**Epicutaneous Immunization with TNP-Ig and Zymosan Induces TCRαβ+ CD4+ Contrasuppressor Cells That Reverse Skin-Induced Suppression via IL-17A**

Monika Majewska-Szczechanik\(^a\)  Anna Strzepa\(^a\)  Katarzyna Marcińska\(^a\)  Li Wen\(^b\)
Marian Szczepanik\(^a\)

\(^a\)Department of Medical Biology, Faculty of Health Sciences, Jagiellonian University College of Medicine, Krakow, Poland;  
\(^b\)Department of Internal Medicine, Section of Endocrinology, Yale University School of Medicine, New Haven, Conn., USA

**Key Words**
Contact hypersensitivity · Epicutaneous immunization · Contrasuppression

**Abstract**

**Background:** Our previous work showed that epicutaneous (EC) immunization with protein antigen e.g. TNP-conjugated mouse immunoglobulin (TNP-Ig) in the form of a patch prior to hapten sensitization inhibits Th1-mediated contact hypersensitivity (CHS) in mice. We also found that suppression of CHS was mediated by TCRαβ+ CD4+ CD8+ T suppressor cells producing TGF-β. The aim of this study was to investigate the role of innate immunity in the suppression of CHS.

**Methods:** Mice were immunized by applying gauze patches containing protein antigen alone or in the presence of zymosan, and were then tested for the CHS response. Adoptive cell transfer experiments were used to study the mechanisms involved in the reversal of skin-induced suppression. The influence of EC immunization on cytokine production by lymph node cells was measured by ELISA.

**Results:** We found that EC immunization with TNP-Ig and zymosan before trinitrophenyl chloride sensitization reverses skin-induced suppression, demonstrated in vivo and in vitro. The reversal of skin-induced suppression was transferable by antigen-specific TCRαβ+ CD4+ T contrasuppressor cells. Furthermore, we showed that the contrasuppression was IL-17A-dependent and TLR2- and MyD88-independent.

**Conclusions:** Our work strongly suggests that EC immunization with protein antigen and zymosan reverses skin-induced suppression and that this approach may be a potential tool to increase the immunogenicity of weakly immunogenic antigens.

**Introduction**

The skin forms an effective barrier between the host and the environment, protects against chemical and physical aggression and prevents the invasion of pathogens. The concept that the skin is a unique and important immune organ is well accepted. It has been known for a long time that immunization via the skin leads to a strong immune response, e.g. contact hypersensitivity (CHS). While the skin is considered an organ where immune responses are easily induced [1], little attention has been given to skin-induced suppression.

Our previous work showed that epicutaneous (EC) immunization of mice with different protein antigens applied on the skin in the form of a gauze patch or cream emulsion induces a state of subsequent tolerance that inhibits both Th1- and Tc1-mediated CHS [2–5]. Further study showed that maneuvering EC immunization with protein antigens also suppresses NK cell-dependent CHS...
This was also found in an animal model of multiple sclerosis, experimental autoimmune encephalomyelitis, where EC immunization with myelin basic protein reduced disease severity and decreased disease incidence [7, 8].

Our findings are in line with the study that used a myelin basic protein-specific TCR transgenic mouse model, in which EC immunization with myelin basic protein peptides induced T suppressor cells; this protects mice from developing experimental autoimmune encephalomyelitis [9]. Furthermore, using an allogeneic skin graft experimental model, we showed that EC immunization with a protein antigen delays graft rejection [10]. It is interesting that EC immunization with the protein antigen TNP-conjugated mouse immunoglobulin (TNP-Ig) can also alleviate TNBS-induced colitis in mice [11]. Finally, study has shown that maneuvering EC immunization inhibits both the onset and progression of collagen-induced arthritis [12]. In summary, EC immunization can effectively suppress undesired immune responses including autoimmunity and graft rejection.

It has been described that EC induced suppression is antigen-nonspecific and transferable with TCRαβ+ CD4+ CD8+ T suppressor (Ts) cells [2, 8] or TCRαβ+ CD4+ CD25+ FoxP3+ T regulatory cells [4] in Th1- and Tc1-mediated immune responses, respectively. The skin constantly interacts with the environment and many commensal bacteria can be found on the skin. We found that EC induced antigen-nonspecific suppression can be reversed by crude bacterial lysates as well by purified TLR2, TLR3, TLR4 and TLR9 ligands [13]. Further experiments showed that EC immunization with TLR4 ligand LPS induces regulatory cells that reverse skin-induced suppression. These regulatory cells, designated as contrasuppressor T (Tcs) cells, belong to the population of TCRαβ+ CD4+ lymphocytes. We also showed that the observed reversal of skin-induced suppression was antigen-specific and IL-17A-dependent but TLR2- and MyD88-independent.

**Material and Methods**

**Mice**

CBA/J (H-2k) mice from the breeding unit of the Department of Medical Biology, Jagiellonian University College of Medicine were used. In some experiments, CBA/J (H-2k) and C57Bl/6 (H-2b) mice purchased from The Jackson Laboratory (Bar Harbor, Me., USA) were also used. TLR2–/–, MyD88–/– and IL-17A–/– mice on C57Bl/6 background were generated as previously reported [15] and were bred at the Yale University School of Medicine. All the mice used in this study were kept in SPF conditions and used at 6–10 weeks of age in groups of 3–9. All the experiments were conducted according to the guidelines of both the Jagiellonian University College of Medicine and Yale University.

**Reagents**

Trinitrophenyl chloride (TNP-Cl) (Chemica Alta, Edmonton, Canada), 4-ethoxymethylene-2-phenyl-2-oxazolin-5-one (OX), Evans blue, formamid, hexadecyltrimethylammonium bromide, o-dianisidine dihydrochloride and hydrogen peroxide were obtained from Sigma-Aldrich (St. Louis, Mo., USA). Protein A was from Pharmacia Fine Chemicals (Piscataway, N.J., USA) and Sepharose 4 Fast Flow was obtained from Pfizer-Pharmacia (LKB Biotechnology AB, Uppsala, Sweden). Low-tox rabbit complement (RC) was from Pel-Freeze Biologicals (Brown Deer, Wisc.). LPS from *Escherichia coli* 026:B6, zymosan from *S. cerevisiae* and curdian from *Alcaligenes faecalis* were obtained from Sigma-Aldrich. Horseradish peroxidase streptavidin was obtained from Vector Laboratories (Burlingame, Calif., USA). Mouse immunoglobulins were prepared from CBA/J mouse sera and conjugated with TNP hapten [16, 17]. A single preparation with a level of substitution of 40 TNP per immunoglobulin molecule (TNP40-Ig) was used throughout the study. Mouse immunoglobulins were conjugated with OX (OX35-Ig), as described by Askenase and Asherson [18]. In addition, a mouse Pan T Cell Isolation Kit II and CD4 MicroBeads were purchased from Miltenyi Biotec (Bergisch Gladbach, Germany).

**Monoclonal Antibodies and Hybridomas**

The following hybridomas were cultured: anti-TCRβ clone (H57–597) from Dr. R. Kubo, Cytel Inc., and anti-CD4 (clone TIB 207) and anti-CD8 (clone TIB 105.3) from the late Dr. C.A. Jane- way, Jr., Yale University, New Haven, Conn. The culture supernatants were then purified on protein A as described previously [19].

**Sensitization and Elicitation of CHS in vivo**

Mice were sensitized by topical application of 0.15 ml of 5% TNP-Cl in an acetone-ethanol mixture (1:3) to the shaved abdomen and chest. Control mice were shaved and painted with the acetone-ethanol mixture alone as a sham sensitization. Four days later, the mice were challenged on both sides of the ears with 10 μl of 0.4% TNP-Cl in an olive oil-acetone mixture (1:1). Resulting ear
thickness was measured prior to testing with a micrometer (Mito-
toyo, Tokyo, Japan) by an observer unaware of the experimental
groups, and then again at 24 h after challenge. The background ear
thickness (±20 μm at 24 h) of littermate sham-sensitized animals
that were similarly challenged was subtracted from each experi-
mental group to yield the net ear swelling expressed in μm ± SE
[2]. Ear swelling was further confirmed by the measurements of
ear weight, vascular permeability, myeloperoxidase (MPO) activ-
ity and IL-17A concentration in ear extracts.

Vascular Permeability Test
To assess very early changes in vascular permeability, TNP-Cl
immunized or naïve mice were challenged with 10 μl of 0.4% TNP-
Cl and then injected with 1% Evans blue dye (83 μg/g body weight)
24 h later. They were anesthetized and sacrificed 1 h after the Evans
blue injection. Ears were removed and a 6-mm-diameter punch of
the ear was made with a biopsy punch (Frey Products Corp., cat#
BP60). Ear punches were transferred to tubes containing 1 ml of
formamid. After 18-hour incubation at 37 °C, the samples were
centrifuged and the optical density of Evans blue in the superna-
tant was read at 565 nm against a blank containing formamid.

MPO Assay
Neutrophil infiltration to the inflamed ears was indirectly
quantitated using an MPO assay, as described previously [20].
Ears were removed 24 h after challenge and a 6-mm-diameter
punch of the ear was made. Biopsies were also collected from the
distal site of the CHS responses and homogenized in 0.5% hexa-
decyltrimethylammonium bromide pH = 6.0 (50 mg of tissue/
ml). The homogenates were freeze-thawed 3 times and centri-
fuged at 40,000 g. Aliquots (0.1 ml) were mixed with 2.9 ml phos-
phate buffer (pH = 6.0) containing 0.167 mg/ml o-dianisidine di-
hydrochloride and 5 × 10⁻⁴% H₂O₂ and incubated at 37 °C for 20 min. The absorbance was measured at 460 nm in 96-well, flat-
bottom plates. MPO activity was expressed in units per protein
concentration (U/mg of protein).

in vitro Measurement of IL-17A in CHS Ear Extracts
To determine the local production of IL-17A in elicited TNP-
Cl CHS, mice were immunized with 5% TNP-Cl or sham-sensi-
itized and challenged with 10 μl of 0.4% TNP-Cl 4 days later. The
ears were removed 24 h after challenge and a 6-mm-diameter
punch of the ear was made. Biopsies were collected from the distal
site of CHS ear responses. The biopsies were frozen rapidly in liq-
uid N₂ and were subsequently thawed and extracted in 300 μl cold
PBS on ice with a tissue microhomogenizer. The concentration of
IL-17A was measured by ELISA with the use of a BD OptEIA set
(BD Biosciences, San Diego, Calif., USA).

Epicutaneous Immunization with TNP-Ig and
Pathogen-Associated Molecular Patterns
EC immunization was performed by applying a gauze patch
(1 cm²) soaked with a solution containing 100 μg TNP-Ig alone or
TNP-Ig plus 100 μg of zymosan in 100 μl PBS to the shaved skin
on the mouse dorsum on day 0. The patch was secured by adhesive
tape wrapped around the midsection of the mouse. Control mice
were EC immunized with zymosan or PBS alone. The patch was
left in place from day 0 until day 4, when it was replaced by a fresh
patch. On day 7, the patches were removed and the mice were ac-
tively sensitized with TNP-Cl. An identical procedure was used for
EC immunization with OX-Ig. In some experiments, mice were
patched with 100 μg TNP-Ig plus 100 μg of curdulan or 100 μg of
curdulan alone prior to TNP-Cl sensitization.

Adaptive Cell Transfer of CHS and Cell-Mixing Assay
to Evaluate Contrasuppression (‘Transfer out’ Protocol)
Donors of CHS effector immune cells were sensitized with 5% TNP-Cl. CHS effector immune cells from auxiliary and inguinal
lymph nodes (ALN) and spleens were harvested on day 4. After
incubation for 30 min at 37 °C in RPMI 1640 medium, immune
cells were washed and injected (7 × 10⁷/mouse) i.v. into naïve syn-
geneic recipients (positive control). For cotransfer, ALN cells, iso-
lated from mice EC immunized with TNP-Ig and harvested on
day 7 [14], were used as the source of Ts cells. CHS effector im-
mune cells (7 × 10⁷) from the TNP-Cl-sensitized donors were co-
transferred with 5 × 10⁷ Ts cells i.v. into naïve recipients (suppres-
sion control). To test if EC immunization with TNP-Ig and zymo-
san induces regulatory cells that could reverse skin-induced
suppression, we ‘generated’ Tcs cells, which were harvested from
mice after EC immunization with TNP-Ig and zymosan on day 7. ALN
cells from these mice were used as Tcs cells, and 7 × 10⁷ of
the CHS effector immune cells from the TNP-Cl-sensitized donors
were mixed with 5 × 10⁷ Ts cells. The cell mix was then incubated
for 30 min at 37 °C with 5 × 10⁷ lymphoid cells from mice that were
EC immunized with TNP-Ig alone and harvested on day 7 (Ts
cells). The final mixture of 3 types of cells was transferred i.v. into
naïve recipients (contrasuppression control). Recipients were sub-
extaneously ear-challenged with TNP-Cl within 30 min of cell trans-
fer and were tested for CHS after 24 h, as described above.

Transfer of Regulatory Cells into EC Tolerized or
Contrasuppressed Mice (‘Transfer in’ Protocol)
To test whether EC-induced Ts or Tcs cells could interfere with
the active induction of contrasuppression or suppression, respec-
tively, we developed a ‘transfer in’ protocol. Recipients were i.v.
injected with 5 × 10⁷ Ts cells or Ts cells 2 h before patching with TNP-
Ig or TNP-Ig plus zymosan, respectively. The suppression control
mice were not transferred with any cells before EC immunization
with TNP-Ig. Mice that were EC immunized with TNP-Ig and zy-
osan but did not receive cell transfer were considered as contra-
suppression controls.

One week after EC immunization, all groups underwent sensi-
tization with 5% TNP-Cl. The positive-control mice (patched with
PBS) were sensitized with TNP-Cl. Four days later, all animals
were challenged and tested for CHS.

Phenotype of Contrasuppressor Cells
To determine the phenotype of skin-induced contrasuppressor
cells in vivo, 5 × 10⁷ cells isolated from mice EC immunized with
TNP-Ig plus zymosan were incubated in PBS on ice with purified
anti-TCRβ or anti-CD4 or anti-CD8 monoclonal antibodies
(mAbs; 1 μg mAb/10⁶ cells) or with PBS alone for 45 min. The cells
were then washed and incubated with a predetermined dilution of
RC for 60 min at 37 °C to lyse the mAb-bound cells. After washing,
the cells were resuspended in PBS. The cell equivalents were trans-
ferred into syngeneic recipients that were EC immunized with a
TNP-Ig patch for 1 week. After removing the patches, the mice
were sensitized with TNP-Cl. Mice patched with PBS before TNP-
Cl sensitization were used as positive controls. In the contrasup-
pression control, recipients received Tcs cells treated with RC

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alone before patching with TNP-Ig and subsequent sensitization with TNP-Cl. The suppressor control group consisted of mice EC immunized with TNP-Ig and painted with TNP-Cl. Four days after sensitization, all mice were challenged and tested for CHS. In addition, in one of the experiments, a pure population of MACS-isolated TCRαβ+ CD4+ lymphocytes was used as Tcs cells.

Reversal of Skin-Induced Suppression in vitro

CBA/J mice were first EC exposed to TNP-Ig alone, TNP-Ig plus zymosan or zymosan alone, as described above. On day ‘+7’ mice were sensitized by topical application of 5% TNP-Cl. As a positive control, we used mice that were EC treated with PBS alone prior to TNP-Cl sensitization. In some experiments, mice were patch with 100 μg TNP-Ig plus 100 μg of curdlan or 100 μg of curdlan alone prior to TNP-Cl sensitization. Four days after TNP-Cl immunization, single-cell suspensions of ALN cells were prepared under aseptic conditions and 3 × 10^6 ALN cells were incubated in U-bottom, 96-well microplates (Falcon, Oxnard, Calif., USA) in triplicate, with 3-fold decreasing dilutions of TNP-1g starting with 300 μg/ml in 200 μl of RPMI 1640 (containing 100 U/ml penicillin, 100 μg/ml streptomycin, 2 mM L-glutamine, 25 mM Hepes, 5 × 10^-5 M 2-mercaptoethanol and 10% fetal calf serum) for 48 h. The cells were incubated for an additional 18 h once 0.5 μCi/well of [3H]-thymidine had been added. Cell proliferation was measured by [3H]-thymidine incorporation [14]. Results are presented as mean c.p.m. ± SD. Background c.p.m. (cells cultured without antigen) were subtracted from all tested groups.

Cytokine (IFN-γ and IL-17A) immunoassays

Mice were EC immunized with TNP-Ig alone, TNP-Ig and zymosan or zymosan alone as described above. Mice EC treated with PBS were used as controls. ALN cells were collected on day 7 from the 4 groups of mice and cultured (3 × 10^6 cells) in 1 ml of RPMI 1640 medium supplemented with 10% FCS in the presence of 100 μg/ml TNPαIg. Cells were distributed in triplicate in flat-bottom, 24-well Falcon plates. The culture supernatants were collected after 48 h and tested for cytokine production using BD OptEIA sets [14].

Statistical Analysis

The data are shown on graphs and as mean ± SE. ANOVA, followed by the Tukey post test, was used for multiple comparisons. p < 0.05 was considered statistically significant.

Results

Skin-Induced Suppression Is Reversed by EC Application of TNP-Ig Together with Zymosan

To test the role of environmental influence on skin-induced suppression, we studied zymosan, which is a major component of a much-used yeast. We EC immunized CBA/J mice with 100 μg of TNP-Ig, TNP-Ig plus 100 μg zymosan, zymosan alone or PBS, as described in Materials and Methods. The mice were further sensitized with 5% TNP-Cl on day 7 and CHS was tested 4 days after sensitization. Figure 1 illustrates the experimental procedure. Figure 2a shows that mice exposed to TNP-Ig prior to TNP-Cl sensitization developed significantly decreased CHS (group B vs. A). However, EC exposure to TNP-Ig plus zymosan before CHS induction reversed skin-induced suppression in vivo (group C vs. B). Patching with zymosan alone prior to TNP-Cl sensitization did not significantly affect CHS (group D vs. A). To confirm the results, we measured the vascular permeability of the above 4 groups of mice in a separate set of experiments. As shown in figure 2b, vascular permeability was significantly reduced by TNP-Ig and this reduction was reversed in the presence of zymosan. The reversal of skin-induced suppression was further confirmed by 3 additional tests including the measurement of ear weight (fig. 2c), MPO activity (fig. 2d) and IL-17A production in the ear extracts (fig. 2e). Finally, the in vivo observations were fully confirmed by proliferation assay in vitro (fig. 3).

EC Immunization with Antigen and Zymosan Induces Tcs Cells

Our results showed that zymosan could reverse immune suppression induced by EC immunization with TNP-Ig. To investigate the mechanism of this suppression reversal, in particular which cell type is involved in this contrasuppression, we used ‘a transfer out’ protocol using 3 cell types: TNP-specific CHS effector cells, TNP-Ig EC immunization-derived suppressor cells and TNP-Ig with zymosan-derived contrasuppressor cells. As shown in figure 4a, CHS can be induced in naïve recipients by transferring effector cells (group A), and the cotransfer of effector cells and Tcs cells resulted in the inhibition of CHS (group B). Importantly, the transfer of all 3 cell types reversed the inhibition (group C), which suggests that zymosan promoted the Tcs cells. However, transferring the Tcs cells alone into naïve recipients that were subsequently challenged did not transfer CHS (group D). Similarly, in a nontransfer experiment, mice were EC immunized with TNP-Ig plus zymosan, and the subsequent challenge with TNP-Cl on the ears did not develop CHS (data not shown). Further, employing ‘transfer in’ protocol, we found that ‘Ts cells blocked EC induced suppression when transferred into the mice just before immunization via skin patching with TNP-Ig alone (fig. 4b; group E vs. B). Moreover, the transfer of Ts cells from donors EC immunized with TNP-Ig did not disturb the induction of contrasuppression when transferred just before EC immunization with TNP-Ig plus zymosan (fig. 4b; group D vs. B). We also showed that the induction of Tcs cells by zymosan...
Fig. 1. CBA/J mice were patched with PBS or 100 μg TNP-Ig alone or TNP-Ig plus 100 μg of zymosan or zymosan alone on days 0 and 4. On day 7, patches were removed and mice were actively sensitized with 5% TNP-Cl. Four days later, the animals were challenged on the ears with 0.4% TNP-Cl and then tested for CHS 24 h later, as described in Materials and Methods.

Fig. 2. a Reversal of skin-induced suppression in vivo. CBA/J mice were EC exposed to PBS (group A) or TNP-Ig (group B) or TNP-Ig plus zymosan (group C) or zymosan alone (group D) prior to TNP-Cl sensitization. Four days later, a CHS response was elicited and then tested 24 h later. Results are shown as mean ± SE (n = 18). *** p ≤ 0.001. b EC immunization with TNP-Ig and zymosan reverses decreased vascular permeability. CBA/J mice were patched with PBS (group A) or TNP-Ig (group B) or TNP-Ig plus zymosan (group C) or zymosan alone (group D) prior to TNP-Cl sensitization, challenge and then injection of 1% Evans blue dye. One hour later, the mice were killed and the ear punches were collected. Evans blue dye was extracted and the optical density was read at 565 nm. For details, see Materials and Methods. Results are shown as mean ± SE (n = 12). ** p ≤ 0.01, *** p ≤ 0.001. c Patching with TNP-Ig and zymosan reversed decreased ear weight when compared to EC tolerized mice. Ears were removed and a punch of the ear was made. The weight of the ear punches was evaluated with the use of an analytical scale, as described in Materials and Methods. Results are shown as mean ± SE (n = 9). * p ≤ 0.05, ** p ≤ 0.01. d EC immunization with TNP-Ig and zymosan reversed inhibited MPO activity. CBA/J mice were EC treated with PBS (group A) or TNP-Ig (group B) or TNP-Ig plus zymosan (group C) or zymosan alone (group D) prior to TNP-Cl sensitization followed by ear challenge. Ears were removed 24 h after challenge and punches were collected. Biopsies were homogenized and MPO activity was measured, as described in Materials and Methods. MPO activity was expressed in units per protein concentration (U/mg of protein). Results are shown as mean ± SE (n = 6). ** p ≤ 0.01, *** p ≤ 0.001. e Patching with TNP-Ig and zymosan reversed IL-17A production in ear extracts. CBA/J mice were EC treated with PBS (group A) or TNP-Ig (group B) or TNP-Ig and zymosan (group C) or zymosan alone (group D) prior to TNP-Cl sensitization and challenge. Ear punches were collected 24 h after challenge. Concentration of IL-17A was measured in tissue homogenates by ELISA. Results are shown as mean ± SE (n = 15). * p ≤ 0.05. n.s. = Non-significant.

(For figure see next page.)
<table>
<thead>
<tr>
<th>Group</th>
<th>EC immunization with TNP-Cl sensitization</th>
<th>% of response</th>
<th>24 h ear swelling (µm ± SE)</th>
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<tbody>
<tr>
<td>A</td>
<td>PBS +</td>
<td>100</td>
<td>***</td>
</tr>
<tr>
<td>B</td>
<td>TNP-Ig +</td>
<td>20</td>
<td>n.s.</td>
</tr>
<tr>
<td>C</td>
<td>TNP-Ig + zymosan +</td>
<td>77</td>
<td>***</td>
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<tr>
<td>D</td>
<td>zymosan +</td>
<td>85</td>
<td>n.s.</td>
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Vascular permeability-concentration of Evans blue in ears (µg/ml ± SE)

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<th>EC immunization with TNP-Cl sensitization</th>
<th>% of response</th>
<th>Ear weight (mg ± SE)</th>
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<tr>
<td>A</td>
<td>PBS +</td>
<td>100</td>
<td>***</td>
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<tr>
<td>B</td>
<td>TNP-Ig +</td>
<td>7</td>
<td>n.s.</td>
</tr>
<tr>
<td>C</td>
<td>TNP-Ig + zymosan +</td>
<td>95</td>
<td>*</td>
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<tr>
<td>D</td>
<td>zymosan +</td>
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MPO activity in ears (U MPO ± SE/1 mg protein)

<table>
<thead>
<tr>
<th>Group</th>
<th>EC immunization with TNP-Cl sensitization</th>
<th>% of response</th>
<th>IL-17A concentration in ears (pg/ml ± SE)</th>
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<tbody>
<tr>
<td>A</td>
<td>PBS +</td>
<td>100</td>
<td>*</td>
</tr>
<tr>
<td>B</td>
<td>TNP-Ig +</td>
<td>39</td>
<td>*</td>
</tr>
<tr>
<td>C</td>
<td>TNP-Ig + zymosan +</td>
<td>124</td>
<td>*</td>
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<tr>
<td>D</td>
<td>zymosan +</td>
<td>150</td>
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was antigen-dependent, as zymosan without TNP-Ig antigen could not induce Tcs cells (fig. 4c; group D vs. C). Our data strongly suggest that EC treatment with protein antigen and zymosan induces regulatory Tcs cells that counterregulate the suppression of Ts cells on CHS effector cells.

**EC Induced Tcs Cells Belong to the Population of TCRαβ+ CD4+ Lymphocytes**

To determine the phenotype of Tcs cells, we used ‘transfer in’ protocol, where Tcs cells were transferred into recipients that were then EC patched with TNP-Ig prior to TNP-Cl sensitization. To test whether nonconventional T cells contribute to Tcs cell function, we depleted conventional TCRαβ cells from ALN cells harvested from animals that were EC treated with TNP-Ig and zymosan. As shown in figure 5a, transfer of non-TCRαβ T cells abolished the capacity of the countersuppressive function (group D vs. C). To further investigate which subset of T cells is responsible for the contrasuppressive function, we transferred CD4- or CD8-depleted ALN cells from ‘pre-conditioned’ mice. Figure 5b shows that CD4+ cells are required for contrasuppression (group D vs. C). Thus, data presented in fig. 5a and 5b suggest that the Tcs cells induced via EC immunization with TNP-Ig and zymosan can belong to the population of TCRαβ+ CD4+ lymphocytes. To verify this possibility, another experiment with positively MACS-isolated TCRαβ+ CD4+ cells was performed. As shown in figure 5c, the transfer of purified TCRαβ+ CD4+ Tcs cells prior to the induction of suppression and subsequent TNP-Cl sensitization reversed skin-induced suppression (group D vs. B). In conclusion, the data presented in figure 5a-c prove that Tcs cells do indeed belong to the population of TCRαβ+ CD4+ lymphocytes.

**Tcs Cells Induced via EC Immunization Are Antigen-Specific**

Our previous study on skin-induced suppression suggested that the suppression is antigen nonspecific. The data shown above suggest that contrasuppression/regulation is antigen-dependent (fig. 4c). To test whether the contrasuppression is also antigen-specific, we performed a new set of experiments, in which we used two different antigens, TNP and OX. In line with our previous study, both TNP-Ig and OX-Ig could induce suppression of TNP-mediated CHS (fig. 6; groups B and D vs. A). However, when mice were EC immunized with OX-Ig and zymosan, this maneuver could not protect TNP-specific CHS from EC induced suppression (fig. 6; group E vs. B).

**EC Immunization with TNP-Ig and Zymosan Induces IL-17A Production**

To determine if cytokines are involved in the reversal of skin-induced suppression, we measured the level of two representative Th1 and Th17 cytokines. We found an increase of IFN-γ production by ALN cells containing Tcs cells (fig. 7a; group C). It is interesting that there was a robust production of IL-17A by ALN cells containing Tcs cells in the same culture supernatants (fig. 7b; group C). Our results suggest that IL-17A is most likely involved in the EC/zymosan-induced contrasuppression.

**Contrasuppression Induced via EC Immunization with TNP-Ig and Zymosan Is IL-17-Dependent and TLR2- and MyD88-Independent**

To confirm whether Tcs cell function is IL-17A-mediated, we performed a ‘transfer out’ experiment, using Tcs cells from IL-17A-/- B6 mice. Tcs cell function was completely abolished in the absence of IL-17A (group D vs. C; fig. 8a). It is known that zymosan is a ligand of TLR2 and
Fig. 4. a Reversal of skin-induced suppression is transferable with ALN cells – ‘transfer out’ protocol: $7 \times 10^7$ of the CHS effector cells from TNP-Cl-sensitized donors were incubated with $5 \times 10^7$ ALN cells used as a source of Ts cells isolated from mice (suppression control group B) patched with TNP-Ig. After incubation, the cell mixture was transferred i.v. into naïve recipients (suppression control group B). In addition, $7 \times 10^7$ of the CHS effector cells were incubated with $5 \times 10^7$ ALN cells (Tcs cells) from mice EC immunized with TNP-Ig plus zymosan prior to incubation with $5 \times 10^7$ lymphoid cells from mice EC immunized with TNP-Ig (Ts cells) (contrasuppression control group C). The resultant cells were transferred into naïve recipients. To determine whether ALN cells from mice EC immunized with TNP-Ig and zymosan could transfer CHS, $5 \times 10^7$ ALN cells were transferred i.v. into naïve recipient mice (group D). Recipients were subsequently ear-challenged with TNP-Cl and tested for CHS at 24 h. Results are shown as mean ± SE (n = 12). ** $p \leq 0.01$, *** $p \leq 0.001$. c EC immunization with zymosan alone did not reverse skin-induced suppression. CBA/J mice were i.v. transferred with $5 \times 10^7$ of ALN cells isolated from donors patched with TNP-Ig plus zymosan (group C) or zymosan alone (group D) and recipient mice were subsequently patched with TNP-Ig. Control groups were EC exposed to PBS (group A), TNP-Ig (group B) or TNP-Ig plus zymosan (group C). In addition, one group of mice before patching with TNP-Ig plus zymosan received $5 \times 10^7$ of EC induced Ts cells (group D) whereas another group of animals was transferred with EC induced Tcs cells before EC immunization with TNP-Ig alone (group E). After patch removal, mice in all the experimental groups were sensitized with TNP-Cl and tested for CHS. Results are shown as mean ± SE (n = 10). *** $p \leq 0.001$. ** Reversal of skin-induced suppression – ‘transfer in’ protocol. CBA/J mice were EC exposed to PBS (group A), TNP-Ig (group B) or TNP-Ig plus zymosan (group C). In addition, one group of mice before patching with TNP-Ig plus zymosan received $5 \times 10^7$ of EC induced Ts cells (group D) whereas another group of animals was transferred with EC induced Tcs cells before EC immunization with TNP-Ig alone (group E). After patch removal, mice in all the experimental groups were sensitized with TNP-Cl and tested for CHS. Results are shown as mean ± SE (n = 10). ** $p \leq 0.01$, *** $p \leq 0.001$. n.s. = Non-significant.
**Fig. 5.**

**a** TCRαβ+ lymphocytes are involved in skin-induced contrasuppression. ALN cells from animals patched with TNP-Ig plus zymosan were treated with anti-TCRβ mAb and RC (group D) or RC alone (group C). Resultant cells were i.v. injected into recipients that were EC immunized with TNP-Ig prior to CHS induction. Animals in control groups were patched with PBS (group A) or TNP-Ig (group B) prior to TNP-Cl sensitization. Four days after sensitization, mice were challenged and tested for CHS. Results are shown as mean ± SE (n=16). *p ≤ 0.05, **p ≤ 0.001.

**b** CD4+ lymphocytes are required to reverse skin-induced suppression. ALN cells from mice patched with TNP-Ig and zymosan were treated with anti-CD4 or anti-CD8 mAbs and RC (groups D and E, respectively). In contrasuppression control, ALN cells were treated with RC alone (group C). Resultant cells were i.v. injected into recipients that were EC immunized with TNP-Ig prior to CHS induction. Mice in control groups were patched with PBS (group A) or TNP-Ig (group B) prior to TNP-Cl sensitization. Four days after sensitization, mice were challenged and tested for CHS. Results are shown as mean ± SE (n=10). *p ≤ 0.05, **p ≤ 0.001.

**c** Contrasuppressor cells belong to the population of TCRαβ+ CD4+ lymphocytes. ALN cells from donors patched with TNP-Ig and zymosan were used to obtain MACS purified TCRαβ+ CD4+ cells (group D). Contrasuppression control contained whole ALN cells (group C). 5 × 10^7 ALN cells or 2.1 × 10^7 MACS-purified TCRαβ+ CD4+ cells were i.v. injected into recipients that were EC immunized with TNP-Ig prior to CHS induction. Mice in control groups were patched with PBS (group A) or TNP-Ig (group B) prior to TNP-Cl sensitization. Four days after sensitization, mice were challenged and tested for CHS. Results are shown as mean ± SE (n=14). **p ≤ 0.001. n.s. = Non-significant.
that the TLR2 signaling pathway is MyD88-dependent. To test the role of TLR2 and MyD88, we also transferred Tcs cells from TLR2–/–B6 or MyD88–/–B6 mice. To our surprise, Tcs cell induction and function are not dependent on TLR2 and MyD88 (fig. 8a; groups E and F, respectively). To further confirm our results from 'transfer out' experiments, we tested the active induction of Tcs cells in the knockout mice. As shown in figure 8b (group C vs. B), Tcs cells can be induced and fully functional in the absence of TLR2; however, they could not be induced in the absence of IL-17 (fig. 8c; group C). To test whether TLR ligands other than zymosan can induce Tcs cells in the absence of IL-17A, we used LPS (the most abundantly present in the environment and a ligand for TLR4) in the Tcs cell induction. Interestingly, LPS could induce contrasuppression in IL-17A–/– mice (fig. 8c; group D vs. C).

Our results suggest that zymosan plays an important role in the induction of Tcs cells that reverse skin-induced suppression, even though TLR signaling does not seem to contribute to the induction of Tcs cells in TLR2–/– mice, nor does it affect IL-17 production (fig. 8d, e).

**Dectin-1 Is Involved in Skin-Induced Contrasuppression**

It is known that zymosan generates innate signals, not only via TLR2, but via Dectin-1 as well. Data presented in figure 8 ruled out the involvement of TLR2 in contrasuppression induced via EC immunization with TNP-Ig and zymosan. These data suggest that receptors other than TLR2 play a role in tested contrasuppression. To verify...
this, animals were EC immunized with TNP-Ig and zymosan or TNP-Ig plus curdlan, which is recognized by Dectin-1. EC immunization with TNP-Ig and curdlan reverses skin-induced suppression similarly to TNP-Ig plus zymosan (group E vs. B and C, respectively; fig. 9a). These in vivo observations were fully confirmed by a proliferation assay in vitro (fig. 9b).

Discussion

Our previous work showed that EC immunization with a protein antigen prior to hapten sensitization could induce antigen-nonspecific suppression of CHS in mice [2–5]. We also found that this antigen-nonspecific inhibition of T cell-mediated immune response could be reversed by crude bacterial lysates [13] as well by purified TLR ligands [14, 21]. Furthermore, experiments with one representative TLR ligand, LPS from E. coli, demonstrated that the observed reversal of skin-induced suppression was antigen-specific, transferable with TCRαβ+ CD4+ Tcs cells and IL-12- and IFN-γ-dependent [14].

In this study, we aimed to investigate the mechanism of contrasuppression induced via EC immunization with the protein antigen TNP-Ig and zymosan. We showed that EC immunization with TNP-Ig together with zymosan prior to TNP-Cl sensitization indeed reversed skin-induced suppression as determined by the measurement of ear swelling. This was further confirmed by vascular permeability, ear weight, MPO activity and IL-17A concentration in ear homogenates, and by proliferation assay using ALN cells. In addition, we found that EC immunization with TNP-Ig and zymosan with subsequent ear challenge, but without previous TNP-Cl sensitization, did not lead to a CHS response. This observation was confirmed by an experiment showing that injection of ALN cells isolated from donors EC treated with TNP-Ig and zymosan did not transfer the CHS reaction. Our data suggest that EC immunization with the protein antigen and zymosan induces regulatory cells but not CHS effector cells. To test our hypothesis, we employed a 'transfer out' experiment where CHS effector cells were incubated with Tcs cells prior to exposure to Ts cells and subsequent adoptive transfer into naïve recipients. The results showed that EC immunization with TNP-Ig and zymosan indeed induced cells that protect from skin-induced suppression. Next, using a 'transfer in' protocol, we demonstrated that ALN cells from donors that were EC immunized with TNP-Ig and zymosan blocked EC induced suppression when transferred into recipients just before patching with TNP-Ig alone. Moreover, ALN cells isolated from mice patched with TNP-Ig alone did not affect the induction of contrasuppression when injected just before EC immunization with TNP-Ig and zymosan. In addition, both negative and positive selection experiments show that these EC induced contrasuppressor cells belong to the population of TCRαβ+ CD4+ lymphocytes. This is consistent with our previous finding that the Tcs cells induced via patching with TNP-Ig and LPS were TCRαβ+ CD4+ cells [14].

Zymosan is a preparation of the S. cerevisiae cell wall and contains β-glucan, mannan, mannoprotein and chitin [22]. Zymosan can be recognized by Dectin-1, TLR1, TLR2 and TLR6 [23, 24]. Studies also show that Dectin-2 and NOD2 might play a role in zymosan recognition, as their presence is required for the development of zymosan-induced arthritis [25, 26]. To test the role of TLR2 in the reversal of skin-induced suppression, we used TLR2–/– mice as a source of contrasuppressor cells. Both the transfer experiment and the active immunization model showed that TLR2 was not required for zymosan-induced suppression.

**Fig. 8.** a Mechanism of skin-induced contrasuppression (‘transfer out’ protocol). 5 × 10⁷ ALN cells isolated from control C57Bl/6 (group C) or IL-17A–/–B6 (group D) or TLR2–/–B6 (group E) or MyD88–/–B6 (group F). Mice EC treated with TNP-Ig and zymosan were incubated with 7 × 10⁶ TNP-specific CHS effector cells and then with 5 × 10⁵ EC induced Ts cells. Positive control recipients were transferred with immune cells alone (group A) whereas the suppression control group received immune cells cotransferred with EC induced Ts cells (group B). The recipients were challenged and tested for CHS. Results are shown as mean ± SE (n = 7). * p ≤ 0.05, ** p ≤ 0.01. b Reversal of skin-induced suppression was TLR2-independent (active immunization protocol). TLR2–/–B6 mice were patched with PBS or TNP-Ig or TNP-Ig plus zymosan (groups A–C), as described in Methods. Seven days later, mice were patched with TNP-Ig and LPS (groups A–D), as described in Materials and Methods. Seven days later, mice were sensitized with TNP-Cl and tested for CHS. Results are shown as mean ± SE (n = 10). ** p ≤ 0.01. c Contrasuppression induced via EC immunization with TNP-Ig and zymosan was IL-17A-dependent (active immunization protocol). IL-17A–/–B6 mice were patched with PBS or TNP-Ig or TNP-Ig plus zymosan or TNP-Ig and LPS (groups A–D), as described in Materials and Methods. Seven days later, mice were sensitized with TNP-Cl and tested for CHS. Results are shown as mean ± SE (n = 8). ** p ≤ 0.01, *** p ≤ 0.001. d, e C57Bl/6 or TLR2–/–B6 mice were patched with PBS or TNP-Ig or TNP-Ig plus zymosan (groups A–C). Seven days later, ALN were isolated and ALN cells (3 × 10⁶) were cultured in the presence of 100 µg/ml of TNP-Ig. After 48 h, supernatants were tested for IL-17A concentration. Results shown as mean ± SE (n = 4). *** p ≤ 0.001. n.s. = Non-significant. (For figure see next page.)
induced contrasuppression. This observation was confirmed in CBA/J wild-type mice, where EC immunization with TNP-Ig and TLR2 ligand Pam3CSK4 did not reverse skin-induced suppression (data not shown). Furthermore, using MyD88−/− mice, we demonstrated that TLR1 and TLR6 were not required for contrasuppression. These observations are in line with published reports that TLR2 and MyD88 play nonessential roles in zymosan-induced arthritis [26]. To test the role of Dectin-1 in skin-induced contrasuppression, mice were EC immunized with TNP-
Ig and curdlan prior to TNP-Cl sensitization. It is known that curdlan is a β-glucan, a ligand for Dectin-1. Both in vivo and in vitro experiments show that when curdlan is applied on the skin together with TNP-Ig, it can reverse skin-induced suppression, similar to the effect induced by EC immunization with zymosan and TNP-Ig. Our data are in line with Ikeda et al. [27], who demonstrate the β-glucan-dependent characteristics of zymosan.

To test if soluble factors are involved in the reversal of skin-induced suppression, we measured the concentration of proinflammatory cytokines in culture supernatants. We found that EC immunization with TNP-Ig and curdlan reversed skin-induced suppression in vitro. CBA/J mice were EC treated with PBS (group A) or TNP-Ig (group B) or TNP-Ig plus zymosan (group C) or zymosan alone (group D) or TNP-Ig plus curdlan (group E) or curdlan alone (group F) prior to TNP-Cl sensitization. Four days later, ALN cells were isolated and 3 × 10^5 ALN cells were cultured in triplicate in the presence of 3-fold dilutions of TNP-Ig for 48 h and then for an additional 18 h with [3H]-thymidine. [3H]-thymidine incorporation was determined by beta liquid scintillation counting. Results are presented as mean ± SE (n = 3). *p ≤ 0.05, **p ≤ 0.01.

**Fig. 9.** EC immunization with TNP-Ig and curdlan reversed skin-induced suppression. a Reversal of skin-induced suppression in vivo. CBA/J mice were patched with PBS (group A) or TNP-Ig (group B) or TNP-Ig plus zymosan (group C) or zymosan alone (group D) or TNP-Ig plus curdlan (group E) or curdlan alone (group F) prior to TNP-Cl sensitization. Four days later, CHS response was elicited and tested 24 h later. Results are shown as mean ± SE (n = 18). **p ≤ 0.01, ***p ≤ 0.001. b EC immunization with TNP-Ig and curdlan reversed skin-induced suppression in vitro. CBA/J mice were EC treated with PBS (group A) or TNP-Ig (group B) or TNP-Ig plus curdlan (group C) or curdlan alone (group D) prior to TNP-Cl sensitization. Four days later, ALN cells were isolated and 3 × 10^5 ALN cells were cultured in triplicate in the presence of 3-fold dilutions of TNP-Ig for 48 h and then for an additional 18 h with [3H]-thymidine. [3H]-thymidine incorporation was determined by beta liquid scintillation counting. Results are presented as mean ± SE (n = 3). *p ≤ 0.05, **p ≤ 0.01.
cytokines and also promotes Th17 responses to fungi [28, 29]. It seems that in our model of contrasuppression, Dectin-1 is involved. Alternatively, Dectin-1 and Dectin-2 may work in concert with innate receptors other than TLR2 or MyD88, e.g. NOD2 [30], to reverse skin-induced suppression.

The observation that the transfer of Tcs cells before the induction of suppression overcomes skin-induced suppression may suggest that these contrasuppressor cells play a regulatory function at a very early inductive stage of negative regulation. Moreover, the 'transfer out' experiment proved that EC induced contrasuppressor cells could work in the effector phase of CHS, as they protected CHS effector cells from negative signals released by suppressor cells. These data suggest that Tcs cells interfere with both the induction and effector phases of skin-induced suppression.

Lastly, the experiments with two different and non-cross-reactive antigens, TNP-Ig and OX-Ig, showed that contrasuppression induced via EC immunization with antigen and zymosan was truly antigen-specific. The induction of Tcs cells required treatment with both antigen and zymosan, as patching with zymosan alone did not result in the reversal of skin-induced suppression. Taken together, our results suggest that EC immunization with antigen and zymosan induces antigen-specific Tcs cells that protect from skin-induced antigen-nonspecific T cells. Our study also supports our previous finding that antigen-specific contrasuppression can be induced via EC immunization with antigen and the TLR4 ligand, LPS [14]. However, contrasuppression induced by LPS and zymosan has a different mechanism. As we showed previously, LPS-induced contrasuppression is IL-12- and IFN-γ-dependent [14] whereas zymosan-induced contrasuppression is mediated by IL-17A. This difference may reflect that LPS and zymosan activate different signaling pathways. LPS is a major ligand for TLR4, which activates both MyD88 and TRIF signaling pathways, which leads to the activation of NF-κB. In our experimental system, zymosan appears to act via Dectin-1. Dectin-1 triggers intracellular signaling via a cytoplasmic ITAM-like motif. Downstream signaling pathways induce a number of innate immune responses including recruitment of Syk, activation of NF-κB via CARD9 and activation of MAPKs and NFAT [31].

In summary, EC immunization with protein antigen TNP-Ig and zymosan induces antigen-specific TCRαβ+ CD4+ Tcs cells that reverse skin-induced suppression. This EC induced contrasuppression is IL-17A-dependent and TLR2- and MyD88-independent.

Our finding suggests that EC immunization with an antigen together with zymosan may play an important role in immunopotentiation and that such maneuvers may be beneficial to enhance the immune response to weakly immunogenic antigens.

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EC Induced Contrasuppressor Cells

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