Inherited human OX40 deficiency underlying classic Kaposi sarcoma of childhood

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Kaposi sarcoma (KS), a human herpes virus 8 (HHV–8; also called KSHV)–induced endothelial tumor, develops only in a small fraction of individuals infected with HHV–8. We hypothesized that inborn errors of immunity to HHV–8 might underlie the exceedingly rare development of classic KS in childhood. We report here autosomal recessive OX40 deficiency in an otherwise healthy adult with childhood–onset classic KS. OX40 is a co-stimulatory receptor expressed on activated T cells. Its ligand, OX40L, is expressed on various cell types, including endothelial cells. We found OX40L was abundantly expressed in KS lesions. The mutant OX40 protein was poorly expressed on the cell surface and failed to bind OX40L, resulting in complete functional OX40 deficiency. The patient had a low proportion of effector memory CD4+ T cells in the peripheral blood, consistent with impaired CD4+ T cell responses to recall antigens in vitro. The proportion of effector memory CD8+ T cells was less diminished. The proportion of circulating memory B cells was low, but the antibody response in vivo was intact, including the response to a vaccine boost. Together, these findings suggest that human OX40 is necessary for robust CD4+ T cell memory and confers apparently selective protective immunity against HHV–8 infection in endothelial cells.

Kaposi sarcoma (KS) is an inflammatory neoplasm affecting cells of endothelial origin (Ganem, 2010) first described by Moritz Kaposi (Kaposi, 1872). The causal infectious agent of all known...
forms of KS is human herpes virus 8 (HHV-8), also known as KS-associated herpes virus (KSHV; Chang et al., 1994). More than 100 million people are infected with HHV-8, with a heterogeneous worldwide distribution (Plancoulaine et al., 2002). Infection with HHV-8 is asymptomatic in the vast majority of cases (Wang et al., 2001; Andreoni et al., 2002). Only a very small proportion of HHV-8–infected individuals develop KS in their lifetime (Davidovici et al., 2001). Acquired immunodeficiency is a strong KS-predisposing factor; HIV coinfection (epidemic KS) and transplantation–related immunosuppression (iatrogenic KS) increase the risk of KS by factors of at least 3,000 and 100, respectively (Jensen et al., 1999; Serraino et al., 2005; Shiel et al., 2011). Idiopathic cases of KS, striking otherwise healthy individuals with no overt immunological deficit, are mostly reported in the Mediterranean Basin (classic KS) and sub-Saharan Africa (endemic KS). Classic KS is typically an indolent disease of the skin occurring predominantly in the elderly (median age of onset: 65 yr; Iscovich et al., 2000). Classic KS is exceedingly rare in children, with fewer than 40 cases reported since 1960 (Dutz and Stout, 1960; Bisceglia et al., 1988; Akman et al., 1989; Zurrida et al., 1994; Landau et al., 2001; Ferrari et al., 2002; Hussein, 2008; Sahin et al., 2010; Salem et al., 2011; Cakir et al., 2013). The data available before the HIV epidemic suggest that endemic KS is usually more aggressive than the classic form, affecting a younger population (median age of onset: 40 yr), with frequent lymph node involvement (Boshoff and Weiss, 2001). Endemic KS in childhood was rare in Africa in the years before the HIV epidemic, although not as rare as the classic form (Taylor et al., 1972). The rarity of childhood KS contrasts with the relatively high seroprevalence of HHV-8 infection in children <15 yr of age in the Mediterranean region and in Sub-Saharan Africa (Mayama et al., 1998; Andreoni et al., 1999; Gessain et al., 1999). Furthermore, childhood KS, whether it is classic or endemic, runs a more aggressive and disseminated course in children than in adults (Dutz and Stout, 1960; Olweny et al., 1976). Thus, inherited or acquired host factors may underlie classic and endemic KS of childhood.

Inherited immunodeficiencies have been described in two unrelated children with classic KS and other, concurrent, infectious phenotypes: autosomal recessive complete IFN-γ receptor-1 (IFN-γR1) deficiency in a Turkish child with KS and mycobacterial disease (Camcioglu et al., 2004) and X-linked recessive Wiskott–Aldrich syndrome (WAS) in a Tunisian child with KS and EBV lymphoma (Picard et al., 2006). The observation that some children with isolated, classic KS were born to consanguineous parents further suggested that single–gene inborn errors of immunity might underlie such cases (Sahin et al., 2010), as seen in children with other isolated life-threatening infectious diseases (Casanova and Abel, 2007; Alcaïs et al., 2010). Accordingly, autosomal recessive complete STIM1 deficiency was found in a Turkish child with fatal, isolated KS (Byun et al., 2010). Collectively, these findings provided evidence that classic KS in childhood, whether isolated or associated with other infections, may result from single–gene inborn errors of immunity to HHV-8.

The three known KS-predisposing genes, IFNγR1, WAS, and STIM1, are pleiotropic and expressed in various cell types, including leukocytes and nonhematopoietic cells. It thus remains unknown which specific cell types and molecular pathways are critical for the effective control of HHV-8 infection. We therefore searched for new genetic etiologies of classic KS in childhood. We report here the discovery and characterization of autosomal recessive complete OX40 deficiency in a young adult with childhood-onset classic KS, suggesting a nonredundant role of OX40 in T cell–mediated defense against HHV-8.

**RESULTS**

**Homozygous TNFRSF4 (OX40) mutation in a patient with childhood-onset classic KS**

We investigated a 19-yr-old Turkish woman diagnosed with classic KS at the age of 14 yr (II.1 in Fig. 1 A). She had a history of curable visceral leishmaniasis (VL) at the age of 9 yr but remained otherwise healthy. The detailed clinical case report for this patient has been published elsewhere (case 3 in Sahin et al. [2010]). The consanguinity of her parents (I.1 and I.2), who are HHV-8 seropositive but KS free, suggested that susceptibility to KS in this patient followed an autosomal recessive mode of inheritance. For the identification of chromosomal intervals linked to KS, we performed genome-wide linkage analysis by homozygosity mapping. Genotype information of a healthy sibling (II.3) with positive HHV-8 serology was also included in the linkage analysis. In total, we identified 37 regions with positive LOD (logarithm of the odds) scores, with a combined size of 96 Mb (Table S1). We also performed whole-exome sequencing with the patient’s genomic DNA. Whole-exome sequencing identified 32,471 variants, most of which were present in public databases (NCBI dbSNP Build 135, 1000 Genomes release 13, and Exome Variant Server data release ESP6500SI) or our own database (>600 whole-exome sequences from patients with conditions other than KS). Of the 520 variants unique to the patient, 25 were homozygous, 5 of which were within the linked intervals (Table S2). Two resulted in amino acid sequence changes: a missense variant of OXA1L and a missense variant of TNFRSF4. The OXA1L variant was considered unlikely to be damaging because the affected residue was not evolutionarily conserved and the missense variation was predicted to be benign by both PolyPhen-2 (Adzhubei et al., 2010) and SIFT (Kumar et al., 2009). In addition, its surrounding region was homozygous in another healthy sibling (II.2). Although the HHV-8 serology for II.2 was unknown, he had likely been exposed to HHV-8 via close contact with the other family members (Plancoulaine et al., 2000). In contrast, the TNFRSF4 variant was heterozygous in II.2 (Fig. 1 A). This variant was a strong candidate because TNFRSF4 encodes OX40 (CD134), a co-stimulatory molecule which has been implicated in long-term T cell immunity (Croit, 2010). OX40 belongs to the TNF receptor (TNFR) superfamily, a group of type I membrane proteins.

The missense variant p.
other types of mouse lymphoid cells, including NK and NKT cells, but data concerning its expression on cells other than T cells in humans remain limited. We investigated the impact of the R65C mutation on production of the protein by determining OX40 levels by flow cytometry with a monoclonal antibody, ACT35. No OX40 expression was detectable on unstimulated circulating T cells from both healthy controls and the patient (not depicted). OX40 expression was strongly induced on PHA-activated CD4+ T cells and to a lesser extent on activated CD8+ T cells from healthy controls (Fig. 2, A and B). OX40 levels were lower, but not abolished, on the surface of activated T cells from the patient. This was not caused by a lack of response to PHA because the induction of CD25 expression was normal on the patient’s T cells (not depicted). OX40 levels on the surface of PHA-activated T cells from R65C heterozygous family members (I.1, I.2, and II.3) were similar to those of healthy controls (Fig. 2, A and B). The ectopic production of OX40 in Jurkat or HEK-293 cells with retroviral expression vectors containing an internal ribosomal entry site (IRES) followed by GFP resulted in lower levels of R65C mutant protein than of WT protein at the cell surface (Fig. 2 C and not depicted). The lower levels of Arg65Cys (R65C), resulting from C to T substitution at nucleotide position 193 (c.193C>T), affects the first of four cysteine-rich domains in the extracellular region of OX40 (Fig. 1, B and C). The arginine residue in position 65 has not been conserved throughout evolution (Fig. 1 D). However, it is adjacent to the highly conserved Cys64 residue, which forms a disulfide bond with another highly conserved residue, Cys46 (Compaan and Hymowitz, 2006). Introduction of an additional cysteine residue in position 65 may impair the correct disulfide bond formation. R65C variation was predicted to be damaging by both PolyPhen-2 and SIFT. The homozygous R65C variation was confirmed by Sanger sequencing (Fig. 1 B). Consistent with an autosomal recessive trait, both parents and two siblings of the patient were found to be heterozygous for the R65C variation (Fig. 1 A). Finally, the R65C variation was not found in 185 Turkish controls and 973 individuals from the HGDP-CEPH Human Diversity Panel (Cann et al., 2002), further suggesting that it was not an irrelevant polymorphism.

Low levels of mutant OX40 proteins on the cell surface

OX40 is expressed predominantly on activated T cells in both humans and mice (Croft, 2010). It has been detected on the surface of both types of mouse lymphoid cells, including NK and NKT cells, but data concerning its expression on cells other than T cells in humans remain limited. We investigated the impact of the R65C mutation on production of the protein by determining OX40 levels by flow cytometry with a monoclonal antibody, ACT35. No OX40 expression was detectable on unstimulated circulating T cells from both healthy controls and the patient (not depicted). OX40 expression was strongly induced on PHA-activated CD4+ T cells and to a lesser extent on activated CD8+ T cells from healthy controls (Fig. 2, A and B). OX40 levels were lower, but not abolished, on the surface of activated T cells from the patient. This was not caused by a lack of response to PHA because the induction of CD25 expression was normal on the patient’s T cells (not depicted). OX40 levels on the surface of PHA-activated T cells from R65C heterozygous family members (I.1, I.2, and II.3) were similar to those of healthy controls (Fig. 2, A and B). The ectopic production of OX40 in Jurkat or HEK-293 cells with retroviral expression vectors containing an internal ribosomal entry site (IRES) followed by GFP resulted in lower levels of R65C mutant protein than of WT protein at the cell surface (Fig. 2 C and not depicted). The lower levels of

![Figure 1](https://example.com/figure1.png)

**Figure 1.** Homozygous TNFRSF4 mutation in a patient with classic KS. (A) Family pedigree with TNFRSF4 allele segregation. The black-filled symbol indicates the patient (P). Anti–HHV-8 antibody titers, assessed in serum samples collected 3 yr after the initial HHV-8 serology (case 3 in Sahin et al. [2010]), are indicated in parentheses. Ag, antigen; NA, data not available; Neg, negative. (B) Confirmation of the single nucleotide substitution c.193C>T (indicated by an arrowhead) by Sanger sequencing. TNFRSF4 exon 2 was amplified by PCR from genomic DNA from a healthy control and the patient. Representative chromatograms for three independent experiments are shown. (C) Schematic representation of OX40 protein structure. Numbers shown below the scheme indicate the amino acid residue number. CRD, cysteine-rich domain; CYT, cytosolic domain; SP, signal peptide; TM, transmembrane domain. (D) Multiple sequence alignment of human TNFRSF4 and its orthologues. The Arg65 residue of human OX40 (top row) and the corresponding residues in the other species are boxed.
OX40 detected at the cell surface with ACT35 did not result from a change in epitope structure because staining with two other antibodies, H-133 (polyclonal antibody) and EPR17Y (monoclonal antibody), yielded similar results (Fig. 2 C). Overall, these data suggest that the R65C mutation impairs OX40 expression on the cell surface.

Intracellular accumulation of mutant OX40 proteins
We then performed immunoblotting for OX40 proteins. In whole-cell lysates of PHA-activated T cell blasts from healthy controls, most of the OX40 proteins migrated with an approximate mol wt of 50 kD, with a minor fraction observed at 35 kD. The patient’s PHA-activated T cell blasts contained a much smaller proportion of the 50-kD form and a much higher proportion of the 35-kD form (Fig. 3 A). Higher levels of the 35-kD form were also observed in R65C heterozygous individuals (I.1, I.2, and II.3). The lower mol wt forms (35 kD) of OX40 were sensitive to endoglycosidase H and resistant to O-glycosidase (Fig. 3 B), indicating they had immature, high-mannose N-linked carbohydrates and no O-linked carbohydrates. Conversely, the higher mol wt forms (50 kD) were endoglycosidase H resistant and O-glycosidase sensitive (Fig. 3 B), indicating they had complex N-linked and O-linked carbohydrates, these modifications being made in the Golgi complex. The biotin labeling of cell surface proteins in HEK-293 cells that expressed OX40 ectopically further demonstrated that the lower mol wt forms were exclusively intracellular (Fig. 3 C). The accumulation of lower mol wt forms of OX40 with immature carbohydrates in the patient’s cells thus suggests that most of the newly synthesized mutant OX40 proteins remained in the ER and did not enter the Golgi complex. However, the unfolded protein response (UPR) was not activated in the patient’s T cell blasts, as assessed with quantitative RT-PCR to measure the induction of BiP/GRP78.
Figure 3. Intracellular accumulation of the mutant OX40 proteins. (A) Immunoblotting of PHA-activated T cell blasts from three healthy controls (C1–C3), R65C heterozygous family members (I.1, I.2, and II.3), and the patient (P) was performed with the anti-OX40 monoclonal antibody EPR17Y. GAPDH served as a loading control. One experiment representative of three performed is shown. (B) Whole-cell lysates of PHA-activated T cell blasts from a healthy control (C) or the patient (P) were treated with the enzymes indicated before immunoblotting. White lines indicate the intervening lanes have been spliced out. One experiment representative of two performed is shown. EndoH, endoglycosidase H; NT, nontreated; O-gly+Neu, O-glycosidase and neuraminidase. (C) HEK-293 cells transduced with retroviral vectors encoding OX40-WT or OX40-R65C were subjected to cell surface biotinylation. Biotinylated proteins were isolated with avidin-coated beads. OX40 immunoblotting was performed on whole-cell lysate (WCL), the flow-through fraction (FT), and the biotinylated fraction (Biot). A GAPDH blot was used to assess whether the biotinylated fraction was free of intracellular proteins. The results of two independent experiments are shown. (D and E) PHA-activated T cell blasts from healthy controls (C1–C3) or the patient (P) were treated with DMSO alone or treated with either Brefeldin A (BFA) or tunicamycin (TM) for 6 h. The results of two independent experiments are shown. (D) The mRNA levels of BiP/GRP78, relative to GAPDH, were assessed by quantitative RT-PCR. Error bars indicate the SEM. (E) PCR was performed using primer pairs that can differentiate various spliced variants of XBP1 mRNA. (F) Whole-cell lysates of HEK-293 cells transduced with retroviral vectors with an empty vector (Mock) or encoding OX40-WT, OX40-R65C, or OX40-R65A were resolved by SDS-PAGE under reducing or nonreducing conditions. The arrowheads indicate protein bands unique to the OX40-R65C sample. One result representative of three independent experiments is shown. (G) Jurkat cells were transduced with bicistronic retroviruses with an empty vector (Mock) or encoding OX40-WT, OX40-R65C, or OX40-R65A, together with IRES-GFP. Cell surface OX40 levels were assessed with ACT35 by flow cytometry. One result representative of three independent experiments is shown.
mRNA and PCR to detect spliced XBP1 mRNA (Fig. 3, D and E). This was not caused by a general defect in the UPR pathway because Brefeldin A– or tunicamycin-induced activation of the UPR was normal (Fig. 3, D and E). Therefore, the accumulation of mutant OX40 proteins does not seem to induce ER stress.

**Introduction of a cysteine at position 65 is deleterious**

We hypothesized that the acquisition of an additional cysteine residue in position 65 might interfere with correct disulfide bond formation. Indeed, multiple bands unique to OX40-R65C were observed in nonreducing conditions, probably corresponding to OX40 molecules with abnormal intra- and/or intermolecular disulfide bonds (Fig. 3 F; arrowheads). We investigated the impact of the cysteine residue in position 65 further by generating an OX40-R65A mutant. No accumulation of lower mol wt forms was observed in HEK–293 cells expressing OX40-R65A, and the migration pattern of OX40-R65A proteins in nonreducing conditions was similar to that of OX40-WT proteins (Fig. 3 F). The much lower levels of OX40 expression at the cell surface observed in OX40-R65C–transduced Jurkat and HEK–293 cells were not observed in OX40-R65A–transduced cells (Fig. 3 G and not depicted). These results suggest that the introduction of a cysteine residue at position 65, rather than the loss of the arginine residue normally present, is responsible for the impaired cell surface expression of the OX40-R65C mutant proteins.

**R65C is a loss-of-function mutation**

We then investigated whether the R65C mutation led to a functional OX40 deficiency. Human OX40 ligand (OX40L), encoded by TNFSF4, is a member of the TNF superfamily and a type II membrane protein. Its expression, which is often inducible, has been observed in professional antigen-presenting cells including dendritic cells, macrophages, and B cells, as well as other cell types such as endothelial cells, mast cells, and lymphoid tissue inducer cells (Croft, 2010). Its expression on endothelial cells is of particular interest, as KS is driven by the proliferation of HHV-8–infected endothelial cells (Ganem, 2010). Despite the expression of residual mutant OX40 on the cell surface, binding to soluble recombinant OX40L by PHA-activated T cells of the patient was completely abolished (Fig. 4 A). Lentiviral expression of a cDNA encoding OX40-WT, but not OX40-R65C, rescued OX40L binding in the patient’s T cells (Fig. 4 B). Moreover, binding to OX40L was severely impaired in Jurkat cells expressing OX40-R65C, as shown by comparison with cells expressing OX40-WT (16-fold decrease; Fig. 4 C). This defect is much more severe than that in surface expression (three- to fourfold decrease; Fig. 2 C). Similar results were obtained with HEK–293 cells (Fig. 4 C). We assessed the impact of the R65C mutation on T cell co-stimulation in response to membrane-bound OX40L by transducing Vero cells with retroviruses encoding human OX40L or an empty vector. An OX40L–dependent enhancement of cell proliferation was observed in anti-CD3–activated T cells from healthy controls but not in those from the patient (Fig. 4 D). This was not caused by an intrinsic defect in T cell proliferation because the patient’s T cells proliferated normally in response to various doses of anti-CD3 antibody alone or PHA (Fig. 4 E). The R65C is thus a loss-of-function allele, and the patient homozygous for this allele displays complete functional OX40 deficiency. These data suggest that two mechanisms underlie the loss of function of the R65C mutant allele: (1) a low level of OX40 protein on the cell surface and (2) an abolition of the binding to OX40L of the residual OX40 proteins on the cell surface.

**Low proportions of nonnaive T cell subsets in the peripheral blood**

The total number of leukocytes and the proportions of the major leukocyte subsets in peripheral blood were normal in the patient (Table 1). The proportions of T, B, NK, NKT, and γδ T cells in PBMCs were also similar to those of age-matched healthy controls (Table 1). Given the T cell–specific expression of OX40, we carefully examined various subsets of peripheral blood T cells. The proportions of CD4+ and CD8+ T cells in the lymphocyte gate were normal in the patient (Table 1). The proportion of naive CD4+ T cells was higher in the patient than in healthy controls (Fig. 5 A). This reflected the lower proportions of nonnaive subsets, particularly for the CD45RA–CCR7+ effector memory (T EM) population, but also for the regulatory T and CXCR3+ (enriched in IFN–γ–producing T cells [Duhene et al., 2012]) subsets (Fig. 5, A and B). The proportions of the CXCR5+ (follicular helper T cells) and CCR6+ (enriched in IL-17–producing T cells) subsets were similar to those of healthy controls. We also examined CD8+ T cell subsets. The proportion of naive CD8+ T cells and that of CD8+ T EM were only higher and lower in the patient than in healthy controls, respectively (Fig. 5 C). We observed no increase in the proportion of the CD45RA+CCR7– T EMRA subset, which is often expanded in individuals with chronic viral infection, despite the patient suffering from KS over 5 yr (Fig. 5 C). Peptide–MHC class I multimer staining showed the patient to have low and normal frequencies of circulating CMV- and EBV-specific CD8+ T cells, respectively (Fig. 5 D), with a T EM phenotype (not depicted). We also investigated the levels of various cell surface markers, 2B4, CD127, CD103, CLA, CD57, CX3CR1, CD28, CD62L, CD11a, CD11b, CD27, CD95, granzyme B, and perforin, the expression of which changes during the differentiation of naive CD4+ and CD8+ T cells into memory and effector subsets. No overt difference was noted between the patient and the healthy controls for these markers (not depicted).

**Impaired memory CD4+ T cell response to recall antigens**

We assessed the function of memory CD4+ T cells in the patient by measuring cytokine production and proliferation in response to a panel of recall antigens in vitro. PBMCs from Bacille Calmette-Guérin (BCG)–vaccinated healthy controls and BCG–vaccinated R65C heterozygous family members secreted large amounts of IFN–γ in response to purified protein derivative (PPD; Fig. 6 A). However, the patient’s PBMCs
**Figure 4.** R65C is a loss-of-function mutation. (A) PHA-activated T cell blasts from healthy controls (C), R65C heterozygous family members (Het), or the patient (P) were incubated with biotinylated recombinant soluble OX40L. Unbound OX40L molecules were washed out, and the levels of cell-bound OX40L were measured by flow cytometry with allophycocyanin-labeled streptavidin (SA-APC). Representative histograms and MFIs (mean fluorescence intensities) of CD3^+^CD4^+^ cells (CD4^+^ T cells) and CD3^+^CD8^+^ cells (CD8^+^ T cells) in three independent experiments are shown. (B) PHA-activated T cell blasts from the patient were transduced with bicistronic lentiviral vectors encoding luciferase (Luc), OX40-WT, or OX40-R65C, together with IRES-RFP. Cell surface OX40 levels measured with ACT35 and binding to OX40L measured with biotinylated recombinant soluble OX40L are shown for CD3^+^CD4^+^RFP^+^ cells. One result representative of two independent experiments is shown. (C) OX40L binding to Jurkat or HEK-293 cells transduced with bicistronic retroviruses with an empty vector (Mock) or encoding OX40-WT or OX40-R65C, together with IRES-GFP, was assessed as in A. The MFIs of OX40L binding for GFP^+^ cells, normalized with respect to those for isotype controls, are shown. The mean of three independent experiments is shown. Error bars indicate the SEM. *, P < 0.05. (D) CFSE-labeled PBMCs from healthy controls (C1–C8) and the patient (P) were incubated for 3 d with 1 ng/ml of plate-bound anti-CD3 antibody and Vero cells infected with retroviruses either with an empty vector (Vero-Mock) or encoding OX40L (Vero-OX40L). Percentages of CFSE-diluted cells in the CD3^+^CD4^+^ population are plotted. One result representative of two independent experiments is shown. (E) CFSE-labeled PBMCs from healthy controls (C1–C8) and the patient (P) were incubated for 3 d with the indicated concentrations of plate-bound anti-CD3 antibody or PHA. Percentages of CFSE-diluted cells in the CD3^+^CD4^+^ population are plotted. One result representative of two independent experiments is shown.
produced only background levels of IFN-γ, similar to those produced by the cells of healthy controls not vaccinated with BCG, despite the patient having been vaccinated twice with BCG (Fig. 6 A). Furthermore, no IFN-γ production by the patient’s PBMCs was detectable in response to other recall antigens, including tetanus toxoid (TT) and common viruses (CMV, varicella zoster virus [VZV], HSV-1, and EBV), although positive serological results indicated that the patient had been exposed to these antigens (Fig. 6 B and Table 2). The responses of healthy controls to these recall antigens were variable, but all mounted positive IFN-γ response to at least two of the five non-PPD recall antigens tested (Fig. 6 B). Similar results were obtained for IL-10 (Fig. 6, C and D). Proliferative responses to these antigens, as assessed by CFSE dilution assay, were also weak in the patient’s CD4+ T cells (Fig. 6 E). Blocking the OX40–OX40L interaction with neutralizing anti-OX40L antibodies in a similar in vitro recall response assay had no effect on the response of healthy control cells, suggesting that a requirement for OX40–OX40L interactions to elicit in vitro recall responses was not the underlying cause of the defect observed for the patient’s cells in this assay (Fig. 6 F). Instead, these results strongly suggest an in vivo deficiency of antigen-specific memory CD4+ T cells in the patient.

**Low frequency of circulating memory B cells but intact antibody response in vivo**

OX40L can be expressed on activated B cells, and OX40–OX40L interactions may promote T cell–dependent B cell responses in some instances (Morimoto et al., 2000). We thus investigated B cell immunity in the patient. The proportion of CD10+CD27+ memory B cells within the total B cell population was lower and that of CD10−CD27− naïve B cells was higher in the patient than in the age-matched healthy controls (Fig. 7 A). This was not caused by a class-switching defect because the proportions of isotype-switched memory B cells (i.e., those expressing IgG and IgA) in the patient were in the normal range (Fig. 7 B). We also studied B cell differentiation in vitro by culturing purified total B cells with CD40L, IL-10, IL-21, and CpG. B cells from the patient gave rise to IgM-secreting plasmablasts with a frequency similar to that observed for healthy controls (Fig. 7 C). In contrast, the frequencies of in vitro–generated total IgG-secreting B cells (Fig. 7 D) as well as the levels of total (Fig. 7 E) and TT-specific (Fig. 7 F) secreted IgG were markedly lower for the patient than the controls. This probably reflects the paucity of memory B cells in the patients’ sample, as these cells are responsible for the production of the vast majority of Ig, especially IgG, in in vivo cultures of total B cells (Bryant et al., 2007). Consistent with this defect in memory B cell generation, boosting the patient with TT vaccine had no appreciable effect on the frequency of antigen-specific memory B cells, as measured 4 wk after immunization (Fig. 7 F). However, a boost TT vaccination in the patient substantially increased the serum titer of TT-specific antibody in the patient (Fig. 7 G). Moreover, the serum concentrations of total IgG, IgA, IgM, and IgE and of IgG subclasses (IgG1: 7,270 mg/liter, IgG2: 3,850 mg/liter, IgG3: 866 mg/liter, and IgG4: 192 mg/liter) were within the normal range in the patient, and the titers of various antigen-specific antibodies in the patient’s serum were similar to those in healthy family members (Table 2). These data, together with the lack of infectious diseases commonly observed in antibody-deficient patients (Conley et al., 2009), strongly suggest that antibody-mediated immunity is intact in the OX40-deficient patient. These data also suggest that OX40–OX40L interaction is differentially required for the generation of antigen-specific plasma cells versus memory B cells.

**OX40L is highly expressed in KS lesions**

The 19-yr-old OX40-deficient patient presented no severe infectious or tumoral phenotype other than KS and VL. The
pathogenesis of KS and VL may involve the systemic CD4+ memory T cell deficiency seen in the patient. The essential and apparently selective role of OX40 in protective immunity to HHV-8, and perhaps to *Leishmania*, may also reflect the tissue tropism of these pathogens. Indeed, the expression of OX40L on endothelial cells has been reported previously both in vitro (Imura et al., 1996; Kunitomi et al., 2000) and in situ (Souza et al., 1999; Papadopoulos et al., 2013). We also documented the expression of OX40L on primary lymphatic and blood endothelial cells (Fig. 8A). Furthermore, we observed an abundant expression of OX40L with immunohistochemistry in tumor biopsies from AIDS-related KS patients (Fig. 8, B–G). A functional OX40–OX40L interaction between CD4+ T cells and HHV-8-infected endothelial cells, as the antigen-presenting cells, may be required for the generation of protective immunity to HHV-8 in endothelial cells and to prevent the development of KS. Interestingly, we found that the expression of OX40L in LANA (HHV-8 latency-associated nuclear antigen)-positive cells was lower than that of LANA-negative cells lining vascular spaces in the vicinity (Fig. 8, B–E). This may be caused by an HHV-8–induced down-regulation of OX40L to avoid immune surveillance. But the down-regulation may not be complete, and/or T cells may receive OX40L stimulation in trans from the neighboring cells. In any case, we speculate that KS may have occurred in the OX40-deficient patient as a result of the combination of the systemic deficiency in CD4+ memory T cells and the local deficiency of OX40–OX40L interaction between T cells and HHV-8–infected endothelial cells at the disease sites.

**DISCUSSION**

We describe here autosomal recessive complete OX40 deficiency in a patient with childhood-onset classic KS. Our data strongly support a causal relationship between OX40 deficiency and the development of KS in this patient. The homozygous missense mutation in OX40 is the only nonsynonymous private variation, found by whole-exome sequencing and genome-wide linkage analysis, that segregated with the KS phenotype in this kindred. We further demonstrated that the patient’s OX40 allele has a complete loss of function with in-depth biochemical and cellular characterization. The immunological phenotype of the patient largely overlaps with that of OX40-deficient mice (Kopf et al., 1999; Pippig et al., 1999), including, in particular, an impairment of memory
developing classic KS in childhood are both exceedingly rare events (each estimated to affect less than one in 10 million individuals [see Materials and methods]). Overall, these data strongly suggest that KS in this patient is a consequence of OX40 deficiency. Interestingly, the patient had a history of curable VL at 9 yr of age. VL is a parasitic disease caused by Leishmania donovani and L. infantum (chagasi). It is fatal if left untreated (Chappuis et al., 2007). The estimated annual incidence of VL is much higher than that of classic KS in childhood ($L. infantum$ alone accounts for $\approx$50,000 cases worldwide). The causal relation between OX40 deficiency and VL in this patient is thus less evident, but nonetheless plausible for the following reasons: (a) most Leishmania-infected individuals do not develop clinical illness, as demonstrated by the relatively high proportions of asymptomatic carriers in endemic areas (0.6–71.3%,
which this patient had been exposed. No higher morbidity or viral burden was reported in OX40- or OX40L-deficient mice than in the corresponding control groups after acute infections with influenza virus, murine CMV, or vaccinia virus, despite apparent defects in T cell memory (Hendriks et al., 2005; Humphreys et al., 2007; Salek-Ardakani et al., 2008). However, OX40-deficient mice infected with the clone 13 isolate of LCMV, which establishes persistent infection in mice, failed to clear the virus efficiently (Boettler et al., 2012). Thus, studies in mice suggest that OX40 may play an important role in controlling immunity to pathogens that persist. This may also be the case in humans, as both HHV-8 and Leishmania establish persistent infections. However, the patient seemed to be able to control other persistent pathogens, including other herpes viruses, such as EBV and CMV, given their low viral loads in the blood (1,724 copies/ml and 102 IU/ml, respectively). The apparent integrity of central memory CD4+ T cells, antigen-specific memory CD8+ T cells, and antibody response may have been sufficient for protective immunity against these pathogens. The patient's specific susceptibility to HHV-8 and Leishmania may be a combinatorial consequence of the systemic deficit of effector memory CD4+ T cells and the absence of OX40–OX40L-mediated depending on the region and the method of detection; Michel et al., 2011); (b) familial clustering and ethnic differences have suggested a role for host genetic factors in determining the outcome of VL (Cabello et al., 1995; Ibrahim et al., 1999; Khalil et al., 2002; Bucheton et al., 2003); (c) IL-12Rβ1 deficiency was reported as the first genetic etiology of familial VL (Sanal et al., 2007), although susceptibility to VL was a rare finding in IL-12Rβ1–deficient patients (1 out of 141 patients; de Beaucoudrey et al., 2010); (d) the risk of developing clinical VL is higher in patients coinfected with HIV (Murray, 1999; Kumar and Nylén, 2012); (e) Leishmania resides in macrophages, which are known to express OX40L; (f) an OX40 agonist has been shown to enhance the killing of amastigotes in C57BL/6 mice infected with L. donovani (Zubairi et al., 2004); and (g) we showed that liver L. infantum load was much higher in infected OX40-deficient mice than WT mice (unpublished data). The discovery of other genetic disorders underlying VL may further elucidate the mechanisms by which OX40 contributes to protective immunity against Leishmania.

The narrow infectious phenotype of the OX40-deficient patient is intriguing. OX40 is probably not essential for immunity to most childhood pathogens, or at least those to which this patient had been exposed. No higher morbidity or viral burden was reported in OX40- or OX40L-deficient mice than in the corresponding control groups after acute infections with influenza virus, murine CMV, or vaccinia virus, despite apparent defects in T cell memory (Hendriks et al., 2005; Humphreys et al., 2007; Salek-Ardakani et al., 2008). However, OX40-deficient mice infected with the clone 13 isolate of LCMV, which establishes persistent infection in mice, failed to clear the virus efficiently (Boettler et al., 2012). Thus, studies in mice suggest that OX40 may play an important role in controlling immunity to pathogens that persist. This may also be the case in humans, as both HHV-8 and Leishmania establish persistent infections. However, the patient seemed to be able to control other persistent pathogens, including other herpes viruses, such as EBV and CMV, given their low viral loads in the blood (1,724 copies/ml and 102 IU/ml, respectively). The apparent integrity of central memory CD4+ T cells, antigen-specific memory CD8+ T cells, and antibody response may have been sufficient for protective immunity against these pathogens. The patient’s specific susceptibility to HHV-8 and Leishmania may be a combinatorial consequence of the systemic deficit of effector memory CD4+ T cells and the absence of OX40–OX40L-mediated

<table>
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<tr>
<th>Parameter</th>
<th>Patient (normal range)</th>
<th>I.1</th>
<th>I.2</th>
<th>I.3</th>
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<td>14</td>
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<td>IgA (g/liter)</td>
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<tr>
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<td>ND</td>
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*Indicates the threshold for each assay.
In animal models, OX40L blockade in vivo has been shown to attenuate harmful diseases in many inflammatory and autoimmune models, including models of asthma, graft versus host disease, multiple sclerosis, rheumatoid arthritis, inflammatory bowel disease, and transplantation (Croft, 2010). Based on these findings, several clinical trials involving biological molecules targeting OX40 or OX40L are currently being considered or are already under way (http://clinicaltrials.gov). However, our findings suggest that a prolonged blockade of the interaction of OX40 with OX40L may lead to a predisposition to KS and perhaps VL. Close monitoring may therefore be warranted in future trials to decrease the likelihood or severity of such adverse events.

MATERIALS AND METHODS

**Ethics statement.** This study was conducted in accordance with the Helsinki Declaration, with written informed consent obtained from the patient and her family. Approval for this study was obtained from the Rockefeller University Institutional Review Board (New York), the Comité de Protection des Personnes (CPP) and Institut National de la Santé et de la Recherche...
Figure 8. **OX40L is abundantly expressed in KS lesions.** (A) Expression of OX40L on the cell surface was assessed with flow cytometry in three types of primary endothelial cells: human dermal microvascular lymphatic endothelial cells (LEC), human dermal microvascular blood endothelial cells (BEC), and HUVECs. One result representative of two independent experiments is shown. (B–G) OX40L (red) and HHV-8 LANA (brown) expression was assessed by immunohistochemistry in frozen tissue sections. Bars, 5 mm. (B–E) AIDS-related KS lesions in lymph node (B), presenting as a submental mass (C), and in skin (D and E) show HHV-8-positive cells (punctate nuclear brown staining) as well as numerous OX40L-positive cells. (F and G) Non-KS tissues used as controls include normal lymph node (F) and skin with mycosis fungoides (G), showing OX40L in cells lining vascular spaces.

Médicale (France), and the human research ethics committees of the St. Vincent’s Hospital and Sydney South West Area Health Service (Australia).

**Case report.** The detailed clinical case report for this patient has been published elsewhere (case 3 in Sahin et al. [2010]). Since the initial KS diagnosis, she has been sequentially treated with IFN-α, vinblastine, etoposide, and pegylated liposomal doxorubicin hydrochloride, with a partial response. The visceral lesions of KS regressed upon treatment, but widespread dermal lesions on the arms and legs remained. Blood samples were taken for leukocyte phenotyping and functional studies at least 6 mo after the last treatment.
HHV-8 viremia, assessed by quantitative real-time PCR using primers specific to HHV-8 vCyclin, was negative in genomic DNA extracted from the patient's PBMCs.

**Genotyping and linkage analysis.** The patient and her parents were genotyped with the Affymetrix Genome-wide SNP 6.0 array, and the unaffected siblings were genotyped with Affymetrix GeneChip Human Mapping 250K array. Genotype calling was achieved with Affymetrix Power Tools. We discarded monomorphic single nucleotide polymorphisms (SNPs), SNPs with a call rate <100%, and SNPs presenting more than one Mendelian inconsistency within the family. SNPs were further filtered with population-based filters. We then used 108,355 high-quality SNP markers to carry out linkage analysis, assuming autosomal recessive inheritance with complete penetrance (homozygosity mapping). Parametric multipoint linkage analysis was carried out with Merlin software (Abecasis et al., 2002). The Turkish family founders and HapMap CEU trios were used to estimate allele frequencies and to define linkage clusters, with an r² threshold of 0.4.

**Whole-exome sequencing.** Exome capture was performed with the SureSelect Human All Exon 50 Mb kit (Agilent Technologies). Paired-end sequencing was performed on a HiSeq 2000 (Illumina) generating 100-base reads. We aligned the sequences with the GRCh37 reference build of the human genome using the BWA aligner (Li and Durbin, 2009). Downstream processing and variant calling were performed with the Genome Analysis Toolkit (GATK; McKenna et al., 2010), SAMtools (Li et al., 2009), and Picard. Substitution and Indel calls were made with GATK Unified Genotyper. All calls with a read coverage ≤2x and a Phred-scaled SNP quality of 10 were discarded.

**Cell lines and retroviral and lentiviral transduction.** Jurkat cells (ATCC) were grown in RPMI 1640 supplemented with 10% FBS, HEPES–293, and Vero cells (ATCC) were grown in Dulbecco's Modified Eagle's Medium supplemented with 10% FBS. The human TNFRSF4 cDNA (OriGene) was inserted into pMSCV-ires-GFP (pMIG-OX40–WT). Site-directed mutagenesis was performed with the following primer pairs (OX40 indicates the base pair mutations in which mutations were introduced): 5′-GAACACCGTGTGCCTGCCTGTCGCTGGGCGG-3′ and 5′-CCCGCAGCCGACGACACCCCTGTTCC-3′ for pMIG-OX40–R65C and 5′-GAACACCGTGTGCCTGCCGGCTGGCCTGTCGCTGGGCGG-3′ and 5′-GCCGGCAGCCAGGGCCACACCTGTTCC-3′ for pMIG-OX40–R65A. Replication-incompetent retroviruses were produced with the GP2–293 packaging cell line (Takara Bio Inc.). Jurkat and HEK-293 cells were transduced with retroviruses in the presence of 3 µg/ml Polybrene (InvivoGen). Successfully transduced cells were identified as having a red fluorescent protein (RFP) signal (30–90% of live cells were RFP positive at the time of analysis). OX40-expressing lentiviral constructs were generated as previously described (Schoggins et al., 2011) with Gateway cloning technology (Life Technologies). In brief, OX40–WT and -R65C coding sequences were amplified by PCR from pMIG-OX40–WT and -R65C, respectively, with oligonucleotides containing attB sites flanking the coding sequence. The destination vector pTRIPCMVIVShsiresTagRFP_Dest was provided by C. Race (The Rockefeller University, New York, NY). Replication-incompetent lentiviruses were produced in HEK-293T cells and concentrated with Lent-X concentrator (Takara Bio Inc.). PBMCs were activated by incubation with 5 µg/ml PHA for 24 h before the addition of lentiviruses. After 72 h of incubation with lentiviruses, cells were analyzed by flow cytometry. Successfully transduced cells were identified as positive for RFP (10–30% of live cells were RFP positive at the time of analysis).

**Western blotting.** Whole-cell lysates were prepared by incubating cell pellets with NP-40 lysis buffer (10 mM Tris HCl, pH 7.5, 150 mM NaCl, and 1% IGEPA-360 [Sigma–Alrich]) for 30 min on ice and then centrifuging at 20,000 g. OX40 immunoblotting was performed with a rabbit monoclonal antibody, EPR17Y (Epitomics). Antibodies against GAPDH (Santa Cruz Biotechnology, Inc.) and actin (Abgent) were used as for loading controls. When indicated, cell lysates were treated with endoglycosidase H or a combination of O-glycosidase and neuraminidase (New England Biolabs, Inc.) according to the manufacturer’s instructions. Cell surface biotinylation was performed with the Pierce Cell Surface Protein Isolation kit (Thermo Fisher Scientific) according to the manufacturer’s instructions.

**Monitoring the UPR activation.** PHA-activated T cell blasts were generated from PBMCs. T cell blasts were treated with DMSO alone, 1 µg/ml Brefeldin A (GolgiPlug, BD), or 1 µg/ml tunicamycin (Sigma–Alrich) for 6 h. mRNA was extracted from the cells with an RNeasy Mini kit (Qiagen), from which we synthesized cDNA with SuperScript III Reverse transcription (Life Technologies). For quantitative RT-PCR of BiP and GAPDH, we used TaqMan gene expression assays for human HSP45 (Hs00607129_gh; Applied Biosystems) and human GAPDH endogenous control (Hs00607129_m1; Applied Biosystems). The following primer pair was used to detect the spliced variant of XBP1 by PCR: 5′-TTACAGAGAGAACACTCATGCGC-3′ and 5′-GGGTCCAAGTTGTCAGAATGC-3′. The PCR products were separated and visualized in a 4% agarose gel.

**Flow cytometry.** Fluorochrome-conjugated antibodies against human CD3, CD4, CD8, CD10, CD20, CD25, CD27, CXCR5, CXCR3, IgM, IgG, IgA (BD), ACT35 clone directed against OX40, CD45RA (eBioscience), CCR7 (R&D Systems), CCR6, CD56, CD126, VYb (BioLegend), Vo24, and VB11 (Beckman Coulter) and the Live/Dead kit (Life Technologies) were used according to the manufacturers’ instructions. Other antibodies against OX40, H-133 (Santa Cruz Biotechnology, Inc.), and EPR17Y (Epitomics) were used with Alexa Fluor 568–conjugated anti–rabbit Ig (Molecular Probes). Biotinylated recombinant human OX40L (Anclind) was binding detected with allopregnanolone-conjugated streptavidin (eBioscience). Virus-specific CD8 T cells were identified by staining PBMCs with specific soluble peptide–MHC multimers. CMV-specific cells were identified with HLA-A*0201–restricted pentamers (ProImmune) loaded with the NLVPMATV peptide derived from p65 (UL83) protein, and EBV-specific cells were identified with HLA-A*0201–restricted dextramers (Immudex) loaded with the GLCTLUVAML peptide, derived from the lytic antigen BMLF1-1. All data were collected on an LSRII flow cytometer (BD) and analyzed with FlowJo software (Tree Star).

**Recall antigen assay.** PBMCs were labeled with 1 µM CFSE at room temperature, in PBS supplemented with 5% FBS. CFSE-labeled PBMCs were incubated with various recall antigens for 5–6 d in RPMI 1640 supplemented with 10% FBS. The final concentrations and the source of recall antigens were as follows: 5 µg/ml tuberculin PPD, 7.5 µg/ml M. leprae 17 (East Coast Biologics), 7.5 µg/ml PPD, 7.5 µg/ml TT (Statens Serum Institut), uninfected normal human dermal fibroblast (NHDF) extract, CMV-infected NHDF extract, VZV-infected NHDF extract, uninfected Vero cell extract, HSV-1–infected Vero cell extract, and 2.5–5 µg/ml EBV-infected human B cell extract (East Coast Bios). PHA (Sigma–Alrich) was used at a concentration of 5 µg/ml as a positive control. ELISA for cytokines was performed according to the manufacturer's protocol for IFN-γ (R&D Systems) and IL-10 (PeliPair).

**In vitro B cell differentiation.** Total B cells were purified from the OX40 patient as well as from six healthy controls using the Dynal negative selection kit (Invitrogen). B cells were cultured for 5 d with CD40L, 100 U/ml IL-10 (gift from DNAx), 50 ng/ml IL–21 (PeptoTech), and 1 µg/ml CpG (Sigma–Alrich). The frequency of IgM– and IgG-secreting cells was then determined by ELISPOT. Levels of total and TT-specific IgG in the 5-d culture were determined by ELISA (Avery et al., 2010).

**Primary endothelial cells.** Lymphatic endothelial cells (Lonza CC-2810), blood endothelial cells (Lonza CC-2811), and HUVECs (Lonza C2517A) were purchased from Lonza and cultured and maintained according to the manufacturer’s instructions. The levels of TNF superfamily members on the cell surface were measured by flow cytometry. All data were collected on an LSRII flow cytometer and analyzed with FlowJo software.
Immunohistochemistry. Double immunohistochemical staining of OX40L/LANA was accomplished by staining the two antibodies sequentially, using a Bond III Autostainer (Leica). For the first antibody OX40L (clone MM0505-8S23, 1:300; Novus Biologicals), after routine preparation (air-drying at room temperature for 1 h, then fixing in acetone for 10 min), frozen sections were subjected to sequential incubation with the dual endogenous enzyme block (Dako) for 5 min, primary antibody at 4°C for 16 h, post primary AP for 20 min, polymer AP for 30 min, and mixed red refine for 15 min (Bond Polymer Refine Red Detection; Leica). For the second antibody LANA (clone LN53, 1:500; Advanced Biotechnologies), OX40L-stained sections were incubated with the endogenous peroxidase block, primary antibody, post primary, polymer, diaminobenzidine, and hematoxylin for 5, 15, 8, 8, 10, and 5 min, respectively (Bond Polymer Refine Detection; Leica). Finally, sections were dehydrated in 100% ethanol and mounted in Cytosol XYL (Thermo Fisher Scientific).

Estimation of the frequency of individuals with biallelic loss-of-function TNFRSF4 mutations. We assumed that essential splice site mutations, frameshift mutations, stop-gain/loss mutations, and missense mutations that are predicted to be damaging by both PolyPhen-2 and SIFT may be deleterious (loss of function). Only two OX40 variants reported in the public databases (1000 Genomes and Exome Variant Server) fall under these categories: C31R (minor allele frequency 2.3 × 10⁻³⁹) and T62M (minor allele frequency 7.7 × 10⁻⁴³). Thus, the estimated frequency of individuals heterozygous for one of the possible loss-of-function OX40 alleles would be 6.2 × 10⁻⁶ (≈ 2 × [(2.3 × 10⁻³⁹ + 7.7 × 10⁻⁴³) × (1 − 2.3 × 10⁻³⁹ − 7.7 × 10⁻⁴³)]), and the frequency of biallelic carriers would be 9.5 × 10⁻¹² (≈ [(2.3 × 10⁻³⁹ + 7.7 × 10⁻⁴³) + 7.7 × 10⁻⁴³]).

Statistics. Prism software (GraphPad Software) was used to perform unpaired, one-tailed Student’s t tests. Values of P < 0.05 were considered significant.

Online supplemental material. Table S1 shows genomic regions with predicted damaging missense mutations. Nat. Methods. 7:248–249.

REFERENCES

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