T^B^NK^ severe combined immunodeficiency caused by complete deficiency of the CD3ξ subunit of the T-cell antigen receptor complex

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CD3ξ is a subunit of the T-cell antigen receptor (TCR) complex required for its assembly and surface expression that also plays an important role in TCR-mediated signal transduction. We report here a patient with T^B^NK^ severe combined immunodeficiency (SCID) who was homozygous for a single C insertion following nucleotide 411 in exon 7 of the CD3ξ gene. The few T cells present contained no detectable CD3ξ protein, expressed low levels of cell surface CD3ξ, and were nonfunctional. CD4^CD8^CD3ξlow, CD4^CD8^CD3ξlow, and CD4^CD8^CD3ξlow cells were detected in the periphery, and the patient also exhibited an unusual population of CD56^CD16^ NK cells with diminished cytolytic activity. Additional studies demonstrated that retrovirally transduced patient mutant CD3ξ cDNA failed to rescue assembly of nascent complete TCR complexes or surface TCR expression in CD3ξ-deficient MA5.8 murine T-cell hybridoma cells. Nascent transduced mutant CD3ξ protein was also not detected in metabolically labeled MA5.8 cells, suggesting that it was unstable and rapidly degraded. Taken together, these findings provide the first demonstration that complete CD3ξ deficiency in humans can cause SCID by preventing normal TCR assembly and surface expression. (Blood. 2007; 109:3198-3206)

Introduction

Severe combined immunodeficiency (SCID) is a syndrome characterized by absent T- and B-lymphocyte function that is uniformly fatal in infancy without successful immune reconstitution.1-3 Several different molecular etiologies of SCID in humans have been described. These include mutations in genes encoding components of lymphocyte cytokine receptors,4-11 gene products responsible for T- and B-cell antigen receptor VDJ recombination,12-17 proteins instrumental in lymphocyte survival18 or function,19,20 and structural subunits of the T-cell antigen receptor (TCR) complex.21,22 The multimeric TCR complex is composed of a clonotypic TCRβ or TCRγδ heterodimer associated with invariant CD3 (CD3γ, CD3δ, CD3ε, and CD3ξ) chains.23-26 TCR complexes are assembled in the endoplasmic reticulum of mature T cells in stepwise fashion, with the final stage being association of CD3 homodimers with incomplete TCRβ-CD3γ-CD3δ-CD3ε complexes.27-35 The addition of CD3ξ subunits is critical for survival and efficient transport of TCR complexes to the plasma membrane,31,34 as incomplete TCRβ-CD3γ-CD3δ-CD3ε complexes are rapidly degraded in lysosomes.34 TCR complex ligand-binding specificity is provided by the clonotypic TCRβ or TCRγδ heterodimer,36 whereas CD3 chains serve as signal transducing subunits via their cytoplasmic immunoreceptor tyrosine-based activation motifs (ITAMs).37-40 CD3γ, CD3δ, and CD3ε chains each contain one ITAM; CD3ξ contains 3. Phosphorylation of CD3 chain ITAMs following TCR ligand engagement results in recruitment and activation of ZAP-70, a protein tyrosine kinase required for normal T-cell signaling.41,42 Hence, surface TCR complex expression is required for both antigen recognition and signal transduction following ligand binding in mature T cells. TCR expression is also vital for T-cell development in the thymus43-46 and murine gene knockout studies have demonstrated the importance of individual components of the TCR complex in this process. Mice lacking expression of TCRβ,47 CD3γ,48 or CD3ε49 exhibited a block at the CD4^-CD8^- stage of thymocyte development, whereas CD4^CD8^- cells accumulated in TCRα-deficient57,50 or CD3δ-deficient animals.51 Development of CD4^CD8+ thymocytes, CD4+ and CD8+ single-positive thymocytes, and peripheral T cells was markedly diminished in mice lacking CD3ξ.52-55 Abnormalities in expression of CD3 subunits of the TCR complex have also been reported in humans. Absence of CD3δ expression led to SCID with a block in T-cell ontogeny prior to the CD4^-CD8^- stage of thymocyte development.21,22 A homozygous CD3ξ gene mutation expected to prevent protein expression has been described in a SCID patient.22 Partial CD3ξ deficiency56,57 and complete CD3γ deficiency58,59 did not completely block T-cell development and resulted in milder immunodeficiency. A complex case of CD3ξ deficiency partially corrected by somatic mutations has recently been reported in a patient with recurrent infections and decreased numbers of peripheral T cells.60 In the present study we describe a unique infant with T^B^ NK^ SCID due to complete CD3ξ deficiency.
Patient, materials, and methods

Patient

The patient was the child of unrelated parents of Chamorro descent from Guam. She presented with pneumonia of unknown etiology at 4 months of age and subsequently developed a chronic cough, recurrent otitis media, failure to thrive, a chronic mild rash, and one episode of Salmonella gastroenteritis. At age 10 months she was hospitalized for thrombocytopenia and found to have a cytomegalovirus (CMV) infection. Initial immune studies in Guam and Hawaii revealed very low numbers of circulating T cells (141/mm$^3$) and absent T-cell proliferative responses. The patient was referred to Duke University Medical Center where the diagnosis of SCID was confirmed, treatment of her CMV infection was begun, and, at age 12 months, she received a T cell-depleted haploidentical bone marrow transplant from her mother without pretransplantation chemotherapy conditioning or posttransplantation graft-versus-host disease (GVHD) prophylaxis. She showed no evidence of immune function over the next 4 months so received a second T cell-depleted maternal marrow transplant without preconditioning or GVHD prophylaxis at 16 months of life. The patient continued to show signs of T-cell chimerism or function after her second transplant perhaps due, at least in part, to ongoing CMV viremia. She received a third stem cell transplant from her father without preconditioning or GVHD prophylaxis and remains clinically stable at 1 month after transplantation.

Immunologic phenotype analysis

Standard 4-color flow cytometry of peripheral blood lymphocytes was performed with labeled antibodies (Abs) to CD3 (clone SK7 or clone BB7.2), CD4, CD8, CD14, CD16, CD20, CD45, CD56 (clone T10B9.1A-31), TCR (clone B1), and HLA-A2 (clone BB7.2) purchased from BD Biosciences (San Jose, CA). A third unshared maternal histocompatibility antigen. To be free of detectable maternal cells by staining with antibody to HLA-A2 (clone SK7; BD Biosciences), an unshared maternal histocompatibility antigen.

Table 1. Patient immunologic phenotype

<table>
<thead>
<tr>
<th>Lymphocyte subpopulation</th>
<th>Patient</th>
<th>Controls</th>
</tr>
</thead>
<tbody>
<tr>
<td>CD3$^{\text{high}}$</td>
<td>17 (0.4)</td>
<td>1111-5183</td>
</tr>
<tr>
<td>CD3$^{\text{low}}$</td>
<td>2688 (63.7)</td>
<td>NA</td>
</tr>
<tr>
<td>CD20$^+$</td>
<td>1401 (33.2)</td>
<td>144-671</td>
</tr>
<tr>
<td>CD16$^+$</td>
<td>174 (4.1)</td>
<td>152-709</td>
</tr>
<tr>
<td>CD56$^+$</td>
<td>0</td>
<td>223-1040</td>
</tr>
</tbody>
</table>

Serum Ig level

- IgG: 2310 (192-515)
- IgA: 213 (12-31)
- IgM: 248 (39-92)
- IgE: 27 (0-150)

Proliferative stimulus

<table>
<thead>
<tr>
<th>Medium</th>
<th>Patient</th>
<th>Controls</th>
</tr>
</thead>
<tbody>
<tr>
<td>PHA</td>
<td>1989</td>
<td>222 330 ± 50 643</td>
</tr>
<tr>
<td>Con A</td>
<td>3695</td>
<td>208 634 ± 52 592</td>
</tr>
<tr>
<td>PWM</td>
<td>7224</td>
<td>135 520 ± 35 077</td>
</tr>
<tr>
<td>Candida</td>
<td>1000</td>
<td>45 431 ± 30 989</td>
</tr>
<tr>
<td>Anti-CD3ε</td>
<td>131</td>
<td>122 953 ± 44 094</td>
</tr>
<tr>
<td>Autologous cells</td>
<td>745</td>
<td>3163 ± 2874</td>
</tr>
<tr>
<td>Allogeneic cells</td>
<td>5520</td>
<td>59 618 ± 38 445</td>
</tr>
</tbody>
</table>

NA indicates not applicable; PHA, phytohemagglutinin; Con A, concanavalin A; PWM, pokeweed mitogen.

Values are expressed as cells/mm$^3$ or (percentage of lymphocytes). Controls values are the 95% confidence intervals for 1550 healthy controls.

Values are expressed as mg/dL (IgG, IgA, IgM) or um/mL (IgE). Normal values are the 95% confidence intervals for 12 6-month-old control subjects.

The patient was receiving IVIG when the IgG level was measured.

Cell sorting

Patient and healthy volunteer CD3ε$^+$ cells used for real-time PCR and immunoblotting were isolated from PBMCs using a fluorescence-activated cell sorting (FACS) Vantage SE (Becton Dickinson, San Jose, CA) by sorting cells stained bright with anti-CD3ε (clone SK7; BD Biosciences) than with isotype control Ab. Patient PBMCs used for sorting were shown to be free of detectable maternal cells by staining with antibody to HLA-A2 (BD Biosciences), an unshared maternal histocompatibility antigen.

Real-time quantitative PCR analysis of CD3ε mRNA expression

RNA isolated from sorted CD3ε$^+$ cells (RNeasy Mini Kit; Qiagen, Santa Clarita, CA) was used for cDNA synthesis (SuperScript II Preamplification System; Invitrogen, Carlsbad, CA) was used for cDNA synthesis (SuperScript II Preamplification System; Invitrogen, Carlsbad, CA). CD3ε and β-actin transcripts were then quantified by real-time PCR with a Roche LightCycler and FastStart DNA Master SYBR Green I Kit (Roche Diagnostics, Indianapolis, IN). The CD3ε primers (5’-CAGCTCTTTCTGAGGAAA and 5’-AGGATCTTCAATC-CAGCAGTA) used for amplification were upstream of both the patient exon 7 mutation site and the region altered in transcript variant 1. CD3ε transcript levels were normalized to those for β-actin in each sample.

CD3ε sequence analysis

gDNA templates isolated from whole blood (RNasey Mini Kit; Qiagen) were PCR-amplified with primer pairs (sequences available on request) spanning each of the 8 CD3ε exons and surrounding intron splice sites. PCR products were purified (Qiagen II Gel Extraction Kit; Qiagen) and used as templates in sequencing reactions (Big Dye Terminator Cycle Sequencing System; Perkin Elmer Life Sciences, Boston, MA). Sequencing reactions representing both strands were analyzed using an ABI 377 Prism DNA (Perkin Elmer) instrument and software. The detected 411insC mutation was further evaluated by sequence analysis of gDNA obtained from the patient’s parents and 50 unrelated healthy individuals of varied ethnic backgrounds (Alzheimer’s Disease Research Center, Duke University Medical Center, Durham, NC). Nucleotide numbers refer to the published cDNA sequence of canonical CD3ε transcript variant 2 (NM000734)$^{16}$ with start codon = 1.

Molecular modeling

Patient mutant CD3ε protein sequence analysis and structure prediction were performed by the Molecular Modeling Facility, Fox Chase Cancer Center (Philadelphia, PA). In these studies, wild-type and D138fsX272 mutant CD3ε sequences were used to perform a multiple-round sequence search using the position-specific iterated basic alignment search tool (PSI-BLAST)$^{15}$ against the National Center for Biotechnology Information (NCBI) nonredundant sequence database. The profiles obtained after each round were used to generate the secondary structure prediction, using...
protein secondary structure prediction program (PSIPRED) for computation and molecular integrated development environment (MolIDE) for graphical rendering. Next, the Protein Data Bank (PDB) was searched for the availability of proteins with known structures that could be used as templates for homology modeling. Transmembrane hidden Markov model (TMHMM) server (Center for Biological Sequence Analysis, Technical University of Denmark, Lyngby, Denmark) was used to predict transmembrane regions in both the wild-type and mutant sequences. A customized database was visualized by fluorography.

**Results**

**Patient immunologic phenotype**

Evaluation of patient peripheral blood lymphocytes by flow cytometry at age 11 months revealed markedly diminished numbers of normal mature T cells expressing high levels of surface CD3ε (clone 145-2C11) and CD3ζ (clone 6B10.2; Santa Cruz Biotechnology) as described. Membranes containing transferred MA5.8 lyses were also probed with anti-Calnexin Ab (David McKean, Mayo Clinic, Rochester, MN). Metabolic labeling

MA5.8 cells were labeled in cysteine- and methionine-free medium containing [35S]methionine for 30 minutes, following which digitonin extracts were either sequentially immunoprecipitated with anti-CD3ε (clone 145-2C11) followed by anti-CD3ζ Ab (clone 6B10.2; Santa Cruz Biotechnology) or immunoprecipitated with anti-CD3ζ Ab alone as described. Isolated TCR subunits were resolved by SDS-PAGE and visualized by fluorography.

**Flow cytometry analysis of retrovirally transduced MA5.8 cells**

Retrovirally transduced cells were isolated by flow cytometry based on eGFP expression as described. Alternatively, mixed populations were stained with anti-TCRβ Ab (clone H57-597) or anti-CD3ε Ab (clone 145-2C11) following which the effect of the transduced construct on TCR expression was evaluated on electronically gated eGFP+ Phoenix-E cells using Flowjo software (Treestar, Ashland, OR).

**Immunoprecipitation and immunoblotting**

Sorted CD3ε+ PBMCs or retrovirally transduced MA5.8 cells were lysed in digitonin (EMD Biosciences, San Diego, CA) lysis buffer (1% digitonin, 50 mM Tris pH 7.4, 150 mM NaCl, 1 mM Na2VO4, 2 mM EDTA), and complete protease inhibitor cocktail (Roche, Basel, Switzerland). Lysates from 8 × 105 CD3ε+ cells were sequentially immunoprecipitated with 2 rounds of anti-CD3ε (clone 145-2C11) followed by anti-CD3ζ Ab (clone 6B10.2; Santa Cruz Biotechnology) as described. Membranes containing transferred MA5.8 lyses were also probed with anti-Calnexin Ab (David McKean, Mayo Clinic, Rochester, MN).

**Plasmids, mutagenesis, and transfection**

Full-length cDNA encoding wild-type CD3ζ transcript variant 2 (NM198053), designated hCD3ζWT, was PCR-amplified from a human cDNA library using CD3ζ primers (5’-GGGATTCCTTCCAGCTGCTTCTGAG and 5’-TCCGAGGCTGACTGAGTGGCGCTTTTCT) and directionally cloned into pBluescript (Stratagene, La Jolla, CA). The patient cytosine insertion mutation (hCD3ζCm) was created by site-directed mutagenesis. Variant constructs hCD3ζWT and hCD3ζCm containing the longer CD3ζ splice variant 1 (NM198053), which includes an additional codon encoding amino acid Q101 due to use of an alternate 5’ splice site in intron 4, were similarly generated using 2 hCD3ζ internal primers (CD3ζ424R: 5’-CCGGTTTTCGCCCCCATCTCA, and CD3ζ425F: 5’-CA-GAGAGAGAACACCCTACAGA) Cloning details will be provided on request. After sequence verification, wild-type and mutant constructs were subcloned into a modified version of p-MSCV-IRES-eGFP (pMiG), with an expanded multiple cloning site (Dario Vignali, St Jude Children’s Research Hospital, Memphis, TN). Phoenix-E retroviral packaging cells were transfected with retroviral vectors using the calcium phosphate transfection method as described. Transfection efficiencies were evaluated by determining the percentage of eGFP+ Phoenix-E cells using flow cytometry (FACSVantage SE; Becton Dickinson). The CD3ζ-deficient murine T hybridoma, MA5.8, was then transduced with retroviral supernatant treated with 8 µg/ml Polybrene.
expressed very low levels of surface CD3ε (CD3εlow) compared with that of normal control cells (Figure 1; Table 1). Similar results were obtained following staining with either of 2 different anti-CD3ε Abs (clone SK7, clone UCHT1; data not shown). In addition, very few patient CD3εlow cells coexpressed detectable surface TCRαβ or TCRγδ (Figure 1) by flow cytometry. CD4+CD3εlow and CD8+CD3εlow patient cells were both present, with the CD8+ population being more prevalent (Figure 1). Further analysis of CD4 and CD8 expression on software-gated peripheral blood CD3εlow cells (Figure 1 bottom panels) revealed a markedly increased frequency of CD4+CD8low CD3εlow cells in the patient (40.7%). Spectratype analysis of patient PBMCs revealed an oligoclonal VB repertoire similar to those previously noted in other SCID patients prior to immune reconstitution (Figure 2B).53 Although not normal, these results nonetheless demonstrate that TCR VB genes had undergone productive rearrangements in patient T cells. TRECs were not detected in patient PBMCs (data not shown). Additional studies revealed that patient T-cell proliferative responses to mitogens, Candida antigen, anti-CD3ε Ab, and allogeneic cells were all profoundly depressed at presentation (Table 1), commensurate with a diagnosis of SCID. Serum IgA and IgM levels were elevated and IgE was present in normal amounts at this time. The patient had received intravenous immunoglobulin (IVIG) shortly before measurement of her serum IgG level. Despite the absence of surface CD3ε expression, patient NK effector cells exhibited detectable, albeit low, lytic activity against labeled K562 (IVIG) shortly before measurement of her serum IgG level. Despite the lack of detectable CD3ζ protein expression in patient CD3εlow cells, we performed CD3ζ sequence analysis and found the patient to be homozygous for a single C insertion in exon 7 following nucleotide 411 of the coding sequence (411insC). Sequencing of patient CD3ζ genes revealed them to be wild type. The patient’s mother and father were both heterozygous for the 411insC CD3ζ mutation, and this sequence abnormality was not detected in gDNA from any of the 50 unrelated healthy individuals analyzed. The C insertion causes a frameshift commencing at amino acid 138 that disrupts expression of the third CD3ζ ITAM (beginning at G139)52 and leads to termination at missense amino acid position 272 (D138fsX272), well beyond wild-type termination codon 164. Hence, unphosphorylated, monomeric, mutant D138fsX272 CD3ζ has a predicted molecular mass (29 kDa) greater than wild-type (16 kDa) CD3ζ (Figure 5). No sequence motifs or regions of homology with known proteins were detected in the mutant CD3ζ sequence and no structure could be confidently assigned to this region by molecular modeling analysis. There were 2 relatively hydrophobic stretches centered around missense amino acids 200 and 250 identified that may represent helices; however, the prediction probability for this was not very high (< 0.5). It is important to note that the epitope recognized by the anti-CD3ζ Ab

Patient CD3ζ expression levels and mutation analysis

Because of the abnormalities in patient cell surface CD3ε expression noted, we next examined expression of CD3ε along with that of another CD3 chain, CD3ζ, in immunoblots of whole-cell lysates prepared from sorted CD3ε+ cells. CD3ε protein levels were similar in patient CD3εlow (Figure 4A lane 7) and healthy control CD3εhigh (Figure 4A lane 3) cells in this experiment. CD3ζ was also abundant in healthy control T cells but was not detected in the equivalent patient sample (Figure 4A lanes 3 and 7). Likewise, we were also unable to detect CD3ζ in permeabilized patient PBMCs by flow cytometry (Figure 4B). Despite the absence of CD3ζ protein, real-time quantitative PCR revealed abundant levels of CD3ζ mRNA in sorted patient CD3εlow PBMCs (Figure 4C). Given the lack of detectable CD3ζ protein expression in patient CD3εlow cells, we performed CD3ζ sequence analysis and found the patient to be homozygous for a single C insertion in exon 7 following nucleotide 411 of the coding sequence (411insC). Sequencing of patient CD3ζ genes revealed them to be wild type. The patient’s mother and father were both heterozygous for the 411insC CD3ζ mutation, and this sequence abnormality was not detected in gDNA from any of the 50 unrelated healthy individuals analyzed. The C insertion causes a frameshift commencing at amino acid 138 that disrupts expression of the third CD3ζ ITAM (beginning at G139)52 and leads to termination at missense amino acid position 272 (D138fsX272), well beyond wild-type termination codon 164. Hence, unphosphorylated, monomeric, mutant D138fsX272 CD3ζ has a predicted molecular mass (29 kDa) greater than wild-type (16 kDa) CD3ζ (Figure 5). No sequence motifs or regions of homology with known proteins were detected in the mutant CD3ζ sequence and no structure could be confidently assigned to this region by molecular modeling analysis. There were 2 relatively hydrophobic stretches centered around missense amino acids 200 and 250 identified that may represent helices; however, the prediction probability for this was not very high (< 0.5). It is important to note that the epitope recognized by the anti-CD3ζ Ab

Figure 3. Patient NK cell function. Values represent the percent specific lysis of 51Cr-labeled K562 target cells by patient or healthy control PBMCs, as indicated, following incubation for 4 hours at effector-to-target (E/T) ratios of 3:1, 6:1, 12:1, 25:1, 50:1 and 100:1.
Lysates were immunoblotted with anti-CD3. In these experiments, lysates prepared from sorted CD3ε+ patient T cells homozygous for the 411insC CD3ε/H9256 mutation were used in our studies lies upstream (amino acids 36-54) of the mutation (Figure 5). Nevertheless, no 29-kDa species was detected used in patient CD3εlow cells (Figure 4A lane 7).

**Patient mutant CD3ε protein fails to support cell surface TCR expression due to rapid degradation**

We next examined steady-state assembly of TCR complexes in patient T cells homozygous for the 411insC CD3ε mutation. Association of CD3ε homodimers with incomplete TCRβ-CD3γ-CD3δ-CD3ε complexes is the final step in TCR assembly and is necessary for survival and efficient transport of complete TCR complexes from the endoplasmic reticulum to the cell surface. In these experiments, lysates prepared from sorted CD3εlow patient and CD3εhigh normal control PBMCs were sequentially immunoprecipitated with 2 rounds of anti-CD3ε Ab to isolate CD3ε-containing complete TCR complexes, followed by anti-CD3ε immunoprecipitation to capture any remaining, unassembled, CD3ε. Immunoblotting of the immunoprecipitated samples from normal control T cells revealed that substantial quantities of CD3ε associated with CD3ε in complete TCR complexes (Figure 4A lane 4), with a smaller amount of free CD3ε (Figure 4A lane 6) remaining in the anti-CD3ε immunoprecipitate. This free CD3ε did not result from failure to capture all of the complete TCR complexes because blotting with anti-CD3ε Ab revealed that all of the CD3ε was captured in the initial immunoprecipitate (Figure 4A; compare lanes 4 and 5). In marked contrast, no CD3ε was detected in the patient samples, either at the 16-kDa size of wild-type CD3ε or at the 29-kDa size predicted for the D138fsX272 mutant (Figure 4A lanes 7-10), despite the presence of immunoprecipitated CD3ε (Figure 4A lane 8).

The absence of detectable mutant CD3ε protein in patient CD3εlow cells in this experiment may have been due to (1) poor expression or rapid degradation of D138fsX272 mutant CD3ε protein prior to assembly with other TCR chains, (2) impaired assembly of D138fsX272 mutant chains with incomplete TCRβ-CD3γ-CD3δ-CD3ε complexes leading to a paucity of complete TCR complexes, or (3) inefficient transport of D138fsX272 mutant CD3ε-containing complete TCR complexes to the plasma membrane. To distinguish among these possibilities, we analyzed TCR assembly and surface expression in MA5.8 cells transduced with patient 411insC mutant or wild-type human CD3ε cDNA retroviral constructs. MA5.8 is a CD3ε-deficient murine T-cell hybridoma in which expression of human CD3ε efficiently restores surface TCR expression. For these studies, retroviral expression constructs containing the patient C insertion mutation in each of 2 prevalent human CD3ε transcript variants were generated by site-directed mutagenesis using PCR. The longer CD3ε transcript variant 1 (NM198053) uses an alternate 5' splice site in intron 4 and encodes a longer CD3ε protein isoform including amino acid Q101 that is not present in canonical transcript variant 2 (NM000734). Assessment of transduced MA5.8 cells by flow cytometry revealed that expression of wild-type CD3ε, but not patient mutant CD3ε, cDNAs effectively restored cell surface TCR expression, as measured by TCRβ and CD3ε staining (Figure 6A). Immunoblot analysis of transduced MA5.8 cell lysates also demonstrated expression of transduced 16-kDa CD3ε protein (Figure 6B upper panel, lanes 2 and 4) and stabilization of endogenous CD3ε expression (Figure 6B middle panel, lanes 2 and 4) by wild-type CD3ε constructs. However, as was the case in patient CD3εlow cells (Figure 4A), we were unable to detect expression of 29-kDa mutant CD3ε protein in transduced MA5.8 cells (Figure 6B upper panel, lanes 3 and 5). Similarly, endogenous CD3ε protein was not stabilized in cells transduced with mutant CD3ε constructs (Figure 6B, compare lanes 3 and 5 to lane 1).

To directly determine if nascent, D138fsX272 mutant CD3ε subunits assemble properly with other TCR chains, MA5.8 cells

**Figure 4. Analysis of CD3ε protein and mRNA expression in patient T cells. (A)** Patient and healthy volunteer sorted CD3ε+ PBMCs were lysed in digitonin lysis buffer and either sequentially immunoprecipitated with 2 rounds of anti-CD3ε Ab followed by anti-CD3ε Ab, or subjected to direct immunoblotting (lanes 3 and 7). Lysates were immunoblotted with anti-CD3ε and anti-CD3δ Abs. Lanes 1 and 2 contain anti-CD3ε- or anti-CD3δ–coated beads incubated with lysis buffer alone, as indicated. (B) Patient and healthy volunteer PBMCs were first stained with PerCP-labeled anti-CD3ε to assess cell surface CD3ε expression. The cells were then fixed and permeabilized, stained with FITC-labeled anti-CD3ε Ab, and analyzed by flow cytometry. Shown are 2-color plots of surface CD3ε and intracellular CD3ε expression. (C) Patient CD3ε+ cells were obtained from 2 different sorted PBMC samples (from different dates). Healthy volunteer sorted CD3ε+ PBMCs were isolated from 2 unrelated individuals (control 1, control 2). cDNA was prepared from sorted cells and CD3ε transcripts were quantified by SYBR green real-time PCR. Levels of CD3ε mRNA were normalized to those of β-actin for each sample. Values represent mean ± SEM of duplicate determinations.

**Figure 5. Diagram of patient CD3ε mutation.** The extracellular (EC), transmembrane (TM), and 3 intracellular ITAMs of wild-type CD3ε protein are depicted along with the location of the homozygous patient mutation and the recognition site of the anti-CD3ε Ab used in the present study. Wild-type (WT) and patient (MUT) CD3ε amino acid sequences in the region affected by the mutation are shown in the expanded view. X indicates stop codon; and fs, frameshift.

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transduced with wild-type or 411insC mutant CD3ε transcript variant 2 constructs were metabolically labeled for 30 minutes and analyzed by sequential immunoprecipitation with anti-CD3ε Ab to detect CD3ε in complete TCR complexes, followed by anti-CD3ζ Ab to capture unassembled CD3ζ, or immunoprecipitation with anti-CD3ζ Ab alone. Only incomplete TCRαβ-CD3ζ-CD8β-CD3ζ complexes lacking detectable CD3ζ were immunoprecipitated with anti-CD3ζ Ab in MA5.8 cells transduced with vector alone (Figure 7 lane 7) as expected, and CD3ζ was not identified in these cells (Figure 7 lanes 7-9). Consistent with the rescue of cell surface TCR expression noted in this study (Figure 6A) and in a previous report,83 newly synthesized CD3ζ assembled in complete TCR complexes was readily apparent in MA5.8 cells transduced with wild-type CD3ζ (Figure 7 lane 1). However, in cells transduced with the 411insC mutant CD3ζ, ectopically expressed CD3ζ was not detected in any form (Figure 7 lanes 4-6), thereby preventing the generation of complete TCR complexes (Figure 7 lane 4). Taken together, these data demonstrate that D138fsX272 mutant CD3ζ protein fails to assemble into complete TCR complexes and is likely to be rapidly degraded.

Discussion

This report describes an infant with T-BNK+ SCID due to a homozygous 411insC mutation in the gene encoding the CD3ζ subunit of the TCR complex. The defects in T-cell development and function noted in this patient are comparable to those previously described in mice lacking CD3ζ expression due to targeted gene disruption.52-55 CD3ζ-deficient mice expressed normal numbers of immature CD4-CD8- thymocytes but markedly diminished development of CD4+CD8+ thymocytes, and very low numbers of CD4+ and CD8+ single-positive TCRαβ thymocytes and peripheral T cells that expressed low levels of surface CD3ε and were nonfunctional.52-55 Productive TCRβ gene rearrangements, which commence at the CD4-CD8-CD44lowCD25+ stage of murine thymocyte development,86 were detected in thymocytes53 and peripheral T cells55 from CD3ζ-deficient mice. These data demonstrated that lack of CD3ζ expression resulted in a severe, but incomplete, block in thymocyte development at the CD4+CD8+ stage in these animals. Thymocytes were not available for analysis from the SCID infant described in this report. However, TCRβ gene rearrangements were detected in the periphery (Figure 2) and the phenotype of circulating patient CD3εlow cells (Table 1; Figures 1 and 4) was nearly identical to that noted in CD3ζ−/− mice.52-55 These findings suggest the defects in T-cell development caused by complete CD3ζ deficiency in mice and humans are similar. CD3ζ plays an important role in TCR signaling via its 3 ITAMs and is also required for assembly and efficient surface expression of the multimeric TCR complex. The observation that T-cell development was restored in CD3ζ-deficient animals by transgenic CD3ζ chains lacking signal transducing ITAMs87 suggested the ability of CD3ζ to promote TCR surface expression was critical for murine thymocyte development but that CD3ζ ITAM signaling was not required for this process. A subsequent analysis of signaling-defective CD3ζ transgenic CD3ζ−/− mice expressing the H-Y TCR transgene did suggest a role for CD3ζ signaling in amplifying TCR signals during thymocyte positive and negative selection.88 The deranged TCR assembly and surface expression noted in patient T cells and MA5.8 cells transduced with patient mutant CD3ζ in the present report suggests the role CD3ζ plays in TCR assembly and expression is also essential for normal human T-cell development.

The homozygous 411insC CD3ζ gene mutation noted in our SCID infant led to complete CD3ζ deficiency with no protein detected in patient CD3εlow cells or retrovirally transduced MA5.8 cells. D138fsX272 chains encoded by 411insC mutant patient CD3ζ alleles are predicted to lack the final 26 residues of wild-type protein that include the third intracellular ITAM, and instead contain 134 C-terminal missense amino acids that exhibit no clear-cut structure or homology with known proteins and lack identifiable sequence motifs. Neither homology searches nor molecular modeling analysis have provided clear insight into how the C-terminal missense extension might prevent stable CD3ζ expression and incorporation into patient TCR complexes. Our

Figure 6. Patient mutant CD3ζ fails to rescue TCR expression in retrovirally-transduced CD3ζ-deficient MA5.8 cells. (A) MA5.8 cells were transduced with individual retroviral vectors containing one of the following human CD3ζ cDNAs: hsCD3ζ WT (wild-type CD3ζ transcript variant 2), hsCD3ζ Cm (patient 411insC mutant CD3ζ transcript variant 2), hlCD3ζ WT (wild-type CD3ζ transcript variant 1), or hlCD3ζ Cm (patient mutant CD3ζ transcript variant 1). Cells were also transduced with pMiG vector alone. Depicted are 1-color histograms of transduced MA5.8 cells analyzed for expression of eGFP indicator protein, surface TCRζ, or surface CD3ζ by flow cytometry. (B) MA5.8 cells retrovirally transduced with the indicated cDNAs were lysed in digitonin lysis buffer, resolved by SDS-PAGE, and immunoblotted with anti-Calnexin Ab (bottom panel) to assess gel loading.

Figure 7. MA5.8 cells transduced with patient mutant CD3ζ cDNA do not express detectable CD3ζ protein and fail to assemble complete TCR complexes. MA5.8 hybridoma cells retrovirally transduced with the indicated human CD3ζ cDNAs or pMiG vector alone were metabolically labeled for 30 minutes and lysed in digitonin lysis buffer. Lysates were either sequentially immunoprecipitated with anti-CD3ζ Ab followed by anti-CD3ζ Ab or immunoprecipitated with anti-CD3ζ Ab alone. Immunoprecipitated material was resolved by SDS-PAGE and TCR chains visualized by fluorography.
inability to identify nascent D138fsX272 mutant protein in metabolically labeled, transduced MA5.8 cells indicated the elongated mutant CD3ζ chains were unstable and rapidly degraded. This finding was unlikely to be simply due to poor expression of mutant CD3ζ cDNA by the pMIG vector in these experiments because (1) endogenous mutant CD3ζ protein was also undetectable in patient CD3ζlow cells, despite normal steady-state levels of CD3ζ mRNA (Figure 4), (2) wild-type and patient mutant CD3ζ cDNAs only differ by one coding region nucleotide, and (3) the observed comparable expression of GFP protein in MA5.8 cells transduced with wild-type or mutant CD3ζ constructs (Figure 6A) demonstrated that the common bicistronic transcriptional unit of the vector encoding both GFP and CD3ζ insert was efficiently expressed in these cells.

The findings in the present study examining complete CD3ζ deficiency also extend those of a recent report describing a patient with recurrent infections and decreased numbers of peripheral T cells due to a complex, partial deficiency of CD3ζ protein expression.60 In that report, 90% of the patient’s T cells exhibited low surface TCRβ and CD3ζ expression, and very low levels of CD3ζ protein, which was only detectable in the cytoplasm. The remaining 10% of patient-origin T cells displayed normal surface TCRβ, CD3ε, and CD3ζ expression but lacked detectable anti-CD3ζ-mediated ZAP-70 phosphorylation.60 Patient T cells with low TCR expression were homozygous for a germline Q70X CD3ζ mutation, whereas cells with normal TCR expression exhibited partial correction of the germline Q70X present on one allele by one of 3 different missense somatic mutations of Q70X found on the other allele.60 An increased frequency of CD4+CD8−CD3ε+ cells was not noted in the patient with CD3ζ deficiency partially corrected by somatic mutations, presumably because partial correction of the CD3ζ mutation returned surface TCR expression to a more normal level. Differences in NK cell development and function were also apparent between the described patient with partial CD3ζ deficiency60 and the SCID infant due to complete CD3ζ deficiency reported herein. This finding is of interest because CD3ζ subunits have been shown to associate with CD16,89,90 NKp46,91 and NKp3092 NK receptors and transduce activation signals. Normal numbers of CD56−CD16+ NK cells that exhibited normal cytoxicity of K562 targets were noted in the reported human case of partially corrected CD3ζ deficiency.60 CD3ζ−/− mice also exhibited normal NK activity against YAC-1 target cells, demonstrating CD3ζ−/− mice was not essential for murine NK cell development and function.55 However, in the present report, normal numbers of CD16+ cells were present but the NK cells lacked detectable coexpression of CD56 on the cell surface (Table 1) and exhibited diminished lytic activity against K562 targets at all E/T ratios tested (Figure 3). The CD56−CD16+ cells present in our patient possibly represent the previously described dysfunctional and rare CD56−CD16+ subset of human NK cells that is greatly expanded in HIV-viremic patients.93,95 CD56−CD16+ NK cells exhibit altered expression of certain inhibitory and activating NK receptors and poor lytic activity against K562 targets.95 Although we have not noted CD56−CD16+ cells in other patients with CMV infection and are not aware of any reports describing a role for this NK subset in the response to CMV, it is conceivable this virus may have been responsible for expansion of CD56−CD16+ NK cells in our patient. Alternatively, CD56−CD16+ cells may be the only NK subset that develops in the setting of complete human CD3ζ deficiency. Studies to further characterize the NK cells in our patient are planned. CD56−CD16+ cells were not noted in the child with partially corrected CD3ζ deficiency reported previously.60 However, that patient exhibited 2 distinct populations of T cells, in terms of CD3ζ expression, and may have likewise expressed more than one population of NK cells.

In conclusion, the present study describes a case of T−B−NK+ SCID due to a homozygous 411insC mutation in the CD3ζ gene that, to our knowledge, is the first reported case of SCID attributable to complete CD3ζ deficiency. In contrast with the findings in a recent report of the first case of partial human CD3ζ deficiency,60 the SCID patient in the present study lacked mature T cells, expressed large numbers of CD3ζlow CD4−CD8− cells in the periphery, and exhibited an unusual population of CD56−CD16+ NK cells. These observations in 2 CD3ζ-deficient patients provide a second example, along with that of CD3ε deficiency,22,57 that differences in the extent of a CD3 chain deficiency can lead to different clinical and immunologic phenotypes.

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Authorship

J.L.R. designed research, collected and analyzed data, and wrote the paper; J.P.H.L. designed and performed research; M.C., R.E.P., E.O.S., C.M.W., and M.D.K. performed research; J.H.C. contributed vital new reagents and analyzed data; M.S.K. contributed analytical tools and analyzed data; M.S. contributed analytical tools and analyzed data; X.-P.Z. contributed analytical tools and analyzed data; J.H.C. performed research and analyzed data; D.L.W. contributed vital new reagents and analyzed data; and R.H.B. designed research and analyzed data.

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T−B+NK+ severe combined immunodeficiency caused by complete deficiency of the CD3 ζ subunit of the T-cell antigen receptor complex

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