T Cell Receptor-induced Nuclear Factor κB (NF-κB) Signaling and Transcriptional Activation Are Regulated by STIM1- and Orai1-mediated Calcium Entry*

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T cell activation following antigen binding to the T cell receptor (TCR) involves the mobilization of intracellular Ca\(^{2+}\) to activate the key transcription factors nuclear factor of activated T lymphocytes (NFAT) and NF-κB. The mechanism of NFAT activation by Ca\(^{2+}\) has been determined. However, the role of Ca\(^{2+}\) in controlling NF-κB signaling is poorly understood, and the source of Ca\(^{2+}\) required for NF-κB activation is unknown. We demonstrate that TCR- but not TNF-induced NF-κB signaling upstream of IκB kinase activation absolutely requires the influx of extracellular Ca\(^{2+}\) via STIM1-dependent Ca\(^{2+}\) release-activated Ca\(^{2+}\)/Orai channels. We further show that Ca\(^{2+}\) influx controls phosphorylation of the NF-κB protein p65 on Ser-536 and that this posttranslational modification controls its nuclear localization and transcriptional activation. Notably, our data reveal that this role for Ca\(^{2+}\) is entirely separate from its upstream control of IκBα degradation, thereby identifying a novel Ca\(^{2+}\)-dependent distal step in TCR-induced NF-κB activation. Finally, we demonstrate that this control of distal signaling occurs via Ca\(^{2+}\)-dependent PKCα-mediated phosphorylation of p65. Thus, we establish the source of Ca\(^{2+}\) required for TCR-induced NF-κB activation and define a new distal Ca\(^{2+}\)-dependent checkpoint in TCR-induced NF-κB signaling that has broad implications for the control of immune cell development and T cell functional specificity.

Activation of T cells following antigen binding to the T cell antigen receptor (TCR) induces diverse lineage- and fate-specific proinflammatory and immune-modulatory responses. Central to these responses is the induction of quantitatively distinct intracellular Ca\(^{2+}\) signals and their selective activation of the key transcription factors NFAT and NF-κB (1–6). The mechanism by which Ca\(^{2+}\) controls NFAT activation in lymphocytes is well established (7). In contrast, although Ca\(^{2+}\) has been implicated in TCR-induced NF-κB signaling (8–10), how Ca\(^{2+}\) regulates NF-κB activity is largely unexplored and represents a significant gap in our understanding of transcriptional control of T cell development, activation, and functional specificity.

In resting T cells, classical NF-κB consists of heterodimers of p50/p65 or p50/c-Rel that are retained in the cytosol by members of the inhibitory family of IκB proteins (11, 12). Following TCR engagement, IκB kinase (IKK)-mediated phosphorylation triggers the ubiquitination and proteosomal degradation of IκBα, releasing p50/p65 and p50/c-Rel, which localize to the nucleus to initiate transcription of crucial immune-regulatory, proinflammatory, and prolipherative genes (13–30). Although TCR-mediated Ca\(^{2+}\) mobilization has been implicated in proximal steps of NF-κB activation (8–10), the precise mechanisms and source of Ca\(^{2+}\) that regulate nuclear localization and transcriptional activation of NF-κB are poorly defined. It is well established that TCR signaling induces inositol 1,4,5-trisphosphate-mediated depletion of Ca\(^{2+}\) from the endoplasmic reticulum (ER). A resulting Ca\(^{2+}\) dissociation from the ER membrane protein stromal interaction molecule 1 (STIM1) triggers its oligomerization and relocation to ER membrane domains juxtaposed to the plasma membrane (31–33), where STIM1 physically gates Orai (also known as Ca\(^{2+}\)-release-activated Ca\(^{2+}\)) channels, allowing extracellular Ca\(^{2+}\) to enter the cell (34, 35). However, it is not known whether Ca\(^{2+}\) control of TCR-induced NF-κB signaling requires STIM1- and Orai1-mediated Ca\(^{2+}\) influx or whether the initial release of Ca\(^{2+}\) from the ER is sufficient for classical NF-κB activation.

In this study, we sought to determine both the source and mechanism of Ca\(^{2+}\) control of antigen receptor-induced NF-κB activation in T cells. We show that influx of extracellular Ca\(^{2+}\) via STIM1 and Orai is critical for TCR- but not TNF-induced IκBα degradation and NF-κB activation. Importantly, we also demonstrate that Ca\(^{2+}\)-dependent, PKCα-mediated phosphorylation of p65 critically regulates its nuclear localization and transcriptional activation following TCR engagement. Thus, our findings define important new proximal and distal Ca\(^{2+}\)-dependent checkpoints in TCR-induced NF-κB signaling that have broad implications for the control of immune cell development and functional specificity.
Novel Calcium-dependent Mechanisms of NF-κB Activation

Materials and Methods

Cells and Cell Culture—Primary human T cells were obtained from the University of Pennsylvania Immunology Core facility. Jurkat T cells were from the ATCC, and Jurkat T cells stably expressing E106A Orai1 were a gift from Dr. Jonathan Soboloff (Temple University, Philadelphia, PA). All cells were cultured in RPMI 1640 medium (BioWhittaker, Walkersville, MD) supplemented with 10% heat-inactivated fetal bovine serum (Hyclone, Thermo Scientific, Logan, UT), 2 mM l-glutamine, penicillin (50 units/ml), and streptomycin (50 units/ml).

Antibodies and Reagents—Antibodies recognizing p65-Ser536 (3033S) and IκBα (48145S) were purchased from Cell Signaling Technology (Danvers, MA). Anti-p65 (372R/G) was purchased from Santa Cruz Biotechnology (Dallas, TX). Anti-α-tubulin (T1568) was from Sigma-Aldrich (St. Louis, MO). Anti-STIM1 (610954) was from BD Transduction Laboratories (Franklin Lakes, NJ). Anti-human TCR (c305 clone) was a gift from Dr. Gary Koretzky (Cornell University, New York, NY), and anti-CD28 was from Invitrogen. Protein A horseradish peroxidase-conjugated antibody (18-160) was from Millipore (Danvers, MA). Alexa Fluor 488 goat anti-rabbit (A11008) and Alexa Fluor 564 goat anti-rabbit (A11010) used for immunofluorescence were obtained from Invitrogen. Recombinant human TNF-α was purchased from R&D Systems (Minneapolis, MN). PMA, and ionomycin were from Sigma-Aldrich, and the luciferase reporter assay system was obtained from Promega (Fitchburg, WI).

Plasmids and Transfections—A cDNA construct expressing full-length p65 N-terminally tagged with EGFP was obtained from Addgene (Cambridge, MA). Mutant p65 constructs were generated using a site-directed mutagenesis kit (Stratagene, La Jolla, CA) to convert serine 536 to alanine or aspartic acid. Short hairpin STIM1 suppression and rescue constructs were generated in the laboratory of Dr. Dan Billadeau (Mayo Clinic, Rochester, MN), as were EGFP-shPKCα and EGFP-pCMS2 (control vector). For transfection, Jurkat T cells were suspended at 20 million cells/ml in RPMI 1640 medium, and 10 million cells were electroporated with 10 μg of DNA (for overexpression or mutant expression) or 40 μg of DNA (for suppression assays) at 315 V for 10 ms using a BTX ECM 830 electroporator (Harvard Apparatus, Holliston, MA). STIM1 and PKCα suppression assays were performed 48 h post-transfection, and EGFP-p65 and p65 mutant expression assays were performed 16–24 h post-transfection.

Immunoblotting—Cells were harvested and lysed without Non-ident P-40 lysis buffer consisting of 50 mM Tris-HCl (pH 7.5), 20 mM EDTA, 1% Nonident P-40, and complete inhibitors (1 mM sodium orthovanadate, 1 mM PMSF, 10 μg/ml leupeptin, and 5 μg/ml aprotinin). Protein concentrations in cell lysates were determined using Bio-Rad reagent and quantified in a Cary 50 Bio UV-visible spectrophotometer. Proteins were resolved by SDS-Polyacrylamide gel electrophoresis (4–15%, Bio-Rad) and then transferred onto PVDF membranes (Millipore, Billerica, MA). Membranes were probed with the respective primary anti-human antibodies and then incubated with protein A HRP secondary antibodies. Blots were developed with enhanced chemiluminescence using Pierce ECL Western blotting substrate. All immunoblots presented are from a single experiment representative of at least three independent experiments.

Luciferase Reporter Analysis—Luciferase-based transcriptional analysis was performed on Jurkat T cells transfected with 2 μg of total DNA (PBXII κB firefly luciferase and pRL TK Renilla luciferase in a 20:1 ratio) per transfection (5 × 10^6 cells in 500 μl of medium) using a square-wave BTX electroporator at 315 V for 10 ms. Twenty-four hours after transfection, cells were treated with PMA (200 nM), PMA (200 nM) and ionomycin (1 μg/ml), anti-TCR (0.5 μg/ml), and anti-CD28 (1:50) or TNF (10 ng/ml) for 4 h. Cells were then lysed in passive lysis buffer (Promega), and luciferase activity was measured using a Luminoscan 96-well automated luminometer (Thermo LabSystems, Franklin, MA). Firefly/Renilla luciferase ratios were calculated using Ascent software (Thermo LabSystems), and the mean ratio from at least three independent experiments (3–4 replicates/experiment) for each condition was compared.

Quantitative Real-time PCR—To quantify IκBα expression, cDNA was synthesized from RNA isolated (RNeasy Plus mini kit, Qiagen) from PMA- or PMA and ionomycin-stimulated cells with a high-capacity cDNA reverse transcription kit (Applied Biosystems, Foster City, CA). cDNA was amplified with IκBα (forward, 5’-CCGAGAC TTTCGAGGA AAT-ACC-3’; reverse, 5’-ACGTGTGGCCATTGAGT-3’), and β-actin-specific primers (forward, 5’-TCAGCAACGAGGAGTATGACGAG-3’; reverse, 5’-ATTGTAACCTTGGGGATGC-3’) on a 7500 Fast real-time PCR system (Applied Biosciences, Warrington, UK) using Power SYBR Green PCR Master Mix (Applied Biosciences). Ct values were obtained in triplicate for each target and analyzed with instrument software v1.3.1 (Applied Biosystems).

Microarray Analysis—RNA was isolated using an RNeasy Plus kit (Qiagen). Biotin-labeled cRNA was generated using the Illumina TotalPrep RNA amplification kit, and a Bioanalyzer (Agilent Technologies, Wilmington, DE) was used to assess total RNA and cRNA quality. Illumina HumanHT-12 version 4 expression bead chips were hybridized with cRNA from two biological replicates per condition and scanned on an Illumina BeadStation 500GX. Scanned images were converted to raw expression values using GenomeStudio v1.8 software (Illumina). Data analysis was performed using the statistical computing environment R (v3.2.3), the Bioconductor suite of packages for R, and RStudio (v0.98). Raw data were background-subtracted, variance-stabilized, and normalized by robust spline normalization using the Lumi package (36). Differentially expressed genes were identified by linear modeling and Bayesian statistics using the Limma package (37, 38). Probe sets that were differentially regulated (≥1.5-fold change between all treatments, false detection rate ≤ 5% after controlling for multiple testing using the Benjamini-Hochberg method (39, 40)) were used for heatmap generation in R. Clusters of co-regulated genes were identified by Pearson correlation using the hclust function of the stats package in R. Differentially expressed genes were identified using a list of validated and putative NF-κB target genes curated by the laboratory of Dr. Thomas Gilmore at Boston University. All microarray data have been deposited in the GEO database for public access (GSE76804).
Chromatin Immunoprecipitation—Jurkat T cells (10 × 10^6) transfected with either EGFP-shPKCα or control EGFP-pCMS2 vector (48 h) were stimulated with PMA (200 nM) and ionomycin (1 μM) for 30 min at 37 °C. Chromatin was prepared using a Covaris truChIP chromatin shearing kit (Covaris Inc., Woburn, MA). Briefly, cells were fixed in 1% methanol-free formaldehyde for 5 min at room temperature, and then fixation was quenched with 0.125 M glycine at room temperature for 5 min. Cells were washed twice with cold PBS and then lysed at 4 °C with rocking for 10 min. Nuclei were then washed and transferred to an AFA milliTUBE for ultrasonication. Samples were sheared using a Covaris S200 focused ultrasonicator for 1500 s in a 6 °C bath at a duty cycle of 5%, an intensity of 4, a peak incident power of 140 W, and at 200 cycles/burst. p65 was precipitated from sheared chromatin (200–1000 bp) with anti-p65 (5 μg, Santa Cruz Biotechnology, rabbit 372X) or normal rabbit IgG (5 μg, Cell Signaling Technology, 2729) for 12–16 h at 4 °C. Immunoprecipitated chromatin was then incubated with protein G Dynabeads (Life Technologies) for 2 h at 4 °C, and chromatin was eluted (50 mM Tris (pH 8.0) and 10 mM EDTA) at 65 °C on a thermomixer (1200 rpm) for 30 min. Cross-linking was reversed by incubating recovered chromatin at 65 °C for 12 h, followed by incubation with RNase A for 2 h at 37 °C and proteinase K for 30 min at 55 °C. DNA was then purified using a ChIP DNA clean and concentrator kit (Zymo Research, Irvine, CA), and quantitative PCR was performed to quantify p65 binding to IkBα, CXCL8, and TNF promoters using IkBα-specific (forward, 5′-TTGGGATCTCAGCAGCCC-GAC-3′; reverse, 5′-GCCACTAGGGTCACGGACAG-3′), CXCL8-specific (forward, 5′-CAGGTGTCCCTGGAGGGATG-3′; reverse, 5′-GGAGTGTCCTCGGTGTTT-3′), and TNF-specific (forward, 5′-GCCGGCATGGGAAAGAAACC-3′; reverse, 5′-GTTCCTTGCTAGGGAGGCT-3′) primers.

Quantitation of p65 Nuclear Translocation—Jurkat T cells transfected with EGFP-shPKCα or EGFP-pCMS2 (48 h) or untransfected cells suspended in medium containing 2 mM Ca^{2+} or Ca^{2+} free equivalent solution were adhered to Cell-Tak-treated coverslips for 10–15 min. Cells were then stimulated at 37 °C as indicated. For PKCα suppression experiments, cells were stimulated in the presence of 2 mM Ca^{2+}. At the indicated times, cells were fixed in formaldehyde (3.7%) for 30 min, permeabilized with 0.2% Triton-X100 for 15 min, and blocked overnight in 2% BSA at 4 °C. Fixed and blocked cells were incubated with rabbit anti-p65 primary antibody (Santa Cruz Biotechnology, catalog no. 372, 1 μg/ml) for 1 h at 37 °C or overnight at 4 °C degrees, washed three times for 5 min each in 1% BSA in PBS, and incubated with Alexa 488 or 546 goat anti-rabbit secondary antibody (4 μg/ml) for 1 h at 37 °C. Nuclei were then labeled with Hoechst 33342 (Life Technologies, catalog no. H3570, 4 μg/ml), washed three times for 5 min each in 1% BSA in PBS, and mounted in Fluoromount (Fisher). Images of p65 localization were obtained with a Yokagawa spinning disk confocal system (Tokyo, Japan) mounted on a Leica DMI4000 microscope (Leica Microsystems, Wetzlar, Germany), and imaging parameters were optimized independently for each channel to maintain fluorescence within the linear range while maximizing intensity resolution. Images of p65 and Hoechst were overlaid, and cytoplasmic/nuclear p65 localization was determined using the Multispectral Cell Scoring application (Molecular Devices, Downingtown, PA). Average nuclear and cytoplasmic p65 fluorescence intensities were quantified within cytoplasmic and nuclear compartments, and intensity ratios were determined for each cell.

Real-time Localization of WT and Mutant p65—Jurkat T cells expressing WT and p65 Ser-536 mutants (16–24 h) were adhered to Cell-Tak-coated (BD Biosciences) coverslips and maintained in culture medium (RPMI 1640 medium, 10% FBS, 1% Glutamax) in a temperature- and CO₂-controlled chamber for 1 h during imaging. GFP-WT and GFP-p65 mutants were visualized every 10 s after stimulation with PMA (200 nM) with ionomycin (1 μM) and/or PMA (200 nM) with and without the delayed addition of ionomycin (1 μM).

Calcium Imaging—Jurkat T cells (3 million cells/ml) were loaded with 3 μM Fura-2 acetoxymethyl ester (Molecular Probes, Eugene, OR) in external solution containing 145 mM NaCl, 4.5 mM KCl, 2 mM CaCl₂, 1 mM MgCl₂, 10 mM glucose, 10 mM HEPES, 2 mM glutamine, and 2% fetal bovine serum (Hyclone, Thermo Scientific) for 10 min at 25 °C. Cells were adhered to coverslips coated with Cell-Tak (BD Biosciences), mounted on the stage of a Leica DMI6000 microscope configured with a Photometrics Evolve 512 camera (Tucson, AZ) using an Olympus ×40 oil objective (Shinjuku, Tokyo, Japan), and images were acquired with MetaFluor software (Molecular Devices). During imaging, cells were perfused with Ca^{2+}-free bath solution before activation with PMA (200 nM) and ionomycin (1 μM), thapsigargin (1 μM), anti-TCR (0.5 μg/ml) and CD28 (1:50) antibodies, or TNF (10 ng/ml) to evaluate stimulus-dependent Ca^{2+} release from the ER. The cells were then perfused with bath solution containing 2 mM Ca^{2+} to assess Ca^{2+} entry via activated Orai channels. In some experiments, cells were pretreated for 15 min with the Orai-1 inhibitor Synta66 (50 μM, Aobious, Gloucester, MA) prior to stimulation. Ca^{2+} mobilization was analyzed by plotting the emission ratio of 340/380-nm excitation for each cell. Each plot is the average ratio from at least 30 cells.

Statistical Analysis—Significance for all statistical tests was determined at p < 0.05 and is shown as *, p < 0.01; **, p < 0.001; and ***, p < 0.001 in all figures. Average firefly/Renilla luciferase ratios were calculated from three to four independent experiments and analyzed using two-tailed Welch’s t test. Western blot protein intensities were quantified using ImageJ (http://imagej.nih.gov/ij/), and average protein intensity values were compared using two-tailed Welch’s t test. p65 nuclear-to-cytoplasmic fluorescence intensity ratios were assessed for normality using probability plots and Kolmogorov-Smirnov test for normality. Normal distributions were compared using standard Student’s t test. Quantitative PCR relative quantification (RQ) values and percent input values were compared using two-tailed Welch’s t test.

Results

Extracellular Ca^{2+} Is Required for TCR-induced NF-κB Signaling—Ca^{2+} regulates proximal TCR signaling upstream of IKK activation (8–10). However, the precise function of Ca^{2+}
and the source of Ca$^{2+}$ required for NF-κB activation is unknown. To address these questions, we first asked whether the initial release of Ca$^{2+}$ from ER stores was sufficient or whether sustained influx of extracellular Ca$^{2+}$ is required for NF-κB activation in T cells. To distinguish between these pools of Ca$^{2+}$, we activated T cells in the presence or absence of extracellular Ca$^{2+}$ with either anti-CD3 and anti-CD28 (3/28) or with the diacylglycerol analog PMA together with ionomycin (P/I), which activate PKC and release ER-stored Ca$^{2+}$, respectively, and we compared these responses to those induced by the proinflammatory cytokine TNF. In Ca$^{2+}$-free medium, 3/28 and P/I, but not TNF, induced a transient rise in cytoplasmic Ca$^{2+}$ concentration because of release from the ER. Reintroduction of extracellular Ca$^{2+}$ led to a sustained secondary increase in cytoplasmic Ca$^{2+}$ levels via entry through activated Orai1/Ca$^{2+}$ release-activated Ca$^{2+}$ channels in 3/28- and P/I-stimulated cells (35) (Fig. 1A). Thus, stimulating cells in the absence of extracellular Ca$^{2+}$ allows us to specifically determine whether release from ER stores alone is sufficient for NF-κB signal activation.

As shown in Fig. 1B, top panels, all three stimuli induced the expected degradation and resynthesis of IκBα in Jurkat T cells, consistent with activation of the IKK complex and the classical NF-κB pathway. In contrast, neither 3/28 nor P/I induced IκBα degradation in Ca$^{2+}$-free medium, whereas IκBα degradation and resynthesis in response to TNF remained intact (Fig. 1B, bottom panels). Consistent with the effects on IκBα degradation, 3/28-stimulated NF-κB transcriptional activity was completely inhibited and P/I-induced transcriptional activation was significantly reduced in Ca$^{2+}$-free medium (Fig. 1C). In contrast, TNF-induced NF-κB reporter activity was unaltered in the presence or absence of extracellular Ca$^{2+}$ (Fig. 1C). A similar regulation of IκBα expression was observed in primary human CD4+ T cells (Fig. 1D). Collectively, these findings reveal that transient release of Ca$^{2+}$ from ER stores is not sufficient and that extracellular Ca$^{2+}$ is required for TCR-induced NF-κB activation.

**TCR-induced NF-κB Activation Requires STIM1 and Orai1**—The extracellular Ca$^{2+}$ requirement for TCR-induced NF-κB activation implies a crucial role for STIM1-operated
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Orai1 channel-mediated Ca\(^{2+}\) influx. To explore this, we expressed a STIM1 shRNA construct or a bicistronic variant for concomitant re-expression of shRNA resistant STIM1 to normal levels in Jurkat T cells (Fig. 2A). STIM1 suppression inhibited 3/28- and P/I-induced extracellular Ca\(^{2+}\) influx, and this was rescued by re-expression of STIM1 (Fig. 2B). Consistent with the lack of TCR-induced NF-κB signaling in Ca\(^{2+}\)-free medium (Fig. 1, B and C), STIM1 suppression prevented 3/28- and P/I-induced IκBα degradation in T cells (Fig. 2C, left versus center panel). In contrast, IκBα degradation and re-expression were normal in STIM1-rescued cells, confirming that the inhibition was due to loss of STIM1 (Fig. 2D, right panel). Furthermore, both 3/28- and P/I-induced NF-κB transcriptional activity was reduced in STIM1-suppressed cells, and this was again rescued by concomitant STIM1 re-expression (Fig. 2C). Notably, TNF-induced IκBα degradation and NF-κB transcriptional activity were unaffected by suppressing STIM1 (Fig. 2, C and D).

To confirm the role of Orai in Ca\(^{2+}\)-dependent NF-κB activation, we examined the consequence of inhibiting Orai-mediated Ca\(^{2+}\) influx with the Orai inhibitor Synta66 (Fig. 3A) and by expressing a mutant Orai1 (glutamic acid at position 106 mutated to alanine, E106A) that exerts a dominant negative effect on the Ca\(^{2+}\) permeability of endogenous Orai channels (Fig. 3C). In the presence of Synta66, the sarcoplasmic/endoplasmic reticulum Ca\(^{2+}\) ATPase inhibitor thapsigargin triggered Ca\(^{2+}\) release from the ER, evident as a small transient increase in cytoplasmic Ca\(^{2+}\) (in Ca\(^{2+}\)-free medium), but no subsequent sustained increase in cytoplasmic Ca\(^{2+}\) (Fig. 3A, compare the top and bottom panels) following perfusion with Ca\(^{2+}\)-containing medium. Consistent with the effects of STIM1 suppression (Fig. 2), Synta66 (Fig. 3B) inhibited 3/28- but not TNF-induced IκBα degradation. A similar block in stimulus-induced Ca\(^{2+}\) entry, and IκBα degradation was observed in permeation-defective Orai1-E106A cells (Fig. 3, C and D) Taken together, these results reveal an obligate role for STIM1-operated Orai1-mediated Ca\(^{2+}\) entry in TCR- but not TNF-induced IκBα degradation and NF-κB activation.

Ca\(^{2+}\) Controls the Transcriptional Activity of TCR-induced NF-κB—P/I treatment mimics TCR signaling upstream of IKK activation because PMA activates the strictly diacylglycerol-dependent and Ca\(^{2+}\)-independent "novel" PKC isoform PKCθ (41), and ionomycin-mediated Ca\(^{2+}\) release from the ER activates STIM1-dependent Orai activation (Figs. 1A and 2B). Thus, both 3/28 and P/I stimulation of T cells induce rapid PKCθ-mediated IKK-dependent degradation of IκBα, followed by resynthesis of IκBα via NF-κB-driven transcription (Figs. 1, 2, 3, and 4A). In seeking to determine the precise contribution of Ca\(^{2+}\) release from the ER to NF-κB activation, we found that PMA alone induces substantial IκBα degradation (Fig. 4A, center panel), suggesting that strong pharmacological activation of PKCθ can circumvent the requirement for Ca\(^{2+}\) upstream of IKK activation. However, treatment with ionomycin alone had no effect on IκBα levels (Fig. 4A, bottom panel), indicating that Ca\(^{2+}\) mobilization in the absence of PKCθ activation is not sufficient to activate the IKK complex.

Strikingly, although PMA in the absence of ionomycin induced IκBα degradation, this was not followed by IκBα resynthesis (Fig. 4A, compare IκBα levels at 60 min). Moreover, the kinetics of PMA-induced IκBα degradation were delayed compared with the response to P/I. We reasoned that the delayed IκBα degradation following stimulation with PMA alone likely

FIGURE 2. STIM1-dependent Ca\(^{2+}\) entry is required for TCR- but not TNF-induced NF-κB activation. A, immunoblot analysis of STIM1 and α-tubulin levels in Jurkat T cells transfected with either vector alone, shSTIM1, or shSTIM1-STIM1 suppression and rescue vectors. **, p < 0.01; n = 4 experiments. B, the role of STIM1 in P/I- and 3/28-mediated Ca\(^{2+}\) entry was similarly assessed in control (vector-transfected) Jurkat T cells, STIM1-suppressed cells, and STIM1 suppression with STIM1 rescue as described in A. Each trace represents the average response of at least 30 cells and is representative of at least three separate experiments. C, Jurkat T cells were transfected with control vector (left panel), shSTIM1 (center panel), or shSTIM1-STIM1 (right panel) and then incubated for the times indicated with 3/28, P/I, or TNF. Lysates were immunoblotted with either anti-IκBα or α-tubulin loading control. D, Jurkat T cells were transfected with the NF-κB luciferase reporter construct together with either vector alone, shSTIM1, or shSTIM1-STIM1 and then activated with 3/28, P/I, or TNF for 4 h. Mean firefly:Renilla luciferase ratios ± S.E. pooled from at least three independent experiments (3 replicates/experiment) are shown and were compared using Welch’s t test. *, p < 0.05; **, p < 0.01; ***, p < 0.001; N.S., not significant.
reflects a cooperative role previously identified for the Ca\(^{2+}\)/H\(_{11001}\) regulated phosphatase calcineurin A (CnA) in CBM complex formation, IKK activation, and IκB\(_{9260}\) degradation (10, 42). Indeed, overexpression of a Ca\(^{2+}\)-independent, constitutively active CnA rescued the delay in PMA-induced IκB\(_{9260}\) degradation so that the rate and extent of degradation were indistinguishable from those in P/I-stimulated cells (Fig. 4B). Importantly, although constitutively active CnA rescued the modulatory role of Ca\(^{2+}\) in proximal steps of NF-κB activation (i.e. IKK activation), it did not rescue IκB\(_{9260}\) re-expression in PMA-stimulated cells, indicating that a separate Ca\(^{2+}\)-dependent mechanism regulates the distal steps of NF-κB activation.

Supporting this conclusion and consistent with the lack of resynthesis of IκB\(_{9260}\) in the absence of Ca\(^{2+}\) (Fig. 4A), analysis of IκB\(_{9260}\) mRNA levels revealed a limited induction of IκB\(_{9260}\) transcription when PKC was activated with PMA or in response to PMA and ionomycin under Ca\(^{2+}\)-free conditions. In contrast, in the presence of extracellular Ca\(^{2+}\), IκB\(_{9260}\) mRNA expression was significantly induced by PMA and ionomycin (Fig. 4C).

Because IκB\(_{9260}\) re-expression is driven by activated NF-κB as a negative feedback loop to limit NF-κB signaling (43), we questioned whether Ca\(^{2+}\)-dependent IκB\(_{9260}\) protein re-expression reflects a global requirement for Ca\(^{2+}\) in NF-κB transcriptional activation. As shown in Fig. 4D, ionomycin alone failed to activate NF-κB reporter activity. PMA alone triggered a small but significant increase in activity compared with baseline; however, PMA and ionomycin together significantly enhanced (4-fold) NF-κB-driven transcriptional activation. Together, these data indicate that Ca\(^{2+}\) controls the distal transcriptional activation of NF-κB following P/I stimulation of T cells.

To extend these findings, we performed an unbiased transcriptional analysis to fully assess the extent of Ca\(^{2+}\) control over NF-κB-regulated gene expression. Microarray analysis was performed on T cells stimulated in the absence or presence of extracellular Ca\(^{2+}\) with either PMA alone or PMA and ionomycin. Transcriptional analyses identified 20, 96, and 112 differentially expressed genes (false detection rate, 0.05, log\(_2\)-fold change, 0.59) at 1, 4, and 8 h, respectively, between unstimulated, PMA-treated, and P/I-treated T cells (Fig. 4E, blue).

This list of differentially expressed genes was further refined to include only validated and putative NF-κB target genes on the basis of known targets (44–47). In total, we found
that 10–20% of all Ca\textsuperscript{2+}-regulated genes were NF-κB targets (Fig. 4E, red). Strikingly, 11 of 12 of these NF-κB target genes were dramatically increased by Ca\textsuperscript{2+} entry (2- to 20-fold increase, false detection rate, <0.05) relative to PMA stimulation (no Ca\textsuperscript{2+} mobilization) or stimulation with P/I in 0 mM Ca\textsuperscript{2+} (ER release but no Ca\textsuperscript{2+} entry). This analysis also revealed that, among this cohort, three classical NF-κB regulated genes (IκBα, CXCL8, and TNF) were among the top differentially expressed Ca\textsuperscript{2+}-dependent genes (Fig. 4, E and F). Together, these data reveal an entirely novel function for Ca\textsuperscript{2+} in regulating TCR-induced, NF-κB-dependent gene activation.

Ca\textsuperscript{2+} Is Required for TCR-induced p65 Nuclear Localization—The failure of PMA alone to induce IκBα resynthesis following its degradation (Fig. 4A) implies that Ca\textsuperscript{2+} controls...
NF-κB activity distal to IKK activation. To address the mechanism of this regulation, we first asked whether Ca\(^{2+}\) entry was required for NF-κB nuclear localization following IκBα degradation. As expected, P/I triggered rapid nuclear translocation of p65, which peaked 30 min after stimulation (Fig. 5, A and B). PMA alone had an effect at 30 and 60 min of stimulation, but the extent was significantly less than that induced by P/I at all time points (Fig. 5B). In contrast, ionomycin alone did not induce p65 nuclear localization at any time point. To confirm this apparent role for extracellular Ca\(^{2+}\) in p65 nuclear localization, we examined the effects of PMA and ionomycin in the presence and absence of extracellular Ca\(^{2+}\). Again, ionomycin alone failed to trigger p65 nuclear localization in either the absence or presence of Ca\(^{2+}\) (Fig. 5, C and D). However, p65 was strongly driven to the nucleus by the combination of P/I in the presence of extracellular Ca\(^{2+}\), whereas, in Ca\(^{2+}\)-free medium, ionomycin failed to synergize with PMA, and the extent of p65 nuclear localization was identical to that triggered by PMA alone (Fig. 5D). Taken together, these findings reveal an essential role for Orai-mediated entry of extracellular Ca\(^{2+}\) in nuclear translocation of p65 following its release from IκBα in response to TCR signaling.

Ca\(^{2+}\) Regulates TCR-induced Phosphorylation of p65 at Ser-536—At least 12 p65 serine or threonine residues have been identified whose phosphorylation regulates its nuclear localization and/or transcriptional activation (13–15, 22, 23, 26–28, 30, 48–52). We therefore asked whether Ca\(^{2+}\) regulates p65 nuclear translocation by controlling the phosphorylation of any of these residues. Among these, we focused on signal-induced phosphorylation of p65 Ser-536 (25, 53) because this has been implicated in TNF-driven NF-κB activation (15). We found that neither PMA nor ionomycin alone induces Ser-536 phosphorylation (Fig. 6A). However, activation with both together (P/I) induced a robust transient increase in phosphorylation at this residue that peaked at 15 min (Fig. 6A). Furthermore, the synergistic effect of PMA and ionomycin on Ser-536 phosphorylation requires extracellular Ca\(^{2+}\) because no phospho-p65 was detected in cells stimulated in Ca\(^{2+}\)-free medium (Fig. 6B). Notably, TNF also induced transient Ser-536 phosphorylation, although this occurred more rapidly than the response to P/I (peak at 5 min), suggesting distinct regulatory mechanisms. Moreover, TNF stimulated robust Ser-536 phosphorylation in the absence of extracellular Ca\(^{2+}\) (Fig. 6C), again consistent with an obligate role for Ca\(^{2+}\) entry in TCR but not TNF-induced NF-κB activation in T cells.

Phosphorylation of p65 at Ser-536 has been shown to alter the kinetics of p65 nuclear translocation (24, 25). To determine whether Ca\(^{2+}\)-dependent Ser-536 phosphorylation regulates p65 nuclear localization, we expressed wild-type p65, a non-phosphorylatable serine-to-alanine (S536A) mutant, and a serine-to-aspartate (S536D) phosphomimic mutant in T cells. We then visualized nuclear translocation in real time by confocal imaging. Cells were first stimulated for 30 min with PMA alone to induce IκBα degradation. Then ionomycin was added, and cells were observed for an additional 30 min. Consistent with the role for Ca\(^{2+}\) in p65 nuclear localization, in cells expressing WT p65, PMA alone did not promote p65 nuclear migration, whereas robust nuclear translocation was observed within 10 min of Ca\(^{2+}\) mobilization by exposure to ionomycin (Fig. 6D, top panel, 40-min time point). In contrast, after PMA treatment, ionomycin did not induce nuclear localization of S536A, whereas S536D exhibited Ca\(^{2+}\)-independent nuclear localization during the initial 30 min of PMA stimulation (Fig. 6D, center and bottom panels) without any requirement for Ca\(^{2+}\) mobilization with ionomycin. Together, these results establish that Ca\(^{2+}\) entry is required for TCR-induced phosphorylation of p65 at Ser-
shown in either Ca\(^{2+}\)-reduced P/I-induced Ser-536 phosphorylation but had no live cells by spinning disk confocal microscopy. In each instance, cells were first stimulated for 30 min with PMA alone to trigger Iκκ and then lysates were immunoblotted to determine the amounts of total p65 and Ser-536 phosphorylation. A, Jurkat T cells were stimulated with P/I, PMA alone, or ionomycin (Iono) alone for the times shown, and then lysates were immunoblotted to determine the amounts of total p65 and Ser-536 phosphorylation. B, cells were treated with P/I in the absence or presence of 2 mM Ca\(^{2+}\), and then p65 or Ser(P)-536 amounts were determined by immunoblotting. C, Jurkat cells were incubated with TNF for the times shown in either Ca\(^{2+}\)-containing or Ca\(^{2+}\)-free extracellular bath solution, and then lysates were prepared and immunoblotted using the antibodies indicated. D, WT p65-GFP, and p65-GFP with serine 536-to-alanine (S536A) and serine 536-to-aspartate (S536D) point mutations were expressed in Jurkat T cells to determine the role of Ser-536 phosphorylation in p65 nuclear localization. WT and mutant p65-GFP localization was visualized over a time course of 60 min in live cells by spinning disk confocal microscopy. In each instance, cells were first stimulated for 30 min with PMA alone to trigger Iκκ degradation and then were treated in the continued presence of PMA with ionomycin after 30 min to assess the role of Ca\(^{2+}\) in p65 nuclear localization. E, Jurkat cells were treated for the times indicated with either P/I (left panel) or PMA alone, followed by addition of ionomycin after 30 min (right panel), and then the amounts of Iκκ were determined by immunoblotting.

PKC\(\alpha\) Regulates Ca\(^{2+}\)-dependent p65-Ser-536 Phosphorylation—Several kinases, including IKK\(\alpha\), IKK\(\beta\), IKKe, TBK1, and PKA, have been implicated in the phosphorylation of p65 at specific serine residues that regulate its transcriptional activation (26, 54), but none of these kinases are known to be Ca\(^{2+}\)-dependent. Because PKC\(\alpha\) is a Ca\(^{2+}\)-dependent Ser/Thr kinase (55, 56) and is the predominant conventional PKC isoform in T cells, we used shRNA knockdown (Fig. 7) to determine whether PKC\(\alpha\) plays a role in TCR-induced NF-κB signaling and, specifically, whether it mediates Ca\(^{2+}\)-dependent phosphorylation of p65 at Ser-536. PKC\(\alpha\) suppression did not affect Iκκ degr-adviation induced by P/I, indicating no apparent role for PKC\(\alpha\) upstream of IKK activation (Fig. 7B). However, similar to incubating cells with PMA alone (Figs. 4A), PKC\(\alpha\) suppression prevented the resynthesis of Iκκ normally observed 60 min after stimulation with P/I (Fig. 7B, compare lanes 5 and 10). In contrast, PKC\(\alpha\) suppression did not affect TNF-induced Iκκ degradation or resynthesis. Consistent with a role in Ser-536 phosphorylation, PKC\(\alpha\) suppression significantly reduced P/I-induced Ser-536 phosphorylation but had no significant effect on phosphorylation induced by TNF (Fig. 7, C and D). Thus, these data confirm that Ca\(^{2+}\)-dependent activation of PKC\(\alpha\) regulates TCR-induced phosphorylation of p65 at Ser-536.

Given the role we have identified for p65 Ser-536 phosphorylation in p65 nuclear localization (Fig. 6D) and the role for PKC\(\alpha\) in p65 Ser-536 phosphorylation, we investigated whether PKC\(\alpha\) controls p65 nuclear localization. Consistent with the role of PKC\(\alpha\)-dependent p65 phosphorylation, PKC\(\alpha\) suppression significantly decreased the extent of p65 nuclear localization 15, 30, 60, and 90 min post-stimulation with P/I in 2 mM Ca\(^{2+}\) (Fig. 7, E and F). Importantly, p65 localization, expressed as the nuclear to cytoplasmic ratio, in unstimulated T cells was identical regardless of the level of PKC\(\alpha\) expression (median nuclear to cytoplasmic ratio: 0.54; median nuclear to cytoplasmic ratio: 0.53).

We next asked to what extent this phosphorylation of p65 impacts p65 binding to promoters of Ca\(^{2+}\)-dependent genes identified in our transcriptional analysis. We performed ChIP analyses to assess p65 binding to three genes found by transcrip-tional analysis to exhibit the strongest Ca\(^{2+}\)-dependent induction (Iκκ, CXCL8, and TNF) and to assess the role of PKC\(\alpha\)-dependent p65 phosphorylation in promoter binding
Quantification of p65 binding to IκBα/H9260B/H9251, CXCL8, and TNF promoters in Jurkat T cells demonstrated that PKCα suppression significantly reduced p65 binding to IκBα/H9260B/H9251, TNF, and CXCL8 promoters (Fig. 7G). These results are also consistent with the observed reduction in IκBα protein re-expression (Fig. 7B), confirming that reduced promoter binding directly impacted subsequent transcription and protein expression. Together, this comprehensive analysis establishes the critical importance of Ca²⁺/H11001 dependent PKCα activation in p65 nuclear localization, promoter binding, and transcriptional activation of a cohort of key NF-κB target genes.

**Discussion**

The need to precisely determine how Ca²⁺/H11001 regulates distinct transcriptional responses in T cells is underscored by the fact that almost 60% of TCR-induced genes are subject to Ca²⁺/H11001-dependent control (57). The notion that Ca²⁺/H11001 regulates NF-κB activation in lymphocytes is rooted in decades-old work dem-
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Demonstrating that NFAT and NF-κB activity is tuned to distinct calcium dynamics (3, 5, 6). Specifically, NFAT activation requires steady-state Ca\(^{2+}\) elevations (2, 3, 5, 6), although the amplitude of steady-state Ca\(^{2+}\) signals may further dictate which NFAT isoform is activated (58, 59). In contrast, selective activation of NF-κB has been linked to low-frequency spikes in cytoplasmic Ca\(^{2+}\) (3, 5). However, little is known about the nature of these Ca\(^{2+}\) signals, and the source of Ca\(^{2+}\) required to activate NF-κB has not previously been explored. Thus, in comparison with the established role and mechanism of Ca\(^{2+}\)-dependent NFAT activation, the mechanisms by which Ca\(^{2+}\) regulates TCR-induced NF-κB activation remain undefined.

We first asked whether the relatively infrequent Ca\(^{2+}\) spikes that selectively activate NF-κB in lymphocytes (3, 5, 6) could be generated from ER release without a need for extracellular influx (60). Unexpectedly, we found that ER release is insufficient and that Ca\(^{2+}\) influx via Orai is required to activate NF-κB. We then focused on the mechanism by which Orai-mediated Ca\(^{2+}\) entry regulates NF-κB activation.

Engagement of the TCR triggers canonical NF-κB activation by PKCθ-driven formation of the CBM complex (containing CARMA1, Bcl10, and MALT1) (8, 9, 61, 62). During formation of the CBM complex, Ca\(^{2+}\) has been implicated in CARMA1 and Bcl10 phosphorylation via Calmodulin kinase II (63–65) and Bcl10 dephosphorylation by Calcineurin A (10, 42). Our data confirm this general modulatory role for Ca\(^{2+}\) in steps proximal to IKK activation and IkBα degradation. However, we also found that pharmacological activation of PKCθ using either PMA stimulation alone (Fig. 4A) or PMA plus ionomycin in Ca\(^{2+}\)-free medium (Fig. 1B) triggers substantial IkBα degradation in the absence of Ca\(^{2+}\) mobilization. Thus, rather than exhibiting an absolute requirement for Ca\(^{2+}\), our data suggest that Ca\(^{2+}\) cooperates with PKCθ to accelerate the rate and, possibly, extent of IkBα degradation. Hence, although PKCθ activation is sufficient for CBM complex formation and IKK activation, Ca\(^{2+}\) serves a modulatory role via CnA upstream of IKK activation. An additional finding here is the oblige role for Ca\(^{2+}\) in NF-κB activation distal to IkBα degradation, where it controls p65 phosphorylation, nuclear localization, target gene promoter binding, and transcriptional activation.

This regulatory role for Ca\(^{2+}\) in IKK-distal signaling is entirely novel and establishes Ca\(^{2+}\) as a critical regulator at multiple checkpoints of NF-κB activity. Notably, although TNF signaling involves IKK activation and induction of p65 phosphorylation, our data establish that this occurs independently of any requirement for Ca\(^{2+}\). Most importantly, we show that TCR-induced p65 phosphorylation on Ser-536 definitively involves Ca\(^{2+}\)–dependent activation of PKCθ and that the kinetics of this phosphorylation are distinct from the regulation of p65 phosphorylation in response to TNF. A number of separate kinases have been described that control TNF-driven and TNF-independent p65 phosphorylation (13, 15, 20, 22–28, 30, 49, 50, 66–69). However, none of these require Ca\(^{2+}\), and our experiments show that PKCθ plays no role in TNF signaling. Thus, we established pathway-specific nodes of control for TCR versus TNF-induced NF-κB signaling in which Ca\(^{2+}\) regulation of PKCθ represents a novel but crucial regulatory step in TCR-induced transcriptional activation of NF-κB in T cells.

Furthermore, we delineated two separate Ca\(^{2+}\)–dependent checkpoints, one proximal and one distal to IKK activation, that modulate TCR-induced NF-κB signaling.

Our results have far-reaching implications concerning the mechanisms controlling T cell development and cell fate specification. In this regard, recent work has highlighted fundamental roles for TCR-induced Ca\(^{2+}\) entry in the development of immunity (34). For example, individuals with functional defects in STIM or Ca\(^{2+}\) release-activated Ca\(^{2+}\)/Orai are profoundly immune-deficient, and mice conditionally lacking STIM in T lymphocytes develop autoimmune disorders in part because of defective thymic natural regulatory T cell induction by high-affinity self-agonists (70). A similar and selective defect in natural regulatory T cell development occurs in mice selectively lacking either c-Rel (71–76) or upstream mediators of NF-κB activation, including BCL10, PKCθ, CARMA1, CnAβ, and IKKβ (77–81). Although our study focuses on p65-dependent transcriptional activation, the dual sensitivity of proximal and distal Ca\(^{2+}\) signals we identified and the role of c-Rel in natural regulatory T cell development raises the intriguing possibility that c-Rel transcriptional activation is also Ca\(^{2+}\)– dependent. If this were the case, then one would speculate that p65 and c-Rel regulation, like NFAT isoforms, might be tuned to quantitatively or qualitatively distinct Ca\(^{2+}\) dynamics. Thus, although we have shown that p65 nuclear localization and transcriptional activation are regulated by Ca\(^{2+}\)–dependent PKCα-mediated phosphorylation of p65 Ser-536, distinct Ca\(^{2+}\)–activated kinases could control c-Rel activity. Hence, critical goals of future studies will be to quantify the Ca\(^{2+}\) threshold of IKK activation and p65 phosphorylation, identify the range of Ca\(^{2+}\)–dependent Ser/Thr kinases activated following TCR engagement, and elucidate the role of Ca\(^{2+}\) in c-Rel activation.

Author Contributions—B. D. F., M. J. M., C. M. G., C. T. B., L. A. M., and U. H. designed the experiments. B. F., M. M., and C. B. wrote the manuscript. X. L., C. B., K. M., L. M., C. G., and K. A. M. performed the experiments. B. F., X. L., C. B., K. M., L. M., C. G., and M. M. performed the transcriptional assays. C. B. and K. A. M. performed the statistical analyses. X. L., C. B., K. M., L. M., C. G., and K. A. M. performed the luciferase assays. G. R. and C. B. performed the luciferase assays.

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