Identification of phosphodiesterase-4 as the therapeutic target of arctigenin in alleviating psoriatic skin inflammation

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A B S T R A C T
Introduction: Arctigenin, derived from Arctium lappa L., has multiple pharmacological activities, including immunoregulatory, anti-diabetic, anti-tumor, and neuroprotective effects. Nevertheless, the potential therapeutic target of arctigenin in modulating inflammation remains undefined.

Objectives: In the present study, we identified that arctigenin was a phosphodiesterase-4 (PDE4) selective inhibitor for the first time. Further investigations were performed to fully uncover the effects and mechanism of arctigenin on experimental murine psoriasis model.

Methods: Crystal structure determination, PDEs enzyme assay, and isothermal titration calorimetry were included to illustrate the binding specialty, inhibitory effects, and selectivity of arctigenin on PDE4D. The anti-inflammatory effects were conducted in LPS-activated human peripheral blood mononuclear cells (PBMCs) and RAW264.7 cells. Imiquimod-induced murine psoriasis was performed to uncover the therapeutic effects and mechanism of arctigenin in vivo.

Results: Arctigenin could bind to the catalytic domain of PDE4D via formation of hydrogen bonds as well as - stacking interactions between the dibenzyl butyrolactone of arctigenin and several residues of PDE4D. Accordingly, arctigenin showed prominent anti-inflammation in human PBMCs and murine RAW264.7 cells. PDE4 inhibition by arctigenin resulted in elevation of intracellular cyclic adenosine monophosphate (cAMP) and phosphorylation of cAMP-response element binding protein (CREB), which were largely blocked through intervention of protein kinase A (PKA) activity by H89 treatment or reduction of protein expression by siRNA transfection. Moreover, we first identified that a topical application of arctigenin ameliorated experimental psoriatic manifestations in imiquimod-induced murine psoriasis model by decreasing adhesion and chemotaxis of several inflammatory cells. Further proteomics analysis revealed that arctigenin could rectify the immune dysfunction and hyperactivation of keratinocytes in the inflamed skin microenvironments, which might be largely related to the expression of Keratins.

Conclusion: The research provided credible clew that inhibition of PDE4 by arctigenin might function as the potential therapeutic approach for the treatment of psoriasis.

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Introduction

Traditional Chinese medicine (TCM) offers a vast treasury of pharmaceutical candidates in modulating immune homeostasis and treating inflammatory diseases [1–3]. Arctigenin, a phenylpropanoid dibenzyl butyrolactone lignan, is one major active ingredient extracted from the fruits and seeds of Arctium lappa L., which has been rifely applied in clinical for anti-influenza and metabolic diseases in Asia [4]. Accumulating evidences have extensively revealed that arctigenin possessed therapeutic efficacies in numerous diseases, such as acute lung and liver injury [5–7], autoimmune hepatitis [8], inflammatory bowel diseases [9–11], and multiple sclerosis [12], which were largely accompanied by suppressing the abnormal activation of immune cells and modulating conventional inflammatory signaling pathways. Nevertheless, the molecular mechanism and potential targets of arctigenin in regulating inflammation remain elusive.
Phosphodiesterases (PDEs) act as the intracellular enzymes, hydrolyzing the cyclic adenosine monophosphate (cAMP) and cyclic guanosine monophosphate (cGMP), which are second messengers and modulators of various physiological processes [13,14]. Significant upregulation of PDE4 activity and distribution was observed in the peripheral immune cells and impaired skin biopsies in patients with psoriasis and other inflammatory diseases [15,16]. Accumulating evidences have demonstrated that inhibition of PDE4 resulted in significant upregulation of cAMP and subsequent activation of protein kinase A (PKA) [17,18]. Thereafter, phosphorylation of nuclear transcriptional factors, such as cAMP-responsive element binding protein (CREB), cAMP responsive element modulator (CREM), and activated transcription factor 1 (ATF-1), collectively regulated inflammation and tissue homeostasis. In the past decades, PDE4-targeted drug discovery has been widely accepted as a competitive and promising strategy for anti-inflammatory agents’ design. Roflumilast, apremilast, and crisaborole have been licensed for patients who suffered from chronic obstructive pulmonary disease (COPD), psoriatic arthritis (PsA), and atopic dermatitis, respectively [19].

Psoriasis, with a high incidence worldwide, is an autoimmune disease, which refers to something wrong with the body’s immune system. It is a multifactorial and inflammatory skin disorder that is distinguished by the local formation of silver plaques and skin abnormal thickness [20]. The pathological mechanism of psoriasis is closely associated with the communication between primary tissue-resident and newly-recruited inflammatory cells, including macrophages, dendritic cells (DCs), and activated T cells, along with the increased production of corresponding cytokines and chemokines [21,22]. Upon exposure to the internal and external harmful stimulus, including physical trauma, bacterial infection, and endotoxins, inflammatory cells are activated to overexpress interleukin-12 (IL-12), IL-17A and IL-23, leading to excessive proliferation of keratinocytes [23,24]. In addition, the abnormal expression of chemokines and adhesion molecules derived from keratinocytes, in turn, will promote the infiltration of an additional variety of immune cells, thereby exacerbating the inflammatory environment of the skin psoriasis.

In the present study, co-crystallization and isothermal titration calorimetry were performed to uncover the binding properties of arctigenin with the catalytic domain of PDE4D. Further functional verifications were performed, including PDEs enzymic inhibition assay, and inflammatory mediator production from human peripheral blood mononuclear cells (PBMCs) and RAW264.7 cells. Moreover, an imiquimod (IMQ)-induced experimental psoriasis model was conducted to explore the efficacies and potential mechanism of arctigenin.

Materials and methods

Crystal structure determination

The protein co-crystallization of the PDE4D catalytic domain in complex with arctigenin was performed by the protocols reported previously [25]. The X-ray diffraction data were collected and analyzed at the Shanghai Synchrotron Radiation Facility (SSRF, BL17U1, PILATUS 6 M detector). The relevant data of refinement statistics are given in Supplementary Table S1.

PDEs enzyme assay

The enzymatic activity of the purified catalytic domain of PDE4D, PDE5A, PDE7A, PDE9A, and PDE10A, and full-length PDE1A, PDE2A, PDE3A, PDE6C, PDE8A, and PDE11A, purchased from BPS Bioscience (San Diego, CA, USA), was performed by determining the biochemical hydrolysis of [3H]-cAMP or [3H]-cGMP into [3H]-AMP or [3H]-GMP, respectively, using the scintillation proximity assay (SPA) as described previously [25]. The enzymatic activities in RAW264.7 cells and skin biopsies were normalized to the concentration of total proteins.

Isothermal titration calorimetry

The binding thermodynamic profile of arctigenin to PDE4D catalytic domain was measured by the isothermal titration calorimetry (ITC) method using the iTC-200 calorimeter (Little Chalfont, Buckinghamshire, UK) as described previously [25]. The enthalpy change and association constant were obtained by nonlinear regression of the data using Origin software (San Clemente, CA, USA).

PDK1 kinase assay

For measurement of the inhibitory effect of arctigenin on PDK1, we used a PDK1 kinase enzyme assay (V2761, Promega (Beijing) Biotech, China) from Promega according to the manufacturer’s instruction. Recombinant full-length human PDK1, ATP, substrate and tested compounds were diluted with kinase buffer that comprised 40 mM Tris (pH 7.5), 20 mM MgCl$_2$, 0.1 mg/mL BSA, and 50 μM DTT to a concentration of 20 nM, 1 ng/μL, 20 μM and from 100 μM to 10 nM, respectively, before adding to the wells of 384 low volume plate. After incubating at room temperature for 60 min, 5 μL of ADP-Glo™ reagent was added to each well of plate and incubated at room temperature for another 40 min. Then adding 10 μL of kinase detection reagent and incubating at room temperature for 30 min before recording luminescence with the Bio-Tek Synergy4 plate reader.

Lipopolysaccharide stimulation of human PBMCs

The inhibitory activity of arctigenin on lipopolysaccharide (LPS, Sigma-Aldrich, St. Louis, MO, USA) stimulated human PBMCs was assayed as previously reported [25]. Human PBMCs (1 × 10$^6$/well) were purchased from SALILY BIO (Shanghai, China) and maintained in the RPMI-1640 medium (Gibco, Grand Island, NY, USA) in the presence of 10% FBS (Gibco) and 2 mM L-glutamine.

Establishment and assessment of IMQ-induced murine psoriatic skin lesions

Female wild-type BALB/c mice (6–8 weeks old, 20–22 g) were acquired from Shanghai Laboratory Animal Center of the Chinese Academy of Sciences. Mice were fed with standard laboratory chow and water under the specific pathogen-free (SPF) facility with 12 h of light/12 h of dark cycle, 24 ± 2 °C and 55 ± 5% relative humidity. All experiments in the present study were conducted as the previous description according to the National Institutes of Health (NIH) Guide for Care and Use of Laboratory Animals in compliance with the ARRIVE guidelines and were approved by the Bioethics Committee of the Shanghai Institute of Materia Medica (SIMM), Chinese Academy of Sciences (CAS) [26].

The dorsal skins of female mice were shaved with an electric razor and depilatory creams. Then, all mice were randomly divided into 3 independent experimental groups (n = 7 mice per group), namely normal, vehicle (IMQ-treated only), and arctigenin treatment (IMQ with 5% arctigenin cream). The psoriatic inflammatory skin lesions were established by topical application with 62.5 mg daily of 0.05% IMQ cream (Sichuan Mingsxin Pharmaceutical, Sichuan, China) on the shaved back skin for 7 successive days. In
the course of arctigenin treatment, murine body weight in each
group and skin inflammatory scores, referring to Psoriasis Area
and Severity Index (PASI, Supplementary Table S2), were mon-
tored blind by 3 investigators, as previous description [27].

Histological examination

Skin sections were fixed in 4% formaldehyde buffered in PBS
solution and then embedded in paraffin. Subsequently, 3 μm sec-
tions were cut and stained with hematoxylin and eosin (H&E)
reagents. The representative pictures were observed and obtained
under a light microscope (Olympus IX73, Tokyo, Japan). Accord-
ingly, the epidermal thickness in all groups was calculated and
recorded by the distance between the interfollicular epidermis by
using the Image-Pro Plus software (Silver Springs, MD, USA).

Protein extraction, quantization, and LC-MS/MS analysis

The skin samples from three groups were transferred into low
protein binding tubes and lysed with digestion buffer. Digested
peptides were centrifugated at 12000 rpm for 20 min and collected
for further Tandem Mass Tags (TMT)-labeling as following: Normal
groups, TMT10-126, TMT10-127 N, TMT10-127C; Vehicle groups,
TMT10-128 N, TMT10-128C, TMT10-129 N; Arctigenin groups,
TMT10-129C, TMT10-130 N, TMT10-130C; Group Mix, TMT10-131.

All proteomic analyses were conducted by the Q-Exactive HF
mass spectrometer (Thermo Fisher Scientific). Samples were then
separated by a C18 column (15 cm × 75 μm) on an EASY-nLC™
1200 system (Thermo Fisher Scientific). ProteomeDiscoverer
(v.2.3, Thermo Fisher Scientific) was used to search all the raw data
thoroughly against the sample protein database (uniprot-reviewed-Mus musculus (Mouse)-201908.fasta). Functional analy-

Flow cytometry analysis

Skin single cell suspensions from the inflamed biopsies were
prepared and isolated as the previous description [28,29]. In brief,
skin samples were minced and digested with collagenase IV
(1 mg/mL, Sigma-Aldrich) in the RPMI-1640 media supplemented
with 10% FBS at 37 °C incubator for 2.5 h, and subsequently filtered
through a 70-μm mesh to acquire the single cells. The skin cell sus-
pensions were washed twice with cold PBS and then incubated
with FVD eFluo™ 780 (Thermo Fisher Scientific) at 4 °C for
30 min to differentiate the viable cells from dead cells. Upon being
blocked with staining with anti-CD16/CD32 mAb (Thermo Fisher Sci-
entific), cells were further incubated with fluorescein isothio-
cyanate (FITC), phycoerythrin (PE), peridinin-chlorophyll
proteins-Cyanine5.5 (Percp-Cy5.5), allophycocyanin (APC),
brilliant violet 421 (BV421), and brilliant ultraviolet 395
(BUV395)-conjugated antibodies. Data were analyzed using FlowJo
software (Tree Star, Ashland, OR, USA).

Cell cultures and treatment

RAW264.7 cells, murine macrophage cell line, were obtained
from the American Type Culture Collection (ATCC, Manassas, VA,
USA) and maintained in Dulbecco’s modified Eagle’s media
(DMEM, Gibco) containing 10% FBS (HyClone), 2 mM of
L-glutamine, 100 U/mL penicillin, and 100 μg/mL streptomycin in
a humidified incubator of 5% CO2 at 37 °C. RAW264.7 cells were
lysed with arctigenin (100 μM), H89 (10 μM, an inhibitor of
PKA, Selleck Chemicals, Houston, TX, USA), and/or forskolin
(10 μM, an activator of adenylyl cyclase, Sigma-Aldrich) for
15 min and then cells were collected for western blotting analysis.

ELISA and Luminox assay

Serum inflammatory cytokines were quantified by the Luminex
200 instrument (Merck Millipore, Billerica, MA, USA) using a Th1/
Th2/Th9/Th17/Th22/Treg Cytokine 17-Plex Mouse ProcartaPlex™
Panel (Thermo Fisher Scientific). Cytokines in RAW264.7 cell
supernatants were analyzed by mouse TNF-α and IL-6 ELISA kits
(BD Biosciences, San Diego, CA, USA) as described previously
[16]. For cAMP detection, cells and psoriatic skin biopsies were
lysed and isolated by 0.1 M HCl and the intracellular cAMP level
was quantified by using the direct CAMP ELISA kits (Enzo Life
Sciences, The Netherlands) according to the directions.

Immunofluorescence and immunohistochemistry

For immunofluorescent evaluation, cells were fixed in 4% for-
maldehyde, blocked with the BSA blocking buffer (Beyotime
Biotechnology, Shanghai, China), and stained at 4 °C overnight with
phospho-CREB antibodies (Cell Signaling Technology). The skin
slices were then incubated with FITC-conjugated goat anti-rabbit
secondary antibodies and counterstained with DAPI to stain the
nuclei of RAW264.7 cells. For immunohistochemistry analysis,
paraffin-embedded skin tissue slices were blocked with 3% H2O2
solution, retrieved by citrate buffer solution and then incubated
at 4 °C overnight with anti-Ki-67 antibodies (Abcam, Cambridge,
UK), and CD3 Abs (Abcam). Then, the positive signals were visual-
ized by biotinylated horse anti-rabbit secondary antibodies and
streptavidin horse peroxidase. The representative pictures
were obtained using the light microscope (Olympus IX73
microscope).

RNA extraction and RT-PCR

Total RNA from the murine skin biopsies and RAW264.7 cells
was isolated and purified by using the RNAsimple total RNA kits
(Tiangen). Subsequently, total RNA (500 ng per unit) was reversely
transcribed by the Hifair™1 1Strand cDNA Synthesis SuperMix
regents (YeaSan, Shanghai, China) with consecutive incubation of
25 °C for 5 min, 42 °C for 30 min. and 85 °C for 5 min. Real-time
quantitative PCR analyses were conducted with the SYBR® Green
Realtime PCR Master Mix regents (Yeasen) according to the direc-
tions on the 7500 Fast Real-Time PCR System (Applied Biosystems,
Foster city, CA, USA) for PCR amplification with holding stage
(95 °C for 30 s), cycling stage (95 °C for 15 s and 60 °C for 40 s
for 40 cycles), and melt curve stage using the gene-specific primers
in Supplementary Table S3 by the ΔΔ Ct method.
Western blotting assay

The protein concentrations from the inflamed skin biopsies and RAW264.7 cells were measured and quantified by the Pierce BCA protein assay kits (Thermo Fisher Scientific). The equal protein amounts were dissolved in the SDS-loading buffer and subjected to 10% electrophoretic SDS-PAGE, then transferred into the nitrocellulose membranes (Bio-Rad, Hercules, CA, USA). The nitrocellulose membranes were blocked for 10 min at room temperature with SuperBlock T20 blocking buffer (Thermo Fisher Scientific) and then incubated at 4 °C overnight with primary antibodies. The positive signals were visualized by using the SuperSignal West Femto Maximum Sensitivity Substrate (Thermo Fisher Scientific) under the exposure of ChemiDoc™ MP Imaging System (Bio-Rad).

Statistical analysis

All data in the present research were shown as mean ± SEM and the statistical variations were performed by using one-way ANOVA (3 and more groups) or Student's t-test (2 groups) with Dunnet's multiple comparisons test with no significant variance inhomogeneity by GraphPad Prism (GraphPad Software, Version 6.0, LLC, CA, USA). P-values of <0.05 (p < 0.05) were regarded as statistical significance.

Results

Arctigenin could specifically bind to the catalytic domain of PDE4D

To discover novel PDE4 inhibitors, we screened our in-house natural products library containing ~ 500 natural products with good solubility and drug-like properties by a scintillation proximity assay (SPA) and found that arctigenin showed an acceptable physicochemical property (Table S4) and moderate inhibitory activity toward PDE4D with an IC₅₀ of 3.76 ± 0.28 μM (Fig. 1A). As illustrated in Fig. 1B, the dimethoxybenzene ring of arctigenin formed two hydrogen bonds with the conserved Q369 and a face-to-face π – π interaction with F372. H160 at the metal binding site formed another hydrogen bond with arctigenin (Fig. 1B). The inhibitory activity as well as the binding mode of arctigenin were well consistent with the thermodynamic properties of arctigenin revealed by the ITC analysis, which suggested the enthalpy contribution is dominant for binding of arctigenin to the catalytic domain of PDE4D with a single digit micromolar binding affinity (Fig. 1C and Table S5).

Arctigenin was highly specific and selective for PDE4 inhibition

Previously, it has been indicated that arctigenin could modulate the PDK1/AKT/PDE4D axis in asthma therapy [30]. We intended to uncover the direct binding of arctigenin with PDK1, whereas the measured inhibition of arctigenin on PDK1 seemed negligible, compared with that of a known PDK1 inhibitor, GSK2334470 (Table S6). The selectivity of arctigenin against other PDE families was also measured. Arctigenin showed weak inhibitory activities against PDE1A, PDE2A, PDE3A, PDE5A, PDE6C, PDE7A, PDE8A, PDE9A, and PDE11A (Table S7). These data indicate that arctigenin is not a PDK1 inhibitor but a PDE4 inhibitor with a relatively high selectivity over other PDEs.

Arctigenin suppressed the secretion of inflammatory mediators in vitro

Since PDE4 was highly expressed in immune cells, arctigenin was further evaluated for the inhibitory activity on the TNF-α production from the LPS-stimulated human PBMCs. As shown in Fig. 1D, arctigenin suppressed the TNF-α production with an IC₅₀ of 35.18 ± 6.01 μM. To further confirm the anti-inflammation effects of arctigenin, murine RAW264.7 cells were included in the current research. Arctigenin could dose-dependently (100, 30, and 10 μM) reduce the production of TNF-α and IL-6 from LPS-induced RAW264.7 cells, which was not associated with the cytotoxicity of arctigenin (Fig. S1). Conformably, arctigenin also decreased the mRNA expression level of TNF-α and IL-6 in RAW264.7 cells (Fig. S2A–B). Moreover, arctigenin downregulated the inflammatory mediators, iNOS and COX-2, in a concentration-dependent manner (Fig. S2C), which was in line with the previous report [31]. Consequently, the accumulation of nitrite, the stable and product of NO by RAW264.7 cells, was dramatically reduced following incubation with arctigenin (Fig. S2D).

Arctigenin activated cAMP-dependent phosphorylation of CREB in RAW264.7 cells

Due to the direct PDE4 inhibition by arctigenin, we further aimed to find out whether arctigenin displayed direct effects of PDE4-mediated signaling pathway in regulating inflammation [32]. Prospectively, arctigenin addition led to significant elevation of intracellular cAMP in RAW264.7 cells in a concentration-dependent manner (Fig. S2E). Linked with previous research, cAMP accumulation by arctigenin subsequently led to the phosphorylation of CREB without any effects on the total expression level of CREB and PKA protein (Fig. 2A and Fig. S3A). The results were confirmed by immunofluorescent staining with phospho-CREB, which showed that localization of phospho-CREB largely co-existed with the nuclei of RAW264.7 cells (Fig. 2B).

We further found that H89 intervention obviously blocked the effects of arctigenin on phosphorylation of CREB, whereas, the adenylyl cyclase activator, forskolin, could further enhance the effects of arctigenin on promoting the phosphorylation of CREB (Fig. 2C and Fig. S3B). Further biological investigation confirmed the function of arctigenin on the secretion of TNF-α (Fig. 2D). Accordingly, siRNA targeting for reducing the expression of PKA were transfected, compared with NC controls, the protein level of PKA was mostly decreased in transfected-RAW264.7 cells (Fig. 2E and Fig. S3C). In line with the effects of H89 treatment, phosphorylation of CREB and inhibition of TNF-α secretion were hindered whether the cells were incubated with arctigenin and/or forskolin (Fig. 2E and 2F). In brief, our research indicated PKA functioned as the crucial factor in the anti-inflammation of arctigenin.

Topical application of arctigenin significantly alleviated IMQ-induced psoriatic skin inflammation

Given the anti-inflammatory activity and intrinsic physicochemical property of arctigenin, we therefore uncovered the potential therapeutic effects of arctigenin in IMQ-induced murine psoriatic skin lesions. As demonstrated in Fig. 3A–D, upon topical application of IMQ cream, severe body weight loss, typical skin lesions, including scales, thickness, and erythema appeared. Therapeutically, topical application of arctigenin treatment (5% specification) significantly reversed the body weight loss and attenuated the experimental manifestations and decreased the cumulative PASI scores of the shaved skin (Fig. 3A–D). Histologically, arctigenin treatment dramatically improved the epidermal thickening and immune cells recruitment in the inflamed tissue microenvironments (Fig. 3E–F). In addition, arctigenin could ameliorate the concomitant splenomegaly (Fig. 3G). Further immunohistochemistry with Ki-67 also elucidated hyperplasia of cuticular layers and the positive staining signals in the epidermis were reduced following arctigenin treatment (Fig. 3H). Moreover, Luminex assay of serum cytokines showed that arctigenin could reduce the production of...
inflammatory cytokines and promote the secretion of IL-10 (Fig. 3I). These data collectively suggested that topical application of arctigenin showed excellent therapeutic capacity for psoriatic inflammation.

Arctigenin rectified the immune dysfunction and hyperactivation of keratinocytes in the skin microenvironment

To fully reveal the pathological changes at molecular levels, the skin samples were collected and analyzed for TMT 10-labeling quantitative proteomic analysis (Fig. 4A). Principle component analysis (PCA) displayed that vehicle group were clearly separated from normal group and arctigenin-treated group by the first principle components which declared 34.82% of the variability of the identified proteins (Fig. 4B). In our analysis, there were 806 differentially expressed proteins between normal and vehicle group; 134 proteins between vehicle and arctigenin-treated groups (Fig. 4C). Among these proteins, by contrast to vehicle group, 41 proteins were upregulated in the arctigenin group, whereas 93 proteins were downregulated (Fig. 4D), which were also demonstrated in the volcano plot (Fig. 4E). Hierarchical cluster analysis of differential protein expression level demonstrated that compared with vehicle group, arctigenin-treated group displayed a specific cluster, which predicting the profound proteomic changes between vehicle and arctigenin-treated group (Fig. 4F).

As illustrated in Fig. 4G, Correlation analysis of differentially expressed proteins, using Pearson algorithm, indicated that there existed an extreme close association between multiple proteins. Further functional analyses of differentially expressed proteins were performed to excavate gene ontology (GO) annotation, including the cellular component, biological process, and molecular function (Fig. S4A). From the top30 GO term, the differential proteins were mainly linked to toll-like receptor signaling, secretion and activity of inflammatory cytokines, regulation of MAPK cascade, expression of keratin filaments (Fig. S4A). Comparison of differential proteins and all proteins in GO level 2 further confirmed that cell adhesion and junction, immune system process, extracellular matrix, antioxidant, and chemoattractant activity largely and differentially appeared in vehicle and arctigenin-treated group (Fig. S4B and Table S8). Moreover, KEGG pathway enrichment of differential proteins also referred to the immune dysfunction and hyperproliferation and abnormal activation of cells in the skin microenvironment (Fig. S4C). The network between vehicle and arctigenin-treated group was visualized by Cytoscape 3.1.1 (Fig. S4D). According to the Score Sequest HT of differentially expressed proteins, Keratin 14 attracted our attention with the highest score (1413.7). Enlightened by the findings, we naturally focused on the proteins possessing the interaction with Keratin 14 in the PPI network (Fig. S4E), which were further confirmed by the mRNA expression level of these proteins, including keratin 6, keratin 8, MMP3, Adgre1, and Integrin M (Fig. S4F).

Arctigenin modulated the PDE4-mediated signaling and attenuated the inflammatory responses in the local skin lesions

Given the critical role of PDE4 in regulating multiple inflammatory processes, we then uncovered the pathological role of PDE4 in IMQ-induced psoriasis. The results demonstrated that increased
activity of cAMP-specific PDEs was observed in the vehicle mice, which could be obviously reversed by arctigenin treatment (Fig. 5A). Consistently, arctigenin could upregulate the skin level of cAMP (Fig. 5B) and promote the phosphorylation of CREB (Fig. 5E). In accordance with the results from RAW264.7 cells, arctigenin could inhibit the enzymatic activity of PDE4 in the concentration-dependent manner (Fig. 5C-D). As a consequence, arctigenin extensively attenuated the expression of several inflammatory cytokines (Fig. 5F).

To confirm the results of proteomics and fully uncover the local inflammatory microenvironments in the inflamed biopsies, the skin single-cell suspensions were determined by flow cytometric analysis. Skin-infiltrated leukocytes (CD45+), antigen-presenting cells (APCs, CD45+MHC II+), T cells (CD45+CD3+), and γδ T cells (CD3+γδTCR+) were reduced upon arctigenin treatment (Fig. S5A-B), which were also found in the skin sections staining with CD3 (Fig. S5C). Further analysis of innate immune cells demonstrated that monocytyc myeloid cells (CD11b+), macrophages (CD11b+ F4/80+), neutrophils (CD11b+Gr-1+), and DCs (CD11b+CD11c+) were largely accumulated in the vehicle group, which were decreased when treated with arctigenin (Fig. S5D). In line with inflammatory recruitment, arctigenin suppressed the protein expression of CCR6 and CXCR3 on CD45+ leukocytes analyzed by flow cytometry (Fig. S5E). Additionally, gene expression levels of multiple inflammatory mediators and chemokines, including RANTES, iNOS, MIP-1α, MIP-1β, MIG, and MIP-3α, were obviously elevated during induction of psoriasis and treatment with arctigenin could suppress the expression of chemokines in inflamed skins (Fig. S5F-G).

**Discussion**

The bioactive lignans, arctigenin and its mono-glycoside, arctiin, are mainly derived from the medicinal herb *Arctium lappa* L. and have been frequently investigated in multiple inflammatory diseases [4]. Previously, arctigenin was effective in relieving the experimental symptoms of pulmonary inflammation and tissue damage, mainly related to inhibition of PI3K/AKT, MAPK, and iNOS signaling pathway [5,7,33]. Recently, a phase I single dose study was designed to determine the maximum tolerated dose (or recommended dose) and bioavailability of arctigenin in healthy volunteers (Clinicaltrials.gov., NCT03703388). Though arctigenin demonstrated broad modulatory effects in various disease models, the potential target remains unclear and ill-defined. In our research, PDE4-targeted ligand screening based on our in-house natural products library led to the discovery of arctigenin as novel inhibitors of PDE4 with a single digit micromolar potency. A fur-
ther crystallography together with thermodynamic study illuminated that arctigenin could bind to the catalytic domain of PDE4D with a typical binding conformation and with a single digit micromolar binding affinity (Fig. 1 and Table S4-S6). To our knowledge, this is the first study which clearly demonstrates the detailed molecular mechanisms of arctigenin binding to a specific target protein in modulating inflammation (Fig. 6).

Over the past few decades, numerous PDE4 inhibitors have been designed in the field of autoimmune and inflammation-related diseases [19]. However, due to the unexpected permeability of blood–brain barrier and other severe adverse effects, a large number of PDE4 inhibitors were impeded and restricted in further applications [17,26]. It is worth informing that the tissue accumulation of arctigenin in brain was lower than that in intestine, liver, kidney, lung, and heart, which might account for the safety profile of arctigenin [34]. In addition, following oral administration, the t1/2 was short (<2h) and the plasma concentration of arctigenin was even undetectable in several animal species, which substantially hindered the therapeutic efficacies in vivo [4,35]. Therefore, there is an urgent need for alternative dosing procedures to improve systemic or target organ exposure. Unlike systemic treatment, topical application might directly touch the inflamed positions and result in higher drug concentration in skin than that by oral administration. Accordingly, we indeed found that topical treatment of 5% arctigenin cream showed a better therapeutic effect than oral administration of 100 mg/kg arctigenin (Fig. S6). Significantly, arctigenin has a good transdermal absorption, and a high concentration of arctigenin could be detected in the inflamed skin tissues (29.67 ± 15.42 μg/g skin samples), which further ensures the therapeutic effect of arctigenin in treating psoriasis.
Psoriasis is an inflammatory skin disease involving immune cells and innate and adaptive immune molecules [36,37]. The pathogenesis of psoriasis is multifactorial and complicated, and is often associated with genetic susceptibility, disorders of the skin’s microbes and immune system, and environmental triggers [20]. To fully understand the histopathological changes at the protein levels, TMT10-labeling quantitative proteomic analysis was performed to uncover the potential mechanism. There existed unique distribution patterns among three groups from PCA analysis and differentially expressed proteins were identified between vehicle and arctigenin-treated groups (Fig. 4B-F). Further functional analyses of differentially expressed proteins, including GO annotation and KEGG PPI network, suggested that the potential mechanism of arctigenin, from the protein level, were mainly related to toll-like receptor signaling, activity of inflammatory cytokines, antioxidant and chemoattractant activity, and expression of keratin filaments (Fig. S4A-B). Keratinocytes, the prominent parenchyma cell in the skin epidermis, function as the impressive barrier to...
separate internal components from external stimuli; Meanwhile, keratins form the predominant intermediate filament cytoskeleton of keratinocytes, in which keratins represent the major constituent of proteins in keratinocytes [38]. Previous reports have indicated hyperproliferation-associated keratins were considered to be the hallmarks of psoriasis [39,40]. Among those proteins, Keratin 14 and its associated proteins, including keratin6, keratin8, MMP3, Adgre1, and Integrin M, mostly functioned in the progression of psoriasis and the process of arctigenin treatment (Fig. S4C-F).

**Conclusion**

The present research indicated that arctigenin could occupy the catalytic binding site of PDE4D and showed inhibitory activity...
against the degradation of cAMP for the first time. As a result, arctigenin displayed dramatic anti-inflammation in LPS-primed human PBMCs and murine RAW264.7 cells, which was largely owing to activation of the cAMP-dependent PKA-CREB signaling. Further proteomics research confirmed that arctigenin could rectify the immune dysfunction and hyperactivation of keratinocytes in the inflamed skin microenvironments, which might be largely associated with the expression of Keratin 14. In sum, our study provided the credible evidence that inhibition of PDE4 by arctigenin might function as the potential therapeutic approach in treating inflammatory disorders.

Author contributions

H Li, XL Zhang, W Tang, and YC Xu designed the study. H Li, XL Zhang, CG Xiang, MT Liu, CL Feng, C Fan, HX Su, Y Zhou, and Q Qi performed the experiments. H Li and XL Zhang analyzed the experimental data and contributed to writing the manuscript. W Tang and YC Xu obtain the funding and revised the manuscript. All authors approved the final version of the manuscript.

Compliance with ethics requirements

Female BALB/c mice (6–8 weeks, 20–22 g) were acquired from Shanghai Laboratory Animal Center of the Chinese Academy of Sciences. All experiments were carried out according to the National Institutes of Health Guide for Care and Use of Laboratory Animals in compliance with the ARRIVE guidelines and were approved by the Bioethics Committee of the Shanghai Institute of Materia Medica (IACUC: 2018–03-ZJP-70).

Declaration of Competing Interest

The authors declared that there is no conflict of interest.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.jare.2021.02.006.

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Fig. 6. Schematic illustration of the effects of arctigenin, as a potent PDE4 inhibitor, involved in attenuating psoriatic skin inflammation.
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