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PKCα/β and CYLD Are Antagonistic Partners in the NFκB and NFAT Transactivation Pathways in Primary Mouse CD3+ T Lymphocytes

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Abstract

In T cells PKCα mediates the activation of critical signals downstream of TCR/CD28 stimulation. We investigated the molecular mechanisms by which PKCα regulates NFκB transactivation by examining PKCα/β single and double knockout mice and observed a redundant involvement of PKCα and PKCβ in this signaling pathway. Mechanistically, we define a PKCα-CYLD protein complex and an interaction between the positive PKCα/β and the negative CYLD signaling pathways that both converge at the level of TAK1/IKK/iκBζ/NFκB and NFAT transactivation. In Jurkat leukemic T cells, CYLD is endoproteolytically processed in the initial minutes of stimulation by the paracaspase MALT1 in a PKC-dependent fashion, which is required for robust IL-2 transcription. However, in primary T cells, CYLD processing occurs with different kinetics and an altered dependence on PKC. The formation of a direct PKCα/CYLD complex appears to regulate the short-term spatial distribution of CYLD, subsequently affecting NFκB and NFAT repressional activity of CYLD prior to its MALT1-dependent inactivation. Taken together, our study establishes CYLD as a new and critical PKCα interactor in T cells and reveals that antagonistic PKCα/β-CYLD crosstalk is crucial for the adjustment of immune thresholds in primary mouse CD3+ T cells.

Introduction

The central role of PKCα in signal transduction pathways during an adaptive immune response has extensively focused on the exact biochemical mechanisms of PKCα function (reviewed in [1–3]). A recent study by Kong et al. identified the structural requirement in PKCα for its localization to the immunological synapse as a prerequisite for activation of downstream signaling [4]. Several transcription factors essential for the T cell activation response (i.e. NFκB, AP1, and NFAT) are regulated by PKCα [5,6]. In vivo analysis of PKCα knockout mice revealed the importance of PKCα for Th2- [7] and Th17-mediated immune responses [8,9] but not for host-protective antiviral responses [10]. Nevertheless, despite a profound understanding of the cellular role of PKCα, little is known about its molecular function, specifically the effector proteins downstream of PKCα during T cell activation.

Ubiquitylation and deubiquitylation are established posttranslational mechanisms for regulating immune responses, as well as the development and activation of immune cells. The tumor suppressor gene CYLD encodes an evolutionary conserved and ubiquitously expressed protein of approximately 120 kDa and was originally discovered as gene mutated in familial cylindromatosis, an autosomal dominant inherited disease characterized by the development of multiple benign skin tumors, principally on the head and neck [11]. Functionally it is a deubiquitylating enzyme (DUB) which removes mainly K63-linked polyubiquitin chains from several specific substrates, influencing in a negative way the activation status and/or spatial distribution of these target proteins in different signaling pathways. Numerous studies both in vitro and in vivo provided us with new insights in its established function as an important negative regulator of inflammatory responses, by counteracting the aberrant activation of NFκB signaling: Cyld−/− animals spontaneously develop intestinal inflammation and autoimmune symptoms due to the constitutive activation of the TAK1/IKK/iκBζ axis [12,13]; the study of Lim et al. described a CYLD dependent negative NFκB regulation during bacteria induced lung inflammation in mice via deubiquitylation of TRAF6 and TRAF7 [14]; moreover, the same scientific group showed that Cyld knockout mice are protected from Streptococcus pneumoniae infection and lethality via a negative crosstalk with p38 MAPK [15]; a synergistic crosstalk between the E3 ligase Itch and CYLD for TAK1 inactivation and termination of tumor necrose factor dependent inflammatory signaling was recently described [16].
Figure 1. Overlapping roles of PKC\(\alpha\) and PKC\(\beta\) in NF\(\kappa\)B and NFAT transactivation processes in primary mouse CD3\(^{+}\) T cells. (A) PKC\(\beta\) expression is upregulated in whole cell extracts of peripheral CD3\(^{+}\) T cells derived from PKC\(\alpha\)-deficient mice. (B) The PKC isoform expression profile in whole cell extracts of naive thymocytes (Thy) and peripheral CD3\(^{+}\) T cells derived from wild-type and PKC\(\alpha\)/\(\beta\)-deficient mice. PKC\(\alpha\)/\(\beta\) inhibition leads to an
increased NF-κB and NFAT transactivation defect in T cells. (C) The nuclear extracts of resting and stimulated (overnight) wild-type, PKCβ−/−, PKCβ+/− and PKCβ+/− CD3+ T cells were probed for DNA binding to radio-labeled probes containing NFκB and NFAT binding site sequences, as indicated. One representative experiment of three is shown. (D) Impaired nuclear import of p50, p65 and NFAT in activated PKCβ+/−−/− T cells. Nuclear extracts of resting and stimulated (overnight) wild-type, PKCβ−/−, PKCβ+/− and PKCβ+/− CD3+ T cells were probed for p65, p50 and NFAT using immunoblot assays. DNA polymerase served as the loading control. One representative experiment of three is shown. (E) Effect of PKCβ inhibition on proximal phosphorylation events after a brief stimulation. Western blot analysis was performed with cytosolic extracts from wild-type, PKCβ−/−, PKCβ+/− and PKCβ+/−−/− CD3+ T cells. CD3+ T cells were stimulated with anti-CD3/anti-CD28 and probed at different time points for the phosphorylation status of (p)S-32 IκBα. Protein phosphorylation levels were relatively quantitated by densitometric analysis. Numbers beneath bands indicate fold change compared to wt control after normalization to FYN.

doi:10.1371/journal.pone.0053709.g001

Table 1. Absolute cell numbers of thymic populations from wild-type and PKCβ+/−/− mice.

<table>
<thead>
<tr>
<th></th>
<th>CD3+</th>
<th>CD4+</th>
<th>CD8+</th>
<th>CD4+CD8+</th>
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<tr>
<td>Thymus</td>
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<tr>
<td>wt</td>
<td>27.1±1.3</td>
<td>8.2±1.6</td>
<td>3.3±2.0</td>
<td>143.8±16.9</td>
</tr>
<tr>
<td>PKCβ+/−</td>
<td>17.2±2.5</td>
<td>4.7±0.9</td>
<td>2.2±0.7</td>
<td>150.6±29.6</td>
</tr>
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Absolute cell numbers of thymic populations (x107). The results shown are the mean±SE of three independent experiments.

doi:10.1371/journal.pone.0053709.t001

Materials and Methods

Mice

PKCβ−/− knockout mice are viable, fertile and were generated by crossing PKCβ [5] and PKCβ−/− [19] single knockout mice. The generation of Cyld-deficient mice was described previously [20]. All mice were on a C57BL/6 background and housed (under SPF conditions) at the mouse facility of the Medical University of Innsbruck. All animal experiments were performed in accordance with the Austrian “Tierschutzgesetz” (GBGL. Nr. 501/1988 i.d.q.f.) and have been granted by the Bundesministerium für Bildung, Wissenschaft und Kultur (bmwfw).

Plasmids and Reagents

Strep-HA-tagged PKCβ and Cyld cDNAs (full-length and R324A mutant) were cloned into pEF-Neo. Vectors expressing full-length Flag-tagged wild-type CYLD or N- or C-terminally truncated forms of CYLD (encoding residues 1–212, 318–956 and 587–986 of CYLD) were described previously [21].

The pan-PKC low molecular weight inhibitor LMWI [22] was provided by NYCOMED GmbH, and the tetrapeptide inhibitor z-VRPR-fmk (MALT1 LMWI) was a gift from Dr. Margot Thome.

Cell Culture and Transfections

Jurkat-TAg cells [23] (a kind gift from G.R. Crabtree, Stanford University, CA) were maintained in RPMI medium supplemented with 10% FCS (Life Technologies, Inc.) and antibiotics. Transient transfection of cells with 20 μg of plasmids encoding GFP, wild-type Cyld or a cleavage-resistant R324A Cyld mutant was performed by electroporation with a BTX-T820 Electro Square Porator (ITC, Biotech, Heidelberg, Germany) apparatus under predetermined optimal conditions: 2×107 cells at 450 V/cm and five pulses of 99 ms.

HEK293T cells were cultured in Dulbecco’s Modified Eagle’s Medium supplemented with 10% FCS, 2 mM L-glutamine, and 100 μg/ml penicillin-streptomycin. HEK293T cells were transfected using MetafecteneTM transfection reagent according to the manufacturer instructions.

Primary human T cells were purified from PBMCs (isolated by standard Hypaque–Ficoll separation from whole blood samples) with the Pan T Cell Isolation Kit (Miltenyi Biotec) according to the manufacturer instructions.

Primary mouse CD3+ T cells were purified from pooled spleens and lymph nodes with mouse T cell enrichment columns (R&D Systems). T cell populations were typically 95% CD3+ as determined by staining and flow cytometry.

doi:10.1371/journal.pone.0053709.t002
Analysis of Proliferative Response and IL-2 Cytokine Production

For in vitro proliferation, $5 \times 10^5$ T cells in 200 µl proliferation medium (RPMI supplemented with 10% FCS, 2 mM L-glutamine and 50 units/ml penicillin/streptomycin) were added in duplicate to 96-well plates precoated with anti-CD3 antibody (clone 2C11, 5 µg/ml) and soluble anti-CD28 (1 µg/ml; BD Bioscience) was added. For TCR-independent T cell stimulation, 10 ng/ml Phorbol 12,13-dibutyrate (PDBu) and 125 ng/ml of the calcium ionophore ionomycin were added to the media. Cells were harvested on filters after a 64 h stimulation period, pulsed with H$^3$-thymidine (1 µCi/well) in the final 16 h and the incorporation of H$^3$-thymidine was measured with a Matrix 96 direct counter system.

For short time stimulation, cells were activated by the addition of anti-CD3 and anti-CD28 (or PDBu and ionomycin), both in soluble form. For crosslinking, anti-hamster IgG1 (clone HIG-632) was used.

IL-2 production in mouse CD3$^+$ T cells after antibody stimulation was determined by BioPlex technology (BioRad Laboratories) from the supernatant.

Western Blot Analysis

Cells were lysed in ice-cold lysis buffer [5 mM Na$_2$VO$_4$, 5 mM Na$_3$P$_2$O$_7$, 5 mM NaF, 5 mM EDTA, 150 mM NaCl, 50 mM Tris (pH 7.3), 2% NP-40, 50 µg/ml aprotinin and leupeptin] and centrifuged at 15,000×g for 15 min at 4°C. Protein lysates were subjected to immunoblotting using antibodies against NFATc1 (Affinity Bioreagents), pS473 AKT, pERK, ERK, p/T183/

Figure 2. PKC\(\alpha/\beta\) and PKC\(\delta\) synergistically regulate TAK1 and JNK activation. (A) Defective Tak1 and JNK activation in PKC\(\alpha/\beta\)-deficient CD3$^+$ cells. Cytosolic extracts of PDBu- and ionomycin-stimulated wild-type and PKC\(\alpha/\beta^{-/-}\) CD3$^+$ T cells were probed for the phosphorylation status of TAK1, JNK and ERK1/2, as indicated. Actin served as loading control. One representative experiment of three is shown. (B) PKC enzymatic activity influences TAK1 and JNK activation status. The cytosolic extracts of PDBu- and ionomycin-stimulated, pan-PKC LMW1 pretreated, or untreated control wild-type CD3$^+$ T cells were probed for the phosphorylation status of Tak1, JNK and ERK1/2, as indicated. Actin served as loading control. One representative experiment of three is shown. doi:10.1371/journal.pone.0053709.g002
Figure 3. T cells from **Cyld**<sup>−/−</sup> mice exhibit hyper-responsiveness to TCR stimulation. (A) Naive wild-type and **Cyld** deficient CD3<sup>+</sup> cells were stimulated overnight (16 h) with the indicated amount of plate-bound anti-CD3 and soluble anti-CD28. Cytokines in the supernatants were measured using Bioplex suspension array technology. (B) Nuclear extracts were isolated from unstimulated and CD3/CD28-stimulated CD3<sup>+</sup> T cells from wild-type control and **Cyld**<sup>−/−</sup> mice. EMSA was performed to determine the activity of NFAT and NFκB. The same nuclear extracts were probed for NFAT and DNA polymerase, using immunoblot assays, the latter one served as the loading control. (C) Purified naive peripheral CD3<sup>+</sup> T cells from wild-type and **Cyld**<sup>−/−</sup> mice were stimulated with anti-CD3 and anti-CD28 for the indicated time periods. Immunoblotting assays were performed using the
Y185 JNK, JNK, p50, p65, (p)Y-783 PLCγ1, (p)T184/T187Tak1, Tak1 (all from Cell Signaling), PKCβ, PKCδ (both from BD Transduction Laboratories), actin, Cyld (E10), DNA polymerase and Fyn (all from Santa Cruz Biotechnology). The Cyld antibody recognizing the NH2-terminal region of Cyld was described previously [20].

Co-immunoprecipitation Analysis

For co-immunoprecipitation, 1 x 10^6 primary mouse T cells (or 1 x 10^6 transiently transfected HEK293T cells) were lysed in 400 μl of immunoprecipitation buffer [5 mM Na3VO4, 5 mM Na2P2O7, 5 mM NaF, 5 mM EDTA, 150 mM NaCl, 1% NP-40, 50 mM Tris (pH 7.4), 50 mg/ml aprotinin and leupeptin]. Lysates were precleared for 1 h at 4°C. Immunoprecipitation was performed at 4°C overnight using 2 μg of the relevant antibodies. Thereafter, lysates were incubated with protein G Sepharose (Amersham-Pharamacia, Vienna) [for Streptactin IP, Streptactin Beads were used] for 1 h at 4°C, extensively washed in lysis buffer, resolved on an SDS–PAGE and immunostained for the relevant protein.

Gel Mobility Shift Assays

Nuclear extracts were harvested from 1 x 10^7 cells according to standard protocols. Briefly, purified CD3^+^ T cells were washed in PBS and resuspended in 10 mM HEPES (pH 7.9) 10 mM KCl, 0.1 mM EDTA, 0.1 mM EGTA, 1 mM DTT and protease inhibitors. Cells were incubated on ice for 15 min. NP-40 was added to a final concentration of 0.6%, cells were vortexed vigorously, and the mixture was centrifuged for 5 min. The nuclear pellets were washed twice and resuspended in 20 mM HEPES (pH7.9), 0.4 M NaCl, 1 mM EDTA, 1 mM EGTA, and 1 mM DTT and protease inhibitors, and the tube was rocked for 30 min at 4°C. After centrifugation for 10 min, the supernatant was collected. Extracted proteins (2 μg) were incubated in binding buffer with [32P]-labeled, double-stranded oligonucleotide probes (NFkB: 5'-GCC ATG GGG GGA TCC CGG AAG TCC-3'; NFAF: 5'-GCC CAA AGA GGA AAA TTT GTT TGT CTC AG-3') (Nushift; Active Motif). In each reaction, 3 x 10^5 c.p.m. of labeled probe was used, and the band shifts were resolved on 5% polyacrylamide gels. All experiments were performed at least three times with similar outcomes.

Flow Cytometry

Single-cell suspensions from the spleen, lymph node and thymus were prepared and incubated for 30 min on ice in staining buffer (PBS containing 2% fetal calf serum and 0.2% NaN3) with FITC, PE or APC antibody conjugates. Surface marker expression was analyzed using a FACScan™ cytometer (Becton Dickinson & Co., Mountain View, CA) and CellQuestPro™ software according to standard protocols. Antibodies against murine CD3, CD4, and CD8 were obtained from Caltag Laboratories; CD19, CD69, CD44, and CD25 were obtained from BD PharMingen.

Retroviral Transduction of Primary Mouse T cells

The packaging cell line platE was transfected with a pMX retroviral vector encoding an EGFP-CyD8 fusion cDNA. Approximately 36 h later, supernatants were collected and used directly to infect 24 h to 48 h preactivated CD3^+^ cells using spin inoculation (1 h, 2000 x g, 32°C), followed by a 5-6 h incubation period at 37°C. Infected cells were washed, resuspended in full supplement- ed medium and incubated for an additional 48 h to 72 h. From these cultures, GFP-expressing cells were analyzed using confocal microscopy to track the subcellular distribution of the protein of interest.

Monitoring CYLD Localization Using Confocal Microscopy

CD3^+^ cells from wild type and PKCδ/β knockout mice that were transduced with a retrovirus expressing an EGFP-CYLD were not stimulated or PDBu- and ionomycstin-stimulated, transferred to a polylysine-coated slide and fixed with 2% paraformaldehyde. After permeabilization (0.1% TritonX-100 in PBS) and a blocking step (5% goat serum in PBS), the cells were stained with Alexa595-CTB (for lipid raft staining) and TOPRO 3 (Nucleus) (Molecular Probes). Immunofluorescence was analyzed with a Zeiss LSM 510 confocal laser scanning microscope and Zeiss LSM software v3.2.

Statistical Analysis

Differences between genotypes were analyzed using the unpaired Student’s t test.

Results

Overlapping Roles of PKCδ and PKCβ in NFκB and NFAT Transactivation Processes in Primary Mouse CD3^+^ T cells

Studies using targeted gene disruption defined a critical role for PKCδ in the activation of the IL-2 promoter in the NFκB and Ca2+/NFAT pathways [5,6]. Surprisingly, the phenotypic characterization of PKCδ-deficient T cells revealed a strong upregulation of PKCβ protein levels in PKCδ single knockout T cells (Fig. 1A). To investigate potentially compensatory and overlapping roles of these two PKC family members in T cell activation processes, PKCδ/β double knockout mice were generated. These mice were viable, fertile and breed at normal Mendelian ratios. The null mutations for PKCδ and PKCβ were confirmed by PCR and immunoblotting of whole cell lysates from naive thymocytes and peripheral CD3^+^ T cells (Fig. 1B).

Flow cytometric analysis of thymocyte populations in PKCδ/β double knockout mice revealed a slightly diminished percentage of CD3^−^, CD4^-^ and CD8^-^ positive cells, comparable to the PKCδ single knockout phenotype and in agreement with previous research [24,25], which might indicate an involvement of PKCδ in the positive selection process during thymocyte development. Nevertheless, in the periphery, PKCδ/β^-^- mice revealed no gross differences in the distribution of CD3^−^, CD4^-^, CD8^-^ positive cells, leading to the conclusion that the concomitant loss of PKCδ and PKCβ did not additively affect T cell development (Table 1+ Table 2).

Examination of the stimulation-dependent upregulation of CD25, CD69 and CD44 surface markers on CD4^+^ and CD8^+^ subsets revealed no gross differences in the total percentage of positive cells between the genotypes, but the total protein amount per cell, monitored by median fluorescence intensity, was strongly reduced in PKCδ/β^-^- and to an intermediate extend in PKC singly-deficient T cells. These data might indicate a possible defect in the upregulation of both the IL-2 receptor chain alpha (CD25) and the activation marker CD69 in PKC-deficient T cells in both CD4^+^ and CD8^+^ T cells (Fig. S1).
Figure 4. Association of CYLD with PKC\(\beta\) and PKC\(\beta\) and CYLD Signaling Crosstalk

(A) CYLD directly interacts with PKC\(\beta\) in primary mouse CD3\(^+\) cells. The complex is formed constitutively and is not affected by TCR activation. One representative experiment of three is shown. The C-terminal region of CYLD is important for interaction with PKC\(\beta\). (B) Co-immunoprecipitation of PKC\(\beta\) using CYLD pulldown. Increased binding of PKC\(\beta\) to the C-terminal region of CYLD was shown in HEK293T cells transiently co-transfected with vectors encoding PKC\(\beta\) (PEFneo) and a full-length Flag-tagged wild-type CYLD, N- or C-terminally truncated forms of CYLD (residues 1–212, 318–956 and 587–986 of CYLD). Untransfected and GFP-transfected controls were included. One representative experiment of three is shown. A schematic representation depicts the CAP-Gly and peptidase domains in wild-type and truncation mutants of CYLD. (C) Co-immunoprecipitation of CYLD using PKC\(\beta\) pulldown. A strep-tagged full-length PKC\(\beta\) construct was co-transfected with Flag-tagged CYLD constructs into HEK293T cells. As previously, the importance of the C-terminal region of CYLD for binding is shown. GFP controls for each CYLD construct were included. One representative experiment of three is shown.

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In contrast to the relatively normal T-cell development observed, the T-cell response of peripheral T cells after TCR stimulation was affected by the single and simultaneous loss of PKC\(\alpha\) and PKC\(\beta\). The H\(^+\)-thymidine uptake and IL-2 secretion response of PKC\(\alpha/\beta\)-deficient T cells stimulated with anti-CD3 and with or without anti-CD28-activated did not significantly exacerbate the defects already observed in the absence of PKC\(\alpha\) alone (Fig. S2A–C). To exclude the proliferative defects being caused by deregulated apoptosis, we analyzed the activation-induced cell death (AICD) of CD4\(^+\) and CD8\(^+\) T-cell blasts derived from wild-type and double knockout animals using CD3 engagement in vitro; in addition also the Fas ligand induced cell death was monitored, but no enhanced apoptotic responses of PKC\(\alpha/\beta\)-deficient were detected (Fig. S3A–B).

However, analysis of the pathways leading to IL-2 transcription revealed additively reduced binding of NF\(\kappa\)B and NFAT to DNA in PKC\(\alpha/\beta\) double-deficient CD3\(^+\) T cells after CD3/CD28 stimulation (Fig. 1C). Immunoblot analysis of nuclear extracts demonstrated that the weaker DNA binding of NF\(\kappa\)B and NFAT was due to the reduced nuclear entry of two NF\(\kappa\)B subunits, p50 and p65, and NFAT upon stimulation (Fig. 1D). Activation of NF\(\kappa\)B involves the phosphorylation of I-kB\(\alpha\) by IKK\(\beta\) and its subsequent proteasomal degradation. Consistent with the additive effect on NF\(\kappa\)B translocation, the double knockout showed a weaker I-kB\(\alpha\) phosphorylation after stimulation with CD3/CD28. Also the activation of the Map kinase pathway was partially affected by PKC\(\alpha/\beta\) deficiency, visible through a reduced ERK phosphorylation, whereas the activation of Akt/PKB was normal (Fig. 1E).

PKC\(\alpha\) and PKC\(\beta\) Synergistically Regulate TAK1 and JNK Activation

In agreement with the defective IKK/I-kB\(\alpha\) axis, PKC\(\alpha/\beta\)-deficient CD3\(^+\) cells revealed a drastic activation defect in TGF\(\beta\) activated kinase 1 (TAK1), which is known to be a key regulator of IKK\(\beta\) signaling. The loss of both PKC isoforms appears to be required to abolish the signal, as TAK1 activation levels were similar between the wild-type and PKC single knockout T cells (not shown). Additionally, the JNK signal was attenuated by the targeted disruption of PKC\(\alpha\) and PKC\(\beta\), whereas ERK1/2 activation was only marginally affected (Fig. 2A). Similar outcomes were observed in PDBu- and ionomycin-stimulated primary mouse wild-type T cells pretreated with 500 nM of a PKC-specific low molecular weight inhibitor (PKC LMW). The stronger effect of the pharmacological pan-PKC inhibitor on MAP kinase activation can be best explained by its described inhibition of additional PKC family members next to PKC\(\alpha\) and PKC\(\beta\) (Fig. 2B).

Cyld\(^{+/−}\) T cells Show a Hyperactive Phenotype in NF\(\kappa\)B and NFAT Transactivation Responses

Since the deubiquitinating enzyme CYLD has been shown to be a negative regulator of Taki [12] and JNK signaling [26], we investigated a possible link between CYLD and the PKC\(\alpha/\beta\) isoforms in NF\(\kappa\)B and NFAT driven IL-2 upregulation.

Despite of an observed thymocyte maturation defect, recent work on T cell signaling in Cyld\(^{−/−}\) mice demonstrated hyperresponsiveness to TCR stimulation by constitutive activation of NF\(\kappa\)B [12]. We confirmed these results as we also observed that Cyld\(^{−/−}\) T cells showed elevated activation-induced IL-2 responses (Fig. 3A). This hyper-responsive IL-2 secretion correlated with an increase of NF\(\kappa\)B DNA binding to the IL-2 promoter in the nuclear fractions of stimulated Cyld-deficient T cells (Fig. 3B) and hyper-phosphorylated I-kB\(\alpha\) levels in the cytosol compared to wild-type controls (Fig. 3C). Interestingly, and in accordance with a previous publication [27], we determined that CYLD also acts as a negative modulator of the NFAT pathway. The examination of NFAT transactivation using immunoblot and EMSA technology revealed increased nuclear translocation and subsequent binding of NFAT to DNA in Cyld-deficient cells (Fig. 3B). Our EMSA result was confirmed by the elevated activation status of phospholipase C\(\gamma\), which has been identified as a key regulator of C2+/Calcineurin/NFAT signaling. However, ERK signaling was not affected by the loss of Cyld (Fig. 3C).

Association of CYLD with PKC\(\alpha\)

Considering the reciprocal phenotypes of Cyld- and PKC\(\alpha/\beta\)-deficient T cells involving NFAT and NF\(\kappa\)B transactivation we investigated a potential direct interaction between these enzymes. Interestingly and indeed, we identified a physical and functional PKC\(\alpha\)/CYLD interaction in the cytosol of primary T cells. The co-immunoprecipitation analysis of CYLD and the PKC\(\alpha\) isotype from cell extracts of unstimulated and CD3/CD28-activated peripheral CD3\(^+\) cells revealed that PKC\(\alpha\) and CYLD physically associate in a complex in resting conditions (Fig. 4A). Next, we mapped the PKC\(\alpha\) interaction domain in the CYLD protein by co-transfection of HEK293T cells with a vector encoding PKC\(\alpha\) and with vectors expressing full-length Flag-tagged wild-type or N- and C-terminally truncated forms of CYLD (encoding residues 1–212, 318–956 and 587–986 of CYLD). The CYLD pull down with a specific Flag antibody revealed increased binding of PKC\(\alpha\) to CYLD mutants containing the deubiquitinae domain (Fig. 4B). We observed identical results when the co-immunoprecipitation was performed to precipitate PKC\(\alpha\). A strep-tagged PKC\(\alpha\) construct was co-transfected with the Flag-tagged CYLD constructs. A GFP control for each CYLD construct was included to identify unspecific binding to the Streptactin-beads. PKC\(\alpha\) precipitation confirmed that the C-terminal part of CYLD is necessary for complex formation between the two interacting protein (Fig. 4C).

In Jurkat Cells, CYLD is Cleaved by MALT1 in a PKC-dependent Mechanism

When Jurkat cells overexpressing an N-terminally HA-tagged CYLD construct were stimulated for 30 min with PDBu and ionomycin, a CYLD fragment of approximately 40 kDa was...
Figure 5. CYLD is endoproteolytically processed in the Jurkat cell line and in primary human T cells. (A) The essential role of both the catalytic activity of PKC and MALT1 for activation-dependent CYLD cleavage. Jurkat cells transfected with HA-tagged Cyld vectors were activated by PDBu and ionomycin in the presence of PKC\textsubscript{h} and MALT1 pharmacological inhibitors. After 20 min, cells were lysed and fractionated into membrane (m), cytosolic (c) and nonsoluble (ns) fractions. CYLD and its clipping product were present in (c), inhibition of PKC\textsubscript{h} and MALT1 activity blocks CYLD cleavage. (B) CYLD processing occurred also under endogenous conditions in stimulated primary human T cells isolated from whole blood samples;
detected using the anti-HA antibody. Because the administration of PDBu mimics TCR signaling by activating PKC family members, we wanted to identify a role for PKC in this cleavage event. Therefore, Jurkat cells were pretreated with the specific pan-PKC pharmacological inhibitor, which resulted in the disappearance of this fragment (Fig. 5A). This finding emphasized that the endoproteolytic cleavage of CYLD is PKC dependent. We also isolated primary T cells from human whole blood and analyzed the CYLD processing under endogenous conditions. In addition, human T cells were treated with the pan-PKC inhibitor to investigate the PKC dependency in this process. Comparable to the results with Jurkat cells, CYLD underwent a stimulation dependent processing also in primary human T cells, which could be blocked by PKC inhibition. The generation of a 40 kDa NH2-terminal and a 70 kDa C-terminal cleavage fragment was confirmed via the use of NH2- and C-terminus recognizing terminal and a 70 kDa C-terminal cleavage fragment was generated a arginine residue at position 324 in the human CYLD protein and terminal 40 kDa proteolytic fragment, we identified the cleavable (Fig. 5A).

Because Coornaert et al. showed that the paracaspase MALT1 directly cleaved the deubiquitinating protein A20 to generate a fragment with a smaller molecular size in stimulated T cells [28], we asked if MALT1 was also responsible for the cleavage of CYLD by treating Jurkat cells with the tetrapeptide inhibitor z-VPRPR-fmk, which has been shown to inhibit specifically the MALT1 protease activity [29]. As a result, cells treated with the MALT1 inhibitor showed a reduced CYLD cleavage after activation (Fig. 5A).

Based on the size and the molecular weight of the CYLD N-terminal 40 kDa proteolytic fragment, we identified the cleavable arginine residue at position 324 in the human CYLD protein and generated a Cyld mutant with alanine substituted for arginine at position 324 (Cyld-R324A). Next, we investigated if this mutant was cleavable when overexpressed in Jurkat cells or was resistant to proteolysis. Wild-type CYLD was processed after stimulation, whereas the mutant could no longer be cleaved (Fig. 5C). This led to permanent inhibition of the NFkB pathway by the inactivation resistant CYLD mutant and result in slightly diminished phosphorylation of IκBα. The MAPK pathway was not affected by the expression of uncleavable CYLD.

Because NFκB is important for IL-2 upregulation in activated T cells, we tested the influence of the protease-resistant Cyld mutant on IL-2 transactivation using an IL-2-promoter-dependent luciferase assay. The diminished NFκB signal observed by immunoblot was correlated with impaired IL-2 transcription (Fig. 5D). This provides experimental evidence that CYLD processing, which leads to an inactivation of its repressor function within a positive feedback loop, is an important prerequisite for robust IL-2 activation. Consistent with our investigation of CYLD cleavage at arginine 324 in the Jurkat tumor cell line, Staal et al. independently published that TCR-induced JNK activation required CYLD proteolysis by MALT1 [18]. Nevertheless, our findings extend the function of CYLD cleavage to NFκB activation. To identify the physiological role of this candidate process, we examined primary CD3ε T cells derived from wild-type and knockout mice.

PKC-dependence, Cleavage Site and Kinetics of CYLD Cleavage Differ in Primary Mouse T cells

Stimulation induced a CYLD fragment not only in human cells but also in primary mouse T cells. Interestingly, activation of primary mouse T cells by CD3 with or without CD28 costimulation generated a NH2-terminal CYLD fragment of approximately 25 kDa, smaller in size than the human fragment (Fig. 6A), implicating a different cleavage site in the mouse CYLD protein. Although we did not determine the exact cleavage site, arginine 235 was the best candidate for the cleavage site. Importantly, the fragment was first detectable after 4 h of stimulation, indicating different kinetics leading to CYLD inactivation in primary mouse cells. Unexpectedly, PKC0/β-deficient T cells showed normal CYLD processing after stimulation, implicating additional protein kinases in this process (Fig. 6B). As a consequence, the activation defects of the TAK1/IKK axis in PKC0/β-deficient T cells cannot solely be explained by CYLD inactivation.

Alternatively, the existence of a constitutive CYLD/PKC0 complex might suggest that PKC0, which shows activation-dependent subcellular translocation, is important for removing CYLD from its NFκB-related targets and attenuates the negative regulatory function of CYLD, enabling feedback control of NFκB activation. To address this hypothesis, we analyzed the subcellular distribution of a retrovirally introduced mutant CYLD-GFP fusion protein in unstimulated and stimulated wild-type and PKC0/β-deficient T cells with confocal microscopy. CD3ε+ T cells from both genotypes were retrovirally infected with a CYLD-GFP fusion construct, and the colocalization of CYLD-GFP with lipid rafts was monitored with Cholera Toxin B. Interestingly, CYLD colocalization with lipid rafts was strongly diminished in PKC0/β-deficient T cells compared to control cells, suggesting PKC0-dependent CYLD membrane shuttling. A statistically significant decrease in CYLD translocation in double knockout cells was detected in unstimulated cells, whereas after 15 min of PDBu and ionomycin stimulation, CYLD translocation to the membrane was observed in both wild-type and knockout cells (Fig. 6C).

Discussion

Numerous studies emphasize PKC0s key role as a regulator of NFκB and Ca2+/NFAT signaling in T cells downstream of the TCR [5,6,30–32]. The PKCβ isotype is also expressed in T cells. However, PKCβ-deficient primary mouse T cells have a fairly normal activation response [33], although Volkov et al. established a major role for PKCβ in LFA1-dependent T-cell locomotion [34,35].

Our recent work defines a redundant role for both the novel PKC0 variant and the classical isotype PKCβ in the NFκB and NFAT signaling pathways. T cells isolated from PKC0/β-deficient mice had a stronger impairment in NFκB and NFAT nuclear entry and DNA binding compared to CD3ε+ T cells from control and single knockout mice. Impaired TAK1 activation in double-deficient T lymphocytes is the best candidate for restricted IκK/IκBα signaling. Functional redundancy of PKC0 with other members of the PKC family in NFκB and/or NFAT activation has been shown in previous studies. For instance, stimulation-
Figure 6. CYLD cleavage in primary mouse T cells shows a different kinetic. (A) CYLD cleavage in primary mouse cells has different kinetics. The activation of primary mouse T cells by CD3 with or without CD28 costimulation leads to the formation of an NH2-terminal CYLD fragment of
dependent colocalization of atypical PKCζ/ι with PKCθ in the lipid raft fraction of T lymphocytes leads to cooperation of these isoforms in modulating the NFκB signaling pathway [36]. The collaborative activity of PKCθ and PKCζ in the NFAT pathway was examined in a PKCζ/θ double knockout mouse strain. Compared to PKCζ and PKCθ single-deficient T cells, double-deficient CD3ζ-cells showed additively reduced IL-2 secretion levels correlated with strongly impaired nuclear translocation and DNA binding of NFAT after stimulation. Of note, the PKCζ/θ double knockout mice showed an impaired alloimmune response, leading to significantly prolonged allograft survival in heart transplantation experiments [37].

Similar to phosphorylation, K63 ubiquitination is a reversible process that influences protein activity, trafficking and signaling complex assembly. The removal of ubiquitin chains is mediated by a family of deubiquitinases, of which the cylindromatous gene product CYLD and the Tumor necrosis factor α-induced protein 3, also called A20, is currently receiving broad scientific attention. Both CYLD and A20 have been implicated as a modulator of the activity of NFκB-related molecules, such as NEMO (IKKγ), TRAF2 and TRAF6 [38–40]. Both enzymes overlap functionally by targeting a similar set of substrates, which was explained by the different expression pattern of A20 and CYLD. A20 function depends on its transcriptional upregulation, whereas CYLD is constitutively expressed, influencing the different time windows of NFκB activation differently. However, a constitutive expression and activity pattern requires a posttranslational regulatory mechanism to inactivate the repressor during signal-induced NFκB signaling.

The cleavage-dependent inactivation of a deubiquitinase as a posttranslational regulatory mechanism in activated T cells was first described by Coornaert et al. [28], in which A20 was defined as a MALT1 substrate, which upon antigen receptor engagement undergoes cleavage for functional NFκB signaling. In our study, we showed that PKCθ and CYLD are constitutively bound in a physical complex in the cytosol of primary mouse CD3ζ-cells. Direct crossstalk between CYLD and a PKC family member has not been described to date; therefore, we aimed to elucidate the biological relevance of this protein-protein interaction in T-cell signaling by examining genetic knockout mouse models in combination with selective pharmacological inhibitors. The reciprocal phenotypes of T-cell signaling pathways in Cyld−/− and PKCθ−/− mice prompted us to analyze the activity of key molecules linked to NFκB and NFAT transactivation to uncover a regulatory mechanism to address the modulation of TAK1 activity. In agreement with Koga et al. [27], who demonstrated negative regulation of NFAT activity by CYLD via the TAK1/MK3/6/p38α/β axis, our experimental data clearly attest to CYLD involvement in NFAT activity modulation downstream of TCR signaling.

Our results show that PKC, particularly the PKCθ/ι isoforms, can influence CYLD repressive activity in different ways. In the human Jurkat leukemic T-cell line, PKC enzymatic activity was important for rapid MALT1-dependent CYLD processing, which is required for TCR-linked NFκB transactivation and leads to functional IL-2 induction. The requirement for MALT1-mediated CYLD cleavage for intact JNK signaling downstream of the TCR has been previously described [18]. Thus, proteolytic inactivation of CYLD affected IL-2 transactivation via the JNK/AP1 pathway; here, we provide experimental evidence that NFκB activity is also specifically dependent on CYLD cleavage, subsequently modulating IL-2 signals. Of note, we independently confirmed arginine 324 as the CYLD cleavage site. Recently, caspase 8 has been shown to cleave CYLD at aspartate 215 in Jurkat cells following TNFα stimulation, generating a pro-survival signal to save the cells from necrotic cell death [41]. Additionally, phosphorylation of CYLD was found to downregulate CYLD activity: transient phosphorylation by IKK in a serine cluster just upstream of the TRAF2 binding site, attenuates DUB function [42]. However, NFκB itself can regulate CYLD expression in a negative feedback loop [43].

In primary T cells isolated from PKCθ−/− deficient mice, CYLD was processed to the same extent as in wild-type control cells. Additionally, the kinetics of CYLD cleavage was different in mouse T cells compared to Jurkat cells, starting approximately 4 hours after stimulation, later then the rapid response through TAK1 activation. The different requirement for PKC and the altered kinetics in the mouse system led to the analysis of the stimulation-dependent spatial and temporal organization of the PKCθ/CYLD complex using immunofluorescence microscopy. Interestingly, we found decreased CYLD lipid raft localization in PKCθ−/− deficient T cells under resting conditions, likely affecting activation-induced signaling.

Conclusion

We observed a direct functional connection between the positive PKCθ/ι and the negative CYLD signaling pathways that fine-tune TCR/CD28-induced signaling responses. Our findings suggest the following scenario: PKCθ/ι are the essential kinases in a physiological signaling cascade that is necessary to counteract CYLD-mediated repression of NFκB and NFAT transactivation. This direct and physical antagonistic crossstalk between the PKG-derived signals and the CYLD-derived signals might represent one mechanism of how antigen-receptor-dependent fine-tuning of the amplitude of T lymphocyte activation is processed.

Supporting Information

**Figure S1** Effect of PKCθ−/− deficiency on CD25, CD44, and CD69 surface expression. T cells were stimulated for 16 h by CD3/CD28 ligation and the surface expression of CD25, CD44, and CD69 for CD4+ and CD8+ subsets were measured by flow cytometry. The relative fluorescence intensities are indicated as the median fluorescence intensity. The results shown are the mean±SE of three independent experiments. (TIF)

**Figure S2** Proliferative and cytokine secretion responses of PKCθ−/− CD3ζ T cells. (A, B) Proliferative responses of PKCθ−/− and PKCθ-deficient CD3ζ T cells were analyzed in comparison to wild-type littermate controls. After incubation using different stimulatory conditions (antibodies or BALB/C splenocytes), cells were analyzed using standard procedures for thymidine incorporation. (C) IL-2 cytokine secretion by knockout CD3ζ T cells was analyzed in comparison to wild-type littermate controls. After stimulation with anti-CD3 with or without soluble
Figure S3  Activation-induced cell death (AICD) of CD4+ and CD8+ T cell blasts derived from double knockout animals was not increased compared to cells from single knockout littersmates. (A, B) AICD was induced by different concentrations of anti-CD3 for 8 hours. The results shown are the means of three independent experiments.

References


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Author Contributions

Conceived and designed the experiments: NT GB. Performed the experiments: NT KW NHK SK FF CLN. Analyzed the data: NT KW SK. Contributed reagents/materials/analysis tools: ML MT RM. Wrote the paper: NT GB.