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BMJ Open Safety of topical corticosteroids in atopic eczema: an umbrella review

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ABSTRACT

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Dr Emma Axon; emma.axon@nottingham.ac.uk **Objective** An umbrella review summarising all safety data from systematic reviews of topical corticosteroids (TCS) in adults and children with atopic eczema.

Methods Embase, MEDLINE, PubMed, Cochrane Database of Systematic Reviews and the Centre of Evidence Based Dermatology map of eczema systematic reviews were searched until 7 November 2018 and Epistemonikos until 2 March 2021. Reviews were included if they assessed the safety of TCS in atopic eczema and searched ≥1 database using a reproducible search strategy. Review quality was assessed using version 2 of 'A MeaSurement Tool to Assess systematic Reviews' (AMSTAR 2 tool).

Results 38 systematic reviews included, 34 low/critically low quality. Treatment and follow-up were usually short (2–4 weeks).

Key findings TCS versus emollient/vehicle: No metaanalyses identified for skin-thinning. Two 2-week randomised controlled trials (RCTs) found no significant increased risk with very potent TCS (0/196 TCS vs 0/33 vehicle in children and 6/109 TCS vs 2/50 vehicle, age unknown). Biochemical adrenal suppression (cortisol) was 3.8% (95% Cl 2.4% to 5.8%) in a meta-analysis of 11 uncontrolled observational studies (any potency TCS, 522 children). Effects reversed when treatment ceased, TCS versus topical calcineurin inhibitors: Meta-analysis showed higher relative risk of skin thinning with TCS (4.86, 95% CI 1.06 to 22.28, n=4128, four RCTs, including one 5-year RCT). Eight cases in 2068 participants, 7 using potent TCS. No evidence of growth suppression. Once daily versus more frequent TCS: No meta-analyses identified. No skin-thinning in one RCT (3 weeks potent TCS, n=94) or biochemical adrenal suppression in two RCTs (up to 2 weeks very potent/moderate TCS, n=129). TCS twice/week to prevent flares ('weekend therapy') versus vehicle: No meta-analyses identified. No evidence of skin thinning in five RCTs. One RCT found biochemical adrenal suppression (2/44 children, potent TCS). Conclusions We found no evidence of harm when TCS were used intermittently 'as required' to treat flares or 'weekend therapy' to prevent flares. However, long-term safety data were limited.

PROSPERO registration number CRD42018079409.

INTRODUCTION

Atopic eczema (also known as atopic dermatitis or eczema) is an itchy inflammatory skin condition. It is most common in children

Strengths and limitations of this study

- Robust Cochrane methodology was followed and a thorough and inclusive literature search was performed to ensure this was a comprehensive overview.
- By extracting data from existing reviews, results are limited to topics for which there is an eligible systematic review.
- Safety was usually reported in less detail than effectiveness in systematic reviews limiting the available data for extraction, therefore potentially missing data included in the original papers.
- Most included reviews were rated low or critically low-quality using AMSTAR 2, and where quality of evidence assessments were reported for individual studies most indicated a high or unclear risk in at least one domain.
- Many randomised controlled trials were only short in duration (2–4 weeks) limiting our ability to assess side effects that take longer to develop such as skin thinning.

with one in five affected worldwide,^{1 2} but often persists into adulthood.³

Topical corticosteroids (TCSs) are first-line therapy for treating inflammatory eczema flares but widespread concerns regarding their safety among patients and healthcare professionals contribute to poor adherence, and subsequent worsening of disease control and quality of life.⁴⁵ Safety concerns include skin thinning and retardation of growth and development. These concerns are thought to mainly originate from what is now considered to be inappropriate use, such as using potent TCS on the face or continual long-term use.⁶ Strategies recommended to minimise exposure to TCS, and hence the risk of adverse events, include reducing frequency of application to once daily during treatment of an inflammatory episode, or TCS used for two consecutive days a week (sometimes referred to as 'weekend therapy') as a strategy to prevent flares.^{7–9} This umbrella review aims

BMJ

to assess safety (local and systemic adverse events) of TCS compared with other topical treatments, placebo or no comparator in people of any age and gender with atopic eczema, and addressed two areas of research prioritised in the James Lind Alliance priority setting partnership for atopic eczema.¹⁰

METHODS

Protocol, registration and study design

This umbrella review includes published systematic reviews of randomised controlled trials (RCTs) and/ or observational studies reporting adverse event data in people with eczema using TCS. The aim of this overview was to summarise data from existing reviews, therefore, meta-analyses and data from individual studies were extracted and presented in this overview in the format and completeness that they were presented in the original systematic reviews. The only exception was for missing p values which were calculated where appropriate. The checklist 'Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA)' was followed.¹¹

Search strategy

Embase, MEDLINE, PubMed, Cochrane Database of Systematic Reviews and Epistemonikos were searched from inception to 7 November 2018 by DJCG (information specialist), with no restrictions on language or publication date. The search strategy is in online supplemental appendix 1. The Epistemonikos search was updated on 2 March 2021, with a publication date restricted to 2018-2021. Epistemonikos is now well established as a comprehensive database of reviews that regularly searches ten major databases including the Cochrane Library, PubMed and Embase¹³ thus making the need to search these individual databases redundant. We also checked the Centre of Evidence Based Dermatology eczema map of systematic reviews,¹⁴ and searched PROSPERO up to 23 March 2021 for any relevant ongoing systematic reviews using the terms 'eczema' and 'dermatitis'.

Eligibility criteria

We included systematic reviews that presented data on the safety of TCS used to treat people of any age and gender with atopic eczema, had clinical outcomes, searched at least one database and provided a reproducible search strategy. Systematic reviews of any types of clinical study design were included. Multiple reviews on the same topic were included, except for 'abridged' versions of the same review where no additional data were reported. To avoid duplication of data, for each comparison, the review that included the highest number of studies on that comparison and therefore appeared the most comprehensive was taken as the primary review and other included reviews were checked for additional studies and data. Conference abstracts were excluded. Reviews that covered multiple skin conditions were only included if they reported data on atopic eczema patients separately.

Interventions and control

Our intervention of interest was any TCS of any preparation and potency used to treat atopic eczema. For RCTs, the comparisons of interest were any other TCS, the same TCS used in a different way, another topical anti-inflammatory treatment, vehicle, no treatment or a combination of any of these. Comparisons with nontopical treatments were excluded as we were interested in clinical practice decisions regarding alternatives to TCS.

Outcomes

Safety outcomes reported during the treatment and follow-up were extracted where reported in the reviews on immediate cutaneous adverse events (eg, burning sensation/stinging), other cutaneous adverse events (eg, skin thinning, telangiectasia, skin infections, folliculitis), systemic adverse events (eg, effects on endocrine system, impact on growth) and rebound symptoms/steroid withdrawal.

Selection of studies and data extraction

Records identified from the database searches were uploaded into Covidence (Veritas Health Innovation, Australia).¹⁵ Two authors (EA and JRC) independently assessed the eligibility of each record, and where unclear the full text was obtained. The number of included and excluded records along with reasons for exclusion were reported in a PRISMA flow diagram.

Two authors (EA and JRC) independently extracted all safety data presented in the included reviews along with other information such as review/participant characteristics, and funding sources. Any disagreements regarding eligibility or data extraction were resolved via discussion or input from a third reviewer (HCW or KST). Where available, we reported results separately for age, filaggrin mutation status, TCS potency, site of application of the TCS, and duration of continuous treatment.

Assessment of quality of included systematic reviews

As this was an overview of reviews, the methodological quality of the evidence was assessed at the systematic review level using version 2 of 'A MeaSurement Tool to Assess systematic Reviews' (AMSTAR 2 tool) and this was conducted in duplicate by EA and JRC.¹⁶ Reviews were considered critically low quality if there was more than one critical flaw. Data on the quality of individual studies (eg, risk of bias) and the quality of evidence (eg, Grading of Recommendations Assessment, Development and Evaluation, GRADE¹⁷) were also extracted where presented in the review, but undertaking these quality assessments for individual studies was not within the remit of this overview.

Measures of treatment effect and data synthesis

Where relevant meta-analyses were presented in the systematic review, the forest plots, relative risk (RR) and 95% CI were extracted. In the absence of any metaanalysis, adverse event data from individual studies were included in this overview based on the data presented in the published systematic review. P values were calculated using Review Manager software,¹⁸ with <0.05 indicating statistically significant results. Where meta-analyses were presented, we assessed the following subgroups where possible: age, TCS potency, anatomical site, treatment duration and genetic predisposition to a disrupted skin barrier (filaggrin status). TCS potency was determined using a hierarchy of sources: UK 'British National Formulary', WHO and USA classifications.^{19–21} A National Health Service classification ranging from very common (>1 in 10 people affected) to very rare (<1 in 10 000) was used to narratively describe the absolute risk of each adverse event.²²

Patient and public involvement

People with eczema and parents of children with eczema were involved in the decision to conduct this overview and in the design. The James Lind Alliance priority setting partnership for atopic eczema involved people with eczema and parents of children with eczema in which two of the identified priority areas were around research into the safety of TCS.¹⁰ Two of the overview authors are patient representatives (AR and AA) and both have been involved in the design of this overview and interpretation of the findings.

Wider patient and parent involvement has been particularly important in identifying important safety outcomes for this overview. We held a workshop involving five patient representatives in which the proposed overview was discussed which highlighted the need to seek out data on long-term TCS use, reversibility of any side effects and TCS withdrawal symptoms. We supplemented this with a survey about safety concerns with TCS at a National Eczema Society meeting of 31 people with eczema or parents of children with eczema and a published qualitative study of patient concerns relating to TCS safety.⁶

Dissemination of the results is underway as part of the wider programme of research of which this overview is a part and our patient representatives are a key stakeholder in this activity.

RESULTS

Search results

After deduplication, 635 records were screened; 127 records underwent full-text screening and 38 systematic reviews met the inclusion criteria (figure 1).⁷⁸²³⁻⁵⁶ The list of excluded reviews is in online supplemental appendix 2. The search of PROSPERO identified five ongoing systematic reviews (online supplemental appendix 3).⁵⁷⁻⁶¹

Characteristics and quality of the included systematic reviews All but three reviews were published in English. Two Chinese reviews and one German review were translated into English.^{32 36 45} Thirty of the included reviews were rated critically low quality according to AMSTAR 2; with four low, two moderate and two high quality (table 1). The most common reasons for downgrading were no

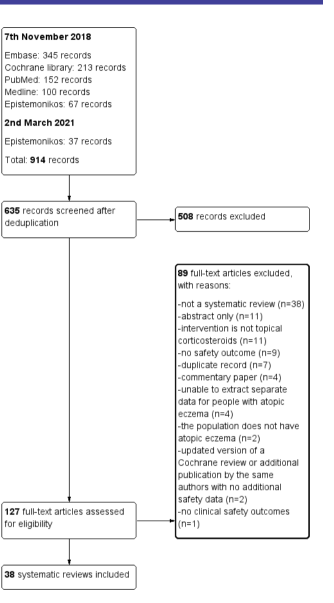


Figure 1 PRISMA flow diagram. PRISMA, Preferred Reporting Items for Systematic Reviews and Meta-Analyses.

protocol, no list of full-text exclusions or a literature search restricted to the English language.

The included reviews identified 106 studies (77 RCTs and 29 observational studies) that included relevant safety data. Risk of bias assessments were available from the reviews for 63 RCTs, of which 42 used the Cochrane risk of bias tool. Most of these assessments rated at least one domain as high or unclear risk, most noticeably selection bias from lack of allocation concealment, performance bias due to lack of blinding of participants and detection bias due to lack of blinding of outcome assessors. The trials included in the reviews usually evaluated use of short bursts of TCS (2-4 weeks) to treat the flare but varied greatly in length of follow-up. Around two-thirds of trials included no post-treatment follow-up, while the remainder included several weeks/months of follow-up generally using TCS intermittently 'as required'. A total of 14 RCTs (5874 participants) and 5 cohort/observational Table 1

First author, publication year

Ashcroft 2005²⁴

Ashcroft 2007²³

Barnes 2015²⁵

Braham 2010²⁶

Broeders 2016²⁷

Callen 2007²⁸

Chen 2010²⁹

Cury Martins 2015³⁰

De Tiedra 1997³¹

Devillers 2006³²

Eichenfield 2014³⁴

Feldman 2005³⁵

Fishbein 2019⁶³

Frangos 2008³⁶

Froeschl 2007³⁷

Gonzalez-Lopez

Green 2004⁷

Gu 2013⁴⁰

Gu 2014³⁹

Hajar 2015⁴¹

Hoare 2000⁴²

Iskedjian 2004⁴³

Juhász 201744

Abędź 2019⁸²

Li 2007⁴⁶

Legendre 2015⁴⁵

Nankervis 201647

2017³⁸

Dong 2017³³

Characteris	tics of included syst	ematic reviews		
thor, tion year	Type of review	Review contained safety data from RCTs for comparisons of interest?	Review contained safety data from observational studies?	AMSTAR 2 rating
: 2005 ²⁴	Non-Cochrane	Yes (TCS vs TCI)	No	Critically low 1367
: 2007 ²³	Cochrane	Yes (TCS vs TCI)	Yes (TCS vs TCI)	Moderate ⁸
2015 ²⁵	Non-Cochrane	Yes (TCS vs vehicle, TCS vs TCl, TCS vs another TCS)	Yes (single arm TCS studies)	Critically low 12346
2010 ²⁶	Non-Cochrane	Yes (occluded TCS vs non- occluded TCS)	Yes (occluded TCS)	Critically low ¹²³⁴⁶
s 2016 ²⁷	Non-Cochrane	Yes (TCS vs TCI)	No	Critically low 1356
007 ²⁸	Non-Cochrane	Yes (TCS vs vehicle, TCS vs another TCS)	Yes (single arm studies or comparing TCS potencies)	Critically low ¹²³⁴⁶
10 ²⁹	Non-Cochrane	Yes (TCS vs TCI)	No	Critically low ¹³⁶
artins 2015 ³⁰	Cochrane	Yes (TCS vs TCI)	Yes (TCS vs TCI)	Moderate ⁸
ra 1997 ³¹	Non-Cochrane	Yes (TCS vs another TCS)	Yes (usually only reported data from one arm of RCTs)	Critically low ¹²³⁴⁶
3 2006 ³²	Non-Cochrane	Yes (occluded TCS vs non- occluded TCS)	Yes (occluded TCS)	Critically low 12346
)17 ³³	Non-Cochrane	Yes (TCS vs TCI)	No	Critically low 12346
eld 2014 ³⁴	Non-Cochrane	No	Yes (different TCS potencies)	Critically low 12346
า 2005 ³⁵	Non-Cochrane	Yes (TCS vs vehicle)	No	Critically low 12346
2019 ⁶³	Non-Cochrane	Yes (TCS vs vehicle/moisturiser)	No	Critically low ³⁴⁵⁶⁷
2008 ³⁶	Non-Cochrane	Yes (TCS vs vehicle)	Yes (single arm studies)	Critically low 12346
l 2007 ³⁷	GMS HTA report	Yes (TCS vs vehicle, TCS vs TCl, TCS vs another TCS)	No	Critically low ¹²⁴⁶
z-Lopez	Non-Cochrane	Yes (occluded TCS vs non- occluded TCS)	No	Critically low ¹³
004 ⁷	HTA report	Yes (once daily vs twice daily TCS use)	No	Low
3 ⁴⁰	Cochrane	Yes (TCS vs topical CHM)	No	High
39	Non-Cochrane	Yes (TCS vs topical CHM)	No	Critically low 1237
15 ⁴¹	Non-Cochrane	No	Yes (case series or case reports)	Critically low ²³⁶
000 ⁴²	NIHR HTA report	Yes (TCS vs vehicle, TCS vs another TCS)	No	Low
ו 2004 ⁴³	Non-Cochrane	Yes (TCS vs vehicle, TCS vs TCI)	No	Critically low ¹³⁶
2017 ⁴⁴	Non-Cochrane	No	Yes (social media analysis)	Critically low 12346
019 ⁸²	Non-Cochrane	Yes (TCS vs TCI)	No	Critically low ¹³⁶⁷
e 2015 ⁴⁵	Non-Cochrane	No	Yes (TCS vs TCI)	Critically low 1236
6	Non-Cochrane	Yes (TCS vs TCI)	No	Critically low ¹³⁶
is 2016 ⁴⁷	NIHR HTA report	Yes (TCS vs vehicle, TCS vs emollients, TCS vs TCl, TCS vs another TCS, once a day vs twice a day use, proactive TCS to prevent flares ('weekend therapy') vs vehicle, occluded TCS vs non- occluded TCS)	No	Low

Continued

First author, publication yearReview contained safety data from subservational studiesKanstare 2 rationBurls 2004thWest Midlands HMSel CS vs TClNoLowSchmitt 2011thNon-CochraneYes (proactive TCS to prevent flares (weekend therapy) vs vehicleNoCritically low 36Sidbury 2014thNon-CochraneYes (proactive TCS to prevent flares (weekend therapy) vs vehicleNoCritically low 1234thSidbury 2014thNon-CochraneYes (proactive TCS to prevent flares (weekend therapy) vs vehicleNoCritically low 1234thSidbury 2014thNon-CochraneYes (proactive TCS to prevent flares (weekend therapy) vs vehicleNoCritically low 1234thSidbury 2014thNon-CochraneYes (TCS vs vehicle, TCS vs vehicleNoCritically low 1234thSingh 2012fhNon-CochraneYes (TCS vs vehicle, TCS vs vehicleNoCritically low 1234thSingh 2012fhNon-CochraneYes (TCS vs TC)NoCritically low 1234thSunsson 2011fhNon-CochraneYes (TCS vs TC)NoCritically low 134thNa Tzuren 2017fhCochraneYes (TCS vs TC)NoHighvan Zuuren 2017fhCochraneYes (TCS vs TC)NoHighYong SingherNon-CochraneYes (TCS vs TC)NoHighYong SingherNon-CochraneYes (TCS vs TC)NoHighYong SingherNon-CochraneYes (TCS vs TC)NoHighYong SingherNon-CochraneYes (TCS vs TC)No <th>Table 1 Continued</th> <th></th> <th></th> <th></th> <th></th>	Table 1 Continued				
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Wood HeickmanNon-CochraneNoYes (single arm cohortCritically low 123467201855studies)	Tang 2014 ⁵³	Non-Cochrane	flares ('weekend therapy') vs	No	Critically low ¹³⁴⁶
2018 ⁵⁵ studies)	van Zuuren 2017 ⁵⁴	Cochrane	Yes (TCS vs emollient)	No	High
Yan 200856Non-CochraneYes (TCS vs TCl)NoCritically low 1367		Non-Cochrane	No		
	Yan 2008 ⁵⁶	Non-Cochrane	Yes (TCS vs TCI)	No	Critically low ¹³⁶⁷

AMSTAR 2 ratings—reasons for downgrading the quality of the review: ¹No protocol; ²Search strategy not comprehensive; ³No list of excluded studies with reasons; ⁴Risk of bias not assessed; ⁵Inappropriate meta-analysis methods; ⁶Risk of bias assessments not included in the interpretation of the results; ⁷Publication bias not explored in the meta-analysis.

Additional data on TCS including potency can be found in online supplemental appendix 6.

CHM, Chinese herbal medicine; GMS, German Medical Science; HTA, Health Technology Assessment; NIHR, National Institute for Health Research; RCTs, randomised controlled trials; TCI, topical calcineurin inhibitors; TCS, topical corticosteroid.

studies (4 438 698 participants) out of a total of 106 studies included follow-up of more than 3 months. One notable trial (the 'PETITE' study) had 5 years follow-up with TCS used 'as required'.⁶²

Characteristics and quality assessments of each systematic review are in table 1, with further detail in online supplemental appendices 4 and 5. Individual study data and quality assessments are in online supplemental appendix 6.

Safety of TCS compared with other topical treatments or corticosteroids

How safe are TCS compared with emollient or vehicle, or no comparison?

Thirteen reviews provided data on this comparison: 1 high⁵⁴, 2low⁴²⁴⁷ and 10 critically low quality.^{25283135–3750515563} Key results can be found in table 2 and additional data in

online supplemental appendix 6.

Reported rates of skin thinning in RCTs were generally very low, with no significant increases seen with TCS compared with emollient/vehicle. No skin thinning or telangiectasia was reported in an RCT, 196 participants aged ≥ 12 years old using very potent TCS twice a day for 2 weeks compared with 33 using vehicle.⁶⁴ Another RCT reported skin thinning in 6/109 participants using very potent TCS for 2 weeks compared with 2/50 using vehicle, p=0.69.⁶⁵

No significant differences in other cutaneous adverse events, such as hypopigmentation, were observed between treatments in five RCTs, and event rates were low. $^{66-70}$

A meta-analysis⁵⁵ of 11 uncontrolled observational studies (up to 4 weeks of treatment) reported biochemical adrenal suppression (cortisol levels) in 20/522 children (3.8%, 95% CI 2.4% to 5.8%) with any potency TCS.^{71–81} This was 2% (3/148 children) when only mild potency TCS were analysed.^{72 74 77 79} No clinical symptoms or signs of adrenal suppression were observed,^{71–81} and the biochemical effects were transient, with cortisol levels returning to normal after TCS were discontinued.⁷¹⁷⁵⁷⁷⁷⁸⁸¹

Two included reviews assessed TCS withdrawal symptoms, mostly from case reports, but no incidence data were reported.^{41 44}

How safe are TCS compared with topical calcineurin inhibitors?

Eight systematic reviews were identified: one moderate²³, one low⁴⁸ and six critically low quality.^{27 30 43 50 52 82} Most RCTs used twice daily TCS to treat the current flare (up to 3 weeks), and where longer-term follow-up was included, TCSs were used 'as required' to treat flares. Key results

 Table 2
 Summary of main findings for key safety outcomes

	Cutaneous adverse events	Systemic adverse events
How safe are TCS compared with emollient or vehicle, or no comparison? 13 reviews: 1 moderate quality 2 low quality 10 critically low quality	 Skin thinning: No significant differences in 2 RCTs of 2–4 weeks compared with emollient vehicle: (1) 0/196 children with very potent TCS and 0/33 vehicle, (2) 6/109 very potent TCS vs 2/50 vehicle, p=0.69. Very low rates. Other cutaneous adverse events: No significant differences in 5 RCTs (2–4 weeks) between TCS (various potencies) and emollient/vehicle (n=172, plus one study, n not specified). Low event rates. 	/ suppression: Meta-analysis (11 observational studies, max 4 weeks)—20/522 children with any potency TCS (3.8%, 95% CI 2.4% to 5.8%), 3/148 children (2%) with mild
How safe are TCS compared with topical calcineurin inhibitors (TCI)? 8 reviews: 1 moderate quality 1 low quality 6 critically low quality	 Skin thinning: Higher with TCS than TCI (meta-analysis of 4 RCTs: RR 4.86, 95% 1.06 to 22.28, n=4128) but very low rate (8/2068, 7 of which were using potent TCS). Other cutaneous adverse events: No difference in skin infections between TCS and TCI (8 RCTs). Skin burning and pruritus lower with TCS than TCI: meta-analysis of 10 RCTs: burning—RR 0.31, 95% CI 0.23 to 0.40 (n=4211), pruritus—RR 0.68, 95% CI 0.56 to 0.82(n=4211). 	 2418 children with 5 years follow-up). Lymphoma: no cases reported in one same large RCT as above. One cohor study (n=1 438 333, approx. 4 years follow-up)—very small non-significant increase with TCI and TCS compared
How safe are once daily TCS compared with twice daily application? 2 reviews: 2 low quality	 Skin thinning: no cases using once daily vs twice daily potent TCS for 3 weeks (1 RCT, 94 adults). Other cutaneous adverse events: no significant difference between groups in telangiectasia, folliculitis, or burning/itching/stinging (4 RCTs, 4–16 weeks follow-up 740 older children/adults). 	Biochemical evidence of adrenal suppression: no significant difference between once and twice daily moderate/potent TCS up to 2 weeks children (2 RCTs, n=129).
How safe are TCS used proactively to prevent flares ('weekend therapy')? 3 reviews: 3 critically low quality	 Skin thinning: no cases with 16–20 weeks of 2 days/week of potent TCS vs vehicle (5 RCTs, n=993). Other cutaneous adverse events: no significant differences between groups, including folliculitis and transient telangiectasia, with potent TCS (16–20 	Biochemical evidence of adrenal suppression: no cases with 16 weeks of 2 days/week of potent TCS (2 RCTs n=129). Possible adrenal suppression in 2/44 children with potent TCS compared with zero using vehicle (1 RCT, 20 weeks).

weeks) compared with either vehicle or

another TCS (2 RCTs, n=423). Events were uncommon in both groups. How safe are TCS used under Skin thinning: no cases in two observational > occlusion? studies (potent TCS +wet wrap, 1-2 weeks, 4 reviews: n=44). 1 high quality Other cutaneous adverse events: One 3 critically low quality case of striae in two observational studies, n-44. More folliculitis with diluted potent TCS (10/19 children) compared with emollient (2/20), both under wet wrap (1 RCT). A meta-analysis (2 RCTs, n=69) of wet wrap vs no wet wrap (mild potency)-no significant difference in cutaneous adverse events.

RCTs, randomised controlled trials; RR, relative risk; TCS, topical corticosteroids.

studies.

up 12 weeks).

Biochemical evidence of adrenal

observational studies (2-14 days of

diluted potent TCS under wet-wraps in

74 children) but rates not specified in

review. Described as transient in two

effect seen in one small short-term

observational study (potent TCS wet-

wrap in eight children, (median follow-

suppression: reported in three

Growth or bone turnover: no

can be found in table 2 and additional data in online supplemental appendix 6.

Meta-analyses of cutaneous adverse events were presented in two reviews.^{27 82} So the more comprehensive review was used to extract the cutaneous adverse event data.²⁷ Some minor modifications were made to the data for this overview shown in online supplemental appendix 7. A meta-analysis of four RCTs (26 weeks to 5 years duration, twice a day or 'as directed') showed a significant increase in the RR of skin thinning with TCS compared with topical calcineurin inhibitors (TCIs) (0.1% tacrolimus or 1% pimecrolimus) (RR 4.86, 95% CI 1.06 to 22.28, p=0.04, n=4128). However, skin thinning was uncommon: 8/2068 participants (0.4%) with TCS vs 0/2060 (0%) with TCIs. Of the eight cases of skin thinning, seven were reported when using potent TCS and one using mild/moderate TCS.^{62 83–85}

The RR of skin burning and pruritus (itching) was significantly lower with TCS compared with TCIs (1% pimecrolimus or 0.1 % / 0.03% tacrolimus) in metaanalyses of 10 RCTs in 4211 participants (skin burning: RR 0.31, 95% CI 0.23 to 0.40, p<0.00001; pruritus: RR 0.68, 95% CI 0.56 to 0.82, p<0.0001). ^{83 85–93} The GRADE assessments for these two adverse events indicated these were of moderate quality. ⁸² There was no significant difference in skin infections with potent, moderate or mild potency TCS compared with TCIs (1% pimecrolimus or 0.1 %/0.03% tacrolimus) ^{62 83–86 88 90 92} or erythema compared with 0.1% tacrolimus (online supplemental appendix 8).^{91 92}

Subgroup analyses of age, TCS potency and specific TCI showed no significant differences for any comparison (online supplemental appendix 9). We were unable to undertake any further subgroup analyses.

No differences in growth were observed in one 5-year RCT ('PETITE' study) in 2418 young children using moderate/mild potency TCS compared with those using TCI (1% pimecrolimus) (rates not given) and no cases of lymphoma were reported.⁶² A large cohort study (n=1 438 333) showed a small non-significant increased risk of lymphoma with TCI and TCS compared with the general population, with a similar risk between treatments.⁹⁴ In addition, one case–control study (294 cases/293 000 controls) found no increased risk of lymphoma with TCS or TCI compared with controls.⁹⁵

Is there any difference in safety of TCS of different potencies?

Six reviews compared the safety of different potency TCS: two low,^{42 47} and four critically low quality.^{28 34 50 53} RCTs were mainly short-term use of TCS (2–3 weeks), used once or twice daily. Results can be found in online supplemental appendix 6.

One RCT reported mild skin thinning in 4/13 children using potent TCS for up to 6 weeks compared with 2/12 using mild TCS (p=0.42),⁹⁶ while another RCT in 37 children found no evidence of skin thinning with mild or moderate potency TCS for 3 weeks.⁹⁷ One study compared 3 weeks of potent and moderate TCS in

40 children and reported 'some' biochemical adrenal suppression (cortisol levels) but no numerical data were provided.⁹⁸

How safe are TCS compared with topically applied Chinese herbal medicine?

Two systematic reviews provided data on TCS compared with topical Chinese herbal medicine: one high quality⁴⁰ and one critically low.³⁹ Results can be found in online supplemental appendix 6.

A meta-analysis of two RCTs^{99 100} was presented in two systematic reviews.^{39 40} More cutaneous adverse events, including application site burning, were observed with 2 weeks of very potent/potent TCS compared with topical Chinese herbal medicine (RR 12.03, 95% CI 1.59 to 91.26, p=0.02; 11/147 vs 0/148 participants). One additional RCT, including 95 young children, reported minor adverse events such as burning with 2 weeks of potent TCS but no numerical data were presented.¹⁰¹

Safety of different strategies for using TCS

How safe are once daily TCS compared with more frequent application?

Two low-quality reviews provided safety data relating to different frequency of application.^{7 47} Key results can be found in table 2 and additional data in online supplemental appendix 6.

No skin thinning was reported with once or twice daily application of potent TCS for 3 weeks in one RCT (94 adults).¹⁰² Four RCTs in 740 older children/adults showed no significant difference between once and twice daily application of moderate/potent TCS in other cutaneous adverse events including telangiectasia,^{103 104} folliculitis¹⁰⁵ and burning, itching or stinging.^{105 106} Two RCTs showed no significant differences in biochemical adrenal suppression (cortisol levels) between once and twice daily very potent/moderate TCS used for up to 2 weeks in 129 children.^{81 107}

How safe are TCS when used proactively to prevent flares ('weekend therapy')?

Two reviews included data on the safety of TCS used proactively 2 days a week ('weekend therapy') to prevent flares, both critically low quality.^{8 53} Key results can be found in table 2 and additional data in online supplemental appendix 6.

There was no evidence of skin thinning in five RCTs comparing 16–20 weeks of weekend therapy with potent TCS versus vehicle in 993 participants.¹⁰³ ^{108–111} Furthermore, two RCTs (n=423) reported no significant differences in other cutaneous adverse events, including folliculitis and transient telangiectasia, with potent TCS compared with vehicle.¹⁰⁸ ¹⁰⁹ Events were uncommon in both groups.

There was no evidence of biochemical adrenal suppression (cortisol levels) in two RCTs (n=129) between potent TCS and vehicle used for 16 weeks.^{$108 ext{ 111}$} In a 20-week

RCT, 2/44 children had possible adrenal suppression with potent TCS compared with zero with vehicle.¹⁰⁹

How safe are TCS used under occlusion?

Four reviews included data on the safety of TCS used under occlusion: one high⁵⁴, and three critically low quality.^{26 32 38} Results can be found in online supplemental appendix 6.

There were no cases of skin thinning and one case of striae in two uncontrolled observational studies of a diluted potent TCS used under wet-wrap for 1–2 weeks in 44 young children.^{112 113} A significant difference in the rate of folliculitis (mostly mild) was observed in one RCT of TCS under wet-wrap for 4 weeks, with more folliculitis in the diluted potent TCS group (10/19 children) compared with emollient (2/20 children) (p=0.02).¹¹⁴ A meta-analysis from one review³⁸ of two RCTs in young children showed no significant difference in the number of participants with cutaneous adverse events between mild potency TCS under wet wrap (7/38 participants) versus not under wet-wrap (0/31 participants) (p=0.08)^{115 116}; this evidence was rated low quality by the systematic review authors using GRADE.¹⁷

Biochemical adrenal suppression (cortisol levels) was reported in three uncontrolled observational studies of 2–14 days of diluted potent TCS under wet-wraps in 74 children.^{112 113 117} Actual rates were not specified in the review, but increases were described as transient in two studies.^{112 117} One short-term uncontrolled observational study of diluted potent TCS under wet-wrap in eight children showed no effect on growth or bone turnover.¹¹⁸

DISCUSSION

This comprehensive overview of systematic reviews which, for the first time, brings together all safety data from systematic reviews on TCS used in eczema from 38 systematic reviews, a topic that was identified as a priority in a James Lind Alliance priority setting partnership on eczema. Skin thinning and effects on growth concern many people with eczema and parents of children with eczema when using TCS. However, we found no evidence of skin thinning when TCS were used intermittently 'as required' to treat flares or as 'weekend therapy' to prevent flares, although the majority of data was from shortterm studies.⁵ Similarly, we found no evidence of growth retardation or clinically significant adrenal suppression but the only data available was from one 5-year study that included 1213 children using TCS.⁶² Other studies only reported biochemical signs of adrenal suppression. Adherence to TCS treatment is known to be poor and these findings, particularly around skin thinning, may encourage appropriate use of TCS and therefore improve treatment effectiveness and patient benefit.¹¹⁹

A thorough literature search was conducted and Cochrane methodology was used. Conclusions were limited by the content of the included reviews because safety was frequently reported in less detail than effectiveness, reviews reported on different adverse events and some adverse events were not described in the reviews. It is not clear whether this is because the trials did not report adverse events in sufficient detail or whether the review authors did not include all the available safety data, perhaps only focusing on a restricted group of adverse events. None of the included systematic reviews presented data on our prespecified subgroup analyses. Furthermore, most of the included reviews were rated low or critically low-quality using AMSTAR 2. The lack of comprehensive search strategies and duplicate screening/data extraction in the included reviews may have resulted in missing studies and safety data, which could have impacted on this overview particularly where there was limited data. In addition, where the quality of evidence assessments (eg, GRADE) were reported in the reviews, most individual studies included in the reviews indicated a high or unclear risk in at least one domain.

Many RCTs did not include follow-up beyond 2-4 weeks of treatment and therefore data on long-term safety are limited. Although short-term TCS use reflects appropriate treatment duration for treating an individual flare, it does not reflect the chronic nature of eczema and the need for TCS use over the long-term. The 'PETITE study' was the notable exception and data published in the correspondence showed there was only one episode of skin thinning in 1213 children using mild/moderate TCS 'as required' with 5-year follow-up.⁶² Trials using intermittent TCS as 'weekend therapy' to prevent flares also provide reassurance for the safety of longer-term use of TCS, as these trials generally included 16-20 weeks of follow-up to assess the prevention of flares. The inclusion of systematic reviews that included observational studies as well as reviews of RCTs also increased the amount of safety data available to report in this overview.

Although this review focused on the safety of TCS as the key issue for patients, treatment decisions are a balance of benefits and harms. For example, although the safety profile of Chinese herbal medicine was better than TCS, in practice this would be considered alongside the relative effectiveness of these treatments. Likewise, although there was no difference in the safety of once vs twice daily TCS, effectiveness of these regimens is also important to consider. A Cochrane review is underway comparing the effectiveness and safety of different ways of using TCS.¹²⁰

In summary, we found no evidence that TCS cause harm when used intermittently 'as required' to treatment eczema flares or as 'weekend therapy' to prevent flares and this should support the use of TCS in the management of eczema. We found that the adverse events of greatest concern to patients and clinicians, such as skin thinning, are uncommon with short-term use of TCS. However, high-quality evidence was limited, particularly for long-term use. Rather than follow-up of perhaps just a few weeks, future RCTs should include lengthier follow-up to enable better safety assessment. However, it should be noted that longer-term prospect observational studies are better placed to explore longer-term safety of TCS and should be designed with years rather than months of follow-up to add useful information to the field. Perhaps equally as important as duration of follow-up in trials is resolution of adverse events which is often not reported. For adverse events such as biochemical signs of adrenal suppression, it is crucial to know if the effect is transient and levels return to normal once the TCS is stopped, particularly as it is not clear how to interpret the clinical relevance of these.

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Appendix 1: Search strategies

Search facets for searches: Topical steroids Eczema Systematic reviews

PubMed search

Uses PubMed Clinical Queries systematic review filter (command systematic[sb]): https://www.nlm.nih.gov/bsd/pubmed_subsets/sysreviews_strategy.html

(steroid* OR corticosteroid* OR glucocorticosteroid* OR glucocorticoid* OR glucocorticoids[MeSH Terms] OR alclometasone OR alclomethasone OR amcinonide OR beclometasone OR beclomethasone OR beclomethasone[MeSH Terms] OR betametasone OR betamethasone OR betamethasone[MeSH Terms] OR clobetasol OR clobetasol[MeSH Terms] OwR clobetasone OR desonide OR desonide[MeSH Terms] OR desoximetasone OR desoximetasone[MeSH Terms] OR diflorasone OR diflucortolone OR diflucortolone[MeSH Terms] OR fludroxycortide OR flumetasone OR flumethasone OR flumethasone[MeSH Terms] OR fluocinolone OR fluocinolone acetonide[MeSH Terms] OR fluocinonide OR fluocinonide[MeSH Terms] OR fluocortolone OR fluocortolone[MeSH Terms] OR flurandrenolide OR flurandrenolone OR flurandrenolone[MeSH Terms] OR fluticasone OR halcinonide OR halcinonide[MeSH Terms] OR halobetasol OR halometasone OR hydrocortisone OR hydrocortisone[MeSH Terms] OR methylprednisolone OR methylprednisolone[MeSH Terms] OR mometasone OR triamcinolone OR triamcinolone[MeSH Terms]) AND ("dermatitis, atopic"[MeSH Terms] OR "eczema"[MeSH Terms] OR "neurodermatitis"[MeSH Terms] OR eczema OR "atopic dermatitis" OR neurodermatitis) AND (systematic[sb] OR "systematic review")

Ovid MEDLINE search

Uses SIGN MEDLINE systematic review filter: http://www.sign.ac.uk/search-filters.html Ovid MEDLINE(R) Epub Ahead of Print, In-Process & Other Non-Indexed Citations, Ovid MEDLINE(R) Daily, Ovid MEDLINE and Versions(R) MEDLINE eczema steroid systematic reviews 1. Meta-Analysis as Topic/ 2. meta analy\$.tw. 3. metaanaly\$.tw. 4. Meta-Analysis/ 5. (systematic adj (review\$1 or overview\$1)).tw. 6. exp Review Literature as Topic/ 7. or/1-6 8. cochrane.ab. 9. embase.ab. 10. (psychlit or psyclit).ab. 11. (psychinfo or psycinfo).ab. 12. (cinahl or cinhal).ab. 13. science citation index.ab. 14. bids.ab. 15. cancerlit.ab. 16. or/8-15 17. reference list\$.ab. 18. bibliograph\$.ab.

19. hand-search\$.ab. 20. relevant journals.ab. 21. manual search\$.ab. 22. or/17-21 23. selection criteria.ab. 24. data extraction.ab. 25. 23 or 24 26. Review/ 27. 25 and 26 28. Comment/ 29. Letter/ 30. Editorial/ 31. animal/ 32. human/ 33. 31 not (31 and 32) 34. or/28-30,33 35. 7 or 16 or 22 or 27 36. 35 not 34 37. steroid\$.mp. 38. corticosteroid\$.mp. 39. glucocorticosteroid\$.mp. 40. glucocorticoid\$.mp. 41. exp Glucocorticoids/ 42. alclometasone.mp. 43. alclomethasone.mp. 44. amcinonide.mp. 45. beclometasone.mp. 46. beclomethasone.mp. 47. exp Beclomethasone/ 48. betametasone.mp. 49. betamethasone.mp. 50. exp Betamethasone/ 51. clobetasol.mp. 52. exp Clobetasol/ 53. clobetasone.mp. 54. desonide.mp. 55. exp Desonide/ 56. desoximetasone.mp. 57. exp Desoximetasone/ 58. diflorasone.mp. 59. diflucortolone.mp. 60. exp Diflucortolone/ 61. fludroxycortide.mp. 62. flumetasone.mp. 63. flumethasone.mp. 64. exp Flumethasone/ 65. fluocinolone.mp. 66. exp Fluocinolone Acetonide/ 67. fluocinonide.mp.

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- 78. halometasone.mp.
- 79. hydrocortisone.mp.
- 80. exp Hydrocortisone/
- 81. methylprednisolone.mp.
- 82. exp methylprednisolone/
- 83. mometasone.mp.
- 84. triamcinolone.mp.
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- 86. or/37-85
- 87. exp dermatitis, atopic/
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- 89. exp neurodermatitis/
- 90. eczema.mp.
- 91. atopic dermatitis.mp.
- 92. neurodermatitis.mp.
- 93. or/87-92
- 94. 36 and 86 and 93

Ovid Embase search

Uses SIGN Embase systematic review filter: http://www.sign.ac.uk/search-filters.html Embase 1974 to 2017 October 23 Embase eczema steroid systematic reviews 1. exp Meta Analysis/ 2. ((meta adj analy\$) or metaanalys\$).tw. 3. (systematic adj (review\$1 or overview\$1)).tw. 4. or/1-3 5. cancerlit.ab. 6. cochrane.ab. 7. embase.ab. 8. (psychlit or psyclit).ab. 9. (psychinfo or psycinfo).ab. 10. (cinahl or cinhal).ab. 11. science citation index.ab. 12. bids.ab. 13. or/5-12 14. reference lists.ab. 15. bibliograph\$.ab. 16. hand-search\$.ab.

17. manual search\$.ab. 18. relevant journals.ab. 19. or/14-18 20. data extraction.ab. 21. selection criteria.ab. 22. 20 or 21 23. review.pt. 24. 22 and 23 25. letter.pt. 26. editorial.pt. 27. animal/ 28. human/ 29. 27 not (27 and 28) 30. or/25-26,29 31. 4 or 13 or 19 or 24 32. 31 not 30 33. steroid\$.mp. 34. corticosteroid\$.mp. 35. exp corticosteroid/ 36. glucocorticosteroid\$.mp. 37. glucocorticoid\$.mp. 38. exp glucocorticoid/ 39. alclometasone.mp. 40. alclomethasone.mp. 41. amcinonide.mp. 42. beclometasone.mp. 43. beclomethasone.mp. 44. betametasone.mp. 45. betamethasone.mp. 46. clobetasol.mp. 47. clobetasone.mp. 48. desonide.mp. 49. desoximetasone.mp. 50. diflorasone.mp. 51. diflucortolone.mp. 52. fludroxycortide.mp. 53. flumetasone.mp. 54. flumethasone.mp. 55. fluocinolone.mp. 56. fluocinonide.mp. 57. fluocortolone.mp. 58. flurandrenolide.mp. 59. flurandrenolone.mp. 60. fluticasone.mp. 61. halcinonide.mp. 62. halobetasol.mp. 63. halometasone.mp. 64. hydrocortisone.mp. 65. methylprednisolone.mp.

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Epistemonikos

(steroid* OR corticosteroid* OR glucocorticosteroid* OR glucocorticoid* OR alclometasone OR alclomethasone OR amcinonide OR beclometasone OR beclomethasone OR betametasone OR betamethasone OR clobetasol OR clobetasone OR desonide OR desoximetasone OR diflorasone OR diflucortolone OR fludroxycortide OR flumetasone OR flumethasone OR fluocinolone OR fluocinonide OR fluocortolone OR flurandrenolide OR flurandrenolone OR fluticasone OR halcinonide OR halobetasol OR halometasone OR hydrocortisone OR methylprednisolone OR mometasone OR triamcinolone) AND (eczema OR "atopic dermatitis" OR neurodermatitis)

Enter search into advanced search and choose "**systematic review**" from drop-down box for "Publication type":

https://www.epistemonikos.org/advanced_search?q=(steroid*%20OR%20corticosteroid*%20OR%20glucocort icosteroid*%20OR%20glucocorticoid*%20OR%20alclometasone%20OR%20alclomethasone%20OR%20amcino nide%20OR%20beclometasone%20OR%20beclomethasone%20OR%20betametasone%20OR%20betamethaso ne%20OR%20clobetasol%20OR%20clobetasone%20OR%20desonide%20OR%20desoximetasone%20OR%20difl orasone%20OR%20diflucortolone%20OR%20fludroxycortide%20OR%20flumetasone%20OR%20flumethasone %20OR%20fluocinolone%20OR%20fluocinonide%20OR%20fluocortolone%20OR%20flurandrenolide%20OR%2 offlurandrenolone%20OR%20fluticasone%20OR%20halcinonide%20OR%20halobetasol%20OR%20halometason e%20OR%20hydrocortisone%20OR%20methylprednisolone%20OR%20mometasone%20OR%20triamcinolone} %20AND%20(eczema%20OR%20%22atopic%20dermatitis%22%20OR%20neurodermatitis)&protocol=no&class ification=systematic-review

Cochrane Library

(steroid* OR corticosteroid* OR glucocorticosteroid* OR glucocorticoid* OR [mh "glucocorticoids"] OR alclometasone OR alclomethasone OR amcinonide OR beclometasone OR beclomethasone OR betametasone OR betamethasone OR clobetasol OR clobetasone OR desonide OR desoximetasone OR diflorasone OR diflucortolone OR fludroxycortide OR flumetasone OR flumethasone OR fluocinolone OR fluocinonide OR fluocortolone OR flurandrenolide OR flurandrenolone OR fluticasone OR halcinonide OR halobetasol OR halometasone OR hydrocortisone OR methylprednisolone OR mometasone OR triamcinolone) AND ([mh "eczema"] OR [mh "dermatitis, atopic"] OR [mh "neurodermatitis"] OR eczema OR "atopic dermatitis" OR neurodermatitis)

"Search all text" option chosen.

Cochrane Reviews, Other Reviews (i.e. DARE), and Technology Assessments (i.e. HTA) chosen.

Appendix 2 - list of excluded studies with reasons

Excluded study	Reason for exclusion
Abramovits 2005 ⁽¹⁾	Not a systematic review
Abramovits 2006 ⁽²⁾	Not a systematic review
Anonymous 1995 ⁽³⁾	Not a systematic review
Anonymous 1999 ⁽⁴⁾	Not a systematic review
Anonymous 2004 ⁽⁵⁾	Not a systematic review
Anonymous 2005 ⁽⁶⁾	Abstract
Anonymous 2007 ⁽⁷⁾	Not a systematic review
Anonymous 2015 ⁽⁸⁾	Abstract
Anonymous 2015 ⁽⁹⁾	Abstract
Aslam 2014 ⁽¹⁰⁾	Not a systematic review
Barfield 2017 (11)	Wrong intervention (not topical corticosteroids)
Batchelor 2010 (12)	Not a systematic review
Bath-Hextall 2010 ⁽¹³⁾	Updated version of a Cochrane review (non-Cochrane) but no additional safety data
Bigby 2001 ⁽¹⁴⁾	Commentary paper
Bonchak 2017 ⁽¹⁵⁾	Wrong intervention (not topical corticosteroids)
Birnie 2008 ⁽¹⁶⁾	Wrong intervention (not topical corticosteroids)
Boucher 2001 ⁽¹⁷⁾	Not a systematic review
Broersen 2015 (18)	Unable to extract separate data for atopic eczema patients
Cameron 2000 ⁽¹⁹⁾	Commentary paper
Carbone 2010 ⁽²⁰⁾	Not a systematic review
Chavigny 2005 (21)	Not a systematic review
Chi 2009 ⁽²²⁾	Unable to extract separate data for atopic eczema patients
Chi 2015 ⁽²³⁾	Unable to extract separate data for atopic eczema patients
Chia 2015 (24)	Not a systematic review
Chu 1995 ⁽²⁵⁾	Not a systematic review
Conroy 2004 ⁽²⁶⁾	Not a systematic review
Das 2017 ⁽²⁷⁾	Not a systematic review
El-Batawy 2009 ⁽²⁸⁾	No safety outcome
Fleischer Jr 2010 ⁽²⁹⁾	Wrong intervention (not topical corticosteroids)
Frohna 2005 ⁽³⁰⁾	Commentary paper
Froschl 2007 ⁽³¹⁾	Duplicate record of an included systematic review
Furue 2006 ⁽³²⁾	Not a systematic review
Furue 2006 ⁽³³⁾	Not a systematic review
Garside 2005 ⁽³⁴⁾	No safety outcome
Ghajar 2019 ⁽³⁵⁾	'Subgroup analysis' of an included review (Wood Heickman 2018) – no additional safety data
Goustas 2003 ⁽³⁶⁾	Not a systematic review
Green 2005 ⁽³⁷⁾	Duplicate record of an included systematic review
Green 2004 ⁽³⁸⁾	Duplicate record of an included systematic review
Halling-Overgaard 2017 ⁽³⁹⁾	Skin atrophy is not assessed clinically in this review
Health Technology Assessment	Not a systematic review
Database 2004 ⁽⁴⁰⁾	· · · · · · · · · · · · · · · · · · ·
Health Technology Assessment	Abstract – unable to find the full publication
Database 2004 ⁽⁴¹⁾	
Health Technology Assessment	Abstract – unable to find the full publication
Database 2001 ⁽⁴²⁾	· · · · · · · · · · · · · · · · · · ·
Health Technology Assessment	Abstract – unable to find the full publication
Database 2004 ⁽⁴³⁾	
Hannuksela 2000 ⁽⁴⁴⁾	Wrong intervention (not topical corticosteroids)
Hebert 2006 (45)	Wrong intervention (not topical corticosteroids)

(46)	Development of the first set of the sector of the sector
Hoare 2000 ⁽⁴⁶⁾	Duplicate record of an included systematic review
Hon 2011 ⁽⁴⁷⁾	Wrong intervention (not topical corticosteroids)
Hulshof 2017 (48)	Wrong intervention (not topical corticosteroids)
Hussain 2016 (49)	Not a systematic review
Kaufman 2016 (50)	Abstract
Legendre 2015 ⁽⁵¹⁾	Abstract
Li 2017 ⁽⁵²⁾	Abstract
Li 2017 ⁽⁵³⁾	No safety outcome
Meffert 1999 ⁽⁵⁴⁾	Not a systematic review
Mooney 2015 (55)	Not a systematic review
Murashkin 2016 ⁽⁵⁶⁾	Not a systematic review
Nankervis 2013 ⁽⁵⁷⁾	Abstract
Nankervis 2016 ⁽⁵⁸⁾	Duplicate record of an included systematic review
Nankervis 2017 ⁽⁵⁹⁾	Duplicate record of an included systematic review
Nowak 2017 ⁽⁶⁰⁾	No safety outcome
Orlow 2007 (61)	Not a systematic review
Pan 2013 ⁽⁶²⁾	No safety outcome
Park-Wyllie 2000 (63)	Unable to extract separate data for atopic eczema patients
Payne 2019 (64)	Wrong intervention (not topical corticosteroids)
Phipatanakul 2006 (65)	Commentary paper
Radovic 2017 ⁽⁶⁶⁾	Wrong intervention (not topical corticosteroids)
Ricci 2007 ⁽⁶⁷⁾	Not a systematic review
Ruzicka 1999 ⁽⁶⁸⁾	Wrong intervention (not topical corticosteroids)
Sanchez 2014 (69)	Not a systematic review
Schiffner 2003 ⁽⁷⁰⁾	No safety outcome
Schmitt 2011 (71)	Not a systematic review
Schmitt 2011 (72)	Duplicate record of an included systematic review
Sher 2012 ⁽⁷³⁾	Abstract
Sher 2012 ⁽⁷⁴⁾	No safety outcome
Siegfried 2013 (75)	Not a systematic review
Siegfried 2018 (76)	Not a systematic review
Silverberg 2014 (77)	Not a systematic review
Simpson 2010 ⁽⁷⁸⁾	Not a systematic review
Spada 2018 (79)	Not a systematic review
Torii 2003 ⁽⁸⁰⁾	No safety outcome
Torley 2013 (81)	Not a systematic review
Uppal 2020 ⁽⁸²⁾	Wrong intervention (not topical corticosteroids)
Van Zuuren 2017 ⁽⁸³⁾	No safety outcome (abridged Cochrane review)
Wat 2014 ⁽⁸⁴⁾	Wrong patient population (not atopic eczema)
Wellington 2004 (85)	Not a systematic review
Williams 2007 ⁽⁸⁶⁾	Not a systematic review
Williams 2008 ⁽⁸⁷⁾	Not a systematic review
Williams 2010 ⁽⁸⁸⁾	Not a systematic review
Wollenberg 2018 ⁽⁸⁹⁾	Not a systematic review

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Appendix 3: Prospero search

Prospero – searched up to 23rd March 2021 – search results from 'Eczema OR dermatitis'

Potentially relevant ongoing systematic reviews:

Prospero ID	Date registered	Title	Status	Anticipated completion date
CRD42015016525 ⁽¹⁾	11 February 2015	Effects of emollients in the management of atopic dermatitis in pediatric patients a systemic review and meta-analysis	Review ongoing	25 February 2016
CRD42015027873 ⁽²⁾	04 November 2015	Interventions to improve quality of life in paediatric atopic dermatitis: a systematic review	Review ongoing	01 January 2016
CRD42020190452 ⁽³⁾	14 July 2020	The association between topical calcineurin inhibitor use and risk of cancer: a systematic review and meta-analysis	Review ongoing	31 August 2020
CRD42020161558 ⁽⁴⁾	28 April 2020	Efficacy of Non-Steroidal Topical Therapies for Atopic Dermatitis: A Systematic Review & Meta-Analysis	Review ongoing	31 May 2020
CRD42021230047 ⁽⁵⁾	31 October 2021	A network meta-analysis of five categories of external therapy of traditional Chinese for common diseases of dermatology	Review ongoing	31 October 2021

1. Tan Q, Tan C, Peng W, Shi Y, Xia L. Effects of emollients in the management of atopic dermatitis in pediatric patients a systemic review and meta-analysis [CRD42015016525] 2015 (accessed: 27/03/21). Available from:

http://www.crd.york.ac.uk/PROSPERO/display_record.php?ID=CRD42015016525.

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3. Lam M, Zhu J, Tadrous M, Drucker A. The association between topical calcineurin inhibitor use and risk of cancer: a systematic review and meta-analysis [CRD42020190452]2020 (accessed: 27/03/2021). Available from: www.crd.york.ac.uk/prospero/display record.php?RecordID=190452.

4. Lee K. Efficacy of Non-Steroidal Topical Therapies for Atopic Dermatitis: A Systematic Review & Meta-Analysis [CRD42020161558] 2020 (accessed: 27/03/2021). Available from: www.crd.york.ac.uk/prospero/display_record.php?RecordID=161558.

5. Ruirui L, Jing G, Dingxi B, Qian Y, Lin Z, Zhi Y, et al. A network meta-analysis of five categories of external therapy of traditional Chinese for common diseases of dermatology [CRD42021230047]2021 (assessed: 27/03/2021). Available from: www.crd.york.ac.uk/prospero/display_record.php?RecordID=230047.

Appendix 4: AMSTAR 2 ratings ⁽¹⁾:

Review ID	1 PICO	2 Protocol*	3 Study designs	4 Search strategy*	5 Duplicate screening	6 Duplicate data extraction	7 Excluded studies*	8 Included studies	9 Risk of bias assessed*	10 Funding of studies	11 Appropriate meta- analysis*	12 Risk of bias in meta- analysis	13 Risk of bias in discussion *	14 Hetero- geneity	15 Publication bias in meta- analysis*	16 Reviewers' conflict of interest	Overall rating
Ashcroft 2005 ⁽²⁾	Yes	No	No	Yes	Yes	Yes	No	Partial yes	Partial yes	No	Yes	No	No	No	No	Yes	Critically low
Ashcroft 2007 ⁽³⁾	Yes	Partial yes	No	Yes	Yes	Yes	Yes	Partial yes	Yes	No	N/A	N/A	Yes	Yes	N/A	Yes	Moderate
Barnes 2015 ⁽⁴⁾	No	No	No	No	No	No	No	No	No	No	N/A	N/A	No	No	N/A	Yes	Critically low
Braham 2010 ⁽⁵⁾	No	No	Yes	No	No	No	No	Partial yes	No	No	N/A	N/A	No	No	N/A	Yes	Critically low
Broeders 2016 ⁽⁶⁾	Yes	No	No	Partial yes	No	No	No	Partial yes	Partial yes	Yes	No	No	No	No	Yes	Yes	Critically low
Callen 2007	Yes	No	No	No	No	No	No	No	No	No	N/A	N/A	No	No	N/A	Yes	Critically low
Chen 2010 (8)	Yes	No	No	Partial yes	No	No	No	Partial yes	Yes	Yes	N/A	N/A	No	No	N/A	Yes	Critically low
Cury Martins 2015 ⁽⁹⁾	Yes	Yes	No	Yes	Yes	Yes	Yes	Partial yes	Yes	No	Yes	Yes	Yes	No	Yes	Yes	Moderate
De Tiedra 1997 ⁽¹⁰⁾	No	No	No	No	No	No	No	Partial yes	No	No	N/A	N/A	No	No	N/A	No	Critically low
Devillers 2006 ⁽¹¹⁾	No	No	Yes	No	No	No	No	Partial yes	No	No	N/A	N/A	No	No	N/A	Yes	Critically low
Dong 2017 (12)	Yes	No	No	No	Yes	Yes	No	No	Yes	No	No	No	No	No	No	No	Critically low
Eichenfield 2014 ⁽¹³⁾	No	No	Yes	No	No	No	No	No	No	No	N/A	N/A	No	No	N/A	Yes	Critically low
Feldman 2005 ⁽¹⁴⁾	No	No	Yes	No	No	No	No	Partial yes	No	No	N/A	N/A	No	No	N/A	No	Critically low
Fishbein 2019 ⁽¹⁵⁾	Yes	Partial yes	Yes	Partial yes	Yes	Yes	No	Yes	No	Yes	No	No	No	No	No	Yes	Critically low
Frangos 2008 ⁽¹⁶⁾	No	No	No	No	No	No	No	No	No	No	N/A	N/A	No	No	N/A	No	Critically low
Froeschl 2007 ⁽¹⁷⁾	No	No	No	No	No	No	Yes	Partial yes	No	Yes	N/A	N/A	No	No	N/A	No	Critically low

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Gonzalez- Lopez 2017	Yes	No	No	Yes	Yes	Yes	No	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Critically low
Green 2004 (19)	Yes	Partial Yes	No	No	Partial yes	Yes	Partial yes	Yes	Partial yes	Yes	N/A	N/A	Yes	Yes	N/A	Yes	Low
Gu 2013 (20)	Yes	Yes	No	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	High
Gu 2014 ⁽²¹⁾	Yes	No	No	No	No	No	No	No	Yes	No	Yes	No	Yes	Yes	No	Yes	Critically low
Hajar 2015 (22)	Yes	Yes	Yes	No	Yes	Yes	No	No	Partial yes	No	N/A	N/A	No	No	N/A	Yes	Critically low
Hoare 2000 (23)	Yes	No	Yes	Yes	No	No	Yes	Partial yes	Partial yes	No	N/A	N/A	Yes	No	N/A	Yes	Low
Iskedjian 2004 ⁽²⁴⁾	Yes	No	No	Partial yes	Yes	Yes	No	Partial yes	Partial yes	No	N/A	N/A	No	No	N/A	No	Critically low
Juhasz 2017 ⁽²⁵⁾	No	No	No	No	No	No	No	No	No	N/A	N/A	N/A	No	No	N/A	Yes	Critically low
Labedz 2019 ⁽²⁶⁾	Yes	No	Yes	Yes	No	No	No	Partial yes	Yes	No	Yes	No	No	No	No	Yes	Critically low
Legendre 2015 ⁽²⁷⁾	Yes	No	No	No	Yes	Yes	No	Partial yes	Yes	No	Yes	No	No	Yes	Yes	Yes	Critically low
Li 2007 ⁽²⁸⁾	Yes	No	No	Partial yes	No	Yes	No	No	Partial yes	No	N/A	N/A	No	No	N/A	No	Critically low
Nankervis 2016 ⁽²⁹⁾	Yes	Partial yes	Yes	Partial yes	No	Yes	No	Partial yes	Partial yes	Yes	N/A	N/A	Yes	No	N/A	Yes	Low
Penaloza Hidalgo 2004 ⁽³⁰⁾	Yes	Partial yes	No	Yes	No	No	Yes	Partial yes	Partial yes	Yes	N/A	N/A	No	Yes	N/A	Yes	Low
Schmitt 2011 ⁽³¹⁾	Yes	Partial yes	No	Partial yes	Yes	Yes	No	Yes	Yes	No	N/A	N/A	No	Yes	N/A	Yes	Critically low
Sidbury 2011 ⁽³²⁾	No	No	Yes	No	No	No	No	No	No	No	N/A	N/A	No	No	N/A	Yes	Critically low
Siegfried 2016 ⁽³³⁾	No	No	No	No	No	No	No	No	No	No	N/A	N/A	No	No	N/A	Yes	Critically low
Singh 2012 (34)	No	No	No	No	No	No	Partial yes	No	Partial yes	No	N/A	N/A	Yes	No	N/A	Yes	Critically low
Svensson 2011 ⁽³⁵⁾	Yes	No	No	Partial yes	Yes	No	No	Partial yes	Partial yes	No	Yes	No	No	Yes	No	Yes	Critically low
Tang 2014 (36)	Yes	No	No	Partial yes	No	No	No	No	No	No	N/A	N/A	No	No	N/A	Yes	Critically low
van Zuuren 2017 ⁽³⁷⁾	Yes	Yes	No	Partial yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Moderate
Wood Heickman 2017 ⁽³⁸⁾	Yes	No	Yes	No	Yes	No	No	No	No	No	Yes	No	No	Yes	No	Yes	Critically low

	Critically low	No	No	No	No	No	Yes	Yes	Partial yes	No	No	No	No	Partial yes	No	No	Yes	Yan 2008 (39)
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Footnotes: AMSTAR 2 domains

1 Did the research questions and inclusion criteria for the review include the components of PICO?

*2 Did the report of the review contain an explicit statement that the review methods were established prior to the conduct of the review and did the report justify any significant deviations from the protocol?

3 Did the review authors explain their selection of the study designs for inclusion in the review?

*4 Did the review authors use a comprehensive literature search strategy?

5 Did the review authors perform study selection in duplicate?

6 Did the review authors perform data extraction in duplicate?

*7 Did the review authors provide a list of excluded studies and justify the exclusions?

8 Did the review authors describe the included studies in adequate detail?

*9 Did the review authors use a satisfactory technique for assessing the risk of bias (RoB) in individual studies that were included in the review?

10 Did the review authors report on the sources of funding for the studies included in the review?

*11 If meta-analysis was performed, did the review authors use appropriate methods for statistical combination of results?

12 If meta-analysis was performed, did the review authors assess the potential impact of RoB in individual studies on the results of the meta-analysis or other evidence synthesis?

*13 Did the review authors account for RoB in primary studies when interpreting/discussing the results of the review?

14 Did the review authors provide a satisfactory explanation for, and discussion of, any heterogeneity observed in the results of the review?

*15 If they performed quantitative synthesis did the review authors carry out an adequate investigation of publication bias (small study bias) and discuss its likely impact on the results of the review?

16 Did the review authors report any potential sources of conflict of interest, including any funding they received for conducting the review?

NB domains marked * in the table and footnotes are critical domains.

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Appendix 5: Characteristics of the included systematic reviews

First author and year of publication	Type of review	Language of publication	Funding source	Conflicts of interest	RCT comparisons	Observational data included	AMSTAR-2 rating
Ashcroft 2005	Non-Cochrane	English	NHS health technology assessment programme.	None known	TCS versus TCI	No	Critically low
Ashcroft 2007	Cochrane	English	School of Pharmacy and Pharmaceutical Sciences, University of Manchester, UK.	None known	TCS versus TCI	Yes – TCI compared with TCS	Moderate
Barnes 2015 (3)	Non-Cochrane	English	No funding	None known	TCS versus vehicle TCS versus TCI TCS versus TCS	Yes (single arm TCS studies)	Critically low
Braham 2010 (4)	Non-Cochrane	English	No funding	One author was a speaker for a number of pharmaceutical companies	Occlusion therapy versus non-occlusion therapy	Yes (occlusive therapy, no comparison)	Critically low
Broeders 2016 ⁽⁵⁾	Non-Cochrane	English	No funding	None known	TCS versus TCI	No	Critically low
Callen 2007 ⁽⁶⁾	Non-Cochrane	English	Funding from EBMed. One author received funding from Novartis Corporation for the project. They declared that "Novartis Corporation played no role in the design and conduct of the study or in data collection, data management, data analysis, interpretation of the data, manuscript preparation, manuscript review or manuscript approval"	Most authors had consultancy fees and/or research support from pharmaceutical companies	TCS versus vehicle TCS versus TCS	Yes (single arm TCS studies or comparing various TCS potencies)	Critically low
Chen 2010 ⁽⁷⁾	Non-Cochrane	English	Not stated	None known	TCS versus TCI	No	Critically low
Cury Martins 2015 ⁽⁸⁾	Cochrane	English	NIHR	None known	TCS versus TCI	Yes – TCI compared to TCS	Moderate
De Tiedra 1997 ⁽⁹⁾	Non-Cochrane	English	Supported by Laboratorios Novag, S.A, Grupo Ferrer.	Not clear	TCS versus TCS	Yes – in most cases they only report data from one arm of an RCT	Critically low
Devillers 2006 (10)	Non-Cochrane	English	Not stated	None known	Occlusive therapy versus non-occlusive therapy	Yes – occlusive therapy (no comparison)	Critically low
Dong 2017 (11)	Non-Cochrane	Chinese	Not stated	Not clear	TCS versus TCI	No	Critically low
Eichenfield 2014 ⁽¹²⁾	Non-Cochrane	English	No funding	Most authors served as consultants, speakers, members of the advisory	None	Yes (comparing different TCS potencies)	Critically low

				board and/or investigators for			
				pharmaceutical companies.			
Feldman 2005 (13)	Non-Cochrane	English	Grant from Galderma Laboratories, LP, Fort Worth, Texas.	Not clear	TCS versus vehicle	No	Critically low
Fishbein 2019	Non-Cochrane	English	No funding	None known	TCS versus vehicle/moisturizer	No	Critically low
Frangos 2008 (15)	Non-Cochrane	English	Not stated	One author is an investigator for Steifel and was an investigator on two of the studies reviewed.	TCS versus vehicle	Yes (single arm studies)	Critically low
Froschl 2007 (16)	GMS HTA report	German (executive summary in English)	Not stated	Not stated	TCS versus placebo/vehicle TCS versus TCS TCS versus TCI	No	Critically low
Gonzalez- Lopez 2017	Non-Cochrane	English	No funding	None known	Occlusive therapy versus non-occlusive therapy	No	Critically low
Green 2004 (18)	HTA report	English	Funded by the HTA Programme on behalf of NICE	None known	Once daily versus twice daily TCS use	No	Low
Gu 2013 ⁽¹⁹⁾	Cochrane	English	RMIT University Nottingham University, UK. NIHR	One author was a principal investigator on one included study (but this study was not relevant for this overview)	Chinese herbal medicine versus TCS	No	High
Gu 2014 ⁽²⁰⁾	Non-Cochrane	English	Not stated	None known	Chinese herbal medicine versus TCS	No	Critically low
Hajar 2015 ⁽²¹⁾	Non-Cochrane	English	No funding	None known	No RCTs found	Yes (case series or case reports on steroid withdrawal)	Critically low
Hoare 2000 (22)	NIHR HTA report	English	HTA programme	One author received payment from Novartis for lectures on the epidemiology of atopic eczema in 1999. Another author has acted as occasional lecturer or consultant for pharmaceutical companies.	TCS versus TCS TCS versus vehicle	No	Low
Iskedjian 2004 (23)	Non-Cochrane	English	Funded by Fujisawa Canada Inc.	Not clear	TCS versus TCI TCS versus placebo	No	Critically low
Juhasz 2017 (24)	Non-Cochrane	English	Not stated	One author had primary contact with the 2nd case and has a blog on the subject matter in this systematic review	No RCTs found	Yes (case reports on steroid withdrawal)	Critically low

Labedz 2019	Non-Cochrane	English	Not stated	None known	TCS versus TCI	No	Critically low
(25)							
Legendre 2015 ⁽²⁶⁾	Non-Cochrane	English	No funding	One author is a consultant and investigator for two pharmaceutical companies. One author is a speaker and/or on the advisory board for five pharmaceutical companies.	Only searched for cohort or case control studies	Yes (comparing TCS and TCI)	Critically low
Li 2007 ⁽²⁷⁾	Non-Cochrane	Chinese	Not stated	Not stated	TCS versus TCI	No	Critically low
Nankervis 2016 ⁽²⁸⁾	NIHR HTA report	English	NIHR	One author reports grants and fees from a number of pharmaceutical companies.	TCS versus placebo/vehicle Proactive treatment versus vehicle TCS versus TCI TCS versus TCS Once a day versus twice a day use of TCS Occlusive therapy versus non-occlusive therapy TCS versus emollients	Νο	Low
Penzaloza Hidalgo 2004 ⁽²⁹⁾	West Midlands HTA report	English	Not stated	None known	TCS versus TCI	No	Low
Schmitt 2011 (30)	Non-Cochrane	English	No funding	One author has served as paid lecturer for a pharmaceutical company.	Proactive treatment versus vehicle	No	Critically low
Sidbury 2014 (31)	Non-Cochrane	English	Not stated	Some authors have served as investigators, consultants, speakers, and on advisory boards for pharmaceutical companies.	Proactive treatment versus vehicle	No	Critically low
Siegfried 2016 (32)	Non-Cochrane	English	Financial support for writing by Valent Pharmaceutical North America LLC. They declared that "Valeant Pharmaceuticals had no role in the design of the literature searches, or analysis and presentation of results."	Authors have either participated in paid contract research, received travel expenses for presentations, consulting fees, speakers, on advisory boards, or on data safety monitoring boards with pharmaceutical companies.	TCS versus TCS TCS versus TCI TCS versus vehicle	Νο	Critically low
Singh 2012 (33)	Non-Cochrane	English	Not stated	None known	TCS versus TCS TCI versus TCS	Yes (single arm TCS study)	Critically low

TCS versus placebo/vehicle

Svensson 2011 ⁽³⁴⁾	Non-Cochrane	English	Funded by Astellas Pharma Europe Ltd.	One author was a paid employee of Astellas Pharma Europe Ltd and one author undertook paid consultancy work for Astellas Pharma Europe Ltd.	TCI versus TCS	No	Critically low
Tang 2014 (35)	Non-Cochrane	English	Not stated	One author has received lecture fees from Astellas.	Proactive treatment versus vehicle	No	Critically low
van Zuuren 2017 ⁽³⁶⁾	Cochrane	English	Oak Foundation, Denmark NIHR	None known	TCS versus emollient	No	Moderate
Wood Heickman 2018 ⁽³⁷⁾	Non-Cochrane	English	No grants, honoraria or royalties were received supporting the writing of the paper.	One author was a consultant with Perrigo, Inc. with regard to topical corticosteroid treatment. All authors have no financial or other potential conflicts of interest.	Two RCTs included but analysed as observational data	Yes – single arm cohort studies	Critically low
Yan 2008 (38)	Non-Cochrane	English	Not stated	Not stated	TCI versus TCS	No	Critically low

Key: TCI=topical calcinueirin inhibitor; TCS=topical corticosteroids; RCT=Randomised Controlled Trial; NIHR= National Institute for Health Research

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	Α	ppendix 6 – Charact	eristics and sa	afety data from the i	ncluded studies	
		How safe are top	ical corticosteroio	ls compared to emollient o	r vehicle?	
Study ID (Systematic review*)	Study design and study duration (Quality assessment)	Intervention and comparator	Participants	Cutaneous adverse events	Systemic adverse events	Unspecified adverse events
			Very potent top	ical corticosteroids		
Breneman 2003 (1) (unpublished) (Feldman 2005 (²⁾ Nankervis ⁽³⁾)	RCT 2 weeks treatment, then followed up for additional 2 weeks Cochrane risk of bias tool: randomisation described, allocation concealment unclear, intention-to- treat unclear.	Intervention: Clobetasol propionate 0.05% lotion (twice a day) (n=96) Intervention: Clobetasol propionate 0.05% emollient cream (twice a day) (n=100) Comparator: Vehicle (n=33)	Severity: moderate to severe Age: ≥ 12 years Sample size: 229 participants	<u>Local application site skin</u> <u>reactions</u> No clinically significant telangiectasia or skin thinning		Unspecified adverse eventsIncidence comparable betweengroups.Treatment-related adverseeventsClobetasol lotion = 4/96 patients(4.2%); Clobetasol cream = 1/100patients (1%)Vehicle = 6/33 patients (18.2%)(Difference between groups: p=0.0006°)
Kimball 2008 ⁽⁴⁾ (trial a) (<i>Frangos 2008</i> (⁵⁾)	RCT Duration not specified in review Risk of bias not assessed in any of the included systematic reviews.	Intervention: Clobetasol propionate emulsion formulation foam 0.05% Comparator: Vehicle	Severity: not specified in the review Age: not specified in the review Sample size: not specified in the review			Incidence of adverse events or treatment related adverse events Clobetasol foam = 8% Vehicle foam = 10% (no significant differences between groups)
Rosso 2009 ⁽⁶⁾ (Barnes 2009 ⁽⁷⁾)	RCT 2 weeks treatment Risk of bias not assessed in any of the included systematic reviews.	Intervention: Fluocinonide 0.1% cream (n=109) Comparator: Vehicle (n=50)	Severity: not specified in the review Age: not specified in the review Sample size: 159 participants	Skin thinning Fluocinonide: 6/109 participants (5.6%) Vehicle: 2/50 participants (4.3%) (Difference between groups: p=0.69°)		
www.olux- e.com (online data) ⁽⁸⁾ (Frangos 2008 (⁵⁾)	Single arm study (observational) 2 weeks treatment	Intervention: Clobetasol propionate emollient foam (twice daily) (n=37) Comparator: No comparator	Severity: ≥30% BSA Age: ≥12 years old Sample size: 37 participants		HPA axis suppression 6/37 patients (16%) (not specified in the review how it was measured)	

Kimball 2008 ⁽⁴⁾ (trial b) (Frangos 2008 ⁽⁵⁾ ; Wood Heickman 2018 ⁽⁹⁾)	Risk of bias not assessed in any of the included systematic reviews. Open label Phase II safety study 2 weeks treatment Risk of bias not assessed in any of the included systematic reviews.	Intervention: Clobetasol propionate emollient foam 0.05% (twice daily) (n=52) Comparator: No comparator	Severity: mild to severe Age: children (from 6 years old) and adults Sample size: 52 participants		 HPA axis suppression 7/30 (23.3%) had adrenal insufficiency (ACTH stimulation testing, measuring serum cortisol levels). 47% of children (aged 6-11) 0% of adolescents (aged 12- 17) 27% of adults (≥18 years) Was reported as transient and reversible. After TCS discontinuation, children with biochemical adrenal insufficiency had complete resolution at retesting. 	
Herz 1991 ⁽¹⁰⁾ (Barnes 2015 ⁽⁷⁾)	Single arm study (observational) (2 weeks treatment) Risk of bias not assessed in any of the included systematic reviews.	Intervention: Clobetasol propionate (n=59) Comparator: No comparator	Severity: not specified in the review Age: not specified in the review Sample size: 59 participants	Skin thinning 1 case of skin thinning reported (not clear if in a psoriasis or eczema patient – but assume its eczema as this is the topic of the systematic review).		
			Potent topica	l corticosteroids		L
Sugarman 2009 (11) (Van Zuuren 2017 ⁽¹²⁾)	RCT (4 weeks treatment) (Cochrane risk of bias tool: low risk of selection, attrition and other biases. Unclear risk of reporting and performance bias, High risk of detection bias. ⁽¹²⁾) (Cochrane risk of bias tool: unclear risk of selection bias, high risk from no blinding. ⁽³⁾)	Intervention: Fluticasone 0.05% cream twice daily (hydrocortisone 2.5% for the face and body folds) (n=62) Comparator: Ceramide- dominant barrier repair formulation (EpiCeram) twice daily (emollient) (n=59)	Severity: moderate to severe Age: children 6 months to 18 years (mean age 7.1 years) Sample size: 121 participants			Serious adverse events The participants did not report any in either group. No further details regarding other possible treatment related adverse events were reported.

Griffiths 2002 ⁽¹³⁾ (Nankervis 2017 ⁽³⁾)	RCT (up to 14 days treatment) (Cochrane risk of bias tool: low risk of selection bias from sequence generation, unclear risk of selection bias from allocation concealment, low risk from blinding. ⁽³⁾)	Intervention: Hydrocortisone 17-butyrate cream (0.1%) maximum application of 2g (four fingertip units) per day (n=49) Comparator: Cipamfylline cream (1.5 mg of cipamfylline per gram of cream) used up to a maximum of 2 g (four fingertip units) of cream per day (emollient) (n=54)	Severity: not specified Age: adults ≥18 years old Sample size: 103 participants	No difference in cutaneous adverse events which were possibly or probably related to treatment in either group (p = 0.13) The adverse events were mostly application site reactions, including itching, stinging or burning, and drug reactions.		Unspecified adverse events Hydrocortisone group: 20/49 (40.8%) participants reported 41 adverse events in total. Emollient: 29/52 (55.8%) participants reported 63 adverse events in total. (Difference between groups: p= 0.14°)
Eichenfield 2006 (14) (Nankervis 2017 (³))	RCT (4 weeks treatment) Risk of bias not assessed	Intervention: Fluticasone propionate four times daily (n=221) Comparator: Vehicle four times daily (n=217)	Severity: moderate to severe Age: children from 3 months old to 16 years old Sample size: 438 children			Withdrawal due to adverseeventsTopical corticosteroids: 4participants in total from thisstudy and from Hebert 2007The number of participantsreporting at least 1 adverseeventFluticasone: 77/221 (34.8%)participantsVehicle: 82/217 (37.8%)participants(Difference between groups:p=0.52a)
Wu 2013 ⁽¹⁵⁾ (Nankervis 2017 ⁽³⁾ , Fishbein 2019 ⁽¹⁶⁾)	RCT (10 days treatment) (Cochrane risk of bias tool: low risk of selection bias from sequence generation. Unclear risk of selection bias from allocation concealment, unclear risk from blinding and other biases: Two out of 60 participants were excluded from the analyses as they used concomitant medication ⁽³⁾	Intervention: Mometasone furoate 0.1% cream, twice a day (n=20) Comparator: placebo of distilled water in 1% dimethyl sulfoxide mixed with the identical cream base as used for the 15(R/S)- methyl-lipoxin A4 (n=20) Comparator: 15(R/S)-methyl- lipoxin A4 0.1% cream (n=20)	Severity: all severities Age: children from 1 month to 1 year old Sample size: 60 participants		None of the safety tests (e.g. full blood count, kidney and liver function test, and electrocardiogram) showed any significant differences compared with baseline for all three treatment groups.	No clinical adverse events were reported.

Pellanda 2005 (17) (Nankervis 2017 (3)	RCT (Duration not specified in the review) Risk of bias not assessed	Intervention: Triamcinolone acetonide Comparator: Vehicle	Severity: mild to moderate Age: not specified in the review Sample size: not specified in the review	Skin changes One report by a participant using placebo (no further details)		
Lebwohl 1996 ⁽¹⁸⁾ (Hoare 2000 ⁽¹⁹⁾)	RCT (29 days treatment) (Moher 1995 quality checklist: method and concealment of randomisation unclear, double blinded, large number of withdrawals and dropouts, no ITT analysis ⁽¹⁹⁾)	Intervention: Fluticasone propionate ointment 0.005% Comparator: Vehicle	Severity: not specified in the review Age: not specified in the review Sample size: 203 participants			The review authors only reported that "Drug related adverse effects were rare"
Lebwohl 1999 ⁽²⁰⁾ (Hoare 2000 ⁽¹⁹⁾)	RCT (29 days treatment) (Moher 1995 quality checklist: method and concealment of randomisation unclear, double blinded, large number of withdrawals and dropouts, no ITT analysis ⁽¹⁹⁾)	Intervention: Fluticasone propionate ointment 0.005% Comparator: Vehicle	Severity: not specified in the review Age: not specified in the review Sample size: 169 participants			The review authors only reported that "Drug related adverse effects were rare"
Abramovitis 2010 ⁽²¹⁾ (Wood Heickman 2018 ⁽⁹⁾ , Fishbein 2019 ⁽¹⁶⁾)	RCT (21 to 29 days treatment) Risk of bias not assessed in any of the included systematic reviews.	Intervention: Hydrocortisone butyrate 0.1% cream, twice daily (n=131) Comparator: Lipocream vehicle, twice daily (n=133)	Severity: Mild to moderate Age: children 3 months to 18 years (mean 7.2 years) Sample size: 264 children		HPA axis suppression (no data for vehicle group) 5/63 (7.9%) children in the hydrocortisone group (measured using ACTH stimulation testing, measuring serum cortisol levels) After TCS discontinuation, children with biochemical adrenal insufficiency had complete resolution at retesting.	The number of participantsreporting at least 1 adverseeventHydrocortisone: 29/131 (22.1%)participantsVehicle: 28/133 (21.1%)participants(Difference between groups: $p=0.83^{a}$)
Matheson 2008 (22)	RCT (28 days treatment)	Intervention: Hydrocortisone butyrate 0.1% lotion, twice daily (n=139)	Severity: Mild to moderate			The number of participants reporting at least 1 adverse event Hydrocortisone: 48/139 (34.5%)

(Fishbein 2019 ⁽¹⁶⁾)	Risk of bias not assessed in any of the included systematic reviews.	Comparator : Vehicle, twice daily (n=145)	Age: children 3 months to 18 years Sample size: 284 children			participants Vehicle: 56/145 (38.6%) participants (Difference between groups: p=0.48°)
Friedlander 2002 ⁽²³⁾ (Callen 2007 ⁽²⁴⁾ ; Wood Heickman 2018 ⁽⁹⁾)	Single arm study (observational) (3 to 4 weeks) Risk of bias not assessed in any of the included systematic reviews.	Intervention: Fluticasone propionate cream 0.05% (n=43) Comparator: No comparator	Severity: not specified in the review Age: children 3 months to 6 years Sample size: 43 participants		HPA axis suppression 2/43 (4.7%) children (measured using ACTH stimulation testing, measuring serum cortisol levels) After TCS discontinuation, children with biochemical adrenal insufficiency had complete resolution at retesting.	
Eichenfield 2007 (25) (Wood Heickman 2018 (9)	Single arm study (observational) (4 weeks treatment) Risk of bias not assessed in any of the included systematic reviews.	Intervention: Hydrocortisone butyrate 0.1% (n=20) Comparator: No comparator	Severity: not specified in the review Age: children (median or mean = 9 years) Sample size: 20 children		HPA axis suppression 0/20 (0%) children (measured using ACTH stimulation testing, measuring serum cortisol levels)	
Hebert 2006 ⁽²⁶⁾ (Wood Heickman 2018 ⁽⁹⁾)	Single arm study (observational) (3 to 4 weeks treatment) <i>Risk of bias not assessed</i> <i>in any of the included</i> <i>systematic reviews.</i>	Intervention: Fluticasone propionate 0.05% lotion (n=42) Comparator: No comparator	Severity: not specified in the review Age: children (median or mean 2.6 years) Sample size: 42 children		HPA axis suppression 0/42 (0%) children (measured using ACTH stimulation testing, measuring serum cortisol levels)	
			Moderate potency	topical corticosteroids		
De Belilovsky 2011 ⁽²⁷⁾ (Van Zuuren 2017 ⁽¹²⁾)	RCT (3 weeks treatment) (Cochrane risk of bias tool: low risk of selection, attrition, reporting and other biases. Unclear risk of performance bias.	Intervention: Hydrocortisone butyric propionate 0.1% twice daily (n=40) Comparator: Stelatopia (2% sunflower oil, fatty acids, ceramides) twice daily (n=40)	Severity: mild to moderate Age: children 4 months to 4 years (mean age 2.3 years) Sample size: 80 participants			No participants reported adverse events

	High risk of detection bias. ⁽¹²⁾)				
	(Cochrane risk of bias tool: unclear risk of selection bias and risk from blinding. ⁽³⁾)				
Rosenthal 1980 (²⁸⁾ (Singh 2012 ⁽²⁹⁾)	RCT (14 days treatment) (Delphi list: method of randomisation not described, allocation not concealed, blinded, no ITT analysis ⁽²⁹⁾)	Intervention: Clocortolone pivalate 0.1% cream (applied thrice daily) Comparator: Vehicle (applied thrice daily)	Severity: not specified in the review Age: not specified in the review Sample size: 100 participants		No adverse events
Binder 1977 (30)	RCT	Intervention: Clocortolone	Severity: not	Irritation and dryness	
(Singh 2012 ⁽²⁹⁾)	(14 days treatment)	pivalate (applied thrice daily) (n=17)	specified in the review	Clinically significant in one patient in each group – did not	
	(Delphi list: method of randomisation not described, allocation not concealed, blinded, no ITT analysis ⁽²⁹⁾)	Comparator: Vehicle (n=12)	Age: mean age 30 years Sample size: 29 participants	result in discontinuation.	
Rauschkol 1981 (³¹⁾ (<i>Fishbein 2019</i> (¹⁶⁾)	Within-participant RCT (14 days treatment) Risk of bias not assessed in any of the included systematic reviews.	Intervention: Halcinonide 0.025% cream, twice daily, on one arm Comparator: Placebo cream unspecified, twice daily on the other arm at the same time	Severity: not reported Age: children 7 months to 15 years (mean age 8 years) Sample size: 86 children		The number of participants reporting at least 1 adverse event Halcinonide: 4/86 (4.7%) participants Placebo: 5/86 (5.8%) participants
Nolting 1991 (32)	RCT (but safety data only	Intervention: Prednicarbate	Severity: Disease		Adverse reactions
(De Tiedra 1997 ⁽³³⁾)	presented for one arm) (21 days treatment)	cream 0.25% (2 applications per day) (n=34)	duration = mean 4.1 years ± 2.7		2/34 patients (5.9%)
	Risk of bias not assessed in any of the included systematic reviews.	Comparator: mometasone cream 0.1% twice daily (no safety data given)	Age: children 2-12 years (mean 6.6 ± 3.6).		
			Sample size: 34 participants (with safety data)		
Rampini 1992 ⁽³⁴⁾	RCT (but safety data only presented for one arm)	Intervention: Prednicarbate cream/unguent 0.25% (2 applications per day) (n=93)	Severity: not specified in the review		Adverse reactions 3/93 patients (3.2%)

(De Tiedra 1997 ⁽³³⁾)	(21 days treatment) (Moher 1995 quality checklist: method and concealment of randomisation unclear, double blinded, two dropouts, no ITT analysis, ⁽¹⁹⁾)	Comparator:methylprednisolone aceponate 0.1% once daily (no safety data given)	Age: children 0.3 to 14 years (mean 6.6). Sample size: 93 participants (with safety data)		
Camacho 1996 (³⁵⁾ (<i>De Tiedra 1997</i> (³³⁾)	RCT (but safety data only presented for one arm) (21 days treatment) (Moher 1995 quality checklist: method and concealment of randomisation unclear, double blinded, no ITT analysis, 14/49 dropouts, ⁽¹⁹⁾	Intervention: Prednicarbate cream 0.25% (2 applications per day) (n=49) Comparator: fluocortolone pivalate cream 0.2% (no safety data given)	Severity: Disease duration= mean 6.2 years ± 8.2 (range 0.25 to 39 years). Age: adults 19 to 65 years (mean 34.1 ± 12). Sample size: 49 participants (with safety data)		Adverse reactions 4/49 patients (8.1%)
Gimenez Camarasa 1994 (36) (De Tiedra 1997 (33)	RCT (but safety data only presented for one arm) (21 days treatment) Risk of bias not assessed in any of the included systematic reviews.	Intervention: Prednicarbate cream 0.25% (2 applications per day) (n=41) Comparator: fluocinolone cream 0.025% twice daily (no safety data given)	Severity: Disease duration = mean 6.4 years ± 8.6 (range 0- 40). Age: adults 18 to 77 years (mean 37.6 ± 15.9). Sample size: 41 participants (with safety data)		Adverse reactions 0/41 patients (0%)
Moshang 2001 ⁽³⁷⁾ (Callen 2007 ⁽²⁴⁾ ; Wood Heickman 2018 ⁽⁹⁾)	Single arm study (observational) (3 weeks treatment) Risk of bias not assessed in any of the included systematic reviews.	Intervention: Prednicarbate emollient cream 0.1%, twice daily (n=55) Comparator: No comparator	Severity: not specified in the review Age: children 4 months to 12 years Sample size: 55 participants	HPA axis suppression All normal (measured using ACTH stimulation testing, measuring serum cortisol levels)	
Conde 2008 ⁽³⁸⁾ (Singh 2012 ⁽²⁹⁾)	Single arm study (observational) (4 weeks treatment)	Intervention: Clocortolone pivalate cream 0.1% twice daily (n=10) Comparator: No comparator	Severity: mild to moderate Age: children, mean age 7.9 years		No adverse events reported

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Crespi 1986 ⁽³⁹⁾ (Callen 2007 ⁽²⁴⁾)	Risk of bias not assessed in any of the included systematic reviews. Single arm study (observational) (4 weeks treatment) Risk of bias not assessed in any of the included systematic reviews.	Intervention: Alclometasone cream, twice daily (n=39) Comparator: No comparator	Sample size: 10 participants Severity: not specified in the review Age: children Sample size: 39 participants		HPA axis suppression All normal (measured via morning cortisol)	
			Mild potency top	pical corticosteroids		
Udompataikul 2011 ⁽⁴⁰⁾ (Van Zuuren 2017 ⁽¹²⁾)	Within-participant RCT (6 weeks treatment) (Cochrane risk of bias tool: unclear risk of selection, performance and attrition bias. Low risk of reporting and other biases. High risk of detection bias. ⁽¹²⁾ (Cochrane risk of bias tool: unclear risk of selection bias from sequence generation and risk from blinding. Low risk of selection bias from allocation concealment, ⁽³⁾)	Intervention: Hydrocortisone acetate 1% cream twice daily, was applied one side of the body for 4 weeks followed by the cream base for 2 weeks. Comparator: Licochalcone (containing <i>Glycyrrhiza inflata</i> root extract, decanediol, menthoxypropanediol and 6- fatty acids) applied twice daily on one side of the body for 6 weeks	Severity: mild to moderate Age: children 2 months to 10 years (mean age 5.8 years) Sample size: 30 participants			No adverse events on either side during the study.
Hebert 2007 ⁽⁴¹⁾ (Nankervis 2017 ⁽³⁾ , Fishbein 2019 ⁽¹⁶⁾)	RCT (28 days) Risk of bias not assessed in any of the included systematic reviews.	Intervention: Desonide 0.05% gel twice daily (n=425) Comparator: Hydrogel vehicle twice daily (n=157)	Severity: mild to moderate Age: children 3 months to 18 years Sample size: 582 children			Serious adverse events One event reported in TCS group but not thought to be related to treatment Withdrawal due to adverse events TCS group: 4 in total from this study and from Eichenfield 2006 The number of participants reporting at least 1 adverse event Desonide: 85/425 (20 %) participants Vehicle: 46/157 (29.3%)

						participants (Difference between groups: p=0.02°)
Udompataikul 2012 ⁽⁴²⁾ (Fishbein 2019 (¹⁶⁾)	Within-participant RCT (4 weeks treatment) Risk of bias not assessed in any of the included systematic reviews.	Intervention: Hydrocortisone 1% ointment twice daily, applied to one arm. Comparator: 5% dexapanthenol ointment twice daily, applied to the other arm at the same time.	Severity: mild to moderate Age: children 2 years to 15 years (mean age 7.2 years) Sample size: 30 participants			No adverse events on either side during the study.
Wananukul 2013 (⁴³⁾ (Van Zuuren 2017 ⁽¹²⁾)	Within-participant RCT (4 weeks treatment) (Cochrane risk of bias tool: low risk of selection bias from sequence generation, performance, detection, attrition, reporting and other biases. Unclear risk of selection bias from allocation concealment. (12)	Intervention: Hydrocortisone acetate 1% cream twice daily on one side of the body Comparator: Licochalcone (containing <i>Glycyrrhiza inflata</i> root extract, decanediol, menthoxypropanediol and 6- fatty acids) twice daily on one side of the body	Severity: mild to moderate Age: children, mean age 3.1 years Sample size: 55 participants			No adverse events on either side during the study
Jirabundansuk 2014 ⁽⁴⁴⁾ (Van Zuuren 2017 ⁽¹²⁾)	Within-participant RCT (4 weeks treatment) (Cochrane risk of bias tool: Unclear risk of selection and performance bias. High risk of detection bias. Low risk of attrition, reporting and other biases. ⁽¹²⁾)	Intervention: Hydrocortisone acetate 1% cream twice daily on one side of the body Comparator: Moisturiser containing spent grain, Vitellaria paradoxa (formerly Butyrospermum parkii) extract plus Argania spinosa kernel oil twice daily on one side of the body	Severity: Mild or moderate Age: children 2-15 years (mean age 4.3 years) Sample size: 31 participants			The investigators stated that "no specific adverse events were reported".
Dolle 2010 ⁽⁴⁵⁾ (Nankervis 2017 ⁽³⁾)	Within-participant RCT (3 weeks treatment) (Cochrane risk of bias tool: unclear risk of selection bias and risk from blinding. ⁽³⁾)	Intervention: 1% hydrocortisone solution once daily for 1 st week then twice daily up to 3 weeks Comparator: 6% miltefosine solution once daily for 1 st week then twice daily up to 3 weeks	Severity: moderate to severe Age: adults (≥18 years old) Sample size: 16 participants	Local topical adverse events related to the treatment Hydrocortisone: 7/16 participants (44%) Emollient: 10/16 participants (63%) These adverse events included pruritus, burning, tingling and dry	No systemic adverse events	No withdrawals because of adverse events

				skin. Dry skin was seen only with emollient treatment.		
Patzelt- Wenczler 2000 (46) (Nankervis 2017 (3)) Paller 2003 ⁽⁴⁷⁾ (Nankervis 2017 (3))	Within-participant RCT (2 weeks treatment) (Cochrane risk of bias tool: unclear risk of selection bias and high risk from no blinding. ⁽³⁾) RCT (2 weeks treatment) Risk of bias not assessed in any of the included systematic reviews.	Intervention: Hydrocortisone 0.5% twice daily Comparator: Kamillosan® cream, containing 2% ethanolic extract of chamomile flowers, twice daily (emollient) Comparator: Vehicle cream applied twice daily Intervention: Fluocinolone acetonide 0.01% twice daily (n=45) Comparator: Vehicle twice daily (n=49)	Severity: at least moderate Age: not specified in the review Sample size: 72 participants Severity: not specified in the review Age: children from 2 to 12 years old Sample size: 94 participants	Mild hypopigmentation Two participants out of 45 reported this event with fluocinolone (4.4%)		Three participants in the emollient group withdrew early because of intolerability.
Patel 1995 ⁽⁴⁸⁾ (Callen 2007 ⁽²⁴⁾)	Single arm study (observational) (3-10 years follow up) Risk of bias not assessed in any of the included systematic reviews.	Intervention: 1% Hydrocortisone ointment (n=14; 9/14 intermittently used moderate to high potency) Comparator: No comparator	Severity: not specified in the review Age: children 3.1 to 10.7 years Sample size: 14 participants		HPA axis suppression Plasma cortisol levels - no change in basal/peak levels but peaked earlier	
Dohil 2009 ⁽⁴⁹⁾ (Wood Heickman 2018 ⁽⁹⁾)	Single arm study (observational) (4 weeks duration) Risk of bias not assessed in any of the included systematic reviews.	Intervention: fluocinolone acetonide 0.01% Comparator: No comparator	Severity: not specified in the review Age: children (median or mean age 1.1 years) Sample size: 24 participants		HPA axis suppression No cases of adrenal insufficiency (measured using ACTH stimulation testing, measuring serum cortisol levels)	
Eichenfield 2007 (⁵⁰⁾ (Wood Heickman 2018 (⁹⁾)	Single arm study (observational) (4 weeks duration) Risk of bias not assessed in any of the included systematic reviews.	Intervention: Desonide hydrogel 0.05% Comparator: No comparator	Severity: not specified in the review Age: children (median or mean age 3.3 years)		HPA axis suppression No cases of adrenal insufficiency (measured using ACTH stimulation testing, measuring serum cortisol levels)	

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Hebert 2008 ⁽⁵¹⁾ (Wood Heickman 2018 ⁽⁹⁾)	Single arm study (observational) (4 weeks duration) Risk of bias not assessed in any of the included systematic reviews.	Intervention: Desonide 0.05% foam Comparator: No comparator	Sample size: 34 participants Severity: not specified in the review Age: children (median or mean age 6.7 years) Sample size: 75 participants		HPA axis suppression Three out of 75 participants had adrenal insufficiency (measured using ACTH stimulation testing, measuring serum cortisol levels)	
		How safe are topical	corticosteroids co	mpared to topical calcineur	rin inhibitors?	
Study ID (Systematic review*)	Study design and study duration (Quality assessment)	Intervention and comparator	Participants	Cutaneous adverse events	Systemic adverse events	Unspecified adverse events
			Potent topica	l corticosteroids	·	
Bieber 2007 ⁽⁵²⁾ (Broeders 2016 ⁽⁵³⁾)	RCT (up to 3 weeks treatment) (Jadad score 4/5 – risk from sequence generation and allocation concealment ⁽⁵³⁾) (Cochrane risk of bias tool: unclear risk of selection bias and from blinding. ⁽³⁾) (Cochrane risk of bias tool: unclear risk of selection bias. Low risk of performance, detection, attrition, reporting and other biases. ⁽⁵⁴⁾)	Intervention: Methyl- prednisolone 0.1% (n=129) once daily in the evening to all affected body surface areas for a minimum of 2 weeks and a maximum of 3 weeks and cleared areas treated for an additional 7 days post clearance. Also applied a vehicle ointment in the morning to maintain blinding. Comparator: Tacrolimus 0.03% (n=136), applied twice daily, morning and evening, to all affected body surface areas for a minimum of 2 weeks and cleared areas treated for an additional 7 days post clearance.	Severity: severe to very severe Age: children 2 to 15 years old Sample size: 265 participants	Adverse events related to treatment Methyl-prednisolone: 0/129 participants (0%) Tacrolimus: 6/136 participants (4.4%) (Difference between groups: p=0.09 ^{a,b})		Severe adverse eventsMethyl-prednisolone: $0/129$ participants (0%)Tacrolimus: $6/136$ participants (4.4%) (Difference between groups: $p=0.09^{a,c}$)Adverse events requiringdiscontinuationMethyl-prednisolone: $0/129$ (0%) Tacrolimus: $4/136$ (3%)(Difference between groups: $p=0.15^{a}$)
Doss 2010 ⁽⁵⁵⁾ (Broeders 2016 ⁽⁵³⁾)	RCT (3 weeks treatment twice daily, plus 3 weeks follow up with once daily treatment)	Intervention: Fluticasone 0.005% ointment applied twice daily to all affected areas except eyelids until clearance, up to 3 weeks. All participants who responded to treatment could apply treatment once a day to	Severity: moderate to severe Age: children 2 to 15 years old	Adverse events related to treatment Fluticasone: 45/239 participants (19%) Tacrolimus: 55/239 participants (23%)		Severe adverse events Fluticasone: 2/239 participants (0.8%) Tacrolimus: 1/239 participants (0.4%) (Difference between groups: p=0.57°)

	(Jadad score 5/5 – risk from allocation concealment ⁽⁵³⁾) (Cochrane risk of bias tool: unclear risk of selection bias and from blinding. ⁽³⁾) (Cochrane risk of bias tool: Low risk of selection, performance, detection, attrition, reporting and other biases. ⁽⁵⁴⁾) Cochrane risk of bias tool: low risk of selection, performance, attrition, reporting and other biases. Unclear risk of performance bias ⁽⁵⁶⁾).	the remaining lesions for another 3 weeks (n=239) Comparator: Tacrolimus 0.03% ointment applied twice daily to all affected areas except eyelids until clearance, up to 3 weeks. All participants who responded to treatment could apply treatment once a day to the remaining lesions for another 3 weeks (n=239)	Sample size: 478 participants	(Difference between groups: $p=0.26^{a}$) Skin burning Fluticasone: 6/239 (2.5%) Tacrolimus: 18/237 (7.6%) (Difference between groups: $p=0.02^{a}$) Pruritus Fluticasone: 8/239 participants (3.3%) Tacrolimus: 10/237 participants (4.2%) (Difference between groups: $p=0.62^{a}$) Skin infection Fluticasone: 49/239 participants (21%) Tacrolimus: 44/239 participants (18%) (Difference between groups: $p=0.56^{a}$)	Adverse events requiring discontinuation Fluticasone: 6/239 participants (2.5%) Tacrolimus: 4/239 participants (1.7%) (Difference between groups: p=0.53°)
Doss 2009 ⁽⁵⁷⁾ (Broeders 2016 ⁽⁵³⁾)	RCT (3 weeks of treatment – then for a further 3 weeks either stop treatment, once daily treatment or switch to other treatment twice daily) (Jadad score 5/5 – risk from allocation concealment ⁽⁵³⁾) (Cochrane risk of bias tool: unclear risk of selection bias and low risk from blinding. ⁽³⁾)	Intervention: Fluticasone 0.005% ointment twice daily on facial eczema lesions for 3 weeks or until clearance (n=279) Comparator: Tacrolimus 0.1% twice daily on facial eczema lesions for 3 weeks or until clearance (n=287) For 21 days after the initial 3 weeks, the participants could stop treatments if the facial lesions had cleared; stay on the same treatment once a day; or swap treatment using it twice daily (still blinded)	Severity: moderate to severe Age: adults Sample size: 566 participants	Adverse events related to treatmentFluticasone: 42/279 participants (15%)Tacrolimus: 75/287 participants (26%) (Difference between groups: $p=0.001^{\circ}$)Skin burning Fluticasone: 9/279 participants (3.2%)Tacrolimus: 47/287 participants (16.4%) (Difference between groups: $p<0.00001^{\circ}$)Pruritus Fluticasone: 9/279 participants (3.2%)Tacrolimus: 12/287 participants (3.2%)Tacrolimus: 12/287 participants (3.2%)(3.2%)Tacrolimus: 12/287 participants (3.2%)(3.2%)Tacrolimus: 12/287 participants (4.2%) (Difference between groups: $p=0.55^{\circ}$)	Severe adverse events Fluticasone: 0/279 participants (0%) Tacrolimus: 1/287 participants (0.3%) (Difference between groups: p=0.51°). Adverse events requiring discontinuation Fluticasone: 8/279 participants (2.9%) Tacrolimus: 7/287 participants (2.4%) (Difference between groups: p=0.75°)

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Luger 2001 ⁽⁵⁸⁾ (Broeders 2016 ⁽⁵³⁾)	RCT (up to 3 weeks treatment) (Jadad score 3/5 – risk from sequence generation and allocation concealment ⁽⁵³⁾) (Cochrane risk of bias tool: unclear risk of selection bias and unclear risk from blinding. ⁽³⁾) (Jadad scale: 3/5 ⁽⁵⁹⁾)	Intervention: Betamethasone valerate 0.1% applied twice daily on all affected areas except for the head and neck for up to 3 weeks or until complete clearance if this was sooner (n=42) Comparator: Pimecrolimus 1% applied twice daily on all affected areas except for the head and neck for up to 3 weeks or until complete clearance if this was sooner (n=45)	Severity: moderate Age: adults ≥ 18 years old Sample size: 87 participants	PruritusBetamethasone: $5/42$ participants (12%)Pimecrolimus: $14/45$ participants (31%) (Difference between groups: $p=0.04^{\circ}$)Skin burningBetamethasone: $4/42$ participants (9.5%)Pimecrolimus: $22/45$ participants (49%) (Difference between groups: $p=0.001^{\circ}$)	Adverse events requiring discontinuation Betamethasone: 1/42 participants (2.4%) Pimecrolimus: 3/45 participants (6.7%) (Difference between groups: p=0.36°)
Luger 2004 ⁽⁶¹⁾ (Broeders 2016	(Cochrane risk of bias tool: unclear risk of selection bias, adequate blinding, inadequate loss to follow up. ⁽⁶⁰⁾) RCT (52 weeks. Twice daily	Intervention: Triamcinolone 0.1% (potent) and Hydrocortisone acetate 1%	Severity: moderate to severe	Skin burning Triamcinolone + hydrocortisone: 36/330 participants (11%)	Severe adverse events Triamcinolone + hydrocortisone: 21/330 participants (6.4%)
(53))	until clearance, restarted with flares) (Jadad score 3/5 – risk from sequence generation and allocation concealment ⁽⁵³⁾) (Cochrane risk of bias tool: unclear risk of selection bias, low risk from blinding. ⁽³⁾)	(face) (Mild potency) twice daily until complete clearance and itching had stopped, then treatment restarted if inflammation recurred (n=330) Comparator: Pimecrolimus 1% twice daily until complete clearance and itching had stopped, then treatment restarted if inflammation recurred (n=328)	Age: adults (age 18 to 79 years) Sample size: 658 participants	Pimecrolimus: 85/328 participants (26%) (Difference between groups: p<0.00001°)	Pimecrolimus: 16/328 participants (4.9%) (Difference between groups: p=0.41°)
	(Jadad scale: 3/5 ⁽⁵⁹⁾) (Cochrane risk of bias tool: adequate allocation generation, unclear allocation concealment, adequate blinding, inadequate loss to follow up. ⁽⁶⁰⁾)			Skin thinning Triamcinolone + hydrocortisone: 3/330 participants (0.9%) Pimecrolimus: 0/328 participants (0%) (Difference between groups: p=20 °) Skin infection Triamcinolone + hydrocortisone: 80/330 participants (24%)	

				Pimecrolimus: 69/328 participants (21%) (<i>Difference between groups:</i> p=0.33°)	
Mandelin 2010 (⁶²⁾ (<i>Broeders 2016</i> (⁵³⁾)	RCT (52 weeks, as prescribed until 7 days after clearance, then restarted with flares) (Jadad score 3/5 – risk from sequence generation and allocation concealment, ⁽⁵³⁾) (Cochrane risk of bias tool: unclear risk of selection bias, risk from no blinding. ⁽³⁾)	Intervention: Hydrocortisone butyrate 0.1% ointment (potent) and Hydrocortisone acetate 1% ointment (face) (Mild potency) twice daily, as prescribed, for a flare until 7 days after clearance, as many times as required in 1 year (n=40) Comparator: Tacrolimus 0.1% ointment twice daily, as prescribed, for a flare until 7 days after clearance, as many times as required in 1 year (n=40)	Severity: moderate to severe Age: adults Sample size: 80 participants	Skin thinningHydrocortisone: 2/40participants (5%)Tacrolimus: 0/40 participants (0%) (Difference between groups: $p=0.29^a$)Skin infectionHydrocortisone: 17/40participants (43%)Tacrolimus: 26/40 participants(65%)(Difference between groups: $p=0.05^a$)	Severe adverse events None in either group

Reitamo 2002 (I) (⁶³⁾ (Broeders 2016 (⁵³); Iskedjian 2004 (⁶⁴))	RCT (3 weeks treatment) (Jadad score 4/5 – risk from allocation concealment ⁽⁵³⁾) (Cochrane risk of bias tool: unclear risk of selection bias, unclear risk from blinding. ⁽³⁾)	Intervention: Hydrocortisone butyrate 0.1% twice daily for 3 weeks (n=186) Comparator: Tacrolimus 0.1% twice daily for 3 weeks (n=191) Comparator: Tacrolimus 0.03% twice daily for 3 weeks (arm not included in Broeders 2016 review) (n=193)	Severity: moderate to severe Age: adults (age 16 to 70 years) Sample size: 571 participants	Skin burning Hydrocortisone: 24/186 participants (13%) Tacrolimus 0.1%: 113/191 participants (59%) (Difference between groups: p<0.00001 °) Pruritus Hydrocortisone: 18/186 participants (9.7%)	Severe adverse eventsHydrocortisone: $0/186$ participants (0%)Tacrolimus 0.1% : $1/191$ participants (0.5%)(Difference between groups: $p=0.51^{\circ}$)Adverse events requiring discontinuation Hydrocortisone: $3/186$
	(Jadad scale: 5/5, ⁽⁵⁹⁾) (Cochrane risk of bias tool: Low risk of selection, performance, detection, attrition, reporting and other biases. ⁽⁵⁴⁾) (Cochrane risk of bias tool: adequate randomisation and allocation concealment, blinding and ITT analysis done. ⁽⁶⁵⁾)			Tacrolimus 0.1%: 29/191 participants (15%) (Difference between groups: p=0.11°) Erythema at application site Hydrocortisone: 1/186 participants (0.5%) Tacrolimus 0.1%: 7/191 participants (3.7%) Tacrolimus 0.03%: 4/193 participants (2.1%) (Difference between groups: tacrolimus 0.1% versus hydrocortisone: p=0.07°) (Difference between groups: tacrolimus 0.03% versus hydrocortisone: p=0.23°)	participants (1.6%) Tacrolimus 0.1%: 8/191 participants (4.2%) (<i>Difference between groups:</i> p=0.15°)

RCT	Intervention: Hydrocortisone	Severity: moderate	Adverse events related to		Severe adverse events
(26 weeks) twice daily		to severe			Hydrocortisone: 9/485
		Age: adults (>18	-		participants (1.9%)
		•	,		Tacrolimus: 5/487 participants
		yearsonay			(1%)
		Sample size: 972	(1.4%)		(Difference between groups:
(Jadad score 5/5; ⁽⁵³⁾)	eczema occurred for 6 months	participants	(Difference between groups:		p=0.29°)
	(n=485)		p=0.34 °)		Adverse events requiring
, ,	Comparatory Togralimus 0.1%		Skin huming		
,	-				discontinuation
					Hydrocortisone: 16/485
from blinding. ⁽³⁾)			,		participants (3.3%)
(Iadad scale: 5/5 (59))					Tacrolimus: 10/487 participants
(suuu scule: 3/3,)	months (n=487)				(2.1%)
(Cochrane risk of bias					(Difference between groups:
tool: High risk of attrition			p<0.00001 °)		p=0.23 °)
bias. Low risk of			Pruritus		
selection, performance,					
detection, reporting and					
other biases. ⁽⁵⁴⁾)			,		
			p=0.05°)		
			Adverse events requiring		
analysis used. ⁽⁶⁵⁾)			discontinuation		
			-		
			$p=0.23^{\circ}$		
			Skin thinning		
			Hydrocortisone: 2/485		
			participants (0.4%)		
			Tacrolimus: 0/487 participants		1
					1
			participants (1.9%)		
			Tacrolimus: 13/487 participants		
			(2.7%)		
			(Difference between groups:		1
			p=0.40°)		
	(26 weeks) twice daily treatment until 7 days after clearance, then whenever a flare occurs) (Jadad score 5/5; ⁽⁵³⁾) (Cochrane risk of bias tool: unclear risk of selection bias,, low risk from blinding. ⁽³⁾) (Jadad scale: 5/5, ⁽⁵⁹⁾) (Cochrane risk of bias tool: High risk of attrition bias. Low risk of selection, performance, detection, reporting and	(26 weeks) twice daily treatment until 7 days after clearance, then whenever a flare occurs)butyrate 0.1%(potent) and Hydrocortisone acetate 1% (face) (Mild potency) twice daily until 7 days after clearance of eczema each time a flare of eczema occurred for 6 months (n=485)(Cochrane risk of bias tool: unclear risk of bias tool: High risk of attrition bias. Low risk of selection, performance, detection, reporting and other biases. (54)butyrate 0.1%(potent) and Hydrocortisone acetate 1% (face) (Mild potency) twice daily until 7 days after clearance of eczema occurred for 6 months (n=485)(Cochrane risk of bias tool: High risk of attrition bias. Low risk of selection bias, unclear if blinded, and unclear if ITTbutyrate 0.1%(potent) and Hydrocortisone acetate 1% (face) (Mild potency) twice daily until 7 days after clearance of eczema occurred for 6 months (n=487)	(26 weeks) twice daily treatment until 7 days after clearance, then whenever a flare occurs)butyrate 0.1%(potent) and Hydrocortisone acetate 1% (face) (Mild potency) twice daily until 7 days after clearance of eczema each time a flare of eczema occurred for 6 months (n=485)to severe(Cochrane risk of bias tool: unclear risk of selection bias, low risk from blinding. ⁽³⁾)Comparator: Tacrolimus 0.1% twice daily until 7 days after clearance of eczema each time a flare of eczema occurred for 6 months (n=487)Sample size: 972 participants(Cochrane risk of bias tool: High risk of attrition bias. Low risk of selection biasy, unclear if blinded, and unclear if ITTComparator: Tacrolimus 0.1% twice daily until 7 days after clearance of eczema each time a flare of eczema occurred for 6 months (n=487)Hage: adults (≥18 years old)	(26 weeks) twice daily treatment until 7 days after clearance, then whenever a flare occurs (ladad score 5/5; (53))butyrate 0.1%(potent) and Hydrocortisone actate 1% (face) (Mild potency) twice daily until 7 days after clearance of eczema acch time a flare of eczema accurred for 6 months (n=485)to servere Age: adults (218 vars old)treatment Hydrocortisone: 11/485 participants (2.3%) Tacrolimus; 7/487 participants (La4%) (Difference between groups: p=0.347)(ladad scale: 5/5, (59))Comparator: Tacrolimus 0.1% twice daily until 7 days after clearance of eczema accurred for 6 months (n=487)Sample size: 372 participantsSkin burning Hydrocortisone: 67/485 participants (1.3%) Tacrolimus: 255/487 participants (52%) (Difference between groups: p=0.30010*)(Cochrane risk of bias tool: unclear risk of selection bias, unclear if tool: unclear risk of selection bias, unclear if tool: unclear risk of selection bias, unclear if (Farsene between groups: p=0.30*)Pruritus Hydrocortisone: 65/485 participants (13%) Tacrolimus: 88/487 participants (13%) (Difference between groups: p=0.35*)(Cochrane risk of bias tool: unclear risk of selection bias, unclear if tool: unclear risk of selection bias, unclear if (Farsene between groups: p=0.35*)Adverse events requiring discontinuation Hydrocortisone: 16/485 participants (0.4%) Tacrolimus: 0/487 participants (2%)(Difference between groups: p=0.30*)Skin Infection Hydrocortisone: 5/485 participants (0.4%) Tacrolimus: 0/487 participants (2%)(Gochrane risk of selection bias, unclear if (Farsene between groups: p=0.30*)Skin Infection Hydrocortisone: 16/485 <br< td=""><td>(26 weeks) twice daily treatment until 7 days after clearance, then whenever a flare occurs, (laded score 5/5.¹⁶¹⁾)but/vate 0.134(potency) twice daily until 7 days after clearance of eczema occurred for 6 months (ned8):to severe severe treatment (Difference between groups: p=0.34)(Cochrone risk of bios. (low risk tool: linkin, low risk tool:</td></br<>	(26 weeks) twice daily treatment until 7 days after clearance, then whenever a flare occurs, (laded score 5/5. ¹⁶¹⁾)but/vate 0.134(potency) twice daily until 7 days after clearance of eczema occurred for 6 months (ned8):to severe severe treatment (Difference between groups: p=0.34)(Cochrone risk of bios. (low risk tool: linkin, low risk tool:

Gradman 2007 (⁶⁷⁾ (Svensson 201 (⁶⁸⁾ 1)	Crossover RCT (2 weeks treatment) (Cochrane risk of bias tool: low risk of selection bias from sequence generation, but unclear for allocation concealment. Low risk from blinding. ⁽³⁾)	Intervention: Mometasone furoate 0.1% once daily Comparator: Tacrolimus 0.1% twice daily	Severity: mild to moderate Age: children 5 to 12 years Sample size: 20 participants		Withdrawal from study Mometasone: 1 patient Tacrolimus: 1 patient
Kawashima 1997 ⁽⁶⁹⁾ (Ashcroft 2005 ⁽⁵⁹⁾)	RCT (3 weeks treatment) (Jadad scale: 5/5, ⁽⁵⁹⁾)	Intervention: Betamethasone valerate 0.12% twice daily for three weeks (n=89) Comparator: tacrolimus 0.1% twice daily for three weeks (n=92)	Severity: mild to moderate Age: adults Sample size: 181 participants	Skin infections Betamethasone: 5/89 participants Tacrolimus: 6/92 participants (Difference between groups: p=0.80°) Skin burning Betamethasone: 3/89 participants Tacrolimus: 25/92 participants (Difference between groups: p=0.0004°)	
			Potent or mild potence	y topical corticosteroids	
Hofman 2006 ⁽⁷⁰⁾ (Broeders 2016 ⁽⁵³⁾ ; Siegfried 2016 ⁽⁷¹⁾)	RCT (2 weeks treatment, 28 weeks follow up) (Jadad score 5/5 – risk from sequence generation and allocation concealment ⁽⁵³⁾)	Intervention: Hydrocortisone ointment 1% (mild potency) twice daily for head/neck and hydrocortisone butyrate ointment 0.1% (potent) for trunk and limbs for 2 weeks then hydrocortisone 1% (mild potency) twice daily for flares. (n=124) Comparator: Tacrolimus 0.03% twice daily for 3 weeks then tacrolimus once daily and vehicle once daily for flares (n=133)	Severity: moderate to severe Age: children 2 to 11 years old (mean 6 years old) Sample size: 257 participants	Adverse events related to treatment Hydrocortisone: 2/124 participants (1.6%) Tacrolimus: 10/133 participants (7.5%) (Difference between groups: $p=0.04^{a}$) Skin burning Hydrocortisone: 0/124 participants (0%) Tacrolimus: 2/133 participants (1.5%) (Difference between groups: $p=0.32^{a}$) Pruritus Hydrocortisone: 4/124 participants (3%) Tacrolimus: 8/133 participants (6%)	Severe adverse events Hydrocortisone: 0/124 participants (0%) Tacrolimus: 2/133 participants (1.5%) (Difference between groups: p=0.32°)

				(Difference between groups: $p=0.30^{a}$) Skin infection Hydrocortisone: 4/124 participants (3.2%) Tacrolimus: 2/133 participants (1.5%) (Difference between groups: $p=0.37^{a}$) Bacterial infection Hydrocortisone: 3/124 participants (2%) Tacrolimus: 33/133 participants (2%) (Difference between groups: p<0.0001) Viral infection Hydrocortisone: incidence not reported Tacrolimus: 1/133 participants (0.8%)	
			Moderate potency t	topical corticosteroids	
Sikder 2005 ⁽⁷²⁾ (Broeders 2016 ⁽⁵³⁾)	RCT (4 weeks treatment) (Jadad score 2/5 – risk from sequence generation and allocation concealment, no blinding of observer or patients, (⁽⁵³⁾) (Cochrane risk of bias tool: Unclear risk of selection and detection bias. Low risk of performance, attrition, reporting and other biases. ⁽⁵⁴⁾) (Cochrane risk of bias tool: low risk of selection bias, from blinding of participants and missing data. Unclear risk from	Intervention: Clobetasone 0.05% twice daily (n=15) Tacrolimus 0.03% twice daily (n=15)	Severity: moderate to severe Age: children 7 to 15 years old Sample size: 30 participants	Skin burning Clobetasone: 1/15 participants (6.7%)Tacrolimus: 7/15 participants (47%) (Difference between groups: $p=0.05^{a,d}$)Pruritus Clobetasone: 2/15 participants (13%) Tacrolimus: 3/15 participants (20%) (Difference between groups: $p=0.63^{a}$)	

Torok 2003 ⁽⁷³⁾ (Svensson 2011 ⁽⁶⁸⁾)	blinding outcome assessors, reporting and other biases ⁽⁵⁶⁾). RCT (3 weeks treatment) (Delphi list: method of randomisation not described, allocation not concealed, blinded assessors but not participants, ITT analysis, ⁽²⁹⁾)	Intervention: Clocortolone pivalate 0.1% twice daily (n=19) Intervention: Clocortolone 0.1% + Tacrolimus 0.1% twice daily (n=19) Comparator: Tacrolimus 0.1% twice daily (n=19)	Severity: not specified in the review Age: adults 16 to 65 years Sample size: 57 participants	Skin irritation Most commonly reported adverse event Skin burning More frequent in those treated with Tacrolimus 0.1%. Pruritus Commonly reported in both arms. (No numerical data provided in the review)		
			Moderate or mild pote	ncy topical corticosteroids		
Sigurgeirsson 2015 ⁽⁷⁴⁾ (Broeders 2016 ⁽⁵³⁾ ; Siegfried 2016 ⁽⁷¹⁾)	RCT (260 weeks used until clearance or according to country's label. Medication reinstated when a flare occurred)) (Jadad score 3/5 – risk from allocation concealment, no blinding of observer or patients, (53))	Intervention: A moderate potency or mild potency TCS used according to the country's label with potency selected by the investigator (n=1213) Comparator: Pimecrolimus 1% twice daily (n=1205)	Severity: mild to moderate Age: children age 3 to 12 months old (mean 7 months) Sample size: 2418 participants	Skin thinning(from online correspondence)Topical corticosteroid: 1/1213 participants (0.08%)Pimecrolimus: 0/1205 (0%) (Difference between groups: $p=0.50^{a}$)Skin infection Topical corticosteroid: 150/1213 participants (12%)Pimecrolimus: 157/1205 participants (13%) (Difference between groups: $p=0.62^{a}$)Cutaneous bacterial infection Topical corticosteroid: 121/1213 participants (10%)Pimecrolimus: 145/1205 participants (10%)Pimecrolimus: 145/1205 participants (23%)Pimecrolimus: 301/12 05 participants (25%)	Systemic bacterial infectionTopical corticosteroid: 206/1213participants (17%)Pimecrolimus: 205/1205participants (17%)(Difference between groups: $p=0.98^a$)Systemic viral infectionTopical corticosteroid: 206/1213participants (17%)Pimecrolimus: 205/1205participants (17%)Difference between groups: $p=0.98^a$)Systemic RTITopical corticosteroid: 388/1213participants (32%)Pimecrolimus: 422/1205participants (35%)(Difference between groups: $p=0.11^a$)Systemic GITopical corticosteroid: 376/1213participants (31%)Pimecrolimus: 386/1205participants (31%)	Severe adverse events Topical corticosteroid: 210/1213 participants (17%) Pimecrolimus: 247/1205 participants (20%) (Difference between groups: p=0.05°) Adverse events requiring discontinuation Topical corticosteroid: 12/1213 participants (1.0%) Pimecrolimus: 7/1205 participants (0.6%) (Difference between groups: p=0.26°)

				(Difference between groups: p=0.25°)	(Difference between groups: p=0.58 ^a) Lymphoma Zero cases in either group Growth rate and immune system No difference between groups	
			Mild potency top	pical corticosteroids		
Reitamo 2002 (II) ⁽⁷⁵⁾ (Broeders 2016 ⁽⁵³⁾ ; Iskedjian 2004 ⁽⁶⁴⁾)	RCT (3 weeks treatment) (Jadad score 5/5 – risk from allocation concealment ⁽⁵³⁾) (Cochrane risk of bias tool: unclear risk of selection bias, unclear risk of blinding ⁽³⁾) (Jadad scale: 5/5, ⁽⁵⁹⁾) (Cochrane risk of bias tool: Low risk of selection, performance, detection, attrition, reporting and other biases. ⁽⁵⁴⁾) Cochrane risk of bias tool: low risk of selection, performance, attrition, reporting and other bias. Unclear risk from blinding outcome assessors ⁽⁵⁶⁾). (Cochrane risk of bias tool: adequate method of randomisation and allocation concealment, blinding done, ITT used. (⁶⁵⁾)	Intervention: Hydrocortisone acetate 1% twice daily (n=185) Comparator: Tacrolimus 0.1% ointment twice daily (n=186) Comparator: Tacrolimus 0.03% ointment twice daily (arm not included in Broeders 2016 review) (n=189)	Severity: moderate to severe Age: children 2 to 15 years old Sample size: 560participants	Skin burningHydrocortisone: 13/185participants (7%)Tacrolimus 0.1%: 38/186participants (20%)(Difference between groups: $p=0.004^{a}$)PruritusHydrocortisone: 14/185participants (7.6%)Tacrolimus 0.1%: 21/186participants (11%)(Difference between groups: $p=0.22^{a}$)Skin infectionHydrocortisone: 4/185participants (2.2%)Tacrolimus 0.1%: 4/186participants (2.2%)Tacrolimus 0.1%: 4/186participants (2.2%)(Difference between groups: $p=0.99^{a}$)Erythema at application siteHydrocortisone: 3/185participants (1.6%)Tacrolimus 0.1%: 1/186participants (2.1%)(Difference between groups,hydrocortisone vs tacrolimus0.1%: $P=0.34^{a}$)(Difference between groups,hydrocortisone vs tacrolimus0.1%: $P=0.34^{a}$)(Difference between groups,hydrocortisone vs tacrolimus0.1%: $P=0.72^{a}$)		Severe adverse events Hydrocortisone: 2/185 participants (1.1%) Tacrolimus 0.1%: 1/186 participants (0.5%) (<i>Difference between groups:</i> (<i>p</i> =0.57°) Adverse events requiring discontinuation Hydrocortisone: 4/185 participants (2.2%) Tacrolimus 0.1%: 3/186 participants (1.6%) (<i>Difference between groups:</i> <i>p</i> =0.70°)

Reitamo 2004 (76) (Broeders 2016 (53))	RCT (3 weeks treatment) (Jadad score 3/5 – risk from sequence generation and allocation concealment ⁽⁵³⁾) (Cochrane risk of bias tool: unclear risk of selection bias. Unclear risk from blinding. ⁽³⁾) (Jadad scale: 4/5, ⁽⁵⁹⁾) (Cochrane risk of bias tool: Unclear risk of selection bias (allocation concealment). Low risk of selection bias (random sequence generation), performance, detection, attrition, reporting and other biases. ⁽⁵⁴⁾) Cochrane risk of bias tool: low risk of selection, performance, attrition, reporting and other bias. Unclear risk from blinding outcome assessors ⁽⁵⁶⁾).	Intervention: Hydrocortisone acetate 1% twice daily (n=207) Comparator: Tacrolimus 0.03% twice daily (n=210)	Severity: moderate to severe Age: children 2 to 15 years old Sample size: 417 participants	Skin burningHydrocortisone: $30/207$ participants (15%)Tacrolimus: $50/210$ participants (24%) (Difference between groups: $p=0.02^a$)PruritusHydrocortisone: $33/207$ participants (16%)Tacrolimus: $45/210$ participants (21%) (Difference between groups: $p=0.15^a$)Skin infectionHydrocortisone: $6/207$ participants (2.9%)Tacrolimus: $6/210$ participants (2.9%) (Difference between groups: $p=0.98^a$)		Severe adverse events Hydrocortisone: $3/207$ participants (1.4%) Tacrolimus: $3/210$ participants (1.4%) (Difference between groups: $p=0.99^{a}$) Adverse events requiring discontinuation Hydrocortisone: $6/207$ participants (2.9%) Tacrolimus: $8/210$ participants (3.8%) (Difference between groups: $p=0.61^{a}$)
		·	Potency of topical co	rticosteroids unknown		
Gutgesell 1998 (77) (abstract only) (Penaloza Hidalgo 2004 (65))	Within-participant RCT (3 weeks treatment) (Cochrane risk of bias tool: randomisation and allocation concealment method inadequate, unclear if blinded ,ITT analysis used ⁽⁶⁵⁾)	Intervention: Topical corticosteroids on one side of the body, twice daily Comparator: Tacrolimus 0.1% on one side of the body, twice daily	Severity: severe Age: adults (22 to 36 years) Sample size: 7 participants	Skin burning Topical corticosteroids: 0/7 (0%) Tacrolimus: 2/7 participants (29%)		
Arellano 2007 (78) (Ashcroft 2007 (⁶⁰⁾ ; Cury Martins 2015 ⁽⁵⁴⁾	Nested case-control (Duration not specified in the review)	Intervention: Topical corticosteroids at different potencies Comparator: pimecrolimus or tacrolimus	Severity: not specified in the review Age: not specified in the review		Lymphoma No increased risk of lymphoma with TCl or TCS when compared against controls. Super potent TCS: OR 1.2, 95% Cl 0.8 to 1.8	

	Risk of bias not assessed in any of the included systematic reviews.	Comparator: controls (not specified in the review)	Sample size: 294 cases/293,000 controls		<i>Low potency TCS: OR</i> 1.1, 95%Cl 0.7 to 1.6 <i>Pimecrolimus:</i> OR 0.8, 95%Cl 0.4 to 1.6 <i>Tacrolimus</i> OR 0.8, 95% Cl 0.4 to 1.7	
Arellano 2009 (⁷⁹⁾ (Cury Martins 2015 ⁽⁵⁴⁾)	Cohort (followed up between 1992 to 2006) Risk of bias not assessed in any of the included systematic reviews.	Intervention: Topical corticosteroids at different potencies Comparator: pimecrolimus or tacrolimus	Severity: not specified in the review Age: not specified in the review Sample size: > 3,000,000		Lymphoma Increased risk with topical corticosteroids (related to potency) but no numerical data given. Insufficient data to assess TCI-related risks.	
Schneeweiss 2009 ⁽⁸⁰⁾ (Cury Martins 2015 ⁽⁵⁴⁾)	Cohort (followed up between the years of 2002 to 2006) Risk of bias not assessed in any of the included systematic reviews.	Intervention: mid to potent topical corticosteroids (n=1,043,025) Comparator: pimecrolimus (n=118,863) or tacrolimus (n=38,757) (also a comparison with untreated dermatitis (n=118,825) and general population (n=118,863) .)	Severity: not specified Age: median 1.3 years Sample size: 1,438,333 participants		Lymphoma Very small non-significant increased risk in TCI and TCS patients when compared with the general population, but with similar risks between the treatment groups	
Reitamo 2000 ⁽⁸¹⁾ (Cury Martins 2015 ⁽⁵⁴⁾)	Open label, single group (6 to 12 months of treatment) Risk of bias not assessed in any of the included systematic reviews.	Intervention: No steroids (except prior to treatment) Comparator: Tacrolimus 0.1%	Severity: not specified in the review Age: adults Sample size: 316 participants	Skin thinning One participant had skin thinning when using TCS prior to treatment with tacrolimus – but this ameliorated after 6 months of treatment with tacrolimus.		
		Is there any difference	e in safety of topi	cal corticosteroids of different	ent potencies?	
Study ID (Systematic review*)	Study design and study duration (Quality assessment)	Intervention and comparator	Participants	Cutaneous adverse events	Systemic adverse events	Unspecified adverse events
		Potent topical o	orticosteroid versus r	noderate potency topical cortico	steroid	
Ulrich 1991 ⁽⁸²⁾ (Hoare 2000 ⁽¹⁹⁾)	RCT (2 weeks treatment) (Moher 1995 quality checklist: method and	Intervention: 0.05% halomethasone cream, twice daily (Assume potent)	Severity: not specified in the review			No adverse events

Smitt 1993 ⁽⁸³⁾ (Callen 2007 ⁽²⁴⁾)	concealment of randomisation unclear, concerns over subgroup analysis ⁽¹⁹⁾ RCT (3 weeks treatment) Risk of bias not assessed in any of the included systematic reviews.	Comparator: 0.25% prednicarbate cream, twice daily (moderate potency) Intervention: Trimaconiolone acetonide 0.1%, twice daily (potent) Comparator: Alclomethasone cream, twice daily (moderate)	Age: not specified in the reviewSample size: 165 participantsSeverity: not specified in the reviewAge: children 1 to 15 yearsSample size: 40 participants		HPA axis suppression There was suppression after 2 weeks, but no further after 3 (no further details).	
		Potent topica	l corticosteroid versu	s mild potency topical corticost	eroid	I
Lebrun-Vignes 2000 ⁽⁸⁴⁾ (Nankervis 2017 (³)	RCT (15 days treatment, 30 days follow up) (Cochrane risk of bias tool: unclear risk of selection bias and unclear risk from blinding. ⁽³⁾)	Intervention: Micronized desonide cream 0.1% (mild potency) 1 to 5 days twice daily (in hospital), days 6 and 7 once daily, then alternate days until day 15 (n=15) Comparator: Betamethasone dipropionate cream 0.05% (potent) 1 to 5 days twice daily (in hospital), days 6 and 7 once daily, then alternate days until day 15 (n=14)	Severity: severe Age: children ≤ 8 years Sample size: 29 participants			There were no adverse events in either group
Prado de Oliveira 2002 ⁽⁸⁵⁾ (Nankervis 2017 ⁽³⁾) Hanifin 1996 ⁽⁸⁶⁾ (Callen 2007 ⁽²⁴⁾)	RCT (Up to 42 days treatment) (Cochrane risk of bias tool: unclear risk of selection bias and unclear risk from blinding. ⁽³⁾) Matched case control (3 weeks treatment) Risk of bias not assessed in any of the included systematic reviews.	Intervention: Mometasone furoate 0.1% once daily after a bath (N=13) (potent) Comparator: Desonide cream 0.05% once a day after a bath (N=12) (mild potency) Intervention: Mometasone cream (potent) Comparator: Hydrocortisone cream (mild potency)	Severity: not specified in the review Age: children 2 to 12 years Sample size: 25 participants Severity: not specified in the review Age: children 6 months to 2 years Sample size: 62 participants	Signs of mild thinning Mometasone furoate: 4/13 participants (31%) Desonide: 2/12 participants (17%) (Difference between groups: p=0.42°)	HPA axis suppression Mometasone: 1 abnormal cotrosyn simulation test	

Kirkup 2003 ⁽⁸⁷⁾ (trial a) (Tang 2014 ⁽⁸⁸⁾ ; Siegfried 2016 ⁽⁷¹⁾) Most safety data presented was combined with Kirkup 2003 (trial b) (see same potency section below)	RCT (16 weeks: twice daily for 2-4 weeks until stabilised then 'as required' for 12 weeks) (Cochrane risk of bias tool: unclear risk of selection bias, unclear risk from blinding. ⁽³⁾)	Intervention: Fluticasone propionate 0.005% ointment (potent), twice daily for 2-4 weeks until stabilised then 'as required' for 12 weeks (n=70) Comparator: Hydrocortisone 1% cream (mild potency), twice daily for 2-4 weeks until stabilised then 'as required' for 12 weeks (n=67)	Severity: moderate Age: children (age 2-14 years old) Sample size (maintenance phase): 137 participants	Ringworm and folliculitis 1 participant but not clear which group Kirkup 2003a and b: Bacterial infection Fluticasone: 1/136 participants (0.7%) Hydrocortisone: 3/129 participants (2%) (Difference between groups: p = 0.32 °) Kirkup 2003a and b: Fungal infection Fluticasone: 1/136 participants (0.7%) Hydrocortisone: 0/129 (0%) (Difference between groups: p = 0.52 °)	Kirkup 2003a and b viral infection Fluticasone: 5/136 participants (4%) Hydrocortisone: 5/129 participants (4%) (Difference between groups: p = 0.93°) Kirkup 2003a and b: Respiratory tract infection Fluticasone: 8/136 participants (6%) Hydrocortisone: 5/129 participants (4%) (Difference between groups: p = 0.45°)	Kirkup 2003a and b: Discontinuation due to adverse events Fluticasone: 1/136 participants (0.7%) Hydrocortisone: 1/129 participants (0.7%) (Difference between groups: p = 0.97°)
		Moderate potency	topical corticosteroid	versus mild potency topical cor	ticosteroid	
Kuokkanen 1987 ⁽⁸⁹⁾ (Hoare 2000 ⁽¹⁹⁾)	RCT, within participant (3 weeks treatment) (Moher 1995 quality checklist: method and concealment of randomisation unclear, double blinded study, three dropouts/ withdrawals, no ITT, ⁽¹⁹⁾)	Intervention: Alclometasone dipropionate 0.05% twice daily (moderate potency) Comparator: Hydrocortisone 1% twice daily (mild potency)	Severity: not specified in the review Age: children Sample size: 37 participants	No evidence of skin thinning		
			Various	potencies		
Ellison 2000 ⁽⁹⁰⁾ (Callen 2007 ⁽²⁴⁾ ; Eichenfield 2014 ⁽⁹¹⁾)	Observational study (Duration 0.7 to 18.7 years) Risk of bias not assessed in any of the included systematic reviews.	Intervention: Mild, moderate, potent topical corticosteroids	Severity: Disease severity score 5-8 Age: children and adolescents (0.7 to 18.7 years) Sample size: 35 participants		HPA axis suppression Mild potency topical corticosteroids: no change in plasma cortisol levels Potent topical corticosteroids: suppression in 4/4 (100%) patients	

(Tang 2014 ⁽⁸⁸⁾ ; Siegfried 2016 ⁽⁷¹⁾)	2-4 weeks until stabilised then intermittently for 12 weeks)	stabilised then intermittently for 12 weeks (n=66) Comparator: Hydrocortisone	14 years old) Sample size: n=128	Kirkup 2003a and b: Bacterial infection Fluticasone: 1/136 participants (0.7%)	(4%) Hydrocortisone: 5/129 participants (4%) (Difference between groups: p =	Fluticasone: 1/136 participants (0.7%) Hydrocortisone: 1/129 participants (0.7%)
Kirkup 2003 ⁽⁸⁷⁾ (trial b)	RCT (16 weeks: twice daily for	Intervention: Fluticasone propionate 0.005% ointment, twice daily for 2-4 weeks until	Severity: moderate Age: children (age 2-	Ringworm and folliculitis None reported	Kirkup 2003a and b viral infection Fluticasone: 5/136 participants	Kirkup 2003a and b: Discontinuation due to adverse events
		· · · · · · · · · · · · · · · · · · ·		another potent topical corticost		
Study ID (Systematic review*)	Study design and study duration (Quality assessment)	Is there any difference in Intervention and comparator	n safety between t Participants	topical corticosteroids of th Cutaneous adverse events	e same potency? Systemic adverse events	Unspecified adverse events
Patel 1998 ⁽⁹⁴⁾ (Eichenfield 2014 ⁽⁹¹⁾)	systematic reviews. Observational study (Duration not specified in the review) Risk of bias not assessed in any of the included systematic reviews.	Intervention: Different potencies of topical corticosteroids	Sample size: not specified in the review Severity: not specified in the review Age: not specified in the review Sample size: not specified in the review		Review authors reported "Also concerns for negative effects on linear growth, although reports have given mixed conclusions"	
Kristmundsdottir 1987 ⁽⁹²⁾ (Eichenfield 2014 ⁽⁹¹⁾) Patel 1997 ⁽⁹³⁾ (Eichenfield 2014 ⁽⁹¹⁾)	Observational study (Duration not specified in the review) Risk of bias not assessed in any of the included systematic reviews. Observational study (Duration not specified in the review) Risk of bias not assessed in any of the included	Intervention: Different potencies of topical corticosteroids Intervention: Four different potency topical corticosteroids	Severity: not specified in the review Age: not specified in the review Sample size: not specified in the review Severity: not specified in the review Age: not specified in the review		Review authors reported "Also concerns for negative effects on linear growth, although reports have given mixed conclusions" Review authors reported "Also concerns for negative effects on linear growth, although reports have given mixed conclusions"	

(trial a) (see different potency section above)		stabilised then intermittently for 12 weeks (n=62)		(Difference between groups: p = 0.32°) Kirkup 2003a and b: Fungal infection Fluticasone: 1/136 participants (0.7%) Hydrocortisone: 0/129 (0%) (Difference between groups: p = 0.52°)	<pre>Fluticasone: 8/136 participants (6%) Hydrocortisone: 5/129 participants (4%) (Difference between groups: p = 0.45 °)</pre>	
		Moderate potency topical	corticosteroid versus	another moderate potency topi	cal corticosteroid	
Aliaga 1994 ⁽⁹⁵⁾ (De Tiedra 1997 ⁽³³⁾)	RCT (21 days treatment) Risk of bias not assessed in any of the included systematic reviews.	Intervention: Prednicarbate ointment 0.25%, twice daily (moderate potency) (n=36) Comparator: Flucortin ointment 0.75%, twice daily (assumed moderate potency) (n=31)	Severity: Disease duration – mean 7.7 years (range 0.1 to 31). Age: adults 18-74 years (mean 33.6) Sample size: 67 participants			Adverse reactions Prednicarbate: 0/36 patients (0%) Flucortin: 2/31 patients (6.5%) (Difference between groups: p = 0.16 ^a)
		Mild potency topical	corticosteroid versus	another mild potency topical co	rticosteroid	
Lucky 1997 ⁽⁹⁶⁾ (Callen 2007 ⁽²⁴⁾ ; Hoare 2000 ⁽¹⁹⁾ ; Wood Heickman 2018 ⁽⁹⁾)	RCT (4 weeks treatment) (Moher 1995 quality checklist: method and concealment of randomisation unclear, open label, five dropouts, no ITT ⁽¹⁹⁾)	Intervention: Desonide 0.05% ointment, twice per day (mild potency) Comparator: Hydrocortisone 2.5% ointment, twice per day (mild potency)	Severity: not specified in the review Age: children (mean or median is 4.7 years) Sample size: 20 participants		HPA axis suppression Normal in both groups (measured using ACTH stimulation testing, measuring serum cortisol levels)	
Jorizzo 1995 ⁽⁹⁷⁾ (Siegfried 2016 ⁽⁷¹⁾ ; Froeschl 2007 ⁽⁹⁸⁾)	RCT (25 weeks: 5 weeks of treatment, 20 weeks follow up) (Moher 1995 quality checklist: method and concealment of randomisation unclear, investigator blind, two dropouts/withdrawals, no ITT ⁽¹⁹⁾)	Intervention: 0.05% desonide twice daily (n=16) (mild potency) Comparator: 1% hydrocortisone ointment twice daily (n=20) (mild potency)	Severity: mild to moderate Age: children 5 years and under Sample size: 36 participants	Skin thinning No cases - measured by a magnifying lamp		

		How safe are topic	al corticosteroids	compared to Chinese herba	I medicine?	
Study ID (Systematic review*)	Study design and study duration (Quality assessment)	Intervention and comparator	Participants	Cutaneous adverse events	Systemic adverse events	Unspecified adverse events
			Very potent top	ical corticosteroids		·
Huang 2010 ⁽⁹⁹⁾ (Gu 2013 ⁽¹⁰⁰⁾ ; Gu 2014 ⁽¹⁰¹⁾)	RCT (2 weeks treatment, followed up for 12 weeks after) (Cochrane risk of bias tool: low risk of selection bias (random sequence generation), and other biases. Unclear risk of selection (allocation concealment), detection and attrition bias. High risk of performance and reporting bias. ^(100, 101)	Intervention: Clobetasol propionate ointment, 3 times daily (n=97) Comparator: Chushi Zhiyang ointment, 3 times daily (n=98)	Severity: not specified in the review Age: children and adults, 3 months to 22 years Sample size: 195 participants	Cutaneous adverse eventsClobetasol: $5/97$ participants (5%) Chinese herbal medicine: $0/98$ participants (0%) (Difference between groups: $p=0.10^{a, e})$ The five events were pigmentation (unclear if hyper- or hypo-)		
			Potent topica	l corticosteroids		1
Chen 2011 ⁽¹⁰²⁾ (Gu 2013 ⁽¹⁰⁰⁾ ; Gu 2014 ⁽¹⁰¹⁾)	RCT (2 weeks treatment) (Cochrane risk of bias tool: unclear risk of selection, detection, attrition, reporting and other bias. High risk of performance bias ^(100, 101))	Intervention: Mometasone furoate cream, once daily (n=50) Comparator: Huanglian Qingdai ointment, 2 to 3 times daily (n=50)	Severity: not specified in the review Age: children 58 days to 2 years Sample size: 100 participants	Cutaneous adverse eventsMometasone: $6/50$ participants (12%) Chinese herbal medicine: $0/50$ participants (0%)(Difference between groups: $p=0.08^{o,f}$)Minor adverse events such asburning, dryness and scaling ofthe skin were reported in the TCSgroups		
Dong 2012 ⁽¹⁰³⁾ (Gu 2014 ⁽¹⁰³⁾)	RCT (2 weeks treatment) (Cochrane risk of bias tool: unclear risk of selection, detection, attrition, and reporting bias. High risk of	Intervention: Hydrocortisone butyrate cream, twice daily (n=47) Comparator: Jingfang mixture solution, twice daily (n=48)	Severity: not specified in the review Age: children 0.5 to 5.5 years Sample size: 95 participants	Minor adverse events such as burning, dryness and scaling of the skin were reported in the TCS groups. (No numerical data provided in the review)		

Xu 2012 ⁽¹⁰⁴⁾ (Gu 2014 ⁽¹⁰¹⁾)	performance bias. Low risk of other biases. ⁽¹⁰¹⁾ RCT (2 weeks treatment) (Cochrane risk of bias tool: unclear risk of selection, detection, attrition, and reporting bias. High risk of performance and other	Intervention: Triamcinolone acetonide acetate cream, twice daily (n=51) Comparator: Kouqiang Xiaoyan powder, twice daily (n=53)	Severity: not specified in the review Age: children 35 days to 2 years Sample size: 104 participants			No adverse events in either group
	biases. ⁽¹⁰¹⁾) How	safe is more frequent topi	cal corticosteroid	application compared with	once daily application?	
Study ID (Systematic review*)	Study design and study duration (Quality assessment)	Intervention and comparator	Participants	Cutaneous adverse events	Systemic adverse events	Unspecified adverse events
			Very potent topi	ical corticosteroid	L	
Schlessinger 2006 ⁽¹⁰⁵⁾ (Nankervis 2017 ⁽³⁾ ; Wood Heickman 2018 ⁽⁹⁾)	Open label RCT (2 weeks treatment) (Cochrane risk of bias tool: unclear risk of selection bias, high risk from no blinding. ⁽³⁾)	Intervention: fluocinonide cream 0.1% applied once daily (n=63) Comparator: fluocinonide cream 0.1% applied twice daily (n=63)	Severity: not specified in the review Age: children, aged 12 to <18 years (cohort 1); 6 to <12 years (cohort 2); 2 to <6 years (cohort 2); 2 to <6 years (cohort 3); and 3 months to <2 years (cohort 4). Sample size: 126 participants		HPA axis suppression Once daily: 0/63 (0%) Twice daily: 3/63 (4.8%) (Difference between groups: P=0.19°) (measured using ACTH stimulation testing, measuring serum cortisol levels) After TCS discontinuation, children with biochemical adrenal insufficiency had complete resolution at retesting.	
			Potent topical	corticosteroids		
Bleehen 1995 (106) (Green 2004 (107)	RCT (4 weeks treatment)) (Quality using NHS CRD criteria: method for randomisation/allocation concealment unknown, adequate blinding, and ITT used. ⁽¹⁰⁷⁾) (Moher 1995 quality checklist: method and	Intervention: Fluticasone propionate 0.05% cream once daily (plus vehicle once daily for blinding) (n=137) Comparator: Fluticasone propionate 0.05% cream twice daily (n=133)	Severity: at least moderate severity Age: children and adults Sample size: 270 participants			Number of events possibility, probably or almost certainly related to study medication Once daily: 26 events Twice daily: 24 events (most were skin disorders)

	concealment of randomisation unclear, Probably investigator blinded but unclear, ITT analysis, ⁽¹⁹⁾)				
GSK report 1995 (108) (Green 2004 (107))	RCT (4 weeks treatment) (Quality using NHS CRD criteria: adequate method of randomisation /allocation concealment, adequate blinding, and ITT used. ⁽¹⁰⁷⁾)	Intervention: Fluticasone propionate 0.005% ointment once daily and placebo only daily (n=123) Comparator: Fluticasone propionate 0.005% ointment twice daily (n=122)	Severity: at least moderate severity Age: children and adults Sample size: 245 participants		Number of adverse events possibly related to medication Once daily: 6 events Twice daily: 8 events Number of adverse events probably related to medication Once daily: 9 events Twice daily: 3 events Number of adverse events almost certain related to medication Once daily: 6 events Twice daily: 3 events (Mainly included skin related disorders including exacerbation of eczema, pruritus and redness of skin)
Koopmans 1995 (109) (Green 2004 (107)	RCT (4 weeks treatment)) (Quality using NHS CRD criteria: method for randomisation /allocation concealment unknown, partial blinding, and no ITT used. (¹⁰⁷)) (Moher 1995 quality checklist: method and concealment of randomisation unclear, double blinded, one dropout, no ITT analysis, (¹⁹)	Intervention: Locoid lipocream (0.1% hydrocortisone 17- butyrate) once daily and locobase once daily (n=75) Comparator: Locoid lipocream twice daily (n=75)	Severity: not specified in the review Age: children aged over 12 years and adults Sample size: 150 participants	Folliculitis in all skin areas after 1 week of treatment – treatment store of treatment – treatment Once daily: 1/75 participants (1.3%) Twice daily: 0/75 participants (0%) Colspan="2">Colspan= 0.50°) Folliculitis - treatment continued Once daily: 0/75 participants (0%) Twice daily: 0/75 participants (5.3%) Difference between groups: p = 0.14°) Burning, itching and stinging sensations – treatment continued Once daily: 3/75 participants (4%)	

				Twice daily: 0/75 participants(0%)(Difference between groups: p =0.20°)	
Tharp 1996 ⁽¹¹⁰⁾ (Green 2004 (107))	RCT (4 weeks treatment) (Quality using NHS CRD criteria: method for randomisation /allocation concealment unknown, adequate blinding, and no ITT used. (¹⁰⁷))	Intervention: Fluticasone propionate cream 0.05% once daily and vehicle once daily (n=77) Comparator: Fluticasone propionate cream 0.05% twice daily (n=77)	Severity: moderate to severe Age: children over 12 years and adults Sample size: 154 participants	Burning Once daily: 2/77 participants (3%) Twice daily: 0/77 participants (0%) (Difference between groups: p = 0.30°) Dryness Once daily: 2/77 participants (3%) Twice daily: 2/77 participants (3%) Twice daily: 0/77 participants (0%) ///fference between groups: p = 0.30°) Pruritus Once daily: 0/77 participants (1%) //Difference between groups: p = 0.50°) Erythema Once daily: 0/77 participants (0%) Twice daily: 1/77 participants (1%) //ifference between groups: p = 0.50°) Irritation Once daily: 0/77 participants (1%) Difference between groups: p = 0.50°) Irritation </td <td>None of adverse events were serious or unexpected</td>	None of adverse events were serious or unexpected

				(Difference between groups: p = 0.50°)	
Hoybye 1991 (111) (Green 2004 (107))	RCT ((3 weeks treatment) (Quality using NHS CRD criteria: method for randomisation /allocation concealment unknown, partial or inadequate blinding, and no ITT used. ⁽¹⁰⁷⁾) (Moher 1995 quality checklist: method and concealment of randomisation unclear, single blind, ten dropouts/withdrawals, no ITT analysis, ⁽¹⁹⁾)	Intervention: Mometasone furoate in fatty cream base (Elocon) once daily (n=49) Comparator : Hydrocortisone 17- butyrate in fatty cream base (Locoid) twice daily (n=45)	Severity: severity score at least 4.5/9 Age: adults (age 18 to 70) Sample size: 94 participants	Treatment related side effects Were only a few and similar in both groups. They included stinging, burning, itching, dryness, acne, folliculitis, and hair growth. Skin thinning No evidence	
Berth-Jones 2003 ⁽¹¹²⁾ (Green 2004 (107)) (This study is also included in the "Topical corticosteroids used proactively to prevent flares", as there is a second phase of the study when participants who have gained control of eczema are randomised to proactive treatment with topical corticosteroid or vehicle. This section only	RCT (four arms) (4 weeks treatment) (Quality using NHS CRD criteria: adequate randomisation /allocation concealment, partial blinding, and ITT used. ⁽¹⁰⁷⁾)	Intervention: Fluticasone propionate cream 0.05% once daily N=95 Intervention: Fluticasone propionate ointment 0.005% once daily N=100 Comparator: Fluticasone propionate cream 0.05% twice daily N=91 Comparator: Fluticasone propionate ointment 0.005% twice daily N=90	Severity: moderate to severe Age: children and adults (12-65 years) Sample size: 376 participants	TelangiectasiaOnce daily cream: 0/95participants (0%)Twice daily cream: 1/91participants (1%)(Difference between groups: p =0.48 °)Once daily ointment: 1/100participants (1%)Twice daily ointment: 0/90participants (0%)(Difference between groups: p =0.54 °)StriaeOnce daily cream: 0/95participants (0%)Twice daily cream: 0/91participants (0%)(Difference between groups: n/a)Once daily ointment: 1/100participants (1%)Twice daily ointment: 0/90participants (0%)(1%)Twice daily ointment: 0/90participants (0%)	

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includes safety				(Difference between groups: p =		
data from the				0.54°)		
induction of remission phase).				For the three events listed above: two of these patients had a previous history of skin changes, and therefore only one report was newly observed (group not specified in the review).		
Marchesi 1994 (113)	RCT (3 weeks treatment))	Intervention: Mometasone furoate ointment 0.1% once	Severity: at least moderate severity	Telangiectasia of mild severity in last 2 weeks Once daily: 4/30 participants	Systemic reactions None – all patients checked for blood test and value varied	
(Green 2004 ⁽¹⁰⁷⁾)	(Quality using NHS CRD criteria: method for randomisation /allocation concealment unknown, partial blinding, and no ITT used. (¹⁰⁷)) (Moher 1995 quality checklist: method and concealment of randomisation unclear, third-party blind evaluator, no dropouts/withdrawals (¹⁹)	daily (n=30) Comparator : Betamethasone dipropionate ointment 0.05% twice daily (n=30)	Age: adults Sample size: 60 participants	(13.3%) Twice daily: 5/30 participants (16.7%) (<i>Difference between groups: p</i> = 0.72°) Possible skin thinning ("Loss of <u>skin marks and reduced</u> <u>elasticity")</u> Once daily: 0/30 participants (0%) Twice daily: 1/30 participants (3.3%) (<i>Difference between groups: p</i> = 0.50°) <u>Local application site reactions</u> Did not occur	within a very narrow range.	
			Moderate potency	topical corticosteroids		
Richelli 1990 (114)	RCT	Intervention: Clobetasone 17-	Severity: not		HPA axis suppression	Adverse effects not reported
(Green 2004 (¹⁰⁷⁾)	(one week treatment	butyrate 0.05% lotion once daily at 9pm (n=9)	specified in the review		No significant difference in serum cortisol and ACTH levels before and after TCS administration in	
	(Quality using NHS CRD criteria: method for	Comparator: Clobetasone 17- butyrate 0.05% lotion twice daily at 8am and 3pm (n=13)	Age: children Sample size: 30 participants		any of the three groups, or any differences between groups	
	randomisation /allocation concealment unknown, inadequate blinding, and no ITT used. (¹⁰⁷⁾)	Comparator: Clobetasone 17- butyrate 0.05% lotion twice daily at 3pm and 8pm (n=8)				
	(Moher 1995 quality checklist: method and concealment of					

	randomisation unclear, blinding unclear, ITT unclear ⁽¹⁹⁾)					
	How	safe are topical corticoste	roids when used j	proactively to prevent flares	s ("weekend therapy")?	
Study ID (Systematic review*)	Study design and study duration (Quality assessment)	Intervention and comparator	Participants	Cutaneous adverse events	Systemic adverse events	Unspecified adverse events
		L	Potent topical cortice	osteroids versus vehicle	L	
Berth-Jones 2003 ⁽¹¹²⁾ (Schmitt 2011 ⁽¹¹⁵⁾ ; Tang 2014 ⁽⁸⁸⁾) (This study is also included under the comparison "Topical corticosteroids applied once a day compared with more frequent application" – where the induction of remission part of the study is included).	RCT (16 weeks maintenance) (Cochrane risk of bias tool: low risk of selection (sequence generation), attrition and other biases, Unclear risk of selection (allocation concealment), bias from blinding and reporting bias. ⁽¹¹⁵⁾)	Intervention: Fluticasone propionate 0.005% ointment on two consecutive days per week, once daily (n=68) Intervention: Fluticasone propionate 0.05% cream on two consecutive days per week, once daily (n=70) Comparator: Vehicle cream or ointment (n=84) Comparator: Vehicle ointment (n=73)	Severity: moderate to severe Age: 12 to 65 years Sample size (maintenance phase): 295 participants	Skin thinning No new visual signs observed in either group during maintenance phase		
Glazenburg 2009 (¹¹⁶⁾ (Schmitt 2011 (^{115]} ; Tang 2014 (⁸⁸⁾)	RCT (16 weeks maintenance) (Cochrane risk of bias tool: low risk of selection (sequence generation), and attrition bias. Unclear risk of selection (allocation concealment), bias from blinding, reporting and other biases. ⁽¹¹⁵⁾)	Intervention: Fluticasone propionate 0.005% ointment (two consecutive days per week, once daily) (n=39) Comparator: Vehicle (n=36)	Severity: moderate to severe Age: children 4-10 years Sample size (maintenance phase): 75 participants	Skin thinningNo evidence in either groupAdverse events related to treatment (cutaneous)Fluticasone: 2 events (flexural hyperpigmentation, folliculitis, transient telangiectasia) (n=39)Vehicle: 1 event (no further details reported) (n=36) (Difference between groups: p=0.61°)	Adrenal suppression No evidence in either group (measured by assessment of urinary overnight cortisol/creatinine ratios) Cancer No cases in either group	

Hanifin 2002 ⁽¹¹⁷⁾ (Schmitt 2011 ⁽¹¹⁵⁾) (Fishbein 2019 ⁽¹⁶⁾)	RCT (20 weeks maintenance) (Cochrane risk of bias tool: low risk of attrition bias. Unclear risk of selection, bias from blinding and reporting bias. High risk of other biases (noncompliance) (115))	Intervention: Fluticasone propionate 0.05% cream (once daily 4 days per week for 4 weeks, then once daily 2 days per week for 16 weeks) (n=229) Comparator: Vehicle (n=119)	Severity: moderate to severe Age: children and adults, 3 months to 65 years Sample size (maintenance phase): 348 participants	Adverse events related to treatment Fluticasone: 1/229 (one case of acne) (0.4%) Vehicle: 0/119 (0%) (Difference between groups: p=0.78°) Skin thinning No evidence (by visual skin assessment)	Possible adrenal suppressionFluticasone: 2/44* children(4.5%)Vehicle: no evidence of adrenalsuppression(measured by cosynthropinstimulation test)*One participant received 345days of treatment and had acortisol stimulation level aftertreatment of 17 ug/dL (normalwas >=18 ug/dL). The otherparticipant was treated for 280days and had a cortisolstimulation level of 9 ug/dL. Nofollow up testing.CancerNo cases	
Van der Meer 1999 ⁽¹¹⁸⁾ (Schmitt 2011 (¹¹⁵⁾)	RCT (16 weeks maintenance) (Cochrane risk of bias tool: low risk of attrition, and other biases. Unclear risk of selection, bias from blinding and reporting bias. ⁽¹¹⁵⁾ ((Moher 1995 quality checklist: method and concealment of randomisation unclear, double blinded, 17 withdrawals/dropouts, no ITT, only data up to first relapse analysed, ⁽¹⁹⁾	Intervention: Fluticasone propionate 0.005% ointment (2 consecutive days per week, once daily) (n=23) Comparator: Vehicle (n=31)	Severity: moderate to severe Age: children and adults, aged 15-50 years Sample size (maintenance phase): 54 participants	Skin thinning No evidence	Adrenal suppression No change in geometric mean cortisol levels at baseline and end of maintenance Cancer No cases	
Peserico 2008 (119) (Schmitt 2011 (115)	RCT (16 weeks maintenance) (Cochrane risk of bias tool: high risk of selection bias (sequence generation). Low risk of attrition bias and bias from blinding. Unclear	Intervention: Prednisolone aceponate 0.1% cream (two consecutive days per week, once daily) (n=112) Comparator: Vehicle (n=108)	Severity: IGA≥ moderate Age: children ≥12 years and adults Sample size (maintenance	<u>Skin thinning</u> No evidence	Cancer No cases.	Adverse events related to treatment Zero in both groups

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	risk of selection (allocation concealment) reporting and other biases. ⁽¹¹⁵⁾)	How safe a	phase): 221 participants re topical corticos	steroids used under occlusio	on?	
Study ID (Systematic review*)	Study design and study duration (Quality assessment)	Intervention and comparator	Participants	Cutaneous adverse events	Systemic adverse events	Unspecified adverse events
,	(,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,		Very potent top	ical corticosteroid		
Volden 1992 ⁽¹²⁰⁾ (Braham 2010 ⁽¹²¹⁾)	Prospective (observational) (8-18 days treatment) Risk of bias not assessed in any of the included systematic reviews.	Intervention: Dry occlusion with clobetasol propionate lotion under dry occlusion (weekly) (n=48) Comparator: No comparator	Severity: therapy resistant atopic eczema Age: adults Sample size: 48 participants	<u>Mild folliculitis</u> 2/48 participants (4%) <u>Skin thinning</u> None		
			Potent topica	l corticosteroids		
Janmohamed 2014 ⁽¹²²⁾ (Van Zuuren 2017 ⁽¹²⁾)	RCT (4 weeks treatment) (Cochrane risk of bias tool: low risk of selection (sequence generation), attrition, reporting and other biases. Unclear risk of selection (allocation concealment), performance and detection bias. ⁽¹²⁾)	Intervention: wet wrap therapy with diluted mometasone furoate 0.1% ointment (n=19) Comparator: 20% petrolatum in cetomacrogol combined with wet wrap (n=20)	Severity: severe Age: children 6 months to 10 years (mean age 3.4 years) Sample size: 39 participants	Folliculitis Mometasone under wet wrap: $9/19 (47\%)$ Emollient under wet wrap: $2/20 (10\%)$ (10%) (Difference between groups: $p = 0.03 \circ$)Severe folliculitis Mometasone under wet wrap: $1/19 (5.2\%)$ Emollient under wet wrap: $1/19 (5.2\%)$ Emollient under wet wrap: $0/0\%$ (Difference between groups: $p = 0.47*$)Secondary infected eczema Mometasone under wet wrap: $0/19 (0\%)$ Emollient under wet wrap: $2/20 (10\%)$ (Difference between groups: $p = 0.30 \circ$)Beginning of decubitus Mometasone under wet wrap: $0/19 (\%)$		

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				Emollient under wet wrap: 2/20 (10%) (Difference between groups: $p = 0.30^{a}$) Decubitus Mometasone under wet wrap: 2/19 (11%) Emollient under wet wrap: 1/20 (5%) (Difference between groups: $p = 0.53^{a}$)		
Schnopp 2002 (123) (Braham 2010 (121))	RCT, within-participant (5 days treatment) (Cochrane risk of bias tool: unclear risk of selection bias, unclear risk from blinding. ⁽³⁾)	Intervention: wet wrap therapy with mometasone furoate 0.1%, twice daily Comparator: wet wrap therapy with vehicle	Severity: exacerbated atopic eczema Age: children aged 2 to 17 years (mean 7.2 years) Sample size: 20 participants	<u>Clinical skin infections</u> None in either group		
McGowan 2003 (124) (Devillers 2006 (125))	Prospective (observational) (Up to 14 days treatment) Risk of bias not assessed in any of the included systematic reviews.	Intervention: wet wrap therapy with diluted beclomethasone dipropionate, once daily (n=8) Comparator: No comparator	Severity: not specified in the review Age: children age 3.3 to 8.8 years Sample size: 8 participants		Short term growth and bone turnover No significant differences found between outcomes before and during a median treatment period of 12 weeks (range 2-18). (assessed safety with knemometry and urinary deoxypyridinoline crosslink excretion and early morning serum cortisol).	
Wolkerstorfer 2000 ⁽¹²⁶⁾ (Braham 2010 (¹²¹⁾) (Fishbein 2019 (¹⁶⁾)	Prospective, side to side (observational) (1 week treatment) Risk of bias not assessed in any of the included systematic reviews.	Intervention: wet wrap therapy with 10-50% dilution fluticasone propionate 0.05% cream (daily) Comparator: emollient (only 2 participants) or no comparator	Severity: severe Age: children 5 months to 13 years Sample size: 18 participants	URI and/or folliculitis Fluticasone: one third of participants Furunculosis Fluticasone: one case Generalized folliculitis One case in both emollient controls Skin thinning No cases	HPA axis suppression "Nearly all" had decreased cortisol, 3 children were HPA suppressed (from Braham 2010 review). Two patients having a 9am serum cortisol < 0.2 umol/L (0.09 and 0.03) after treatment for 7 days. Those participants used 957 ug/m ² and 1125 ug/m ² of steroid cream. There was no follow up	

					testing (from Fishbein 2019 review).	
Tang 2000 ⁽¹²⁷⁾ (Braham 2010 ⁽¹²¹⁾)	Prospective (observational) ("Few days" treatment) Risk of bias not assessed in any of the included systematic reviews.	Intervention: wet wrap therapy with 10% dilution mometasone furoate 0.1% (daily for 2 to 3 hours) (n=10) Comparator: No comparator	Severity: review only reports 'facial eczema flare' Age: children (mean 8.4 years) Sample size: 10 participants	Skin thinning None Infections None		
Goodyear 1991 (128) (Braham 2010 (121))	Prospective (observational) (2 to 5 days treatment) Risk of bias not assessed in any of the included systematic reviews.	Intervention: wet wrap therapy with 25% dilution betamethasone or hydrocortisone 1%, twice daily (potent or mild potency) (n=30) Comparator: No comparator	Severity: acute erythrodermic eczema Age: children aged 9 months to 2 years (mean 5.5 years) Sample size: 30 participants	Bacterial infections Some during follow up at home	HPA axis suppression Transient low morning cortisol. During the follow up at home some adrenal suppression.	
Mallon 1994 ⁽¹²⁹⁾ (Braham 2010 (¹²¹))	Prospective (observational) (up to 5 days treatment) Risk of bias not assessed in any of the included systematic reviews.	Intervention: wet wrap therapy with 10% dilution betamethasone 0.1% cream or hydrocortisone 0.5% cream(daily) (potent or mild potency) (n=21) Comparator: No comparator	Severity: chronic severe eczema Age: children aged 4 months to 10 years (5.1 years) Sample size: 21 participants	No infections.		
Devillers 2002 (130) (Braham 2010 (121))	Retrospective side to side (observational) (1 week treatment) Risk of bias not assessed in any of the included systematic reviews.	Intervention: wet wrap therapy with diluted fluticasone propionate 0.05% (daily re-wet every 2 to3 hours) (n=26) Comparator: No comparator	Severity: refractory atopic eczema Age: children (mean 3 years), adults (mean 30 years) Sample size: 26 participants (14 children, 12 adults)	Infections 38% (n=10) had localized folliculitis, impetigo, pseudomonas, cellulitis, or purulent conjunctitivitis Skin thinning One case of striae in a patient taking inhaled steroids.	HPA axis suppression Transient low morning cortisol, 12.5% with HPA suppression	
			Moderate potency	opical corticosteroids		
Foelster-Holst 2006 ⁽¹³¹⁾	Within-participant RCT (48 to 72 hours treatment)	Intervention: wet wrap therapy with prednicarbate ointment	Severity: local SCORAD >10, severe	Zero adverse events in either group. Did not observe severe cutaneous events.	Did not observe systemic events such as growth retardation or HPA suppression – but these	

(Gonzalez-Lopez 2017 ⁽¹³²⁾)	(Cochrane risk of bias tool: unclear risk of selection and performance bias. High risk of performance bias. Unclear risk of attrition, reporting and other biases. ⁽¹³²⁾ (Cochrane risk of bias tool: unclear risk of selection bias, high risk from no blinding. ⁽³⁾)	Comparator: Prednicarbate ointment	Age: children and adults, aged 6-63 years Sample size: 24 participants		events were not actively investigated.	
	·		Mild potency to	pical corticosteroid	·	
Beattie 2004 ⁽¹³³⁾ (Gonzalez-Lopez 2017 ⁽¹³²⁾)	RCT (2 weeks treatment) (Cochrane risk of bias tool: low risk of selection, reporting and other biases. High risk of performance and attrition bias. Unclear risk of detection bias. Gonzalez-Lopez 2017) (Cochrane risk of bias tool: low risk of selection bias (sequence generation), unclear risk of selection bias (allocation concealment), unclear risk from blinding. ⁽³⁾)	Intervention: wet wrap therapy with hydrocortisone 1% twice daily then overnight the second week(n=10) Comparator: Hydrocortisone 1% twice daily then daily (n=9)	Severity: moderate Age: children < 5 years Sample size: 19 participants	Cutaneous adverse events Wet wrap therapy with hydrocortisone: 2/10 participants (20%) (2 events were folliculitis, one child withdrew) Hydrocortisone only: 0/9 participants (0%) (Difference between groups: (p=0.31 °) Did not observe severe cutaneous events.	Did not observe systemic such as growth retardation or HPA suppression – but these events were not actively investigated.	
Hindley 2006 (134) (Gonzalez-Lopez 2017 ⁽¹³²⁾)	RCT (4 weeks – not clear if treatment given for whole 4 weeks) (Cochrane risk of bias tool: low risk of selection (random sequence generation) and reporting bias. Unclear risk of selection (allocation concealment),	Intervention: wet wrap therapy with hydrocortisone 1% for 24 hours – could be reduced to 12 hours per day after first week (n=28) Comparator: Hydrocortisone 1% twice day (n=22)	Severity: SCORAD >15, moderate to severe Age: children 3 months to 5 years Sample size: 50 participants	Cutaneous adverse events Wet wrap therapy with hydrocortisone: 5/28 participants (18%) (five cases of infected eczema) Hydrocortisone only: 0/22 participants (0%) (Difference between groups: p = 0.14°)	Did not observe systemic events such as growth retardation or HPA suppression – but these events were not actively investigated.	

detection and other	Did not observe severe cutaneous	
biases. High risk of	events.	
performance and		
attrition bias. ⁽¹³²⁾)		
(Cochrane risk of bias tool: unclear risk of selection bias, low risk from blinding. ⁽³⁾)		

Footnotes:

*This column refers to the systematic review in which the safety data was extract from. The trial may have also been included in other systematic reviews, but no additional safety data was reported. Abbreviations: RCT = randomised controlled trial; TCS = topical corticosteroids; TCI = topical calcineurin inhibitors; HPA = hypothalamic pituitary adrenal, WWT = wet wrap therapy; RR = risk ratio; OR: odds ratios; 95% CI = 95% confidence interval; CHM = Chinese herbal medicine; IPA = Investigator's Global Assessment; BSA = Body Surface Area

^a P value calculated by review authors using RevMan software.

^b The P value calculated from Fisher's Exact Test was significant: 0.0298 (but in the overview, this study is included in a meta-analysis)

^c The P value calculated from Fisher's Exact Test was significant: 0.0298 (but in the overview, this study is included in a meta-analysis)

^d The P value calculated from Fisher's Exact Test was significant: 0.0352 (but in the overview, this study is included in a meta-analysis)

^e The P value calculated from Fisher's Exact Test was significant: 0.0289 (but in the overview, this study is included in a meta-analysis)

^f The P value calculated from Fisher's Exact Test was significant: 0.0267 (but in the overview, this study is included in a meta-analysis)

Where studies include "diluted" topical corticosteroids and we aren't sure how this affects the potency, we have put the topical corticosteroids in the potency classification based on the undiluted version. The terms skin atrophy and skin thinning were both used in the included reviews – for consistently we have used skin thinning throughout.

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Appendix 7: edits made to meta-analyses and data from Broeders et al 2016¹

- 1) Switched the forest plot labels around so topical corticosteroids are the intervention and topical calcineurin inhibitors are is the comparator.
- 2) Amended a data error given in the skin infection forest plot where the number of events and participants were given the wrong way round for topical corticosteroids and topical calcineurin inhibitors in Luger et al 2004².
- 3) Added skin atrophy data from Sigurgeirsson et al 2015³ into the forest plot this is not provided in the publication but is given in online correspondence on the journal website
- 4) Changed to random effects instead of fixed effects as the decision was based on whether the I² value which is not appropriate.
- 5) Bieber et al 2007 ⁴ was listed as "least potent" in table I of the publication but according to the Australian potency classification it should be classified as potent.
- 6) In table I, the topical calcineurin inhibitors given for Mandelin et al 2010 5 is tacrolimus 1% this should be 0.1%.
- 7) In table I, the therapy given for Hofman et al 2006 ⁶ was hydrocortisone acetate 0.1%. However, patients used hydrocortisone ointment 1% (mild potency) twice daily for head/neck and hydrocortisone butyrate ointment 0.1% (potent) for trunk and limbs for 2 weeks then hydrocortisone 1% twice daily for flares.

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Appendix 8 – meta-analysis of TCS versus TCI – cutaneous adverse events

		Corticosteroid	I (TCS)	Calcineurin inhibit	or (TCI)		Risk Ratio		Risk Ratio
	Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl		M-H, Random, 95% Cl
	Luger 2004 (1)	3	330	0	328	26.5%	6.96 [0.36, 134.17]		
1) Skin	Mandelin 2010 (2)	2	40	0	40	25.7%	5.00 [0.25, 100.97]		
	Reitamo 2005 (3)	2	485	0	487	25.2%	5.02 [0.24, 104.30]		
thinning	Sigurgeirsson 2015 (4)	1	1213	0	1205	22.7%	2.98 [0.12, 73.08]		
	Total (95% CI)		2068		2060	100.0%	4.86 [1.06, 22.28]		
	Total events	8		0					
	Heterogeneity: Tau ² = 0.0 Test for overall effect: Z =		f=3(P=	0.99); I² = 0%				0.01	0.1 1 10 100 More events with TCI More events with TCS

		Corticosteroid (1	CS)	Calcineurin inhibitor (1	CI)		Risk Ratio	Risk Ratio
	Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% Cl
	Doss 2009 (5)	9	279	47	287	9.1%	0.20 [0.10, 0.39]	_
	Doss 2010 (6)	6	239	18	237	6.4%	0.33 [0.13, 0.82]	
2) Skin	Hofman 2006 (7)	2	124	10	133	2.9%	0.21 [0.05, 0.96]	
	Luger 2001 (8)	4	42	22	45	5.8%	0.19 [0.07, 0.52]	
burning	Luger 2004 (1)	36	330	85	328	15.6%	0.42 [0.29, 0.60]	
	Reitamo 2002(l) (9)	24	186	113	191	14.9%	0.22 [0.15, 0.32]	
	Reitamo 2002(II) (10)	13	185	38	186	10.7%	0.34 [0.19, 0.62]	
	Reitamo 2004 (11)	30	207	50	210	14.5%	0.61 [0.40, 0.92]	
	Reitamo 2005 (3)	67	485	255	487	18.4%	0.26 [0.21, 0.33]	+
	Sikder 2005 (12)	1	15	7	15	1.8%	0.14 [0.02, 1.02]	
	Total (95% CI)		2092		2119	100.0%	0.31 [0.23, 0.40]	•
	Total events	192		645				
	Heterogeneity: Tau ² = 0	.09; Chi ² = 21.66, d	f= 9 (i	° = 0.01); I² = 58%				
	Test for overall effect: Z	= 8.44 (P < 0.0000	1)					More events with TCI More events with TCS

		Corticosteroid (T	CS)	Calcineurin inhibitor	(TCI)		Risk Ratio	Risk Ratio
	Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% Cl
	Doss 2009 (5)	9	279	12	287	4.7%	0.77 [0.33, 1.80]	+
	Doss 2010 (6)	8	239	10	237	4.1%	0.79 [0.32, 1.98]	
2) Danualture	Hofman 2006 (7)	4	124	8	133	2.5%	0.54 [0.17, 1.74]	
3) Pruritus	Luger 2001 (8)	5	42	14	45	3.9%	0.38 [0.15, 0.97]	
	Luger 2004 (1)	6	330	18	328	4.1%	0.33 [0.13, 0.82]	
	Reitamo 2002(I) (9)	18	186	29	191	11.2%	0.64 [0.37, 1.11]	
	Reitamo 2002(II) (10)	14	185	21	186	8.2%	0.67 [0.35, 1.28]	
	Reitamo 2004 (11)	33	207	45	210	20.7%	0.74 [0.50, 1.12]	
	Reitamo 2005 (3)	65	485	88	487	39.3%	0.74 [0.55, 1.00]	
	Sikder 2005 (12)	2	15	3	15	1.3%	0.67 [0.13, 3.44]	
	Total (95% CI)		2092		2119	100.0%	0.68 [0.56, 0.82]	•
	Total events	164		248				
	Heterogeneity: Tau ² = 0	.00; Chi ² = 4.80, df	= 9 (P :	= 0.85); I² = 0%				
	Test for overall effect: Z	= 4.11 (P < 0.0001))					More events with TCI More events with TCS

	Study or Subgroup	Corticosteroid Events	I (TCS) Total	Calcineurin inhibito Events	· ·	Weight	Risk Ratio M-H, Random, 95% Cl	Risk Ratio M-H, Random, 95% Cl
	Doss 2010 (6)	49	239	44	239	14.6%	1.11 [0.77, 1.60]	
	Hofman 2006 (7)	4	124	2	133	0.7%	2.15 [0.40, 11.51]	
4) Skin	Luger 2004 (1)	80	330	69	328	24.2%	1.15 [0.87, 1.53]	-
4) SKIII	Mandelin 2010 (2)	17	40	26	40	10.7%	0.65 [0.43, 1.00]	
infections	Reitamo 2002(II) (10)	4	185	4	186	1.0%	1.01 [0.26, 3.96]	
	Reitamo 2004 (11)	6	207	6	210	1.6%	1.01 [0.33, 3.09]	
	Reitamo 2005 (3)	9	485	13	487	2.8%	0.70 [0.30, 1.61]	
	Sigurgeirsson 2015 (4)	150	1213	157	1205	44.5%	0.95 [0.77, 1.17]	•
	Total (95% CI)		2823		2828	100.0%	0.98 [0.85, 1.12]	•
	Total events	319		321				
	Heterogeneity: Tau² = 0.0 Test for overall effect: Z =		= 7 (P =	0.45); I² = 0%				0.01 0.1 1 10 100 More events with TCI More events with TCS

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Appendix 9: subgroup analyses of TCS versus TCI – cutaneous adverse events

By different topical corticosteroid potencies

Skin thinning

	Corticosteroids	(TCS)	Calcineurin inhibito	r (TCI)		Risk Ratio		Risk	Ratio		
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl		M-H, Rand	lom, 95% Cl		
2.6.1 Potent											
Luger 2004 (1)	3	330	0	328	26.5%	6.96 [0.36, 134.17]			-		
Mandelin 2010 (2)	2	40	0	40	25.7%	5.00 [0.25, 100.97]			-		
Reitamo 2005 (3) Subtotal (95% CI)	2	485 855	0	487 855	25.2% 77.3%	5.02 [0.24, 104.30] 5.61 [0.99, 31.67]					
Total events	7		0								
Heterogeneity: Tau ² = 0.00; Test for overall effect: Z = 1.		: 2 (P = 0.	98); I² = 0%								
2.6.2 Moderate or mild pot	ency										
Sigurgeirsson 2015 (4) Subtotal (95% Cl)	1	1213 1213	0	1205 1205	22.7% 22.7 %	2.98 [0.12, 73.08] 2.98 [0.12, 73.08]					
Total events Heterogeneity: Not applicat	1 Die		0								
Test for overall effect: Z = 0.	67 (P = 0.50)										
Total (95% CI)		2068		2060	100.0%	4.86 [1.06, 22.28]					
Total events Heterogeneity: Tau ² = 0.00; Test for overall effect: Z = 2. Test for subgroup differenc	03 (P = 0.04)						0.01	0.1 More events with TCI	1 More even	10 ts with TC	100

Skin burning

	Corticosteroids		Calcineurin inhibitor			Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% Cl
2.2.1 Potent							
Doss 2009 (5)	9	279	47	287	9.1%	0.20 [0.10, 0.39]	_ _
Doss 2010 (6)	6	239	18	237	6.4%	0.33 [0.13, 0.82]	
Luger 2001 (7)	4	42	22	45	5.8%	0.19 [0.07, 0.52]	
Luger 2004 (1)	36	330	85	328	15.6%	0.42 [0.29, 0.60]	
Reitamo 2002(l) (8)	24	186	113	191	14.9%	0.22 [0.15, 0.32]	
Reitamo 2005 (3) Subtotal (95% CI)	67	485 1561	255	487 1575	18.4% 70.2 %	0.26 [0.21, 0.33] 0.27 [0.21, 0.35]	→
Total events	146		540				
Heterogeneity: Tau² = 0 Test for overall effect: Z			0.13); i² = 41%				
2.2.2 Potent or mild po	tency						
Hofman 2006 (9) Subtotal (95% Cl)	2	124 124	10	133 133	2.9% 2.9 %	0.21 [0.05, 0.96] 0.21 [0.05, 0.96]	
Total events Heterogeneity: Not app	2 licable		10				
Test for overall effect: Z							
2.2.3 Moderate potenc	-						
Sikder 2005 (10) Subtotal (95% CI)	1	15 15	7	15 15	1.8% 1.8 %	0.14 [0.02, 1.02] 0.14 [0.02, 1.02]	
Total events Heterogeneity: Not app	1 licable		7				
Test for overall effect: Z							
2.2.4 Mild potency							
Reitamo 2002(II) (11)	13	185	38		10.7%	0.34 [0.19, 0.62]	
Reitamo 2004 (12) Subtotal (95% Cl)	30	207 392	50	210 396	14.5% 25.1 %	0.61 [0.40, 0.92] 0.48 [0.27, 0.83]	•
Total events Heterogeneity: Tau² = 0 Test for overall effect: Z			88 0.12); I² = 59%				
Total (95% CI)		2092		2119	100.0%	0.31 [0.23, 0.40]	•
Total events	192		645			,,	Ŧ
Heterogeneity: Tau² = 0 Test for overall effect: Z	.09; Chi ² = 21.66,						0.01 0.1 1 10 1
			P = 0.27). I ² = 23.4%				More events with TCI More events with TCS

Pruritus

	Corticosteroids		Calcineurin inhibito			Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% Cl
2.3.1 Potent							
Doss 2009 (5)	9	279	12	287	4.7%	0.77 [0.33, 1.80]	
Doss 2010 (6)	8	239	10	237	4.1%	0.79 [0.32, 1.98]	
_uger 2001 (7)	5	42	14	45	3.9%	0.38 [0.15, 0.97]	
_uger 2004 (1)	6	330	18	328	4.1%	0.33 [0.13, 0.82]	
Reitamo 2002(l) (8)	18	186	29	191	11.2%	0.64 [0.37, 1.11]	
Reitamo 2005 (3) Subtotal (95% Cl)	65	485 1561	88	487 1575	39.3% 67.4%	0.74 [0.55, 1.00] 0.67 [0.53, 0.84]	•
Fotal events	111		171				
Heterogeneity: Tau² = (Fest for overall effect: Z			0.49); I² = 0%				
2.3.2 Potent or mild po	tency						
Hofman 2006 (9) Subtotal (95% Cl)	4	124 124	8	133 133	2.5% 2.5 %	0.54 [0.17, 1.74] 0.54 [0.17, 1.74]	
Fotal events Heterogeneity: Not app	4 Jicable		8				
Fest for overall effect: Z							
2.3.3 Moderate potenc	y						
Sikder 2005 (10) Subtotal (95% Cl)	2	15 15	3	15 15	1.3% 1.3 %	0.67 [0.13, 3.44] 0.67 [0.13, 3.44]	
Fotal events	2		3				
Heterogeneity: Not app Fest for overall effect: Z							
2.3.4 Mild potency							
Reitamo 2002(II) (11)	14	185	21	186	8.2%	0.67 [0.35, 1.28]	
Reitamo 2002(ii) (11)	33	207	45	210	20.7%	0.74 [0.50, 1.12]	
Subtotal (95% CI)	55	392	45	396	28.9%	0.72 [0.51, 1.02]	•
Fotal events	47		66				•
Heterogeneity: Tau² = (Fest for overall effect: Z	0.00; Chi² = 0.07, d	lf=1 (P=					
fotal (95% CI)		2092		2119	100.0%	0.68 [0.56, 0.82]	•
Fotal events	164		248				
Heterogeneity: Tau ² = (0.00; Chi ^z = 4.80, d	f= 9 (P =	0.85); I² = 0%				
Fest for overall effect: Z	= 4.11 (P < 0.000	1)					0.01 0.1 1 10 1 More events with TCI More events with TCS

Skin infections

	Corticosteroid		Calcineurin inhibit			Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% Cl
2.7.1 Potent							
Doss 2010 (6)	49	239	44	239	14.6%	1.11 [0.77, 1.60]	
Luger 2004 (1)	80	330	69	328	24.2%	1.15 [0.87, 1.53]	
Mandelin 2010 (2)	17	40	26	40	10.7%	0.65 [0.43, 1.00]	
Reitamo 2005 (3)	9	485	13	487	2.8%	0.70 [0.30, 1.61]	
Subtotal (95% CI)		1094		1094	52.2%	0.94 [0.70, 1.26]	•
Total events	155		152				
Heterogeneity: Tau² = 0.04		= 3 (P = 0	.12); I² = 49%				
Test for overall effect: Z = 0	0.41 (P = 0.68)						
2.7.2 Potent or mild poter	ю						
Hofman 2006 (9)	4	124	2	133	0.7%	2.15 [0.40, 11.51]	
Subtotal (95% CI)		124		133	0.7%	2.15 [0.40, 11.51]	
Total events	4		2				
Heterogeneity: Not applica	able						
Test for overall effect: Z = (0.89 (P = 0.37)						
2.7.3 Moderate or mild po	tency						
Sigurgeirsson 2015 (4)	150	1213	157	1205	44.5%	0.95 [0.77, 1.17]	
Subtotal (95% CI)		1213		1205	44.5%	0.95 [0.77, 1.17]	+
Total events	150		157				
Heterogeneity: Not applica	able						
Test for overall effect: Z = (0.49 (P = 0.62)						
2.7.4 Mild potency							
Reitamo 2002(II) (11)	4	185	4	186	1.0%	1.01 [0.26, 3.96]	
Reitamo 2004 (12)	6	207	6	210	1.6%	1.01 [0.33, 3.09]	
Subtotal (95% CI)		392		396	2.6%	1.01 [0.43, 2.40]	
Total events	10		10				
Heterogeneity: Tau ² = 0.00); Chi² = 0.00, df:	= 1 (P = 0	.99); I² = 0%				
Test for overall effect: Z = 0	0.02 (P = 0.98)						
Total (95% CI)		2823		2828	100.0%	0.98 [0.85, 1.12]	•
Total events	319		321				
Heterogeneity: Tau ² = 0.00); Chi [≈] = 6.80, df÷	= 7 (P = 0	.45); I² = 0%				0.1 0.2 0.5 1 2 5 10
Test for overall effect: Z = (0.1 0.2 0.5 1 2 5 1 More events with TCI More events with TCS
Test for subaroup differen		4f = 0.7D	- 0.003 17 - 004				More events with FCF more events with FCS

By age of participants (children or adults)

Skin thinning

	Corticosteroids	(TCS)	Calcineurin inhibito	r (TCI)		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% Cl
3.6.1 Adults							
Luger 2004 (1)	3	330	0	328	26.5%	6.96 [0.36, 134.17]	
Mandelin 2010 (2)	2	40	0	40	25.7%	5.00 [0.25, 100.97]	
Reitamo 2005 (3) Subtotal (95% CI)	2	485 855	0	487 855	25.2% 77.3%	5.02 [0.24, 104.30] 5.61 [0.99, 31.67]	
Total events	7		0				
Heterogeneity: Tau ² = 0.00 Test for overall effect: Z = 7 3.6.2 Children							
Sigurgeirsson 2015 (4) Subtotal (95% CI)	1	1213 1213	0	1205 1205	22.7% 22.7 %	2.98 [0.12, 73.08] 2.98 [0.12, 73.08]	
Total events Heterogeneity: Not applica Test for overall effect: Z = (0				
Total (95% CI)		2068		2060	100.0%	4.86 [1.06, 22.28]	
Total events Heterogeneity: Tau ² = 0.00 Test for overall effect: Z = 2 Test for subgroup differen	2.03 (P = 0.04)						0.005 0.1 1 10 20 More events with TCI More events with TCS

Skin burning

	Corticosteroid	s (TCS)	Calcineurin inhibito	r (TCI)		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% Cl
3.2.1 Adults							
Doss 2009 (5)	9	279	47	287	9.1%	0.20 [0.10, 0.39]	_ -
Luger 2001 (7)	4	42	22	45	5.8%	0.19 [0.07, 0.52]	
Luger 2004 (1)	36	330	85	328	15.6%	0.42 [0.29, 0.60]	
Reitamo 2002(l) (8)	24	186	113	191	14.9%	0.22 [0.15, 0.32]	
Reitamo 2005 (3) Subtotal (95% CI)	67	485 1322	255	487 1338	18.4% 63.7%	0.26 [0.21, 0.33] 0.27 [0.20, 0.36]	→
Total events	140		522				-
Heterogeneity: Tau ² = 0. Test for overall effect: Z			0.08); I² = 52%				
3.2.2 Children							
Doss 2010 (6)	6	239	18	237	6.4%	0.33 [0.13, 0.82]	_
Hofman 2006 (9)	2	124	10	133	2.9%	0.21 [0.05, 0.96]	
Reitamo 2002(II) (11)	13	185	38	186	10.7%	0.34 [0.19, 0.62]	
Reitamo 2004 (12)	30	207	50	210	14.5%	0.61 [0.40, 0.92]	
Sikder 2005 (10)	1	15	7	15	1.8%	0.14 [0.02, 1.02]	
Subtotal (95% CI)		770		781	36.3%	0.40 [0.26, 0.62]	◆
Total events	52		123				
Heterogeneity: Tau² = 0. Test for overall effect: Z			0.22); I ^z = 30%				
Total (95% CI)		2092		2119	100.0%	0.31 [0.23, 0.40]	•
Total events	192		645				
Heterogeneity: Tau ² = 0.	.09; Chi ² = 21.66,	df = 9 (P	= 0.01); I² = 58%				
Test for overall effect: Z	= 8.44 (P < 0.000	01)					0.01 0.1 1 10 11 More events with TCI More events with TCS
Test for subgroup differ	ences: Chi ² = 2.4	8, df = 1 (i	P = 0.11), I² = 59.8%				More events with FGT More events with FGS

Pruritus

	Corticosteroids	s (TCS)	Calcineurin inhibito	r (TCI)		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% Cl
3.3.1 Adults							
Doss 2009 (5)	9	279	12	287	4.7%	0.77 [0.33, 1.80]	
Luger 2001 (7)	5	42	14	45	3.9%	0.38 [0.15, 0.97]	
Luger 2004 (1)	6	330	18	328	4.1%	0.33 [0.13, 0.82]	
Reitamo 2002(I) (8)	18	186	29	191	11.2%	0.64 [0.37, 1.11]	
Reitamo 2005 (3)	65	485	88	487	39.3%	0.74 [0.55, 1.00]	
Subtotal (95% CI)		1322		1338	63.2%	0.65 [0.50, 0.84]	◆
Total events	103		161				
Heterogeneity: Tau ² = (0.01; Chi ² = 4.28, d	if = 4 (P =	0.37); I² = 7%				
Test for overall effect: 2	z = 3.34 (P = 0.000	8)					
3.3.2 Children							
Doss 2010 (6)	8	239	10	237	4.1%	0.79 [0.32, 1.98]	
Hofman 2006 (9)	4	124	8	133	2.5%	0.54 [0.17, 1.74]	
Reitamo 2002(II) (11)	14	185	21	186	8.2%	0.67 [0.35, 1.28]	
Reitamo 2004 (12)	33	207	45	210	20.7%	0.74 [0.50, 1.12]	
Sikder 2005 (10)	2	15	3	15	1.3%	0.67 [0.13, 3.44]	
Subtotal (95% CI)		770		781	36.8%	0.71 [0.53, 0.97]	\bullet
Total events	61		87				
Heterogeneity: Tau ² = (0.00; Chi ^z = 0.36, d	if = 4 (P =	0.99); I² = 0%				
Test for overall effect: 2	Z = 2.17 (P = 0.03)						
Total (95% CI)		2092		2119	100.0%	0.68 [0.56, 0.82]	◆
Total events	164		248				
Heterogeneity: Tau ² = (0.00; Chi ² = 4.80, d	if = 9 (P =	0.85); I² = 0%				0.02 0.1 1 10 5
Test for overall effect: 2	4.11 (P < 0.000	1)					U.U2 U.1 1 1U 5 More events with TCI More events with TCS
Test for subaroup diffe	rences: Chi ² = 0.2	3, df = 1 (l	P = 0.63), I ² = 0%				More events with FCF. More events with FCS

Skin infections

	Corticosteroids	s (TCS)	Calcineurin inhibito	or (TCI)		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% Cl
3.7.1 Adults							
Luger 2004 (1)	80	330	69	328	24.2%	1.15 [0.87, 1.53]	-
Mandelin 2010 (2)	17	40	26	40	10.7%	0.65 [0.43, 1.00]	
Reitamo 2005 (3)	9	485	13	487	2.8%	0.70 [0.30, 1.61]	
Subtotal (95% CI)		855		855	37.7%	0.85 [0.55, 1.32]	•
Total events	106		108				
Heterogeneity: Tau ² = 0.0)9; Chi ² = 5.35, df :	= 2 (P = 0.	07); I² = 63%				
Test for overall effect: Z =	0.70 (P = 0.48)						
3.7.2 Children							
Doss 2010 (6)	49	239	44	239	14.6%	1.11 [0.77, 1.60]	- - -
Hofman 2006 (9)	4	124	2	133	0.7%	2.15 [0.40, 11.51]	
Reitamo 2002(II) (11)	4	185	4	186	1.0%	1.01 [0.26, 3.96]	
Reitamo 2004 (12)	6	207	6	210	1.6%	1.01 [0.33, 3.09]	
Sigurgeirsson 2015 (4)	150	1213	157	1205	44.5%	0.95 [0.77, 1.17]	+
Subtotal (95% CI)		1968		1973	62.3%	1.00 [0.84, 1.19]	•
Total events	213		213				
Heterogeneity: Tau ² = 0.0	00; Chi ² = 1.37, df :	= 4 (P = 0.	85); I² = 0%				
Test for overall effect: Z =	0.04 (P = 0.97)						
Total (95% CI)		2823		2828	100.0%	0.98 [0.85, 1.12]	•
Total events	319		321				
Heterogeneity: Tau ² = 0.0	00; Chi ² = 6.80, df:	= 7 (P = 0.	45); I ² = 0%				
Test for overall effect: Z =	0.33 (P = 0.74)						0.01 0.1 1 10 100 More events with TCI More events with TCS
Test for subgroup differe	nces: Chi ² = 0.41,	df = 1 (P :	= 0.52), I² = 0%				More events with rich More events with richs

By individual topical calcineurin inhibitor (TCI)

Skin thinning

	Corticosteroid	s (TCS)	Calcineurin inhibite	or (TCI)		Risk Ratio		Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl		M-H, Random, 95% Cl
4.6.1 Tacrolimus 0.1%								
Mandelin 2010 (2)	2	40	0	40	25.7%	5.00 [0.25, 100.97]		
Reitamo 2005 (3)	2	485	0	487	25.2%	5.02 [0.24, 104.30]		
Subtotal (95% CI)		525		527	50.9 %	5.01 [0.59, 42.37]		
Total events	4		0					
Heterogeneity: Tau ² = 0.0	10; Chi² = 0.00, df:	= 1 (P = 1.	.00); I ² = 0%					
Test for overall effect: Z =	1.48 (P = 0.14)							
4.6.2 Pimecrolimus 1%								
Luger 2004 (1)	3	330	0	328	26.5%	6.96 [0.36, 134.17]		
Sigurgeirsson 2015 (4)	1	1213	0	1205	22.7%	2.98 [0.12, 73.08]		
Subtotal (95% CI)		1543		1533	49.1%	4.71 [0.54, 41.33]		
Total events	4		0					
Heterogeneity: Tau ² = 0.0	10; Chi ² = 0.15, df:	= 1 (P = 0.	.70); I² = 0%					
Test for overall effect: Z =	1.40 (P = 0.16)							
Total (95% CI)		2068		2060	100.0%	4.86 [1.06, 22.28]		
Total events	8		0					
Heterogeneity: Tau ² = 0.0	10; Chi² = 0.15, df:	= 3 (P = 0.	.99); I² = 0%				-	
Test for overall effect: Z =	2.03 (P = 0.04)						0.01	0.1 1 10 100 More events with TCI More events with TCS
Ta ak ƙasa sa ka sana sa si ƙƙasar		-16 4 (D	0.070.17.000					More events with For More events with Fos

Test for subgroup differences: Chi² = 0.00, df = 1 (P = 0.97), l² = 0%

Skin burning

	Corticosteroids	s (TCS)	Calcineurin inhibitor	(TCI)		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% Cl
4.2.1 Tacrolimus 0.1%							
Doss 2009 (5)	9	279	47	287	9.1%	0.20 [0.10, 0.39]	_
Reitamo 2002(I) (8)	24	186	113	191	14.9%	0.22 [0.15, 0.32]	
Reitamo 2002(II) (11)	13	185	38	186	10.7%	0.34 [0.19, 0.62]	_ -
Reitamo 2005 (3)	67	485	255	487	18.4%	0.26 [0.21, 0.33]	+
Subtotal (95% CI)		1135		1151	53.0%	0.25 [0.21, 0.31]	•
Total events	113		453				
Heterogeneity: Tau ² = 0	.00; Chi ² = 2.19, d	if = 3 (P =	0.53); I ² = 0%				
Test for overall effect: Z	= 14.49 (P < 0.00	001)					
4.2.2 Tacrolimus 0.03%	1						
Doss 2010 (6)	6	239	18	237	6.4%	0.33 [0.13, 0.82]	-
Hofman 2006 (9)	2	124	10	133	2.9%	0.21 [0.05, 0.96]	
Reitamo 2004 (12)	30	207	50	210	14.5%	0.61 [0.40, 0.92]	
Sikder 2005 (10)	1	15	7	15	1.8%	0.14 [0.02, 1.02]	
Subtotal (95% Cl)		585		595	25.6%	0.40 [0.22, 0.73]	◆
Total events	39		85				
Heterogeneity: Tau ² = 0	.13; Chi ² = 4.60, d	if = 3 (P =	0.20); I ² = 35%				
Test for overall effect: Z	= 3.00 (P = 0.003)					
4.2.3 Pimecrolimus 1%							
Luger 2001 (7)	4	42	22	45	5.8%	0.19 [0.07, 0.52]	
Luger 2004 (1)	36	330	85	328	15.6%	0.42 [0.29, 0.60]	
Subtotal (95% CI)		372		373	21.4%	0.33 [0.16, 0.67]	◆
Total events	40		107				
Heterogeneity: Tau ² = 0	.16; Chi ² = 2.12, d	if = 1 (P =	0.15); I² = 53%				
Test for overall effect: Z	= 3.08 (P = 0.002)					
Total (95% CI)		2092		2119	100.0%	0.31 [0.23, 0.40]	•
Total events	192		645				
Heterogeneity: Tau ² = 0	.09; Chi ² = 21.66,	df = 9 (P	= 0.01); I² = 58%				0.01 0.1 1 10 1
Test for overall effect: Z	= 8.44 (P < 0.000	01)					More events with TCI More events with TCS
Fest for subaroun differ	oncos: Chiž – 24	3 df = 27	P = 0.30\ IZ = 17.9%				more events with for more events with fos

Test for subgroup differences: Chi² = 2.43, df = 2 (P = 0.30), l² = 17.8%

Pruritus

	Corticosteroids	s (TCS)	Calcineurin inhibi	tor (TCI)		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% Cl
4.3.1 Tacrolimus 0.1%							
Doss 2009 (5)	9	279	12	287	4.7%	0.77 [0.33, 1.80]	
Reitamo 2002(l) (8)	18	186	29	191	11.2%	0.64 [0.37, 1.11]	
Reitamo 2002(II) (11)	14	185	21	186	8.2%	0.67 [0.35, 1.28]	
Reitamo 2005 (3)	65	485	88	487	39.3%	0.74 [0.55, 1.00]	.
Subtotal (95% CI)		1135		1151	63.4%	0.71 [0.57, 0.90]	•
Total events	106		150				
Heterogeneity: Tau² = 0.			0.96); I² = 0%				
Test for overall effect: Z =	= 2.84 (P = 0.005)					
4.3.2 Tacrolimus 0.03%							
Doss 2010 (6)	8	239	10	237	4.1%	0.79 [0.32, 1.98]	
Hofman 2006 (9)	4	124	8	133	2.5%	0.54 [0.17, 1.74]	
Reitamo 2004 (12)	33	207	45	210	20.7%	0.74 [0.50, 1.12]	
Sikder 2005 (10)	2	15	3	15	1.3%	0.67 [0.13, 3.44]	
Subtotal (95% CI)		585		595	28.5%	0.73 [0.51, 1.03]	•
Total events	47		66				
Heterogeneity: Tau² = 0.		lf = 3 (P =	0.96); I² = 0%				
Test for overall effect: Z =	= 1.81 (P = 0.07)						
4.3.3 Pimecrolimus 1%							
Luger 2001 (7)	5	42	14	45	3.9%	0.38 [0.15, 0.97]	
Luger 2004 (1)	6	330	18	328	4.1%	0.33 [0.13, 0.82]	
Subtotal (95% CI)		372		373	8.1%	0.36 [0.19, 0.68]	◆
Total events	11		32				
Heterogeneity: Tau² = 0.			0.83); I² = 0%				
Test for overall effect: Z =	= 3.11 (P = 0.002)					
Total (95% CI)		2092		2119	100.0%	0.68 [0.56, 0.82]	•
Total events	164		248				
Heterogeneity: Tau ² = 0.	00; Chi ² = 4.80, d	f= 9 (P =	0.85); I² = 0%				0.01 0.1 1 10 10
Test for overall effect: Z =	= 4.11 (P < 0.000	1)					0.01 0.1 1 10 10 More events with TCI More events with TCS
est for subaroup differe	ences: Chi ² = 4.1:	3, df = 2 (l	P = 0.13), P = 51.69	6			more events with FGF more events with FCS

Skin infections

	Corticosteroids		Calcineurin inhibitor			Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% Cl
4.7.1 Tacrolimus 0.1%							
Mandelin 2010 (2)	17	40	26	40	10.7%	0.65 [0.43, 1.00]	
Reitamo 2002(II) (11)	4	185	4	186	1.0%	1.01 [0.26, 3.96]	
Reitamo 2005 (3)	9	485	13	487	2.8%	0.70 [0.30, 1.61]	
Subtotal (95% CI)		710		713	14.5%	0.68 [0.47, 0.98]	\bullet
Total events	30		43				
Heterogeneity: Tau ² = 0.0	0; Chi ² = 0.36, df =	= 2 (P = 0	.83); I² = 0%				
Test for overall effect: Z =	2.05 (P = 0.04)						
4.7.2 Tacrolimus 0.03%							
Doss 2010 (6)	49	239	44	239	14.6%	1.11 [0.77, 1.60]	+-
Hofman 2006 (9)	4	124	2	133	0.7%	2.15 [0.40, 11.51]	
Reitamo 2004 (12)	6	207	6	210	1.6%	1.01 [0.33, 3.09]	
Subtotal (95% CI)		570		582	16.8%	1.13 [0.81, 1.59]	*
Total events	59		52				
Heterogeneity: Tau ² = 0.0	0; Chi ² = 0.60, df =	= 2 (P = 0	.74); I ² = 0%				
Test for overall effect: Z =	0.73 (P = 0.47)						
4.7.3 Pimecrolimus 1%							
Luger 2004 (1)	80	330	69	328	24.2%	1.15 [0.87, 1.53]	+
Sigurgeirsson 2015 (4)	150	1213	157	1205	44.5%	0.95 [0.77, 1.17]	+
Subtotal (95% CI)		1543		1533	68.7%	1.02 [0.85, 1.23]	◆
Total events	230		226				
Heterogeneity: Tau ² = 0.0	0; Chi ² = 1.17, df =	= 1 (P = 0	.28); I² = 15%				
Test for overall effect: Z =	0.22 (P = 0.83)						
Total (95% CI)		2823		2828	100.0%	0.98 [0.85, 1.12]	•
Total events	319		321				
Heterogeneity: Tau ² = 0.0	0; Chi² = 6.80, df =	7 (P = 0	.45); I² = 0%				0.01 0.1 1 10 100
Test for overall effect: Z =	0.33 (P = 0.74)						U.U1 U.1 1 10 100 More events with TCI More events with TCS
Test for subgroup differer	nces: Chi² = 4.65,	df = 2 (P	= 0.10), I² = 57.0%				More events with LCL More events with LCS

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