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New role for the (pro)renin receptor in T-cell development

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Key Points

- PRR deletion in T cells drastically reduces the number of peripheral and thymic CD3+ T cells.
- We identify multiple stages of thymocyte development that require PRR expression.

Introduction

The (pro)renin receptor (PRR) was originally thought to be important for regulating blood pressure via the renin-angiotensin system. However, it is now emerging that PRR has instead a generic role in cellular development. Here, we have specifically deleted PRR from T cells. T-cell-specific PRR-knockout mice had a significant decrease in thymic cellularity, corresponding with a 100-fold decrease in the number of CD4+ and CD8+ thymocytes, and a large increase in double-negative (DN) precursors. Gene expression analysis on sorted DN3 thymocytes indicated that PRR-deficient thymocytes have perturbations in key cellular pathways essential at the DN3 stage, including transcription and translation. Further characterization of DN T-cell progenitors leads us to propose that PRR deletion affects thymocyte survival and development at multiple stages; from DN3 through to DN4, double-positive, and single-positive CD4 and CD8. Our study thus identifies a new role for PRR in T-cell development. (Blood. 2015;126(4):504-507)

Results and discussion

The cellularity of peripheral lymphoid organs and blood were analyzed by flow cytometry. In PRR cKO mice, a significant decrease in CD3+ T cells was observed in all tissues, with no difference in the number of B, macrophage, or dendritic cells (Figure 1A; supplemental Figures 2-3).
A striking reduction in both CD4<sup>+</sup> and CD8<sup>+</sup> naïve T cells was evident, with no change in the number of peripheral T-cell receptor (TCR) γδ cells observed (Figure 1B). As PRR is found to be essential for homeostasis<sup>7-11</sup> we asked if the remaining peripheral T cells had the PRR gene excised. We performed polymerase chain reaction (PCR) that specifically amplified a segment of the PRR gene only after it was mutated by Cre recombination (supplemental Figure 4). Using this approach, we could detect the recombined gene (ie, with the PRR gene excised) in thymic but not peripheral CD90.2<sup>+</sup>T cells, which expressed similar levels of PRR protein as control T cells (supplemental Figure 4). Together, these results indicate that peripheral cKO T cells simply do not develop or survive. Instead, those few cells that do not recombine the PRR gene have an enormous selective advantage and reach the periphery.

The striking reduction in peripheral naïve T cells is consistent with a defect in thymic T-cell development, so we turned our focus to this organ. A severe atrophy of PRR cKO thymi was observed, evidenced by a reduction in weight and cellularity (Figure 1C). PRR cKO thymi had a 10-fold increase in the proportion of double-negative (DN) thymocytes (Figure 1D). A significant increase in the proportion of intermediate single-positive (ISP) and TCRγδ cells was observed, whereas all other populations were decreased (Figure 1E). The number of DN cells was unchanged, whereas the numbers of ISP, DP, CD8<sup>+</sup> SP, CD4<sup>+</sup> SP, and regulatory T cells were reduced (Figure 1E). Histologic analysis indicated that the loss of SP T cells was associated with a reduction in the thymic medulla of cKO mice but preserved corticomedullary junction (supplemental Figure 5), consistent with other models of severe SP T-cell loss.<sup>20</sup> No change in the number of TCRγδ cells was evident. This is unsurprising as the lineage commitment for TCRγδ cells begins at the DN2 stage, prior to Lck-driven Cre recombinase expression.<sup>16</sup> Taken together, the almost 100-fold reduction of peripheral naïve T cells is consistent with a defect in thymic T-cell development.
Figure 2. PRR cKO impairs thymocyte survival and development from DN3 and beyond. (A) Representative flow cytometry of CD25 and CD44 expression in DN (CD4^+ CD8^-) thymocytes from 39-day-old mice.

(B) The frequency and actual number of DN3 and DN4 cells from panel A. (C) TCRβ gene rearrangement by PCR of sorted DN4 cells from control and cKO mice. DNA from bone marrow (BM) cells is shown as a nonrearranged control. (D) Expression of intracellular TCRβ by PCR of sorted DN4 thymocytes as from panel B. (E) Scatter plot showing the comparison of gene expression from microarray analysis of DN3 sorted cells from control and cKO mice. Significantly different genes (by 1-way analysis of variance [ANOVA]; \(q < 0.05\)) are labeled (black circles). For clarity, the gene 5730437N04Rik is abbreviated as “5…Rik,” and the gene 1600029D21Rik is abbreviated as “1…Rik.” Inset shows PRR expression by real-time quantitative PCR. N = 3 biological replicates. (F) GO pathways that were significantly enriched in cKO DN3 cells (false-discovery rate [FDR] < 0.5) are shown. Details of the genes in each pathway are listed in supplemental Table 1. (G) Representative PCR from sorted thymocytes from 39-day-old control and cKO mice to detect the unexcised PRR gene (“floxed”) and mutated/deleted gene in cKO (“excised”). Cre recombinase expression is also shown. (H) TCRα gene rearrangement was determined in sorted DP cells from control and cKO mice by quantitative PCR with primers specific for V\(\alpha\)8, V\(\alpha\)2, and V\(\alpha\)10 in conjunction with different J\(\alpha\) primers. The \(P\) value shown for the effect of PRR deletion was calculated by 2-way ANOVA. \#\(P < .01\) by Sidak’s multiple comparison test. N = 3-4. (I) Thymocytes from control and cKO mice were cultured in vitro with interleukin 7, and the proportion of live cells was determined by flow cytometry at the desired time points. The \(P\) value shown for the effect of PRR deletion was calculated by 2-way ANOVA. **\(P < .01\); ****\(P < .0001\) by Sidak’s multiple comparison test. N = 5-9. (J) Schematic of T-cell development stages affected by PRR deletion.
reduction in SP cells indicates that PRR cKO mice have drastically altered thymocyte development.

We next analyzed early T-cell developmental events (DN1-4). There was an increase in the proportion of DN3 cells in cKO mice, which was correlated with a decreased proportion of DN4 (Figure 2A). These proportional changes were not associated with differences in the number of DN3 or DN4 (Figure 2B). At the DN3 stage, cells commit to the T-cell lineage and rearrange the TCR β-chain gene locus. PCR products corresponding to rearranged TCRβ loci were detected in both control and cKO cells, indicating that this process was not disturbed (Figure 2C). There was no difference in intracellular TCRβ levels (Figure 2D).

We next performed gene expression analysis of sorted DN3 cells (Figure 2E; supplemental Table 1A). In cKO, 615 and 518 genes were up- and downregulated >1.5-fold. Of these, 9 genes were significantly different and encoded molecules important for processes including the cell cycle (Mctsl, Cenm2), mitochondria (Tom7, Uqcrh) and vesicular trafficking (Tcr3ipl, Rabac1). We next performed gene ontology (GO) analysis on the differentially expressed genes (Figure 2F; supplemental Table 1B). Several GO pathways were enriched in cKO DN3 cells, indicating disturbances in transcription, translation, and the mitochondria. Collectively, these represent cellular activities essential for this developmental period, providing an explanation for why PRR deletion has such a profound effect on T-cell development.

Comparing our findings to conditional deletion of the Wnt pathway in T cells, several similarities are evident: reduction in peripheral T cells and a change in the proportion of DN3:DN4.16 However, overall it is clear that PRR deletion induces a more profound phenotype. Wnt cKO led to an increase in the proportion of DN cells to 12%, and a partial block at the DN3-DN4 transition, leading to an increased number of DN3 cells and decreased DN4. In the periphery, there was a fourfold reduction in T cells, which did have the β-catenin deleted allele.16 In contrast, PRR cKO leads to an increase in the proportion of DN cells to ~50% of thymocytes, we detect no increase in the number of DN3 cells, and we find that DN4 numbers are decreased only in old mice (data not shown). Furthermore, we are unable to detect the deleted allele in peripheral T cells, indicating that PRR-deficient T cells do not develop/survive. To understand these results further, we sorted thymocytes and again performed PCR to specifically identify cKO cells. This revealed the presence of cKO cells beyond the DN3 stage, including DN4, DP, and CD4 SP (Figure 2G). We analyzed DP cells further and observed an impaired survival and, subsequently, rearrangement of the TCR α chain (Tcra) (Figure 2H-I). Thus, given the striking severity of our phenotype, that no single precursor thymocyte population accumulates, and that we also observe defects in DP cells, we propose that deletion of PRR affects thymocyte survival and development from DN3 and beyond (Figure 2J).

Acknowledgments

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Authorship


Conflict-of-interest disclosure: The authors declare no competing financial interests.

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References

ONLINE SUPPLEMENTAL DATA

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Contains:
Supplemental Figures 1 - 6
Full experimental procedures
Supplemental references
Supplemental Figure 1. Schematic of study.

A) Female mice carrying the loxP-flanked target allele encoding PRR were bred to Lck-cre transgenic male mice. As PRR is located on the X chromosome, first-generation knockout male offspring with PRR specific deleted in T cells (ATP6AP2flox/y;Lck-Cre).

B) Body weight between control and cKO mice was unchanged. N=20 (control); 9 (cKO).
Fig S2

A

Lymph node

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Gating strategy:

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**Supplemental Figure 2. Cellularity of peripheral lymph and circulating lymphocytes.**

A) The cellularity of peripheral lymphoid organs (spleen and pooled lymph nodes) and PBMC’s was determined in control and cKO mice. A significant decrease in the cellularity of lymph nodes was observed.

B) Gating strategy for flow cytometry analysis of the proportion of T, B, macrophages and dendritic cells (Fig 1A). Single celled lymphocytes were first gated as CD45⁺, and then for CD3, CD19, CD11c and F4/80 expression.

C) The actual number of T cells (CD3⁺), B cells (CD19⁺), macrophages (CD3⁻CD19⁻CD11c⁻ F4/80⁺) and dendritic cells (CD3⁻CD19⁻CD11c⁺) was determined in lymph nodes and PBMC’s from 7-8 week old control and cKO mice.
Supplemental Figure 3. Reduction in peripheral T cells.

A) Representative CD4 and CD8 expression in spleens of 6-8 week old control and cKO mice by flow cytometry.

B) The frequency (top row) and actual number (bottom row) of cell populations from B.

TCR γ8 numbers are presented in Fig 1B.
Supplemental Figure 4. Remaining peripheral T cells in cKO mice express normal levels of PRR.

A) Schematic for PCR’s to detect the unexcised PRR gene (“floxed”) and mutated/deleted gene in cKO (“excised”).

B) From the same mouse, thymocytes (sorted DN4 cells) and peripheral T cells (CD90.2^+ MACS-enriched splenic T cells) were subjected to the PCR as in A. Six biological replicates are shown, where “Cre” and “Flox” corresponds to control mice, and “cKO” indicates PRR cKO mice.

C) PRR expression in sorted peripheral (pooled spleen and lymph node) CD90.2^+ T cells. Loading control eIF4E is shown.
Supplemental Figure 5. Histological analysis of control and cKO thymi. Thymic architecture of control and cKO mice was determined by staining thymi with haeomotoxylin and eosin. A reduction in thymic medulla of cKO mice is observed; a characteristic secondary effect due to a lack of epithelial cell interaction with single positive T cells. Scale bars, 200 μm
Supplemental Figure 6. Microarray quality control and ANOVA

A) Hierarchical clustering of averages of biological replicates (NIA array tool). N=3 control, N=3 cKO.

B) Differential gene expression between control and cKO samples, genes plotted as log-ratio versus log-intensity. Genes which significantly differentially regulated were identified by ANOVA (FDR<0.0.5). Genes that were significantly down-regulated are in green, up-regulated in red.
**Full experimental procedures**

**Mice.**

We generated mice with a conditional deletion of PRR from T cells by breeding female C57Bl/6 mice with loxP sites flanking exon 2 of the ATP6AP2 genes with male mice expressing Cre recombinase under the distal Lck promoter (founder line 3779)\(^1\). In this model, Cre is expressed specifically in early T cell development, beginning at the DN3 stage (**Fig S1A**). It has previously been reported that in this model no Cre is expressed at DN2, ~20-40% at DN3, and 90-100% at the DP stage\(^2\). As gamma-delta T cell receptor recombination begins at the DN2 stage of T cell development, alpha-beta T cell development is predominantly affected in this Lck-Cre model. Also, as PRR is located on the X chromosome, first-generation male offspring with PRR conditionally deleted in T cells were generated (PRR\(^{\text{flox/y}}\);Lck-Cre\(^{+/−}\)) and are henceforth referred to as “cKO”. Littermate control males (PRR\(^{\text{flox/y}}\) or Lck-Cre\(^{+/−}\)) were pooled. Animal experiments were approved by the local authorities (LaGeSo, Berlin, Germany). Mice had normal weight (**Fig S1B**) and appeared healthy. They were housed under standard SPF-conditions and monitored daily to minimize harm.

**Preparation of single-cell suspensions for flow cytometry.**

Spleens and lymph nodes (two axial and two inguinal, pooled) were dissected from 7-8 week-old control and cKO mice. Thymuses were from 6 week-old mice, unless indicated otherwise. Organs were disrupted with forceps and scissors, and then passed through a 70 µM cell strainer (BD Bioscience) to generate single-cell suspensions. Peripheral blood mononuclear cells (PBMC’s) were isolated by centrifugation through a Ficoll cushion (Ficoll-Paque PLUS, GE Healthcare). All samples were kept on ice. Red blood cells were removed from all single-cell suspensions by osmotic lysis. Lymphocytes were washed into FACS buffer (PBS with 0.5% (v/v) fetal calf serum (FCS) (S0615, Biochrom), and 2 mM EDTA). Cells were stained with trypan blue and the number of live cells was counted using a haemocytometer. For intracellular staining, cells
were first permeabilized using the CytoFix/CytoPerm Solution Kit (BD Bioscience), according to the manufacturer’s protocol. For the staining of extracellular surface molecules, cells were resuspended into FACS buffer with the desired fluorescently labelled antibodies. Antibodies used were: CD3-APC-Cy7 (BD Biosciences, 557596), CD3-APC-eFluor660 (eBiosciences, 50-0032-82), CD4-FITC (BD Biosciences, 553729), CD4-PacificBlue (BD Biosciences 558107), CD8-PerCP-Cy5.5 (eBiosciences, 45-0081-82), CD8-eFluor450 (eBiosciences, 48-0081-82), CD8-PE (BD Biosciences, 553032), CD11c-Alexa647 (own polyclonal), CD19-PE (BD Biosciences, 12-0193-81), CD25-PE (BD Biosciences 553866), CD44-APC (eBiosciences, 17-0441-81), CD44-FITC (BD Biosciences, 561859), CD45-FITC (BD Biosciences, 553079), CD62L-APC (BD Biosciences, 561919), CD90.2-APC (BD Biosciences, 553037), F4/80-PacificBlue (eBiosciences, 48-4811), FOXP3-PerCP-Cy5.5 (eBiosciences, 75-5775-80), TCRβ (eBiosciences, 48-5961-82), TCRγδ-PE (BD Biosciences, 55378). After staining, cells were analysed on a BD FACS Canto II (BD Biosciences), and data was analyzed with the FlowJo software version 10 (Tree Star). Cell doublets were excluded by analysis of FSC-H vs FSC-A. Cell populations are expressed as a percentage of the respective organ, and the absolute number was also determined by multiplying by the total cell number of each organ. Cell populations were defined by the following gating strategies:

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<td>TCRγδ</td>
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Naïve (CD4⁺ or CD8⁺) | CD44⁺ CD62L⁺
---|---
**Thymus**
CD4⁺ single positive | CD4⁺ CD8⁻ CD25⁻ CD44⁺
CD8⁻ single positive | CD4⁻ CD8⁺ CD25⁻ CD44⁺
T regulatory | CD3⁺ CD4⁺ CD25⁻ icFOXP3⁺
TCRγδ | CD3⁺ CD4⁻ CD8⁻ TCRγδ⁺
Double positive (DP) | CD4⁺ CD8⁺ CD25⁻ CD44⁺
Double negative (DN) | CD4⁻ CD8⁻ CD25⁻ CD44⁺
Intermediate single positive (ISP) | CD4⁻ CD8⁻ CD25⁻ CD44⁺
DN1 | CD4⁻ CD8⁻ CD25⁻ CD44⁺
DN2 | CD4⁻ CD8⁻ CD25⁻ CD44⁺
DN3 | CD4⁻ CD8⁻ CD25⁻ CD44⁺
DN4 | CD4⁻ CD8⁻ CD25⁻ CD44⁺

*Enrichment of CD90.2 peripheral T cells (MACS).*

Single cell suspensions were prepared from pooled lymph nodes and spleens as above. CD90.2⁺ T cells were enriched by positive selection with CD90.2 microbeads, according to the manufacturer’s directions (Miltenyi Biotec, 130-049-101). MACS-enriched samples were stained with CD3 antibodies and analysed by flow cytometry. All samples had T cell purities of >91%.

*Flow-assisted cell sorting (FACS).*

Single cell suspensions were prepared from thymic tissue and peripheral organs as above. After staining with the desired antibodies, cells were sorted on a FACSaria (BD Biosciences). To sort, doublets were first excluded by FSC-A vs. FSC-H analysis, and then gated as desired. Sorted samples were re-analysed and all had 96-99% purity.
In vitro thymocyte survival assay.

The survival of control and cKO thymocytes was measured as published previously\(^3\). Briefly, single-cell suspensions of thymocytes from 3-week old mice were prepared as before, and 1 \(\times\) 10\(^5\) cells were cultured in RPMI with 10% FCS, 55 \(\mu\)M \(\beta\)-mercaptoethanol, 1%P/S and 10 ng/\(\mu\)L IL-7 (R&D Systems). At the desired time points, the proportion of live cells was determined by staining with a Live/Dead Fixable Near IR stain (Life Technologies), as per the manufacturer’s directions.

PCR.

T cell populations were isolated by FACS-sorting as described above. Genomic DNA was extracted from FACS-sorted cells by phenol/ethanol according to the protocol of Hansmann \(et\ al\)\(^4\). The analysis of TCR\(\beta\) gene rearrangements was performed in DN4 sorted cells (>98% purity) according to the protocol of Hoshii \(et\ al\)\(^5\). The analysis of the excision of the PRR gene was from various sorted thymic and peripheral T cell populations, all with a purity >95%. For detection of the PRR floxed gene (i.e. non-excised), DNA samples were incubated with the forward primer \#1 (5’-AGCAGCTCTCTTCCAGGTATGTTGTG-3’), and reverse primer \#1 (5’-CTGGATCCCCGAGCATGGGGTAAAGG-3’). This resulted in the production of a 330 bp product for the floxed gene, or a 280 bp product for non-floxed/wildtype genes. For detection of the mutated gene after Cre-recombination “cKO” (i.e. excised), DNA samples were incubated with the forward primer \#1 and reverse primer \#2 (5’-GCCCTCTCTTACAGTTCTATCAGT-3’). This PCR product was 326 bp in size. Primers for detection of the Cre recombinase were as described previously\(^6\). Bands were resolved on an agarose gel, and visualised with Ethidium bromide under ultra-violet light. Quantitative RT-PCR of Tcra rearrangement of genomic sorted DP samples was performed as described previously\(^3\).
**Western blotting.**

Single cell thymocyte suspensions were stained with anti-CD90.2 and sorted by FACS as above. The purity of CD90.2$^+$ cells was $>$95%. Sorted cells were then washed once with cold PBS and lysed in RIPA buffer (#9806S, Cell Signaling) containing protease inhibitors. Total protein was determined by a Bradford assay and an equal amount of total protein was loaded per lane of a 15% SDS-PAGE. Proteins were blotted onto nitrocellulose membrane, blocked with 5% BSA and incubated with antibodies against PRR (Sigma) and eIF4E (Cell signaling) overnight at 4°C. Blots were then incubated with infrared labelled secondary antibodies and bands visualised with an Odyssey system (Li-COR Biotechnology). Quantification of bands was performed with the program ImageJ.

**Haemotoxylin and Eosin staining of thymii**

Thymi were isolated from 6-week old control and PRR cKO mice. After fixing and embedding in paraffin, thymi were sectioned to a thickness of 8 $\mu$m for further analysis. Hematoxylin and eosin staining was performed by routine procedures. Sections were analyzed with a Zeiss Axioplan 2 imaging microscope (Carl Zeiss) at 5X magnification.

**Isolation of total RNA and real-time qPCR.**

FACS isolated DN3 cells (>96% purity) were washed once with PBS and then resuspended in Qiazol for the isolation of total RNA using an RNeasy RNA isolation kit (Qiagen), according to the manufacturer’s protocol. The synthesis of cDNA and quantitative analysis of mRNA expressions by real-time PCR was performed as described previously. Real-Time PCR was performed on a 7500 Fast Real-Time PCR System using TaqMan Fast Universal PCR Master Mix (both Applied Biosystems). Data were evaluated with the 7500 Fast System Software (Applied Biosystems). Samples were measured in triplicate and normalized against 18s. The primer and oligonucleotides for real-time PCR were designed using Primer Express®ABI PRISM software to
be species specific (except for the eukaryotic 18s primer and probe) and exon overlapping. 18s (fw 5’ACATCCAAGGAAGGCAGCAG 3’, rev 5’ TTTTCGTCACTACCTCCCCG 3’, probe 5’ FAM-CGCGCAAATTACCCACTCCCCGAC-TAMRA 3’); mouse PRR (fw 5’ CCAGTTTGTTGTCTCGTCAATAAC 3’, rev 5’ ACCTGCCAGCTCCAATGAAT 3’, probe 5’ FAM-TCTAGCCAAGGACCATTCCACCCGACTT-TAMRA 3’).

Microarray.

Total RNA was isolated from FACS-sorted DN3 cells as above. Due to the low RNA content in these samples, amplified cDNA was prepared with the Ovation PicoSL WTA System V2 (NuGEN Technologies, CA, USA) according to the manufacturer’s directions. Three μg of amplified and purified cDNA was then labelled with biotin by the Encore BiotinIL Module (also NuGEN), according to the manufacturer’s directions. Finally, 900 ng of labelled target was analyzed using the Illumina Mouse ref 8v2.0 array. Microarray data are available in the ArrayExpress database (www.ebi.ac.uk/arrayexpress) under accession number E-MTAB-12345. Data was processed on the Illumina GenomeStudio V2011.1 Platform (Gene Expression Module 1.9.0), quantile normalized on a probe level, without background correction (Table S1A). The data was then analyzed with the NIA Array Analysis tools (http://lgsun.grc.nia.nih.gov/ANOVA/index.html). First, hierarchical clustering of averages was performed for assessment of the quality of biological replicates (Fig S6A). Genes which were significantly differentially regulated were identified by ANOVA–false discovery rate (FDR) statistics (http://lgsun.grc.nia.nih.gov/ANOVA) (FDR<0.05; Fig S6B). Fold-changes were calculated from mean expression values in control to cKO samples. A list containing 1133 genes were identified as having a difference in expression >1.5-fold. This list was then analyzed for the enrichment of gene ontology (GO) pathways with the online tool DAVID (http://david.abcc.ncifcrf.gov/) (Table S1B). Due to the high overlap of gene names within these pathway lists, for presentation purposes only 10 representative GO pathways are shown. Full details are provided in Table S1B.
Statistics.

All data is presented as mean ± standard error. Outliers were first excluded using a Grubbs' statistical test. Normality of data was assessed according to the Kolmogorov-Smirnov test, and then analyzed by either two-sided Student’s t-tests (parametric) or a Mann-Whitney U-tests (for non-parametric data). Analysis of Tcra rearrangement in DP cells, and thymocyte survival over time was by ordinary two-way ANOVA with Sidak’s multiple comparison test.

Supplemental references


