Plasma Membrane Calcium ATPase Regulates Stoichiometry of CD4+ T-Cell Compartments

Maylin Merino-Wong, Barbara A. Niemeyer and Dalia Alansary*

Molecular Biophysics, Saarland University, Homburg, Germany

Immune responses involve mobilization of T cells within naïve and memory compartments. Tightly regulated Ca2+ levels are essential for balanced immune outcomes. How Ca2+ contributes to regulating compartment stoichiometry is unknown. Here, we show that plasma membrane Ca2+ ATPase 4 (PMCA4) is differentially expressed in human CD4+ T compartments yielding distinct store operated Ca2+ entry (SOCE) profiles. Modulation of PMCA4 yielded a more prominent increase of SOCE in memory than in naïve CD4+ T cell. Interestingly, downregulation of PMCA4 reduced the effector compartment fraction and led to accumulation of cells in the naïve compartment. In silico analysis and chromatin immunoprecipitation point towards Ying Yang 1 (YY1) as a transcription factor regulating PMCA4 expression. Analyses of PMCA and YY1 expression patterns following activation and of PMCA promoter activity following downregulation of YY1 highlight repressive role of YY1 on PMCA expression. Our findings show that PMCA4 adapts Ca2+ levels to cellular requirements during effector and quiescent phases and thereby represent a potential target to intervene with the outcome of the immune response.

Keywords: Plasma Membrane Calcium ATPase, Yin Yang 1, CD4+ compartments, calcium signaling, stoichiometry

INTRODUCTION

The spatiotemporal characteristics of Ca2+ signals in T cells tightly control the outcome of an immune response as well as the fate of the immune cells involved. A local or global rise of intracellular [Ca2+]i is achieved by activation of Ca2+ channels or by mobilization from intracellular stores. Once the Ca2+ signal is conducted, extrusion pumps restore the resting concentration. Alternatively, resting concentration is restored by Ca2+ uptake into intracellular stores: mitochondria or endoplasmic reticulum (ER).

Plasma membrane Ca2+ ATPase (PMCA) is the major extrusion pump playing a pivotal role in the regulation of Ca2+ dynamics during activation of T cells (1–4) in co-operation with sodium calcium exchanger (NCX) (5) and the sarco-endoplasmic reticulum Ca2+ ATPase (SERCA) (6). In addition to its global role, PMCA is able to generate local gradient in its vicinity thereby influencing Ca2+ dependent interaction partners (7). Furthermore, PMCA interacts with the inhibitory domain of CD147 in a Ca2+ independent manner which is necessary for CD147-mediated inhibition of IL-2 production (8). PMCA is a type-P ATPase of the class PII (9). In mammals PMCA contains a C-terminal auto-inhibitory domain, by which the inhibition is alleviated by binding to calmodulin or acidic phospholipids which can also bind to a basic domain in the first cytosolic loop [reviewed...
in (10)]. There are four genes expressing PMCA (ATP2B1–4) with two possible splice sites resulting in a sum of about 30 variants belonging to four isoforms (11). All isoforms of PMCA have Ca²⁺-dependent activity with PMCA4b being the slowest isoform despite a higher affinity to Ca²⁺/calmodulin than PMCA4a (12). The immunoglobulin proteins basigin and neuroplastin were identified as obligatory subunits of PMCA isoforms (13, 14). Co-assembly into a complex with either protein is necessary for stable PMCA surface localization in rat brain derived neurons (14), but also murine cells lacking neuroplastin showed reduced expression of PMCA (13).

The past four decades witnessed the development of models describing the differentiation of naïve T cells following antigen encounter into different compartments of effector and memory cells. These models are based on flow cytometric phenotypic analysis, and their outcome provides the basis for a nomenclature of subpopulations and describes function and persistence of populations arising during antigen encounter and clearance [for comprehensive reviews see (15–17)]. There are three main models describing memory cell generation. The first is based on epigenomic profiling and proposes linear development of naïve into stem central memory (TSCM) then CM, followed by effector memory (EM) and eventually effector (E) cells (15). The bifurcative model suggests that naïve cells asymmetrically divide to directly develop into E cells and in parallel an EM that gives rise to a CM lineage. The third model overlaps with the linear model describing memory cell generation. The name of the model (18). Despite the intensive efforts, the mechanisms regulating naïve-to-memory cell transition are not entirely understood. One important aspect of this transition is the transcriptional reprogramming of the cells (19, 20). Through iterative interactions, transcription factors selectively combine genes to be expressed to define cytokine production and survival potential profiles characteristic of cell populations. Ca²⁺-dependence is a common feature of many of the transcription factors involved in T cell differentiation such as NFAT (21, 22), CREB (23) and the transcription repressor DREAM (24). The main pathway for Ca²⁺ influx in T cells is the store operated Ca²⁺ entry (SOCE) constituted by the channel forming ORAI proteins (25, 26) and the activator ER-resident Ca²⁺ sensor STIM proteins (27–29). While Ca²⁺ levels influence the activity of transcription factors, conversely, the expression of SOCE components (30) and also expression of proteins involved in Ca²⁺ homeostasis are subjected to transcriptional control (31). Transcriptional control of PMCA is, however, poorly understood and whether this control is altered during naïve-to-memory cell transition is yet to be explored. Therefore, we set out in the current study to investigate how PMCA alters Ca²⁺ signals in T cells to promote activation and differentiation; and to gain insight into mechanisms regulating PMCA expression during T cell activation.

MATERIALS AND METHODS

T-Cell Isolation and In Vitro Polarization

Blood samples were collected from healthy donors in the Institute of Clinical Hemostaseology and Transfusion Medicine, Saarland University, Homburg. The research was approved by the local ethical committee (83/15; FOR2889-TP6, Niemeyer/Alansary), and blood donors provided their written consent. Following thrombocyte apheresis, peripheral blood mononuclear cells (PBMCs) were isolated from leukocyte reduction chambers (LRS, Trima Accel®) according to (32, 33). Lymphocytes where enriched by overnight incubation of PBMCs to remove adherent monocyte and macrophages. On the next day CD4⁺ cells were stained and naïve (CD4⁺CD127highCD25⁻CD45RO⁻) or memory (CD4⁺CD127highCD25⁻CD45RO⁺) cells were sorted using FACS Aria III (BD). Isolated CD4⁺ cells were seeded at density of 2 × 10⁶ cells/ml in AIMV medium containing 10% FCS and 1% Pen/Strep and cultured for at least 24 h before conducting the experiments. Where indicated, naïve cells were polarized in vitro into Th1 or Treg according to (34). Alternatively, where indicated, total CD4⁺ T cells were obtained from PBMC using negative isolation Kit (Miltenyi, #130-096-533) and automated cell isolation system (Miltenyi, AutoMACS).

Transfection

For downregulation of PMCA, 2 × 10⁵ cells were seeded in 96-well plate in Accell siRNA Delivery Media (ADM, Horizon) supplemented with 10 ng/ml of recombinant IL-2 and transfected with 1 µM accel siRNA targeting ATP2B4 (Accell Human ATP2B4 siRNA-SMARTpool, E-006118-00-0005, Horizon) or non-silencing control RNA (Accell Non-targeting Pool, D-001910-10-05, Horizon) according to the manufacturer’s instructions. Seventy-two hours later cells were harvested for mRNA or functional analysis.

Flow Cytometric Analysis

For flow cytometric analysis, cells were harvested, washed then stained with a viability dye (Zombie Aqua Fixable Viability Kit, Biolegend) followed by surface staining. Antibodies used for staining were supplied by Biolegend and are listed in Supplemental Table 2. Flow cytometric analysis was performed with FACSVerse (BD).

Single Cell Ca²⁺ Imaging

T cells were loaded in suspension with 1 µM Fura 2-AM in AIMV medium at room temperature for 20–25 min and seeded on polyornithine coated glass coverslips. All experiments were at room temperature as in (34). For stimulation of store operated Ca²⁺ influx 1 µM Thapsigargin (Tg) was used. Where indicated cells were treated 5 or 20 µM caloxin1 C2. Analyzed parameters include: maximum [Ca²⁺], (Peak), steady state [Ca²⁺], analyzed as the average [Ca²⁺], 25 s before removal of [Ca²⁺], (Plateau), and the fraction of retained [Ca²⁺] (Plateau/Peak) calculated for each cell independently. Analysis of the efflux rates was performed using Origin software. To test the influence of [Ca²⁺] on PMCA activity, cells were binned according to their peak or plateau [Ca²⁺], (iso-cells) to facilitate comparison of the corresponding efflux rates in the presence or absence of extracellular Ca²⁺, respectively.

Transcription Factors Binding Site Analysis

A 5,000 bp long sequence upstream to start codon of ATP2B4 was used as a potential promoter sequence for ATP2B4.
Prediction of transcription factor potentially regulating PMCA was done using the algorithms provided by three different data bases: ChIP-Seq (35), HOCOMOCO (36), and Promo V3 (37). Identification of transcription factor binding site (TFBS) was done by Emsembl Transcription Factor tool [Enembl Regulatory Build (38)], and making use of the matrix-based transcription factor binding site in JASPAR database (39). Results obtained from the independent data bases were subject to Venn analysis using the online tool Venny 2.1 (40) to detect common results, listed in Supplementary Table 1.

**Chromatin Immunoprecipitation**

To test binding of transcription factors to PMCA4b promoter region, antibodies against transcription factors (YY1 or NFAT, cell signaling, 3 µg each) or isotype control antibodies (rabbit IgG or mouse IgG, respectively) were coupled to protein A/G agarose beads (Santa Cruz) in coupling buffer (0.01 M sodium phosphate, 0.15 M NaCl, pH 7.2) overnight at 4°C then cross-linked to the beads using 5 mM BS3 (Thermoscientific). In parallel, human CD4+ T cells were harvested, washed then cross-linked with PBS then containing 5 mM DSP (Thermoscientific) and 3% PFA for 20 min at 4°C. Excess cross-linker and PFA were quenched by 125 mM Glycerine in PBS. Cell pellets were lysed in 20 mM Tris, 100 mM KCl, 10% glycerine (v/v), 0.5% n-Octyl-β-D-glucopyranoside (Merk Millipore). Finally lysates were subjected to chromatin shearing with Qsonica Sonicator Q700 (Thermoscientific) in total processing time of 4 min, 20 s laps of 40% amplitude and 10 s pauses. Clear lysates were diluted to 1 ml with binding buffer [20 mM Tris, 100 mM KCl, 10% glycerine (v/v)]. An aliquot of 50 µl was kept as an input control. Lysates were incubated with preclearing beads (isotype IgG bound beads) for 3 h at 4°C; unbound lysate was allowed to bind to control IgG (a fresh aliquot of isotype IgG bound beads) for 3 h at 4°C, and finally unbound lysate was bound to chromatin IP beads (transcription factor bound beads) for 3 h at 4°C. Beads collected at different steps were washed with standard low salt, high salt, LiCl buffer, and finally with TE according to Thermoscientific protocol. Finally, DNA was eluted with 1% SDS in 100 mM NaHCO3, purified using DNA purification kit (Qiagen), and used as template to test for binding of the PMCA4b promoter region using Q5 hot start polymerase (NEB biolabs) and the primers: forward 5′-CACACCCGTGCCAGCTAA-3′, reverse 3′-GAAGGGTGCTAGTTGGACA-5′. Amplicons were visualized by DNA agarose, and identity was confirmed by Sanger sequencing.

**Dual Luciferase Reporter Assay**

To estimate alterations in PMCA4 promoter activity, the region (Gene ID NG_029589.1, bp 663-1052) identified by ChIP assay to be potentially bound by YY1 was amplified with similar primers as above mentioned but adding SacI and NheI recognition sequences. The corresponding cDNA was subcloned into pGL3 basic (Promega, no. E1751) upstream to sequence encoding red firefly luciferase. Three days before conducting the assay, HEK293 cells were transfected with siRNA targeting YY1 (siYY1) (Accel Human YY1 siRNA SMART pool, no. E-011796-00-0010, Horizon) or with non-silencing control RNA (ns). Forty-eight hours later, cells were transfected with the created plasmid (PMCA Promoter, PP) or pGL3 basic (c) together with plasmid encoding green renilla luciferase under thymidine kinase promoter control (pTK-Green Renilla Luciferase, ThermoFisher Scientific, no. 16154) in a DNA ratio of 2:1. Finally, 24 h later, cells were stimulated with 5 ng/ml phorbol 12-myristate 13-acetate (PMA) and 500 ng/ml ionomycin for 6 h then lysed in passive lysis buffer commercially provided in the dual luciferase reporter assay kit (Promega, no. E1910). Assay was conducted according to the manufacturer’s instructions, and luminesence measurements were done using CLARIOstar (BMG, LABTEC) at gain of 3,600. For analysis, red firefly luciferase luminescence was normalized to the green renilla luciferase and data presented as average ± SEM of the relative change to the control condition of each independent experiment.

**RNA Isolation, cDNA Synthesis, and Quantitative Real-Time PCR**

The indicated cell types were harvested and stored at −80°C until RNA was isolated using RNeasy kit (Qiagen) following the manufacturer’s instructions. SuperScriptTMII Reverse Transcriptase (Life technologies) was used to generate complementary DNA (cDNA), and subsequent qRT-PCR was conducted using Quantitect SYBR Green Kit (Qiagen) and a CFX96 Real-Time System (Biorad). For quantification, threshold cycle (Cq) values of a gene of interest were normalized to that of either RNA polymerase or to TATA box binding protein (TBP) using the ΔCq method. For comparison of mRNA levels between two or more populations, fold change was calculated with 2−ΔΔCq method.

**Western Blot**

For protein expression analysis, cells were harvested, washed with PBS then lysed in buffer containing 20 mM Tris, 100 mM KCl, 10% glycerine (v/v), 0.5% n-Dodecyl beta Maltoside (DDM), pH 7.4. Lysates were stored at −80° until analyzed. Standard SDS-PAGE was performed followed by electrotransfer to nitrocellulose membranes. Immunoblots were probed with anti-Orai1 (Sigma, catalogue number O8264), anti-STIM1 (Protienech, # 11565-1-A), anti-STIM2 (Sigma, # S8572), anti-GAPDH (Cell Signaling, clone 14C10) anti-β-actin (Abcam, clone AC-15) or anti-PMCA4 (Thermoscientific, clone JA9) at a 1:1,000 dilution. For protein detection, an enhanced chemiluminescence detection reagent was used (Clarity Western ECL Substrate, Biorad). Densitometric quantification of detected protein bands was done with Quantity one software (Biorad).

**Statistical Analysis**

For analysis of Ca2+ imaging data, 100–250 cells were measured per experiment; one to two independent experiments were done per donor up to 14 donors. Otherwise, data is presented as mean ± S.E.M. Data was tested for normal distribution. When comparing two groups, statistical significance was tested by
performing unpaired, two-tailed Student t-test for normally distributed data sets and Mann–Whitney test when samples are not normally distributed or when the sample size was not sufficient to test for normality. Comparing more than two groups was done using Kruskal–Wallis one-way analysis of variance (ANOVA) or conventional ANOVA when data showed normal distribution and equal variance. Asterisks indicate significant differences for different p values as follows: * p < 0.05, ** p < 0.01, *** p < 0.001. Statistical analysis was performed using Graphpad Prism.

RESULTS

Differential Expression of PMCA4b Results in Distinct SOCE Profiles in CD4+ Naive and Memory Cells

In our previous work we characterized the SOCE profiles of antiCD3/CD28-activated cells, in vitro polarized into CD4+ T cell subtypes (Th1, Th2, Th17, and Treg) and observed that the control naïve and memory cells that were activated but cultured in non-polarizing conditions showed differential SOCE profiles with concomitant upregulation of PMCA in the activated memory cells (34). Therefore, we set out in the current work to investigate whether in resting cells PMCA plays a differential role in Ca2+ homeostasis and thus regulates the differentiation process of human CD4+ T cells. First, we measured SOCE profiles in resting naïve (CD4+CD127hiCD25lo) and memory (CD4+CD127hiCD25 hi) CD4+ T cells. We isolated the desired populations from human peripheral blood mononuclear cells (PBMCs) using the fluorescence activated cell sorting (FACSorting) strategy shown (Figure S1) where we excluded regulatory cells (Treg, CD4+CD127loCD25hi) and sorted the desired populations from conventional CD4+ T cells (CD4+CD127hiCD25lo). One day later, we measured changes of intracellular Ca2+ concentration following thapsigargin (Tg)-induced store depletion. The resulting SOCE profiles of naïve and memory human CD4+ cells were significantly distinct. Naïve cells showed a higher peak and plateau of [Ca2+]i than memory cells (Figures 1A, B). The observation that the peak [Ca2+]i of memory cells was reduced by 13.5% while the steady state plateau [Ca2+]i, and the fraction of retained Ca2+ (Plateau/Peak) were reduced by more than 40% each compared to naïve cells indicates that Ca2+ signals in memory cells exhibit faster decay kinetics. To determine whether the difference in naïve and memory profiles is due to differential expression of Ca2+ influx and/or extrusion mechanisms, we analyzed the expression of the relevant genes. First, we analyzed the expression levels of ORAI1–3 and STIM1–2, the main constituents of Ca2+ influx pathways in T cells. Analysis of mRNA (Figure 1C) and protein (Figures 1D, E) levels showed that both naïve and memory CD4+ cells express comparable levels of SOCE components. The specificity of the antibodies was validated by cells where ORAI1 (41), STIM1, or STIM2 (42) was deleted by CRISPR-Cas9 mediated gene targeting. Using commercially available antibodies for ORAI2 and ORAI3 proteins we were unable to reliably detect endogenous proteins. These results indicate that reduced [Ca2+]i, is likely not explained by reduced Ca2+ influx through ORAI channels and imply involvement of Ca2+ efflux mechanisms. Analysis of expressed isoform of plasma membrane Ca2+ dependent ATPase (PMCA, ATP2B1–4) revealed that the main isoforms expressed in CD4+ cells are PMCA1 (ATP2B1) and the more predominant PMCA4 (ATP2B4) in agreement with (12) while isoforms PMCA2 (ATP2B2) and PMCA3 (ATP2B3) were not detected (Figure 1F). Furthermore, using splice-specific primers, mRNA analysis showed that PMCA4b (ATP2B4b) is the main variant in CD4+ T cells (Figure 1G). Interestingly, memory cells showed a significantly higher expression level of PMCA4b (ATP2B4b) on both mRNA level (Figure 1H) and protein (Figures 1I, J) level, using an antibody with tested specificity [(43) and Figures S2A, B]. Moreover, we analyzed Ca2+ efflux rates in the presence (Figure 1K) and after removal (Figure 1L) of extracellular Ca2+, taking into consideration the Ca2+ dependence of PMCA activity (1, 2). Binning the cells according to [Ca2+]i allowed comparison of cells with similar [Ca2+]i, (iso-cells) and revealed that memory cells extrude Ca2+ with higher rates (Figures 1K, L) and shorter time constants (τ) (Figures S2C, D) compared to naïve cells independent of the cytosolic [Ca2+]i. Significant differences are observed at physiologically relevant concentrations but not at higher concentrations probably due to exceeding pumping capacity of PMCA but also due to the small fraction of cells exhibiting this concentration, which hampers statistical significance.

Pharmacological Inhibition or Downregulation of PMCA Reverses SOCE Phenotypes of Memory Cells

To directly test whether PMCA has a differential role in Ca2+ homeostasis in resting CD4+ T cell subsets, we tested the effect of pharmacological inhibition of PMCA by caloxin1C2 (C1C2) (44) on SOCE profiles of naïve and memory cells. Treatment with 5 µM or 20 µM caloxin1C2 only mildly altered SOCE profile of the naïve cells resulting in no or a slight increase that did not exceed 14% of the tested parameters (Figures 2A, B). On the other hand, memory cells treated acutely with the PMCA blocker showed a significantly increased SOCE resulting in a 58% increase of the plateau and 47% increase of the fraction of retained Ca2+ compared to untreated cells (Figures 2C, D). Noteworthy is that the application of 20 µM, a concentration shown to block all PMCA isoforms (44), did not result in additional effects on the plateau levels compared to the application of the PMCA4-specific inhibitory concentration of 5 µM (Figures 2C, D compare blue to gray traces and dots) indicating that PMCA4 is indeed the main functional isoform in T cells and in agreement with the expression analyses (Figures 1F–J).

In a second approach we transfected naïve or memory CD4+ T cells using an ATP2B4 specific siRNA (siPMCA) that resulted in significant downregulation on RNA (Figure S2A) and protein levels (Figure S2B). Similar to the pharmacological manipulation by C1C2, downregulation of PMCA4 led to no or subtle alteration of analyzed parameters of SOCE profile of the naïve cells (Figures 3A, B). Memory cells treated with siPMCA
showed a 15% increased influx peak, while the plateau was increased by 49% compared to memory cells transfected with non-silencing RNA (ns) resulting in significant increase in fraction of Ca\(^{2+}\) retained in the cells (Plateau/Peak ratio) (Figures 3C, D). These results indicate a differential role of PMCA in Ca\(^{2+}\) homeostasis of naïve and memory cells.
In Vivo Developed Effector and Memory CD4+ Cellular Compartments Differentially Express PMCA4 and Exhibit Corresponding SOCE Profiles

The findings presented so far compared the Ca2+ signals and expression profiles of naïve and memory cells. Memory cells (CD4+CD45RO+), however, represent a heterologous population that is commonly divided into different compartments depending on effector and proliferative capacities of the cells. Therefore, our next aim was to answer the question whether in vivo developed memory compartments show distinct PMCA expression profiles or do they homogenously share higher expression levels of PMCA. To address this question, we FACSorted CD4+ T cell compartments from human PBMCs based on the surface markers (CD45RO/CCR7) as illustrated in (Figure 4A). Purity of the sorted populations was tested by post-sorting staining (Figure 4B). Cells that were CCR7−CD45RO−CD45RA+ are defined as naïve cells (N), CCR7+CD45RO+CD45RA− as central memory cells (CMs), CCR7+CD45RO+CD45RA− as effector memory (EM) and CCR7+CD45RO−CD45RA+ as terminally differentiated effector memory cells (EMRA) (45–47). TEMRA cells are a poorly characterized population of T cells expressing CD45RA but are commonly used due to the sharing functional properties with the short lived effector cells (15, 48, 49). For further characterization, we measured the expression of IFNγ which showed the expected pattern where the EMRA and EM populations had higher expression of IFNγ than the CM while it was lacking in naïve cells (Figure 4C).

Interestingly, analysis of PMCA expression in the sorted populations revealed differential expression profiles: with the terminally differentiated effector (EMRA) and EM cells showing the higher expression than CM and naïve cells (Figure 4D). Finally, we tested whether SOCE profiles of the in vivo differentiated cellular compartments recapitulate the differentially expressed PMCA levels. Indeed, measurements of Tg-induced Ca2+ influx showed that CD4+ compartments have distinct SOCE profiles with comparable Ca2+ peaks (Figures 4E, F) but distinct increase in decay rates resulting in the lowest plateau [Ca2+]i and retained Ca2+ fractions for the effector cells (EMRA and EM), intermediate levels for CM cells while the naïve cells have the highest plateau [Ca2+]i and retained Ca2+ fractions (Figures 4E, F) in line with the observed PMCA expression levels (Figure 4D). These differences are underlined by parallel differences in the efflux rates in presence of (Figure S3A) and after removal (Figure S3B) of extracellular Ca2+ indicating that the differences in the plateau and retained Ca2+ fractions are indeed due to differential expression and function of PMCA.

PMCA4 Regulates Compartment Stoichiometry Upon Activation

The outcome of an immune response depends on the dynamics of mobilization of T cells within naïve and memory...
compartments. Furthermore, our findings show that CD4+ cellular compartments exhibit differential SOCE profiles and PMCA expression levels. Therefore, we sought to address if and how regulation of [Ca2+]i by PMCA4 contributes to fate-decision making concerning compartment distribution of CD4+ T cells upon activation. To this end, we isolated naïve CD4+ T cells from PBMCs to exclude effects of previous antigen exposure and on the following day transfected naïve cells with non-silencing RNA (ns) or with siRNA targeting ATP2B4 (siPMCA). Three days later, naïve cells were replenished with fresh siRNA, subjected to stimulation using antiCD3/CD28 coated beads, and monitored for compartment distribution 24 h later. The experimental design and the surface markers used to assign activated cells into different compartments are shown in Figure 5A. Downregulation of ATP2B4 did not alter stoichiometry of cellular compartments as defined by CD45RO or CD45RA and CCR7 (Figures S3C, D). Because, classically, T cell compartments are defined based on CD45RO, CD45RA, CCR7, and CD62L (45–47), and CD62L is commonly used as an alternative marker for CCR7 for compartment definition (50, 51), we analyzed compartment stoichiometry based on CD45RO and CD62L expression. Interestingly, this analysis showed that downregulation of ATP2B4 resulted in a significant decrease of the fraction of cells in terminally differentiated effector memory CD45RA+ compartment T_{EMRA} (CD4+CD62L−CD45RO−), while the fraction of cells remaining in the naïve compartment (CD4+CD62L+CD45RO+) was significantly increased compared to cells treated with non-silencing RNA (Figures 5B, C).

The effects on EM and CM compartments showed similar tendencies as the effects observed on EMRA and naïve compartments, respectively, albeit not significant (Figures 5B, C).

To test whether downregulation of PMCA4 is accompanied by altered potential of naïve CD4+ cells to be activated, we measured the expression level of activation markers CD69 and CD154 under the same experimental conditions as above. Despite the reduced fraction of effector cells, treatment of naïve cells with siPMCA resulted in increased expression of activation marker CD154 (CD25) (Figures 5D, E), while expression of CD25 was not altered (Figure S3E). These results imply that PMCA4 downregulation does not inhibit activation per se but rather regulates the compartment stoichiometry.

Transcriptional Control of PMCA Results in Biphasic Expression Following Activation

The experiments described above provide evidence that PMCA4 expression level correlates with the cellular compartment of CD4+ T cells and contributes to regulation of compartment stoichiometry. To explore potential underlying mechanisms, we set out to investigate transcriptional regulation of PMCA expression. To this end, we explored potential transcription factor binding sites (TFBSs) in the promoter region of PMCA (ATP2B4) using available ChIP seq information (35) combined with TFBS prediction data bases HOCOMOCO (36) and Promo 3 (37, 52). The resulting analyses (Figure 6A) predicted 33 transcription factors (listed in Supplementary Table 1) that were detected in all three algorithms with the transcription
factor YY1 having the highest predicted score. Consequently, we explored whether we can experimentally confirm binding of YY1 to the promoter region of PMCA. Because our preliminary experiments showed that a conventional protocol of chromatin immune precipitation (ChIP) experiments was prone to misleading positive results, we applied a rigorous protocol that eliminates all unspecific binding using isotype control antibody-bound beads in a "pre-clearing" step (Figure 6B first lane). The efficiency of elimination of unspecific binding was demonstrated by applying the pre-cleared lysate to fresh isotype control antibody-bound beads (Figure 6B second lane). Finally, the ability of YY1 to bind to the promoter region was tested by using a YY1-specific antibody to immunoprecipitate YY1 and the bound genomic DNA which was then amplified with specific primers, designed based on the predicted binding domains. In parallel, we used an anti-NFAT antibody to serve as an additional control for the specificity of binding. Results of the ChIP experiments showed that YY1, but not NFAT, was able to bind the promoter region of PMCA4 as visualized on agarose gels (Figure 6B). The identity of the amplicon was further confirmed by sequencing.

To investigate whether the levels of YY1 are altered in CD4+ cell compartments similar to PMCA, we analyzed expression levels in naïve and memory CD4+ cells. Surprisingly,
expression level of YY1 was comparable to the resting naïve and memory CD4+ cells (Figures 6C, D). We hypothesized that this finding might be due to performing the analysis in resting cells where YY1 expression is adapted to steady state levels. To test this hypothesis, we stimulated total CD4+ cells for 24 h using antiCD3/CD28 coated beads, and indeed we observed an increased level of YY1 concomitant to decreased expression of PMCA (Figures 6E, F), indicating that YY1 represses PMCA levels upon stimulation, resulting in reduced Ca2+ clearance compared to unstimulated cells (Figures S4A, B). Because PMCA levels were decreased after acute stimulation (Figures 6E, F) contrary to the increased levels we observed in resting memory compared to naïve cells (Figures 1H–J), we extended the expression analysis to examine whether YY1 alters PMCA4 expression in association with long term stimulation and full differentiation of T cells. To this end, we subjected the naïve cells to in vitro polarization protocol providing stimulatory, co-stimulatory, and cytokine mediated signals necessary for differentiation of naïve cells into Th1 or Treg cells (34). This experiment not only aims to monitor changes of PMCA and YY1

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**FIGURE 5** | Downregulation of PMCA alters compartment stoichiometry upon activation of naïve CD4+ T cells. (A) Schematic representation of the experimental design: naïve cells were isolated from human PBMCs using FACSorting, transfected one day later with control non-silencing RNA (ns) or with siRNA targeting PMCA4 (siPMCA), stimulated with antiCD3/CD28 coated beads 72 h following transfection and 24 h later analyzed for compartment distribution using flow cytometry and the shown surface markers. (B) Representative flow cytometric analysis images showing compartment distribution in resting naïve cells (left image) or naïve cells transfected with control non-silencing RNA (ns, middle image) or transfected with siRNA targeting PMCA4b (siPMCA, right image) then activated following time scheme shown in (A). (C) Frequency of cells (% Cells) measured in the EM (CD62L−CD45RO+), CM (CD62L+CD45RO+), N (CD62L+CD45RO−) and EMRA (CD62L−CD45RO−) compartments following treatment with non-silencing RNA (ns, black) or siRNA targeting PMCA4 (siPMCA, blue) according to time scheme shown in (A). (D) Representative flow cytometric analysis images showing expression of the activation markers CD154 and CD69 in cells treated as in (A, B). (E) Frequency of cells (% Cells) measured in (D) and expressing activation markers following treatment with non-silencing RNA (ns, black) or siRNA targeting PMCA4b (siPMCA, blue) according to time scheme shown in (A). Data represent average ± s.e.m obtained from eight independent experiments. Asterisks indicate significance at *p < 0.05, **p < 0.01 using Kruskal–Wallis one-way analysis of variance (ANOVA).
expression levels upon longer stimulation but also monitors correlation of PMCA expression to developing effector functions. The polarized cells showed the expected differential expression of IFNG (Figure S4C) and FOXP3 (Figure S4D), the hallmarks of Th1 and Treg populations, respectively. Importantly, terminally differentiated effector cells (Th1, Treg) showed a significant upregulation of PMCA4, while the control cells which were stimulated in non-polarizing conditions (Th0)
underwent milder upregulation of PMCA (Figures 6G, H and Figure S4E). Strikingly, levels of YY1 followed a reverse pattern: while YY1 underwent transient upregulation within 24 h of activation (Figures 6E, F) YY1 expression decreased upon longer activation (Figures 6G, I and Figure S4F), in line with the equal levels analyzed in resting cells (Figures 6C, D).

Finally, we aimed to confirm that altered expression of PMCA is due to direct regulation by YY1. To achieve this correlation, we constructed a plasmid expressing red firefly luciferase under transcriptional control of the identified PMCA promoter region where YY1 is predicted to bind (Figures 6A, B). To facilitate analysis of promoter activity independent of transfection efficiency, the red firefly luciferase activity was normalized to the co-transfected green renilla luciferase expressed under the transcriptional control of TK promoter. To address whether the PMCA promoter activity is directly regulated by YY1, we downregulated YY1 using siRNA (Figures S4G, H) then conducted the dual luciferase assay (Figures 6J). Measurements (Figure 6K) show that in cells transfected with non-silencing RNA (ns), the PMCA promoter (ns + PP) was able to induced expression of the firefly luciferase compared to control cells (ns + φ). More importantly, the PMCA promoter activity is significantly increased following downregulation of YY1 (siYY1 + PP). Collectively, these results point towards YY1 being part of the transcriptional factor complex regulating PMCA promoter activity to allow T cells to adapt the expression of PMCA4 to fulfill cellular requirements of Ca^{2+} during different phases of an immune response.

**DISCUSSION**

In the current work we investigated how Ca^{2+} signals are fine-tuned during activation of CD4{sup}+ T cells to regulate the stoichiometry of CD4{sup}+ T cell compartments and thus the outcome of an immune response. Using fura-2 Ca^{2+} imaging we show that quiescent memory cells accommodate a lower [Ca^{2+}], and are able to clear the Ca^{2+} signals faster than their naïve counterpart parts (Figures 1A, B). Comparable levels of expressed SOCE genes (Figures 1C–E) but significantly more expressed PMCA4b (Figures 1F–J) and higher efflux rates independent of [Ca^{2+}], (Figures 1K, L and Figures S2C, D) present strong evidence that SOCE phenotype of memory cells is mostly a result of more efficient clearance machinery. Furthermore, analysis of the contribution of PMCA to Ca^{2+} homeostasis in naïve and memory cells by applying an acute pharmacological inhibition approach (Figure 2) as well as by directly downregulating ATP2B4 (PMCA4) (Figure 3) presents evidence that clearly supports the hypothesis that PMCA4 plays a functionally more significant role in Ca^{2+} clearance in memory compared to naïve CD4{sup}+ T cells. Further analysis of the CD4{sup}+ T cell subpopulations revealed that the quiescent cells in the naïve and central memory compartments express less PMCA and exhibit slower Ca^{2+} signal decay kinetics than compartments associated with more effector functions (EMRA and EM) (Figure 4). In vitro generation of effector populations (Th1, Treg) able to express effector hallmarks (Figures S4C, D) and the concomitant significant increase in PMCA (Figures 6G, H) strongly support the hypothesis that fully differentiated effector cells acquire higher expression of PMCA to guard against apoptosis resulting from activation induced [Ca^{2+}], overloads. Noteworthy is that, before cells reach full differentiation state, PMCA levels transiently decrease after short term stimulation (Figures 6E, F) allowing [Ca^{2+}], levels required to fulfill cellular demands during initial proliferation.

CD4{sup}+ T cell differentiation is a complex process in which a plethora of genes are rigorously orchestrated, and we hypothesized that PMCA4 is an important regulator of this process. To gain a direct insight into the regulatory role of PMCA4 in CD4{sup}+ T cell differentiation, we monitored differentiation and compartment distribution upon activation of naïve cells while downregulating PMCA4 (siPMCA). We found that siPMCA-treated cells exhibit a skewed compartment stoichiometry following activation with more cells remaining in the naïve compartment parallel to a decreased fraction of cells able to reach the effector compartments (EM and EMRA) (Figures 5B, C). Interestingly, although CD62L and CCR7 are often co-expressed and are commonly considered equivalent for population definition (46, 50), we observed an effect of PMCA4 downregulation on the expression of CD62L (Figures 5B, C) but not of CCR7 (Figure S3C). This might be explained by the different kinetics of expression of both molecules where activated T cells rapidly shed CD62L of their surface and gain new set of selectins which allow them to migrate to non-lymphoid tissues (53). Moreover, a CD62L CCR7{sup}+ population has been identified as a frequently existing memory population (54) and was detected in cancer studies where it has been shown to have homing and effector functions that are distinct from conventional CM and EM, though much less understood (55). These results implicate an importance of PMCA4 in regulating early events of activation.

Our analysis of the effect of downregulation of PMCA on activation markers revealed differential effects. Transcriptional upregulation of the activation marker CD69 is triggered immediately (30–60 min) upon activation and reaches measurable levels already after 2–3 h (56) then declines within 6 h (57). These fast kinetics could indicate that CD69 is regulated by mechanisms independent of PMCA and [Ca^{2+}], and thus explain our finding that total frequency of cells expressing CD69 was not altered by siPMCA treatment (Figures 4D, E). On the other hand, downregulation of PMCA4 resulted in a significant increase of fraction of cells expressing CD154 (CD40L). CD154 belongs to the TNF gene family, and its upregulation follows a biphasic pattern with the first phase being TCR dependent and taking place within the first 24 h after activation. The later phase is dependent on the cytokine composition in the cellular milieu (58, 59). Furthermore, CD154 transcription is enhanced by Ca^{2+}, -NFAT pathway (60, 61) which together with an increased NFAT activity upon downregulation of PMCA4 (62) might explain the increase observed in CD154 expression in siPMCA-treated cells (Figures 4D, E). Together, these findings
support our hypothesis that PMCA is essential for regulating differentiation processes in functional CD4+ T cells. By virtue of their ability to activate, repress, or modify gene expression, transcription factors regulate cell division, differentiation, and function (63). Transcriptional control of PMCA isoforms has been addressed (64, 65) but remains to time, however, poorly understood. Here we applied in silico analyses (Figure 6A) and identified YY1, a ubiquitously expressed transcription factor (66), to be likely involved in the modulation of PMCA4 expression. Indeed, we confirmed that YY1 binds specifically to a predicted sequence in the PMCA4 promoter region (Figure 6B). Furthermore, our results suggest a repressive effect of YY1 on PMCA expression, as indicated by the inverse correlation between expression of PMCA4 and that of YY1 over the course of differentiation (Figures 6E–I) and more importantly, by the increased PMCA promoter activity upon downregulation of YY1 (Figures 6J, K). There are several lines of evidence for the interaction between YY1 and c-myc (67–69), c-Myc inhibits the DNA binding capacity of YY1 (69) in a direct, binding-independent (67) but c-Myc concentration dependent manner (69). Moreover, Habib and co-workers have shown that in B-cells c-Myc regulates differentiation and function by mechanisms including downregulation of PMCA (64). Therefore, we hypothesize that PMCA4 is regulated by a complex of transcription factors including c-Myc and YY1 (Zeller et al., 2006, Habib et al., 2007). If and how individual members of this complex have reciprocal effects needs further investigation. By regulating PMCA4 levels and thereby shaping Ca2+ signals, the activities of calcineurin and NFAT, and thus T cell differentiation are modulated. With an upregulation approach, the importance of PMCA to clear near plasma membrane Ca2+ thus allowing for NFAT activation was also demonstrated (70). Our results suggest a scenario in which another layer of transcriptional control regulates T cell differentiation and activity in which upstream to the PMCA-Calcineurin/NFAT line, expression of PMCA is regulated by YY1/c-Myc. Furthermore, it is plausible to speculate that the net outcome of transcriptional regulation depends on the expression of the individual components of such a transcription factor complex which makes it difficult to establish direct causal relationships in studies addressing a single transcription factor.

Together, our data suggest a model in which PMCA4 plays a pivotal role in shaping Ca2+ signals in different activation states of CD4+ T cells. This role is under a biphasic transcriptional regulation by YY1. The first phase starts once CD4+ T cells are activated where YY1 is upregulated and represses PMCA expression so that CD4+ T cells can accommodate higher cytosolic [Ca2+]i levels to support gene expression and proliferation. In the second phase, the initial repression by YY1 is alleviated, and PMCA expression increases to guard against [Ca2+]i overloads and promote survival of effector cells and formation of long lasting memory cells that are able to efficiently mediate faster immune response upon subsequent antigen encounter.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary Material.

ETHICS STATEMENT

Blood samples were collected from healthy donors in the Institute of Clinical Hemostaseology and Transfusion Medicine, Saarland University, Homburg. Research was approved by the local ethical committee (83/15; FOR2289-TP6, Niemeyer/Alansary), and blood donors provided their written consent. The patients/participants provided their written informed consent to participate in this study.

AUTHOR CONTRIBUTIONS

All authors conceptualized the study and edited the manuscript. MM-W and DA performed the experiments and data analysis. BN and DA secured funding and resources. DA wrote the manuscript. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fimmu.2021.687242/full#supplementary-material
Supplementary Figure 2 | siRNA-mediated downregulation of PMCA and analysis of time constants of Ca2+ efflux rates of naïve and memory cells (A) Quantitative RT-PCR analysis showing relative expression level of PMCA4 in naïve (black) and memory (red) cells transfected with non-silencing RNA (ns) or with siRNA targeting PMCA4 (siPMCA). (B) Representative WB showing PMCA4 expression in total CD4 T cells transfected with non-silencing RNA (ns) or with siRNA targeting PMCA4 (siPMCA). (C) Analysis of time constants (Tau) of Ca2+ efflux rates in presence of [Ca2+]o in cells measured in Figure 1A as function of peak [Ca2+]i. (D) Analysis of time constants (Tau) of Ca2+ efflux rates after removal of [Ca2+]o in cells measured in in Figure 4E as function of plateau [Ca2+]i.

Supplementary Figure 3 | Ca2+ efflux rates in CD4 compartments and effects of downregulation of PMCA on surface markers. (A) Analysis of Ca2+ efflux rates in presence of [Ca2+]o in cells measured in Figure 4E as function of peak [Ca2+]i (B) Analysis of Ca2+ efflux rates after removal of [Ca2+]o in cells measured in in Figure 4F as function of plateau [Ca2+]i. (C–E) Average fractions of cells expressing the indicated surface markers as obtained by flow cytometry analysis of naïve cells transfected with non-silencing RNA (ns, black) or with siRNA targeting PMCA4 (siPMCA, blue) for 72 h then stimulated with antiCD3/CD28 coated beads for further 24 h as in Figure 5.

Supplementary Figure 4 | SOCE profile and expression analysis of genes of interest in activated cells (A) Average traces showing changes of [Ca2+]i over time in response to changes of extracellular solutions containing the shown [Ca2+]o indicated in the bar above the traces. SOCE was measured in resting total human CD4+ cells (Unstim., black) or following a 24-h antiCD3/CD28 coated bead stimulation (24h., red). (B) Bar graphs showing analyzed parameters of SOCE measured in (A), (C–E) Quantitative RT-PCR analysis of NFAT (C), FOXP3 (D), ATP2B4 (E) and YY1 (F) measured in resting naïve CD4+ T cells (naïve) or cells cultured for 7 d under control conditions (Th0) or conditions to polarize cells into Th1 or Treg. (G) Quantitative RT-PCR analysis showing relative expression level of YY1 in total CD4+ T cells transfected with non-silencing RNA (ns) or with siRNA targeting YY1 (siYY1). (B) Representative WB showing YY1 expression in total CD4+ T cells transfected with non-silencing RNA (ns) or with siRNA targeting YY1 (siYY1).

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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