Cross-linking Proteomics Indicates Effects of Simvastatin on the TLR2 Interactome and Reveals ACTR1A as a Novel Regulator of the TLR2 Signal Cascade

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In Brief
The TLR2 interactome was defined in cells upon exposure to the TLR2 agonist Pam3CSK4 and the HMG CoA reductase inhibitor simvastatin, singly and in combination, by applying co-immunoprecipitation-based cross-linking proteomics. Two different crosslinkers with different spacer chain lengths were used for fixing protein complexes. The results indicate that ACTR1A is a potential interactor of TLR2. Functional studies using RNA interference confirm that ACTR1A is required for TLR2 signaling to induction of pro-inflammatory cytokines.

Highlights
- An innovative co-IP crosslinking proteomics study was designed for the TLR2 interactome.
- Proteomic profiling revealed combinatorial effects of simvastatin and Pam3CSK4 on the TLR2 interactome.
- ACTR1A and MARCKSL1 proteins were identified as potential interactors of TLR2 during the immune response.
- ACTR1A has important modulatory actions on the TLR2 pro-inflammatory signaling cascade.
Cross-linking Proteomics Indicates Effects of Simvastatin on the TLR2 Interactome and Reveals ACTR1A as a Novel Regulator of the TLR2 Signal Cascade*

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Toll-like receptor 2 (TLR2) is a pattern recognition receptor that, upon ligation by microbial molecules, interacts with other proteins to initiate pro-inflammatory responses by the cell. Statins (hydroxymethylglutaryl coenzyme A reductase inhibitors), drugs widely prescribed to reduce hypercholesterolemia, are reported to have both pro- and anti-inflammatory effects upon cells. Some of these responses are presumed to be driven by effects on signaling proteins at the plasma membrane, but the underlying mechanisms remain obscure. We reasoned that profiling the effect of statins on the repertoire of TLR2-interacting proteins might provide novel insights into the mechanisms by which statins impact inflammation. In order to study the TLR2 interactome, we designed a coimmunoprecipitation (IP)-based cross-linking proteomics study. A hemagglutinin (HA)-tagged-TLR2 transfected HEK293 cell line was used to precipitate the TLR2 interactome upon cell exposure to the TLR2 agonist Pam3CSK4 and simvastatin, singly and in combination. To stabilize protein interactors, we used two different chemical cross-linkers with different spacer chain lengths. Proteomic analysis revealed important combinatorial effects of simvastatin and Pam3CSK4 on the TLR2 interactome. After stringent data filtering, we identified alpha-cenactin (ACTR1A), an actin-related protein and subunit of the dynactin complex, as a potential interactor of TLR2. The interaction was validated using biochemical methods. RNA interference studies revealed an important role for ACTR1A in induction of pro-inflammatory cytokines. Taken together, we report that statins remodel the TLR2 interactome, and we identify ACTR1A, a part of the dynactin complex, as a novel regulator of TLR2-mediated immune signaling pathways. *Molecular & Cellular Proteomics 18: 1732–1744, 2019. DOI: 10.1074/mcp.RA119.001377.

Protein interactions have an important role in biological and cellular systems, including gene expression, signaling, and immune responses. The challenges associated with identifying specific protein-interacting partners in complex biological samples (1-3) have led to the development of several methodological approaches. Coimmunoprecipitation (IP)-based identification of protein interactions is a gold standard technique for defining protein complexes in native biological systems (4). In this method, a protein of interest is subjected to affinity- or antibody-based purifications along with its interacting partners. Optimization of wash conditions that remove nonspecific interactions but preserve transient and weak interactions is a major challenge that renders this method most amenable to identifying stable protein-protein interactions. In order to improve co-IP proteomics, protein cross-linking methods that covalently attach proximal protein binding partners have recently been employed (5, 6). Cross-linking theoretically captures transient and weak protein interactions, permitting the subsequent use of strong denaturing washing conditions that preserve specificity. A further advantage of cross-linking methods is that interactions can be defined either through identifying the proteins or in some cases through specifically examining cross-linked peptides. Although domain-specific cross-linking data analysis is hindered because of the complexity of bioinformatics software, several software packages are currently available for specific cross-linkers. Nonetheless, because confident protein identification is still very challenging for large-scale data sets, identifying the interaction of cross-linked proteins by examining unmodified peptides has become a very popular approach.

The Toll-like receptors (TLRs) are a family of type I trans-membrane proteins of the innate immune system that trigger a stereotypical pro-inflammatory cytokine induction response upon ligation. The 10 TLRs of the human innate immune system are localized to either the plasma membrane (TLR1, 2, 4, 5, 6) or endosome (TLR3, 7, 8, 9), and are activated by signature molecular patterns present in microbes (7-11). Of
the TLRs, TLR2, activated by lipoteichoic acid, synthetic lipo-peptides (Pam3CSK4 (P3C)), and glycans from Gram-positive bacteria, Gram-negative bacteria, and mycobacteria (12, 13), plays a pivotal role in the early innate immune response to microbial infections through triggering a signaling cascade that leads to the activation of the pro-inflammatory transcription factor nuclear factor-κB (8, 13, 14). Moreover, TLR2-dependent signaling contributes to the pathogenesis of a wide variety of diseases, such as antiphospholipid syndrome, sepsis, tuberculosis, chronic obstructive pulmonary, cytomegalovirus infection, rheumatic heart disease, cerebral injury, cutaneous leishmaniasis, atherosclerosis and leptospirosis (16–22). Given this, an improved understanding of TLR2-dependent signaling and how it is altered by immunomodulatory medications may provide novel insights relevant to human disease.

Statins, inhibitors of 3-hydroxy-3-methylglutaryl-CoA (HMG-CoA) reductase, are widely prescribed to reduce serum cholesterol in hyperlipidemic patients. Recently, statins have been shown to have additional immunomodulatory activities that are relevant to the pathogenesis of cardiovascular and other diseases (23). For example, statins suppress maturation of human monocyte-derived dendritic cells (24), and have been shown to ameliorate inflammation in a wide range of animal models of immunological disorders such as autoimmune encephalomyelitis (25, 26), sepsis (27), and graft arterial disease (GAD). Paradoxically, in other settings, proinflammatory effects of statins have also been identified on several cell types, including endothelial cells, peripheral blood mononuclear cells, and dendritic cells (28). The mechanisms that determine the pro- versus anti-inflammatory actions of statins in different settings remain poorly understood, but in many cases have been proposed to stem from indirect effects on membrane proteins (e.g. receptors).

We reasoned that P3C activation of TLR2 would serve as a well-defined model system amenable to unbiased proteomic-based profiling of the receptor-proximal effects of statins upon inflammatory signaling. To study the combinatorial effects of P3C and statins on the TLR2 protein interactome, we designed a cross-linking-based co-IP MS strategy. HEK293 cells stably expressing HA-tagged TLR2 were used to pull down TLR2 along with its interactors following crosslinker treatment. Given that smaller cross-linking agents may miss covalent attachment of surface receptors and cytosolic proteins near the membrane, we used two cross-linker agents with different spacer-chain lengths, our recently developed a dual cleavable cross-linker (DUCCT; spacer chain distance, 11.4 Å). After cross-linking and affinity pull-down, proteins were separated by SDS-PAGE, trypsin-digested, and the resultant peptides were analyzed by liquid chromatography-tandem mass spectrometry (LC-MS/MS) followed by data filtering.

Two proteins, alpha-centractin (ACTR1A) and myristoylated alanine-rich protein kinase C substrate-like protein 1 (MARCKSL1), were identified as novel interactors of TLR2 exclusively in statin-treated cells under DUCCT cross-linker treatment. We followed this discovery up with biochemical validation studies. We show that ACTR1A has important modulatory actions on the TLR2 pro-inflammatory signaling cascade. Taken together, these data identify for the first time that ACTR1A is a statin-responsive protein that serves to modulate TLR2-mediated signaling. Given the prevalence of statin use in human populations, these mechanistic studies may have important translational implications.

**EXPERIMENTAL PROCEDURES**

**HA-TLR2-MD2-CD14-293 Cell Line**—A plasmid expressing the HA-tagged human TLR2 gene (Catalogue # puno-htlr2ha, Invivogen, San Diego, CA) was transiently transfected into HEK293-hMD2-CD14 (Invivogen) using Lipofectamine 2000. Standard antibiotic selection procedures using Blasticidin S hydrochloride (Invivogen) were used together with Western blotting and immunostaining verification, to generate stable cell lines that strongly expressed (driven by a composite promoter hEF1/hTLV) the hTLR2-HA protein. In order to maintain selective pressure, the cell line was grown and maintained in DMEM containing 10% FBS, 10 U/ml penicillin, and 100 μg/ml streptomycin, supplemented with Blasticidin 10 μg/ml and Hygromycin 50 μg/ml.

**Cell Culture and Protein Preparation**—Hemagglutinin (HA) tagged-TLR2-MD2-CD14 human embryonic kidney (HEK)293 cells were maintained in DMEM supplemented with 10% fetal bovine serum, 1% penicillin/streptomycin in a humidified atmosphere of 5% CO2, and antibiotics (50 μg/ml hygromycin and 10 μg/ml blasticidin). Cells were treated with 10 μM simvastatin (Sigma) for 24 h, then stimulated with 1 μg/ml Pam3CSK4 (P3C; InvivoGen) for 24 h in fresh medium. The cells were then treated with Dual Cleaveable Cross-linking Technology (DUCCT) (29) or commercial bissulfosuccinimidyl suberate (BS3) cross-linker (XL), added at a final concentration of 1 μM/ml for 30 min, followed by quenching the reaction with 50 mM Tris-HCl, pH 8.0. For IP-pull down for proteomics, the cells were lysed with immunoprecipitation (IP)- lysis buffer supplemented with protease inhibitors at 4 °C for 15 mins, then sonicated for another 15 mins. Finally, the suspended cells were kept at 4 °C for 30 mins, then centrifuged (20,000 × g, 4 °C, 30 min). The supernatant was collected for measuring the protein concentration with a BCA protein assay kit, using bovine serum albumin as a standard.

**Separating the TLR2-interacting Partners Using Immunoprecipitation**—Anti-HA magnetic beads (Thermo Scientific, MA) were washed with 0.05% TBS-T buffer and gently vortexed. Suspended magnetic beads were collected using magnetic stand for 5 min at room temperature (RT). HA-tagged protein samples were mixed into the pre-washed beads and gently rotated at 4 °C overnight. The beads were then collected with a magnetic stand and washed with TBS-T buffer and ultrapure-water twice, followed by elution in Laemmli buffer (95 °C, 5 min). After centrifugation, the reduced samples were loaded onto SDS-PAGE gels (12%) for separation, followed by staining with Sypro Ruby in the dark for 12 h (supplemental Fig. S1).

For reverse coimmunoprecipitation (IP), protein samples were mixed with 5–10 μg anti-ACTR1A or -MARCKSL1 antibody, after volume adjustment to 500 μl with IP lysis buffer. The samples were incubated for overnight at 4 °C with continuous mixing, then exposed

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1 The abbreviations used are: HEK293T, HA-TLR2-MD2-CD14-HEK293 cells; XL, cross-linker; P3C, Pam3CSK4; ACTR1A, alpha-centractin; DUCCT, dual cleavable cross-linking technology.
to washed protein G magnetic beads (Thermo Scientific, MA) and incubated overnight at 4 °C with continuous mixing. Beads were collected using a magnetic stand, washed with washing buffer and ultra-pure water, then eluted in Laemmli buffer (95 °C, 5 min). The protein eluent was then separated by SDS-PAGE for immunoblotting.

In-gel Digestion and Mass Analysis (nano-LC-MS/MS)—SDS-PAGE gel bands were excised and minced (six pieces for each gel band), squeezed with acetonitrile, and dried at room temperature. Proteins were then reduced and alkylated and digested with trypsin (porcine) (MS Grade) at 37 °C for overnight (30). Formic acid to pH < 3 was added to the resulting peptides, followed by drying by speed vacuum, and then dissolution in 0.1% formic acid. Finally, the peptides were centrifuged at 20,000 × g for 30 min at 4 °C.

Digested peptides were analyzed by nano LC-MS/MS using a Velos Pro Dual-Pressure Linear Ion Trap Mass Spectrometer (ThermoFisher Scientific, MA) coupled to an UltiMate 3000 UHPLC (ThermoFisher Scientific, MA). Peptides were loaded onto the analytical column and separated by reverse-phase chromatography using a 15-cm column (Acclaim PepMap RSLC) with an inner diameter of 75 μm and packed with 2 μm C18 particles (Thermo Fisher Scientific, MA). The peptide samples were eluted from the nano column with multi-step gradients of 4–90% solvent B (A: 0.1% formic acid in water; B: 95% acetonitrile and 0.1% formic acid in water) over 70 min with a flow rate of 300 nL/min with a total run time of 90 min. The mass spectrometer was operated in positive ionization mode with nano spray voltage set at 2.50–3.00 kV and source temperature at 275 °C. The three precursor ions with the most intense signal in a full MS scan were consecutively isolated and fragmented to acquire their corresponding MS2 scans. Full MS scans were performed with 1 micro scan at resolution of 3000, and a mass range of m/z 350–1500. Normalized collision energy (NCE) was set at 35%. Fragment ion spectra produced via high-energy collision-induced dissociation (CID) was acquired in the Linear Ion Trap with a resolution of 0.05 FWHM (full-width half maximum) with an Ultra Zoom-Scan between m/z 50–2000. A maximum injection volume of 5 μL was used during data acquisition with partial injection mode. The mass spectrometer was controlled in a data-dependent mode that toggled automatically between MS and MS/MS acquisition. MS/MS data acquisition and processing were performed by Xcalibur™ software, ver. 2.2 (ThermoFisher Scientific, MA).

Database Search—Proteins were identified through Proteome Discoverer software (ver. 2.1, Thermo Fisher Scientific) using UniProt human (Homo sapiens) protein sequence database (120,672 sequences, and 44,548,111 residues). The reviewed protein sequences of human were downloaded from UniProt protein database (www.uniprot.org) on August 12, 2016. The considerations in SEQUEST searches for normal peptide sequences were used with carbamidomethylation of cysteine as the static modification and oxidation of methionine as the dynamic modification. Trypsin was indicated as the proteolytic enzyme with two missed cleavages. Peptide and fragment mass tolerance were set at ± 1.6 and 0.6 Da and precursor mass range of 350–3500 Da, and peptide charges were set excluding +1 charge state. SEQUEST results were filtered with the target PSM validator to improve the sensitivity and accuracy of the peptide identification. Using a decoy search strategy, target false discovery rates for peptide identification of all searches were < 1% with at least two peptides per protein, a maximum of two missed cleavage, and the results were strictly filtered by ΔCn (< 0.01), Xcorr (≥ 1.5) for peptides, and peptide spectral matches (PSMs) with high confidence, that is, with q-value of ≤ 0.05. Proteins quantifications were conducted using the total spectrum count of identified proteins. Additional criteria were applied to increase confidence that PSMs must be present in all three biological replicates samples. Normalization of identified PSMs among LC-MS/MS runs was performed by dividing individual PSMs of proteins with total PSMs and average of % PSM count was used for calculating fold changes for different treatment conditions (30, 31). For contrasting relative intensities of proteins between control, P3C, statin-P3C, and statin groups, samples were evaluated using cumulative confident normalized PSMs value.

Gene Ontology and Protein Interaction Analysis—Protein-encoding genes were functionally categorized using gene ontology systems by PANTHER classification system-based biological processes, molecular activities, and cellular components (32). Protein abundances were visualized as a heat map. The cluster was generated by MeV software (ver. 4.9; http://www.tm4.org/) (33). The proteomic data set, which included UniProt identifiers and fold changes of total identified protein, was submitted into Ingenuity Pathway Analysis (IPA) for core analysis (Ingenuity Systems, Redwood City, CA). The matched proteins with submitted dataset in Ingenuity Knowledge Base generated TLR2 protein interaction networks according to biological as well as molecular functions. The core analysis was performed with the settings of indirect and direct relationships between molecules based on experimentally observed data, and data sources were considered in human databases in the Ingenuity Knowledge Base (34). For generating the protein interaction networks in proteins exclusively identified upon treatment with DUCCT- and BS3-XLs, identified protein-coding genes were submitted into the Cytoscape ver. 3.6.1 according to affinity purification-mass spectrometry protein network analysis methods (35).

Immunoblotting—For immunoblotting, cells were washed with 1× PBS twice and then lysed with RIPA buffer (same as protein preparation). Protein samples were prepared in 2× Laemmli buffer and were heated for 5 min at 95 °C. Proteins were separated on a 12% polyacrylamide gel. The proteins were transferred to a 0.45 μm nitrocellulose membrane for 1 h at 100 V. The nitrocellulose membrane was then blocked in skim milk (5%) in TBST buffer for 2 h at room temperature (RT) and incubated with primary antibodies against ACTR1A (ab203833; Abcam), MARCKS-L1 (ab184546; Abcam), or TLR2 (ab191458; Abcam) in bovine serum albumin (5%) at 4 °C overnight. Goat anti-rabbit IgG secondary antibody conjugated to HRP (Abcam) was then used for 2 h at RT. β-actin (ab8227; Abcam), and GAPDH (ab9485; Abcam) antibodies were used as loading controls. The targeted protein bands were visualized using clarity Western enhanced chemiluminescent substrate (BioRad).

Immunocytochemistry—Cells were grown on 1m HCl-treated glass slides, and then fixed with chilled methanol for 5 min at RT. Cells were subsequently permeabilized with 0.1% Triton X-100 in 1× PBS for 10 min, blocked with bovine serum albumin and glycine in 1× PBS for 30 min at RT in the dark, and then incubated with anti-ACTR1A (ab11009, Abcam) or anti-TLR2 (PA5-20020, ThermoFisher scientific, IL) antibodies at 4 °C overnight in the dark. They were then incubated with secondary antibody (goat anti-rabbit IgG H&L, Alexa Flour 488, ab15007, Abcam) (2 h, dark, room temperature), and imaged with a Leica DMi8 confocal microscope (Leica, IL). The images were analyzed using Lax X (Leica, IL). DAPI was used for nuclear staining.

Transfection with ACTR1A siRNA and qRT-PCR—ACTR1A-targeted small interfering RNAs (siRNA; SR306823 for human) and nonsense siRNA were purchased from OriGene (OriGene, MD). For transfections, HEK293 cells were seeded in 6-well plates with DMEM medium supplemented with 10% FBS, 1% penicillin/streptomycin, and selective antibiotics (see cell culture methods). After 50–70% confluence, cells were transfected according to the manufacturer’s instructions. After 48 h, cells were treated with statin for 24 h and P3C for an additional 24 h. Following this, the cells were collected for mRNA analysis. Total RNA was extracted by TRizol® (Invitrogen) (30). First-strand cDNA synthesis was performed using one-step cDNA synthesis kit (OriGene, MD). Real-time PCR was performed on the CFX96 real-time system (Bio-Rad) using the SsoAdvanced™ Universal
sal SYBR® Green Supermix (Bio-Rad). Each assay was performed in triplicate, and the mean value was used to calculate the mRNA expression for the gene of interest and the housekeeping reference gene (GAPDH). The abundance of the gene of interest in each sample was normalized to that of the reference control using the comparative ($2^{-\Delta\Delta CT}$) method (36). Sequences of the primers are provided in the supplementary information (supplemental Table S1).

**Experimental Design and Statistical Rationale**—All Co-IP studies was performed in three biological replicates. Each biological replicate has four exposure conditions, control, P3C, Statin, and statin P3C. Each treatment condition was also treated with cross-linker or no cross-linker (control). Three replicates of SDS-PAGE gel were ran. Each gel lane was excised in six pieces and in gel tryptic digestion was performed.

The quantitative analysis of proteins as PSMs was performed using built-in-statistical packages in Proteome Discoverer (Ver. 2.1). Results were considered statistically significant if $q \leq 0.05$ ($n = 3$). Scatter plots and pairwise correlation matrices were generated using the R package, where results were considered if correlation coefficient ($R^2$) was $> 0.80$. The data are depicted in the graphs as mean ± S.E. (S.E.). Statistical significance was determined using one-way ANOVA with $p \leq 0.05$ ($n = 3$) considered as significant. GraphPad Prism version 6 was used (GraphPad Software, Inc).

**RESULTS**

**Identification of TLR2-interacting Proteins**—To identify the effect of P3C and statins on the TLR2 interactome, we performed co-IP proteomics on HA-TLR2-MD2-CD14-HEK293 cells from four exposure conditions (control; P3C; statin; statin-P3C) following post-exposure treatment with DUCCT or BS3 cross-linker (Fig. 1). Control samples untreated with crosslinker were also analyzed. After pulldown with anti-HA magnetic beads, precipitated proteins were separated by SDS-PAGE (supplemental Fig. S1) and the resulting gel bands were digested using trypsin and then analyzed by nano-LCMS/MS and database searching (UniProt). On-bead digests were also explored. However, we found that in-gel digestion after Laemmli elution yielded better recovery of HA-tagged bait TLR2, likely because of improved solubilization/denaturation of this transmembrane protein. To improve protein recovery from the gel, we also minced the gel bands into six pieces. Peptides were quantified using Peptide Spectrum Matches (PSMs). Correlation matrix comparisons among three biological replicates are shown in supplemental Fig. S2. Pairwise correlation coefficients among the biological replicates showed high correlation with a $R^2$ value of $> 0.80$.

Overall, 1153 proteins were identified and quantified across all conditions. The data set was filtered using two unique peptides per protein and a false discovery rate of 1%. Detailed information about the identification of proteins and peptides is shown in supplemental Table S2–S4. First, we examined proteins that were identified across the four cell exposure conditions, but in the absence of crosslinker treatment. Out of 803 proteins, 249 proteins were common to all four exposures, whereas 249 proteins were exclusive to a single exposure (16 in control, 78 in P3C, 149 in statin-P3C, and 6 in statin) (supplemental Fig. S3A). We next evaluated the impact of the two cross-linkers on protein discovery in the setting of HA-TLR2 pulldowns from P3C, statin, and P3C-statin-treated cells. In samples treated with DUCCT in combination with P3C, 220 proteins were commonly shared across control, P3C, P3C-DUCCT and DUCCT conditions, whereas 288 proteins were exclusively identified in individual conditions. Consistent with improved protein recovery with DUCCT, 16 more proteins were identified in P3C-DUCCT samples (total, 589 proteins) than in P3C-stimulated samples without DUCCT (total, 605 proteins) (supplemental Fig. S3B and supplemental Table S2). Of these, 147 proteins were exclusive to P3C-DUCCT (i.e. not detected under P3C, control, or DUCCT conditions) (supplemental Fig. S3B). Regarding statin-P3C-cotreated samples, 167 proteins were identified exclusively in statin-P3C samples, whereas, 28 proteins were exclusive to statin-P3C-DUCCT samples (supplemental Fig. S3C). In comparison of statin and statin-DUCCT treated samples (supplemental Fig. S3D), 15 and 221 proteins were exclusively identified, respectively.

Distinct effects on the TLR2 interactome were noted with BS3 cross-linker. Contrary to the increase in protein recovery
in P3C-treated cells enforced by DUCCT, BS3 treatment led to 224 fewer proteins identified under P3C treatment conditions (supplemental Fig. S3E). Because of this, remarkably, 240 more proteins were identified in DUCCT-treated P3C samples than in BS3-treated P3C samples (compare supplemental Figs. S3B and S3E). Similarly, 285 fewer proteins were identified in statin-P3C-BS3 samples than in statin-P3C samples (supplemental Fig. S3F). In this case, however, 77 more proteins were identified in BS3 samples compared with DUCCT samples after statin-P3C (supplemental Fig. S3C and S3F). Finally, in the case of statin-treated cells, BS3 led to identification of 107 more proteins (supplemental Fig. S3G). Because of a more marked improvement in protein recovery with DUCCT, 208 more proteins were identified in DUCCT-treated samples than in BS3-treated samples following statin exposure (supplemental Fig. S3D and S3G). Taken together, we conclude that, overall, compared with BS3, the DUCCT cross-linker led to improved recovery of the TLR2 interactome.

Visualization by heat map of the relative expression (normalized PSMs) of TLR2-interacting proteins (Fig. 2A) suggests that treatment with P3C, statins, and P3C-statins induce distinct biological states of the cells (Fig. 2A). Different proteomic expression patterns were evident among 1082 proteins identified under DUCCT treatment (Fig. 2B) and the 975 protein intensities identified under BS3 treatment (Fig. 2C), confirming that the two cross-linkers enforce a significant effect on the detected TLR2 interactome.

All 803 proteins identified in the absence of cross-linker treatment were categorized based on gene ontology using PANTHER gene classification systems (32). Overall, 8, 11, and 8 ontology pathways were represented within the ontology categories of cellular components, biological processes, and molecular functions, respectively. Several ontology pathways were particularly enriched by P3C and statin treatment. These are “cell part” and “organelle” under the “cellular components” category, “metabolic process” and “cellular process” under the “biological processes” category, and “catalytic activity” and “binding” under the “molecular functions” category (supplemental Fig. S4).

IPA-based TLR2-targeted Protein Interactions Network—Core analysis was next performed by IPA (Ingenuity Systems) (37) in order to construct hypothetical interaction networks, canonical pathways, functions and disease pathways, and putative upstream regulators among the TLR2 interactome proteins identified. We found 25 classified networks according to top diseases and functions, within which a TLR2 protein-interacting network was ranked number eight (supplemental Table S5). The TLR2 protein network centered on cellular compromise, cellular function and maintenance, and cell death and survival. Given that we had by design targeted
TLR2 in our interactome strategy, we next performed a TLR2-based network analysis (Fig. 3). Here, we found 32 interacting partner proteins. Expression of TLR2 itself was slightly increased by treatment with P3C, but sharply decreased during treatment with statin-P3C and statin (Figs. 3 and supplemental Fig. S5A–S5B). In this putative network, TLR2 interacted directly with heat shock protein 90-beta (HSP90B1) and the PI3-kinase p85 (PI3K p85) complex, and indirectly with vascular endothelial growth factor (VEGF). In the network, VEGF, in turn, is proposed to be regulated by interleukin enhancer-binding factor 3 (ILF3), which, in turn, is interconnected to heterogeneous nuclear ribonucleoprotein R (HNRNPD), far upstream element-binding protein 2 (KHSRP), ELAV-like protein 1 (ELAVL1), and ILF2. Of interest, ELVL1 protein is densely interconnected with 10 proteins experimentally observed in our pulldown. Further, ELVL1 connects to PI3K family, HNRNPD, and KHSRP, highlighting the potential for complex inter-communications between proteins in the proposed TLR2 interactome (Fig. 3 and supplemental Fig. S3A–S3B). In this study, we found that ATP citrate lyase (ACLY) was highly increased (10.8-fold) upon treatment with P3C compared with statin-P3C and statin treatment (Figs. 3, supplemental Fig. S3A–S3B). ACLY has recently been reported to support macrophage inflammatory activation (38), but has not previously been connected to TLR2 to our knowledge. Its suppression by statin treatment in our study interestingly identifies it as a putative mediator of the effect of statins on the inflammatory response.

Protein Identification and Interactions After Cross-linking—In this study, we used two chemical crosslinkers, DUCCT and BS3, for covalent attachment of nearby proteins, aiming to improve recovery of low abundance and weakly interacting proteins. After DUCCT, we identified 605, 285, and 618 proteins under P3C, statin-P3C, and statin exposure conditions. After BS3, 365, 362, and 410 proteins were correspondingly identified under these three exposures (supplemental Table S2–S4). After stringent filtering among cross-linker control, treatment controls, and cross-linked samples, we exclusively identified 166 proteins in P3C-DUCCT, 47 proteins in statin-P3C-DUCCT, and 225 proteins in statin-DUCCT-treated samples (Figs. 4 and supplemental Fig. S6, supplemental Table S3). Correspondingly, we exclusively identified 32 proteins in P3C-BS3, 43 proteins in statin-P3C-BS3, and 40 proteins in statin-BS3-treated samp-
Consequently, considering total and exclusively identified proteins, DUCCT cross-linker enriched more TLR2-interacting proteins compared with BS3. After stringent filtering of the identified proteins among all exposure and crosslinking conditions individually and in combination, the data indicates that DUCCT exhibits superior efficiency to couple proteins across different treatment conditions compared with BS3 (supplemental Fig. S6). A protein-protein interaction network was constructed using the exclusively identified proteins because DUCCT and BS3 treatments among the four cell exposure conditions (control, P3C, statin-P3C, statin), using the UniProt database via Cytoscape software (Figs. 4 and supplemental Fig. S7). A total of 325 DUCCT-exclusive proteins were used to generate the networks, containing 218 nodes and 320 edges (Fig. 4). As is evident, the highest node degree genes were RNA binding motif protein 8A (RBM8A; 35 edges), endoplasmic reticulum lipid raft-associated protein 2 (ERLIN2; 28 edges), eukaryotic translation initiation factor 4A3 (EIF4A3; 19 edges), RuvB like AAA ATPase 2 (RUVEL2; 16 edges), eukaryotic translation initiation factor 3 subunit B (EIF3B; 14 edges), splicing factor proline and glutamine rich (SFPQ; 14 edges), and transmembrane P24 trafficking protein 9 (TMED9; 13 edges). In parallel, 92 BS3-exclusive proteins were used to generate a network containing 121 nodes and 141 edges, within which G3BP stress granule assembly factor 1 (G3BP1) protein-coding gene showed high interaction with 3 node genes (supplemental Fig. S7).

Validation of Selected Proteins and Their Interacting Partners—To verify the mass spectrometry-based protein data, we performed IP and immunoblot analysis on selected candidate proteins. Among the TLR2-interacting proteins identified, we focused our attention on alpha-centractin (ACTR1A) and myristoylated alanine-rich protein kinase C substrate-like protein 1 (MARCKSL1), based on their expression as well as their potential functions. In the HA-TLR2 interactome proteomics pulldown, ACTR1A was identified exclusively in the DUCCT-treated samples under the two exposure conditions.
of P3C and statin (Fig. 5A), whereas MARCKSL1 protein was detected only in statin-P3C and statin exposure conditions in the absence of crosslinker treatment (Fig. 5A), suggesting distinct patterns of responsiveness of these two proteins to P3C and statin. For validation, first, we performed immunoblot analysis of whole cell lysates to evaluate the expression status of these two proteins. Both ACTR1A and MARCKSL1 were highly up-regulated in statin-P3C- and statin-treated samples compared with control and P3C-treated samples (Fig. 5B), suggesting that statins induce the expression of these two proteins in HEK293 cells. Next, HA-TLR2 IP samples were analyzed by immunoblot. We found that levels of ACTR1A coprecipitating with HA-TLR2 were significantly decreased in statin-treated cells (Fig. 5B). To further validate interactions of TLR2 with ACTR1A and MARCKSL1, we performed a reverse co-IP (i.e. immunoblot of TLR2 after ACTR1A IP) (supplemental Fig. S8). This revealed that TLR2 was highly increased in P3C- and statin-P3C-treated ACTR1A pull-down samples compared with control and statin-treated samples (Fig 5B). TLR2 was increased in P3C-, statin-P3C-, and statin-treated MARCKSL1 pull-down samples compared with control (Fig. 5B). Taken together, these findings suggest that P3C and statins enforce differential changes in the interaction of TLR2 with ACTR1A and MARCKSL1 in HEK293 cells. For further cross-validation, we performed immunocytochemistry on ACTR1A and TLR2 in the HEK293 cells. Here, we noted that in HEK293 cells TLR2 protein expression was inhibited by statin treatment, whereas ACTR1A protein was increased by statins (Fig. 6).

**ACTR1A Knockdown Changes the Levels of Cytokines**—To test for a possible function of ACTR1A in the TLR2 inflammatory response, we used siRNA to silence ACTR1A in HEK293 cells. After confirmation of siRNA efficiency in untreated cells (Fig. 7A), we analyzed expression of ACTR1A and of the pro-inflammatory genes tumor necrosis factor (TNFα), interleukin 6 (IL-6), and interleukin 8 (IL-8) in cells exposed to P3C, statin, and P3C-statin (Fig. 7). ACTR1A gene expression was successfully silenced by the siRNA under all treatment conditions (Fig. 7B). As expected, P3C induced robust TNFα (Fig. 7C). Of interest, statin treatment by itself did not change TNFα from control levels, but augmented the TNFα induction response to P3C. Whereas the TNFα response to P3C was not modified by silencing of ACTR1A, the TNFα response to combined P3C-statin treatment was significantly inhibited by ACTR1A silencing, suggesting that statins augment TLR2-dependent TNFα through a mechanism that requires ACTR1A.

Under our experimental conditions, P3C did not induce IL-6 in HEK293 cells, although, interestingly, statin treatment itself modestly increased IL-6 (Fig. 7D). Finally, as with TNFα, statins modestly augmented P3C induction of IL-8. Induction of IL-8 by P3C and P3C-statin treatment were both significantly reduced by ACTR1A silencing, suggesting that ACTR1A is required for both TLR2 responses of the cell.

**DISCUSSION**

Protein-protein interactions play a central role in signal transduction. Currently, IP coupled to high throughput mass spectrometry is popular for identifying target protein complexes (11, 39). Co-IP-based mass spectrometry has become a gold standard to study these interactions in a large-scale experimental design. Cross-linking before affinity or antibody pulldown aids in protein discovery but has been hindered in many cases by the low abundance of target proteins and also by the analytic challenges of cross-linked peptides in large-scale proteomic studies. We have developed DUCCT, a dual cleavable crosslinker with a spacer chain distance of ~16 Å

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**Fig. 5. Validation of TLR2 protein interactors.** A, ACTR1A and MARCKSL1 proteins expression in HEK293 cells by LC-MS/MS. B. ACTR1A and MARCKSL1 and their interactions were validated using immunoblotting (IB) and coimmunoprecipitation (IP) in HEK293 cells. All samples were treated with statin drug and bacterial ligand Pam3CSK4 except control.
and a software package is currently under development to search the resulting cross-linked products.

In the present study, for the first time, we used two cross-linkers with different spacer chain lengths to study TLR2-interacting proteins. Aiming to test for possible effects of statins on the TLR2 interactome, we evaluated cells treated with the TLR2 ligand P3C, with statin, or with P3C following statin treatment. Novel interactors were identified through pulldown of TLR2 by a HA epitope tag, and further validated with biochemical approaches. Importantly, our data indicate that DUCCT enhances recovery of the TLR2 interactome and does so in a superior fashion to BS3. Of interest, gene ontology analysis of the interactors using PANTHER gene function classification showed nearly half or one-third of the identified

![Fig. 6. Cross-validation of candidate proteins. ACTR1A and TLR2 protein expressions were cross-validated using immunocytochemistry upon the treatment of Pam3CSK4 and statin in HEK293 cells.](image)

![Fig. 7. Silencing of ACTR1A modulates cytokine expression.](image)

A, HEK293 cells were transfected with scramble siRNA and siRNA targeting ACTR1A, and then ACTR1A expression was analyzed by qRT-PCR. B–D, After 48 h with or without siRNA treatment, the relative mRNA expression of ACTR1A (B), TNF-alpha (C), IL6 (D), and IL8 (E) were analyzed by qRT-PCR upon treatment with statin and Pam3CSK4, as shown. All data are showed as mean ± S.E. (n = 3 in each group) with *, p < 0.05.

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proteins were involved in the protein binding category (33.96%) under molecular functions, cellular processes category (30.65%) under biological processes, and cell part category (39.69%) under cellular components (supplemental Fig. S4).

Computational biology techniques, IPA and Cytoscape, were used to predict interaction partners and targeted protein-encoding genes following treatment of cells with P3C and/or statin. We predicted the TLR2-targeted proteins network in Fig. 3 using the identified proteins in the pull-down samples. In supplemental Table S5, IPA analysis predicted a TLR2 protein network involved in cell integrity including maintenance the cellular function, cell death and survival (40). Interestingly, IPA networks predicted direct interactions between TLR2 and HSP90B1, a protein detected by us in the TLR2 interactome (Fig. 3). HSP90B1 protein expression increased in P3C-stimulated samples, while decreasing in statin-P3C and statin-treated samples (Figs. 3 and supplemental Fig. S5A–S5B), suggesting that statin treatment may modify the proximal TLR2 interactome. The endoplasmic reticulum HSP90 isoforms are commonly involved in cell survival and proliferation (41). A previous study reported that HSP90B1 up-regulation is associated with epithelial ovarian cancer and cancer cell survival (42). Moreover, bacterial infection and inflammation were increased in HSP90B1-deficient macrophages, which interfered with TLR signaling as well as function in the affected cells (43). Given this, we speculate that inhibition of HSP90B1 could be a potential strategy in cancer and anti-inflammatory therapy. Our interactome results further suggest that statins may be a potential tool for targeting HSP90B1 in the TLR2 cascade.

Protein-protein interaction network analysis upon treatment with P3C and statins along with DUCCT and BS3 revealed several proteins and protein complexes which are potential TLR2 interactors. RBM8A proteins were exclusively identified in statin-treated cells following DUCCT (Fig. 4). RBM8A proteins belong to the RNA-binding protein family. This family of proteins is believed to be involved in protein synthesis, cell cycle, apoptosis, and RNA splicing (44–47) and has also been shown to regulate intercellular immunity and production of inflammatory and anti-inflammatory cytokines (48). RBM8A is reported to be associated with the regulation of apoptosis, as inhibition of this protein increased cell apoptosis and reduced cell proliferations (45, 49). We speculate, based on our findings, that RBM8A may play a particularly important role in TLR2-mediated cell responses.

We validated two potential interacting partners of TLR2, namely ACTR1A and MARCKSL1, using immunoblot, reciprocal co-IP, and immunocytochemistry (Figs. 5–6). ACTR1A and MARCKSL1 proteins were remarkably up-regulated by statin treatment in HEK293 cells (Figs. 5–6). The related protein MARCKS is known to be a regulator of actin cytoskeleton dynamics and MARCKSL1 protein is thought to share similar molecular functions. MARCKSL1 is expressed in a variety of different cells including astrocytes, fibroblasts and microglial cells and is up-regulated in macrophages by LPS (50–53). MARCKS proteins crosslink actin filaments and maintain mechanical and structural properties of the actin cytoskeleton. During macrophage activation, MARCKS joins with actin or actin-binding proteins at sites of the cytoskeleton and plasma membrane (51, 54, 55). Both actin crosslinking and formation of actin bundles are inhibited by protein kinase C-dependent phosphorylation and binding of calmodulin to MARCKS proteins (56). MARCKS and MARCKSL1 proteins migrate from the cell membrane to cytosol upon phosphorylation or binding to calmodulin and thereby activate various signal transduction pathways by influencing actin dynamics and vesicular trafficking (56). Moreover, MARCKS proteins are involved in cell migration, adhesion, motility, membrane trafficking, mitogenesis, and spreading because of actin filament and microtubule reorganization (57, 58). Copurification of MARCKSL1 with ACTR1A suggests their cooperation in actin cytoskeletal dynamics in the TLR2-mediated signaling response.

ACTR1A is cytoplasmic protein that is a part of the cytoskeleton, particularly microtubules. It appears as short filaments at the center of the dynactin complex that actively associate to dynein motor activity (59, 60). Cytoplasmic ACTR1A protein has also been found to play a vital role in actin filament assembly and organization (61) and intracellular vesicle trafficking. Prior studies have reported that macrophage activation by mycobacteria or cell wall lipoprotein p19 (TLR2 agonist) induces cytoskeletal rearrangement by TLR2-mediated phosphatidylinositol 3-kinase (PI3K) activation pathways (62). This activation of PI3K was necessary for the actin assembly and reorganization that underlies macrophage spreading and polarization (62). The actin cytoskeleton is also assembled in P3C-stimulated dendritic cells which enhances antigen generation and capture (63). Taken together, a key role for the actin cytoskeleton has been identified in TLR2-dependent immune responses, and our findings lead us to speculate that ACTR1A, as a novel TLR2 interactor, may play an important role in mediating this connection.

Interestingly, in our studies, ACTR1A was highly expressed following statin treatment and also copurified with TLR2 in HEK293 cells, suggesting that it is a statin-sensitive TLR2 interactor (Fig. 5–6). Our co-IP-based mass spectrometry studies revealed that TLR2 interacts with ACTR1A in HEK293 cells upon statin and P3C treatment. The TLR2-ACTR1A interaction was confirmed with biochemical approaches. Further, to study the functional involvement of ACTR1A in TLR2 signaling pathways, we knocked down the expression of ACTR1A. Silencing of ACTR1A interestingly reduced pro-inflammatory cytokine expression in HEK293 cells, confirming an important role of ACTR1A in transducing the TLR2 pro-inflammatory signal. Future studies may be warranted to determine whether ACTR1A supports TLR2 signaling through linking TLR2 to the underlying cortical cytoskeleton.
ACTR1A is a Potential Regulator of the TLR2 Signal Cascade

Of note, we identified ACTR1A after cross-linking with DUCCT but not BS3. DUCCT has a larger cross-linker spacer chain compared with BS3 and may thus be superior for conjugating larger protein complexes and penetrating the plasma membrane. A part of large dynactin complex, ACTR1A function is not clearly known. This is the first time to our knowledge that ACTR1A and TLR2 interaction has been revealed by using co-IP cross-linking proteomics. MARCKS in the plasma membrane binds to dynamitin, a subunit of dynactin complex in cytoplasm (57). Copurification of MARCKSL1 and membrane binds to dynamitin, a subunit of dynactin complex using co-IP cross-linking proteomics. MARCKS in the plasma edge that ACTR1A and TLR2 interaction has been revealed by membrane. A part of large dynactin complex, ACTR1A functions as a junctioning larger protein complexes and penetrating the plasma membrane. Due in part to the suboptimal performance of IP antibodies targeting native TLRs, stable epitope tagged-TLR-HEK293 cell model systems have been used for many years to define TLR interactions and signaling events (66). Parental HEK293 cells are quite suitable for these controlled stable expression studies as they do not express native TLR2 proteins. Moreover, stable TLR2-HEK293 cells are functional, producing pro-inflammatory cytokines after TLR2 ligand stimulation, like native TLR2-expressing cells (67). Nonetheless, it is possible that interactions of native TLR2 could differ from those that we have defined in our model cell system, and that TLR2 interactions could even be cell type specific. Moreover, it is conceivable that different static preparations (e.g. hydrophobic versus hydrophilic) may trigger distinct effects on the TLR2 interactome. Given this, although our study has the strength of enriching the TLR2 interactome under well-defined conditions and generating novel predictions for the TLR field, we feel that future studies are now warranted to investigate the role of these new interactors in primary cells, in relation to other TLRs, and in the context of additional clinically relevant static preparations.

DATA AVAILABILITY

The mass spectrometry data have been deposited to the ProteomeXchange consortium via the pride (68) partner repository with the dataset identifier PXD012759.

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