and proteasomal degradation of the inhibitor IkBα and release of NF-κB. Thus, it is not surprising that defects in NEMO, CARMA1 and MALT1 have been found to cause human CID [18–23].

We describe a new patient in whom CID and immune dysregulation due to MALT1 compound heterozygous mutations was successfully treated by allogeneic hematopoietic cell transplantation (HCT). This case in a non-consanguineous family, combined with 2 prior reports [22–24], broadens the spectrum of MALT1 deficiency disease and suggests an effective treatment.

Methods

Patient

After informed consent, as approved by the University of California San Francisco Committee on Human Research, the patient, his parents and 2 healthy siblings were studied with whole exome sequencing and immunological assessments.

DNA Studies

Genomic DNA from the patient, obtained prior to HCT, and from his parents and siblings was subjected to WES. Analysis tools were similar to [25], with modifications detailed in the Supplementary methods. DNA variants were confirmed by Sanger sequencing. With parental consent, residual dried blood spots obtained in the newborn nursery were recovered from the California Department of Public Health Newborn Screening Program, and T cell receptor excision circles (TRECs) were analyzed as described [26].

Cell Separations and Reagents

After HCT from an unrelated donor differing at a single HLA-C locus, the patient developed mixed chimerism of the hematopoietic system. Patient alleles were HLA-C *08:01, *03:04; donor alleles were *08:01, *07:02. Staining cells with monoclonal human IgM antibody (clone ID: TRA2G9) recognizing antigens encoded by C*01/*03/*04:01/*03:04; donor alleles were *08:01, *07:02. Staining cells with anti-human IgM (clone MHM-88), permitted separation of autologous patient lymphocytes from those of the donor by flow cytometry. For specific antibodies see Supplementary methods.

PCR and Western Blotting

RNA was isolated from sorted autologous patient PBMCs obtained post-HCT (RNeasy kit, Qiagen), and expression of MALT1 transcripts (primers in Suppl Table 1) was detected by PCR (Superscript III system, Life Technologies) followed by Sanger sequencing. The sorted cells were also lysed with 1 % NP-40 and analyzed by Western blotting using antibodies recognizing MALT1 (EP603Y, Abcam) and BCL-10 (H-197, Santa Cruz Biotechnologies).

Intracellular Signaling Assays

For phosphorylation assays PBMCs or Epstein-Barr virus (EBV)-transformed B cells were stimulated with 400 nM PMA and 250 ng/ml ionomycin at 37 °C, for 10 min. For cytokine assays PBMCs were stimulated for 6 h with PMA plus ionomycin: 200 ng/ml supernatant staphylococcal enterotoxin E (SEE, Toxin Technology, Inc.) plus 4 µg/ml anti-CD28 clone 9.3; or 1:500 anti-CD3 clone Leu-4 ascites plus 4 µg/ml anti-CD28. The cells were then fixed, permeabilized (Invitrogen Solutions A and B, Life Technologies) and incubated with either phospho-
specific unconjugated antibodies followed by anti-rabbit-PE, or anti-mouse-FITC labeled secondary antibodies, or antibodies against IL-2 or IFN-γ. Fluorescent antibodies to relevant surface markers were included.

Fig. 1 a, Skin lesions, showing (i) facial erythroderma (ii) dorsal trunk and (iii) close-up with linear distribution of lesions corresponding to excoriations. b, Schematic representation of MALT1 protein (isoform a, NP_006776.1), illustrating death domain, 3 immunoglobulin (Ig)-like domains, and paracaspase domain [30]. Mutations are shown for our patient (black) and prior homozygous cases (blue) [22, 23]. c, Browser view of patient stacked DNA sequence tracks, demonstrating >30X coverage with individual reads homozygous for either the splice disrupting variant c.1019-2A>G or the single nucleotide deletion cDNA c.1060delC (the former also present in maternal DNA, not shown) (ClinVar number SCV000196510). Colored blocks, non-reference nucleotides; black blocks, deleted nucleotides. Below, genomic sequence of MALT1 exon 10 (isoform a, NM_006785.3) and protein translation with reference sequence (black) and de novo deleted sequence (red).
Lentivirus Transduction

*MALT1* cDNA (Genecopia) was ligated into lentiviral vector MP-283: pSicoR-BstXI-EF1a-puro-T2A-mCherry, (kindly provided by Michael McManus, Lentiviral RNAi Core, UCSF). MP-283-MALT1 lentiviral supernatant prepared by transient transfection of 293 cells in DMEM medium with 10 % fetal calf serum (FCS) was used to transduce EBV cells in plates pre-coated with Rectronectin (Takara, Japan). After two 24 h infections, the EBV cells were washed and expanded in RPMI 1640 medium with GlutaMax (Invitrogen), 20 % FCS, penicillin, and streptomycin. MP-283 lentivirus prepared as above without the insert was used in parallel. Transduced cells were detected by mCherry fluorescence.

## Results

### Patient History

The infant, born at term to non-consanguineous parents with negative family history, developed blood-streaked diarrhea and a desquamating, erythematous pruritic rash, the latter evolving into firm erythematous papules affecting trunk, palms and soles (Fig. 1a, Table 1). There were no indications of environmental atopy; IgE was undetectable, and steroids were ineffective. Skin biopsies showed perivascular lymphocytic infiltration in the dermis. Following monthly otitis media infections, an immune workup showed expansion of CD4 and CD8 T cell populations with high proportions of naïve CD45RA T cells, but impaired in vitro proliferative responses (Tables 1 and 2). Maternal T cell engraftment was absent, and the newborn dried blood spot, retrieved from the time of the patient’s birth, had 627 T cell receptor excision circles (TRECs)/μL (normal >40). B and NK cells were present, but low immunoglobulin levels and absent antibody responses to vaccines necessitated immunoglobulin infusions. The patient experienced poor weight growth, stomatitis, oral thrush, RSV bronchiolitis, and CMV viremia and CMV pneumonitis.

At 18 months of age, following non-ablative conditioning with 120 mg/kg cyclophosphamide, 140 mg/m² melphalan and 8 mg/kg rabbit anti-thymocyte globulin, the patient received peripheral mobilized CD34-selected cells from a 9 of 10 HLA antigen-matched unrelated donor (differing at one HLA-C locus). The patient received 4.85 × 10⁶ CD34 cells/kg and 14.72 × 10⁶ CD3 cells/kg. Graft versus host disease (GVHD) prophylaxis included methotrexate and cyclosporine, and no GVHD was observed. The patient’s rash resolved within 4 weeks and did not return. Donor T cell function was established by 5 months post-HCT. However, diarrhea and CMV viremia, refractory to gancyclovir and foscarnet treatment, continued until both resolved promptly following infusion 12 months post-HCT of donor T cells that had been pulsed in vitro with CMV peptide and expanded. Immunoglobulin replacement has been given post-HCT, but donor B cell chimerism, IgA, IgM and anti-A IgM isohemagglutinins are now detectable.

### Identification of Mutations in MALT1

Sequencing of a comprehensive panel of known SCID genes revealed no mutations. To search for disease-causing variants,

| Age | Clinical manifestation |
|-----|------------------------|
| 1–3 months | Bloody stool, erythroderma (later biopsy showing lymphocyte infiltration) |
| 9 months | Poor growth (<5 % weight, 5 % height), hospitalization for prolonged fever, presumed bacterial infections responding to systemic antibiotics; *S. aureus* superinfection of rash |
| 10 months | IgG infusions instituted |
| 13 months | Thrush, candida esophagitis |
| Continuous antibiotic prophylaxis started |
| DNA isolated from PBMCs, later used for whole exome sequencing |
| 15 months | Persistent CMV >3,000 copies by PCR from blood, lung washings despite gancyclovir and foscarnet treatment; ground glass pneumonitis; self limited RSV bronchiolitis; diarrhea with C. difficile |
| 18 months | Hematopoietic cell transplant from 9/10 HLA matched unrelated donor |
| 19 months | Rash resolved, donor T cells detected; no graft vs. host disease |
| 23 months | Graft vs. host disease prophylaxis discontinued |
| Lymphocyte proliferation to PHA >50 % normal, persistent CMV viremia 1,500 copies |
| PBMCs isolated, separated into autologous patient and donor populations for in vitro functional studies |
| 28 months | Antibiotic prophylaxis discontinued |
| 30 months | Donor T cell infusion for persistent CMV viremia |
| CMV viremia resolved, gaining weight (25 % for age) |
| 6 years | Donor B cell function detected with normal IgM and IgA, positive IgM isohemagglutinin |
Table 2  Clinical and laboratory findings of patients with MALT1 deficiency

|                          | New patient, this report                  | Jabara H, et al.                  | McKinnon M, et al.                  |
|--------------------------|------------------------------------------|-----------------------------------|-----------------------------------|
|                          | Pre-transplant | Post-transplant | 2 siblings | 1 patient                          |
| Age at immune evaluation | 9 – 13 months | 2.5 years       | 4 years, 2.25 years | 15 years |
| Consanguinity            | No                        | Yes                      | Yes                          | Yes |
| Infections               |                           |                          |                             |     |
| *S. aureus*              | Cellulitis                | Resolved                 | Cellulitis, pneumonia         |     |
| CMV                      | Blood, bronchial lavage   | Resolved                 | Lung, duodenum                |     |
| Candida                  | Oral thrush               | Resolved                 | Repeated urine isolations     |     |
| *C. difficile*           | Diarrhea                  | Resolved                 | Pneumonia                      |     |
| RSV                      | Bronchiolitis             | Resolved                 |                               |     |
| *S. pneumoniae*          | Yes                       | Pneumonia and meningitis  |                               |     |
| Other pulmonary isolates | No                        |                           |                               |     |
| Other skin isolates      | No                        | Not reported             |                               |     |
| Clinical manifestations  |                           |                          |                               |     |
| Poor or delayed growth   | Yes                       | Resolved (10 % height, 30 % weight for age) | Yes | Yes |
| Oral lesions (aphthous ulcers, cheilitis, gingivitis, thrush) | Yes | Resolved | Yes | Yes |
| Eczematous rash          | Yes, erythroderma         | Resolved                 | Not reported                   |     |
| Inflammatory bowel disease | Yes                      | Resolved                 | Yes, severe dermatitis         |     |
| Neurologic development   | Normal                    | Normal                   | Normal                         |     |
| Bronchiectasis           | No                        | No                       | Yes, respiratory failure       |     |
| Other findings           | None                      | None                     | Mastoiditis                    | Dysorphic facies, bone fractures, granulation tissue on vocal cord, larynx, ear canal |
| Treatments               |                           |                          |                               |     |
| Immunoglobulin infusions | Yes                       | Yes                      | Yes                            | Yes |
| Antibiotics              | Yes                       | No                       | Yes                            | Yes |
| Additional measures      | Hematopoietic cell transplant |                        | None reported                  | Nissan fundoplication, jejunostomy |
| Outcome                  | Alive, well, 7 years      | Deceased, respiratory failure, 13 years, 7 years | Alive                        |     |
| Immunologic Parameters   |                           |                          |                               |     |
| Cells X 10^9/L (normal range for age) |         |                          |                               |     |
| WBC (4.5–17.5)           | 17.0                      | 8.9                      | N.A.*                          | N.A. |
| Lymphocytes (2–8)        | 10.6 **                   | 4.5                      | Normal                         | Normal |
| Eosinophils (0–1.1)      | 2.43 ↑                    | 0.2                      | N.A.                           | N.A. |
| Table 2 (continued) | New patient, this report | Jabara H, et al. | McKinnon M, et al. |
|---------------------|--------------------------|-----------------|-------------------|
|                     | Pre-transplant | Post-transplant | 2 siblings | 1 patient |
| Lymphocyte subsets, cells/μL | | | | |
| CD3 (1,610–4,230) | 9,133 ↑ | 3,743 | Normal | Elevated |
| CD4 (900–2,860) | 4,142 ↑ | 2,300 | Normal | Elevated |
| CD8 (630–1,900) | 4,460 ↑ | 1,128 | Normal | N.A. |
| CD4:CD8 ratio (1–2.1) | 0.9 ↓ | 2.0 | N.A. | 3 ↑ |
| CD3 CD4 CD45RA | 2,526 | 1,426 | Normal | N.A. |
| CD3 CD8 CD45RA | 1,561 ↑ | 823 | N.A. | N.A. |
| CD19 (700–1,300) | 1062 | 226 | Normal | 50 ↓ |
| NK (130–1,300) | 425 | 451 | Low, normal | Normal |
| T regulatory cell % | | | | |
| CD25% of CD4 cells (11–20) | 5 ↓ | 30 | N.A. | N.A. |
| FoxP3% of CD4 CD25<sup>+</sup>CTLA-4 (79–91) | 56 ↓ | N.A. | N.A. | Normal |
| CD4 CD25 CD45RA (4–67) | N.A. | 7 | N.A. | N.A. |
| CD4 CD25 CD45RO (4–25) | N.A. | 23 | N.A. | N.A. |
| B cell subset % (normal range) | | | | |
| CD27<sup>+</sup>IgM<sup>+</sup>IgD<sup>+</sup> of CD19<sup>+</sup> (0.2–12) | N.A. | 2.1 | N.A. | Absent |
| CD27<sup>+</sup>IgM-IgD<sup>-</sup> of CD19<sup>+</sup> (1.9–30.4) | N.A. | 1.9 | N.A. | Reduced |
| CD18<sup>+</sup>IgM<sup>+</sup> of CD19<sup>+</sup> (7.6–48.6) | N.A. | 70.6 ↑ | N.A. | N.A. |
| CD18<sup>+</sup>IgM<sup>-</sup> of CD19<sup>+</sup> (2.9–51.8) | N.A. | 7 | N.A. | N.A. |
| Lymphocyte proliferation | | | | |
| PHA % of lower limit of CD4 response | N.A. | 57 % | Reduced | Absent |
| PHA % of lower limit of normal CD3 response (>50 %) | 46 % ↓ | 53 % | Reduced | N.A. |
| PWM % of lower limit of CD4 response (>50 %) | 52 % | 100 % | Reduced | N.A. |
| ConA % of lower limit of CD3 response (>50 %) | 39 % ↓ | N.A. | Reduced | N.A. |
| Serum immunoglobulin concentrations | | | | |
| IgA mg/dL (7–13 m: 16–100; >6 y: 70–312) | 15 | 378 | Normal | Normal |
| IgM mg/dL (7–13 m: 25–115; >6 y: 56–352) | 2 ↓ | 18 | Normal | Normal |
| IgG mg/dL (7–13 m: 300–1500; >6 y: 639–1344) | 160 ↓ | 1520 (on IgG) | Normal | Normal |
| IgE IU/L (<100) | <2 | <1 | Normal | 9,856 ↑ |
| Specific antibody titers | | | | |
| Isohemagglutinins | 1:2 | 1:4 | Negative | Positive |
| Pneumococcal panel, 14 serotypes | 0 of 14 protective | 0 of 4 protective | N.A. | N.A. |
| Tetanus toxoid | Negative | Negative | Positive | Positive |
WES was performed on genomic DNA from the patient (pre-HCT) and nuclear family, with analysis under the hypothesis of an autosomal or X-linked recessive single gene disorder. The patient was a compound heterozygote with 2 non-synonymous variants affecting MALT1 exon 10 (isoform a, NM_006785.3) (Fig. 1b and c) (ClinVar number SCV000196510) [30]. As shown, the variants were sufficiently close to one another to determine that they resided on opposite haplotypes; all sequence reads showed one or the other, but not both variants: a splice acceptor defect, cDNA c.1019-2A > G (also in maternal DNA); and a single nucleotide deletion, cDNA c.1060delC, leading to a frameshift within the MALT1 paracaspase domain and truncation after 18 missense codons, p.Y353fs*18 (not in either parent or siblings, thus de novo in the patient).

Patient Mixed Donor Chimerism

Following HCT, mixed chimerism in the blood was established with 11.2% of peripheral blood nucleated cells and 16.6% of T cells being donor derived (Table 2). Using the TRA2G9 monoclonal antibody to label the patient’s, but not the donor’s, HLA-C antigens, the autologous PBMCs were distinguished by flow cytometry (Fig. 2a), showing 15.5% donor T cells and 3.4% donor B cells, in agreement with clinical determinations. Only 0.25% of the donor CD4 T cells were naïve CD45RA positive cells. Pre-HCT patient CD4+ CD25+ regulatory T cells (Tregs) and FOXP3 expressing T cells were low, but were restored after HCT (Table 2). Almost all Tregs and all active Tregs in the patient’s PBMCs were donor-derived (Fig. 2b).

Expression of MALT1 mRNA and MALT1 Protein

Autologous, TRA2G9 antibody-positive patient lymphocyte subsets were sorted from post-HCT patient PBMCs; MALT1 cDNA was prepared from these cells and from blood samples from the mother and a healthy control. The wild type MALT1 exon 10 contains an even multiple of 3 nucleotides encoding a predicted nuclear export signal and allowing for the possibility of in-frame exon skipping. Indeed, short cDNA amplicons with exon 9 joined to exon 11 were detected from patient and mother, but not controls, using the internal primers listed in Supplementary Table 1. However, larger amplicons lacking exon 10 were never observed, despite robust amplification of wild type cDNA including exon 10 from controls and even from the patient in trace amounts (not shown). No cDNA from the patient c.1060delC allele could be amplified, consistent with nonsense-mediated decay of mRNA from this early truncation allele.

Whole cell lysates from autologous, sorted patient PBMCs had no detectable MALT1 protein, while maternal heterozygous cellular MALT1 levels were reduced compared to the control (Fig. 2c), further indicating that c.1060delC is a null allele.
Functional Consequences of MALT1 Mutations in PBMCs

Phosphorylation of NF-κB and degradation of IκB following stimulation were analyzed to indicate the status of the CBM signalosome in PBMCs from the patient, his mother and a control. The patient’s autologous cells, identified by staining with the TRA2G9 antibody, were unable to phosphorylate NF-κB or degrade IκB (Fig. 3a, left and right panels, respectively), while cells from the mother and control cells demonstrated equivalent levels of NF-κB phosphorylation and IκB degradation in naïve and memory T cells as well as B cells. As noted above, too few donor-derived naïve T cells were detected in the patient’s blood to analyze. Phosphorylation of Erk, P38 and S6, each independent of the CBM signalosome, was intact in autologous patient cells, demonstrating selectivity of the MALT1 defect (Suppl Figure 1).

We also examined downstream effects of MALT1-dependent NF-κB signaling. After activation with PMA and ionomycin, donor-derived memory T cells and naïve and memory T cells from the mother and control had abundant IL-2 and IFN-γ, but the patient’s autologous, MALT1-defective T cells had little of either cytokine (Fig. 3b). Similar results were observed in PBMCs stimulated by the superantigen SEE or anti-CD3 plus anti-CD28 (Suppl Figure 2).

Reconstitution of Impaired NF-κB Signaling in MALT1 Defective Epstein-Barr Virus Transformed B Cell Lines

A pure TRA2G9-positive MALT1-defective sub-line was expanded from sorted patient EBV cells. This and a control EBV line were analyzed at rest and following stimulation, showing the same inability as T cells to phosphorylate NF-κB or degrade IκB in response to PMA and ionomycin (Fig. 3c and not shown). To conclusively implicate MALT1 in the functional deficits observed in CBM complex formation and downstream signaling, patient EBV cells were transduced with MP283 lentiviruses expressing mCherry alone or WT MALT1 cDNA and mCherry. While the former had no effect (Fig. 3c, middle panels), the vector restoring MALT1 normalized both NF-κB phosphorylation and IκB destruction (Fig. 3c, bottom panels).
Discussion

Our patient with profound CID and dysregulation adds to the 2 prior reports and extends our understanding of MALT1-associated disease and its therapy [22, 23]. Like the prior cases (Table 2), our patient had functionally impaired T and B cells leading to recurrent bacterial and viral infections from early life, notably with CMV, which in our patient was not controlled by antiviral therapy and required post-HCT donor T cell infusions for resolution. While HCT was recently postulated as treatment for MALT1-deficient CID [24], our report is the first of a MALT1-deficient patient cured by HCT. Two prior siblings died in childhood and a surviving teenager suffers significant multi-organ disease, including T cell inflammation of the skin and bowel similar to that in our patient.

MALT1 mutant patients reported to date had variable B cell numbers, serum immunoglobulin levels and ability to make specific antibodies (Table 2). Immune dysregulation consisting of prominent rash and suspected inflammatory bowel disease was shared between our patient and the living girl with W580S mutation [23]. In contrast to that patient, however, ours had normal B cell numbers, no IgE elevation and (like the deceased children with S89I mutation) absent protective antibody production.

Our patient’s MALT1 compound heterozygous mutations resulted in undetectable protein and MALT1 function. As with the previous reports, after introduction of wild type MALT1 cDNA, our patient’s mutant cells had MALT1 expression and NF-κB signaling reconstituted. Our case also highlights how MALT1 mutations may lead to immune dysregulation and autoimmunity. After HCT, almost all Tregs and all active...
Tregs in our patient’s PBMCs were donor-derived (Fig. 2b), accounting for the pre-HCT failure of autologous T cells to control auto-reactive attack on the skin and possibly the intestinal tract. The mixed chimerism exhibited by our patient post-HCT allowed us to evaluate both wild type and MALT1 deficient lymphocytes that had developed from hematopoietic progenitors in vivo. Moreover, our patient’s disease resolved even though he received a non-myeloablative preparative regimen (due to his high risk status with ongoing CMV infection) that did not result in a substantial circulating naïve T cell population. Whether the donor T cells originated from limited thymic development from CD34 progenitors vs. expanded donor T cells, or a combination of both, cannot be determined. However, the patient’s successful outcome indicates that only 15% of donor T cells were sufficient to reconstitute functional immunity and immune regulation.

The two heterozygous mutations in our patient abrogated MALT1 protein expression. Whereas the p.Y353fs*18 transcript was, as expected, degraded by nonsense-mediated mRNA decay, full-length transcripts skipping exon 10, due to the splice acceptor site defect, were also not detectable. MALT1 protein normally shuttles between the cytoplasm and nucleus with the aid of a nuclear export signal (NES1) [31, 32] and a regulatory region for NES1 encoded in exon 10, as well as second NES (NES2) in the C-terminal region of the protein [33, 34]. If a transcript missing exon 10 had been stable in our patient, the resulting protein might have been trapped in the nucleus. However, no nuclear or cytoplasmic protein from autologous patient cells was detected.

The high degree of consanguinity in the previously reported patients may indicate that loci other than MALT1 modified their immune phenotype as well as contributing to delayed bone age, fractures, short stature, and dysmorphia in the surviving patient and poor growth in the deceased siblings; these features were absent in our outbred patient who had compound heterozygosity at the MALT1 locus and has regained normal weight and stature following HCT.

Malt1−/− mice have defects in TCR activation and cytoplasmic production similar to those observed in humans with MALT1 deficiency [10, 35]. Malt1−/− mice demonstrate diverse B cell defects, as have humans, with NF-κB activity in B cells reduced in one report [36], but only marginally altered after immunoglobulin receptor engagement in another [35]. Mice deficient in CBM complex proteins have reduced thymic Treg cells [37, 38], but this has not been described in mice lacking Malt1. While the recently discovered human patients with CARD11 mutations had Treg deficiency, similar to the mouse model [20, 21], and Tregs were either normal or not studied in the previously described patients with MALT1 mutations, our patient’s autologous PBMCs had nearly absent Tregs and very low levels of Foxp3 expression. Thus, although Malt1 appears not to be required for development of Tregs in mice, it may be important in humans, in keeping with our patient’s resolution of dysregulated immune phenotypes following HCT that provided donor-derived Tregs.

When determining the underlying cause of immune deficiencies without lymphopenia and with normal TREC numbers, signaling molecules downstream of the antigen receptors have become candidates for analysis. As shown by our patient and others with defects in MALT1, as well as patients lacking CARMA1, NF-κB essential modulator, and IKK2 (IKBKB) [39], proteins important in antigen receptor and NF-κB signaling should be investigated in patients with combined immunodeficiency.

Conclusions

MALT1 deficiency can cause infantile combined immunodeficiency and immune dysregulation without T cell lymphopenia, but with impaired lymphocyte signaling through NF-κB, failure to generate memory and regulatory T cells, and hypogammaglobulinemia. Hematopoietic cell transplantation can be curative.

Acknowledgments

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