

Imiquimod-induced psoriasis model: induction protocols, model characterization and factors adversely affecting the model

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Imiguimod-induced psoriasis is widely-employed to study disease pathogenesis and to screen drugs. While the original protocol was published more than a decade ago and has been rigorously used in research since then, a modified protocol was described recently with several advantages including milder systemic manifestations although the disease morphology is highly conserved. Being a toll-like receptor 7 and 8 agonist, IL-23/IL-17 axis predominates in imiquimod-induced psoriasis. In addition, different immunocytes were described to aggravate or supress the disease. This article aims to review the currently available protocols of imiquimod-induced psoriasis in vivo, to characterize the model as described in literature and to define the five important independent factors adversely influencing the model which researchers should pay attention to.

Keywords: animal model, B-cell, IL-17, IL-23, imiquimod-induced psoriasis, T-cell

Received: 14 July, 2022; revised: 15 May, 2023; accepted: 18 July, 2023; available on-line: 22 November, 2023

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Abbreviations: PASI, Psoriasis Area and Severity Index

INTRODUCTION

Animal models of psoriasis are categorized into four categories: the first category represents models resulting from spontaneous mutations, such as homozygous asebia (Scd1ab/Scd1ab) and flaky skin (Ttcfsn/Ttcfsn) mice where the latter model shares with human psoriasis acanthosis, parakeratosis, and corneal neutrophils infiltration. The second category represents genetically-engineered models where epidermal proteins like keratins and/or cytokines are modified to produce features resembling psoriasis. The third category includes humanized models or models generated by xenotransplantation where lesional skin biopsy or skin equivalent is transplanted to mice. A recently-described but frequently used model is the directly induced model, where imiquimod is used to produce an immunological reaction resembling the one seen in psoriasis, largely mediated by IL-23/IL-17 axis and TNF-α (Guerrero-Aspizua et al., 2020; Jean & Pouliot, 2010; Jeong & Lee, 2018).

This article aims to review the model of imiquimodinduced psoriasis, highlighting the currently published two induction protocols and characterizing the model in terms of dominant cytokines and cellular infiltrate. It also tackles the five factors that may influence disease modelization.

THE ORIGINAL AND MODIFIED PROTOCOLS OF IMIOUIMOD-INDUCED PSORIASIS

Imiquimod was first described in the mid-1990s as an immunomodulatory agent that augments the innate and adaptive immune systems. It is a toll-like receptor-7 and 8 agonist. It obtained the US Food and Drug Administration approval to treat anogenital warts, facial actinic keratoses and superficial basal cell carcinoma (Hanna *et al.*, 2016). Topical application of imiquimod may induce psoriasis (Wu & Strutton, 2004).

Van der Fits and others (van der Fits et al., 2009) were the first to employ imiquimod to modelize psoriasis in vivo. They applied 62.5 mg of imiquimod 5% cream (Aldara) daily, equivalent to 3.125 mg of active ingredient, on the shaved back and right ear of BALB/c and C57BL/6 mice for five or six consecutive days and they assessed the severity of psoriasis using modified Psoriasis Area and Severity Index (PASI). Signs of psoriasis start to appear within the first three days and severity steadily increases till the end of the experiment. Authors reported lack of difference between the two strains of mice. Imiquimod-treated skin shows the cardinal histopathological features of psoriasis such as acanthosis (Singh et al., 2019), parakeratosis and hypogranulosis. Immunohistochemical staining shows infiltration of dendrocytes, neutrophils and CD4+ cells. Imiquimod also induces IL-23 and augments IL-17A, IL-17F and IL-22 production (van der Fits et al., 2009).

Horvath and others (Horvath *et al.*, 2019) further modified the original protocol. They applied 25 mg of imiquimod 5% cream in Finn chambers on the back of C57BL/6 mice daily. This protocol results in erythema after the second application and scaling and skin thickening after the third application. The modified protocol was reported to be comparable to the original one. On histological examination, features of psoriasis such as parakeratosis, acanthosis, Munro microabscesses and dilated blood vessels in the dermal papillae were observed among both groups. Consistently, immunohistochemical examination reveals overexpression of Ki-67 in both. However, the modified protocol minimizes systemic manifestations and allows for prolonged imiquimod treatments.

Imiquimod-induced psoriasis model is largely used to study disease pathogenesis and to screen drugs. The model was employed for drug screening in more than 100 publications. In the majority of studies, the experimental drug of interest was administered concurrently on the same day of psoriasis induction. In a limited number of studies, it was started a few days before (1–21 days) or after induction. Regarding the protocol of induction, majority of studies complied with the original duration described as five or six consecutive days; few, howev-

er, applied the cream for shorter (three to four days) or longer periods (vast majority for seven days, up to 15 days). Prolonged daily application of imiquimod cream results in tachyphylaxis where Ki-67 expression diminishes in alignment with spontaneous attenuation and disappearance of erythema and scaling at three to four weeks despite continuous application of the cream (Kataoka et al., 2018).

Some studies lack a positive control; but where a positive control was used, it is a topical or systemic preparation that matches the route of administration of the experimental agent. Topical control preparations include betamethasone, calcipotriol, clobetasol, dexamethasone, dithranol, methotrexate and tacrolimus. Systemic control preparations include cyclosporine orally, dexamethasone orally and intraperitoneally, etanercept, methotrexate orally and intraperitoneally and tacrolimus.

CHARACTERIZATION OF IMIQUIMOD-INDUCED PSORIASIS MODEL

Jabeen and others (Jabeen *et al.*, 2020) characterized the model of imiquimod-induced psoriasis where they applied 62.5 mg of imiquimod 5% cream for eight days. Cutaneous concentration of imiquimod approaches 100 μg/g on day 2 and it doubles by six folds on day 8 corresponding with a pronounced worsening of redness, thickness, scaling and total modified PASI. Acanthosis is evident on day 8 compared with day 2, explaining the clinically apparent thickness. Dermal hypervascularity is also marked, explaining the progressive redness. In terms of cytokine profile, elevation of IL-1β, IL-6 and IL-17A was observed in skin and TNF-α and IL-17A in serum. Disease progression associates with elongation of spleen, enlargement of total area of lymph nodes, and loss of weight independently of food intake (Zhang *et al.*, 2020).

Macrophages and dendrocytes were investigated in the current model of psoriasis. While plasmacytoid dendrocytes are absent in imiquimod-induced lesions, the model shows a biphasic cellular behaviour. During the early phase, neutrophils infiltrate the epidermis and monocytes predominate in the dermis. Whereas in the late phase, Langerhans cells are pronounced in the epidermis and macrophages in the dermis. Depletion of Langerhans cells results in massive neutrophil infiltrate during the late phase, suggesting a potential anti-inflammatory role of Langerhans cells (Terhorst et al., 2015). On the contrary (Xiao et al., 2017) concluded that Langerhans cell depletion attenuates psoriasis and downregulates psoriasis-associated cytokine gene expression. (Lee et al., 2018) found that resident and monocyte-derived Langerhans cells secrete IL-23. Depletion of these cells inhibits IL-22 and IL-17A secretion (Lee et al., 2018), diminishes gamma-delta T-cell infiltration (Lee, 2016) and ultimately, attenuates psoriasis (Lee et al., 2018). Parallelly (Yoshiki et al., 2014) found IL-23-secreating Langerhans cells to induce IL-17A-producing gamma-delta T-cells. Depletion of Langerhans cells decreases Th-17-related cytokines and ameliorates psoriasis. In contrast, Kusuba and others (Kusuba et al., 2016) found that depletion of neutrophils early during psoriasis induction inhibits the infiltration of dermal monocytes, whereas depletion of both, neutrophils and monocytes, significantly attenuates psoriasis (Kusuba et al., 2016).

While IL-17 receptor is expressed on different cells, including T-cells and keratinocytes, its importance is cell-specific. For instance, deletion of keratinocyte's IL-17 receptor reduces neutrophil infiltration and abolishes

psoriasis; yet, this is not the case with T-cell-expressed receptor, emphasizing on keratinocytes' role in neutrophil chemoattraction (Moos *et al.*, 2019). Likewise, IL-17 abrogation inhibits imiquimod-induced psoriasis (Ha *et al.*, 2013). On the contrary (El Malki *et al.*, 2013) found that in IL-17A receptor-knockout mice, imiquimod may still induce psoriasis independently of IL-17 pathway. The C-X-C motif chemokine receptor type-2 is involved in neutrophil chemoattraction as well. It promotes neutrophil-produced leukotriene-B₄ and augments neutrophil chemotaxis and infiltration (Sumida *et al.*, 2014). Likewise, kallikrein-related peptidase-8 is elevated in psoriasis. If knocked out, the severity of imiquimod-induced psoriasis is comparable to wildtype, however, lesions lack neutrophil microabscesses (Iinuma *et al.*, 2015).

IL-1 and IL-36α chemoattract neutrophils. Both molecules mediate human generalized pustular psoriasis which is accompanied by systemic symptoms such as fever and malaise. In the current model of psoriasis, mice also display systemic symptoms such as weight loss and generalized malaise, suggesting the contribution of IL-1 and IL-36α to model development. Deficiency of IL-1 receptor-1 or IL-36α variably attenuates psoriasis; however, deficiency of both absolutely abolishes the disease (Alvarez & Jensen, 2016). IL-36 role is further verified in IL-36 receptor-knockout mice where these are resistant to imiquimod (Goldstein *et al.*, 2019).

Imiquimod-treated mice exhibit antihistamine-resistant itching that is largely driven by μ-opioid receptor located in the epidermis, the dorsal root ganglia, and the spinal cord. In alignment, naloxone, a μ-opioid antagonist successfully inhibits itching in imiquimod-treated mice (Takahashi *et al.*, 2017). Itching is also mediated by sphingosine 1-phosphate receptor-3, which if knocked out, scratching behaviour improves (Hill *et al.*, 2020). In addition, Oishi and others (Oishi *et al.*, 2019) found imiquimod treatment to associate with expansion of mastocytes and overexpression of the nerve growth factor, the neurotrophic factor neurotrophin 3 and enkephalin precursor preproenkephalin (Oishi *et al.*, 2019).

REGULATION OF IMIQUIMOD-INDUCED PSORIASIS

Imiquimod-induced psoriasis is negatively regulated by B-cells (Yanaba *et al.*, 2013), regulatory T-cells (Choi *et al.*, 2020; Oka *et al.*, 2017), matrix remodelling associated-7 (Ning *et al.*, 2018), indoleamine 2, 3-dioxygenase 2 (Elizei *et al.*, 2018; Fujii *et al.*, 2020), IFN regulatory factor-2 (Kawaguchi *et al.*, 2018), IFN regulatory factor-5 (Nakao *et al.*, 2020), dermokine β/γ (Tokuriki *et al.*, 2016), IL-10 (Jin *et al.*, 2018), IL-27 (Chen *et al.*, 2017; Shibata *et al.*, 2013), poly(ADP-ribose) polymerase-1 (Kiss *et al.*, 2020), endogenous n-3 polyunsaturated fatty acids (Qin *et al.*, 2014), L-selectin and ICAM-1 (Mitsui *et al.*, 2015).

The regulatory role of B-cells, regulatory T-cells and IL-10 is evident in different studies. In a model of CD19-/- mice, exacerbation of psoriasis is attributed to the loss of IL-10-secreting regulatory B-cell subset (Yanaba *et al.*, 2013). Likewise, depletion of regulatory T-cells disturbs the closely regulated gamma-delta T-cells, augments TNF-α and IL-17A secretion and aggravates the disease (Choi *et al.*, 2020). Neutralization of IL-10 in imiquimod-induced psoriasis promotes epidermal thickening, increases neutrophil infiltration and accentuates IL-23/IL-17 axis (Xu *et al.*, 2018). Likewise, knocking out IL-10 aggravates psoriasis macroscopically and mi-

croscopically, emphasising on its anti-inflammatory role in the disease (Jin et al., 2018).

FACTORS INFLUENCING IMIQUIMOD-INDUCED PSORIASIS

Five factors adversely modify the model of imiquimod-induced psoriasis: the brand of imiquimod 5% cream, mouse strain, mouse sex, stress and obesity.

The brand of the commercially available imiquimod 5% cream may interfere with the model. While (Singh *et al.*, 2019) claimed generic formulations of imiquimod to produce a psoriasiform inflammation that is comparable to Aldara, Luo and others (Luo *et al.*, 2016) found that in comparison with Aldara, Likejie creams mediates a milder form of psoriasis with a modified PASI of 3.25±1.56 (compared with 9.81±0.84 in Aldara), a less pronounced acanthosis with a Backer's score of 2.93±1.07 (compared with 6.47±1.50 in Aldara) and an epidermal thickness of 49.79±14.16 µm (compared with 85.62±17.55 µm in Aldara), concluding that different brands may adversely affect the successful establishment of the model (Luo *et al.*, 2016).

In terms of the employed strain of mice, although van der Fits and others (van der Fits et al., 2009) described their protocol in two different strains, (Swindell et al., 2017) reported variation in modelization across six different strains of mice using a five-day course of 62.5 mg imiquimod 5% cream (Aldara). Microarray showed gene expression of imiquimod-induced psoriasis to largely overlap with that of human psoriasis. C57BL/6 mice, in particular, show the highest consistency, in contrast to MOLF/EiJ and 129X1/Sv mice where gene expression is opposite to human psoriasis. In terms of IL-17 gene expression, C57BL/6 mice highly express IL-17A, IL-17B, IL-17C and IL-17F. D'Souza and others (D'Souza et al., 2020) examined the psoriatic changes induced by imiquimod in two different strains: BALB/c and the Swiss mice and concluded that imiquimod induces psoriatic changes macroscopically and microscopically among both strains, although these are more pronounced in the Swiss mice.

In terms of sex differences, and compared with male mice, female mice develop severe psoriasis in response to imiquimod, resulting in a greater weight loss, significant distress and unexpected early death. Inductions in females may also mandate euthanization (Alvarez & Jensen, 2016). In contrast, the influence of patient's sex on the severity of psoriasis is controversial. While female patients were found to significantly display milder psoriasis than male patients in two studies conducted in Swaziland and Sweden (Guillet *et al.*, 2022; Hagg *et al.*, 2017), this was contradicted by a third study (Goldburg *et al.*, 2022).

Wang and others (Wang *et al.*, 2020) investigated the effect of stress on imiquimod-induced psoriasis in a model of mice with emotional stress. In comparison with a control group with psoriasis kept off stress, stress was found to prolong the disease, to upregulate IL-1β, IL-17 and IL-22 gene expression and to increase IL-1β, IL-12, IL-17 and IL-22 secretion. This should further explain the role of stress in human psoriasis. For instance, stressful events were found to proceed psoriasis onset and were reported to trigger the disease in 31-88% of patients. Stress was also observed to aggravate psoriasis where daily stressors may expand the disease and worsens pruritus (Rigas *et al.*, 2019; Rousset & Halioua, 2018). This is evident in pediatrics as well, where child-

hood trauma is commoner in patients with psoriasis, and likewise, children with psoriasis score higher in anxiety scores (Wintermann *et al.*, 2022).

Obesity is known to exacerbate psoriasis in humans. This is also evident in imiquimod-induced psoriasis model where obese mice display thicker psoriatic lesions compared with non-obese subjects. Diet restriction partially improves psoriasis and cytokine profile (Hong et al., 2019; Kanemaru et al., 2015) and consistently, leptin deficiency attenuates the disease (Stjernholm et al., 2017). The relationship between human psoriasis and obesity was vigorously studied. A metanalysis found the odd ratio of obesity in psoriasis is 1.66, and it can approach 2.23 in patients with severe disease (Armstrong et al., 2012). A systematic review did also conclude that seven out of nine studies found a statistically significant association between increased psoriasis severity and increased body mass index (Fleming et al., 2015). Such an association is attributed to a shared mechanism involving inflammatory mediators and adipokines (Jensen & Skov,

CONCLUSIONS

Imiquimod-induced psoriasis serves as an acceptable model to study IL-23/IL-17 axis and to screen pharmaceutical agents in psoriasis. While the model could be induced using two protocols, the original protocol described by van der Fits and others (van der Fits *et al.*, 2009) is widely employed in different studies. To ensure consistency of results, researchers should take into account that variation in the brand of imiquimod 5% cream, strain of mice, sex of mice, exposure to stress and obesity may adversely modify the course of disease.

Declarations

Interest statement. Author declares no conflict of interest.

REFERENCES

Alvarez P, Jensen LE (2016) Imiquimod treatment causes systemic disease in mice resembling generalized pustular psoriasis in an IL-1 and IL-36 dependent manner. *Mediators Inflamm* **2016**: 6756138. htt-ps://doi.org/10.1155/2016/6756138

Armstrong AW, Harskamp CT, Armstrong EJ (2012) The association between psoriasis and obesity: a systematic review and meta-analysis of observational studies. *Nutr Diabetes* 2: e54. https://doi.org/10.1038/nutd.2012.26

Chen W, Gong Y, Zhang X, Tong Y, Wang X, Fei C, Xu H, Yu Q, Wang Y, Shi Y (2017) Decreased expression of IL-27 in moderate-to-severe psoriasis and its anti-inflammation role in imiquimod-induced psoriasis-like mouse model. *J Dermatol Sci* 85: 115–123. https://doi.org/10.1016/j.idermscj.2016.11.011

https://doi.org/10.1016/j.jdermsci.2016.11.011
Choi CW, Kim BR, Yang S, Kim Y, Kang JS, Youn SW (2020) Regulatory T cells suppress skin inflammation in the imiquimod-induced psoriasis-like mouse model. *J Dermatol Sci* **98**: 199–202. https://doi.org/10.1016/j.jdermsci.2020.04.008

D'Souza L, Badanthadka M, Salwa F (2020) Effect of animal strain on model stability to imiquimod-induced psoriasis. *Indian J Physiol Pharmacol* 2 **64**: 83–91

El Malki K, Karbach SH, Huppert J, Zayoud M, Reissig S, Schuler R, Nikolaev A, Karram K, Munzel T, Kuhlmann CR, Luhmann HJ, von Stebut E, Wortge S, Kurschus FC, Waisman A (2013) An alternative pathway of imiquimod-induced psoriasis-like skin inflammation in the absence of interleukin-17 receptor a signaling. *J Imest Dermatol* 133: 441–451. https://doi.org/10.1038/jid.2012.318

Elizei SS, Pakyari M, Ghoreishi M, Kilani R, Mahmoudi S, Ghahary A (2018) IDO-expressing fibroblasts suppress the development of imiquimod-induced psoriasis-like dermatitis. *Cell Transplant* 27: 557– 570. https://doi.org/10.1177/0963689718757482

Fleming P, Kraft J, Gulliver WP, Lynde C (2015) The relationship of obesity with the severity of psoriasis: a systematic review. *J Cutan Med Surg* 19: 450–456. https://doi.org/10.1177/1203475415586332

- Fujii K, Yamamoto Y, Mizutani Y, Saito K, Seishima M (2020) Indoleamine 2,3-dioxygenase 2 deficiency exacerbates imiquimodinduced psoriasis-like skin inflammation. Int J Mol Sci 21: 5515. https://doi.org/10.3390/ijms21155515
- Goldburg S, Chen R, Langholff W, Lafferty KP, Gooderham M, Jong EMd, Strober B (2022) Sex differences in moderate to severe psoriasis: analysis of the psoriasis longitudinal assessment and registry. J Psoriasis Psoriatis Arthritis 7: 132–139. https://doi. org/10.1177/24755303221099848
- Goldstein J, Bassoy E, Palomo J, Rodriguez E, Gabay C (2019) IL-36 signaling in keratinocytes is mandatory in imiquimod-induced psoriasis in mice. ARD 78: A1–A83. https://doi.org/10.1136/annrheumdis-2018-EWRR2019.82
- Guerrero-Aspizua S, Carretero M, Conti CJ, Del Rio M (2020) The importance of immunity in the development of reliable animal models for psoriasis and atopic dermatitis. *Immunol Cell Biol* **98**: 626–638. https://doi.org/10.1111/imcb.12365 Guillet C, Seeli C, Nina M, Maul LV, Maul JT (2022) The impact of
- gender and sex in psoriasis: What to be aware of when treating women with psoriasis. Int J Womens Dermatol 8: e010. https://doi. IW9.0000000000000000000010 org/10 1097/
- Ha H-L, Wang H, Pisitkun P, Kim J-C, Morasso M, Udey M, Siebenlist U (2013) Critical cell-type specific functions of the II-17 receptor signaling adaptor CIKS/ACT1 in imiquimod-induced psoriasis. Cytokine 63: 268
- Hagg D, Sundstrom A, Eriksson M, Schmitt-Egenolf M (2017) Severity of psoriasis differs between men and women: a study of the clinical outcome measure psoriasis area and severity index (PASI) in 5438 Swedish register patients. Am J Clin Dermatol 18: 583–590. https://doi.org/10.1007/s40257-017-0274-0
 Hanna E, Abadi R, Abbas O (2016) Imiquimod in dermatology: an
- overview. Int J Dermatol 55: 831-844. https://doi.org/10.1111/
- Hill RZ, Rifi Z, Vuong C, Bautista DM (2020) Loss of S1PR3 attenuates scratching behaviors in mice in the imiquimod model of psoriasis, but not in the MC903 model of atopic dermatitis. Itch 5: e35
- Hong SM, Kim JU, Cho GJ, Jin WJ, Park SH, Park IH, Seol JE, Jung SY, Wang HY, Kim H (2019) Obesity exacerbates imiquimod-induced psoriasis by enhancing IL-6 production and Th17 cell differentiation in C57BL/6 mice. 프로그램북(구초록집) 71: 389 (in
- Horvath S, Komlodi R, Perkecz A, Pinter E, Gyulai R, Kemeny A (2019) Methodological refinement of Aldara-induced psoriasiform dermatitis model in mice. Sci Rep 9: 3685. https://doi.org/10.1038/ s41598-019-39903-x
- Iinuma S, Kishibe M, Saito N, Igawa S, Honma M, Takahashi H, Bando Y, Yoshida S, Iizuka H, Ishida-Yamamoto A (2015) Klk8 is required for microabscess formation in a mouse imiquimod model of psoriasis. Exp Dermatol 24: 887-889. https://doi.org/10.1111/
- Jabeen M, Boisgard AS, Danoy A, El Kholti N, Salvi JP, Boulieu R, Fromy B, Verrier B, Lamrayah M (2020) Advanced characterization of imiquimod-induced psoriasis-like mouse model. Pharmaceutics 12. https://doi.org/10.3390/pharmaceutics12090789
- Jean J, Pouliot R (2010) In vivo and in vitro models of psoriasis. In Eberli D ed. *Tissue Engineering*: InTech Jensen P, Skov L (2016) Psoriasis and obesity. *Dermatology* **232**: 633–
- 639. https://doi.org/10.1159/000455840
- Jeong I, Lee HJ (2018) Psoriasis skin models as promising tools in psoriasis research. Biomed J Sci Tech Res 2. https://doi.org/10.26717/ BISTR.2018.02.000760
- Jin SP, Koh SJ, Yu DA, Kim MW, Yun HT, Lee DH, Yoon HS, Cho S, Park HS (2018) Imiquimod-applied Interleukin-10 deficient mice better reflects severe and persistent psoriasis with systemic inflammatory state. Exp Dermatol 27: 43–49. https://doi.org/10.1111/ exd.13403
- Kanemaru K, Matsuyuki A, Nakamura Y, Fukami K (2015) Obesity exacerbates imiquimod-induced psoriasis-like epidermal hyperplasia and interleukin-17 and interleukin-22 production in mice. Exp Dermatol **24**: 436–442
- Kataoka S, Yamamoto M, Ohko K, Nakajima K, Sano S (2018) Distinct kinetics of two pathologies induced in mice by topical treatment with imiquimod cream: Psoriasis-like inflammation and systemic autoimmunity. *J Dermatol Sci* **91**: 225–228. https://doi.or-g/10.1016/j.jdermsci.2018.05.001
- Kawaguchi M, Oka T, Sugaya M, Suga H, Kimura T, Morimura S, Fujita H, Sato S (2018) IRF-2 haploinsufficiency causes enhanced imiquimod-induced psoriasis-like skin inflammation. J Dermatol Sci 90: 35–45. https://doi.org/10.1016/j.jdermsci.2017.12.014
- Kiss B, Szanto M, Hegedus C, Antal D, Szodenyi A, Marton J, Mehes G, Virag L, Szegedi A, Bai P (2020) Poly(ADP-ribose) polymerase-1 depletion enhances the severity of inflammation in an imiquimodinduced model of psoriasis. Exp Dermatol 29: 79-85. https://doi. org/10.1111/exd.14061
- Lee M (2016) Functional role of epidermal Langerhans cells in imiquimod-induced psoriasis-like dermatitis model [Master's thesis]. Yonsei University

- Lee M, Kim SH, Kim TG, Park J, Lee JW, Lee MG (2018) Resident and monocyte-derived Langerhans cells are required for imiquimodinduced psoriasis-like dermatitis model. J Dermatol Sci 91: 52-59. https://doi.org/10.1016/j.jdermsci.2018.04.003
- Luo DQ, Wu HH, Zhao YK, Liu JH, Wang F (2016) Original Research: Different imiquimod creams resulting in differential effects for imiquimod-induced psoriatic mouse models. Exp Biol Med (Maywood) 241: 1733–1738. https://doi.org/10.1177/1535370216647183 Mitsui A, Tada Y, Shibata S, Kamata M, Hau C, Asahina A, Sato S
- (2015) Deficiency of both L-selectin and ICAM-1 exacerbates imiquimod-induced psoriasis-like skin inflammation through increased
- infiltration of antigen presenting cells. *Clin Immunol* **157**: 43–55. htt-ps://doi.org/10.1016/j.clim.2014.12.011

 Moos S, Mohebiany AN, Waisman A, Kurschus FC (2019) Imiquimod-induced psoriasis in mice depends on the IL-17 signaling of keratinocytes. *J Invest Dermatol* **139**: 1110–1117. https://doi. org/10.1016/j.jid.2019.01.006
- Nakao M, Miyagaki T, Sugaya M, Sato S (2020) Exacerbated imiquimod-induced psoriasis-like skin inflammation in IRF5-deficient mice. *Int J Mol Sci* **21**: 3681. https://doi.org/10.3390/ijms21103681
- Ning J, Shen Y, Wang T, Wang M, Liu W, Sun Y, Zhang F, Chen L, Wang Y (2018) Altered expression of matrix remodelling associated 7 (MXRA7) in psoriatic epidermis: Evidence for a protective role in the psoriasis imiquimod mouse model. Exp Dermatol 27: 1038-1042. https://doi.org/10.1111/exd.13687
- Kusuba N, Kitoh A, Miyachi Y, Kabashima K (2016) Role of neutrophils in the pathogenesis of imiquimod-induced psoriasis-like skin lesions. J Dermatol Sci 84 e73
- Oishi N, Iwata H, Kambe N, Kobayashi N, Fujimoto K, Sato H, Hisaka A, Ueno K, Yamaura K (2019) Expression of precipitating factors of pruritus found in humans in an imiquimod-induced psoriasis mouse model. Heliyon 5: e01981. https://doi.org/10.1016/j. helivon.2019.e01981
- Oka T, Sugaya M, Takahashi N, Takahashi T, Shibata S, Miyagaki T, Asano Y, Sato S (2017) CXCL17 attenuates imiquimod-induced psoriasis-like skin inflammation by recruiting myeloid-derived suppressor cells and regulatory T cells. J Immunol 198: 3897-3908. htts://doi.org/10.4049/jimmunol.1601607
- Qin S, Wen J, Bai XC, Chen TY, Zheng RC, Zhou GB, Ma J, Feng JY, Zhong BL, Li YM (2014) Endogenous n-3 polyunsaturated fatty acids protect against imiquimod-induced psoriasis-like inflammation via the IL-17/IL-23 axis. Mol Med Rep 9: 2097-2104. https://doi. org/10.3892/mmr.2014.2136
- Rigas HM, Bucur S, Ciurduc DM, Nita IE, Constantin MM (2019) Psychological stress and depression in psoriasis patients - a dermatologist's perspective. Maedica (Bucur) 14: 287-291. https://doi. org/10.26574/maedica.2019.14.3.287
- Rousset L, Halioua B (2018) Stress and psoriasis. Int J Dermatol 57: 1165-1172. https://doi.org/10.1111/ijd.14032
- Shibata S, Tada Y, Asano Y, Yanaba K, Sugaya M, Kadono T, Kanda N, Watanabe S, Sato S (2013) IL-27 activates Th1-mediated responses in imiquimod-induced psoriasis-like skin lesions. *J Invest Dermatol* **133**: 479–488. https://doi.org/10.1038/jid.2012.313
- Singh TP, Zhang HH, Hwang ST, Farber JM (2019) IL-23- and imiquimod-induced models of experimental psoriasis in mice. Curr Protoc Immunol 125: e71. https://doi.org/10.1002/cpim.71
 Stjernholm T, Ommen P, Langkilde A, Johansen C, Iversen L, Rosada
- C, Stenderup K (2017) Leptin deficiency in mice counteracts imiquimod (IMQ)-induced psoriasis-like skin inflammation while leptin stimulation induces inflammation in human keratinocytes. Exp Der-
- matol 26: 338–345. https://doi.org/10.1111/exd.13149
 Sumida H, Yanagida K, Kita Y, Abe J, Matsushima K, Nakamura M, Ishii S, Sato S, Shimizu T (2014) Interplay between CXCR2 and BLT1 facilitates neutrophil infiltration and resultant keratinocyte activation in a murine model of imiquimod-induced psoriasis. J Immunol 192: 4361–4369. https://doi.org/10.4049/jimmunol.1302959
- Swindell WR, Michaels KA, Sutter AJ, Diaconu D, Fritz Y, Xing X, Sarkar MK, Liang Y, Tsoi A, Gudjonsson JE, Ward NL (2017) Imiquimod has strain-dependent effects in mice and does not uniquely model human psoriasis. Genome Med 9: 24. https://doi.org/10.1186/ :13073-017-0415-3
- Takahashi N, Tominaga M, Kosaka R, Kamata Y, Umehara Y, Matsuda H, Sakaguchi A, Ogawa H, Takamori K (2017) Involvement of micro-opioid receptors and kappa-opioid receptors in itchrelated scratching behaviour of imiquimod-induced psoriasis-like dermatitis in mice. Acta Derm Venereol 97: 928-933. https://doi. org/10.2340/00015555-2704
- Terhorst D, Chelbi R, Wohn C, Malosse C, Tamoutounour S, Jorquera A, Bajenoff M, Dalod M, Malissen B, Henri S (2015) Dynamics and transcriptomics of skin dendritic cells and macrophages in an imiquimod-induced, biphasic mouse model of psoriasis. J Immunol 195:
- 4953–4961. https://doi.org/10.4049/jimmunol.1500551
 Tokuriki A, Chino T, Luong V, Oyama N, Higashi K, Saito K, Hasegawa M (2016) Dermokine β/γ deficiency enhances imiquimod-induced psoriasis-like inflammation. J Dermatol Sci 84: e89-e180

- van der Fits L, Mourits S, Voerman JS, Kant M, Boon L, Laman JD, Cornelissen F, Mus AM, Florencia E, Prens EP, Lubberts E (2009) Imiquimod-induced psoriasis-like skin inflammation in mice is mediated via the IL-23/IL-17 axis. *J Immunol* 182: 5836–5845. https://doi.org/10.4049/jimmunol.0802999
- Wang Y, Li P, Zhang L, Fu J, Di T, Li N, Meng Y, Guo J, Zhao J (2020) Stress aggravates and prolongs imiquimod-induced psoriasis-like epidermal hyperplasis and IL-1beta/IL-23p40 production. J Leukoc Biol 108: 267–281. https://doi.org/10.1002/JLB.3MA0320-363RR
- Wintermann GB, Bierling AL, Peters EMJ, Abraham S, Beissert S, Weidner K (2022) Childhood trauma and psychosocial stress affect treatment outcome in patients with psoriasis starting a new treatment episode. Front Psychiatry 13: 848708. https://doi.org/10.3389/ fpsyt.2022.848708
- Wu JK, Siller G, Strutton G (2004) Psoriasis induced by topical imiquimod. Australas J Dermatol 45: 47–50. https://doi.org/10.1111/j.1440--0960.2004.00030.x
- Xiao C, Zhu Z, Sun S, Gao J, Fu M, Liu Y, Wang G, Yao X, Li W (2017) Activation of Langerhans cells promotes the inflammation in imiquimod-induced psoriasis-like dermatitis. J Dermatol Sci 85: 170– 177. https://doi.org/10.1016/j.jdermsci.2016.12.003

- Xu X, Prens E, Florencia E, Boon L, Asmawidjaja P, Otten-Mus A-M, Lubberts E (2018) IL-10 Regulates skin thickness and scaling in imiquimod-induced psoriasis-like skin inflammation in mice. Ann Rheum Dis 77: A1–A77
- Yanaba K, Kamata M, Ishiura N, Shibata S, Asano Y, Tada Y, Sugaya M, Kadono T, Tedder TF, Sato S (2013) Regulatory B cells suppress imiquimod-induced, psoriasis-like skin inflammation. *J Leukoc Biol* **94**: 563–573. https://doi.org/10.1189/jlb.1112562
 Yoshiki R, Kabashima K, Honda T, Nakamizo S, Sawada Y, Sugita
- Yoshiki R, Kabashima K, Honda T, Nakamizo S, Sawada Y, Sugita K, Yoshioka H, Ohmori S, Malissen B, Tokura Y, Nakamura M (2014) IL-23 from Langerhans cells is required for the development of imiquimod-induced psoriasis-like dermatitis by induction of IL-17A-producing gammadelta T cells. J Invest Dermatol 134: 1912-1921. https://doi.org/10.1038/jid.2014.98
- Zhang J, Yang X, Hong Qiu, Chen W (2020) Weight loss may be unrelated to dietary intake in the imiquimod-induced plaque psoriasis mice model. *Open Life Sci.* **15**: 79–82