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Second-site mutation in the Wiskott-Aldrich syndrome (WAS) protein gene causes somatic mosaicism in two WAS siblings

Taizo Wada,1 Akihiro Konno,1 Shepherd H. Schurman,1 Elizabeth K. Garabedian,2 Stacie M. Anderson,1 Martha Kirby,1 David L. Nelson,3 and Fabio Candotti1

Introduction
The Wiskott-Aldrich syndrome (WAS) is an X-linked disorder that is characterized by thrombocytopenia, eczema, impaired cellular and humoral immunity, and increased susceptibility to lymphoid malignancies and autoimmunity (1). The gene responsible for WAS was identified by positional cloning and encodes the WAS protein (WASP), a 502–amino acid proline-rich peptide (2). WASP is constitutively expressed in cytoplasm of all nonerythroid hematopoietic cells (3, 4) and appears to be of central importance for signal transduction and cytoskeleton reorganization events of hematopoietic cells (5).

Revertant mosaicism due to true back mutations or second-site mutations has been identified in several inherited disorders. The occurrence of revertants is considered rare, and the underlying genetic mechanisms remain mostly unknown. Here we describe somatic mosaicism in two brothers affected with Wiskott-Aldrich syndrome (WAS). The original mutation causing disease in this family is a single base insertion (1305insG) in the WAS protein (WASP) gene, which results in frameshift and abrogates protein expression. Both patients, however, showed expression of WASP in a fraction of their T cells that were demonstrated to carry a second-site mutation causing the deletion of 19 nucleotides from nucleotide 1299 to 1316. This deletion abrogated the effects of the original mutation and restored the WASP reading frame. In vitro expression studies indicated that mutant protein encoded by the second-site mutation was expressed and functional, since it was able to bind to cellular partners and mediate T cell receptor/CD3 downregulation. These observations were consistent with evidence of in vivo selective advantage of WASP-expressing lymphocytes. Molecular analysis revealed that the sequence surrounding the deletion contained two 4-bp direct repeats and that a hairpin structure could be formed by five GC pairs within the deleted fragment. These findings strongly suggest that slipped mispairing was the cause of this second-site mutation and that selective accumulation of WASP-expressing T lymphocytes led to revertant mosaicism in these patients.

Polymerase slippage is also the most commonly invoked mechanism to explain triplet repeat expansion in human diseases (e.g., Huntington disease, fragile X syndrome, and Friedreich ataxia) (23).

Here, we described two WAS siblings with revertant mosaicism due to the same second-site mutation, likely caused by slipped mispairing. The original mutation responsible for the disease in this family is single base G insertion predicted to cause frameshift and premature termination of the protein. A second-site mutation was found in T lymphocytes from both brothers that carried a deletion of a 19 bp fragment, including the original mutation. This deletion resulted in restoration of the reading frame and the generation of an internally truncated, yet functional, WASP protein. Revertant T lymphocytes showed WASP expression and selective growth advantage in vivo over the WASP-negative (WASP–) cells. The occurrence of the same second-site mutation in two subjects from the same family strongly suggests a common underlying mechanism, and the characteristics of nucleotide sequence involved in the mutation are clearly compatible with slipped strand mispairing. From our observations we hypothesize that slipped mispairing is commonly occurring in the contest of this particular sequence, but only recognized in the rare cases in which selective advantage permits the accumulation of revertant cells above the detection threshold.

Methods

Case presentation. Figure 1 shows the pedigree of the patients’ family. Patient II-1 is an 18-year-old male who developed petechiae early after birth when his platelet count was in the range 7,000–20,000/µl. In the first year of life, he had eczema, otitis media, and conjunctivitis. After a clinical diagnosis of WAS was made at the age of 6 months, he underwent a splenectomy because of persistent thrombocytopenia. Between the age of 2 and 3 years, he had multiple episodes of pneumonia and recurrent leukocytoclastic vasculitis. Frequent ear infections and one episode of bronchitis are described until the age of 6, when he suffered from streptococcal pneumonia and sepsis. Between the ages of 12 and 16 years, he had documented episodes of medium vessel vasculitis leading to recurrent skin ulcers.

Patient II-1’s younger brother (II-2) also developed thrombocytopenia early after birth and underwent a splenectomy at the age of 6 months. The postoperative clinical history is remarkable for occasional otitis media and continued severe eczema until the age of 9 years, when he developed molluscum contagiosum, which progressed unremittingly for 3 years until all extremities and truncal surfaces accessible to hands were affected. At age 13, he developed non-Hodgkin B cell lymphoma, which relapsed after chemotherapy at the age of 15. Both clinical histories are consistent with severe WAS phenotype (score of 5) (24). No clinical improvement with age has yet been observed in either patient. No history of other affected males was noted in the maternal side of the family.

Cell preparation. PBMCs were isolated by Ficoll-Hypaque (Mediatech Inc., Washington, DC, USA) gradient centrifugation from the patients and healthy controls. For T lymphocyte cultures, PBMCs were maintained in the presence of 100 ng/ml of anti-CD3 (OKT3; Ortho Diagnostic Systems Inc., Raritan, New Jersey, USA), 5 µg/ml of anti-CD28 mAb (PharMingen, San Diego, California, USA), and 100 IU/ml of recombinant IL-2 (rIL-2) (gift of C. Reynolds, National Cancer Institute, Frederick, Maryland, USA) for 3 days in RPMI-1640 medium (Life Technologies Inc., Rockville, Maryland, USA) containing 10% FBS (Gemini Biological Products, Woodland, California, USA) and antibiotics. Cells were then cultured in RPMI-1640 plus 10% FBS medium supplemented with 100 IU/ml of rIL-2 for 12 days. Human T lymphotropic virus type 1–transformed WAS T cell lines carrying the 57delG “null” mutation (25) were maintained in the same medium. MOLT-4 cells were obtained from American Type Culture Collection (Manassas, Virginia, USA).

Flow-cytometric analysis of WASP and cell sorting. Flow cytometric analysis of WASP was performed as described previously (15) with minor modifications. PBMCs were stained for cell surface antigens prior to cell membrane permeabilization using the following mAb’s: CyChrome-conjugated anti-CD3 or anti-CD20 (PharMingen); TRIC-color–labeled anti-CD8 (Caltag Laboratories Inc., Burlingame, California, USA); FITC-conjugated anti-CD4 (Caltag Laboratories Inc.); FITC-conjugated anti-CD45RA and anti-CD45R0 (PharMingen); and FITC-conjugated anti-CD14 or anti-CD36 (Becton Dickinson Immunocytometry Systems, San Jose, California, USA). After washing, cells were fixed and permeabilized with CytoStain kit (PharMingen) and were incubated with 1:200 diluted anti-WASP (3F3-A5) mAb (3) or purified mouse IgG-1 (PharMingen) at 4°C for 20 min. Cells were then reacted with biotin-conjugated anti-mouse IgG-1 at 4°C for 20 min, followed by further incubation with phycoerythrin-conjugated (PE-conjugated) streptavidin (both from PharMingen). Stained cells were analyzed with a FACScan flow cytometer and the CellQuest software (Becton Dickinson Immunocytometry Systems). For sequencing analysis, WASP-expressing (WASP+) and WASP– cells were purified from cultured T lymphocytes using a FACSVantage cell sorter (Becton Dickinson Immunocytometry Systems). The purity of

Figure 1

Pedigree of the study family. Solid symbols represent affected individuals, and the carrier status of subject I-1 is indicated by the central dot.
sorted WASP+ and WASP– T lymphocytes was more than 98%, as determined by a flow cytometric analysis.

For mutation detection in lineage-specific cells using fluorescent genotyping, PBMCs and polymorphonuclear cells (PMNs) were separated by Ficoll-Hypaque gradient centrifugation. CD3+ T cells were isolated from PBMCs and PMNs with immunomagnetic beads (Dynal Inc., Lake Success, New York, USA), following manufacturer’s instructions. CD3-depleted PMNs were frozen and stored until use. The CD3-negative PBMC fraction was divided into three aliquots that were separately stained with PE-conjugated anti-CD14, CyChrome-conjugated anti-CD20 or PE-conjugated anti-CD20, or CyChrome-conjugated anti-CD56 (all from PharMingen). Both aliquots were also stained with FITC-conjugated anti–TCR-αβ and anti–TCR-γδ (both from Becton Dickinson Immunocytometry Systems) for detection of residual T lymphocytes. CD14+/TCR-αβ, TCR-γδ, CD20+/TCR-αβ, TCR-γδ, and CD56+/TCR-γδ cells were then isolated using a FACS Vantage cell sorter (Becton Dickinson Immunocytometry Systems).

Western blot analysis and in vitro binding assay. Western blot analysis of WASP and in vitro binding assay were performed as described (25, 26). Briefly, for Western blot analysis, cell lysates (10–100 µg) were boiled, subjected to 8% SDS-PAGE, and electrotransferred onto nylon membranes (Immobilon-P; Millipore Corp., Bedford, Massachusetts, USA). For the in vitro binding assay, 100 µg of cell lysate protein was incubated with 5 µg of Fyn-Src homology 3 (SH3) glutathione S-transferase (GST) fusion protein bound to agarose beads (Santa Cruz Biotechnology Inc., Santa Cruz, California, USA) and Grb2-GST agarose (Upstate Biotechnology Inc., Lake Placid, New York, USA) in 500 µl of lysis buffer (300 mM NaCl, 50 mM Tris-HCl, 2 mM EDTA, 0.5% Triton X-100, 2.5 mM p-nitrophenyl ρ′-guanidino-benzoate, and 10 µg/ml aprotinin and leupeptin) for 2 h at 4°C. Immunoprecipitates were then washed, boiled, and resolved using 12% SDS-PAGE. Detection of WASP was performed using the anti-WASP (3F3-A5), or B-9 (Santa Cruz Biotechnology Inc.) mAb’s, a rabbit polyclonal Ab purchased from Upstate Biotechnology Inc. A rabbit polyclonal Ab specific for STAT5a was used for reprobing of membranes as described previously (27).

DNA extraction and mutation analysis of the WASP gene. DNA was extracted from blood samples and cell lines using the QIAamp DNA blood mini kit (QIAGEN Inc., Valencia, California, USA). Each exon of the WASP gene was amplified from genomic DNA using specific primers (15). Sequencing was performed on gel-purified PCR products or PCR products subcloned in the pCR2.1 vector (Invitrogen Corp., Carlsbad, California, USA) using the ABI Prism BigDye Terminator Cycle sequencing kit on an ABI Prism 310 Genetic Analyzer (Perkin-Elmer Applied Biosystems, Foster City, California, USA), as described previously (15).

For fluorescent genotyping (GeneScan) analysis, the nucleotide 1150–1358 fragment of the WASP genomic DNA was amplified from 300 ng of DNA extracted from PBMCs, immunomagnetic bead-purified CD3+ T lymphocytes, CD3+ cell-depleted PMNs and sorted CD20+ B lymphocytes, CD56+ NK cells, and CD14+ monocytes, using 20 pmol each of the 5′-TCC AGC TAC TGG AGC TTC TG-3′ forward primer and the FAM-labeled 5′-TTC CTC GCC GGA TTT GAT-3′ reverse primer in a 50-µl volume reaction containing 1.25 U Ex Taq polymerase, 1× Ex Taq Buffer, 2 mM Mg2+, and 200 µM dNTPs (all from Takara Bio Inc., Shiga, Japan). Thirty-five amplification cycles (30 s at 94°C, 30 s at 60°C, 30 s at 72°C) were carried out. One microliter of the amplification product was then combined to 12 µl of deionized formamide and 0.5 µl of GeneScan size standard (Perkin-Elmer Applied Biosystems) and analyzed on an ABI Prism 310 Genetic Analyzer using the GeneScan Analysis Software (Perkin-Elmer Applied Biosystems). Titration studies showed that this procedure allowed detection of one copy of the Δ19bp mutation present within 5,000 cells (data not shown).

Analysis of T cell receptor-β chain gene rearrangements. T cell receptor-β (TCR-β) gene rearrangements were detected by family-specific PCR in combination with the GeneScan technique as described recently (28), with minor modifications. Briefly, 50 ng of DNA extracted from sorted WASP+ and WASP– T lymphocytes was subjected to PCR amplification using a TCR βv consensus primer [βv pan: 5′-CTC GAA TTC T(T/G) T(A/T) (C/T)T GGT A(C/T) C(G/A)T/A/CA-3′, 30 pmol] and a TCR-β family-specific primer (βv 2.1: 5′-ACG GTG GCC GGT-CC-3′, 10 pmol) (28) in a 50-µl volume reaction containing 1.25 U Ex Taq polymerase, 1× Ex Taq buffer, 2 mM Mg2+, and 200 µM dNTP (all from Takara Bio Inc.). Thirty-five amplification cycles (30 s at 94°C, 40 s at 57°C, 40 s at 72°C) were carried out. One microliter of the amplification product was then subjected to a seminested reaction using the same amplification conditions, but using a FAM-labeled βv pan primer in combination with a nested βv 2.1–specific primer (5′-AGG-CTG CTC GCC-GCC-GGA-3′) (28). One microliter of the second PCR reaction product was combined with 12 µl of deionized formamide and 0.5 µl of GeneScan size standard (Perkin-Elmer Applied Biosystems) and analyzed on an ABI Prism 310 Genetic Analyzer using the GeneScan Analysis Software (Perkin-Elmer Applied Biosystems). DNA extracted from MOLT-4 cells with known clonal TCR-β2.1 rearrangement was used as positive control.

WASP vector constructs and transductions. The GCWASP retroviral vector expressing the wild-type WASP and the neomycin resistance (Neo) genes and the transduction procedures used for gene transfer into WAST cells have been described previously (25). Briefly, GCWASP viral particles pseudotyped with the gibbon ape leukemia virus envelope were produced using the PG13 cell line (29). To construct retroviral vectors carrying the patients’ mutations, the exon 10 fragment of wild-type WASP was replaced at the XhoI and EcoRI sites with sequence-verifying PCR products of the mutated exon 10 sequences amplified from patient II-1. The SAM-EN vector expressing only the neomycin resistance gene (30) was used as
negative control. For transductions, WASP-null T cell lines were incubated in phosphate-free RPMI-1640 medium (Life Technologies Inc.) for 6 h, resuspended in retroviral supernatant supplemented with 8 µg/ml polybrene (Sigma-Aldrich, St. Louis, Missouri, USA), and then transferred to 24-well plates. After centrifugation at 1,800 g for 30 min, cells were incubated at 37°C overnight. The following day, the procedure was repeated. After transduction, cells were selected in 1.0 mg/ml of G418 (Geneticin; Life Technologies Inc.) for 12 days.

**Analysis of the inserted proviral genome by semiquantitative PCR.** WASP exon 3 and 4 sequences were amplified from genomic DNA using sense (5′-GGA ACA GGA GCT GTA CTC AC-3′) and antisense (5′-TCC ACT TTG CCT CTG ATT CC-3′) primers and 30 cycles of amplification (94°C for 30 s, 60°C for 30 s, and 72°C for 30 s). The endogenous WASP gene fragment containing the intron 3 sequence (100 bp) and the retroviral-derived integrants generate fragments of 272 bp and 172 bp, respectively. Amplified products were separated on a 3% agarose gel and visualized by ethidium bromide staining. The intensity of each band was quantified using the BIT Image software (Aladdin Systems Inc., Watsonville, California, USA) and the NIH Image software. The ratio between the intensity of the endogenous WASP gene and the retroviral transgene products was then calculated as described (31).

**TCR/CD3 downregulation assay.** T cells (10⁶) were incubated with 1 µg/ml anti-CD3 mAb (Leu4; Becton Dickinson Immunocytometry Systems) for 30 min on ice. After washing, cells were reacted with 5 µg/ml biotinylated goat anti-mouse Ab (BioSource International, Camarillo, California, USA) for 30 min on ice and then warmed at 37°C for 60 min to allow downregulation of CD3. Cells were then transferred to ice, washed, and stained with PE-conjugated streptavidin (PharMingen). After fixation in 2% paraformaldehyde, CD3 expression was analyzed by FACS. Data were reported as the percentage of mean fluorescent intensity of control cells not subjected to incubation at 37°C (25). Analysis of differences among data groups was performed using the Student t test for paired samples. P values less than 0.05 were considered significant.

**Results**

**Patients’ immunophenotype and mosaic expression of WASP.** The immunophenotypic characteristics of the patients’ T lymphocytes are described in Table 1. As part of the clinical evaluation of our WAS patients, we analyzed WASP expression in PBMCs by multicolor flow cytometry. As shown in Figure 2, total lymphocytes as well as T, B, and NK cell subsets and CD14+ monocytes from a

![Figure 2](image-url)
healthy control expressed WASP. In contrast, patients II-1 and II-2 exhibited no detectable expression of WASP in CD14+ monocytes, but showed a sizable fraction of WASP+ cells among their lymphocytes, suggesting the presence of somatic mosaicism. WASP+ cells were found to include CD3+ T cells, but were not detected among CD20+ B cells and CD56+ NK cells in either patient. Both CD4+ and CD8+ T cells were involved in the mosaicism for WASP expression; however, WASP+ cells were detected only in less than 10% of the patients’ CD8+ T cells. Interestingly, more than half of CD45R0+CD3+ memory T lymphocytes expressed WASP in both patients, whereas WASP+ cells were detected in only less than 10% of CD45RA+CD3+ naïve T cells.

Restoration of the WASP gene reading frame by a second-site mutation in revertant cells. The mutation causing WAS in this family was described previously as a single base G insertion after nucleotide 1305 in exon 10 of the WASP gene (insG) (32). To determine whether the same mutation was present in WASP+ T cells, direct sequence analysis was performed on genomic DNA obtained from sorted WASP+ and WASP– T cells from patient II-1. As shown in Figure 3a, we found the insG mutation in DNA from the WASP– T cells, whereas DNA from the WASP+ T cells showed a second-site mutation, represented by a 19-bp deletion (Δ19bp) encompassing the original insG mutation. When we analyzed the sequence of WASP exon 10 in subcloned PCR products obtained from unsorted cultured T lymphocytes of patient II-1, the Δ19bp mutation was detected in 18.2% of 22 clones. In patient II-2, the coexistence of the Δ19bp mutation was also demonstrated in DNA obtained from his PBMCs (4.5% of 22 clones). Sequence analysis of 100 normal chromosomes demonstrated the absence of the Δ19bp mutation in the general population. Analysis of the DNA sequence surrounding the Δ19bp mutation showed the presence of two 4-bp (CCTG) direct repeats and structural characteristics compatible with the formation of a hairpin loop containing five GC pairs within the deleted fragment (Figures 3, a and b).

Figure 3c shows the predicted structure of the mutated WASP molecules. The insG mutation causes a frameshift resulting in a novel sequence beginning at AA 424 and concluding in a premature termination at position 493. The Δ19bp mutation leads to an internal deletion of 6-AA at the end of the proline-rich region, but to restoration of the WASP reading frame beginning at AA 427.

To determine the characteristics of WASP in the revertant cells, Western blot analysis was performed using the patients’ cultured T lymphocytes. As shown in Figure 3d, T lymphocytes from a healthy control showed the presence of the expected approximately 59-kDa WASP band. In contrast, lysates from the patients’ lymphocytes showed a WASP band of lower molecular weight that was consistent with the predicted effects of the Δ19bp mutation.

Genotypic analysis of lymphohematopoietic lineages. Purity of magnetic bead-isolated CD3+ T cells and CD3-depleted polymorphonuclear cells was more than 98%, while sorted CD20+ B lymphocytes, CD56+ NK cells, and CD14+ monocytes contained less than 0.05% contaminating TCR-αβ+/TCR-γδ+ cells (data not shown). Fluorescent genotyping of purified lymphocyte and myeloid cells showed clear evidence of the Δ19bp mutation only in WASP+ cells.

Figure 3
Characterization of WASP gene mutations and expression. (a) The WASP gene exon 10 sequence was amplified from DNA extracted from normal PBMCs, as well as WASP− T and WASP+ T lymphocytes of patient II-1. Direct sequencing was performed using an automated sequencer. A thick bar highlights the position of the original insG mutation and thin bars show the 4-bp repeat sequences. (b) Model for the generation of hairpin structure facilitating the second-site mutation by slipped mispairing and deletion of 19 bp. (c) Predicted structures of mutated WASP molecules. PH, pleckstrin homology; WH, Wiskott homology; GBD, GTPase-binding domain; PPPP, proline rich. del, deletion. (d) Western blot analysis of WASP was performed using lysates of cultured T lymphocytes obtained from a healthy control and the patients. Arrows indicate the position of the WASP band and of a nonspecific protein reactive with the secondary Ab (NS). IB, immunoblotting.
among CD3+ T cells (Figure 4a). Genotyping and sequence analysis of DNA extracted from PBMCs of the patients’ mother (subject I-1) showed that she was heterozygous carrier of the insG mutation, but did not show detectable evidence of the \(\Delta 19\)bp mutation (Figure 4b and data not shown).

**Policlonality of WASP+ T cells.** High-resolution (Gene Scan) analysis of TCR-\(\beta\) gene rearrangements showed a Gaussian-like distribution of the sizes of the TCR-\(\beta\) amplificates in T lymphocytes from a healthy control and from sorted WASP+ T cells from patients II-1 and II-2 (Figure 5). TCR-\(\beta\) rearrangements from the patients’ WASP+ cells were characterized by a lower number of peaks, suggesting that although polyclonal, these cells carried a TCR repertoire of reduced complexity. As expected, leukemic MOLT-4 cells showed a single peak indicative of monoclonal TCR-\(\beta\) rearrangement (Figure 5).

**Retrovirus-mediated expression of mutated WASP.** To assess the expression and functional characteristics of the WASP mutants, retroviral vectors carrying wild-type WASP cDNA, the insG, and the \(\Delta 19\)bp mutants were used to transduce WAS T cell lines carrying a null mutation of the WASP gene with no detectable WASP expression. As shown in Figure 6a, T cells untransduced or transduced with the Neo control vector did not express WASP, whereas WASP expression was well reconstituted in T cells transduced with the vector carrying the wild-type WASP cDNA. T cells transduced with the insG mutant showed a very faint WASP band of lower molecular weight, as predicted by the early termination at codon 493. In contrast, T cells transduced with the \(\Delta 19\)bp mutant showed near normal levels of a WASP molecule with size characteristics compatible with the predicted internal deletion of 6 AA. As shown in Figure 6b, transgene-specific sequences were detected in T cells transduced with each of the retroviral vectors (wild type, insG, and \(\Delta 19\)bp), but not in untransduced T cells or cells transduced with the Neo control vector. Semi-quantitative estimation of transgene copy number showed that all T cells transduced with the vectors carrying the WASP sequence had approximately one copy of the provirus per cell after G418 selection. These results indicate that low expression of WASP in T cells transduced with the insG mutation was neither due to low transduction efficiency nor to a failure of G418 selection, but more likely to instability of the WASP mRNA or protein carrying the insG mutation. Even when different anti-WASP Ab’s were used for the detection of WASP (see Methods), T cells transduced with the insG mutant showed only minimal WASP expression of the mutated WASP (data not shown).
We next analyzed the binding ability of the mutant proteins to the SH3-domain containing proteins Fyn and Grb2. SH3 partners have been shown to interact with WASP through its proline-rich region (32), which is affected by the mutations found in these patients (Figure 3c). As expected and consistent with our previous experience (25), Fyn and Grb2 were able to bind in vitro to WASP from lysates of T cells transduced with wild-type WASP cDNA, and no product was precipitable from lysates of cells transduced with the Neo control vector (Figure 6c). In contrast, the amount of WASP precipitated by SH3-GST agarose beads was extremely low in lysates from T cells transduced with the insG mutant. Importantly, T cells transduced with the Δ19bp mutation showed restored SH3-interaction comparable with the effect of wild-type WASP.

To further assess the functionality of the mutant proteins, we examined CD3-mediated downregulation of TCR/CD3 in transduced cells. As shown in Figure 6d and consistent with our previous results (25), the levels of TCR/CD3 expression decreased 1 h after simulation in T cells transduced with wild-type WASP cDNA and was impaired in T cells transduced with the Neo control vector. Interestingly, the insG mutant failed to correct the downregulation defect, while T cells transduced with the Δ19bp mutation showed significant improved responses to anti-CD3 stimulation.

**Detection of circulating WASP<sup>+</sup> cells at various time points.** To evaluate whether the accumulation of the WASP<sup>+</sup> T cells varied with time, a flow-cytometric analysis of WASP was performed using PBMCs of patient II-1 collected at 8, 11, and 18 years of age. As shown in Table 2, WASP<sup>+</sup> cells were detected in lymphocytes from all analyzed samples, and, importantly, the percentage of WASP<sup>+</sup> T cells was shown to increase with the age of the patient.

### Discussion

Somatic revertant mosaicism due to back mutations or second-site mutations has been reported recently in several genetic diseases: tyrosinemia type 1 (6), Bloom syndrome (7), epidermolysis bullosa (9, 16), Fanconi anemia (10, 11, 17), and primary immunodeficiencies, including adenosine deaminase deficiency (12, 13, 18), X-linked severe combined immunodeficiency (8), and WAS (14, 15). The molecular mechanism of such events remains mostly unclear, except for the few cases where intragenic recombination (7, 10), gene conversion (9), or DNA polymerase slippage (11, 15, 18) has been shown. The real frequency of the occurrence of revertant mosaicism also remains to be determined, although it is assumed to be extremely low regardless of the mutagenic mechanism. Here we describe the unusual case of two WAS siblings presenting with somatic mosaicism due to the same second-site mutation, a deletion of 19 bp, including the original single nucleotide insertion. The Δ19bp mutation was observed only in the patients’ T lymphocytes and led to reconstitution of WASP expression. Sequencing analysis of the WASP gene in the general population excluded the possibility that the Δ19bp mutation could be a functional polymorphism. The Δ19bp mutation was not detectable in PBMC samples from the patients’ mother, although somatic mosaicism would be difficult to detect in women due to X-chromosome inactivation.

### Table 2

<table>
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<tr>
<th>Age (yr)</th>
<th>8.4&lt;sup&gt;a&lt;/sup&gt;</th>
<th>11.4&lt;sup&gt;a&lt;/sup&gt;</th>
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<td>T lymphocyte subsets (%)</td>
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<td>CD4&lt;sup&gt;+&lt;/sup&gt;CD8&lt;sup&gt;+&lt;/sup&gt;CD3&lt;sup&gt;+&lt;/sup&gt;</td>
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<td>WASP&lt;sup&gt;+&lt;/sup&gt; cells in lymphocyte subsets (%)</td>
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<td></td>
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<td>43.6</td>
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<sup>a</sup>Immunophenotyping and flow cytometric analysis of WASP were performed on frozen samples.
lack of selective advantage. Both patients, however, had the normal 46 XY male karyotype, thus ruling out the possibility that the WASP \(^{19}\) T lymphocytes could have arisen from maternal T cell engraftment. It is also extremely unlikely that both patients acquired the same mutant T cells postnataally from infusions of blood products. We therefore conclude that the T lymphocytes carrying the \(\Delta 19bp\) mutation originated in both patients in vivo due to a second-site mutation.

Revertant mosaicism due to a second-site mutation has been reported previously (16–18). In particular, the occurrence of the same second-site mutation in two brothers with Fanconi anemia has been reported by Waisfisz et al. (17), who proposed the presence of the same second-site mutation having resulted from the same specific molecular mechanism in the two subjects. Similarly, our observations strongly suggest that a common underlying mechanism resulted in the occurrence of the same second-site mutation in the two brothers with WAS, described here. Slipped mispairing is a well-accepted mechanism for genomic instability, resulting in insertions and deletions of repetitive sequences in many genetic disorders (19–21). Analysis of the sequence surrounding the \(\Delta 19bp\) second-site mutation (Figure 3c) leads us to propose that slipped mispairing is at the basis of our findings. We argue that the presence of two 4-bp direct sequences (CCTG) and the hairpin structure created by five GC pairs within the deleted fragment are able to induce slippage and mispairing of the nascent and template strands during DNA replication.

The slipped mispairing event leading to the \(\Delta 19bp\) deletion may have occurred in a primitive pluripotent hematopoietic progenitor or in a committed T cell precursor. The latter possibility is supported by the notion that WAS-carrier women preferentially inactivate the X chromosome carrying the mutated WASP gene in hematopoietic progenitors (33, 34) and all mature hematopoietic cells (35, 36). WASP \(^\ast\) cells deriving from a revertant pluripotent hematopoietic progenitor would therefore be expected to show selective advantage among all lineages, whereas we could only detect WASP \(^\ast\) cells within the CD3\(^{+}\)/CD4\(^{+}\) and CD3\(^{+}\)/CD8\(^{+}\) T lymphocytes from our two patients.

On the other hand, the occurrence of the revertant mosaicism in both patients suggests that the second-site mutation is not an extremely rare event. It seems unlikely, therefore, that such occurrence would coincidentally target only the same cell-differentiation stage in both patients. On the contrary, slipped mispairing could be a relatively common event with respect to nucleotide 1299–1316 of the WASP sequence and, therefore, that the \(\Delta 19bp\) deletion may occur in a variety of cell types, but become apparent only in cells characterized by a long life span that can accumulate over time. To the detection limits of our genotyping assay, we could clearly demonstrate the presence of the \(\Delta 19bp\) deletion only among patients’ purified T lymphocytes, however. These findings, taken together with the detection of WASP \(^\ast\) cells among both CD4\(^{+}\) and CD8\(^{+}\) T lymphocytes and the observation of polyclonal TCR-\(\beta\) gene rearrangements in WASP \(^\ast\) T cells, allow placement of the second-site mutation event at the level of a T cell progenitor cell before CD4/CD8 lineage commitment and TCR rearrangement.

Consistent with our previous observations (15), there is evidence of in vivo selective advantage of WASP \(^\ast\) T lymphocytes in these two patients. Flow-cytometric studies showed that 2.4–6.2% WASP \(^\ast\) cells appeared among CD45RA\(^{+}\) naive T cells, while more than half of the CD45RO\(^{+}\) memory T cells were WASP \(^\ast\) in both patients. These findings clearly indicate WASP \(^\ast\) T cells had selective growth advantage over WASP \(^\ast\) counterparts during the differentiation and proliferation steps leading to the acquisition of the immunological memory. The reasons why the patients showed higher proportion of revertant CD4\(^{+}\) T lymphocytes than CD8\(^{+}\) T cells are presently unclear. Immunophenotyping of T lymphocyte subsets demonstrated an increased number of activated memory CD4\(^{+}\) T lymphocytes in both patients (Table 1). It is therefore possible that peripheral expansion of these CD4\(^{+}\) T lymphocytes had favored the selective advantage of revertant cells within this subset of lymphocytes.

Our data also suggest that the expression of the internally deleted WASP mutant generated by the \(\Delta 19bp\) mutation was responsible for the in vivo selective advantage of revertant T cells through reconstitution of WASP function. It has been recognized recently that WASP function is regulated through an autoinhibitory conformation involving intramolecular interaction between the GTPase-binding domain and the verprolin homology, cofilin homology, and acidic (VCA) region of WASP (37). The original insG mutation destroys the VCA region, thus likely interfering with the protein tertiary structure and resulting in instability. On the contrary, the \(\Delta 19bp\) mutation results in the deletion of 6-AA at the end of the proline-rich region but restores the sequence of the VCA domain, which can regenerate its ability to interact to the GTPase-binding domain, thus resulting in a more stable protein (38). Accordingly, our in vitro reconstitution experiments showed that WASP expression, binding to SH3 partners, and TCR/CD3 downregulation in T cells carrying the \(\Delta 19bp\) mutant were compatible with those observed in the presence of wild-type WASP and indicated that the \(\Delta 19bp\) mutant is functional.

The availability of archived PBMC samples from patient II-1 offered us a unique opportunity to study the accumulation of WASP \(^\ast\) T lymphocytes over time in this patient. We were able to detect WASP expression in his lymphocytes at age 8, and the percentage of WASP \(^\ast\) cells increased with age, suggesting that the accumulation of WASP \(^\ast\) cells in peripheral blood had started more than 10 years ago. Neither of the two patients in this study, however, have yet shown improvement of their clinical phenotype. This is in contrast to a previously described case of somatic mosaicism in a WAS patient carrying more than 80% of revertant T lymphocytes (15), but is consistent with the finding of Ariga et al.
who did not observe clinical improvement in another case of WAS with revertant mosaicism (14). One explanation for the lack of clinical benefit from the occurrence of the second-site mutation in the patients described here is that the absolute numbers of WASP-negative lymphocytes, and especially revertant naive T lymphocytes, were simply not high enough to correct the WASP phenotype. If this is the case, it is possible that an improvement will be observed in the future because of the selective accumulation of WASP-positive lymphocytes. Alternatively, the function of the mutant WASP carrying the A19bp mutation may not be as effective in vivo as the wild-type WASP.

In summary, our studies provide evidence for potential beneficial outcomes of slipped mispairing events and tie this genetic mechanism to the generation of somatic mosaicism. Although our current knowledge regards back mutations or second-site mutations as extremely rare events, the observation of the present two cases raises the possibility that genetic reversions may take place more often than is commonly accepted, but remain undetected because they do not necessarily result in modification of the clinical phenotype. Because revertant mosaicism is a natural form of gene therapy, the identification of revertants and characterization of mechanism underlying back mutations or second-site mutations is obviously of great importance for the development of new therapeutic strategies for genetic disorders and deserve active investigation.

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