Clinical efficacy and IL-17 targeting mechanism of *Indigo naturalis* as a topical agent in moderate psoriasis

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**Abstract**

**Background:** *Indigo naturalis* is a Traditional Chinese Medicine (TCM) ingredient long-recognized as a therapy for several inflammatory conditions, including psoriasis. However, its mechanism is unknown due to lack of knowledge about the responsible chemical entity. We took a different approach to this challenge by investigating the molecular profile of *Indigo naturalis* treatment and impacted pathways.

**Methods:** A randomized, double-blind, placebo-controlled clinical study was conducted using *Indigo naturalis* as topical monotherapy to treat moderate plaque psoriasis in a Chinese cohort (*n* = 24). Patients were treated with *Indigo naturalis* ointment (*n* = 16) or matched placebo (*n* = 8) twice daily for 8 weeks, with 1 week of follow-up.

**Results:** At week 8, significant improvements in Psoriasis Area and Severity Index (PASI) scores from baseline were observed in *Indigo naturalis*-treated patients (56.3% had 75% improvement [PASI 75] response) compared with placebo (0.0%). A gene expression signature of moderate psoriasis was established from baseline skin biopsies, which included the up-regulation of the interleukin (IL)-17 pathway as a key component; *Indigo naturalis* treatment resulted in most of these signature genes returning toward normal, including down-regulation of the IL-17 pathway. Using an in vitro keratinocyte assay, an IL-17-inhibitory effect was observed for tryptanthrin, a component of *Indigo naturalis*.

**Conclusions:** This study demonstrated the clinical efficacy of *Indigo naturalis* in moderate psoriasis, and exemplified a novel experimental medicine approach to understand TCM targeting mechanisms.

**Trial registration:** NCT01901705.

**Keywords:** *Indigo naturalis*, Gene expression, Mechanism of action, Psoriasis

**Background**

Psoriasis is a chronic inflammatory skin disease afflicting 1–3% of the world’s population [1] and 0.12–0.47% of the Chinese population [2–4]. Psoriasis is characterized by hyperproliferation of keratinocytes and dermal infiltration by activated T cells, neutrophils, and dendritic cells. The infiltrated psoriatic lesions have significantly increased levels of the CD4 T-cell subsets T helper (Th)1 and Th17 [5]. Clinical studies have shown that neutralizing tumor necrosis factor-alpha (TNF-α), interleukin (IL)-12/23, or IL-17A is efficacious, demonstrating the central role of Th1/Th17 cells in psoriasis pathogenesis [6–12]. Several classes of therapy are available to treat patients with psoriasis including topical agents (e.g., corticosteroids, vitamin D₃, salicylic acid, retinoids) and systemic agents (e.g., methotrexate, actretin, cyclosporine, biologics). Topicals are regularly used as first-line options to treat mild-to-moderate psoriasis; however, long-term use of topical agents, such as corticosteroids, has been associated with adverse events such as atrophy and striae formation [13]. Despite the availability of these treatments, topical Traditional Chinese Medicines (TCM), with *Indigo naturalis* as a key
component, are commonly used as alternate therapies, especially in the Asia Pacific region [14–16].

*Indigo naturalis*, a Chinese herb known as Qing Dai, is a dried pigment prepared from several plant species including *Baphicacanthus cusia*. Recently, clinical studies have demonstrated that *Indigo naturalis* used as topical monotherapy is efficacious in treating patients with mild-to-moderate psoriasis [17–22], although the validity of these studies is challenged by intra-patient treatment comparison designs [23]. One component of *Indigo naturalis*, indirubin, was reported to inhibit cyclin-dependent kinase [24, 25] and signal transducer and activator of transcription-3 (STAT3) activities [26], and keratinocyte proliferation in vitro [17, 20, 27]. Another compound, tryptanthrin, was reported to inhibit interferon-γ production by lymphocytes from Peyer’s Patches [28], and nitric oxide and prostaglandin E2 synthesis by murine macrophages [29]. Moreover, *Indigo naturalis* extract has been shown to inhibit oxygen generation and elastase release in formyl-methionyl-leucyl-phenylalanine-induced human neutrophils in vitro [21]. Still, as with most TCM, the therapeutically-active chemical entities of *Indigo naturalis* are poorly defined. Other plant extracts have also been investigated as topical agents for the treatment of psoriasis in controlled clinical trials [30], including a topical cream with *mahonia* extracts (*Mahonia aquifolium*), in which the potential active chemical ingredient was speculated to have an anti-inflammatory mechanism [31].

We report the first randomized, double-blind, placebo-controlled clinical trial to evaluate the pharmacological effect of *Indigo naturalis* as a single topical agent (topical ointment) in Chinese patients with moderate plaque-type psoriasis. Additionally, global gene expression analysis of affected skin was conducted to establish a molecular signature of moderate psoriasis (in our Chinese cohort and in a predominantly White cohort [32]), and used to assess the impact of *Indigo naturalis* treatment. Finally, major chemical components of *Indigo naturalis* were evaluated for anti-IL-17-induced cytokine release in cultured keratinocytes.

**Methods**

**Clinical study**

This randomized, double-blinded, placebo-controlled study was registered with the Institutional Review Board of China Medical University Hospital, Taiwan on December 13, 2012. All patients provided written informed consent prior to enrollment. Patients were eligible to be included in the study if at baseline they were 20–65 years, had a diagnosis of plaque-type psoriasis for ≥6 months, a Physician’s Global Assessment (PGA) score of 2–3, <20% total body surface area (BSA) involvement, and a target plaque of ≥4 cm².

Patients were excluded if they had non-plaque psoriasis, rebound/flare of chronic psoriasis, history of psoriatic arthritis, current drug-induced psoriasis, were pregnant/nursing/planning pregnancy (men and women), used biologics within 3 months or 5 times the half-life, received phototherapy/systemic treatment within 4 weeks, topicals within 2 weeks, any systemic immunosuppressants within 4 weeks, lithium/antimalarial/intramuscular gold within 4 weeks, tested positive for HIV/hepatitis B/C, had a history of alcohol/drug abuse, clinically-significant laboratory abnormality, sensitivity to Chinese herbs/olive oil/microcrystalline wax/Vaseline, had current signs/symptoms of severe, progressive, or uncontrolled medical conditions, or were participating concurrently in an investigational study. Patients were asked to maintain a daily diary to document treatment compliance and adverse events.

**Study agents**

*Indigo naturalis* (Qing Dai) was extracted from the aerial part of *Baphicacanthus cusia* and stored as dried powder. The *Indigo naturalis* ointment was composed of a 1:10 mixture of *Indigo naturalis* powder and a vehicle consisting of Vaseline: microcrystalline wax: olive oil (5:6:9 ratio) (Fig. 1c). The placebo was a mixture of blue dye powder (54.8% Indigo carmine aluminum lake [Blue #2] and 45.2% Allura Red AC aluminum lake [Red #40] powders), Vaseline, microcrystalline wax, and olive oil. Across three batches of extracts, consistent indigo (2.83%) and indirubin (0.24%) concentration levels were maintained. To maintain the blind, both the *Indigo naturalis* and the placebo-control agents had a blue color (Fig. 1c). The blue dye of *Indigo naturalis* is not considered an active ingredient. Both study agents were manufactured at the same Good Manufacturing Practice facility (Sheng Chang Pharmaceutical Co., Ltd., Taoyuan, Taiwan). High-performance liquid chromatography was used for quality control of the final product.

**Sample size**

Sample size was calculated using the hypothesis that a decrease in Psoriasis Area and Severity Index (PASI) scores at week 8 would be observed following *Indigo naturalis* treatment. It has been reported that a sample size of 7 placebo patients and 12 *Indigo naturalis* patients was needed to achieve a statistical power of 90% at 0.05 level of significance [19]. Assuming a patient follow-up rate of 0.9, it was determined that at least 8 placebo patients and 16 *Indigo naturalis* patients would be needed to reach statistical significance.

**Study design**

A total of 24 patients with moderate psoriasis were randomized 1:2 at week 0 to placebo (*n* = 8) or *Indigo naturalis* (*n* = 16) (Fig. 1a). Prior to week 8, one patient...
(placebo) discontinued due to gout. Patients were given and instructed to use the topical ointment twice daily for 8 weeks or until complete skin clearance was achieved, whichever occurred first. The amount of ointment used was assessed by estimating the surface area of each psoriatic lesion. To calculate the number of fingertip units (FTUs) to be used, 1 FTU = 0.5 g = 100 cm$^2$ of surface area. In the 4-week screening period prior to treatment, whole blood, lesional, and nonlesional biopsy samples were collected. At week 0, patients received
their first treatment. Subsequent visits occurred at weeks 1, 2, 4, 6, 8, and 9. Final whole blood and lesional samples were collected at week 8 (Fig. 1b).

**Study assessments**

Safety and tolerability were evaluated through week 9. Safety evaluations included adverse event assessments, physical exams, electrocardiogram measurements, and hematology and blood chemistry analyses. Tolerability was evaluated through week 9 using: PASI to measure the severity of psoriasis ranging from a score of 0 (no psoriasis) to 72 (very severe); BSA involvement with psoriasis; PGA scores ranging from 0 to 5: cleared (0), minimal (1), mild (2), moderate (3), marked (4), and severe (5); and Overall Target Plaque Severity Scores (OTPSS) that measures plaque severity ranging from a score of 0 (no evidence of disease) to 8 (very severe).

**Skin biopsy and gene expression microarray**

Two adjacent 4-mm punch biopsy specimens of lesional and nonlesional skin were obtained from each patient at week 0. A repeat biopsy was taken from the same lesional area 8 weeks after treatment with *Indigo naturalis* or placebo. Ribonucleic acid (RNA) was amplified using the NuGen Ovation® RNA Amplification System (NuGen, San Carlos, CA). The fragmented cDNA was hybridized to Affymetrix HT HG-U133 + PM Array (Affymetrix, Santa Clara, CA).

Microarray data were analyzed using ArrayStudio (Omicsoft, Cary, NC). Overall, 68 samples passed quality control analysis including 23 nonlesional samples and 24 lesional samples at baseline, and 21 lesional samples at week 8. Group comparisons were assessed using a general linear model (fold-change cutoff = 1.5) followed by multiple test correction using false discovery rate methodology (cut-off = 0.05). Pathway level analysis of differentially expressed genes (DEG) was conducted using enrichment analysis in Ingenuity Pathway Analysis (http://www.ingenuity.com).

**Inhibition of IL-17-induced cytokine release in keratinocytes**

Normal human adult keratinocytes (Lonza) were seeded into 48-well culture plates and cultured to passage three with an approximate density of 70–80%. Keratinocytes were stimulated with IL-17A (100 ng/mL) (R&D Systems, Inc., Minneapolis, MN) and treated with varying concentrations of isatin, tryptanthrin, and indirubin (Sigma-Aldrich, St. Louis, MO), and a control compound, PD 0325901 [33]. PD 0325901 is a mitogen-activated protein kinase kinase (MEK) inhibitor; a similar compound has been demonstrated to inhibit IL-17-induced cytokine release [34]. Cell viability was confirmed using the PrestoBlue® assay (Thermo Fisher Scientific, Waltham, MA). Tryptanthrin showed no impact on cell viability at all testing concentrations. Tissue culture supernatants were analyzed for pro-inflammatory cytokines IL-6 and IL-8 using MesoScale Discovery V-plex assay (MesoScale Discovery, Rockville, MD). Data were analyzed using GraphPad Prism v5.0 (http://www.graphpad.com).

**Statistical analysis**

Descriptive statistics such as mean, median, standard deviation, and proportion were used. No imputation of missing observations was performed. Safety was assessed by summarizing the incidence and type of adverse events through week 9. Bivariate statistical analysis such as the Chi-square test or Fisher’s exact test for categorical variables and two-sample t test for continuous variables were employed to evaluate the differences in baseline demographic and outcomes variables between active treatment and placebo groups.

**Results**

**Baseline demographics and disease characteristics**

Of the 31 patients screened, 24 patients were randomized 2:1 to receive *Indigo naturalis* (n = 16) or placebo (n = 8); 23 patients completed the study (Fig. 1a). All patients were Asian, and 70.8% were men (Table 1). At baseline, the extent of BSA involvement with psoriasis was approximately 9%; the mean PASI score was approximately 10; and the mean PGA score was approximately 3 (Table 1), consistent with moderate psoriasis [35].

**Efficacy**

PASI scores were measured at baseline and each visit through week 9. Baseline PASI scores were similar to placebo; however, there was a significant improvement in the placebo group at week 9 (Table 3). Baseline PGA scores were similar to placebo through week 9. Baseline OTPSS scores were similar to placebo through week 9. Baseline BSA involvement scores were similar to placebo through week 9.

| Table 1 | Baseline demographics and disease characteristics |
|---------|--------------------------------------------------|
|         | Placebo | *Indigo naturalis* | Total |
| Patients randomized, n | 8 | 16 | 24 |
| Men, n (%) | 7 (87.5) | 10 (62.5) | 17 (70.8) |
| Race, n (%) | | | |
| Asian | 8 (100) | 16 (100) | 24 (100) |
| Age, years | 40.1 ± 10.9 | 39.3 ± 10.1 | 39.6 ± 10.1 |
| Weight, kg | 74.1 ± 16.7 | 73.2 ± 16.0 | 73.5 ± 15.9 |
| Body mass index (kg/m²) | 25.2 ± 4.6 | 25.9 ± 3.9 | 25.7 ± 4.0 |
| Duration of psoriasis, years | 14.9 ± 12.1 | 13.1 ± 11.1 | 13.7 ± 11.2 |
| Age at diagnosis, years | 25.3 ± 13.3 | 26.2 ± 12.0 | 25.7 ± 11.2 |
| BSA involvement (%) | 9.6 ± 5.8 | 8.4 ± 5.8 | 8.8 ± 5.7 |
| PASI score (0–72) | 11.1 ± 3.7 | 10.1 ± 4.3 | 10.4 ± 4.0 |
| ≥ 10, n (%) | 4 (50.0) | 8 (50.0) | 12 (50.0) |
| < 10, n (%) | 4 (50.0) | 8 (50.0) | 12 (50.0) |
| PGA score | 3.3 ± 0.5 | 3.0 ± 0.5 | 3.1 ± 0.5 |
| OTPSS | 3.8 ± 0.5 | 3.6 ± 0.5 | 3.6 ± 0.5 |

Values are mean ± SD, unless otherwise indicated

BSA body surface area, OTPSS Overall Target Plaque Severity Score, PASI Psoriasis Area and Severity Index, PGA Physician’s Global Assessment, SD standard deviation
between the two groups (Table 1). As early as week 2, *Indigo naturalis*-treated patients showed a statistically significant difference in mean PASI score compared with placebo-treated patients (*p* = 0.02; Fig. 2a). At week 8, the time of the last *Indigo naturalis* treatment, the mean PASI score of *Indigo naturalis* patients was 2.64 ± 1.5 representing a significant improvement from baseline (*p* = 0.01) and a significant difference compared with placebo patients (8.30 ± 4.0; *p* = 0.01; Fig. 2a). Moreover, 56.3% of *Indigo naturalis* patients achieved at least a 75% improvement in PASI (PASI 75 response) at week 8 compared with none of the placebo patients (0/7; 0.0%; *p* = 0.02; Fig. 2b). Consistent with PASI results, baseline PGA scores were similar between the two groups; however, a significant difference in mean PGA score was observed between *Indigo naturalis* patients (1.31 ± 0.6) and placebo patients at week 8 (2.29 ± 0.8; *p* < 0.003; Fig. 2c). Similarly, a significant difference in mean was
observed between *Indigo naturalis* (1.31 ± 0.9) and placebo patients (2.86 ± 1.5) at week 8 (*p* = 0.003; Fig. 2d). An example of a skin lesion pre- and post-*Indigo naturalis* treatment is shown in Fig. 2e. At week 9 however, the mean PASI score for the *Indigo naturalis* patients was 3.74 ± 2.3, indicating diminished efficacy following termination of treatment.

**Safety**

*Indigo naturalis* was well-tolerated. With an average duration of follow-up of approximately 9 weeks, the proportion of patients with at least one adverse event was comparable between treatment groups (placebo: 50.0%; *Indigo naturalis*: 44.0%; Table 2). The most common adverse event in the *Indigo naturalis* group was pruritus (25.0%), followed by rash and nasopharyngitis (each 12.5%) and single events of pruritus, gout, allergies, and pyrexia were reported in the placebo group (all 12.5%). One placebo-treated patient discontinued treatment prior to study completion (at week 3) due to an adverse event of gout (Fig. 1a).

**Gene expression signature of moderate psoriasis**

To gain a comprehensive understanding of the molecular characteristics of moderate psoriasis represented by this cohort, a global gene expression profile was generated from skin biopsies. Baseline lesional skin samples were compared with nonlesional samples from the same patients (*n* = 24). A total of 6845 probe sets with significant modulation (3384 up and 3461 down) were identified. The corresponding 4320 DEG define a disease signature of moderate psoriasis. A hierarchical clustering analysis showed distinct expression patterns of these genes in psoriatic lesions compared with nonlesional skin (Fig. 3a). To gain functional insight to the disease signature, a pathway enrichment analysis was conducted that resulted in 193 enriched pathways (see Additional file 1). The top 15 pathways (Fig. 3b) included the “Role of IL-17A in psoriasis,” indicating that the IL-17 pathway is a key component of moderate psoriasis.

Because our study population was composed entirely of Asian patients, we performed a comparison to another psoriasis population [32] that was predominantly White. That study population [32] consisted of a similar sample size (*n* = 25 vs. *n* = 24), with slightly higher baseline PASI scores (average 14.1 vs. 10.1), and comparable baseline PGA scores (average 2.7 vs. 3.1). The same statistical approach used for comparing nonlesional samples to lesional samples at baseline resulted in 4585 DEG. The two signatures shared approximately 62% of genes that were modulated in the same direction. Pathway enrichment analyses showed that 78% of the underlying pathways were shared (see Additional files 1 and 2). “Role of IL-17A in psoriasis” ranked high in both signatures, supporting its role in both populations studied.

**Gene expression signature of *Indigo naturalis* treatment**

To investigate the pharmacological impact of *Indigo naturalis* treatment, gene expression patterns of post-treatment lesional skin were compared to pre-treatment lesional skin in the scope of the disease signature. By hierarchical clustering, 8/14 *Indigo naturalis*-treated week-8 samples were clustered together with baseline nonlesional samples, indicative of a return toward normal gene expression patterns, compared with only one placebo-treated sample (Fig. 4a). Five *Indigo naturalis* and two placebo-treated samples showed a mixed pattern between baseline lesional and nonlesional samples, indicating heterogeneity of the treatment effects.

A gene signature of *Indigo naturalis* treatment was generated by comparing week-8 lesional skin to the matched baseline samples from nine patients who achieved PASI 75 responses. The signature consisted of 5324 probe sets that corresponded to 3427 DEG (data not shown). When compared with the disease signature, 69% of genes were in common, but regulated in the opposite direction, indicating that a majority of the disease signature genes were impacted by the treatment (Fig. 4b). Enrichment analysis for this treatment signature also supports an impact on the majority of disease pathways (4 of 5 top pathways were identical to that of the disease signature, but in the opposite direction of gene regulation) (Fig. 4c; see Additional file 3). The pathway “Role of IL-17A in psoriasis” ranked even higher in this analysis than in the disease signature, and all pathway genes were reversely regulated (Fig. 4d). For example, IL-

| Table 2 Patient safety data through week 9 |
|------------------------------------------|
|                                      | Placebo | *Indigo naturalis* |
|----------------------------------------|---------|--------------------|
| Patients treated, n                   | 8       | 16                 |
| Patients with >1 adverse event, n (%) | 4 (50.0) | 7 (44.0)          |
| Average duration of follow-up (weeks) | 9.14    | 9.13               |
| Average exposure (weeks)              | 8.16    | 8.13               |
| Common adverse events, n (%)          |         |                    |
| Pruritus                               | 1 (12.5)| 4 (25.0)           |
| Rash                                   | 0 (0.0 )| 2 (12.5)           |
| Nasopharyngitis                        | 0 (0.0 )| 2 (12.5)           |
| Abdominal distension                  | 0 (0.0 )| 1 (6.3)            |
| Constipation                           | 0 (0.0 )| 1 (6.3)            |
| Cough                                  | 0 (0.0 )| 1 (6.3)            |
| Dizziness                              | 0 (0.0 )| 1 (6.3)            |
| Oropharyngeal pain                     | 0 (0.0 )| 1 (6.3)            |
| Gout                                   | 1 (12.5)| 0 (0.0)            |
| Allergies                              | 1 (12.5)| 0 (0.0)            |
| Pyrexia                                | 1 (12.5)| 0 (0.0)            |
17A expression was decreased by 4.8-fold following *Indigo naturalis* treatment (*p* = 4.6 E-11) versus a non-significant 1.7-fold decrease with placebo treatment (Fig. 5a). Together, these data provide evidence that expression of IL-17A and its pathway genes was upregulated in psoriatic lesions of our study patients and down-regulated after *Indigo naturalis* treatment.

**Inhibition of IL-17-induced cytokine release by chemical compounds in *Indigo naturalis***

To further understand the mechanism of action, the three major chemical components of *Indigo naturalis*—isatin, indirubin, and tryptanthrin—were evaluated for IL-17 inhibition activity using an IL-17-induced cytokine release assay. Cultured human keratinocytes were stimulated with IL-17A (100 ng/mL), which resulted in a 4.5- and 5.5-fold increase in secretion of the pro-inflammatory cytokines IL-6 and IL-8, respectively. This elevation in production of these cytokines was inhibited by the MEK inhibitor PD 0325901 (Fig. 5b & c), but was not affected by the presence of isatin and indirubin (data not shown). On the contrary, tryptanthrin showed moderate, but significant inhibition, reducing IL-6 secretion by nearly half at the highest concentration tested (Fig. 5b). More moderate reduction of IL-17-stimulated IL-8 secretion was also observed (Fig. 5c).

**Discussion**

*Indigo naturalis* is a TCM long recognized as a therapy for several inflammatory conditions [20, 36, 37]. Previous clinical studies have shown that monotherapy as a topical agent can be an effective treatment for mild-to-moderate psoriasis [17, 22]; however, interpretation of these studies is limited due to the lack of proper placebo controls. There also remains little understanding of the mechanism of action, although there is anecdotal evidence attributed to the anti-proliferation activity of indirubin [27]. Our study was designed to address these critical questions at both the clinical and molecular level.

Previous clinical studies have included intra-patient placebo controls: placebo was applied to one side of the body, and *Indigo naturalis* to the other side of the same patient. Therefore, it was not administered in a truly blinded manner, and there was the possibility of exposure of active agent to the contra-lateral, placebo-exposed side via potential percutaneous absorption [17, 19]. To avoid bias, our study was designed as a double-blinded, placebo-control trial in which each patient was assigned to either active agent or placebo. Furthermore, the placebo was prepared using a blue color to match the active agent in appearance and to maintain the study blinding to prevent any possible bias.
We generated, for the first time, a comprehensive profile of molecular changes associated with moderate psoriasis and with the treatment effect of Indigo naturalis in skin. As in more severe disease, deregulation of IL-17 expression was observed in the more moderate population in this study. We presented evidence that topical Indigo naturalis significantly down-regulated the IL-17 pathway in affected skin, similar to other therapies that successfully target this pathway [9, 11, 38–40]. Moreover, we showed that one of the chemical components in Indigo naturalis, tryptanthrin, possesses moderate anti-IL-17 activity. To our knowledge, this is the first report that associates Indigo naturalis, in particular one of its chemical ingredients, tryptanthrin, with anti-IL-17...
activity. Although an earlier study found that indirubin derivatives can inhibit STAT3 [41], which was shown to be a required transcription factor for Th17 differentiation [42], the link between indirubin-STAT3 and IL-17 has not been directly proven.

Although there has been a great deal of interest in understanding the pharmacologic mechanism of TCMs, limited progress has been made. One challenge is identifying the active components within TCM, as there may be numerous chemical components in a single TCM. Here, we have employed a different approach by first generating a treatment molecular signature and then identifying the specific pathways that are affected by TCM treatment, resulting in a hypothesis (IL-17 inhibition) that could be tested using purer chemical components.

We have made an effort to compare the disease signatures generated from our study population to a similar study [32] with a predominantly White population. The two signatures are mostly overlapping, indicating that the underlying driver of the disease may be the same in both psoriasis populations and suggesting that *Indigo naturalis* should be effective in broader populations of psoriasis patients.

There were a few limitations in this study, including small sample size and no systemic marker measurement. The latter limited the ability to address the reasons behind the short benefit duration of topical *Indigo naturalis* treatment. It is unknown whether this could be due to the lack of deeper skin penetration, leading to lower systemic exposure, or a relatively short half-life of the effective chemical ingredients. Future studies with a larger sample size and additional markers, such as serum tryptanthrin and IL-17 levels, would be more informative.

**Conclusion**

In summary, this experimental medicine study in patients with moderate psoriasis identified IL-17 as a key pathway that can be modulated by treatment with *Indigo naturalis*. This study exemplified a novel approach to understanding the mechanism of action of a TCM, which may be applied to understanding the therapeutic effect of other forms of TCMs.

**Additional files**

Additional file 1: Enrichment of ingenuity pathways by moderate psoriasis gene signature from our study population. (DOCX 22 kb)

Additional file 2: Enrichment of ingenuity pathways by gene signature from a predominately White psoriasis population [27]. (DOCX 18 kb)
Additional file 3: Enrichment of ingenuity pathways by gene signature of Jasplo natural treatment of moderate psoriasis from our study population. (DOCX 23 kb)

Abbreviations
BSA: Body surface area; DEG: differentially expressed genes; FTU: fingertip units; IL: interleukin; MEK: mitogen-activated protein kinase kinase; OTPSS: Overall Target Plaque Severity Score; PASI 75: 75% improvement in PASI score from baseline; PS1: Psoriasis Area and Severity Index; PGA: Physician’s Global Assessment; RNA: ribonucleic acid; STAT-3: signal transducer and activator of transcription-3; TCM: Traditional Chinese Medicine; Th: T helper; TNF-α: tumor necrosis factor-alpha

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Availability of data and materials
The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Authors’ contributions
H-MC was the Principal Investigator of the clinical study, provided and characterized patient samples, conducted the clinical study, wrote the Clinical Study Report, and was involved in the study concept development, study design, protocol synopsis and manuscript development; Y-CW was the Principal Investigator of this clinical project and involved in the study design, data analysis, and manuscript development; QW was involved in the study design, protocol synopsis development, study coordination, data analysis, and manuscript development; MS was involved in the study design, protocol synopsis development, study coordination (clinical), data analysis, and manuscript development; JW was involved in the study design concept, study design, and protocol development; DC was involved in the study design, protocol synopsis development, analysis of the data, and manuscript development; KL was involved in the study design, protocol synopsis development, and manuscript development; EW was involved in the in vitro keratinocyte cytokine release assay development and manuscript development; S-TK participated in the study design, protocol synopsis development, and data interpretation; T-CL was the chief biostatistician and was responsible for the data management and analysis, and manuscript development; FL was the Janssen-responsible officer for this study, involved in the study concept development, interpretation of data, and manuscript development; KH was involved in the mRNA isolation and manuscript development; CB was involved in the manuscript development; CCH was involved in the study concept development, study design, protocol synopsis development, data analysis, manuscript development, and communication with all co-authors. All authors read and approved the final manuscript.

Ethics approval and consent to participate
This study was approved by the Institutional Review Board of China Medical University Hospital, Taichung, Taiwan, and conducted at this institute. The protocol followed Declaration of Helsinki and detailed information can be found at the ClinicalTrials.gov website (NCT01901705). All patients provided written informed consent prior to any study-related procedures, and agreed to return for follow-up visits and provide all required biopsies.

Consent for publication
Patients provided written consent for their photographs to be used in this study and publication.

Competing interests
Q. Wang, M. Song, D. Chen, K. Li, E. Wadman, F. Leon, K. Hayden, C. Brodmerkel, and C. C. Huang are all employees of Janssen Research & Development, LLC, a subsidiary of Johnson & Johnson, and C. Wang, M. Song, D. Chen, K. Li, F. Leon, C. Brodmerkel, and C. C. Huang own stock in Johnson & Johnson.

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