Interleukin-22 participates in the inflammatory process of vitiligo

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ABSTRACT

Vitiligo is an acquired depigmentary skin inflammatory disorder. The pathogenesis of inflammatory skin disease involves the release of cytokines from keratinocytes, including interleukin (IL)-1β. IL-22 belongs to a family of cytokines structurally related to IL-10, including IL-19, IL-20, IL-24, and IL-26. In contrast to IL-10, IL-22 has proinflammatory activities. Among skin cell populations only keratinocytes are the major targets of IL-22. In the present study, we demonstrated that IL-22 promoting IL-1β secretion from keratinocytes via the Reactive oxygen species (ROS)-NOD-like receptor family, pyrin domain containing 3 (NLRP3)-caspase-1 pathway. It inhibited the expression of protease-activated receptor-2 (PAR-2) of keratinocytes. However, IL-22 had no direct effect on normal human foreskin-derived epidermal melanocytes (NHEM). Considering the closely connection between keratinocytes and melanocytes, and the ability of keratinocytes to produce a plethora of cytokines, in the present work, we examined whether IL-22 could regulate melanocytes functions by keratinocytes participation. Keratinocytes were exposed to IL-22 and the conditional medium was collected. The effect of conditional medium on melanocytes was studied. The expressions of relative proteins were assessed by western blot. Influence of conditional medium on NHEM migration was assessed by Transwell method and the apoptosis by flow cytometry analysis. The IL-22-treating keratinocytes conditional medium inhibited melanogenesis and restrained the expressions of Rab GTPases of NHEM. In addition, the conditional medium suppressed melanocytes migration and induced apoptosis. Our results collectively indicated that IL-22 may potentiate IL-1β-mediated skin inflammation and result in participating in the inflammatory pathogenesis of vitiligo.

INTRODUCTION

Skin works as a bio-barrier against outside influences and also as pigmentary system [1, 2]. As playing protective functions, the skin mobilizes a large amount of cell populations to act in a coordinated fashion. Skin's neuroimmunoendocrine axes is established by epidermal keratinocytes, melanocytes, and dendritic Langerhans cells (LCs), combining with dermal fibroblasts, macrophages, mast cells and others [3, 4]. These communication and interaction among cells are passed through the multiply kind of molecules termed cytokines. Cytokines play
an important role in modulation of body homeostatic response. They are classically secreted “messenger” proteins that allow for cell-cell communication [5, 6].

Vitiligo is an intriguing depigmentary disorder affecting approximately 0.5-2% of the world population. Numerous factors have participated in the development of vitiligo, histologic evidence indicates that vitiligo is an inflammatory disease with development of a lichenoid tissue reaction [7]. It develops due to progressive disappearance of epidermal melanocytes. Melanocytes can detect and decode the solar or thermal energy, or respond to biological of physicochemical signals to sense the environment. All these functions of melanocytes promote forming a hypothesis that melanocytes are ‘neurons of the skin’ [8].

In the skin, inflammasome protein NLRP3 can be activated by UV exposure and sensitizers, which induces IL-1β production [9, 10]. IL-1β mRNA expression was elevated in the lesioned edge of vitiligo [11]. Upregulation of IL-1β transcript in patients advocates its possible role in autoimmune pathogenesis of vitiligo [12]. When co-cultured with activated T cells, IL-1β secretion and caspase-1 expression was increased of keratinocytes [13]. IL-22 belongs to IL-10 family, which plays a vital role in various inflammatory and infectious diseases. IL-22 is primarily produced by obvious immune cells, and Th22 cells are a major source of IL-22 in many diseases [14–16]. When treated with IL-22, keratinocytes could be triggered to induce inflammatory responses [17]. The functional heterodimeric receptor of IL-22 comprised of IL-10RB and IL-22RA1 subunit [18–20]. IL-10RB subunit is the ubiquitously expressed one while the IL-22RA1 subunit is more restricted. IL-22RA1 subunit is mainly expressed on epithelial cells, keratinocytes, and hepatocytes, but are mostly absent from cells of hematopoietic origin [21–23]. The restricted expression of IL-22RA1 defines that the target tissues of IL-22 are special correspondingly, including skin, small intestine, lung, kidney and so on [24, 25]. The downstream signaling of IL-22–IL-22R system is mediated through the Janus kinase (JAK) –signal transducer and activator of transcription (STAT) pathway, primarily activate STAT3 [26, 27].

Considering the closely connection between keratinocytes and melanocytes, and the ability of keratinocytes to produce a plethora of cytokines [28], we evaluated the influence of IL-22 on melanocytes functions. We demonstrated that IL-22 could regulate melanocytes functions by keratinocytes participation.

RESULTS

IL-22 induces the anti-microbial peptide and chemokines from HaCaT cells

To explore the functional role of IL-22 in the production of proinflammatory mediators, some of IL-1β-induced inflammatory mediators were determined, including CXCL1, CXCL5, CXCL8, BD1, LL37, S100A7, S100A8, S100A9 and proteins of the matrix metalloproteinase 3 (MMP3). IL-22 stimulation increased transcription of these mediators (Figure 1A). According to the literature, the IL-
IL-1β facilitates specific inflammatory mechanism of IL-22 in the skin disorders. As shown in Figure 1B, IL-22 increased expression level of NLRP3, caspase-1, and mature IL-1β. This revealed that IL-22 might provoke NLRP3-mediated pathway to increase IL-1β secretion.

**IL-22–IL-22R system participates in the process of IL-1β production via the NLRP3 pathway**

The anti-IL-22R antibody was applied to block downstream signaling. The expression of NLRP3 and caspase-1 (Figure 2A) were decreased, as well as the IL-1β production following IL-22R blockage (Figure 2B). NLRP3 knockdown experiments were performed to confirm IL-22 induced IL-1β increase via the NLRP3 pathway (Figure 2C). Caspase-1 and IL-1β expression levels stimulated with IL-22 were reduced by NLRP3 siRNA transfection, as well as IL-1β secretion in culture supernatants examined by Elisa (Figure 2D). The downstream signaling of IL-22–IL-22R system was also detected, the representative molecule, phosphorylated STAT3 expression level was not influenced obviously by NLRP3 siRNA-transfection (Figure 2E). Therefore, our data showed that the activation of caspase-1 and accompanying IL-1β production was mediated by NLRP3 participation.

**Figure 2: IL-22–IL-22R system participates in the process of IL-1β production via the NLRP3 pathway.** (A) To block IL-22R-mediated signaling, pre-treatment of HaCaT cells with 50 μg/ml of anti-IL-22R blocking antibody for 2 h prior to IL-22 stimulation resulted in reduced NLRP3 and caspase-1 expression (B) as well as a consequent reduction in IL-1β secretion as measured by ELISA. (C) To reduce NLRP3 expression in HaCaT cells, we transfected NLRP3 siRNA or control siRNA into HaCaT cells. The level of protein was quantified to evaluate the knockdown efficiency of the endogenous NLRP3 protein in HaCaT cells. Data from three independent experiments for protein expression are shown. (D) The active form of IL-1β in culture supernatants was quantitated by ELISA. (E) NLRP3 knockdown in HaCaT cells resulted in a reduction in the levels of caspase-1 and the active form of IL-1β in IL-22-treated cells. Representative blots of three independent experiments are shown. Results were normalized against β-actin expression. The data are expressed as means ± SD (i.e. n=3) (*P < 0.05). Ab, anti-IL-22R blocking antibody; IgG, isotype antibody for the negative control of IL-22R antibody.
IL-22 induces IL-1β secretion by activating NF-κB to generate ROS

It has been reported that ROS can activate the NLRP3 inflammasome [29]. The ROS level was measured after IL-22 treatment in this study. As results showed that IL-22 promoted ROS formation significantly (Figure 3A). The role of ROS in activating NLRP3-caspase-1 pathway to produce active IL-1β was determined by the application of NAC (Figure 3A). NAC blocked the generation of ROS induced by IL-22, accompanying the decrease of NLRP3, caspase-1 and IL-1β expressions (Figure 3B and 3C). Additionally, IL-22 induces NF-κB activation (Figure 3D). Pretreated with the NF-κB inhibitor SN50 resulted in the reduction of NF-κB levels (Figure 3D) and the generation of ROS (Figure 3E). These results demonstrate that IL-22 induces ROS production via NF-κB activation.

IL-22 inhibits PAR-2 expression in HaCaT keratinocytes

PAR-2 is expressed on keratinocyte, which can regulate melanosome transfer [30]. We examined the effect of IL-22 on PAR-2 expression of keratinocytes. As shown in Figure 4A, treatment of IL-22 for 24 h decreased PAR-2 expression. We studied the response of keratinocytes to treatment with IL-22 (20 ng/mL) over time ranging from 5 min to 60 min and observed rapid phosphorylation of Stat3 beginning at 20 min after IL-22 treatment (Figure 4B). These results demonstrated that melanocytes responded directly to IL-22 stimulation by activating the JAK/Stat3 signaling pathway. IL-22 also engaged the PI3K/Akt pathway, as the phosphorylation of Akt with peak activation at 60 min following IL-22 treatment (Figure 4B). The effect of IL-22 on Akt was partially suppressed by pre-treatment of keratinocytes with LY294002 (Figure 4C). The inhibition of IL-22 on PAR-2 expression can be interfered by Stat3 inhibitor Stattic, but not LY294002 (Figure 4C and 4D). From these results, we drew a conclusion that the activation of Stat3 was essential for the effect of IL-22 on PAR-2 expression, but not the invoking of Akt.

Effect of IL-22 on melanogenesis of NHEM

As shown in Figure 5A, the purity of cells was identified by immunohistochemical staining with two markers of Tyr and Mitf, and 99% of primary melanocytes were obtained. To determine the effect of IL-22 on melanogenesis, MTT assay was performed as the first step to examine whether IL-22 was cytotoxic to NHEM. As shown in Figure 5B,
IL-22 was not cytotoxic to NHEM within certain limits. NHEM was then exposed to different concentrations of IL-22, tyrosinase activity was measured by the rate of L-DOPA oxidation. As comparing with control group, IL-22 made no effect on the activity of Tyr (Figure 5C), as well as the melanin synthesizing level in NHEM (Figure 5D). The pigment cell-specific melanosomal proteins Mitf, Tyr, TRP-1 and DCT were not responding to IL-22 stimulation (Figure 5E).

**Effect of conditional medium on melanogenesis of NHEM**

Among skin cell populations, keratinocytes are the unique major targets of IL-22. In consideration of the interactive relationship between keratinocytes and melanocytes, we examined the effect of IL-22–treated keratinocytes conditional medium on melanocytes. Three control groups were set, including medium treated only group (Ctrl), 20 ng/ml of IL-22 treated group and conditional medium with no IL-22 treated group (CM-0). Two administered groups were 12 h–IL-22–treated conditional medium group (CM-12) and 24 h–IL-22–treated conditional medium group (CM-24). After 48 h of treatments, MTT assay was performed as the first step. CM-12 and CM-24 showed certain extent suppression on melanocytes (Figure 6A). Then TYR activity and melanin content were measured. CM-12 and CM-24 inhibited the activity of Tyr (Figure 6B) and restrained melanogenesis (Figure 6C). Furthermore, CM-12 and CM-24 suppressed expressions of the melanin synthesis related proteins (Figure 6D).

**Conditional medium inhibits NHEM migration**

The effects of conditional medium on NHEM migration were observed. As shown in Figure 7A,B, CM-12 and CM-24 inhibited the cells migration. It has been reported that Rab GTPases play roles in melanosome maturation or trafficking in melanocytes. Researches have been published showing that Rab7 mainly regulates early and intermediate stage melanosomes and Rab27a primarily work on intermediate and mature melanosomes [31]. In addition, Rab17 acts on melanosomes downstream of Rab27a [32]. Our data shown that CM-12 and CM-24 suppressed expressions of the Rab7, Rab 27a and Rab17 (Figure 7C).

**Conditional medium induces NHEM apoptosis**

The conditional medium induced NHEM apoptosis, as shown in Figure 8A, the percentage of apoptotic cells increased significantly in CM-12 and CM-24 group by flow cytometry analysis. In our studies, CM-12 and CM-24 increased the expressions of Bax, cleaved-caspase 3 and cytochrome c, at the same time decreased Bcl-2 (Figure 8B), hence, Bcl-2/Bax ratio significantly decreased (Figure 8C).

**DISCUSSION**

In mammals, the epidermis forms the top protective covering of skin to provide photoprotection and thermoregulation. This function relies on the principal trait of epidermal melanocytes——‘pigmentation’. Melanin, the pigment responsible for most pigmentation, is most contained by epithelial cells [33]. Except those of the eye producing their own melanin, most pigmented epithelial cells acquire their pigment from melanocytes, such as keratinocytes. On the basis of physiological structure, melanocytes are located in the basal layer of the epidermis, surrounding...
by approximately 36 keratinocytes. Melanocytes and keratinocytes establish a division of activities, that one cell producing pigment, another cell using it [34, 35].

There were literatures reported that Rab7 is involved in the transport of late endosomes/lysosomes [36, 37]. The activated Rab27a can be found on mature melanosomes, which locate in the tips of the dendrites of melanocytes [38–40]. Rab17 resides on melanosome and endosomes (REs) membranes in melanocytic cells, which also regulates melanosome trafficking. It has been proved that Rab27a works as upper stream of Rab 17 on melanosome transport [32]. Multiple enzymatic and structural proteins participate in regulating the maturation of melanosomes, including Tyr, TRP-1, DCT, OA1, antigen recognized by T cells (MART-1), Pmel17/gp100, GPNMB and others [41–43]. The transcription of GPNMB is regulated by MITF [44, 45], which also regulates other pigmentation genes, such as Tyr, TRP-1, DCT, Pmel17, OA1 and so on.

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**Figure 5: Effect of IL-22 on melanogenesis of NHEM.** (A) Identification of primary melanocytes. Cells were labeling with Mitf (green) and Tyr (red) antibodies. Nuclei were labeled with DAPI (blue). (B) After incubation with IL-22 (0, 1, 10, 20 ng/ml) for 48 h, cell viability was determined using a MTT assay. (C) Tyrosinase activity was determined by L-DOPA oxidation as described in ‘Materials and methods’. Stimulation of tyrosinase activity of NHEM with IL-22 (0, 10, 20 ng/ml) for 48 h. (D) Melanin content was performed as described in ‘Materials and methods’ the same for 48 h. (E) Western blot assays were performed to examine Mitf, Tyr, TRP-1 and DCT expression levels, Total cellular proteins (20 μg/lane of NHEM) were subjected to 10% SDS-PAGE. Results were normalized against β-actin expression. Results shown are means ± SD (i.e. n=4). Data were analyzed by One-Way Analysis of Variance (ANOVA) followed by post hoc Tukey test.
**Figure 6: Effect of conditional medium on melanogenesis of NHEM.** (A) After incubation with IL-22 and different conditional medium for 48 h, cell viability was determined using a MTT assay. (B) Tyrosinase activity was determined by $L$-DOPA oxidation as described in ‘Materials and methods’. (C) Melanin content was performed as described in ‘Materials and methods’. (D) Western blot assays were performed to examine GPNMB, Mitf, Tyr, TRP-1, DCT, OA1 and Pmel17 expression levels, Total cellular proteins (20 μg/lane of NHEM) were subjected to 10% SDS-PAGE. Results were normalized against β-actin expression. Results shown are means ± SD (i.e. n=3). Data were analyzed by One-Way Analysis of Variance (ANOVA) followed by post hoc Tukey test. *P < 0.05, **P < 0.01, compared with control.

**Figure 7: Conditional medium inhibits NHEM migration.** (A) NHEM cells were seeded in the upper chamber of transwell. CM-12 and CM-24 inhibited NHEM migration. Scale bar, 50 μm. (B) After 24 h, the migrated NHEMs were quantified. (C) Western blot assays were performed to examine Rab7, Rab27a and Rab17 expression levels, Total cellular proteins (20 μg/lane of NHEM) were subjected to 12 % SDS-PAGE. Results were normalized against GAPDH expression. Results shown are means ± SD (i.e. n=3). Data were analyzed by One-Way Analysis of Variance (ANOVA) followed by post hoc Tukey test. **P < 0.01, compared with control.
Vitiligo is an acquired depigmentary skin disorder, which is not only a disease of skin melanocytes. Recent research in vitiligo suggests that skin immune innate system participates in the pathogenetic process and many triggers precede adaptive immune responses targeting melanocytes [46]. Furthermore, various cytokines released from keratinocytes chronic involve in the immune responses of the epidermal layer. It has been reported that the mRNA levels of IL-1β in vitiligo patients were found to be significantly higher than controls suggesting its possible role in inflammatory pathogenesis of vitiligo [47]. It has been reported that in comparison with controls, tissue level of IL-22 in lesional skin were significantly higher, being further augmented in perilesional skin. IL-22 increased IL-1β level via ROS production which led to NLRP3 activation in HaCaT cells (Figure 3). These results indicate that IL-22 regulates IL-1β production via NLRP3-caspase-1 activation (Figure 2). IL-22 also increased the production of anti-microbial peptides and chemokines from HaCaT cells (Figure 1). We demonstrated that IL-22 played little parts in cell proliferation and melanogenesis of melanocytes (Figure 5), and were identical with those reported in literature that melanocytes showed no coexpression of IL-22RA1 and IL-10R2 [48]. We treated HaCa T keratinocytes with IL-22 (20 ng/ml) for 12h and 24h, and then collected the conditional medium (CM-

![Figure 8: Conditional medium induces NHEM apoptosis.](image)

**Figure 8: Conditional medium induces NHEM apoptosis.** (A) Apoptosis of different groups of NHME were measured using flow cytometry with double staining of Annexin V and PI. (B) Western blot of cytosol fraction of melanocytes demonstrated increase of Bax, cleaved-caspase 3 and cytochrome c protein levels and decrease of Bcl-2. Results were normalized against β-actin expression. (C) The band densities of interest proteins were measured by Tanon 5200 Multi analysis software program, and then the Bcl-2/Bax ratio were got. Results shown are means ± SD and are representative of 3 independent experiments. Data were analyzed by one-way analysis of variance (ANOVA) followed by post hoc Tukey test. *P < 0.01, compared with control.
0, CM-12 and CM-24). CM-12 and CM-24 suppressed expressions of the melanin synthesis related proteins at different levels, including Mitf, GPNMB, Tyr, TRP-1, DCT, OA1, and Pmel 17 (Figure 6). Considering the involvement of the keratinocyte receptor PAR-2 in melanosome transfer, we examined the effect of IL-22 on PAR-2 expression of keratinocytes. As shown in Figure 4A, treatment of IL-22 for 24 h decreased PAR-2 expression. We studied the response of keratinocytes to treatment with IL-22 (20 ng/mL) over time ranging from 5 min to 60 min and observed rapid phosphorylation of Stat3 beginning at 20 min after IL-22 treatment. These results

Figure 9: A proposed scheme shows the indirect effect of IL-22 on melanocytes by keratinocytes participation.
demonstrated that keratinocytes responded directly to IL-22 stimulation by activating the JAK/Stat-3 signaling pathway. IL-22 also engaged the PI3K/Akt pathway, as the phosphorylation of Akt with peak activation at 60 min following IL-22 treatment (Figure 4B). The effect of IL-22 on Akt was partially suppressed by pre-treatment of keratinocytes with the PI3K inhibitor, LY294002 (Figure 4C). The inhibition of IL-22 on PAR-2 expression can be interfered by Stat-3 inhibitor Stat3, but not LY294002 (Figure 4C and 4D). We speculated that IL-22 modulated PAR-2 activity to decrease melanosome transfer and affected pigmentation. This effect further illustrated the suppression on melanosomes mature and trafficking. As shown in Figure 7, CM-12 and CM-24 inhibited the cells migration. The effect of conditional medium on NHEM apoptosis was the next focus of our study. As shown in Figure 8A, the percentage of apoptotic cells increased significantly in CM-12 and CM-24 group by flow cytometry analysis. CM-12 and CM-24 induced the expressions of Bax, cleaved-caspase 3 and cytochrome c, at the same time reduced Bcl-2 (Figure 8B), hence, the Bcl-2/Bax ratio was significantly decreased (Figure 8C).

In conclusion, our data demonstrated that IL-22 could indirectly regulate melanocytes functions by keratinocytes participation and mediation (Figure 9). The components released by keratinocytes contained in the conditional medium and their action mode, maybe individual or synergistic, need intense researches. This is another evidence of cell-cell connections to regulate the body’s function.

MATERIALS AND METHODS

Reagents

Recombinant human interleukin-22 was from R&D (USA). Dimethylsulfoxide (DMSO), L-3,4-dihydroxyphenylalanine (L-DOPA), melamin, LY294002, β-actin primary antibody and horseradish peroxidase-conjugated secondary antibodies were purchased from Sigma-Aldrich (USA). Stat3 was from TOCRIS (USA). Medium 254 and Human Melanocyte Growth Supplement (HMGS) were from Invitrogen (USA). GAPDH primary antibody was from Proteintech (USA). Enhanced BCA protein assay Kit, phenylmethylsulfonyl fluoride (PMSF), cell lysis buffer for Western and IP, nuclear and cytoplasmic protein extraction Kit, LDH cytotoxicity assay Kit were from Beyotime Institute of Biotechnology (China).

Cell culture

The studies on human material were approved by local ethic committee. Normal human foreskin-derived epidermal melanocytes (NHEM) were derived from young male adult foreskins (ethnic Han/aged 18 to 22 years) obtained at circumcision following standard protocols [49]. Briefly, foreskins were cut into strips and digested with 0.25% trypsin at 4°C for 20 h. Epidermis was separated from dermis. The NHEM suspension was filtered and cells were washed twice at 1,500 rpm for 5 min prior to resuspension in Medium 254 (containing the HMGS). NHEM were grown in a humidified atmosphere with 5% CO₂ at 37°C.

HaCaT human keratinocytes cell was obtained from CAS (Chinese Academy of Sciences, China). The cells were grown in Medium 245 supplemented with 10% fetal bovine serum (GIBCO, USA), 100 U/ml penicillin and 100 mg/ml streptomycin (GIBCO, USA) in a humidified atmosphere with 5% CO₂ at 37°C.

Conditional medium collection

HaCaT cells were grown in Medium 245 supplemented with 10% fetal bovine serum in 60 mm dish. After growing adhering to the wall, HaCaT cells were cultured sequentially to about 80% confluent state. Then cells were rinsed twice with PBS and divided into three groups. Cells of group one were cultured in vehicle treatment for 24 hours; cells of group two were firstly cultured in serum-free Medium 245 for 12 hours and then IL-22 (20 ng/ml) was added, for another 12 hour; cells of group three were cultured in 3 ml of serum-free Medium 245 containing IL-22 (20 ng/ml) for 24 h. The different time-point cultural treated supernatant was collected and centrifuged at 1,500 rpm for 5 min. Aspirating 2 ml of the liquid supernatant and storing in 4°C (we used it immediately after collection, otherwise, it need to be frozen and stored at −70°C until use). According to the different treated time-point, the conditional medium was termed CM-0 (from group one), CM-12 (from group two) and CM-24 (from group three).

Immunofluorescence

NHMC were grown on coverslips that had previously placed in 24-well plates. After the cells grew adhering to the wall, they were fixed in 4% paraformaldehyde PBS for 30 min at RT and permeabilized with 0.03% Triton X-100 in PBS for 5 min. The cells were rinsed with 0.1% TBST, blocked for 30 min, and incubated for 2 h at room temperature with anti-Tyr and anti-Mitf primary antibodies. Cells were rinsed three times with TBST and incubated for 60 min at room temperature with FITC-conjugated and TRITC-conjugated secondary antibodies. After further rinsing, the coverslips were sealed and the nuclei were stained with DAPI. Images were captured using a CCD camera (Olympus CKX41, Tokyo, Japan).

Tyrosinase activity and melanin contents assay

Tyrosinase activity, as the dopa oxidase here, was measured by the rate of L-DOPA oxidation as reported [50]. NHEM were treated with IL-10 for 48 h, washed with ice-
cold PBS, lysed by incubation in cell lysis buffer [1 mM PMSF] at 4°C for 20 min, and then lysates were centrifuged at 14,000 rpm for 10 min to obtain the supernatant for activity assay and centrifugation for melanin contents assay. Protein concentrations were determined by BCA kit with bovine serum albumin (BSA) as a standard. 100 μL of supernatant containing the same 10 μg total proteins was added to each well in 96-well plate, and then mixed with 100 μL 0.1% L-DOPA in 0.1 M PBS (pH 6.8) (M/V). After incubation at 37°C for 0.5 h, the dopachrome was monitored by measuring the absorbance at 475 nm.

Total melanin in the cell pellet was dissolved in 100 μL of 1 N NaOH/10% DMSO for 1 h at 80°C, and solubilized melanin was measured at 405 nm. Melanin content was calculated as a percent of the control.

Real time–PCR

The primers employed in real-time PCR were as follows: 5’-TGAGATGGCCCTCAGGTGTA-3’ and 5’-CGGAGGCAGAATAGAGAC-3’ for human b-defensin (BD1); 5’-GGGGCTCTTTGTACATCAGT-3’ and 5’-TGGTACAGATTCCGCAA-3’ for human LL37; 5’-GCTATGACATTTCCACAAACTAC-3’, and reverse, 5’-TGTTATCGCTGGCTATGTCTCCC-3’ for human S100 A7; 5’-GCCTTCTACAGGATGACCT-3’ and 5’-TTTTGTGGCTTCTCTCATGC-3’ for human S100A8; 5’-GCCGTCTACAGGGATGACCT-3’ and 5’-TTTGTGGCTTTCTTCATGGC-3’ for human CXCL1; 5’-GCCGTCTACAGGGATGACCT-3’ and 5’-TTTGTGGCTTTCTTCATGGC-3’ for human CXCL5; 5’-GCCGTCTACAGGGATGACCT-3’ and 5’-TTTGTGGCTTTCTTCATGGC-3’ for human CXCL8; 5’-TGGCATTCAGTCCCTCTATGG-3’, 5’-AGGACAAAGCAGGATCACAGTT-3’ for human matrix metalloproteinase 3 (MMP-3).

Western blot analysis

The protein suspension was obtained as the method mentioned above. Western blot was performed as described previously [51]. The primary antibodies used were p-Stat-3 (CST 9145), MITF (BS1550), TYR (C-19) (SC7833), TRP-1 (SC10443), DCT (ab74073), GPNMB (ab125898), PAR-2 (CST 6976), Rab7 (ab137029), Rab27a (ab55667), Rab17 (ab118998), NLRP3 (ab16097), caspase-1 (CST 2225), IκBα (CST 2022), IL-22R (ab5984), cytochrome c (CST 11940), Bel-2 (CST2870), Bel-2 (CST 2764), Bax (CST 2772), cleaved-caspase 3 (CST 9661), GAPDH (60004) and β-actin (CST 3700). Proteins were visualized using an enhanced chemiluminescence detection system. Densitometric analysis was again performed by using the Tanon 5200 Multi to scan the signals. Western blot assay results reported here are representative of at least 3 experiments.

Reactive oxygen species measurement

ROS generation was detected by 2’,7’-dichlorodihydrofluorescein (D6883, Sigma), which can be oxidized to fluorescent dichlorodihydrofluorescein due to ROS. N-acetyl-L-cysteine (NAC, 3 mM, A9165, Sigma) was applied for 4 h after stimulation with IL-22 to inhibit ROS generation. In a separate experiment, HaCaT cells were treated with 20 ng/ml of IL-22 in a time dependent manner, for 6, 12, 18 and 24 h. Cells were incubated with 10 μM 2’,7’-dichlorodihydrofluorescein for 30 min, then the cell lysis buffer was added. After 5 min, 150 μL of the mixture was used for the fluorescence emission at 485 nm. SN50 is a nuclear factor-kappa B (NF-κB) inhibitor (10 μM, SML1471, Sigma). HaCaT cells were treated with 20 ng/ml of IL-22 for 24 h, with or without pretreatment with SN50.

IL-1β content determination

The culture supernatants of HaCaT cells were collected and the levels of secreted IL-1β were determined using an ELISA kit in accordance with the manufacturer’s recommended protocols (557955; BD Bioscience). The detection limit of IL-1β was 4 pg/ml.

NF-κB luciferase assay

HaCaT cells (2×10^5) were plated in 24-well tissue culture plates and transfectioned with 2 μg of NF-κB-dependent luciferase reporter plasmid (Stratagene, USA). The cells were treated with IL-22 for 24 h at 37°C and then lysed in 100 μl of passive lysis buffer (Promega). 40 μl of cell lysates were assayed for both firefly and Renilla luciferase activities using the Dual-Luciferase Reporter Assay System (Promega) in accordance with the manufacturer’s instructions, and the relative light units (RLU) were measured on a luminometer. RLU from firefly luciferase was normalized for transfection efficiency to the Renilla luciferase RLU in each lysate (normalized RLU = RLUfirefly luciferase/RLU Renilla luciferase).

Blocking of IL-22 receptor signaling

The cells were pretreated with either rabbit anti-human IL-22R antibody (AF2770; R&D systems) or with the isotype control (AB-108-C; R&D systems) at a concentration of 50 μg/ml for 2 h to block IL-22R in HaCaT cells.

siRNA silencing of NLRP3 expression

To reduce endogenous NLRP3 expression, siRNA oligonucleotides (80 pmol, sc-45469; Santa Cruz) was used according to the manufacturer’s instructions. The cells were transfected with either NLRP3 siRNA oligonucleotide or a non-targeted control siRNA
oligonucleotide and incubated for 6h at 37°C under standard culture conditions. Then the cells were cultured with fresh culture media for further 14 h. In the next moment, the cells were treated with 20 ng/ml of IL-22 for another 24h. Thereafter, the cell lysates were harvested for immunoblotting, the supernatants for Elisa assay.

### Flow cytometry analysis

NHMC (2 × 10⁵/well) grown were in 6-well plate were treated according to above description. After the treatment, NHMC were harvested and analyzed for cell apoptosis by Annexin-V and propidium iodide (PI) staining, using FITC Annexin-V apoptosis detection kit (Life technology) according to the manufacturer’s instructions with a flow cytometry (BD, USA).

### Transwell migration assay

The bottom chambers of Transwell were set as five different groups (Ctrl, IL-22, CM-0, CM-12 and CM-24 group), and the top chambers were seeded inactivated 5 × 10⁴ cells per well NHEM in 200 μl Medium 254 (containing the HMGS). After 24 h of migration, the cells on the top surface of the membrane (non-migrated cells) were scraped with a cotton swab and the cells spreading on the bottom sides of the membrane (migrated cells) were fixed with cold 4% paraformaldehyde for 30 min. After that, those migrated cells were stained with 0.1% hexamethyl pararosaniline. Images were taken using Olympus inverted microscope and migrated cells were quantified by manual counting.

### Statistical analysis

All data were expressed as means ± SEM. Statistical analysis was performed with one-way ANOVA followed by Tukey’s post hoc test for multiple comparisons tests. Significant differences were accepted when \( P < 0.05 \).

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### CONFLICTS OF INTEREST

The authors declare no conflicts of interest.

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