AIOLOS Variants Causing Immunodeficiency in Human and Mice

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AIOLOS is encoded by IKZF3 and is a member of the IKAROS zinc finger transcription factor family. Heterozygous missense variants in the second zinc finger of AIOLOS have recently been reported to be found in the families of patients with inborn errors of immunity. The AIOLOS<sup>G159R</sup> variant was identified in patients with B-lymphopenia and familial Epstein–Barr virus-associated lymphoma. Early B-cell progenitors were significantly reduced in the bone marrow of patients with AIOLOS<sup>G159R</sup>. Another variant, AIOLOS<sup>N160S</sup>, was identified in the patients presented with hypogammaglobulinemia, susceptibility to Pneumocystis jirovecii pneumonia, and chronic lymphocytic leukemia. Patients with AIOLOS<sup>N160S</sup> had mostly normal B cell counts but showed increased levels of CD21<sup>lo</sup> B cells, decreased CD23 expression, and abrogated CD40 response. Both variants were determined to be loss-of-function. Mouse models harboring the corresponding patient’s variants recapitulated the phenotypes of the patients. AIOLOS is therefore a novel disease-causing gene in human adaptive immune deficiency.

Keywords: AIOLOS, IKZF3, IKAROS, IKZF transcription factor, inborn errors of immunity

INTRODUCTION

AIOLOS, encoded by the IKZF3 gene, is a member of the IKAROS zinc finger (IKZF) transcription factor family. AIOLOS was first discovered as a binding partner of the pioneer molecule of the IKZF family IKAROS (1). The early studies of Aiolos-deficient mice revealed that Aiolos regulates B cell activation and maturation, Th17 differentiation, and natural killer (NK) cell maturation (2–4). Ikaros and Aiolos cooperatively regulate the expression of λ5, a pre-B cell receptor component, in pre-B cells (5). They also suppress c-Myc expression in pre-B cells, inhibiting pre-B cell proliferation (6). Aiolos, together with Ikaros, is also involved in T cell development. They regulate CD8α expression in thymocytes and regulate Bcl-6 expression in follicular helper T cells (7, 8).

IKZF family transcription factors share the N-terminal C2H2 zinc finger (ZF) domain that is required for DNA binding and the C-terminal C2H2 ZF domain that mediates homo- and heterodimerization (Figure 1A) (9). IKZF proteins comprise the nucleosome remodeling, histone deacetylase complex and polycomb repressive complex 2 in the form of homo- and heterodimers (10, 11). They modulate the chromatin status of the target genes and mediate the activating as well
as the repressive effects. Moreover, these proteins play pivotal roles in maintaining and developing various hematopoietic cells, but are especially important for the development of lymphocytes.

IKAROS was the first member of IKZF family protein to be discovered as the causative gene in the patients with inborn errors of immunity (IEI) (12–14). Heterozygous loss-of-function variants of \( \text{IKZF1} \) result in IKAROS deficiency, which is characterized by a reduction in the levels of B cells and hypogammaglobulinemia often associated with recurrent sinopulmonary infections and other infections (9). Autoimmunity and hematologic malignancies are frequently observed to occur. In particular, IKAROS N159 missense variants exert dominant-negative effect against wild-type IKAROS (14–16). Even though the wide range of clinical phenotypes is observed in the patients with IKAROS loss-of-function variants, the patients with IKAROS N159 missense variants present with a profound combined immunodeficiency (9, 17). All patients developed \( \text{Pneumocystis jirovecii} \) pneumonia (PjP) in the first years of life. Various bacterial, viral, fungal, and parasitic infections were also observed. The patients with IKAROS N159 missense variants manifest profound B-lymphopenia associated with severe hypogammaglobulinemia. T cells were present but manifested predominantly naïve phenotype, and Th polarization defect was observed. The heterozygous and homozygous germline variants of HELIOS have been recently identified in the patients with recurrent sinopulmonary infections, chronic \( \text{Candida} \) infection, hypogammaglobulinemia, and lymphoma (18, 19). Heterozygous HELIOS variants have also recently been reported in patients with autoimmune diseases such as systemic lupus erythematosus, and Epstein–Barr virus (EBV)-associated hemophagocytic lymphohistiocytosis (20). In addition, heterozygous PEGASUS variants have been identified in a cohort with hereditary thrombocytopenia (21).

Somatic mutations of AIOLOS have been reportedly found in hematologic malignancies, including B-cell acute lymphoblastic leukemia (ALL) and chronic lymphocytic leukemia (CLL) (22–25). Two recent studies revealed the germline variants of AIOLOS in human diseases (Figure 1B) (26, 27). Therefore, we reviewed the clinical characteristics and murine models of the AIOLOS variants in these studies.

AIOLOS VARIANTS ASSOCIATED WITH HUMAN IEI

AIOLOS G159R Variant

Yamashita et al. reported the case of three patients in a family with profound B-lymphopenia and familial lymphoma (presented in Figure 2 and Table 1) (26). The index patient was a 42-year-old woman with recurrent sinopulmonary infections since childhood, requiring home oxygen therapy. She presented with splenomegaly in her 30s and subsequently developed cervical non-Hodgkin lymphoma (follicular
lymphoma and EBV-positive diffuse large B-cell lymphoma (DLBCL) at the age of 42 years. Laboratory studies revealed hypogammaglobulinemia, pancytopenia, and profound reduction in B-cell levels. After chemotherapy for lymphoma, she underwent bone marrow transplantation (BMT) and subsequently died of multiple organ failure after transplantation. Her son presented with the same clinical characteristics: recurrent sinopulmonary infection and adult-onset B-cell lymphoma (EBV-positive DLBCL). Additionally, he experienced recurrent perianal abscesses in his childhood. Moreover, his B-cell levels in the peripheral blood lymphocytes were remarkably low. He underwent allogeneic BMT and survived. Conversely, the daughter of the index patient exhibited susceptibility to EBV infection. She experienced recurrent infectious mononucleosis due to EBV at 4 years of age, repeated EBV-associated hemophagocytic syndrome, and contracted chronic active EBV disease (CAEBV) in the following year. B-lymphopenia was noted upon examination. She died of gastrointestinal bleeding associated with CAEBV at the age of 6 years. Autoimmunity was not observed in any of the patients with the AIOLOS\textsuperscript{G159R} variant.

All patients manifested B-cell lymphopenia (0.63%–3% of total lymphocyte). B cell lineage-committed progenitors and immature B cells were almost absent in the index patient, suggesting a defect in B cell differentiation. However, hypogammaglobulinemia (low IgG and IgM, normal IgA) was only found in one patient (index patient, IgG 605 mg/dL, IgA 272 mg/dL, IgM 20 mg/dL). In other patients, IgG and IgA levels were relatively high (IgG 1328–1714 mg/dL, IgA 251–719 mg/dL) and IgM levels were normal to low (IgM 25–123 mg/dL). IgE level was assessed in one patient and found to be high (1940 IU/L).

T cell abnormality in the patients was characterized by CD4\textsuperscript+ T cells skewing to memory and activated phenotypes, as 98% and 30% of CD4\textsuperscript+ T cells were CD45RO\textsuperscript+ and CD38\textsuperscript+HLA-DR\textsuperscript+, respectively. In addition, there was an abnormal Th subset balance, i.e., decreased Th17 and increased Th1* (CD4\textsuperscript+CD45RO\textsuperscript+CD161\textsuperscript+CCR6\textsuperscript+CXCR3\textsuperscript+ ) levels. CD8\textsuperscript+ T cells in the patients also demonstrated an activated phenotype. In addition, reduced expression levels of CD3 and T-cell receptor (TCR) \alpha/\beta on T cells were observed.

**AIOLOS N160S Variant**

Kuehn et al. reported four patients from a single family with susceptibility to PjP and hypogammaglobulinemia as summarized in Figure 2 and Table 1 (27). The index patient died at the age of 42 years. Laboratory studies revealed hypogammaglobulinemia, pancytopenia, and profound reduction in B-cell levels. After chemotherapy for lymphoma, she underwent bone marrow transplantation (BMT) and subsequently died of multiple organ failure after transplantation. Her son presented with the same clinical characteristics: recurrent sinopulmonary infection and adult-onset B-cell lymphoma (EBV-positive DLBCL). Additionally, he experienced recurrent perianal abscesses in his childhood. Moreover, his B-cell levels in the peripheral blood lymphocytes were remarkably low. He underwent allogeneic BMT and survived. Conversely, the daughter of the index patient exhibited susceptibility to EBV infection. She experienced recurrent infectious mononucleosis due to EBV at 4 years of age, repeated EBV-associated hemophagocytic syndrome, and contracted chronic active EBV disease (CAEBV) in the following year. B-lymphopenia was noted upon examination. She died of gastrointestinal bleeding associated with CAEBV at the age of 6 years. Autoimmunity was not observed in any of the patients with the AIOLOS\textsuperscript{G159R} variant.

All patients manifested B-cell lymphopenia (0.63%–3% of total lymphocyte). B cell lineage-committed progenitors and immature B cells were almost absent in the index patient, suggesting a defect in B cell differentiation. However, hypogammaglobulinemia (low IgG and IgM, normal IgA) was only found in one patient (index patient, IgG 605 mg/dL, IgA 272 mg/dL, IgM 20 mg/dL). In other patients, IgG and IgA levels were relatively high (IgG 1328–1714 mg/dL, IgA 251–719 mg/dL) and IgM levels were normal to low (IgM 25–123 mg/dL). IgE level was assessed in one patient and found to be high (1940 IU/L).

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| Clinical phenotypes | AIOLOSG159R/AiolosG158R | AIOLOSN160S/AiolosN159S |
|---------------------|--------------------------|--------------------------|
| **Infection**       | Recurrent sinopulmonary infection (2/3) | Recurrent sinopulmonary infection (3/4) |
|                     | Recurrent EBV-IM (1/3)    | P. jiroveci pneumonia (3/4) |
|                     | Recurrent EBV-VAHS (1/3)  | Bacterial meningitis (1/4), and sepsis (1/4) |
|                     | CAEBV (1/3)               | Extensive cutaneous warts (1/4) |
|                     |                          | M. avium complex lung disease (1/4) |
| **Autoimmunity**    | None                     | None                     |
| **Malignancy**      | EBV-associated lymphoma (2/3) | CLL (1/4)                |
|                     |                          | Metastatic melanoma (1/4) |
| **Immunological phenotypes** | **B cells** | **T cells** |
|                     | ↓ B cell (3/3)            | CD4+ T cells skewed to memory (1/1 evaluated) |
|                     | ↓ B cell-lineage committed progenitor in BM (1/1 evaluated) | CD4+ T cells skewed to memory (1/1 evaluated) |
|                     | ↓ splenic B cells         | CD4+ T cells showed activated phenotypes (1/1 evaluated) |
|                     | Hypogammaglobulinemia (1/3) | Imbalanced Th subsets (¶ Th1 and ¶ Th17) (1/1 evaluated) |
|                     | ↓ FO B cells and MZ B cells | ↓ CD8+ T cells |
|                     |                         | ↓ levels of CD3 and TCRβ (1/1 evaluated) |
|                     |                         | CD4+ T cells skewed to memory (only in homozygous mice) |
|                     |                         | CD4+ T cells showed activated phenotypes (only in homozygous mice) |
|                     |                         | § CD8+ T cells |
|                     |                         | ↓ levels of CD3 and TCRαβ (1/1 evaluated) |
| **In vitro characterization** | LOF                    | LOF                    |
| **Key molecular mechanism** | Interference against IKAROS via formation of heterodimer | Interference against IKAROS via AIOLOSWT |

¶ decreased; § increased; →, normal.
contrasted PjP at infancy, and hypogammaglobulinemia was discovered upon examination. Her father, paternal aunt, and paternal grandfather were also hypogammaglobulinemic, and two of them had histories of PjP. The medical history of her aunt was remarkable, showing oligoarticular arthritis, and development of CLL and metastatic melanoma in her 30s. All the patients underwent immunoglobulin replacement therapies.

Patients with the AIOLOS<sup>N160S</sup> variant are characterized by susceptibilities to opportunistic infections. Three out of four patients contracted PjP, whereas recurrent sinopulmonary infections were observed in all the patients. In addition, meningitis in the aunt and sepsis, extensive cutaneous warts, and Mycobacterium avium complex lung disease in the grandfather were also observed. No EBV infection was identified within this group of patients. Autoimmunity was not observed in these patients with the AIOLOS<sup>N160S</sup> variant.

In contrast to the patients with the AIOLOS<sup>G159R</sup> variant, the patients in this group had normal to high B-cell levels in their peripheral blood (13%–28% of total lymphocytes/234–990 cells/μL) except for the index patient’s grandfather (3% of total lymphocytes/38 cells/μL). Immunoglobulin levels were low in all patients (IgA < 5–10 mg/dL; IgM < 5 mg/dL; IgG 914–1420 mg/dL with immunoglobulin supplementation). Phenotypes of B cells in the patients with AIOLOS<sup>N160S</sup> were characterized by increased levels of CD21<sup>lo</sup> B cells, nearly absent memory B cells, and minimal CD23 expressions. Of note, IgM expressions on B cells were decreased in two of the four patients, and the kappa–lambda light chain ratio was skewed toward the kappa chain in three of the four patients. T cell numbers were normal in the patients with AIOLOS<sup>N160S</sup>. Additionally, naive T cells and recent thymic emigrant levels increased, whereas memory T cells and follicular helper T cell levels decreased in the adult patients. Among all the patients, reduction in Th1 levels was common, whereas the levels of Th2 and Th17 did not reduce.

**MURINE MODELS OF AIOLOS VARIANTS**

IKZF family of transcription factors is evolutionally well-conserved among species (28), with human and murine Aiolos sharing 87% of the amino acid sequences. In the studies of AIOLOS variants in IEI, knock-in mouse models were produced using CRISPR/Cas9 gene-editing technology to further investigate detailed immunophenotypes and elucidate the pathogenic mechanisms caused by these AIOLOS variants. AIOLOS<sup>G159R</sup> and AIOLOS<sup>N160S</sup> in humans correspond to Aiolos<sup>G158R</sup> and Aiolos<sup>N159S</sup> in mice, respectively.

**Aiolos-Deficient Mice**

Aiolos-deficient mice demonstrated increased pre-B cell fraction and a decrease in mature recirculating B cells in the bone marrow, suggesting the partial differentiation defect of B cells (2). Most splenic B cells in Aiolos-deficient mice were IgM<sup>−</sup>IgD<sup>−</sup> follicular B cells, and marginal zone B cells were absent (29). CD21 expression was downregulated in the B cells. These B cells exhibited activated phenotype, and B-cell receptor (BCR)-stimulated proliferation was also strengthened. Increased germinal center formation and elevated serum IgG and IgE levels were observed in the absence of immunization. Autoantibodies were frequently produced.

In contrast to B cells, T cell development in Aiolos-deficient mice was relatively unaffected. Thymocyte and peripheral T cell populations were comparable to wild-type mice (2). However, augmented TCR-mediated proliferation of T cells was observed. Naïve CD4<sup>+</sup> T cells of Aiolos-deficient mice showed impairment in their differentiation into Th17 cells, producing more IL-2 which is directly regulated by Aiolos. Conversely, Aiolos-deficient CD4<sup>+</sup> T cells produced more interferon-γ when activated under Th1-polarizing conditions (3).

NK cells were present in the bone marrow and spleen of Aiolos-deficient mice. CD11b<sup>+</sup>CD27<sup>+</sup> and CD11b<sup>+</sup>CD27<sup>−</sup> NK cell subsets were increased, whereas terminally matured CD11b<sup>+</sup>CD27<sup>−</sup> subset was decreased in Aiolos-deficient mice. Aiolos-deficient NK cells showed hyperresponsiveness to activation and proliferation in response to IL-15 stimulation (4).

Spontaneous development of B-cell lymphoma was observed in aging Aiolos-deficient mice (2). Six out of twenty mice developed lymphoproliferations in the peripheral lymphoid organs. Four out of five examined mice had developed B-cell lymphoma, whereas one developed T-cell lymphoproliferation.

**Aiolos G158R Mice**

Aiolos<sup>G158R</sup> mutant mice showed defects in B-cell developmental with a differentiation blockade at the pre-pro-B cells to pro-B cell and pre-B-cell stages in heterozygous and homozygous mutant mice (26). Splenic B-cell levels were significantly reduced, and all the splenic B-cell subsets, i.e., follicular B cells (B220<sup>+</sup>AA4.1<sup>−</sup>CD23<sup>hi</sup>CD21<sup>hi</sup>), marginal zone B cells (B220<sup>+</sup>AA4.1<sup>−</sup>CD23<sup>hi</sup>CD21<sup>−</sup>), and germinal center B cells (B220<sup>+</sup>GL7<sup>+</sup>CD95<sup>+</sup>CD38<sup>−</sup>), were absent in Aiolos<sup>G158R/G158R</sup> mice. Splenic B cells decreased in Aiolos<sup>+/G158R</sup> as observed in homozygous mutant mice but exhibited a milder phenotype, and B-cell developmental defects in the heterozygous mutant mice was exacerbated with aging.

T-cell phenotypes of Aiolos<sup>G158R</sup> mice were also similar to the T cells abnormalities in the patients. CD3ε and TCRβ expression levels on postselection thymocytes and in the later stages of T cells in the thymus and lymph nodes decreased in heterozygous and homozygous Aiolos<sup>G158R</sup> mice. Furthermore, CD8<sup>+</sup> T cells were decreased in the lymph nodes of Aiolos<sup>G158R</sup> mice. Additionally, in Aiolos<sup>G158R/G158R</sup> mice, CD4<sup>+</sup> T cells were skewed to memory phenotype, and an increase of CD4<sup>+</sup>CD8<sup>−</sup> T cells were detected.

**Aiolos N159S Mice**

Aiolos<sup>N159S</sup> mutant mice showed different B-cell developmental defects than those in Aiolos<sup>G158R</sup>, reflecting the different phenotypes of the patients. In contrast to Aiolos<sup>G158R</sup> mice, the number of B cells in secondary lymphoid organs and peripheral blood did not decrease in Aiolos<sup>N159S/N159S</sup> mice (27). However, B-cell levels within total lymphocytes decreased in Aiolos<sup>N159S/N159S</sup> mice. Characteristic aberrant expression patterns of CD21 and
CD23 on mature B cells were recapitulated in the heterogeneous and homozygous AiolosN159S mice. CD23hiCD21int follicular B cells also decreased in the spleen and peripheral blood of Aiolos+/N159S and AiolosN159S/N159S mice. The cell surface expressions of IgM and IgD were altered on the B cells of heterozygous and homozygous AiolosN159S mutant mice, but the aberrant pattern differed from the human patients. IgG expression was decreased in the B cells of AiolosCd19-L161R/N159S mice, whereas AiolosN159S/N159S B cells showed completely abrogated IgD expression. In addition, the B cells in the Peyer’s patches were almost absent in AiolosN159S/N159S mice, although residual germinal center B cells were observed. A progressive decrease in IgM and IgA was observed in Aiolos+/N159S mice. Immunoglobulin of all classes (IgG, IgA, and IgM) decreased in Aiolos+/N159S/N159S mice. T-cell phenotypes of Aiolos+/N159S mice showed increased T-cell frequency in the peripheral blood in a gene-dosage dependent manner. CD4+ T cells were relatively increased, but the naive CD8+ T cells decreased in AiolosN159S/N159S mice. In addition, follicular helper T cells were reduced in Peyer’s patches of Aiolos+/N159S and almost absent in the Aiolos+/-N159S mice; this may be related to the defects of antibody production in Aiolos+/N159S mice.

Aiolos L161R Mice

Another Aiolos mutant mouse (30), a somatic hotspot mutation in AIOLOS (L162R) found in CLL patients, has been recently reported as a human disease model (30) (Figure 1B) (25). The B-cell-conditional knock-in mouse model harboring the homologous variant L161R (Cd19-Cre; Aiolos+/Cd19-L161R) was hereafter designated as Aiolos+/Cd19-L161R and AiolosCd19-L161R mice, respectively) was generated and studied. In terms of binding to canonical consensus sequence and pericentromeric heterochromatin (PC-HC) targeting, the variant was not identified as loss-of-function (27, 30).

B-cell progenitors in the bone marrow were not affected and total splenic B-cell numbers were comparable to wild-type controls. However, there was a decrease in the marginal zone B cells and increase in follicular B cells in Aiolos+/Cd19-L161R and AiolosCd19-L161R mice. After immunization, an increased fraction of germinal center B cells (B220+CD95+CD38hi) was observed in AiolosCd19-L161R mice. Hyperactivation of BCR signaling and overexpression of nuclear factor-κ B (NF-κB) were observed in AiolosCd19-L161R B cells in a gene-dosage dependent manner.

In elderly AiolosCd19-L161R mice (18–24 months of age), 10% of AiolosCd19-L161R mice and 36% of Aiolos+/Cd19-L161R mice developed CLL-like disease and occasional splenomegaly.

MOLECULAR MECHANISMS UNDERLYING IMMUNODEFIciENCY CAUSE BY AIOLOS VARIANTS

AIOLOS G159R/Aiolos G158R

The AIOLOS G159R variant is characterized in vitro as loss of binding to canonical consensus sequence of AIOLOS as shown by electrophoretic mobility shift assay (EMSA) and defective PC-HC foci formation. Furthermore, genome-wide binding of the AIOLOS G159R variant was assessed using AIOLOS G159R ChIP-seq in the AIOLOS-knockout human pre-B cell line NALM-6. In addition to the loss of binding to its physiological binding sites containing the consensus sequences, novel binding sites were detected for the AIOLOS G159R variant enriched with aberrant motifs.

In a study involving AIOLOS G159R, a murine model was used to investigate the underlying molecular pathogenesis of immunodeficiency caused by the AIOLOS variant. Given that AIOLOS G159R mice exhibited differentiation defects in B cells at the pre-B cell stage and Aiolos is a transcription factor, the pre-B cells of Aiolos+/-G159R mice were subjected to transcriptome analysis. Genes related to B-cell differentiation were downregulated in the pre-B cells of Aiolos+/-G159R mice. Interestingly, dysregulated genes in the Aiolos+/-G159R pre-B cells had higher frequency of Ikaros bindings in their promoter regions than expected frequency calculated from genome-wide Ikaros bindings.

These findings led to the hypothesis that AIOLOS G159R interferes with IKAROS through the formation of heterodimers, resulting in the dysregulation of IKAROS-regulated genes that are essential for B-cell development. Genome-wide bindings of AIOLOS G159R and Ikaros were examined by ChIP-seq experiments in the thymus of AiolosG158R/G158R mice. Ikaros was unable to bind to the canonical Aiolos binding motifs; it preferentially bound to aberrant motifs. Genome-wide binding of Ikaros had changed in AiolosG158R/G158R thymus as hypothesized. Ikaros was often found to co-bind with Aio1os G158R, and the binding motifs of Ikaros changed as Aio1os G158R. In vitro PC-HC formation assay showed that the co-transfection of AIOLOS G159R inhibited PC-HC targeting by IKAROS. Additionally, the transcriptional activities of IKAROS and wild-type AIOLOS were inhibited by the co-transfection of AIOLOS G159R in a dominant-negative manner.

Therefore, this evidence suggests that the immune defect caused by AIOLOS G159R/Aiolos G158R variants results from the inhibition of IKAROS through the formation of a heterodimer with mutant AIOLOS.

AIOLOS N160S/Aiolos N159S

In vitro characterization of the AIOLOS N160S variant suggested a loss-of-function mutant since EMSA indicated that AIOLOS N160S abolished binding to its canonical consensus sequence. When AIOLOS N160S was co-transfected with wild-type AIOLOS for EMSA, the binding of wild-type AIOLOS was reduced to 20% of the wild-type AIOLOS suggesting the dominant-negative effect of AIOLOS N160S against wild-type AIOLOS. PC-HC formation assay demonstrated that the AIOLOS N160S variant is loss-of-function mutant, and it exerted a dominant-negative effect against wild-type AIOLOS. However, AIOLOS N160S had no dominant-negative effect against IKAROS in PC-HC formation assay.

Considering the clinical similarities between the patients with AIOLOS N160S and those with CD40 ligand (CD40L) deficiency,
CD40/CD40L signaling was evaluated in the patients’ B and T cells. CD40 expressions on B cells were not decreased in the patients; however, decreased B cell proliferation upon CD40L plus either anti-IgM, IL-4, or IL-21, was identified in all patients. Moreover, B cells did not differentiate into plasmablasts by the stimulation of CD40L and IL-21. In addition, following PMA and ionomycin stimulation, CD40L expression on activated T cells decreased compared with that in healthy control. These findings suggest that defective CD40L upregulation contributed to the CD40L deficiency-like phenotype, such as susceptibility to P. jirovecii infection. The characteristic CD21\(^{hi}\) B cells in the patients are likely associated with impaired B-cell activation upon CD40L/CD40 stimulation.

RNA-seq analysis of the naive B cells and T-cell blasts showed several dysregulated genes in the naive B cells of patients with AIOLOS\(^{N160S}\). The expressions of several genes related to B-cell development and survival were decreased, including TNFRSF17 (BCMA), TNFRSF13 (APRIL), and TNFSF13B (BAFF). Transcriptomic changes in the T-cell blasts of the patients were associated with immune cell signaling. Additionally, genes involved in the inflammation of respiratory system components, cellular infiltration, and antibody production were downregulated in the patients’ T-cell blasts. Conversely, genes involved in cell survival, homing of cells, chemotaxis, and leukopoiensis were upregulated.

T cells from two patients with AIOLOS\(^{N160S}\) were subjected to AIOLOS ChIP-seq to further characterize the effects of N160S mutation. Clustering analysis revealed that the genome-wide binding of AIOLOS in the patients clustered away from that of healthy controls. However, there was no change in the AIOLOS binding motif among the patients and healthy controls. Furthermore, no direct correlation of change in AIOLOS binding in the T cells and differences in the transcriptome of the T-cell blasts were observed among the patients and healthy controls.

**DISCUSSION AND FUTURE DIRECTION**

The heterozygous AIOLOS\(^{G159R/AiolosG158R}\) and AIOLOS\(^{N160S/AiolosN159S}\) variants caused adaptive immune defects in humans and mice (Figure 2). Both of G159R and N160S variants reside in the ZF2 of the AIOLOS protein, one of four N-terminal ZF that mainly mediates DNA binding. The roles of each ZF are extensively studied in IKAROS. ZF2 and ZF3 of IKAROS are essential in binding to the core IKZF motif, whereas ZF1 and ZF4 determine the specificity to the flanking sequences (31, 32). Within ZF2 and ZF3 of IKAROS, substitutions of the amino acids at -1, 2, 3, and 6 positions in reference to the beginning of the α-helix result in abrogated DNA binding (31, 33, 34). Regarding the positions of substituted amino acids in the AIOLOS variants, G159 and N160 correspond to 2 and 3 position of helical positions, respectively. Thus, missense variants of these amino acids would alter the DNA binding ability of AIOLOS. In vitro and in silico analyses conducted in the studies of AIOLOS\(^{G159R}\) and AIOLOS\(^{N160S}\) supported this hypothesis. The fundamental question lies in understanding the clinical and immunological phenotypes of the patients and mouse models; the mutated amino acid is next to each other, but why were there so many differences? For example, the AIOLOS\(^{G159R}\) variant is associated with a profound decrease in B cells and possible EBV susceptibility, but two of the three patients were not hypogammaglobulinemic. AIOLOS\(^{N160S}\) variant is associated with P. jirovecii infection and severe hypogammaglobulinemia, but B cells were not decreased in the patients’ peripheral blood. The phenotypes of the patients were well recapitulated in the mouse models with the homologous variants. Several hypotheses can explain the difference between AIOLOS\(^{G159R/AiolosG158R}\) and AIOLOS\(^{N160S/AiolosN159S}\), such as 1) the severities of functional defect are different among variants; 2) neomorphic attributes of the variants affect the phenotype; or 3) protein interactions differ among the variants. Both variants were loss-of-function in terms of binding to the canonical consensus sequence and formation of PC-HC and acted in negative-dominance against wild-type AIOLOS. AIOLOS\(^{G159R}\) exerted the dominant-negative effect against IKAROS as demonstrated by PC-HC formation and luciferase reporter gene assays but not AIOLOS\(^{N160S}\). Thus, the degree of functional damage may be greater in the G159R variant. However, patients with AIOLOS\(^{N160S}\) did not necessarily exhibit milder phenotypes. The neomorphic aspects of the mutations may also explain the phenotypic differences. ChIP-seq analyses of AIOLOS\(^{G159R}\) and Aiolos\(^{G158R}\) detected the novel binding motifs preferentially bound by these variants. In contrast, AIOLOS ChIP-seq in the T cells of patients with AIOLOS\(^{N160S}\) did not alter the binding motifs but might have been affected by the bindings of wild-type AIOLOS as half of the AIOLOS protein in the cells would have been the wild-type. In this regard, it is important to compare genome-wide bindings of Aiolos and Ikaros between Aiolos\(^{G158R/G158R}\) and Aiolos\(^{N159S/N159S}\) mice to evaluate how differential bindings by the different AIOLOS variants affect the phenotypes of the patients and mouse models. The current evidence suggests that the newly acquired bindings by AIOLOS\(^{G159R}\) and Aiolos\(^{G158R}\) do not have cis-regulatory effects on surrounding genes but rather work as sequestration of IKAROS to non-physiological binding sites. Although the N-terminal ZFs (ZF1-4) of AIOLOS is known to mediate DNA binding, there is a possibility that the variants in these ZFs also affect protein–protein interaction. The alteration of protein–protein interactions by each variant will be the subject of the future research.

Alternative miRNA splicing of IKZF molecules results in multiple isofoms with differential patterns of domains present in each isoform. The isoforms lacking the whole DNA binding domain or core binding ZFs (ZF2 and ZF3 for Ikaros) are designated as dominant-negative isoforms. Dominant-negative isoforms lack DNA binding ability while maintaining dimerizing capacity, and they inhibit the DNA binding of dimerizing partners. The dominant-negative isoforms of IKZF proteins can be detected in normal lymphocytes (35–37). However, the specific roles of these dominant-negative isoforms in the lymphocyte development are still not known. The expressions of dominant-negative isoforms of IKZF molecules are also observed in various hematologic malignancies (37–41).
**IKAROS**\textsuperscript{N159S} somatic mutation is also found in ALL (42). Contribution of dominant-negative IKAROS to the leukemogenesis is further supported by the observations of spontaneous development of T-cell leukemia and lymphoma in the heterozygous *Ikaros* dominant-negative (AZF1-3) mice and the mice harboring heterozygous *Ikaros*\textsuperscript{H191R} mutation (43, 44). Conditional deletion of *Ikaros* exon 5 (coding ZF2 and ZF3) in common lymphoid progenitors (*Cd2-Cre; Ikaros E5\textsuperscript{fl/fl}* ) also developed T-lymphoid malignancies (45). Pre-B cells of Ikaro *Cd2-Cre; Ikaros E5\textsuperscript{fl/fl}* mice underwent leukemic transformation when transplanted to immunodeficient NOD SCID gamma (NSG) mouse. Ikaro-deficient mice lack B cells, NK cell, and fetal T cell, while T cells develop postnatally (46). On the other hand, *Ikaros* dominant-negative (AZF1-3) mice and mice with ENU-mediated missense mutation in ZF3 (H191R) demonstrate severe lymphocyte developmental defect characterized by absent T, B, and NK cells (44, 47). *Ikaros*\textsuperscript{H191R/H191R} fetal liver cells demonstrated defect in terminal myeloopoiesis and erythropoiesis. Moreover, *Ikaros*\textsuperscript{H191R} mice were fatally lethal (44). The severe phenotypes observed in *Ikaros* dominant-negative mice when compared to Ikaro-deficient mice have been hypothesized as the results of dominant-negative effect against other IKZF molecules by the mutant Ikaros. The studies of AIOLOS variants provided counterpart of this hypothesis. Heterozygous and homozygous *Aiolos*\textsuperscript{G158R} and *Aiolos*\textsuperscript{N159S} variants resulted in severe lymphocyte developmental defects when compared to Aiolos-deficient mice. These observations indicate dominant-negative effect of Aiolos\textsuperscript{G158R} and Aiolos\textsuperscript{N159S} variants against other IKZF molecules. Precisely, it is not clear whether the phenotypes of these mutant mice solely resulted from the dominant-negative effects against other molecules or partially due to neomorphic function by the variants. Also, the roles of each ZF of Aiolos in lymphocyte development have not been studied. In this regard, it is essential to compare the phenotypes of *Aiolos* dominant-negative (AZF1-4) mice, mice lacking individual Aiolos ZFs, and the missense *Aiolos*\textsuperscript{G158R} and *Aiolos*\textsuperscript{N159S} mutant mice.

One interesting feature of the AIOLOS\textsuperscript{N160S} variant is its susceptibility to PJP. The homologous variant in IKAROS (N159S) is associated with similar clinical and immunological phenotypes, including PJP susceptibility (14, 15). B cell levels in patients with AIOLOS\textsuperscript{N160S} were normal to high, whereas class-switched memory B cell and plasmablast levels were low. In addition, CD23 and CD21 expressions were decreased on the B cells. In contrast, B cells in IKAROS\textsuperscript{N159S} patients were almost absent. Both variants were associated with severe hypogammaglobulinemia. T cell abnormalities were characterized by increased naïve CD4\textsuperscript{+} T cells and impaired Th polarization (decreased Th1 in AIOLOS\textsuperscript{N160S} and decreased Th1, Th2, and Th17 in IKAROS\textsuperscript{N159S} patients) in both AIOLOS\textsuperscript{N160S} and IKAROS\textsuperscript{N159S} patients. Impaired CD40/CD40L signals were the cause of PJP susceptibility in patients with AIOLOS\textsuperscript{N160S} but no CD40L signal defects were reported in patients with IKAROS\textsuperscript{N159S}. The precise mechanism underlying PJP susceptibility in patients with AIOLOS\textsuperscript{N160S} and IKAROS\textsuperscript{N159S} should be investigated in future studies.

The development of hematologic malignancies is a common clinical characteristic of IEI caused by AIOLOS variants. EBV-associated lymphoma was observed in two of the three patients with AIOLOS\textsuperscript{N159R} and CLL was observed in one of the four patients with AIOLOS\textsuperscript{N160S}. The tendency to develop hematologic malignancy in patients with the AIOLOS variants is plausible as they were observed in patients with IKAROS and HELIOS deficiencies (17, 19). Additionally, germline variants of IKAROS were also detected in cohorts of patients with B-ALL (48). Although, the precise mechanism underlying tumorigenesis in patients with the AIOLOS variants remains unknown. Two patients with AIOLOS\textsuperscript{G158R} developed EBV-associated lymphoma, whereas another patient presented with chronic and recurrent EBV infections. Thus, the underlying cause of lymphomagenesis is possibly EBV susceptibility. On contrast, somatic loss-of-function AIOLOS mutations were reported in B-ALL (22). Moreover, aged Aiolos-null mice develop B-cell lymphoma (2). These studies and reports of IEI caused by AIOLOS variants indicate a tumor-suppressive role of AIOLOS and the development of hematologic malignancies following the functional deterioration of AIOLOS.

Alternatively, altered interaction with other IKZF molecules and heteromerizing partners or neomorphic features of the AIOLOS variants may have led to cancer development in the patients. The somatic AIOLOS\textsuperscript{L162R} mutation was repeatedly reported in CLL (25). AIOLOS\textsuperscript{N162R} variant was not proven to be loss-of-function in DNA binding and PC-HC targeting abilities (27, 30). Studies on the murine homologous variant *Aiolos*\textsuperscript{L161R} suggested that this variant enhanced BCR and NF-kB signaling, contributing to the CLL-like disease development in aged mice. It is not clear if CLL development in patients with AIOLOS\textsuperscript{N160S} shared the same molecular mechanism as in AIOLOS\textsuperscript{L162R/Aiolos\textsuperscript{L161R}}.

Studies on AIOLOS\textsuperscript{G158R}/Aiolos\textsuperscript{G158R} demonstrated that these variants can cause pathogenicity through heterodimerization with Ikaros, interfering with the physiological function of Ikaros. Seeking therapeutic potential for the patients with IEI caused by AIOLOS variants, we performed gene therapy for AIOLOS\textsuperscript{G158R} mice. A small indel was introduced before the dimerizing ZF (ZF5) of the mutant Aiolos allele, designated as Aiolos\textsuperscript{G158R\Delta-ZF}, using CRISPR/Cas9 gene-editing technology (26). This mutant theoretically had a DNA-binding domain, but it could not form a dimer with wild-type Aiolos or Ikaros and other dimerizing partners. The *Aiolos*\textsuperscript{G158R\Delta-ZF} mice recovered the B-cell levels in the peripheral lymphoid tissues and early B-cell development in the bone marrow was normalized. T cell abnormalities that were observed in Aiolos\textsuperscript{G158R} mice were also normalized. These findings suggest the therapeutic potential of gene therapy and small molecules that interfere with the AIOLOS–IKAROS interaction. However, applying this strategy to the patients will be challenging. Germinal center B cells increased in mice with biallelic C-terminal deletion in the Aiolos, as previously reported in Aiolos-deficient mice. Considering autoimmunity and spontaneous lymphoma development in Aiolos-knockout mice, selective inhibition of mutant AIOLOS allele/protein would be ideal strategy.

Increasing number of novel IEIs is being reported with the advancement of genetic diagnostic technologies. However, most newly described IEIs are sporadic and are often confined to a handful of patients. For example, the first reports of IEI caused by AIOLOS variants include only seven patients in two families.
Although these studies address extremely rare disease, detailed phenotyping of the patients and thorough understanding of underlying mechanism using animal models can provide insights into the unknown but essential aspects of Aiolos in the development and function of lymphocytes. Future studies should address the differences in the phenotypes of immunodeficiency caused by AIOLOS variants and novel therapeutic approaches that specifically inhibit mutant AIOLOS proteins without interfering with the unaffected allele and other heterodimerizing partners should be developed.

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