Ig class switch recombination (CSR) deficiencies are rare primary immunodeficiencies, characterized by a lack of switched isotype (IgG, IgA, or IgE) production, variably associated with abnormal somatic hypermutation (SHM). Deficiencies in CD40 ligand, CD40, activation–induced cytidine deaminase, and uracil–N-glycosylase may account for this syndrome. We previously described another Ig CSR deficiency condition, characterized by a defect in CSR downstream of the generation of double-stranded DNA breaks in switch (S) regions. Further analysis performed with the cells of five affected patients showed that the Ig CSR deficiency was associated with an abnormal formation of the S junctions characterized by microhomology and with increased cell radiosensitivity. In addition, SHM was skewed toward transitions at G/C residues. Overall, these findings suggest that a unique Ig CSR deficiency phenotype could be related to an as-yet-uncharacterized defect in a DNA repair pathway involved in both CSR and SHM events.

Ig class switch recombination (CSR) deficiencies are rare primary immunodeficiencies, usually called hyper-IgM syndromes, whose frequency is 1 in 100,000 births. They are characterized by a defective Ig CSR, as shown by serum IgM levels that are normal or increased, contrasting with a marked decrease, or absence, of IgG, IgA, and IgE (1). As a consequence of the molecular defect, the defective CSR may be associated with defective generation of somatic hypermutations (SHMs) in the Ig variable (V) region. The definition of several Ig CSR deficiencies made possible a better description of the mechanisms underlying CSR and SHM, both required for the maturation of antibody responses (2).

The maturation of the antibody repertoire produces several antibody isotypes with high affinity for antigen, a necessary feature for an efficient humoral response. Antibody maturation occurs mostly in the germinal centers of the secondary lymphoid organs after antigen and T cell–driven activation: CSR results in the production of antibodies of different isotype (IgG, IgA, or IgE) with the same V(D)J specificity and, therefore, the same antigen affinity (3, 4). SHM commonly introduces stochastic mutations (1/10^3 bp/cell cycle), mainly in the V region of the Ig, a genetic modification that is followed by the positive selection of B cells harboring a B cell receptor (BCR) with high antigen affinity (5, 6). CSR and SHM occur together in germinal centers under BCR/CD40 activation, but neither is a prerequisite for the other because IgM may be mutated, whereas IgG or IgA may not (7–9).

Mutations in the gene encoding the CD40 ligand molecule (CD40L and CD154; references
A CSR DEFICIENCY ASSOCIATED WITH IMPAIRED DNA REPAIR | Péron et al.

RESULTS

Skewed pattern of SHM in patients’ CD19+/CD27+ B cells

All five patients presented with a defective in vivo and in vitro CSR defect (26). The pattern of SHM was assessed in the IgVH 3–23 region of IgM on purified CD19+CD27+ B lymphocytes (Fig. 1). SHM frequency was either slightly decreased (patients P1 and P5) or normal (P2, P3, and P4), and the ratio of mutated clones to total analyzed clones was generally lower than in controls. G/C nucleotides were preferentially targeted in the two patients with a low SHM frequency, with, respectively, 80 and 74% of mutations on G/C residues (control, 65 ± 4%; range, 55–71%), whereas G/C targeting was normal in P2, P3, and P4. However, in all five patients, although there were variations from one patient to another, including in siblings P2 and P3, SHM showed a skewed pattern of nucleotide substitution on G/C residues; 63–92% mutations were transitions (G > A, C > T), compared with 56 ± 5% (range, 49–62%) in the controls (P < 0.005; nonparametric Mann–Whitney U test). Except for P1, transitions at A/T residues were within normal ranges (Fig. 1).

The CSR defect associated with a biased SHM pattern found in these patients was reminiscent of that seen in UNG deficiency (24). The in vitro uracil-DNA glycosylase activity of extracts from patients’ cell lines was therefore studied on a double-stranded probe containing a U:G mismatch. Cell-free extracts from all patients were similarly able to undergo base excision, leading to the cleavage of the probe (Fig. 2). As expected, an UNG-deficient B cell line had no detectable activity, and no cleavage was observed using control or patients’ cell extracts on a double-stranded DNA probe that did not contain a U/G mismatch. These results, associated with the observation of normal UNG sequences and CSR–induced DSB in Sp regions (26), excluded an UNG deficiency as the basis of the CSR deficiency found in these patients.

Another pathway used for bypassing the UNG-induced abasic site consists of translesion synthesis, involving the Rev1 polymerases (27). Its role in antibody maturation has been recently reported, based on the observation of normal frequency but a skewed pattern of SHM in Rev1-deficient mice (28).

Figure 1. SHMs in V regions of patients’ CD27+ B cells. SHMs of VH3–23 IgM region were assessed in CD19+CD27+ sorted B cells by RT-PCR using Pfu Taq. Cloned products were sequenced. Nucleotide substitutions are shown as absolute numbers, and SHM frequency is shown as a percentage of mutations occurring among all analyzed nucleotides (at least 2,400). Numbers of mutated clones among all different studied clones are also noted.

Figure 2. Normal Uracil incision activity in patients’ cell lines. Protein extracts from control fibroblasts (control 1); control EBV B cell line (control 2); P2, P3, and P4 fibroblasts; and P1 and P5 EBV B cell lines were mixed with double-stranded fluorescein-labeled oligonucleotide substrate with or without a single dU/dG mismatch. Cleavage of the probe containing a dU/dG mismatch revealed an efficient base excision activity in patients and controls. No detectable base excision activity was observed with UNG−/− cells or in the presence of protein extracts (−).
Rev1-deficient lines have an increased sensitivity to γ-irradiation and other DNA damaging agents, including methyl methane sulfite (29). Although unlikely, as Rev1−/− mice are not as affected by a CSR defect, REV1, REV3, and REV7 genes were sequenced and found to be normal (unpublished data). The skewed SHM pattern is also reminiscent of mismatch

Figure 3. Abnormal pattern of Sμ-Sα junctions in patients’ B cells. (A) Sequences of Sμ-Sα junctions. Two sequences from each patient are shown. The Sμ and Sα1 or Sα2 sequences are aligned above and below the recombination switch junctional sequences. Microhomology was determined by identifying the longest region at the switch junction of perfect uninterrupted donor/acceptor identity (boxed with solid lines). Imperfect repeat was determined by identifying the longest overlap region at the switch junction by allowing one mismatch on either side of the breakpoint (the extra nucleotide identified beyond the perfect-matched sequence identity is boxed by dotted lines). The Sμ and Sα breakpoints for each switch fragment are indicated by ▼ and ▲, respectively, and their positions in the germ line sequences are indicated on top of or below the arrowheads. The number of base pairs involved in microhomology and imperfect repeat for each junction is shown at the bottom right of each switch junction. (B) Pie charts demonstrate the perfectly matched short homology usage at Sμ-Sα junctions in controls and patients. The proportion of switch junctions with a given size of perfectly matched short homology is indicated by the size of the slices. (C) Comparison of Sμ-Sα junctions in controls, patients described herein, and A-T and DNA ligase IV-deficient patients. *, P < 0.05; **, P < 0.01; ***, P < 0.001 (χ² test).
repair (MMR) defect (30). Such a defect, although unlikely because of the absence of cancer during early life (31, 32), was excluded because MSH2, MSH6, MSH5, EXO1, MLH1, and PMS2 RNA transcripts were normally expressed and gene sequence was normal (unpublished data).

Abnormal switch junctions in patients’ B cells
Switch junctions are generated after the processing of DNA ends produced in S regions and several DNA repair defects lead to abnormal structure of these junctions. We therefore characterized switch junctions from patients to detect potential abnormalities. We cloned and sequenced 44 switch fragments (43 Sµ–Sα and 1 Sµ–Sγ–Sα) from B cells of patients P1, P3, P4, and P5. All the switch fragment sequences were unique and therefore represent independent CSR events. Two sequences from each patient are shown in Fig. 3 A. The Sµ–Sα junctions from controls (n = 154), used for comparison, have been previously published (33, 34). There was a significant increase in the extent of donor-acceptor homology at the Sµ–Sα junctions from patients B cells (the mean length of overlap was 7.2 ± 4.7 bp in patients vs. 1.8 ± 3.2 bp in controls; Student’s t test, P = 1.2 × 10−9). The majority of junctions (39 out of 43; 91%) from patients displayed a perfectly matched homology (microhomology) of ≥1 bp (i.e., at least one nucleotide is shared by both the Sµ and Sα regions), whereas the remaining four junctions showed a 1-bp insertion and no junction showed precisely joined blunt ends (Fig. 3, B and C). Moreover, 60% of the junctions exhibited a long microhomology of ≥7 bp. When one mismatch was allowed at either side of the switch junction, most of the switch junctions (38 out of 43; 88%) from the patients were flanked by ≥7–8 bp of imperfect repeats (unpublished data). The dramatic shift in using long microhomologies or imperfect repeats in the Sµ–Sα junctions from these patients have previously only been observed in patients with ataxia telangiectasia (A-T) or DNA ligase IV deficiency (Lig4D; references 33, 35). Interestingly, in our patients, the shift was causing by homologies encompassing 7–9 bp or longer, whereas in A-T and Lig4D patients, it was mainly due to an increased usage of microhomologies of ≥10 bp (Fig. 3 C). Of note, a significantly reduced rate of insertions, but not mutations, was observed at or close to the switch junctions of B cells from patients as compared with controls (Fig. 3 C).

Increased radiosensitivity of patients’ cell lines
To search for a possible DNA repair defect, the radiosensitivity of patients’ fibroblasts was first tested. Fibroblast lines from three patients (P2, P3, and P4) were submitted to increasing doses of γ-irradiation, and their survival was assessed by clonogenic assay. A reproducibly increased radiosensitivity was observed in these three cell lines, including those from the two siblings (P2 and P3). Although the increased radiosensitivity

![Image](78x276 to 282x424)

**Figure 4.** Increased radiosensitivity of patients’ cell lines. (A) Fibroblasts from patients [P2 [X], P3 [ ], and P4 [ ])] were irradiated at 0.5–3 Gy. Survival was assessed after 14 d of culture as the number of colony-forming cells compared with nonirradiated cells. Fibroblasts from two age-matched controls and fibroblasts from two Ig CSR-deficient patients with normal SHM (diagonal lines), one ARTEMIS−/−, and one A-T cell line (gray) were positive and negative controls. Results are expressed in log scale. *, P < 0.05; **, P < 0.005; ***, P < 0.001 (unpaired two-tailed Student’s t test). (B) EBV B cell lines from patients [P1 [ ], P2 [X], P4 [ ], and P5 [ ])] were irradiated at 0.5–1.5 Gy. After 10 d of culture, survival was assessed as the number of positive wells [defined as viable cell colonies containing >32 cells] for each plate containing irradiated cells compared with number of positive wells for each plate containing unirradiated cells. EBV B cell lines from four controls, three AID-deficient patients, two UNG-deficient patients, and six patients with Ig CSR deficiency with normal SHM (diagonal lines), and two A-T-EBV B cell lines (gray) were used as positive and negative controls. *, P < 0.05; **, P < 0.005 (unpaired two-tailed Student’s t test). Radiosensitivity of cell lines was assessed two to four times each. Results are expressed as mean ± SD.

![Image](318x160 to 558x347)

**Figure 5.** Normal irradiation-induced cell cycle progression arrest in patients’ fibroblasts. (A) The G1/S cell cycle checkpoint was assessed by BrdU incorporation and DNA content quantification of fibroblasts from a control patient, an A-T patient, and P3 after 5 Gy of irradiation or no irradiation. (B) The G2/M cell cycle checkpoint was assessed by FACS analysis of phosphorylation of histone H3 and DNA content in either untreated or 5 Gy–irradiated fibroblasts from a control patient, an A-T patient, and P3. The same results were obtained in P2 and P4 fibroblasts. Percentages of G1/S and G2/M cells are indicated.
was less marked than that of fibroblasts from patients suffering a DNA repair defect, such as A-T or Artemis deficiency used as controls, it was significantly different from that of healthy fibroblasts (Fig. 4 A: 0.5 and 3 Gy, P < 0.05; 2 Gy, P < 0.005; 1 Gy, P < 0.001 [unpaired two-tailed Student’s t test]). These results were confirmed by using Epstein-Barr virus (EBV) B cell lines from P1, P2, P4, and P5 that were more radiosensitive than the control cell lines (Fig. 4 B; 0.5 and 1 Gy, P < 0.05; 2 Gy, P < 0.005 [unpaired two-tailed Student’s t test]). In contrast, increased radiosensitivity of AID or UNG-deficient cell lines or cell lines from patients with other forms of Ig CSR deficiency was not observed (Fig. 4). These results suggest a defect in double-stranded DNA break repair.

Normal irradiation-induced cell cycle progression arrest in patients’ fibroblasts

Another event occurring rapidly after DNA damage sensing in dividing cells is cell cycle progression arrest. We therefore studied the irradiation-induced inhibition of cycle progression in fibroblasts from patients P2, P3, and P4. Arrest of entry into S phase (G1/S checkpoint) was studied 10 h after a 5-Gy irradiation, whereas entry into mitosis (G2/M checkpoint) was assessed after 1 h. Arrested cell cycle progression was observed in the patients’ cells in contrast to A-T fibroblasts, which exhibited, as expected, a drastic defect in both checkpoints (Fig. 5). These results show that the increased sensitivity of cells to γ-irradiation does not result from a defect in the cell cycle checkpoints induced by DNA damage.

Normal irradiation-induced foci formation in patients’ cell lines

Excessive radiosensitivity could be due to a DNA repair defect caused by an impaired recruitment of proteins to double-stranded DNA break sites. We, therefore, studied one of the earliest responses to DNA damage, namely, the induction of histone H2AX phosphorylation (γH2AX; references 36, 37). γH2AX is essential for keeping DNA ends together and for stabilizing the association of DNA repair factors, such as the MRE11–RAD50–NBS1 complex, 53BP1 (tumor protein. p53-binding.protein.1), and mediator of DNA damage checkpoint 1 (MDC1), at the site of the damage (38). DNA repair foci, including γH2AX, MRE11, 53BP1, and MDC1, were equally recruited 2 h after a 2-Gy irradiation in control and patients’ cells, both in fibroblasts (P2, P3, and P4) and in EBV B cell lines (P1, P2, P4, and P5; Fig. 6 and not depicted). FACS analysis did not demonstrate the persistence of γH2AX in nuclei of fibroblasts (P2 and P4) or of EBV B cell lines (P1, P2, P4, and P5) as analyzed at different time points after irradiation (unpublished data). These results indicate that the increased sensitivity to irradiation does not result from a defect in the initial DNA damage sensing, or in a major DNA repair, pathway. Alternatively, the molecular defect may lead to the unrepair of only a fraction of irradiation-induced DNA damage, not detectable in these experiments, as observed in Artemis deficiency (39).

Normal nonhomologous end joining (NHEJ) in patients’ cell lines

The major DSB DNA repair pathway used in mammals is the NHEJ pathway. Some NHEJ factors have been shown to be necessary during CSR (40, 41). To study the ability of patients’ cells to join double-stranded DNA ends by this pathway, we analyzed the in vitro end joining of linearized plasmid DNA by using patients’ fibroblast and/or EBV B cell line extracts by the methods of Baumann and West (42) and Buck et al. (43). The DNA-end ligation assay resulted in the formation of DNA concatemers when extracts from both patient and controls were used (Fig. 7 A). Expectedly,
Cernunnos-deficient fibroblast extracts did not lead to concatemer formation. We next examined the ability of patients’ cells to join incompatible DNA DSB formed in vivo and, thus, requiring DNA break processing before ligation. P2 and P4 fibroblasts were transfected with restriction enzyme–digested, linearized plasmids containing incompatible 3′–3′ overhang ends. Recircularized plasmids were recovered 72 h after transfection, and their junctions were studied by DNA sequencing (43). Most junctions in plasmids recovered from both patient and control fibroblasts showed similarly accurate repair, in contrast to what is observed in Cernunnos-deficient fibroblasts (Fig. 7 B). Because a subtle defect may be undetectable in these assays, we excluded hypomorphic defects in DNA-dependent protein kinase, Ku complex, Cernunnos, Artemis, XRCC4 (X-ray repair complementing defective in Chinese hamster), and ligase IV either by gene sequencing in patients or by study of segregation with polymorphic markers in both informative families (P2, P3, and P5; unpublished data).

DISCUSSION

We have described here a new primary immunodeficiency, characterized by a defect in Ig class switching associated with a reduced memory B cell population and a skewed nucleotide substitution in SHM. The observation of a female born to consanguineous parents and of two affected siblings suggests an autosomal recessive inheritance pattern. Although all patients exhibited the same CSR defect with increased radiosensitivity, there were some differences between patients in regard to SHM pattern and frequency. Because the underlying molecular basis of this condition is still unknown, we do not know so far whether all patients are affected by distinct molecular defects or by distinct mutations of the same gene. Other genetic and/or environmental factors could also be involved, according to P2 and P3 phenotypic differences. However, the overall phenotype was unique in these five patients. Several hypotheses, including one of defective survival of switched B cells, or of a defect in the CSR/SHM machinery involving DNA repair, could account for the condition.
Abnormalities in survival signaling of switched B cells could underlie this condition. Molecular interactions are known to be essential for B cell survival, including that of B cell activating factor (BAFF) with its receptor on B cells, BAFF-R. The observation of fewer switched CD27+ B cells could fit this model. However, the observed in vitro defective CD40-dependent CSR, cannot be accounted for by a BAFF-R or BAFF abnormality. A response to DNA damage leading to inappropriate cell death should also be considered. In cells other than germinal center B cells, DSBs activate p53 and p21, resulting in cell cycle arrest and apoptosis. In contrast, in germinal centers, the p53 response to DNA damage is directly inhibited by the highly expressed transcriptional regulator B cell lymphoma 6 (BCL6), whereas p21-induced cell cycle arrest is suppressed through interaction of its transcriptional activator Miz1 (protein inhibitor of activated STAT2) with BCL6. Both these events enable intense proliferation of B cells undergoing CSR. Fitting in with this observation, BCL6-deficient mice are depleted of germinal centers because of a strong B cell apoptosis (48). Such a defect in transcriptional repression of proteins involved in cell cycle arrest induced by DNA damage could also underlie this Ig CSR deficiency. Another interesting hypothesis is related to the recently described role of phosphoinositide-3 kinase (PI3K) acting as a negative regulator of CSR (49). PI3K-induced CSR inhibition is dependent on the inhibition of AICDA transcription via B lymphocyte-induced maturation protein 1 overexpression and inactivation of the Forkhead Box family (FOXO) of transcription factors by the serine threonine kinase Akt. Activated B cells from the patients in this study expressed the AICDA gene transcripts and AID protein normally. However, PI3K also exerts a downstream function in AID activity regulation, as shown by the observation that overexpressed AID does not fully compensate for the inhibitory effect of PI3K on CSR. Thus, any abnormal signal leading to increased PI3K activity in patients’ B cells could induce a CSR deficiency.

Alternatively, a defect in the complex machinery underlying the SHM and the CSR processes, and especially the DNA repair, appears more likely because of abnormal S junctions and increased radiosensitivity of fibroblasts and EBV B cell lines. AID and UNG-induced DNA lesions are repaired differently in S and V regions. The CSR-induced DSB repair requires phosphorylation of the H2AX histone (γH2AX), as well as the MRE11–RAD50–NBS1 complex and the 53BP1 and MDC1 proteins, as shown by the phenotype of mice or B cells depleted of each of these molecules (50–54). The observed normal accumulation of γH2AX, MRE11–RAD50–NBS1 complex, and 53BP1 and MDC1 proteins on DNA repair foci of patients’ cells excluded a defect in the first step after DNA damage sensing, leading to DNA repair. In addition, the H2AX, 53BP1, and MDC1 genes were sequenced and found to be normal. MRE11 and NBS1 mutations in man lead to well-known syndromes, A-T–like disease (55) and Nijmegen syndrome (56), respectively. ATM (A-T mutated) is involved in DNA repair of S regions (53, 57). Nevertheless, as A-T patients exhibit an Ig CSR deficiency (reference 58; unpublished data), a potential role for ATM could be envisaged. The observation of a normal irradiation-induced cell cycle progression arrest excludes an abnormal ATM-mediated cell cycle checkpoint pathway. Moreover, the switched junctions, although based on microhomologies, were slightly different from those observed in A-T. The NHEJ enzymes have been shown to be required for CSR (35, 40, 41, 59). However, such deficiency of these factors is unlikely because of normal TCR and BCR expression and function in patients’ cells, as well as the absence of γH2AX persistence after irradiation and the normal results obtained for both NHEJ assays. However, we cannot definitively rule out a defect in a NHEJ factor redundant with other factors in mediating the V(D)J recombination process that remains undetectable in the experimental assays used. It should be stressed that the nature of the switch junctions and the bias to transitions at G/C residues do not fit well with defective NHEJ. The SHM abnormality could be the consequence of a defective repair of V region breaks. The nature of the DNA breaks in V regions and their repair have not been completely elucidated (60). The recent observation that the MRE11–RAD50–NBS1 complex localizes on V regions is compatible with DSB generation in V regions during SHM introduction (61). The MMR enzymes play a role in CSR and SHM in mice, as shown by the slightly defective CSR and skewed pattern of SHM observed in MMR-deficient mice (62, 63). In addition, abnormality of S junctions observed in the B cells from our patients is reminiscent of that observed in Mlh1- or Pms2-deficient mice (30), both defects, however, that are excluded by gene sequencing. Thus, an as-yet-uncharacterized defect in a DNA repair pathway can be postulated to account for a unique phenotype characterized by defective CSR and SHM, associated with an abnormality of the switch junction repair and increased cell radiosensitivity. This factor could be required for efficient NHEJ in S regions and DNA repair of V regions. It could also be NHEJ independent. Of note, AID-dependent illegitimate recombination events occurring between the IgH locus and c-myc in B cell lymphomagenesis have been shown to be mediated by an as-yet-unknown NHEJ-independent process (64). It is, thus, attractive to consider that this as-yet-uncharacterized DNA repair pathway might be physiologically involved in the CSR and SHM processes.

**MATERIALS AND METHODS**

**Patients.** We studied five patients (4 males and 1 female), 4–8 yr of age, from four unrelated families. P2 and P3 were siblings, and P5 was born to a consanguineous family. All patients suffered from recurrent bacterial infections. P4 also presented with enlarged lymph nodes, and P4 and P5 had severe autoimmune hemolytic anemia. P3 died from severe hepatitis, and P5 suddenly died at 6 yr of age. A diagnosis of defective Ig CSR was made on the basis of low-serum IgG, IgA, and IgE concentrations and high-serum IgM concentrations (Table I). Cases P1, P2, P3, and P4 were previously reported (26). B cell and T cell counts were similar to those of age-matched controls, and T cell functions were found to be normal (unpublished data). However, the percentage of memory B cells was strongly decreased, and switched B cells were virtually absent (Fig. 1), whereas the in vitro IgE
production of B cells upon soluble CD40L+ IL-4 activation was negative, though B cells proliferated normally in that setting. The in vitro CSR defect was shown to occur downstream of DNA DSBs (26). AID and Ung deficiencies were excluded by normal gene sequence. AID protein was detectable by Western blot in activated B cells whenever tested (EBV B cell lines from P1, P2, P4, and P5 and soluble CD40L+IL-4-activated B cells from P1 and P4 (unpublished data). Immunological and genetic studies were performed after the informed consent of parents. The study was approved by the ethics committee.

Analysis of SHM in variable gene of Ig. SHM in VH3-23 IgM gene was assessed as described previously (18). RT-PCR was performed with 0.5 U Phu polymerase (Stratagene) and the primers V3-23 leader exon and Cα (5′-CAGTGAGTTGAGTGAAGATT-3′). The amplification of Sα fragments was visualized by FluorImager (FLA3000; Fujifilm). Products were cloned and sequenced.

Uracil incision assay. Whole-cell extracts (WCEs) were prepared by cells lysis in buffer 1 (10 mM TRIS, pH 8, 1 mM EDTA, 5 mM dithiothreitol [DTT], and protease inhibitors) followed by the addition of 0.5 volume of buffer 2 (50 mM Tris, pH 8.0, 1 M KCl, 2 mM EDTA, and 2 mM DTT) and submitted to three freeze-thaw cycles. After centrifugation, supernatants were dialyzed against dialysis buffer (20 mM Tris, pH 8.0, 20% glycerol, 0.1 M K [OAc], 0.5 mM EDTA, and 1 mM DTT).

For the uracil incision assay, adapted from Di Noia (65), 10 μg WCEs were mixed in reaction buffer with 1 pmol of double-stranded, FITC-labeled oligonucleotide containing or not containing a single dU/dG mismatch in 20 μl. After 2 h at 37°C, the reaction was stopped with 10 μl of formamide loading dye (GE Healthcare) and products run on a denaturing gel were visualized via FluorImager (FLA3000; Fujifilm).

Study of switch junctions. The amplification of Sα-Sα fragments from in vivo switched B cells was performed as described previously (33, 66). The PCR-amplified switch fragment sequences were gel purified, cloned, and sequenced. The switch breakpoints were determined by aligning the switch fragment sequences with the Sα (X54713), Sα1 (L191219), or Sα2 (AF030305) sequences. Analysis of microhomology usage at the junctions and mutations in the IV–deficient patients was performed as described previously (33, 58). Data from controls, A-, and DNA ligase IV–deficient patients were described previously (33–35).

Ionizing radiation sensitivity assay. Primary fibroblasts were irradiated (137Cs source) with different doses (0, 0.5, 1, 2, or 3 Gy), and serial dilutions were cultured for 14 d in a 10-cm culture dish. The number of colonies for each dose was assessed, and ratios referring to the dilutions of the nonirradiated cells were determined (67).

Irradiated (0, 0.5, 1, or 1.5 Gy) EBV B cell lines were serially diluted and cultured in a 96-well plate for 10 d (68). Each well containing viable cell colonies, dark blue stained after a 4-h incubation in 1 mg/ml 3-(4,5-dimethylthiazol-2-4)-2,5-diphenyloxazole bromide (Sigma-Aldrich), with >32 cells was scored as positive.

Table 1. Phenotype of patients

| Characteristic | P1  | P2a | P3a | P4  | P5 | Age-matched controls |
|----------------|-----|-----|-----|-----|----|---------------------|
| Age at diagnosis (yr) | 6   | 6   | 8   | 4   | 4  | –                   |
| Serum Ig levels (g/l) |     |     |     |     |    |                     |
| IgM            | 16  | 4.1 | 2.6 | 2.8 | 0.9 | 0.5–1.2             |
| IgG            | 0.6 | 3.6 | 5.7 | 0.7 | 1.4 | 6.8–12.5            |
| IgA            | 0.3 | <0.03 | <0.03 | <0.03 | <0.03 | 0.6–1.6           |
| T lymphocyte counts/μl | 2,400 | 1,420 | 1,238 | 1,612 | 1,300 | 1,200–2,600 |
| B lymphocyte counts/μl | 452  | 120 | 170 | 462 | 440 | 110–570             |
| IgG+/CD19+ (%) | 99.9 | 99.2 | ND  | 99.8 | ND  | 83–93               |
| CD27+/CD19+ (%) | 1.5  | 3.3 | 3.1 | 4.7 | 3.1 | 13–58              |

aP2 and P3 are siblings.
bGated on CD19 + B cells.

gDNA repair foci detection. 2 h after 2–5 Gy of γ-irradiation, primary fibroblasts or EBV B cell lines were labeled with the mouse monoclonal anti-γH2AX (Ser139; clone JBW103; Upstate Biotechnology), mouse monoclonal anti-MRE11 (clone 12D7; Abcam), mouse monoclonal anti-53BP1 (provided by I. Ward, Mayo Clinic College of Medicine, Rochester, MN), or rabbit polyclonal anti-MDC1 (provided by S. Grant, Baylor College of Medicine, Houston, TX) antibodies. Cells were incubated with the secondary antibodies goat anti-mouse Alexa Fluor 488, goat anti-mouse Alexa Fluor 546, and goat anti-rabbit Alexa Fluor 488 (Invitrogen). Slides were counterstained with DAPI and analyzed by epifluorescence microscopy (Axioplan; Carl Zeiss MicroImaging, Inc.).

In vitro and in vivo NHEJ assay. Functional activity of the NHEJ system repair was assessed in vitro (43) by incubation of 5 μg WCEs with 25 ng EcoRI-digested pEFPPL2 in ligation buffer for 1 h at 37°C. Reactions were treated with 1 mg/ml RNase followed by deproteination. Samples were run on agarose gels and stained by SYBR gold (Invitrogen), and fluorescence was detected via a FluorImager.

In vivo NHEJ assay was performed as described previously (43), and 5 μg Sα–SαII linearized 3′-3′ overhang-ends EGFP-N2 plasmid (CLON-TECH Laboratories, Inc.) were introduced into primary fibroblasts by electroporation. After 72 h, recircularized plasmids were extracted. The junctions were PCR amplified, cloned, and sequenced.

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