Leaky severe combined immunodeficiency in mice lacking nonhomologous end joining factors XLF and MRI

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ABSTRACT

Non-homologous end-joining (NHEJ) is a DNA repair pathway required to detect, process, and ligate DNA double-stranded breaks (DSBs) throughout the cell cycle. The NHEJ pathway is necessary for V(D)J recombination in developing B and T lymphocytes. During NHEJ, Ku70 and Ku80 form a heterodimer that recognizes DSBs and promotes recruitment and function of downstream factors PAXX, MRI, DNA-PKcs, Artemis, XLF, XRCC4, and LIG4. Mutations in several known NHEJ genes result in severe combined immunodeficiency (SCID). Inactivation of *Mri, Paxx* or *Xlf* in mice results in normal or mild phenotype, while combined inactivation of *Xlf/Mri, Xlf/Paxx*, or *Xlf/Dna-pkcs* leads to late embryonic lethality. Here, we describe three new mouse models. We demonstrate that deletion of *Trp53* rescues embryonic lethality in mice with combined deficiencies of *Xlf* and *Mri*. Furthermore, *Xlf^{/-}Mri^{-/-}Trp53^{+/-}* and *Xlf^{/-}Paxx^{-/-}Trp53^{+/-}* mice possess reduced body weight, severely reduced mature lymphocyte counts, and accumulation of progenitor B cells. We also report that combined inactivation of *Mri/Paxx* results in live-born mice with modest phenotype, and combined inactivation of *Mri/Dna-pkcs* results in live-born mice with modest phenotype, and combined inactivation of *Mri/Dna-pkcs* results in embryonic lethality. Therefore, we conclude that XLF is functionally redundant with MRI and PAXX during lymphocyte development *in vivo*. Moreover, *Mri* genetically interacts with *Dna-pkcs* and *Paxx*.

INTRODUCTION

Non-homologous end-joining (NHEJ) is a DNA repair pathway that recognizes, processes and ligates DNA double-stranded breaks (DSB) throughout the cell cycle. NHEJ is required for lymphocyte development; in particular, to repair DSBs induced by the recombination activating genes (RAG) 1 and 2 in developing B and T lymphocytes, and by activation-induced cytidine deaminase (AID) in mature B cells [1]. NHEJ is initiated when Ku70 and Ku80 (Ku) are recruited to the DSB sites. Ku, together with DNA-dependent protein kinase, catalytic subunit (DNA-PKcs), forms the DNA-PK holoenzyme [2]. Subsequently, the nuclease Artemis is recruited to the DSB sites to process DNA hairpins and overhangs [3]. Finally, DNA ligase IV (LIG4), X- ray repair cross-complementing protein 4 (XRCC4) and XRCC4-like factor (XLF) mediate DNA end ligation. The NHEJ complex is stabilized by a paralogue of XRCC4 and XLF (PAXX) and a modulator of retroviral infection (MRI/CYREN) [4, 5].

Inactivation of *Ku70*, *Ku80*, *Dna-pkcs* or *Artemis* results in severe combined immunodeficiency (SCID) characterized by lack of mature B and T lymphocytes [2, 3, 6–8]. Deletion of both alleles of *Xrcc4* [9] or *Lig4* [10] results in late embryonic lethality in mice, which correlates with increased apoptosis in the central nervous system (CNS). Inactivation of *Xlf* (*Cernunnos*) only results in modest immunodeficiency in mice [11– 13], while mice lacking *Paxx* [14–17] or *Mri* [5, 18] display no overt phenotype.

The mild phenotype observed in mice lacking XLF could be explained by functional redundancy between XLF and multiple DNA repair factors, including Ataxia telangiectasia mutated (ATM), histone H2AX [19], Mediator of DNA Damage Checkpoint 1 (MDC1) [20, 21], p53-binding protein 1 (53BP1) [17, 22], RAG2 [23], DNA-PKcs [20, 24, 25], PAXX [4, 14, 15, 20, 26– 28] and MRI [5]. However, combined inactivation of Xlf and Paxx [4, 14, 15, 20], as well as Xlf and Mri [5], results in late embryonic lethality in mice, presenting a challenge to the study of B and T lymphocyte development in vivo. It has also been shown that both embryonic lethality and increased levels of CNS neuronal apoptosis in mice with deficiency in Lig4 [9, 10, 29, 30], Xrcc4 [9, 31], Xlf and Paxx [20], or Xlf and Dna-pkcs [24, 25] is p53-dependent.

In this study, we rescue synthetic lethality from *Xlf* and *Mri* by inactivating one or two alleles of *Trp53*. We also show that both $Xlf'Mri''Trp53^{+/-}$ and Xlf'Paxx''. *Trp53*^{+/-} mice possess a leaky SCID phenotype with severely reduced mature B and T lymphocyte counts in the spleen, low mature T cell counts in the thymus, and accumulated progenitor B cells in the bone marrow. Finally, we demonstrate that MRI is functionally redundant with DNA-PKcs and PAXX.

RESULTS

Inactivation of *Trp53* gene rescued embryonic lethality in mice lacking XLF and MRI

Combined inactivation of *Xlf* and *Mri* has previously been shown to result in synthetic lethality in mice [5]. To generate XLF/MRI deficient mice with altered expression of *Trp53*, we intercrossed an *Mri^{-/-}* strain [18] with an *Xlf^{-/-}Trp53^{+/-}* [20] strain. Next, we selected and intercrossed triple heterozygous $(Xlf^{+/-}Mri^{+/-}Trp53^{+/-})$, and later, *Xlf^{-/-}Mri^{+/-}Trp53^{+/-}* mice. With PCR

screening, we identified $Xlf^{/-}Mri^{'-}Trp53^{+/-}$ (n=11), $Xlf^{/-}Mri^{'-}Trp53^{+/-}$ (n=2), and $Xlf^{/-}Mri^{'-}Trp53^{+/+}$ (n=1) (Figure 1A) among the resulting offspring. Mice lacking both XLF and MRI possessed reduced weight (12 g on average, p<0.0001) when compared with gender- and age-matched WT (19 g), $Xlf^{/-}$ (19 g) and $Mri^{-/-}$ (20 g) controls (Figure 1B and 1C). In addition, $Xlf^{/-}Mri^{'/-}Trp53^{+/-}$ and $Xlf^{/-}Mri^{'/-}Trp53^{-/-}$ mice were viable up to 63 days and died for unknown reasons. We used $Xlf^{/-}Mri^{'/-}Trp53^{+/-}$ mice to further characterize the development of B and T lymphocytes *in vivo*.

Leaky SCID in *Xlf^{-/}Mri^{-/-}Trp53^{+/-}* mice

To determine the roles of XLF and MRI in lymphocyte development in vivo, we isolated the thymus, spleen, and femur from $Xlf^{-}Mri^{-/-}Trp53^{+/-}$ mice, as well as from $Xlf^{/-}$, $Mri^{-/-}$, $Trp53^{+/-}$ and WT controls. Combined deficiency for XLF and MRI resulted in a 3-fold reduction in thymus size (32 mg on average, p < 0.0001) and a 9-fold reduction in thymocyte count (1.9×10^7) , p < 0.0001) when compared to single deficient or WT controls (Figure 1D). Similarly, both average spleen weight (22 mg, p<0.0001) and splenocyte count $(2.0 \times 10^7, p < 0.0001)$ in $X l f'^{-} M r i^{-7} T r p 53^{+/-1}$ mice decreased approximately 4-5 fold when compared with WT and single deficient controls (Figure 1E). The reduced number of splenocytes in XLF/MRI doubledeficient mice could be explained by decreased populations of B and T lymphocytes observed in the Xlf $Mri^{-T}Trp53^{+/-}$ mice (Figure 1F–1H and Supplementary Tables 1-4). Specifically, CD3+ T cells were reduced 6fold (p < 0.0001), while CD19+ B cells were reduced 50fold (p < 0.0001) when compared with single deficient and WT controls (Figure 1F-1H). Likewise, counts of CD4+ and CD8+ T cells in the spleen (Supplementary Tables 3 and 4), were all dramatically reduced when compared with single deficient and WT controls (about 4-fold, p < 0.0001; Figure 1F, 1H) as well as counts of CD4+, CD8+ and CD4+CD8+ T cells in the thymus (Figure 1F, 1I and Supplementary Tables 5-7). From these observations, we conclude that XLF and MRI are functionally redundant during B and T lymphocytes development in mice.

Leaky SCID in mice lacking XLF and PAXX

Combined inactivation of XLF and PAXX has been shown to result in embryonic lethality in mice [4, 14, 15, 20]. To determine the impact of XLF and PAXX on B and T cell development *in vivo*, we rescued the synthetic lethality by inactivating one allele of *Trp53*, as described previously [20]. We did not detect any direct influence of altered *Trp53* genotype on lymphocyte development (Supplementary Tables 1–9). The resulting *Xlf^{/-}Paxx^{-/-}Trp53^{+/-}* and *Xlf^{/-}Paxx^{-/-}Trp53^{-/-}* mice possess

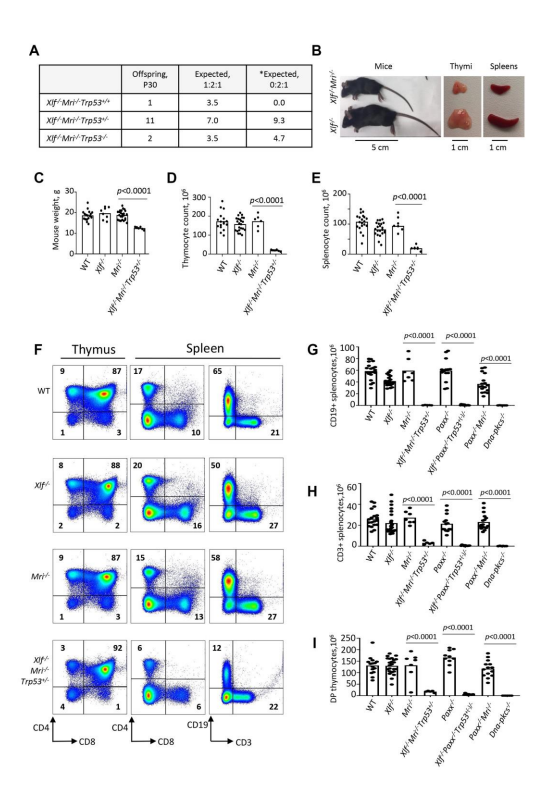


Figure 1. Development of B and T lymphocytes in *Xlf^{-/}Mrī^{/-}Trp53^{+/-}* **mice.** (A) Number of thirty-day-old mice (P30) of indicated genotypes. *Expected distribution assuming lethality. (B) Comparison of body size, thymi and spleens of XLF/MRI-deficient and XLF-deficient mice of the same age. (C) Weights of WT, *Xlf^{/-}, Mrī^{/-}, Xlf^{/-}Mrī^{/-}Trp53^{+/-}* mice. (D, E) Number (×10⁶) of thymocytes (D) and splenocytes (E) in WT, *Xlf^{/-}, Mrī^{/-}, Xlf^{/-}Mrī^{/-}Trp53^{+/-}* mice. (D, E) Number (×10⁶) of thymocytes (D) and splenocytes (E) in WT, *Xlf^{/-}, Mrī^{/-}, Xlf^{/-}Mrī^{/-}Trp53^{+/-}* mice. (F) Flow cytometric analysis of thymic and splenic T cell subsets and splenic B cells. (G, H, I) Number (×10⁶) of splenic CD19+ B cells (G), splenic CD3+ T cells (H) and thymic CD4+CD8+ double positive (DP) T cells (I) in WT, *Xlf^{/-}, Mrī^{/-}, Xlf^{/-}Mrī^{/-}* Trp53^{+/-} mice. *Dna-pkcs^{-/-}* mice were used as an immunodeficient control. Comparisons between every two groups were made using one-way ANOVA, GraphPad Prism 8.0.1. *Xlf^{/-}Paxx^{-/-}Trp53^{+(-)/-}* is a combination of *Xlf^{/-}Paxx^{-/-}Trp53^{+/-}* and *Xlf^{/-}Paxx^{-/-}Mrī^{/-}*, *p*<0.0001 (****), Antī^{/-}, *p*<0.0001 (****), *Mrī^{/-}*, *p*<0.0001 (****), and *Xlf^{/-}Paxx^{-/-}Trp53^{+(-)/-}* vs *Paxx^{-/-}Mrī^{/-}*, *p*<0.0001 (****).

30- to 40-fold reduced thymocyte count (4.0×10^6) , p < 0.0001) when compared to WT (1.3x10⁸), Xlf^{/-} (1.4×10^8) and $Paxx^{-/-}$ (1.7×10^8) mice. This is reflected in decreased levels of double-positive CD4+CD8+ cells, as well as decreased levels of single-positive CD4+ and CD8+ T cells (Figure 1, Supplementary Figure 1, and Supplementary Tables 5–7). Spleen development was dramatically affected in mice lacking XLF and PAXX compared to WT and single-deficient controls, due to the lack of B cells and decreased T cell count (Figure 1, Supplementary Figure 1, and Supplementary Tables 1-4). When compared with the WT and single knockout controls, Xlf⁻Paxx^{-/-}Trp53^{+/-} and Xlf^{/-}Paxx^{-/-}Trp53^{-/-} mice had a 100- to 600-fold reduction in CD19+ B splenocyte count $(0.7 \times 10^6, p < 0.0001)$ and a 50- to 90fold reduction in CD3+ splenocyte count (to $0.5 \times 10^{\circ}$) (Figure 1F-1H and Supplementary Figure 1). From these results, we concluded that XLF and PAXX are functionally redundant during the B and T lymphocyte development in vivo.

Early B cell development is abrogated in mice lacking XLF and MRI, or XLF and PAXX

Reduced counts and proportions of mature B lymphocytes in Xlf' Mri'' $Trp53^{+/-}$ mice suggest a blockage in B cell development in the bone marrow. To

investigate this further, we isolated the bone marrow cells from femora of mice lacking XLF. MRI or both XLF/MRI, and analyzed the proportions of B220+CD43+IgM- progenitor B cells and B220+CD43-IgM+ immature and mature B cells. We detected only background levels of B220+CD43-IgM+ B cells in bone marrows isolated from $Xlf^{-}Mri^{-}Trp53^{+/-}$ mice (Figure 2A, 2B and Supplementary Table 8). However, these mice exhibited a 2- to 3-fold higher proportion of pro-B cells when compared with WT, Xlf^{-/-} and Mri^{-/-} controls (Figure 2A, 2C and Supplementary Table 9). Similarly, $Xlf^{-}Paxx^{-}Trp53^{+/-}$ and $Xlf^{-}Paxx^{-}Trp53^{-/-}$ mice also possess background levels of IgM+ B cells (p < 0.0001; Figure 2A, 2B and Supplementary Table 8) while having 3- to 4-fold higher proportion of pro-B cells when compared with WT, Xlf^{\prime} and $Paxx^{\prime}$ controls (*p*<0.0001; Figure 2A, 2C and Supplementary Table 9). Therefore, we conclude that B cell development is blocked at the pro-B cell stage of $Xlf^{-}Mri^{-}Trp53^{+/-}$ and $Xlf^{-}Paxx^{-}Trp53^{+}$ mice.

Paxx^{-/-}Mri^{-/-} mice possess a modest phenotype

Both PAXX and MRI are NHEJ factors that are functionally redundant with XLF in mice. Combined inactivation of *Paxx* and *Xlf* [4, 14, 15, 20], or *Mri* and *Xlf* [[5]; this study) results in synthetic lethality in mice,

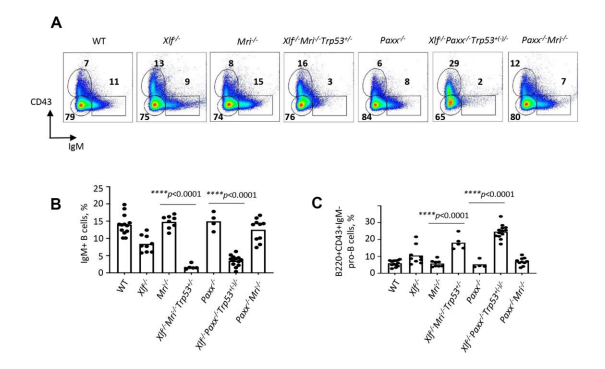
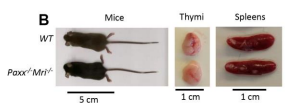


Figure 2. Development of B cells is abrogated in bone marrow of *Xlf^{/-}Trp53^{+/-}* **and** *Xlf^{/-}Paxx^{-/-}Trp53^{+/-}* **mice.** (A) Flow cytometric analysis of developing B cells. Upper left boxes mark B220+CD43+lgM- progenitor B cell populations, and lower right boxes mark the B220+CD43-lgM+ B cells. (B, C) Frequencies (%) of B220+CD43-lgM+ B cells (B) and B220+CD43+lgM- progenitor B cells (C) in WT, *Xlf^{/-}*, *Xlf^{/-}Mri^{/-}*, *Xlf^{/-}Paxx^{-/-}*, *Xlf^{/-}Paxx^{-/-}*, *Xlf^{/-}Paxx^{-/-}*, *Xlf^{/-}Paxx^{-/-}*, *Xlf^{/-}* and *Paxx^{-/-}*, *Trp53^{+(-/-}* and *Paxx^{-/-}*, *Trp53^{+(-/-}* and *Paxx^{-/-}*, *Trp53^{+(-/-}* and *Paxx^{-/-}*, *Trp53^{+/-}*.

Α		Offspring, P30	Expected, 1:2:1
	Paxx ^{-/-} Mri ^{+/+}	2	3.25
	Paxx ^{-/-} Mri ^{+/-}	4	6.50
	Paxx ^{-/-} Mri ^{-/-}	7	3.25
	Total	13	13.00



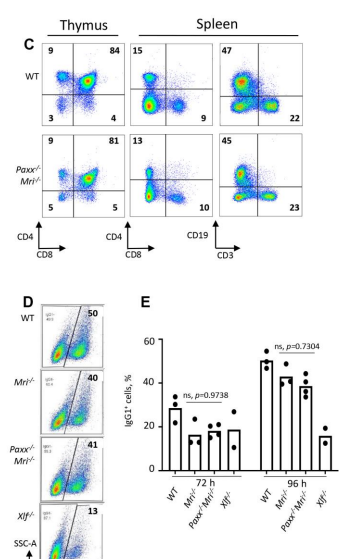


Figure 3. Development of B and T cells in $Paxx^{-/-}Mri^{-/-}$ **mice.** (A) Number of thirty-day-old mice (P30) of indicated genotypes. Parents were $Paxx^{+/-}Mri^{+/-}$ and $Paxx^{-/-}Mri^{+/-}$. (B) Example of thirty-day-old $Paxx^{-/-}Mri^{-/-}$ and WT male littermates with their respective thymi and spleens. (C) Example of flow cytometry analyzes of B and T cells in $Paxx^{-/-}Mri^{-/-}$ and WT mice. (D, E) Class switching analyzes of *in vitro* activated naïve B cells of indicated genotypes.

►lgG1

as well as in abrogated V(D)J recombination in vAbl pre-B cells [4, 5, 14, 15, 27]. To determine if Paxx genetically interacts with Mri, we intercrossed mice that are heterozygous or null for both genes (such as Paxx^{-/-} $Mri^{+/-}$ and $Paxx^{+/-}Mri^{+/-}$). We found that resulting $Paxx^{-/-}Mri^{-/-}$ mice are live-born, fertile, and are similar in size to WT littermates (17 g, p>0.9999) (Figure 3A and 3B). Specifically, we observe that Paxx^{-/-}Mri^{-/-} mice have normal thymocyte and splenocyte counts. Furthermore, *Paxx^{-/-}Mri^{-/-}* mice underwent normal T cell development that was indistinguishable from the WT, Paxx^{-/-}, and Mri^{-/-} controls (Figures 1H, 1I and 3C). However, Paxx⁻ $^{/}Mri^{/-}$ mice had reduced CD19+ B cell counts (Figure 1G) when were compared to WT, Paxx^{-/-} and Mri^{-/-} controls (p<0.0025). Moreover, CD19+ B cell counts were similar in $Paxx^{-/-}Mri^{-/-}$ and $Xlf^{/-}$ mice (p>0.9270), suggesting that combined depletion of PAXX and MRI has modest phenotype similar to the one in XLFdeficient mice. CSR to IgG1 was performed in order to determine if DNA repair-dependent immunoglobulin production is affected in mature B cells lacking PAXX and MRI [16, 18]. Paxx inactivation did not affect Ig switch to IgG1 in MRI-deficient B cells (Figure 3D and 3E). The quantity of IgG1+ cells after CSR stimulation was similar between $Paxx^{-/-}Mri^{-/-}$ and $Mri^{-/-}$ naïve B cells (p>0.73). From this, we can conclude that there is a genetic interaction between Paxx and Mri in vivo, and it is only detected in B cells.

Synthetic lethality between *Mri* and *Dna-pkcs* in mice

Both MRI and DNA-PKcs are functionally redundant with XLF in mouse development [5, 24]. Combined inactivation of Paxx and Mri (this study), or Paxx and Dna-pkcs [20] genes results in live-born mice that are indistinguishable from single deficient controls. To determine if Mri genetically interacts with Dna-pkcs, we crossed $Mri^{+/-}$ and $Dna-pkcs^{+/-}$ mouse strains, then intercrossed the double-heterozygous Mri^{+/-}Dna-pkcs^{+/-}, and then *Mri^{-/-}Dna-pkcs^{+/-}* mice (Figure 4A). We identified 12 *Mri^{-/-}Dna-pkcs^{+/+}* and 12 *Mri^{-/-}Dna-pkcs^{+/+}*, but no *Mri^{-/-}Dna-pkcs^{-/-}* mice (out of 6 expected). To determine if double-deficient Mri^{-/-}Dna-pkcs^{-/-} embryos are present at day E14.5, we intercrossed Mri-Dna $pkcs^{+/-}$ mice, extracted and genotyped the embryos (Figure 4B). We identified two *Mri^{-/-}Dna-pkcs^{-/-}* mice at E14.5 (63mg), which were about 40% lighter than Mri^{-1} littermates (108mg) (Figure 4C and 4D). A Chi-Square test (χ^2) was performed to determine if the embryonic distribution data fits the mendelian ratio of 1:2:1 that is expected from Mri^{-/-}Dna-pkcs^{+/-} parents. With DF=2 and $\chi^2 = 1.8$, the corresponding p-value lies within the range 0.25 . This affirms that our data fit the expected1:2:1 distribution and suggests that *Mri⁻⁻Dna-pkcs^{-/-}* is synthetic lethal. Therefore, we can conclude that there is genetic interaction between Mri and Dna-pkcs in vivo.

Recent findings by our and other research groups suggest that MRI forms heterogeneous complexes involving PAXX or XLF, which function during DNA DSB repair by NHEJ [5]. Furthermore, genetic inactivation of Xlf [11], Paxx [4, 14–16], or Mri [5, 18] in mice leads to development of modest or no detectable phenotype. However, combined inactivation of Xlf and Mri [5] or Xlf and Paxx [4, 14, 15] results in embryonic lethality, which correlates with increased levels of neuronal apoptosis in the CNS (Figure 5). Here, we show that synthetic lethality produced by combined inactivation of Xlf and Mri can be rescued by altered Trp53 expression, similar to our previous Xlf⁻Paxx^{-/-} $Trp53^{+(-)/-1}$ [20] mouse model. Furthermore, we have developed and presented here Paxx^{-/-}Mri^{-/-} and Mri^{-/-} Dna-pkcs^{-/-} double deficient models.

Our findings have demonstrated that mice lacking XLF, MRI and p53, although live-born, possess a leaky SCID phenotype. $Xlf'^{-}Mri^{-}Trp53^{+/-}$ mice have a clear fraction of mature B cells in the spleens (CD19+) and bone marrow (B220+CD43-IgM+) (Figures 1 and 2), as well as clear fractions of double- and single-positive T cells in the thymus (CD4+CD8+, CD4+, CD8+) and singlepositive T cells in the spleen (CD4+ and CD8+) (Figure 1). However, the cell fractions from these mice are noticeably smaller than those of WT or single-deficient mice. Strikingly, we were able to identify one Xlf^{-/-}Mri^{-/-} $Trp53^{+/+}$ mouse at day P30 post-birth. This mouse resembled Xlf^{/-}Mri^{-/-}Trp53^{+/-} mice of similar age with respect of B and T cell development (Supplementary Table 10), although this mouse was generally sicker than its littermates and had to be euthanized. Similarly, one live-born Xlf' Paxx' mouse was reported by Balmus et al. 2016 [15], indicating that, exceptionally, embryonic lethality in NHEJ ligation-deficient mice can be overcome, likely due to activity of alternative endjoining. Previously, in 2018, Hung et al. [5] reported that combined inactivation of *Xlf* and *Mri* in vAbl pre-B cells results in a severe block in V(D)J recombination and accumulation of unrepaired DSBs in vitro, although it was unclear whether this combined inactivation would lead to a deficiency in B lymphocytes when translated to a mouse model [5]. Similarly, double deficient vAbl pre-B cells lacking Xlf and Paxx are also unable to sustain V(D)J recombination. Importantly, the lack of a progenitor T cell model system left the question of T cell development in Xlf⁻Mri⁻⁻ and Xlf⁻⁻ $Paxx^{-/-}$ mice completely unexplored.

Previously, we showed that mice lacking XLF, PAXX and p53 were live-born and had nearly no B and T cells, reduced size of spleen and hardly detectable thymus [20] (Figure 5). Consistent with this model, a conditional

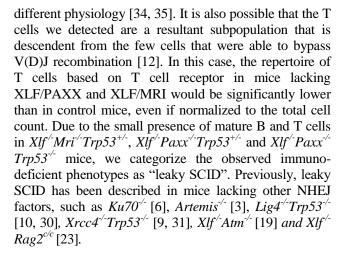
knockout mouse model, which results in doubledeficiency of XLF/PAXX in early hematopoietic progenitor cells, was also able to overcome the embryonic lethality of $Xlf^{-}Paxx^{-}$ mice [33]. With this model, impairment of V(D)J recombination in Xlf⁻Paxx^{-/-} cells, as well as the resulting depletion of mature B cells and lack of a visible thymus could also be observed in *vivo* [33]. Our new data provide evidence that Xlf⁻Paxx^{-/-} $Trp53^{+/-}$ and $Xlf^{/-}Paxx^{-/-}Trp53^{-/-}$ mice possess a very small number of mature B cells in the spleen and bone marrow, as well as very minor fractions of single positive T cells in thymus and spleen (Figures 2, 5 and Supplementary Figure 1). Therefore, both mature B and T cells are present in mice lacking XLF/PAXX and XLF/MRI. This can be explained by incomplete blockage in NHEJ and V(D)J recombination, in which the process is dramatically reduced but still possible. We also detected more mature T cells than B cells in these double-deficient mice. Potential explanations include longer lifespan of T cells, which accumulate over time following low efficiency of V(D)J recombination, while B cells are eliminated faster from the pool due to the

Α

	Offspring, P30	Expected, 1:2:1	*Expected, 1:2:0
Mri ^{-/-} Dna-pkcs⁺/+	12	6	8
Mri ^{-/-} Dna-pkcs ^{+/-}	12	12	16
Mri ^{-/-} Dna-pkcs ^{-/-}	0	6	0

В

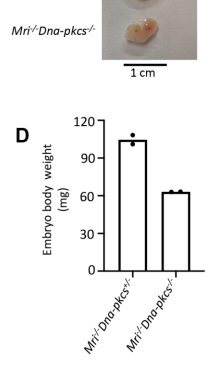
	Offspring, E14.5	Expected, 1:2:1	*Expected, 1:2:0
Mri ^{-/-} Dna-pkcs ^{+/+}	1	2.5	3.3
Mri ^{-/-} Dna-pkcs ^{+/-}	7	5.0	6.7
Mri ^{-/-} Dna-pkcs ^{-/-}	2	2.5	0.0



In addition to XLF/MRI and XLF/PAXX deficient mice, inactivation of one or two alleles of *Trp53* also rescues the embryonic lethality of $Xrcc4^{-/-}$ [9, 31], $Lig4^{-/-}$ [10, 30] and $Xlf^{-/-}Dna-pkcs^{-/-}$ [20] mice. We propose a model (Figure 5), when single deficiency for DNA-PKcs, PAXX or MRI results in no or modest

Mri^{-/-}

С



Embryos 14.5

Figure 4. Genetic interaction between *Mri* and *Dna-pkcs in vivo*. (A) No live-born $Mri^{-/-}Dna-pkcs^{-/-}$ mice were detected. (B, C) $Mri^{-/-}Dna-pkcs^{-/-}$ embryos were detected at day E14.5. (D) Body weight in milligrams (mg) from two E14.5 $Mri^{-/-}Dna-pkcs^{-/-}$ and $Mri^{-/-}Dna-pkcs^{+/-}$ embryos from the same litter. The mendelian ratio 1:2:1 in embryos was verified by the Chi-Square test (χ^2). The χ^2 was 1.8 and its corresponding probability was between 25 and 50%. *Expected distribution assuming lethality.

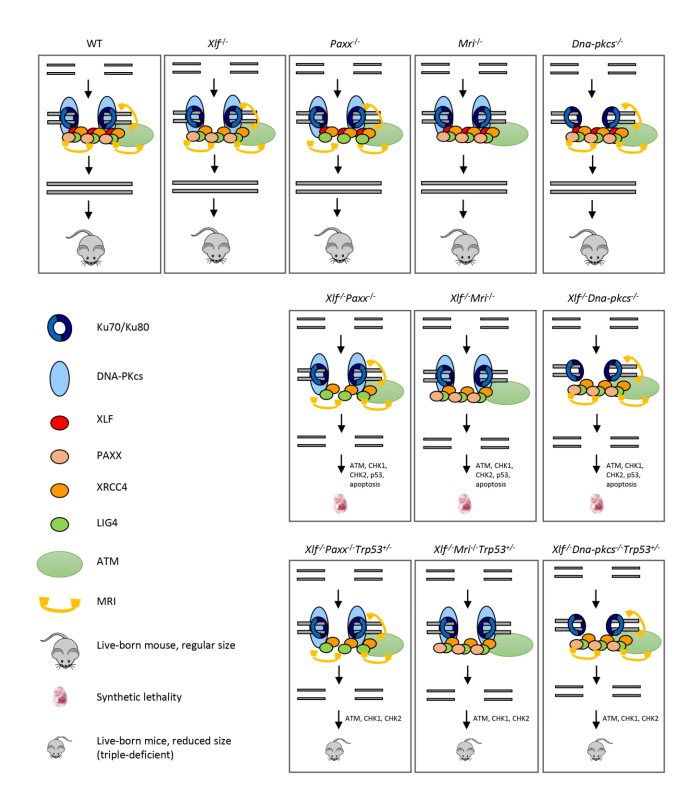


Figure 5. Mutations in NHEJ genes result in distinct phenotypes. Suggested models. Inactivation of *Paxx* or *Mri* results in live-born mice with nearly no DNA repair defects. Inactivation of *Xlf or Dna-pkcs* results in live-born mice with increased levels of genomic instability due to reduced NHEJ activity. Combined inactivation of *Xlf/Paxx*, *Xlf/Mri* or *Xlf/Dna-pkcs* leads to embryonic lethality in mice that correlate with high levels of genomic instability and nearly no NHEJ. Accumulated DSBs activate the ATM-dependent DNA damage response (DDR) pathway; ATM phosphorylates CHK checkpoint proteins that further trigger cell cycle arrest and apoptosis. Alternative end-joining is blocked by presence of Ku70/Ku80. Inactivation of one or two alleles of *Trp53* rescues embryonic lethality of *Xlf/Paxx*, *Xlf/Mri* and *Xlf/Dna-pkcs* mice. While in these mice the levels of DSBs are increased and ATM-dependent DDR response is activated, lack of p53 prevents massive apoptosis and thus results in alive mice. Sizes of the triple-deficient mice are reduced, as one option, due to DNA damage-dependent cell cycle arrest in multiple cells of the body. The embryonic lethality in mice lacking *Xlf/Paxx* and *Xlf/Mri* is likely to be rescued by inactivation of *Ku70* or *Ku80*.

phenotypes, and DSBs are efficiently repaired. Combined inactivation of Xlf/Dna-pkcs, Xlf/Paxx and results in inefficient DSB Xlf/Mri ligation, accumulation of DNA breaks, activation of ATMdependent DDR, checkpoint protein CHK2, stabilization of p53 and massive apoptosis. This results embryonic lethality in mice. Furthermore, in inactivation of Trp53 results in Xlf/Dna-pkcs/Trp53, Xlf/Paxx/Trp53 and Xlf/Mri/Trp53 triple-deficient mice. While DNA breaks in these mice are not repaired, ATM-dependent DDR response and activation of CHK proteins takes place. However, without p53, apoptosis is not activated, allowing survival of mice (Figure 5). Moreover, we propose that inactivation of Atm will also rescue embryonic lethality of Xlf/Paxx and Xlf/Mri mice due to the mechanisms proposed above. However, inactivation of Atm will not rescue embryonic lethality of Xlf/Dna-pkcs mice, due to synthetic lethality between Atm and Dna-pkcs.

It is important to note that altered *Trp53* expression is not always sufficient to rescue embryonic lethality in mice; for example, PLK1-interacting checkpoint helicase (PICH)-deficient mice possess developmental defects in the presence or absence of p53 [36], and ATR mutants (Seckel syndrome) are not completely rescued from embryonic lethality with the inactivation of Trp53 [37]. Embryonic lethality of XLF/PAXX and XLF/MRI double-deficient mice can be explained by the presence of Ku70/Ku80 heterodimer at the DSBs sites, which blocks DNA repair by alternative end-joining pathway(s), leading to massive apoptosis and cell cycle arrest [38]. Previously, it was shown that embryonic lethality of LIG4-deficient [39] and XLF/DNA-PKcs double-deficient mice [25] could be rescued by inactivating Ku70 or Ku80 genes. Similarly, we propose that inactivation of either Ku70 or Ku80 gene will rescue the embryonic lethality of XLF/PAXX and XLF/MRI double-deficient mice and will result in mice indistinguishable from Ku70- or Ku80-deficient controls (Figure 5).

Recent studies have shown that *Xlf* genetically interacts with *Rag2* [23] and DDR factors, such as *Atm*, *53bp1*, *H2ax*, and *Mdc1* [17, 19–22, 38]. *Xlf^{-/}Rag2^{c/c}* mice almost completely lack mature B cells and have significantly fewer mature T cells than single deficient controls [23]. *Xlf^{-/}Atm^{-/-}* and *Xlf^{/-53bp1^{-/-}* mice are liveborn and exhibit reduced body weight, increased genomic instability, and severe lymphocytopenia as a result of V(D)J recombination impairment in developing B and T cells [1, 17, 19, 22]. *Xlf^{/-}H2ax^{-/-}* and *Xlf^{/-}Mdc1^{-/-}*, on the other hand, are embryonic lethal [19–21]. There are several possible explanations for the functional redundancy observed between DNA repair genes. For instance, the two factors could have identical} (e.g., if both proteins are involved in ligation or DNA end tethering) or complementary (e.g., if one protein stimulates ligation while the other is required for DNA end tethering) functions. To date, XLF has been shown to genetically interact with multiple DNA repair factors [1, 4, 5, 14, 15, 19, 20, 24, 25], and this list is likely to grow [38, 40]. However, no clear genetic interaction has been shown between *Xlf* and *Artemis* or *Xrcc4* in the context of mouse development and V(D)J recombination [24], meaning that it remains difficult to predict genetic interactions without developing and characterizing genetic models.

We found that mice with combined inactivation of *Paxx* and *Mri* (*Paxx^{-/-}Mri^{-/-}*) are live-born, fertile, and undergo almost normal B and T cell development (Figure 3), where only the number of splenic B cells is affected, giving rise to a modest phenotype. Moreover, inactivation of *Paxx* did not affect the CSR efficiency in *in vitro* stimulated MRI-deficient B cells (Figure 3), thereby confirming our observations *in vitro*. It has been also shown that combined inactivation of *Paxx* and *Mri* genes in vAbl pre-B cells lead to similar V(D)J recombination efficiency to single-deficient *Mri^{-/-}*, *Paxx^{-/-}* and WT controls [5]. Thus, we conclude that there is a genetic interaction between *Paxx* and *Mri*, which results in a modest phenotype.

Lastly, we found that combined inactivation of Mri and Dna-pkcs ($Mri^{-/}Dna-pkcs^{-/-}$) leads to embryonic lethality, and that E14.5 $Mri^{-/}Dna-pkcs^{-/-}$ murine embryos were about 40% smaller than single-deficient siblings (Figure 4). DNA-PKcs is associated with the N-terminus of the MRI and Ku heterodimer in the process of recognizing DSBs [5], which may account for genetic interaction between Mri and Dna-pkcs. Thus, inactivation of Trp53, Ku70 or Ku80 may be a viable method to rescue synthetic lethality from $Mri^{-/}Dna-pkcs^{-/-}$ mice.

In conclusion, we have developed and described several complex genetic mouse models (Figure 5). $Xlf'^{-}Mri^{-/-}Trp53^{+/-}$ and $Xlf'^{-}Paxx^{-/-}Trp53^{+(-)/-}$ mice possessed severely impaired B and T lymphocyte development, leaky SCID; $Paxx^{-/-}Mri^{-/-}$ mice develop a modest B cell phenotype; and $Mri^{-/-}Dna-pkcs^{-/-}$ mice are embryonic lethal.

MATERIALS AND METHODS

Mice

All experiments involving mice were performed according to the protocols approved by the Comparative Medicine Core Facility (CoMed) at the Norwegian University of Science and Technology (NTNU, Trondheim, Norway). $Xlf^{+/-}$ [11] and $Dna-pkcs^{+/-}$ [2] mice were imported from the laboratory of Professor Frederick W. Alt at Harvard Medical School. $Trp53^{+/-}$ mice [32] were imported from Jackson Laboratories. $Paxx^{+/-}$ [16] and $Mri^{+/-}$ [18] mice were generated by the Oksenych group and described previously.

Lymphocyte development

Lymphocyte populations were analyzed by flow cytometry [16, 18, 19, 22]. In summary, cells were isolated from the spleen, thymus, and femur of 5-7week-old mice and treated with red blood cell lysis buffer Hybri-MaxTM (Sigma Aldrich, St. Louis, MO, USA; #R7757). The cells were resuspended in PBS (Thermo Scientific, Basingstoke, UK; #BR0014G) containing 5% Fetal bovine serum, FCS (Sigma Life Science, St. Louis, Missouri, United States; #F7524), and counted using a CountessTM II Automated Cell Counter (Invitrogen, Carlsbad, CA, United States; #A27977). Then, the cell suspension was diluted with PBS to get a final cell concentration of 2.5 x 10^7 cells/mL. Finally, surface markers were labeled with fluorochrome-conjugated antibodies and the cell populations were analyzed using flow cytometry.

Class switch recombination (CSR)

Spleens were isolated from 5-7-week-old mice and stored in cold PBS. Splenocytes were obtained by mincing the spleens, and naïve B cells were negatively selected using an EasySep Isolation kit (StemcellTM, Cambridge, UK; #19854). Lipopolysaccharide (LPS; 40 µg/mL; Sigma Aldrich, St. Louis, MO, USA; #437627-5MG) and interleukin 4 (IL-4; 20 ng/mL; PeproTech, Stockholm, Sweden; #214-14) were used to induce CSR to IgG1. Expression of IgG1 was analyzed by flow cytometry.

Antibodies

The following antibodies were used for flow cytometric analysis: rat anti-CD4-PE-Cy7 (BD PharmingenTM, Allschwil, Switzerland, #552775; 1:100); rat anti-CD8-PE-Cy5 (BD Pharmingen[™], Allschwil, Switzerland, #553034; 1:100); anti-CD19-PE-Cy7 (Biolegend, San Diego, CA, USA, #115520; 1:100); hamster anti-mouse PharmingenTM, anti-CD3-FITC (BD Allschwil, Switzerland, #561827; 1:100); rat anti-mouse anti-PharmingenTM, (BD CD43-FITC Allschwil, Switzerland, #561856; 1:100); rat anti-mouse anti-CD45R/B220-APC (BD PharmingenTM, Allschwil, Switzerland; #553092; 1:100); rat anti-mouse anti-IgM-PE-Cy7 (BD PharmingenTM, Allschwil, Switzerland, #552867; 1:100); rat anti-mouse IgG1-APC (BD PharmingenTM, Allschwil, Switzerland; #550874; 1:100). A LIVE/DEADTM fixable violet dead cell stain

kit (ThermoFisher Scientific, Waltham, MA, USA; #L34955; 1:1000) was used to identify dead cells.

Statistics

Statistical analyses were performed using one-way ANOVA, GraphPad Prism 8.0.1.244 (San Diego, CA, USA). In all statistical tests, p<0.05 were taken to be significant (*p<0.05; **p<0.01; ***p<0.001; ***p<0.001;

AUTHOR CONTRIBUTIONS

VO, SCZ, QZ, AL and MFB designed the study, analyzed and interpreted the results. SCZ, QZ, AL and MFB performed most of the experiments. VO wrote the paper with the help of SCZ and RY. All the authors contributed to writing of the final manuscript.

CONFLICTS OF INTEREST

The authors declare no conflicts of interest.

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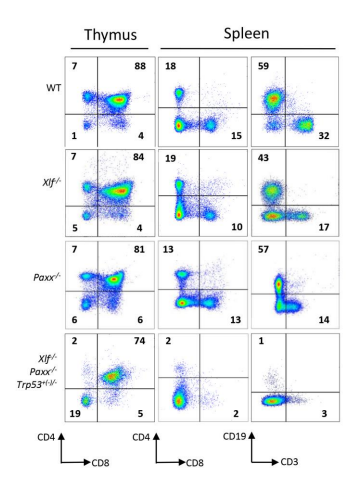
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SUPPLEMENTARY MATERIALS

Supplementary Figure



Supplementary Figure 1. B and T cell development in *Xlf^{/-}Paxx^{-/-}Trp53^{+(-)/-}* **mice.** Examples of flow cytometric analysis of thymic and splenic T cell subsets and splenic CD19+ B cells. *Xlf^{/-}Paxx^{-/-}Trp53^{+(-)/-}* is a combination of *Xlf^{/-}Paxx^{-/-}Trp53^{+/-}* and *Xlf^{/-}Paxx^{-/-}Trp53^{+/-}*.

Supplementary Tables

WT	Xlf'-	Mri ^{-/-}	Xlf ^{-/-} Mri ^{-/-} Trp53 ^{+/-}	Paxx ^{-/-}	Xlf ⁻ Paxx	-/- <i>Trp53</i> +(-)/-	– Paxx ^{-/-} Mri ^{-/-}	Dna-pkcs ^{-/-}
VV I	лу	MITI	лу мн прээ	1 илл	<i>Trp53</i> ^{+/-}	Trp53 ^{-/-}	- Faxx Mri	Dna-рксs
55.82	38.91	41.56	0.48	28.22	0.18	0.42	21.71	0.02
60.49	59.69	56.91	0.23	29.15	0.11	1.53	34.40	0.36
79.07	41.87	42.70	0.80	91.10	0.05	0.93	42.52	0.03
63.94	39.23	92.59	0.48	58.69	0.20		55.20	0.08
36.16	54.24	79.30	0.19	55.26	0.16		21.26	0.02
56.50	41.89	48.03		61.57	2.21		31.01	0.05
63.69	36.46	55.14		56.28	1.82		25.84	
42.85	37.87			39.59	0.58		15.51	
75.05	28.68			55.36	0.08		32.73	
67.60	39.08			61.29			58.43	
38.27	29.73			80.28			64.75	
79.47	47.11			61.29			36.96	
29.43	56.65			93.23			37.92	
52.58	44.10			61.86			63.63	
65.47	34.65						31.99	
62.80	30.79						22.31	
56.68	34.55						28.62	
57.03	50.48							
54.05	52.23							
75.79	36.41							

Supplementary Table 1. Summary of splenic CD19+ B cells.

CD19+ splenocytes (×10⁶) in WT, Xlf^{/-}, Mri^{-/-}, Xlf^{/-}Mri^{-/-}Trp53^{+/-}, Paxx^{-/-}, Xlf^{/-}Paxx^{-/-}Trp53^{+(-)/-} and Paxx^{-/-}Mri^{-/-} mice. Dna-pkcs^{-/-} mice were used as an immunodeficient control. Xlf^{/-}Paxx^{-/-}Trp53^{+(-)/-} is a combination of Xlf^{/-}Paxx^{-/-}Trp53^{+/-} and Xlf^{/-}Paxx^{-/-}Trp53^{-/-}.

	VIC/-	N/	Xlf Mri ^{-/-} Trp53 ^{+/-}	D	Xlf ^{/-} Paxx ^{-/-}	Trp53 ^{+(-)/-}	D	D
WT	Xlf'-	Mri ^{-/-}	Xlf Mri Trp53	<i>Paxx</i> ^{-/-} –	<i>Trp53</i> ^{+/-}	<i>Trp53^{-/-}</i>	- Paxx ^{-/-} Mrï ^{/-}	Dna-pkcs ^{-/-}
41.61	15.06	23.01	5.55	34.34	0.13	0.41	26.93	0.38
37.93	36.01	36.61	3.31	21.89	0.52	1.42	24.53	0.23
42.64	33.33	23.12	3.05	38.84	0.42	0.84	36.66	0.03
29.44	39.67	32.84	2.36	23.51	0.26		41.18	0.09
15.25	30.11	30.74	1.55	31.11	0.92		15.22	0.02
22.62	15.19	19.45		16.26	0.25		18.83	0.07
20.41	49.15	25.38		14.86	0.71		20.21	
18.50	24.24			10.45	0.38		13.65	
27.19	17.04			14.62			17.85	
24.49	14.83			16.18			25.19	
13.86	15.41			21.20			28.55	
28.79	11.67			16.18			20.70	
19.05	15.90			24.62			16.90	
23.72	12.10			16.34			32.74	
22.75	19.17						17.44	
20.53	23.05						19.22	
27.01	17.94						18.25	
25.60	14.10							
35.90	12.52							
	14.06							
	20.54							
	21.25							
	42.82							
	14.81							

Supplementary Table 2. Summary of splenic CD3+ T cells.

CD3+ splenocytes (×10⁶) in WT, Xlf^{-/-}, Mri^{-/-}, Xlf^{/-}Mri^{-/-}Trp53^{+/-}, Paxx^{-/-}, Xlf^{/-}Paxx^{-/-}Trp53^{+(-)/-} and Paxx^{-/-}Mri^{-/-} mice. Dna-pkcs^{-/-} mice were used as an immunodeficient control. Xlf^{/-}Paxx^{-/-}Trp53^{+(-)/-} is a combination of Xlf^{/-}Paxx^{-/-}Trp53^{+/-} and Xlf^{/-}Paxx^{-/-}Trp53^{+/-}.

WT	Xlf'-	Mri [≁]	Xlf ^{/-} Mri ^{-/-} Trp53 ^{+/-}	Paxx ^{-/-} -	Xlf' ⁻ Paxx ^{-/-} Trp53 ^{+(-)/-}		- Paxx ^{-/-} Mri ^{-/-}	Dna-pkcs ^{-/-}
VV I	ЛIJ		Alj Miri Trp55	Paxx -	<i>Trp53</i> ^{+/-}	<i>Trp53^{-/-}</i>	- Paxx Mri	Dna-pkcs
19.51	5.46	10.36	1.14	12.17	0.03	0.49	8.3	0.08
17.39	11.43	15.61	1.66	9.96	0.01	0.34	12.54	0.05
18.17	18.99	13.88	1.09	24.65	0.49	0.56	17.64	0.09
9.23	15.66	19.32	0.96	15.97	0.59		18.51	
13.63	14.9	17.57		19.38	0.35		6.66	
12.38	6.88	13.25			0.37		8.13	
11.89	21.81	14.42			0.37		8.92	
9.844					0.27		5.51	
8.961							9.67	
9.96							15.61	
15.85							18.01	
19.29							10.36	
16.42								
12.85								

Supplementary Table 3. Summary of splenic CD4+ T cells.

CD4+ splenocytes (×10⁶) in WT, Xlf^{/-}, Mri^{-/-}, Xlf^{/-}Mri^{-/-}Trp53^{+/-}, Paxx^{-/-}, Xlf^{/-}Paxx^{-/-}Trp53^{+(-)/-} and Paxx^{-/-}Mri^{-/-} mice. Dna-pkcs^{-/-} mice were used as an immunodeficient control. Xlf^{/-}Paxx^{-/-}Trp53^{+(-)/-} is a combination of Xlf^{/-}Paxx^{-/-}Trp53^{+/-} and Xlf^{/-}Paxx^{-/-}Trp53^{-/-}.

WT	V1¢/-	Mri ^{-/-}	Xlf Mrī Trp53 ^{+/-}	Paxx ^{-/-}	Xlf ^{/-} Paxx ⁻	^{-/-} Trp53 ^{+(-)/-}	- Paxx ^{-/-} Mri ^{-/-}	D
WT	Xlf ^{/-}	Miri	лу мн трээ	Γαλλ	<i>Trp53</i> ^{+/-}	<i>Trp53^{-/-}</i>	- Paxx Mri	Dna-pkcs ^{-/-}
14.39	4.03	12.59	0.74	6.83	0.32	0.40	5.86	0.08
18.35	12.06	14.79	1.72	9.30	0.07	0.41	10.45	0.05
12.31	12.73	10.57	1.19	15.76	0.22	0.29	15.26	0.06
7.13	13.11	18.08	0.66	12.05	0.24		16.02	
9.70	9.67	15.62		14.11	0.02		5.60	
9.64	6.39	9.96			0.39		7.69	
7.96	14.92	12.09			0.20		7.16	
14.26							4.86	
12.96							7.28	
13.91							13.26	
11.62							14.87	
12.17							7.78	
10.88								
7.88								

Supplementary Table 4. Summary of splenic CD8+ T cells.

CD8+ splenocytes (×10⁶) in WT, Xlf^{-/-}, Mri^{-/-}, Xlf^{/-}Mri^{-/-}Trp53^{+/-}, Paxx^{-/-}, Xlf^{/-}Paxx^{-/-}Trp53^{+(-)/-} and Paxx^{-/-}Mri^{-/-} mice. Dna-pkcs^{-/-} mice were used as an immunodeficient control. Xlf^{/-}Paxx^{-/-}Trp53^{+(-)/-} is a combination of Xlf^{/-}Paxx^{-/-}Trp53^{+/-} and Xlf^{/-}Paxx^{-/-}Trp53^{-/-}.

W/T	VIC/-	Xlf'- Mri ^{-/-}	Xlf ^{/-} Mri ^{-/-} Trp53 ^{+/-}	Paxx ^{-/-}	Xlf ^{/-} Paxx ⁻	/- <i>Trp53</i> +(-)/-	- Paxx ^{-/-} Mri ^{-/-}	Dna-pkcs ^{-/-}
WT	ХIJ			Paxx	<i>Trp53</i> ^{+/-}	Trp53 ^{-/-}	- Paxx Mri	Dna-pkcs
9.88	8.45	8.63	1.32	6.01	0.66	0.13	5.53	0.02
9.06	3.41	10.16	0.48	8.77	0.07	0.06	8.40	0.001
10.48	11.88	6.74	0.89	12.88	0.07	0.19	4.31	0.01
16.33	7.05	7.95	0.65	11.10	0.11		6.77	0.02
7.50	6.67	15.23	0.50	9.80	0.40		6.85	
10.64	11.66	17.57			0.29		13.08	
7.12		15.94			0.19		9.52	
4.33		10.99			0.13		6.86	
2.55		15.88					12.67	
11.74		9.21					12.97	
13.15								
14.54								
14.41								
12.07								
11.39								
9.70								

Supplementary Table 5. Summary of thymic CD4+ T cells.

CD4+ thymocytes (×10⁶) in WT, Xlf^{/-}, Mri^{-/-}, Xlf^{/-}Mri^{-/-}Trp53^{+/-}, Paxx^{-/-}, Xlf^{/-}Paxx^{-/-}Trp53^{+(-)/-} and Paxx^{-/-}Mri^{-/-} mice. Dna-pkcs^{-/-} mice were used as an immunodeficient control. Xlf^{/-}Paxx^{-/-}Trp53^{+(-)/-} is a combination of Xlf^{/-}Paxx^{-/-}Trp53^{+/-} and Xlf^{/-}Paxx^{-/-}Trp53^{+/-}.

WT	Xlf'-	Mri ^{-/-}	Xlf ^{/-} Mri ^{-/-} Trp53 ^{+/-}	Paxx ^{-/-}	Xlf ^{/-} Paxx ⁻	^{-/-} Trp53 ^{+(-)/-}	Paxx ^{-/-} Mri ^{-/-}	Dna-pkcs ^{-/-}
WT	ХIJ	Mri	Alj Miri Trp55	Paxx	<i>Trp53</i> ^{+/-}	<i>Trp53^{-/-}</i>	Paxx Mri	Dna-pkcs
2.70	2.39	3.48	0.64	4.34	0.18	0.19	1.08	0.02
2.00	1.83	3.43	0.33	2.93	0.06	0.09	1.85	0.001
3.84	3.40	1.82	0.91	5.51	0.28	0.21	1.33	0.01
4.69	2.60	2.72	0.29	4.52	0.51		2.24	0.01
1.47	2.60	7.14	0.67	4.7	0.15		3.07	
2.11	3.59	6.40			0.23		6.2	
5.07		5.45			0.60		4.03	
3.26		4.28			0.49		3.16	
1.98							5.14	
3.91							5.11	
13.59								
11.68								
13.78								
3.47								
5.36								
4.90								
2.25								
4.56								

Supplementary Table 6. Summary of thymic CD8+ T cells.

CD8+ thymocytes (×10⁶) in WT, Xlf^{/-}, Mri^{-/-}, Xlf^{/-}Mri^{-/-}Trp53^{+/-}, Paxx^{-/-}, Xlf^{/-}Paxx^{-/-}Trp53^{+(-)/-} and Paxx^{-/-}Mri^{-/-} mice. Dna-pkcs^{-/-} mice were used as an immunodeficient control. Xlf^{/-}Paxx^{-/-}Trp53^{+(-)/-} is a combination of Xlf^{/-}Paxx^{-/-}Trp53^{+/-} and Xlf^{/-}Paxx^{-/-}Trp53^{+/-}.

Supplementary Table 7. Summary of thymic CD4+CD8+ double positive T cells.

W / D	v 1c/-	Mri ^{-/-}	Xlf Mri ^{-/-} Trp53 ^{+/-}	Paxx ^{-/-}	Xlf ^{/-} Paxx	-/-Trp53+(-)/-	D	D
WT	Xlf'-	Mri	Alf Mrt Trp55	Paxx	<i>Trp53</i> ^{+/-}	<i>Trp53^{-/-}</i>	Paxx ^{-/-} Mri ^{-/-}	Dna-pkcs ^{-/-}
154.05	48.40	160.60	20.69	132.56	2.68	6.22	75.37	0.17
141.52	73.77	133.62	11.40	161.22	7.21	6.54	184.27	0.002
230.74	163.07	14.22	21.18	151.30	6.79	5.14	122.11	0.0002
147.74	95.47	165.78	17.78	208.39	3.39		105.44	0.001
138.62	115.71	154.74	17.71	202.99	3.72		88.37	0.002
98.78	115.77	193.15		161.06	4.43		168.00	
115.10	174.36	102.72		171.03	11.90		122.32	
66.71	144.88			100.87	5.18		87.20	
102.13	160.88			175.18			136.82	
162.51	105.29			197.59			153.47	
126.04	155.83						114.36	
79.43	90.46						136.39	
140.22	118.17						55.65	
146.71	183.46						96.22	
119.15	158.35							
	172.50							
	146.98							
	101.50							
	143.61							
	114.38							
	132.49							
	105.96							
	136.45							
	162.14							

CD4+CD8+ thymocytes (×10⁶) in WT, Xlf^{-/-}, Mri^{-/-}, Xlf^{-/-}Mri^{-/-}Trp53^{+/-}, Paxx^{-/-}, Xlf^{-/-}Paxx^{-/-}Trp53^{+(-)/-} and Paxx^{-/-}Mri^{-/-} mice. Dnapkcs^{-/-} mice were used as an immunodeficient control. Xlf^{-/-}Paxx^{-/-}Trp53^{+(-)/-} is a combination of Xlf^{-/-}Paxx^{-/-}Trp53^{+/-} and Xlf^{-/-}Paxx^{-/-}Trp53^{-/-}.

WT	V16/-	lf'- Mrī'-	Xlf ^{/-} Mri ^{-/-} Trp53 ^{+/-}	Paxx ^{-/-}	Xlf ^{/-} Paxx ^{-/}	⁻ <i>Trp53</i> ^{+(-)/-}	Paxx ^{-/-} Mri ^{-/-}
VV I	ЛIJ		Alf Mrt Trp55	Paxx	<i>Trp53</i> ^{+/-}	<i>Trp53^{-/-}</i>	Paxx Mri
19.80	7.82	17.10	3.04	17.8	2.19	3.26	15.90
16.70	11.00	16.30	1.47	14.3	2.53	4.01	16.70
18.60	8.47	16.00	1.42	15.9	4.09	3.55	14.30
10.10	6.04	15.80	1.10	11.9	0.37		14.00
14.10	7.68	13.80	1.57		4.60		10.10
11.60	6.06	11.50			4.35		9.82
12.40	12.40	14.90			1.50		7.34
13.90	10.60	13.00			3.76		8.22
14.10	5.79				5.19		14.60
14.50					6.20		14.10
13.70							
10.10							
12.10							

Supplementary Table 8. Summary of IgM+ B cells in bone marrow.

Frequencies (%) of B220+CD43-IgM+ B cells in WT, Xlf^{/-}, Mri^{-/-}, Xlf^{/-}Mri^{-/-}Trp53^{+/-}, Paxx^{-/-}, Xlf^{-/-}Paxx^{-/-}Trp53^{+(-)/-} and Paxx^{-/-}Mri^{-/-} mice. Xlf^{/-}Paxx^{-/-}Trp53^{+/-} is a combination of Xlf^{/-}Paxx^{-/-}Trp53^{+/-} and Xlf^{/-}Paxx^{-/-}Trp53^{-/-}. W

Supplementary Table 9. Summary of progenitor B cells in bone marrow.

WT	Xlf'-	Mri⁻∕-	Xlf ^{/-} Mri ^{-/-} Trp53 ^{+/-}	Paxx ^{-/-} -	Xlf ^{/-} Paxx ^{-/}	Trp53 ^{+(-)/-}	– Paxx ^{-/-} Mri ^{-/-}
VV I	ЛIJ		лу мн прээ	Paxx -	<i>Trp53</i> ^{+/-}	<i>Trp53^{-/-}</i>	- Paxx Mri
6.00	21.6	3.74	24.90	8.92	33.70	20.10	7.53
7.47	8.73	4.31	14.50	4.40	28.40	24.00	7.04
3.38	17.5	9.53	17.20	4.07	23.20	25.60	8.64
6.96	10.9	7.03	14.90	3.81	21.90		10.9
5.02	9.61	6.73	19.00		26.80		9.00
5.49	7.99	6.58			27.10		6.28
2.75	5.31	4.17			25.10		6.76
4.25	7.16	4.28			17.30		6.20
7.58	6.08				22.20		3.32
7.47					25.50		4.06
5.03							
7.72							
8.29							

Frequencies (%) of B220+CD43+IgM- pro-B cells in WT, Xlf^{/-}, Mri^{-/-}, Xlf^{/-}Mri^{-/-}Trp53^{+/-}, Paxx^{-/-}, Xlf^{/-}Paxx^{-/-}Trp53^{+(-)/-} and Paxx^{-/-} Mri^{-/-} mice. Xlf^{/-}Paxx^{-/-}Trp53^{+(-)/-} is a combination of Xlf^{/-}Paxx^{-/-}Trp53^{+/-} and Xlf^{/-}Paxx^{-/-}Trp53^{+/-}.

Supplementary Table 10. Lymphocytic development in the *Xlf^{/-}Mri^{/-}Trp53^{+/+}* mouse.

Splenocytes (×10 ⁶)				Thymocytes (×10 ⁶)			Cell populations (%) in bone marrow	
CD19+ B cells	CD3+ T cells	CD4+ T cells	CD8+ T cells	CD4+ T cells	CD8+ T cells	CD4+CD8+ T cells	IgM+ B cells	Pro-B cells
0.11	0.80	0.53	0.41	0.42	0.41	15.70	2.71	19.40

Summary of splenic (×10⁶) B- and T cells; and T cell subpopulations in the thymus (×10⁶). Frequencies (%) in bone marrow of B220+CD43-IgM+ B cells and B220+CD43+IgM- pro-B cells.