Generation of induced pluripotent stem cell-derived mice by reprogramming of a mature NKT cell

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Abstract

NKT cells are characterized by their expression of an NKT-cell-specific invariant antigen-receptor α chain encoded by Va14Ja18 gene segments. These NKT cells bridge the innate and acquired immune systems to mediate effective and augmented responses; however, the limited number of NKT cells in vivo hampers their analysis. Here, two lines of induced pluripotent stem cell-derived mice (NKT-iPSC-derived mice) were generated by reprogramming of mature NKT cells, where one harbors both rearranged Va14Ja18 and Vβ7 genes and the other carries rearranged Va14Ja18 on both alleles but germline Vβ loci. The analysis of NKT-iPSC-derived mice showed a significant increase in NKT cell numbers with relatively normal frequencies of functional subsets, but significantly enhanced in some cases, and acquired functional NKT cell maturation in peripheral lymphoid organs. NKT-iPSC-derived mice also showed normal development of other immune cells except for the absence of γδ T cells and disturbed development of conventional CD4 αβ T cells. These results suggest that the NKT-iPSC-derived mice are a better model for NKT cell development and function study rather than transgenic mouse models reported previously and also that the presence of a pre-rearranged Va14Ja18 in the natural chromosomal context favors the developmental fate of NKT cells.

Keywords: iPSC, NKT cell, TCR rearrangement

Introduction

NKT cells are characterized by their expression of an invariant TCR, Va14Ja18 paired with Vβ8.2, Vβ7 or Vβ12 in mice and Va24Jx18/Vβ11 in humans. These stereotypic TCRs recognize glycolipid antigens such as α-galactosylceramide (α-GaICer) in conjunction with the monomorphic MHC class I-like molecule, CD1d (1, 2). Recent studies have demonstrated that NKT cells can be classified into three different functional subtypes, IL-4/IL-13-producing, IL-17A-producing and IFN-γ-producing NKT cells, even though they all express the same invariant antigen receptor (3). These NKT cell subtypes mediate bystander immune regulatory functions, activating various immune effector cell types, including NK cells, macrophages, granulocytes, dendritic cells (DCs), basophils and eosinophils in the innate system as well as CD4 T and CD8 T cells in the acquired system. Therefore, NKT cells participate in the regulation of various disease states, including infection, autoimmunity, allergy, antitumor responses as well as maintenance of transplantation tolerance (1, 2). In fact, X-linked lymphoproliferative disease patients who lack NKT cells die from uncontrolled Epstein–Barr virus infection (4), and NKT-deficient mice also show a shorter survival time when infected with Streptococcus pneumoniae (5). Thus, NKT cells are essential to achieve effective immune responses. On the basis of these multiple functions, NKT cells are considered a promising target for immunotherapy, although there are still some limitations preventing the broad application of NKT cells, especially their extremely low frequency. To overcome this problem, the induced pluripotent stem cell (iPSC) technology is potentially a very powerful tool for analysis and application of NKT cells.
Generation of NKT-iPSC-derived mice

Another important issue in NKT cell biology is to understand the molecular mechanisms of their fate determination. In previous studies, rearranged Vα14Jα18 and Vβ8.2 genes were introduced into RAG-knockout (KO) mice and there was preferential generation of NKT cells but no NK cells, B cells or conventional T cells (6). Moreover, iPSC cell lines obtained by reprogramming of mature NKT cells preferentially generate NKT cells but no γδT, γδT cells, NK cells, DCs or B cells in vitro (7). These results suggest that the pre-rearranged Vα14Jα18/Vβ8.2 TCR genes determine the NKT cell fate. On the other hand, Serwold et al. (8) have demonstrated abnormal T-cell development in the thymus of mice generated by nuclear transfer from mature conventional T cells. Thus, it is of interest to determine whether TCRα or TCRβ chain gene rearrangements are involved in cell fate determination in vivo, particularly in NKT cells.

Here, we successfully generated NKT-iPSC-derived mice by using an NKT cell-derived iPSC (NKT-iPSC) line that was reprogrammed from mature NKT cells of a wild-type (WT) C57BL/6 (B6) mouse. Two types of NKT-iPSC-derived mice are generated from this NKT-iPSC line: one carries a rearranged Vα14Jα18 on both alleles with the TCR Vβ loci in germline configuration (Vα14/WTβ mice) and the other harbors rearrangements of both Vα14Jα18 and Vβ7 on both chromosomes (Vα14/Vβ7 mice). Both types of NKT-iPSC-derived mice had an increased number of NKT cells with relatively normal frequencies of the functional subsets, as well as normal development of all other cell types except for the absence of γδT cells and disturbed development of conventional CD4 γδT cells. Although NKT cells in Vα14/Vβ7 mice had immature phenotypes in the thymus, they acquired functional maturation in the peripheral lymphoid organs.

Methods

Generation of NKT-iPS-derived mice

Two lines of NKT-iPSC-derived mice (Vα14/WTβ and Vα14/Vβ7) were generated as described in Results and Supplementary Figure 1, available at International Immunology Online. In brief, the NKT-iPS line, designated iPSC-58 3E7, was established by reprogramming of mature splenic NKT cells from B6 mice with the Yamanaka factors (9) as previously described (7). The NKT-iPSCs were injected into BALB/c blastocysts to produce chimeric mice. After mating chimeras with B6 mice and genotyping their offspring, pups that harbored rearranged Vα14Jα18 and/or Vβ7 genes in their genomes were chosen for further breeding to generate the NKT-iPSC-derived mice. Germline transmission of the rearranged Vα14Jα18 and Vβ7 loci was achieved with two male chimeras, designated as iPSC-58 3E7-1 and iPSC-58 3E7-2. Genotyping PCR primer sequences are listed in Supplementary Table S1, available at International Immunology Online.

Mice

B6 mice were purchased from Charles River Laboratories or CLEA Japan, Inc. Two lines of NKT-iPSC-derived mice (Vα14/WTβ mice and Vα14/Vβ7 mice) and all other mice were kept under specific pathogen-free conditions and were used at 8–16 weeks of age unless otherwise indicated. All procedures were conducted according to protocols approved by the RIKEN Animal Care and Use Committee.

Flow cytometry

Antibodies (BD Biosciences, eBioscience and BioLegend) were used: APC-Cy7 and Brilliant Violet 421 anti-TCRβ (H57-597), FITC and Pacific Blue anti-CD4 (RM4-5), PE-Cy7 and FITC anti-CD8a (53–6.7), PE anti-CD8b.2 (53–5.8), PE-Cy7 anti-CK1.1 (PK136), PerCP-Cy5.5 anti-CD25 (PC61), FITC anti-CD25 (7D4), FITC anti-Vβ8.1/8.2 (MR5-2), FITC anti-Vβ7 (TR310), FITC anti-Vβ2 (B20.6), PE anti-TCRγδ (GL3), APC anti-TCRδ (eBioGL3), APC anti-CD11c (HL3), FITC anti-CD19 (eBio1D3), PerCP-Cy5.5 anti-B220 (RA3-6B2), FITC and PE anti-CD3e (145-2C11), PE anti-CD1d (B1B), PE anti-Ly108 (330-AU), PE anti-CD150 (9D1), Pacific Blue anti-CD62L (MFL-14), FITC anti-CD24 (M1/69), PE anti-rat IgG1 (A110-1) and APC-eFluor 780 anti-CD117 (2B8). PE anti-FOXP3 (FJK-16s) was used for intracellular staining with BD Cytofix/Cytoperm™ Fixation/Permeabilization Kit (BD Biosciences) according to the manufacturer’s directions. Biotinylated anti-mouse IL-17RB (B5F6) was prepared as previously described (10) and detected by staining with PE-Avidin. α-GalCer-loaded CD1d (α-GalCer/CD1d) dimer (BD Biosciences) for NKT cell enrichment and detection was prepared by the method described previously (11). Cells were analyzed with FACSCanto II (BD Biosciences) or FACSARia II (BD Biosciences). Cell sorting was done using FACSAria II (BD Biosciences). Data were analyzed by FlowJo software (Tree Star).

Quantitative real-time PCR and reverse transcription—PCR

RNA was isolated from FACS-sorted cells using the RNeasy Micro Kit (Qiagen), and cDNAs were generated with the High capacity cDNA Reverse Transcription Kit with RNAse Inhibitor (Applied Biosystems) according to the manufacturer’s instructions. Quantitative real-time PCR (qPCR) primers and probes were designed with Universal Probe Library Assay (Roche) or TaqMan® Gene Expression Assay (Applied Biosystems). The primer sequences and probes are listed in Supplementary Table S3, available at International Immunology Online. qPCR was performed using the ABI PRISM 7900HT system (Applied Biosystems) with FastStart Universal Probe Master (Roche) or by LightCycler 480 (Roche Applied Science) with LightCycler 480 Probes Master (Roche). Results were analyzed using the ΔΔCt method with Gapdh as the internal control with the Universal Probe Library Mouse GAPD Gene Assay (Roche Applied Science). Reverse transcription—PCR (RT–PCR) was done with primers listed in Supplementary Table S3, available at International Immunology Online, using Ex Taq (Takara). PCR consisted of initial denaturation at 96°C for 3 min, followed by 28 cycles of amplification for Gapdh and 35 cycles for Vα–Cα variants with the thermal cycler conditions: 96°C for 30 s, 55°C for 20 s and 72°C for 1 min, and final extension at 72°C for 5 min. PCR products were visualized with SYBR Safe DNA gel stain (Molecular Probes) and visualized with LAS-4000 image analyzer (Fuji Film, Japan).
**Intracellular cytokine staining**

FACS-sorted NKT cells (5 x 10^5) were stimulated with Cell Stimulation Cocktail (plus protein transport inhibitors; eBio-science) for 4 h, followed by intracellular cytokine staining using BD Cytofix/Cytoperm™ Fixation/Permeabilization Kit (BD Biosciences) according to the manufacturer’s instructions. Antibodies (BD Biosciences or eBio-science) used for intracellular cytokine staining were FITC anti-IFN-γ (XMG1.2), PE anti-IL-4 (11B11) and eFluor 450 anti-IL-17A (eBio17B7). Cells were analyzed by FACSCanto II (BD Biosciences). Data were analyzed by FlowJo software (Tree Star).

**Isolation of intraepithelial lymphocytes**

After Peyer’s patches and connective tissues were removed from small intestines, they were opened longitudinally, washed three times in ice-cold PBS buffer to clean their contents, then cut into 0.5 cm pieces and incubated with gentle stirring for 30 min at 37°C in RPMI 1640 containing 1% FCS (Gibco) and 1 mM EDTA. After incubation, tubes were shaken vigorously and the content was passed through a 100 µm cell strainer (BD Falcon), centrifuged at 400 x g for 5 min and washed with PBS buffer containing 2% FCS. Then cell pellets were resuspended in 5 ml of 40% Percoll, and centrifuged at 2000 x g for 10 min. The resulting cell pellet was washed two times with PBS containing 2% FCS and used for flow cytometry analysis.

**Statistical analysis**

Data were presented as mean ± SD or mean ± SEM from three independent experiments. The statistical significance of differences was assessed by the Student’s t-test. Any difference with a P value of <0.01 was considered significant (*P < 0.01).

**Results**

**Generation of NKT-iPSC-derived mice**

We succeeded in reprogramming B6 mature splenic NKT cells with the Yamanaka factors (9) and established an NKT-iPSC line, designated iPSC-58 3E7. As already demonstrated in previous in vitro studies (7), the NKT-iPSCs preferentially generated α-GaICer/CD1d dimer+ TCRβ+ NKT cells but no αβT, γδT, DC, NK or B cells when co-cultured on either OP9 or OP9/Dl1 for 25 days in the presence of Flt3 ligand (Flt3L) and IL-7.

By using this NKT-iPSC line, germline transmission of the rearranged Vα14.Jα18 and Vß7 loci was achieved with two male chimeras, designated as iPSC-58 3E7-1 and iPSC-58 3E7-2 (Supplementary Figure 1A, available at International Immunology Online). Pups with rearranged Vα14.Jα18 and/or Vß7 (Supplementary Figure 1B, available at International Immunology Online) were further bred with B6 mice to generate two lines of NKT-iPSC-derived mice harboring Vα14.Jα18 on both alleles with germline TCRVß (Vα14/WTß mice) or Vα14.Jα18 and Vß7-Dß2-Jß2.3 on both respective alleles (Vα14/Vß7 mice). These mice were used for the present study.

**Preferential development of NKT cells in Vα14/WTß and Vα14/Vß7 mice**

Both Vα14/WTß and Vα14/Vß7 mice showed increased numbers and percentages of α-GaICer/CD1d+ NKT cells in the thymus, spleen and liver compared to B6 mice (Fig. 1A and B). In B6 mice, the absolute number of NKT cells was 4.5 x 10^6 in the thymus, 5.7 x 10^5 in the spleen and 2.9 x 10^5 in the liver. By contrast, there was a threefold increase in NKT cell numbers in the thymus of Vα14/WTß (14.7 x 10^5) and Vα14/Vß7 (15.1 x 10^5) mice, while NKT cells in the spleen (91 x 10^5) in Vα14/WTß, 133 x 10^5 in Vα14/Vß7 mice) and liver (21 x 10^5 in Vα14/WTß, 44 x 10^5 in Vα14/Vß7 mice) were about 10- to 20-fold higher than in B6 mice (Fig. 1B). Under physiological conditions, NKT cells have a strong bias for Vß8, Vß7 and Vß2 usage due to the positive selection mediated by CD1d (12). A similar bias was seen in the Vß repertoire of NKT cells from Vα14/WTß mice (Fig. 1C). On the other hand, the NKT cells in Vα14/Vß7 mice were monoclonal and all cells exclusively used Vß7 due to allelic exclusion by the Vß chain gene locus (13). These results suggest that a normal selection process is taking place during NKT cell development in the thymus. In fact, genes and molecules important for NKT cell development, including CD1d (7, 11) and SLAM family receptor Ly108 (14) (encoded by Slambf6), SAP (4, 15, 16), Fyn (17, 18), c-Mycb (19), c-Myc (20, 21), Runx1 (22), HEB (23) and Egr2 (24), were expressed at normal levels in the thymic CD4/CD8 double-positive (DP) populations in NKT-iPSC-derived mice (Fig. 1D and E), where positive selection of NKT cells is believed to take place (25).

Collectively, the rearranged gene directs robust differentiation towards the NKT cell lineage, and there is preferential development of NKT cells in mice generated from NKT-iPSC.

**Lymphocyte development in Vα14/WTß and Vα14/ Vß7 mice**

We also investigated development of other thymocyte subsets in the NKT-iPSC mice. The absolute number of total thymocytes was severely decreased in both Vα14/WTß and Vα14/Vß7 mice, indicating the disturbed development of conventional T cells, whereas the number of spleen cells and liver mononuclear cells was comparable to B6 mice (Fig. 2A). Among thymocytes, the number of CD4 single-positive (SP) thymocytes in Vα14/WTß mice was 30% that of B6 mice and was only 4% of normal in Vα14/Vß7 mice, whereas the number of CD8 SP thymocytes was similar to B6 in both Vα14/WTß and Vα14/Vß7 mice (Fig. 2B and C). The absolute number of DP thymocytes was severely diminished to 10 and 3% of B6 in Vα14/WTß and Vα14/Vß7 mice, respectively (Fig. 2C). These decreases were observed despite the fact that genes important for the generation of T cells by secondary rearrangement events, such as RAG (26), RORγt (27), the antiapoptotic factors Bcl-xL (28) and Bcl2 (29) (encoded by Rag1, Rag2, Rorc, Bc1l1 and Bcl2, respectively), were highly expressed by DP thymocytes in both Vα14/WTß and Vα14/Vß7 mice (Fig. 2D). To investigate the secondary TCRα chain rearrangement underlying the appearance of non-NKT cells in NKT-iPSC-derived mice, we sorted conventional αβT cells defined as the α-GaICer/CD1d+ TCRα+ NKT-iPSC-derived mice.
CD1d− TCRβ+ population shown in Fig. 1 (A) from the thymus of Vα14/WTVβ and Vα14/Vβ7 mice and analyzed their TCRα chain usage by RT–PCR analysis. Results showed that these conventional αβ T cells expressed Trav6, 8D-1 and 10 located upstream of Vα14 (Trav11 according to the nomenclature by IMGT, the international ImMunoGeneTics information system, http://www.imgt.org) but did not express Trav1 and Trav2 (Supplementary Figure 2, available at International Immunology Online). As for the control, the Trav21, located downstream of Vα14, was not detected in these αβ T cells because the pre-rearranged Vα14Jα18 locus in Vα14/WTVβ mice should leave intact genome upstream of Vα14 (Trav11) and downstream of Jα18 (Traj18) but should have deletion of the genomic region downstream of Vα14 and upstream of Jα18. These data demonstrate that the α-GalCer/CD1d− but TCRβ+ cells are conventional T cells that are derived by the secondary TCRα chain rearrangement events and furthermore suggest that conventional αβ T cells developed in NKT-iPSC-derived mice have limited TCRα chain repertoire.

The number of CD4/CD8 double-negative (DN) thymocytes was normal in Vα14/WTVβ mice, but was decreased to 40% of WT in Vα14/Vβ7 mice (Fig. 2C), indicating disturbed development as early as the thymic DN stage in Vα14/Vβ7 but not in Vα14/WTVβ mice. In order to investigate the development of DN thymocytes, we analyzed the DN stages of thymic ontogeny using CD117 and CD25 mAbs to distinguish the DN1 (CD117+CD25−), DN2 (CD117+CD25+), DN3 (CD117−CD25+), and DN4 (CD117−CD25−) stages. Results from flow cytometry analyses showed severely decreased DN3 stage thymocytes in heterozygous Vα14/Vβ7 mice while Vα14/WTVβ mice showed less dramatic changes although DN4 stage thymocytes were decreased compared with those from B6 mice (Supplementary Figure 3A and B, available at...
Fig. 2. Profiles of lymphocytes in NKT-iPSC-derived mice. (A) Absolute number of total mononuclear cells (MNCs) in the indicated organs of B6, Vα14/WTVβ and Vα14/Vβ7 mice. Data are presented as mean ± SD from three independent experiments. *P < 0.01 versus B6 group. ns, not statistically significant. (B) Representative FACS profiles of CD4/CD8 expression in α-GalCer/CD1d dimer negative thymocyte populations of B6, Vα14/WTVβ and Vα14/Vβ7 mice. (C) Absolute number of CD4 SP, CD4/8 DP, CD4/8 DN and CD8 SP cells in B6, Vα14/WTVβ and Vα14/Vβ7 mice. (D) Relative expression of Rag1, Rag2, Rorc, Bcl2l1, and Bcl2. (E) Absolute number of CD4 SP, CD4/8 DP, CD4/8 DN and CD8 SP cells in B6, Vα14/WTVβ and Vα14/Vβ7 mice. (F) αβ T cell and γδ T cell numbers. (G) Absolute number of NK1.1+, CD3ε+ and CD11c+ splenocytes. (H) Relative expression of NK cell, B cell, and DC genes.
**International Immunology Online**. These data suggest the pre-rearranged state of TCR genes in NKT-iPSC-derived mice causes the disturbed differentiation of DN thymocytes with more severe disturbances observed when both TCRα and TCRβ chains are pre-rearranged.

The frequency and cell number of other immune cells, such as αβT, γδT, NK, B and DCs in lymphoid organs of Vα14/WTVβ and Vα14/Vβ7 mice was also investigated. As shown in Fig. 2 (E and F), no γδT cells were detected in the thymus of either Vα14/WTVβ or Vα14/Vβ7 mice, due to the loss of TCRδ locus by the Vα14Jx18 gene rearrangement events on both chromosomes (Supplementary Figure 1C, available at International Immunology Online). When the frequency and absolute number of splenic DC, B and NK cells was compared to that of B6 mice, no significant difference was observed in either Vα14/WTVβ or Vα14/Vβ7 mice (Fig. 2G and H).

We also analyzed T lymphocyte subsets in NKT-iPSC-derived mice, such as FOXP3+ Treg cells and intraepithelial CD8αα T lymphocytes. Flow cytometry analyses revealed that both Treg and CD8αα cells are detected in Vα14/WTVβ and heterozygous Vα14/Vβ7 mice (Supplementary Figure 4A and B, available at International Immunology Online).

Taken together, these data suggest that both Vα14/WTVβ and Vα14/Vβ7 mice have preferential development of NK cells and also grossly normal development of all immune cell types except for the absence of γδT and disturbed development of conventional CD4 αβT lymphocyte compartments, likely due to the pre-rearrangement of Vα14Jx18 gene loci.

**Development of NKT cell subtypes in Vα14/WTVβ and Vα14/Vβ7 mice**

It has been shown that IL-17 receptor B (IL-17RB) and CD4 are reliable and distinct surface markers to distinguish IL-4-producing (CD4IL-17RB+), IL-17A-producing (CD4IL-17RB+) and IFN-γ-producing (IL-17RB−) NKT cells (3). Therefore, the expression of CD4 and IL-17RB was investigated to define the frequency and absolute number of NKT cell subtypes in various tissues of the NKT-iPSC-derived mice.

The absolute number of all NKT cell subsets was increased about 3.2-fold in the thymus, 12.2-fold in the spleen and 9.0-fold in the liver of Vα14/WTVβ compared to B6 mice (Fig. 3A and B). Similarly, in Vα14/Vβ7 mice, the frequency and total NKT cell numbers were increased 3.4-fold in the thymus, 23.6-fold in the spleen and 18.4-fold in the liver, indicating a significant increase in all subsets of functional NKT cells in NKT-iPSC-derived mice. The frequency of NKT cell subsets was variable in the thymus of NKT-iPSC-derived mice but became relatively normal in the periphery.

Among the three functionally different NKT cell subsets, the CD4IL-17RB−IL-17A-producing subset was dramatically increased in the Vα14/WTVβ and Vα14/Vβ7 mice (Fig. 3B and C). The increase was 45-fold in the thymus, 98-fold in the spleen and 73-fold in the liver of Vα14/WTVβ mice, and in Vα14/Vβ7 mice was 22-fold in the thymus, 57-fold in the spleen and 55-fold in the liver, compared to B6 mice.

There were also increases in the CD4IL-17RB−NKT cells, which are characterized by IL-4 production. There was a 5- to 20-fold increase in their absolute number; in Vα14/WTVβ mice, their increases were 14- to 15-fold in all tissues, while in Vα14/Vβ7 mice, the fold increase was 5-fold in the thymus, 21 in the spleen and 16 in the liver, compared to B6 mice (Fig. 3B and C).

On the other hand, the IFN-γ-producing IL-17RB−NKT cells were increased only 2-fold in the thymus in Vα14/WTVβ mice and 3-fold in Vα14/Vβ7 mice, but were significantly increased in the peripheral tissues, 10-fold in the spleen and 8-fold in the liver of Vα14/WTVβ mice, and 24- and 18-fold in the spleen and liver in the Vα14/Vβ7 mice, compared to those of B6 mice (Fig. 3B and C).

**Functional properties of NKT cell subtypes in Vα14/WTVβ and Vα14/Vβ7 mice**

To investigate functional properties of the NKT cell subtypes in NKT-iPSC-derived mice, cytokine profiles were analyzed by intracellular staining to detect IFN-γ, IL-4- and IL-17A-expressing NKT cells after activation with phorbol 12-myristate 13-acetate and ionomycin in vitro (Fig. 4A and B). The frequency and number of functional NKT cells in Vα14/WTVβ mice were increased 3.6-fold in the thymus, 8-fold in the spleen and 4.7-fold in the liver compared to B6 mice (Fig. 4A−C). Among these, the fold increases were 2.1 for IL-17A-producing NKT cells, 5.1 for the IL-4-producing NKT cells, and 2.8 for the IFN-γ-producing NKT cells, suggesting that there was no impairment in the normal development of functional NKT cells in Vα14/WTVβ mice (Fig. 4D).

On the other hand, in the thymus of Vα14/Vβ7 mice, total numbers of functional NKT cells were decreased by 20% compared to B6 mice, which is significantly lower than that in Vα14/WTVβ mice, where there was a 3.6-fold increase (Fig. 4C). Among the functional NKT cell subsets in the thymus of Vα14/Vβ7 mice, the IL-17A-producing and IL-4-producing NKT cells were 24 and 44%, respectively, of those in B6 mice, whereas the IFN-γ-producing NKT cells were increased 1.1-fold (Fig. 4D), demonstrating the significant developmental arrest in the thymus of Vα14/Vβ7 mice. Interestingly, however, in the peripheral tissues of these mice (Fig. 4C), there were global increases in total numbers of functional NKT cells (28.3-fold increase in the spleen, 5.6-fold in the liver) as well as in numbers of NKT subset cells (2.2-fold increase in IL-17A-producing NKT cells, 10 times in IL-4-producing NKT cells and 44-fold in IFN-γ-producing NKT cells in the spleen and 1.6-fold for IL-17A- and IL-4-producing NKT cells and 7.3-fold for IFN-γ-producing NKT cells in the liver) (Fig. 4D).
Fig. 3. NKT cell subsets in NKT-iPSC-derived mice analyzed by cell surface markers. (A) Representative FACS profiles of CD4/IL-17RB expression on the gated α-GalCer/CD1d dimer+ TCRβ+ NKT cell population in the indicated organs of B6, Vα14/WTVβ and Vα14/Vβ7 mice. (B) Absolute cell number of CD4− IL-17RB+ (blue), CD4+ IL-17RB+ (red) and IL-17RB− (green) fractions of NKT cells in the indicated organs from B6, Vα14/ WTVβ and Vα14/Vβ7 mice. Actual absolute cell numbers are shown in the tables below the figures. Data are presented as mean ± SD from three independent experiments. (C) Fold changes in the number of cells in the different subsets of NKT cells over those of B6 mice are shown.
Fig. 4. Functional NKT cell subsets in NKT-iPSC-derived mice analyzed by intracellular staining. (A and B) Intracellular staining for IFN-γ, IL-4 and IL-17A. NKT cells from the indicated organs were sorted and stimulated with Cell Stimulation Cocktail (plus protein transport inhibitors) for 4h. Negative controls were nonstimulated B6 thymic NKT cells. Representative FACS profiles of IFN-γ/IL-4 and IFN-γ/IL-17A expression.
indicating that the developmental arrest occurring in the thymus of Vα14/Vβ7 mice could be recovered in the peripheral tissues.

In agreement with the above findings shown in Fig. 4 (A–D), expression levels of CD24 (30) and CD62L (31), markers for the immature stage, were lower in liver NKT cells of Vα14/Vβ7 mice, while thymic NKT cells appeared immature based on their expression of high levels of CD24 and CD62L in comparison to Vα14/WTβ and B6 mice (Fig. 4E and F). These data suggested that thymic NKT cells of Vα14/Vβ7 mice are rather immature but can mature in the peripheral tissues.

Discussion

Here, we have shown the successful generation of mice from NKT-iPSC made by reprogramming mature NKT cells of WT B6 mice. The two lines of NKT-iPSC-derived mice (Vα14/WTβ) and Vα14/Vβ7) both preferentially generate large numbers of NKT cells (Fig. 1A and B), a result consistent with previous reports showing that a preponderance of NKT cells resulted from introduction of the rearranged Vα14/xα, either by a transgenic mouse approach (32) or by cloning mice from NK-ES cells (NT-ESC) established by direct transfer of NKT cell nuclei into unfertilized eggs (33–35). The preferential development of NKT cells in NKT-iPSC-derived mice recapitulates results of in vitro experiments (7), indicating that the pre-rearranged Vα14 gene loci in the natural chromosomal context predispose developing thymocytes to an NKT cell fate. Alternatively, this outcome may be interpreted as being the result of higher proliferative responses of β7-expressing NKT cells upon interaction with naturally expressed endogenous ligands (12).

Different from the preferential development of NKT cells, the number of total thymocytes other than NKT cells was strikingly decreased in both Vα14/WTβ and Vα14/Vβ7 mice because of the severe reduction of conventional CD4 T cells associated with decreased numbers of DP cells and perturbed differentiation of DN stage thymocytes, which may be due to the existence of pre-rearranged TCRα loci on both alleles in the NKT-iPSC-derived mice and is reminiscent of results with TCR transgenic models (36, 37). By contrast, CD8 T-cell development was normal in the NKT-iPSC-derived mice, and one possible mechanism is that there exists a bypass mechanism, as proposed by Park et al. using MH Class I-null mice (38). According to this model, the cytotoxic CD8 T-cell developmental pathway is not dependent per se on positive selection mechanisms with cognate MHC. Instead, signals emanating from the IL-7R can bypass TCR-MHC selection, resulting in the differentiation of CD8 T cells in the thymus. In addition, the limited TCR repertoire observed in T cells from NKT-iPSC-derived mice might result in the skewed development of CD8 T cells over CD4 T cells.

The most significant difference between the NKT-iPSC-derived mice and the previous in vitro studies using the NKT-iPSC was in the development of other immune cell lineages. In the in vitro studies, only NKT cells and a few macrophages, but no conventional T, B or DCs, could be generated from the NKT-iPSC line in either the OP9 or OP9/Dll1 culture system (7). However, the NKT-iPSC-derived mice, although they had disturbed development of conventional CD4 αβT lymphocytes, could generate all types of immune cells, except for γδ T cells (Fig. 2E and F), which was due to the elimination of TCRα loci during the pre-rearrangement of Vα14 and Jκ18 gene segments (see Supplementary Figure 1C, available at International Immunology Online). In addition, the NKT-iPSC-derived mice were fully pluripotent by virtue of their normal fertility and ability to reproduce progeny.

Thus, the NKT-iPSC-derived mouse appears to be a better model for NKT cell development and function study over transgenic mouse models, as abnormal development of immune cell types presented in transgenic mouse models (32, 39), as well as other problems might be caused by random integration of artificial promoters or unknown influences by the near-by genes in the integrated chromosome. Moreover, the large number of functional peripheral NKT cells residing in NKT-iPSC-derived mouse might represent another advantageous point for experiments involving cell transfer and measuring levels of molecules.

The number and frequency of the three subsets of NKT cells, CD4– IL-17RB+, CD4+ IL-17RB+ and IL-17RB– NKT cells corresponding to IL-17A-, IL-4- and IFN-γ-producing NKT cell subsets, respectively, are all increased in both Vα14/WTβ and Vα14/Vβ7 mice (Fig. 3A–C). However, the functional maturation of NKT cells was different among these NKT-iPSC-derived mice. In fact, the number and frequency of all functional NKT cell subsets in the Vα14/WTβ mice are high in all tissues, and they are functionally mature in terms of their production of IL-17A, IL-4 and IFN-γ cytokines, while those in Vα14/Vβ7 mice, particularly the IL-17A- and IL-4-producing NKT cell subsets, are significantly lower in the thymus but increased in spleen and liver (Fig. 4A–D). These results suggest that NKT cells in Vα14/Vβ7 mice undergo functional maturation in the periphery. Consistently, thymic NKT cells from Vα14/Vβ7 mice are phenotypically immature based on higher expression levels of CD24 and CD62L, markers of very immature NKT cells (30) and naive conventional T cells (31), whereas these markers are downregulated in the peripheral NKT cells in Vα14/Vβ7 mice (Fig. 4E and F).

The immature state of thymic NKT cells from Vα14/Vβ7 mice might be related with their abnormally rapid development in the thymus, possibly induced by the pre-rearranged TCRβ loci in the chromosome. In agreement with this notion, the decreased number of DN thymocytes with severely diminished DN3 stage thymocytes observed in Vα14/Vβ7 mice (Fig. 2C and Supplementary Figure 3, available at of the gated α-GalCer/CD1d dimer– TCRβ– NKT cells. (A) Absolute cell number of IFN-γ-producing (green), IL-4-producing (red) and IL-17A-producing (blue) NKT cells in thymus, spleen and liver of B6, Vα14/WTβ and Vα14/Vβ7 mice. Data are presented as mean ± SD from three independent experiments. (B) Fold changes in the cell numbers of subsets of NKT cells compared to B6 mice are shown. (C) Representative FACS profiles of CD24 (left) and CD62L (right) expression by gated α-GalCer/CD1d dimer– TCRβ– NKT cells. The open histograms indicate staining with monoclonal antibodies. Each color represents the cells of the indicated organs. The grey histograms indicate the isotype control. (F) Percentage of CD24 (left) and CD62L (right) positive NKT cells in the indicated organs of B6, Vα14/WTβ and Vα14/Vβ7 mice.
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International Immunology Online) indicates the possibility of DN development stage skipping, due to the TCRβ pre-rearrangement, similar to a mouse model with a pre-rearranged TCRβ in a previous study (8).

It is important to compare the quality and function of the in vitro differentiated NKT cells with the in vivo differentiated NKT cells as the in vitro system using iPSC holds a potential to generate unlimited numbers of IFN-γ-producing functional NKT cells. Moreover, these in vitro differentiated NKT cells demonstrated promising characteristics of NKT cells when transferred into recipient mice by inducing adjuvant activity, which resulted in the augmented development of antigen-specific CD8 cytotoxic T cells and the bystander-activation of NK cells in vivo, although their surface phenotypes detected in vitro was reminiscent of immature NKT cells based on their high expression of CD24, double-positivity for CD4, CD8 expression etc. (7). In contrast to the in vitro differentiated NKT cells, the full functional maturation of NKT cells occurs in the peripheral organs, such as spleen and liver. Another noticed difference between the in vitro and the in vivo differentiated NKT cells is that the in vitro differentiated NKT cannot produce IL-17 (data not shown), while the in vivo differentiated NKT cells do produce IL-17 as shown in the present study. Therefore, it is tempting to conclude that the in vitro differentiated NKT cells are rather immature but would be functionally matured upon delivery to the tissue specific microenvironments in vivo.

Finally, it seems that the Vα14WTβ NKT-iPSC-derived mice described here and NKT clone mice with the rearranged Vα14Jα18 and germline ββ generated by NT-ESC (34) are similar in terms of increased number of NKT cells even though they have an abnormality in thymocyte development. However, we demonstrate here that NKT cells from the NKT-iPSC-derived mouse show functional maturation in the peripheral organs by employing systematic analyses of surface phenotypes, gene expression and functional properties of NKT cells from the thymus, spleen and liver, which were not reported by Wakao et al. (34). It is also important to note that the main advantage of iPSC technology over the NT-ESC method is relatively convenience and ethical acceptability, because the iPSC technology does not require human oocytes for the generation of unlimited numbers of functional NKT cells with potential for clinical applications.

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