SPECIAL ARTICLE

Consensus on the use of oral isotretinoin in dermatology - Brazilian Society of Dermatology

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Abstract

Background: Isotretinoin is a synthetic retinoid, derived from vitamin A, with multiple mechanisms of action and highly effective in the treatment of acne, despite common adverse events, manageable and dose-dependent. Dose-independent teratogenicity is the most serious. Therefore, off-label prescriptions require strict criteria.

Objective: To communicate the experience and recommendation of Brazilian dermatologists on oral use of the drug in dermatology.

Methods: Eight experts from five universities were appointed by the Brazilian Society of Dermatology to develop a consensus on indications for this drug. Through the adapted DELPHI methodology, relevant elements were listed and an extensive analysis of the literature was carried out. The consensus was defined with the approval of at least 70% of the experts.

Results: With 100% approval from the authors, there was no doubt about the efficacy of oral isotretinoin in the treatment of acne, including as an adjunct in the correction of scars. Common and manageable common adverse events are mucocutaneous in nature. Others, such as growth retardation, abnormal healing, depression, and inflammatory bowel disease have been

KEYWORDS

Acne vulgaris; Dermatitis, seborrheic; Isotretinoin; Rosacea; Vitamin A


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Introduction

Oral isotretinoin (13-cis-retinoic acid) is a retinoid, derived from vitamin A. It was synthesized in 1955, but it was only in 1973 that studies on its use in psoriasis, genetic disorders of keratinization, cystic acne, and basal cell carcinoma began. In the 1980s, it became the most effective option for treating nodular-cystic acne, and is currently indicated for moderate forms resistant to other treatments. It was approved for acne in the United States in 1982, in 1983 in Europe, and in 1990 in Brazil, revolutionizing the treatment of severe forms of acne.1–5

Clinical (mucocutaneous) and laboratory (liver function and lipid profile) side effects are dose-dependent, predictable, manageable and reversible, except for teratogenicity. Acne is the only approved indication, although many off-label uses have been reported.6–9

Isotretinoin acts as a prodrug, being converted into all-trans-retinoic acid (ATRA) in the cytoplasm of cells to be transported to the nucleus, where it binds to the nuclear retinoic acid receptor (RAR and RXR), isoforms a, b, and g.9 The known mechanisms of action are normalization of infundibular hyperkeratinization, inhibition of the production of cytokeratins 1, 10, and 14, filaggrin and matrix metalloproteinases (MMPs), and increase of cytokeratins 7, 13, and 19, lamin B1, and IL-1. Effects on proliferation, differentiation, apoptosis, and cell renewal, in addition to immunomodulation, are related to the regulation of gene expression, influencing nuclear transcription factors. There is activation of some genes (tumor suppressors or apoptotic, such as p53 and BAX and coding for collagen and fibronectin production) and inhibition of others (involved in lipid metabolism).10–12 On apoptosis, ATRA increases the expression of the forkhead box O3 transcription factor (FOXO3), activates the tumor necrosis factor-related apoptosis-inducing ligand (TRAIL) pathway and produces FOXO1 caspases, interrupting the cell cycle, by expressing the genes p21, 27 and 53.13–16 Activation of FOXO1, a negative co-regulator of the androgen receptor, peroxisome proliferator-activated receptor-gamma (PPAR gamma), and liver X receptor-[alpha] sterol response element binding protein-1c (SREBP-1c), reduces lipogenesis. The attenuation of the mechanistic target of rapamycin complex 1 (mTORC1) stimulates the expression of PPAR gamma and SREBP-1c.16 Due to the negative regulation of genes related to insulin-like growth factor 1 (IGF1)/phosphatidylinositol 3-kinase (PI3K)/AKT (protein kinase B)/mTORC1 pathway and positive regulation of those responsible for FOXO1 and FOXO3/TRAIL/caspase pathways, there is suppression of sebogenesis and apoptosis of sebocytes.16–18 The activation of the p53 pathways represents the interconnection between the signaling pathways regulated positively or negatively by isotretinoin. The BAX protein induces apoptosis of keratinocytes with mutations induced by UV radiation; its expression is reduced by isotretinoin due to its anti-carcinogenic action.14 It is the only drug that alone acts on the four etiopathogenic factors of acne: it reduces acroinfundibular hyperkeratinization and comedogenesis; suppresses sebogenesis, by reducing the size and activity of sebaceous glands by up to 90%; decreases the population of Cutibacterium acnes (C. acnes), formerly called Propionibacterium acnes (P. acnes) due to changes in the follicular microenvironment; and modulates inflammation by the negative regulation of toll-like 2 and 4 membrane receptors (TLR-2 and 4) in keratinocytes, sebocytes, monocytes, corneal cells, and immune cells.18–22 These receptors are activated by identification of the molecular patterns of C. acnes and, when inhibited, there is downregulation of the nuclear factor kappa B (NF-κB) pathways, which triggers the production of cytokines (IL-8, IL-1β, IL-17, IFNγ) and activator protein 1 (AP-1) responsible for the synthesis of MMPs.13,20 After 40 years, it is believed that not all mechanisms of action on the skin and other organs are elucidated. However, existing knowledge explains the efficacy in acne vulgaris and adverse events, justifying unapproved, off-label indications.23–25

The actions of isotretinoin in different cells, explaining its beneficial effects and adverse events, are summarized in Table 1.

The purpose of this article is to present a consensus on the effects of oral isotretinoin on the skin and its indications for acne and other conditions not yet approved, despite the existence of relevant data in the consulted literature.

Methods

Eight dermatologists, experts in isotretinoin, were nominated to reach a consensus on the use of oral isotretinoin in dermatology, following the adapted DELPHI methodology. In the first phase, relevant topics were discussed and the text was structured, each author responsible for a different topic; in the second phase, a bibliographic review and drafting of the texts was carried out. The databases consulted were as follows: Cochrane Skin Group Specialized
Register, Cochrane Library, MEDLINE, PubMed, Embase, and LILACS. The literature in Portuguese, English, and Spanish was searched using the following keywords isotretinoina, aliança terapêutica, dermatologia, acne vulgar, dermatite seborreica, rosácea, psoriase, ceratose actínica, envelhecimento da pele, dermatology, oral isotretinoin, off label use, off label prescribing, acne, skin diseases, rosacea, photoaging of skin, actinic keratosis, seborrheic dermatitis, psoriasis, alopecia. The first author and the mentor were responsible for compiling a single text and sending it for review by the others. In the third phase, both authors assessed the consensus on the texts; what reached 70% consensus remained in the final version.

Results/Discussion

Two topics on oral isotretinoin in dermatology were defined: acne vulgaris and relevant, off-label indications (inflammatory diseases of the skin and scalp, photoaging, and field cancerization). The results and discussion of each topic are presented below.

Acne vulgaris

Isotretinoin is the only drug that acts on all etiopathogenic factors of acne vulgaris, remaining the only monotherapy capable of providing prolonged remission or cure in up to 80% of patients, with one treatment cycle.

Acne vulgaris, a chronic, immune-mediated, multifactorial inflammatory disease that affects the pilosebaceous unit is among the three most prevalent dermatoses worldwide. It can generate physical (scarring) and psychological sequelae, second only to eczema. It affects 80% to 90% of the world population at some stage in life, with a peak prevalence between 16–20 years. According to a survey among members of the Brazilian Society of Dermatology and other epidemiological studies, acne vulgaris is the leading cause of dermatology consultations.

The clinical effectiveness of oral isotretinoin is superior to other acne treatments, promoting healing or prolonged remission, improving quality of life, and reducing psychosocial damage; however, adverse effects are observed in up to 90% of patients. Figs. 1 and 2 illustrate patients treated with oral isotretinoin, with healing of acne and absence of recurrence after two years of follow-up. Some controversies about rare and serious events, particularly depression, suicide, and inflammatory bowel disease (IBD), have not been proven to be causally associated. It was approved for severe acne (conglobata and nodular-cystic), but evidence demonstrated in controlled randomized clinical trials (CRCTs) since 1980, in systematic reviews (SRs), consensus, and recommendations of dermatology societies allowed to expand the indication for nodular-cystic and moderate papulopustular forms resistant to other treatments, with tendency to scarring, and emotional and social functional impairment. A single course of the drug leads to cure in two-thirds of patients. Recurrences may take place, but they are milder and manageable with topical treatments. Some characteristics of the disease favor recurrence and retreatment.

Table 2 presents indications, contraindications, and warnings for use, in addition to the acne characteristics related to the need for retreatment. Despite 31 CRCTs, two SRs concluded that studies with better methodology and less heterogeneous as to the efficacy outcomes are needed, particularly comparative studies between the use of oral isotretinoin and the use of oral antibiotics associated to topical agents (combinations of retinoid and benzoyl peroxide).

In addition, studies should include a greater number of participants, mainly females, prepubescents, and patients with trunk involvement; the long-term efficacy outcomes must be assessed, especially the superiority of this drug in terms of cure or prolonged remission of acne.

The approved dose in the package insert is 0.5 to 1 mg/kg/day, taken after meals, due to the lipophilic character of the molecule, except for the isotretinoin-lidose variant, not available in Brazil, which can be administered while fasting. A CRCT compared the daily dose at two meals vs. single dose; no difference was observed in efficacy, but adverse effects were more frequent with the use of a single dose. However, there is a preference for a single dose, due to greater treatment adherence. An SR analyzed CRCT with different daily doses and therapeutic regimens. The effectiveness was greater among groups that received a conventional or low dose (<0.5 mg/kg/day) daily when compared with intermittent use, in monthly pulses or alternate days.

Mild adverse effects were more frequently observed with daily and continuous use, under low or conventional doses. Intermittent use was less effective and is not recommended. Recently, studies have shown a tendency toward lower daily doses (0.1–0.5 mg/kg, up to 5 mg) for moderate acne, with a longer duration, up to 18 months, presenting less adverse events, better tolerability, and recurrence rates similar to those observed with conventional dose, maintaining treatment for two to four months after total lesion resolution. Prolonged duration is necessary in severe cases and in extra-facial involvement. The approved total dose (120 to 150 mg/kg) is maintained in clinical studies, consensus, and dermatological practice; however, there has never been a CRCT-based rationale. Studies with a better methodology have shown that a fixed total dose is not the best reference for the duration of treatment, which must con...
The and with cants, are maintained, with improvement and recurrence after two to three months. During the last cycle, the clinical picture worsened. The patient was treated with oral isotretinoin, 40 mg/kg/day (0.6 mg/kg/day), with total lesion regression after four months and maintenance for another month (total dose = 100 mg/kg/day) – regimen based on recent publications.\textsuperscript{77,80-82} Photos before and after treatment with oral isotretinoin. Maintenance treatment with adapalene 0.1% gel, for 12 months. There was no recurrence.

Figure 1 18-year-old teenager, with moderate inflammatory acne on the face and trunk for four years, presenting scars, with a relevant negative impact on quality of life. The patient had been submitted to four cycles of oral cyclin, associated with topical combination of benzoyl peroxide and adapalene, with improvement and recurrence after two to three months. During the last cycle, the clinical picture worsened. The patient was treated with oral isotretinoin, 40 mg/kg/day (0.6 mg/kg/day), with total lesion regression after four months and maintenance for another month (total dose = 100 mg/kg/day) – regimen based on recent publications.\textsuperscript{77,80-82} Photos before and after treatment with oral isotretinoin. Maintenance treatment with adapalene 0.1% gel, for 12 months. There was no recurrence.

Figure 2 A 22-year-old patient with acne conglobata on the face alone for 15 months. Previously treated with oral antibiotics and topical products (whose names the patient was unable to report), without improvement. Treatment with isotretinoin 20 mg/day (0.3 mg/kg/day) and prednisone 40, 30, 20, and 10 mg/day every seven days was initiated. The duration of treatment, always with the same daily dose, was 18 months (160 mg/kg), until complete resolution of the lesions. A maintenance treatment with benzoyl peroxide 5% was maintained for 12 months. There was no recurrence.

consider individual conditions, regression of the disease, and maintenance for two to four months after total resolution of the lesions.\textsuperscript{40,71,77,81,82,84,85}

Dose-dependent mucocutaneous clinical adverse events are common, such as cheilitis, which affects 90–100% of patients, and cutaneous, ocular, and nasal mucosa xerosis. They are manageable with the use of lip lubricants, and ocular and nasal emollients, and they regress with dose reduction or treatment suspension.\textsuperscript{16} Other rare events include alopecia, pyogenic granuloma, photosensitivity, arthralgia, myalgia, headache, anorexia, insomnia, and irritability.\textsuperscript{70} The most serious risk is teratogenicity, which is dose-independent.\textsuperscript{4,40} Pregnancy can have a normal course in 65–85% of cases, but there is a risk of miscarriage (10.9–20%) and embryopathies (18–28%), with craniofacial, central nervous system, thymus, and cardiovascular anomalies.\textsuperscript{4-7} The possibility of pregnancy must be ruled out (by testing and waiting for menstruation); prescription of oral contraceptives or intrauterine devices associated with condoms is mandatory for women of child-bearing age, unless hysterectomized.\textsuperscript{4,83} The measurement of blood chorionic beta-gonadotropin should be requested beforehand and monthly, during treatment. There are no risks for future pregnancies, which are authorized one month after the end of treatment.\textsuperscript{83,86}

The hypothesis of triggering psychiatric disorders and IBD has caused numerous lawsuits in the United States. However, no CRCT has demonstrated these associations.\textsuperscript{40,70} Qualitative analysis of 14 non-randomized studies on serious adverse events, nine on psychiatric adverse events, and seven on IBD did not demonstrate an increased risk.\textsuperscript{40,87-95} Two other SR with meta-analysis assessed depression and IBD, and did not detect an increased risk due to exposure to isotretinoin. In contrast, reduced levels of depression have been demonstrated in comparison with topical therapy.\textsuperscript{96}
Table 2  Indications, contraindications, warnings, and characteristics of isotretinoin in the treatment of acne, and conditions related to the need for retreatment

<table>
<thead>
<tr>
<th>Indications for use</th>
<th>Conditions related to the need for more than one course of treatment</th>
<th>Contraindications for use</th>
<th>Warnings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Severe acne (conglobata and nodular-cystic)</td>
<td>Age &lt; 16 years and family history of severe acne</td>
<td>Concomitant treatment with antibiotics from the tetracycline group (risk of intracranial hypertension)</td>
<td>Oral isotretinoin may reduce serum levels of carbamazepine and phenytoin (caution when using concomitantly)</td>
</tr>
<tr>
<td>Moderate acne (nodular-cystic or papular-pustular) with resistance to initial treatments, tendency to scarring, significant emotional impairment or impaired social functions</td>
<td>Long-lasting acne</td>
<td>Liver failure</td>
<td>Progesterone microdoses (&quot;mini-pills&quot;) are unsuitable for necessary contraception</td>
</tr>
<tr>
<td></td>
<td>Female sex and hormonal changes (polycystic ovary syndrome)</td>
<td>Pregnancy and breastfeeding</td>
<td>Do not donate blood during treatment and up to 30 days after termination (risk of accidental exposure of pregnant women to the drug)</td>
</tr>
<tr>
<td></td>
<td>Drug intake outside mealtimes (except isotretinoin-lidose, not available in Brazil)</td>
<td>Pre-existing hypervitaminosis A</td>
<td>Avoid sun exposure</td>
</tr>
<tr>
<td></td>
<td>Presence of macrocomedones</td>
<td>Severe dyslipidemia</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Treatment interruption before total resolution</td>
<td>Allergic to soy and parabens (excipient and preservative in capsule formulation, respectively)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>&quot;Hormonal athletes&quot;: patients with hyperandrogenism secondary to the use of androgen-containing drugs</td>
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</tbody>
</table>

Acne is related to psychosocial damage, increased risk of depression, and suicide, conditions already present in adolescence. Some subgroups may be more susceptible to depression and psychosis induced by isotretinoin in an idiosyncratic manner. Personal and/or family history of depression are not contraindications to the use of the drug in low daily doses and monitoring of mood and behavior in the daily routine, with the help of a psychiatrist. Thus, a history of IBD is not a contraindication for isotretinoin.

Acne flares in the first eight weeks of treatment are related to sebocyte apoptosis, antigen release, and intense inflammatory response, being observed in 15–18% of patients, with spontaneous resolution. However, they can mimic fulminant acne, without systemic symptoms, and with intense inflammation, ulceration, scabs, and scars. The drug should be kept at a low dose and associated with prednisone, 0.5–1 mg/kg/day for two to four weeks or until resolution. Severe and extensive acne (face, chest, and back), macrocomedones, and family history indicate initiation of treatment with low daily dose (0.1–0.2 mg/kg), associated with prednisone in the first two to four weeks; low dose is maintained for eight weeks and may or may not be increased gradually, along with fractional corticosteroid withdrawal.

Laboratory alterations correspond to 2% of the detected adverse events. The serum dosages most frequently altered, according to SRs and meta-analyses, are as follows: triglycerides (44%), total cholesterol, LDL-cholesterol (33%), and liver enzymes (11%). There is no evidence that these elevations increase cardiovascular risk. Previous liver and lipid profiles are recommended, repeated after one month and every three months. Analysis of laboratory monitoring concluded that tests requested less frequently are safe and economical, since changes are rare or discreet and reversible. Thus, a lipid and hepatic profile is recommended at baseline and after two months; subsequently, only the altered exams should be repeated, according to the patient’s medical history. Thus, a lipid and hepatic profile is recommended at baseline and after two months; subsequently, only the altered exams should be repeated, according to the patient’s medical history. However, some authors and even the Brazilian Unified Health
Table 3  Dermatological procedures for patients currently using or recently having used oral isotretinoin

<table>
<thead>
<tr>
<th>Safe</th>
<th>Not recommended</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manual dermabrasion</td>
<td>Mechanical dermabrasion</td>
</tr>
<tr>
<td>Microdermabrasion</td>
<td>Deep peels</td>
</tr>
<tr>
<td>Microneedling</td>
<td>Ablative lasers</td>
</tr>
<tr>
<td>Ablative and non-ablative fractional lasers</td>
<td>Deep dermatological excisions, muscle flap</td>
</tr>
<tr>
<td>Medium and superficial peels</td>
<td>Q-switched lasers and vascular lasers</td>
</tr>
<tr>
<td>Micro-needled fractional radiofrequency</td>
<td>Biopsies, superficial excisions</td>
</tr>
</tbody>
</table>

System (Sistema Único de Saúde [SUS]) still recommend frequent monitoring. A very recent study also concluded that the quality of care for patients with acne can be improved by reducing the frequency of assessment of lipids and hepatic function and eliminating the blood count assessment. The possibility of interference with strength, fatigue, and muscle endurance was investigated and no difference was observed in a study that compared patients with individuals who did not use isotretinoin. Thus, CPK measurements are only indicated if the patient has severe muscle pain.

The risk of abnormal scarring with the use of isotretinoin was assessed; five recently published guidelines concluded that there was no evidence to delay superficial cosmetic procedures, biopsies, and dermatological surgeries without involvement of muscle planes (Table 3). A retrospective observational study demonstrated no tendency to hypertrophic scarring and keloid among acne patients who used oral isotretinoin. On the contrary, some recent publications have emphasized that the use of lasers is safe, even producing better results in the case of scars, if started in the last month of treatment with isotretinoin.

Off-label prescriptions

Inflammatory diseases

Rosacea. It is believed that isotretinoin may act on rosacea by modulating innate immunity and reducing the inflammatory response through the negative regulation of TLR-2 expression in keratinocytes. Off-label use is indicated for moderate to severe papule-pustular rosacea, at a low daily dose (0.25–0.3 mg/kg), for four months, with a slow and progressive reduction. Maintenance treatment is mandatory, with topical medication (metronidazole, azelaic acid, or tretinoin) or isotretinoin in microdoses (20 mg/week), with laboratory control and assessment of pregnancy risk.

The use of oral isotretinoin for severe rosacea was first reported in 1981 in a German study that demonstrated efficacy and longer periods of remission when compared with usual treatments. Daily doses of 0.05 mg/kg, 0.5 mg/kg, or 1 mg/kg were used for 12–28 weeks. There was a 50% regression of inflammatory lesions in two weeks and 95% in eight weeks. Only telangiectasias and chronic conjunctivitis showed little improvement. Remissions were observed for more than 12 months. Side effects were mild cheilitis and a slight increase in triglycerides and cholesterol. A multicenter study, including 92 patients, lasting 20 weeks and using the same doses, concluded that isotretinoin is effective in rosacea refractory to previous recommended treatments. A CRCT compared isotretinoin, at a dose of 10 mg/day, with 0.025% tretinoin cream or both, for 16 weeks and another 16 weeks of maintenance with tretinoin or placebo cream in severe rosacea, with no differences and no advantage of the association. Adverse events were minimal and well tolerated.

The use of oral isotretinoin in the treatment of rosacea has been reported since the 1980s, in most cases by European and American authors. It is worth mentioning the first publication in Latin America in 1994, by a Chilean author who observed, in a series of six cases treated for three to six months with a dose of 0.5 mg/kg/day, rapid remission of papules and pustules, improvement in ocular manifestations, few side effects, and maintenance of results for approximately 15 months.

A multicenter, double-blinded, randomized study included 573 patients with papule-pustular and phymatous rosacea comparing different doses (0.3; 0.5; 1 mg/kg/day) vs. doxycycline 100 mg/day, 14 days and then 50 mg/day vs. placebo. After 12 weeks, the dose of 0.3 mg/kg/day was more effective than placebo, with efficacy equal to or greater than doxycycline (reduction of 90% vs. 83% of lesions) and fewer side effects.

Another multicenter, randomized study, including 156 patients, compared the dose of 0.25 mg/kg/day (n = 108) vs. placebo (n = 48), for four months. The primary outcome (90% reduction in the number of lesions) was observed in 57% vs. 10% of the patients. Four-month recurrence was observed in 58% of patients. Studies have been suggested to investigate the minimum dose to maintain remission.

To control recurrences, continuous microdoses have been proposed. Twelve patients with recurrent rosacea were treated with 10–20 mg/day for four to six months and subsequently received a maintenance dose of 0.03–0.17 mg/kg/day (mean: 0.07 mg/kg/day) for up to 33 months. There was an improvement in quality of life, suggesting that microdosing would be a better option than multiple cycles of antibiotic therapy. In another study, 25 patients were treated with a dose of 20 mg/day for four months, with rapid reduction of erythema and inflammatory lesions; subsequently, a slow dose reduction was performed for six months, up to 20 mg/week. At 11 months, 45% of cases presented recurrence.

Fulminant rosacea is a unique, rare, highly inflammatory form in the center of the face, with an abrupt onset and the presence of papules, pustules, nodules, and sinus tracts draining sero-purulent, coalescent secretion. The treatment of choice is isotretinoin associated with prednisone, 40–60 mg/day. The initial daily dose of 0.2–0.5 mg/kg is recommended, increasing to 0.5–1 mg/kg for three to four months.

There are no controlled and randomized studies on the use of oral isotretinoin in phymatous rosacea. A Singapore author reported, in a letter, a reduction in rhinophyma in one patient, after six months of treatment with isotretinoin, 20 mg/day, with a tendency to recurrence after eight
months. That author remarked that it is an option to reduce the lesion for later surgical or laser procedures.\textsuperscript{121} In the last published SR, it was not possible to include studies on phyma.\textsuperscript{122} By suppressing the sebaceous gland and decreasing sebogenesis, isotretinoin could delay the progression of the phyma when used in the pre-fibrotic phase, with better results in young patients, but recurrence is observed after drug discontinuation.\textsuperscript{122,123} The global consensus panel, ROSaceaConsensus (ROSCO), indicates isotretinoin as a therapeutic option in the severe inflammatory (papulopustular) form and in inflamed phyma, in an early stage, with a high degree of recommendation.\textsuperscript{124}

Regarding ocular rosacea, a review article indicated the benefit and safety of isotretinoin.\textsuperscript{125} A recent comparative study with doxycycline, published by Brazilian ophthalmologists and dermatologists, showed that although doxycycline was more effective, isotretinoin, at a dose of 10 mg/day, also improved blepharitis and conjunctivitis, without adverse events.\textsuperscript{126}

An SR, using the Cochrane methodology, concluded that isotretinoin has a high degree of recommendation for moderate to severe papular-pustular rosacea, relapsing cases or those unresponsive to antibiotic therapy, and for inflamed phymas. The dose of 0.25 mg/kg/day for 12–16 weeks is greater than that of doxycycline, 50–100 mg/day. Topical maintenance is always recommended.\textsuperscript{122}

The Ibero-Latin American Rosacea Studies Group has published a treatment algorithm including low daily dose isotretinoin for the papule-pustular and hyperplastic/phymatous gland subtypes.\textsuperscript{127} A Canadian guideline presented the same recommendation.\textsuperscript{128} A review article highlights the excellent results of this drug for rosacea and recommends that dermatologists consider this option, since its safety has been determined after more than 30 years of use, reducing the use of oral antibiotics for chronic disease.\textsuperscript{129,130} The American Society of Acne and Rosacea, in its consensus, suggests isotretinoin for diffuse mid-facial erythema with papules and pustules, granulomatous rosacea, and early phyma.\textsuperscript{31} A low dose is effective, with fewer side effects and good adherence. There is a need for clinical and laboratory control and attention to teratogenicity.\textsuperscript{131} As it affects the face, rosacea has a negative impact on quality of life and its control provides benefits in patients’ emotional, social, and professional lives.\textsuperscript{132}

**Seborrheic dermatitis (SD).** SD is a chronic, recurrent inflammatory dermatosis, located in areas of high concentration of sebaceous glands: face (88%), retroauricular region, scalp (70%), anterior chest (27%), lower limbs (2%), upper limbs (1%), and flexures (5%).\textsuperscript{133}

Despite little knowledge about its etiopathogenesis, it is admitted that the efficacy of isotretinoin in SD is explained by the sebo-suppressive action and modulation of innate immunity and inflammatory response, \textit{i.e.}, downregulation of TLR-2 and the NF-κB pathway, with reduction in cytokine production. In SD, TLR-2 is activated by lipophilic fungi of the genus \textit{Malassezia}, in adults and \textit{Candida} spp. in infants, present in the normal skin microbiota,\textsuperscript{134,135} explaining the option of topical treatment of SD with antifungals.\textsuperscript{136} Topical immunomodulators and corticosteroids are also used\textsuperscript{137}; in extensive conditions resistant to topical treatment, systemic treatment with corticosteroids or isotretinoin may be necessary. This drug is a second-line treatment, used in clinical practice, but there is no definition of dose and duration of treatment. The need for laboratory control and pregnancy prevention is emphasized.\textsuperscript{138}

The first report of successful use of isotretinoin in SD, in a low daily dose, was published in Germany in 2003.\textsuperscript{139} Subsequently, a 14-year-old adolescent with pityriasis versicolor (PV) on the back and severe acne was treated with 40 mg of isotretinoin, twice daily (1 mg/kg/day) for five months. Clinical and mycological cure of PV was observed, suggesting a role against \textit{Malassezia} directly or by reducing the skin’s lipid content due to the xerosis caused by the drug, interfering with the microbiota’s conditions, since this fungus is lipophilic.\textsuperscript{140}

A patient with severe facial SD for 22 years was treated with isotretinoin, 0.3 mg/kg/day, with improvement after 30 days; the dose was reduced to 0.15 mg/kg, every other day for two months, with complete remission.\textsuperscript{141} In 2017, a review of 46 cases, 40 associated with acne, 57 women, mean age 26 years, with non-responsive SD, treated with doses of 0.05–0.51 mg/kg/day (mean: 33 weeks), associated with topical ketoconazole and hydrocortisone, showed total regression or excellent response in 89% of the cases; one patient presented no improvement.\textsuperscript{142} A study that compared 10 mg/day, on alternate days, with salicylic acid and piroctone olamine topical treatment (shampoo and soap) for six months, in parallel groups, observed a reduction in the clinical score in both groups; however, this reduction was greater in the isotretinoin group, with reduction in the rate of sebaceous secretion and no effect on the quantity and species of \textit{Malassezia}.\textsuperscript{143,144} The role of \textit{Malassezia} in the pathogenesis of SD remains controversial.

**Psoriasis.** Isotretinoin, as well as etretinate and acitretin, act in the control of psoriasis by converting keratinocytes in the cytoplasm into all-trans retinoic acid, which penetrates the nucleus, binds to nuclear receptors, and activates specific regions of DNA, involved in regulating growth and cell differentiation and apoptosis. Thus, it reduces the hyperproliferation of keratinocytes, which is one of the events involved in the pathogenesis of psoriasis.\textsuperscript{24}

The report of four cases of extensive psoriasis in women treated with 0.6 mg/kg/day of isotretinoin associated with phototherapy, with oral 8-methoxypsoralen and exposure to UVA (PUVA), showed reduction in the number of PUVA sessions.\textsuperscript{145} Two randomized clinical studies described the benefit of this drug, at a dose of 0.5 mg/kg/day, associated with narrowband ultraviolet B (NB-UVB) or PUVA, for disseminated plaque psoriasis, reducing the number of phototherapy sessions. The option for isotretinoin is due to the shorter period of contraception, due to its shorter half-life in relation to etretinate or acitretin.\textsuperscript{146,147} For the same reason, isotretinoin was used, with excellent results, in a female teenager with generalized pustular psoriasis at a dose of 1.0 mg/kg/day and in two other adult patients at doses of 1.5–2.0 mg/kg/day, for four months.\textsuperscript{148,149} In a recent SR on the treatment of palmoplantar pustulosis, it was not possible to demonstrate evidence for any treatment, except for potent or systemic topical corticosteroids.\textsuperscript{150}

Other systemic treatment options for psoriasis are available, such as methotrexate, cyclosporine, and a large number of immunobiologicals. Retinoid monotherapy has limited efficacy, but can be useful when combined with
corticosteroids in pustular psoriasis and phototherapy in HIV-positive individuals, as it has no immunosuppressive effect.\textsuperscript{151}

**Hidradenitis suppurativa (HS)**

Isotretinoin in HS is not the treatment of choice; effectiveness is variable and can be explained by anti-inflammatory actions (TLR-2 modulation), and reduced expression of genes related to keratinocyte hyperproliferation.\textsuperscript{24}

HS is a chronic inflammatory disease, difficult to treat, with a negative impact on quality of life, with nodules, fistulas, abscesses, and scars. Deep excision of the lesions is the curative treatment. The use of isotretinoin, alone or in association with other treatments, has been mentioned in the literature, with variable results, in the most severe forms, as an option to reduce lesions and facilitate surgery later.\textsuperscript{152} In a retrospective study including 209 patients, 39 treated with isotretinoin, at a dose of 0.5–1.2 mg/kg/day, for four to 12 months, 14 (36%) patients presented improvement, with benefit for performing surgery.\textsuperscript{153} Another recent study assessed drug combinations for HS in 31 patients and demonstrated the benefit of isotretinoin associated with spironolactone, in milder, initial cases, an ideal time to introduce treatment and prevent disease progression.\textsuperscript{154}

**Photoaging**

Oral isotretinoin can improve the clinical, histological, and molecular characteristics of photodamage in the skin, possibly due to its conversion to all-trans retinoic acid or tretinoin.\textsuperscript{24} Topical use of tretinoin is the treatment of choice, with the highest level of evidence for moderate to severe photoaging.\textsuperscript{155–157} Its mechanisms of action are as follows: reversal of mutations in the p53 gene, reduced MMPs, increased tissue inhibition of metalloproteinase (TIMPs), and reduced loss and accelerated recovery of nuclear retinoid receptors after exposure to UV radiation.\textsuperscript{24,158}

Regarding its use in photoaging, an author from El Salvador reported, for the first time, his experience on the use of isotretinoin as an adjunct to cosmetic procedures. Despite being randomized, the study was open and uncontrolled, including 120 patients. The dose was 10 or 20 mg/day, without reference to the criterion used, three times a week, for only two months, in a group of patients undergoing varied procedures (chemical peels, botulinum toxin, collagen filling, blepharoplasty, liposuction, fat graft, facelift), without explaining whether the use was previous or concomitant. The clinical outcomes, which are difficult to assess, were as follows: pore size, pigmentation, wrinkles, thickness, elasticity, and skin color. The results were compared to those of the group that did not receive the drug and were considered better with the association.\textsuperscript{159} Another five studies were published by Brazilian authors (details in Table 4).\textsuperscript{160–164} One of them, which included 188 patients, only compared the clinical and histological effects of doses of 10 or 20 mg, on alternate days for two to six months, and found no differences.\textsuperscript{161} The two randomized studies used isotretinoin, at a dose of 20 mg, on alternate days, for three and six months, and were compared to the use of only photoprotector and moisturizer or topical tretinoin, respectively. In both cases, there was no superiority of isotretinoin in terms of clinical, histological, and immunohistochemical outcomes, except for the expression of the epidermal p53 protein, which had a significant reduction with the use of the evaluated oral drug. As for safety, no adverse clinical or laboratory events were observed, except for mild cheilitis and xerosis.\textsuperscript{162,163}

**Field cancerization – multiple actinic keratoses**

The concept of field cancerization is old, and was based on histopathological studies of multifocal neoplasms of the oral mucosa that can coalesce, relapse, and develop new lesions. It was later extended to the skin, where UV radiation causes mutations in the p53 gene, resulting in multiple actinic keratoses and non-melanoma skin cancer.\textsuperscript{164–167} Oral isotretinoin improves the clinical, histological, and immunohistochemical parameters of field cancerization, notably reducing the epidermal p53 protein.\textsuperscript{152,163}

The mechanism of action of retinoids in the prevention and treatment of non-melanoma skin cancer is not fully understood. They are known to have antiproliferative and anti-apoptotic actions, regulate keratinocyte differentiation and apoptosis, interfere with tumor initiation, reduce regulation of proto-oncogenes, and alter the expression of p53 and pro-apoptotic caspases.\textsuperscript{156-170} They work by preventing the proliferation of human papillomavirus (HPV), a known co-carcinogen.\textsuperscript{171}

Studies involving oral retinoids have focused on the prevention and treatment of non-melanoma skin tumors that are only part of the cancerization process. Details of the studies that used it for treatment are presented in Table 5.\textsuperscript{172–175}

The reported indications for prevention include: multiple non-melanoma skin cancers (> 5 per year); multiple actinic keratoses (AKs); eruptive keratoacanthomas or occurring in transplanted and/or immunosuppressed patients, xeroderma pigmentosum, exposure to chronic phototherapy, and verruciform epidermodysplasia.\textsuperscript{171,176–183}

Little is known about the use of retinoids in multiple AKs and field cancerization in immunocompetent or immunodepressed patients, at risk of developing non-melanoma skin cancer. The delimitation of field cancerization and the methodology to evaluate the effectiveness of therapies for its control is challenging. The most used method is treatment in a restricted, well-defined area, and counting of AKs with primary resolution. The recommended doses range from 0.25 to 6 mg/kg/day, lasting from months to years.\textsuperscript{184,185}

Considering that AKs are early signs of field cancerization and studies on oral retinoids are scarce,\textsuperscript{186–188} the present authors highlight the most recent study with oral isotretinoin, 10 mg/day vs. 0.05% cream tretinoin, on alternate nights. The results were similar, with a 28% decrease in the number of new AKs, after destruction of all visible AKs with cryotherapy. An improvement was observed in the immunohistochemical parameters with reduced expression of epidermal p53 and BAX proteins. The genes that encode these proteins undergo mutations induced by UV radiation, and start to act as tumor inducers instead of inducing apoptosis of keratinocytes, which were also mutated as a protective mechanism against carcinogenesis.\textsuperscript{189}
Acitretin is mostly used in immunocompromised individuals, while isotretinoin is preferred for immunocompetent patients and women with the potential to become pregnant, due to its shorter half-life. Low doses are less effective to justify its use in the treatment of non-melanoma skin cancer. However, for prevention in high-risk patients, high doses and long-term treatment should be discouraged, due to the risk of adverse events; low doses are justified to stabilize field cancerization.\textsuperscript{190–192}

<table>
<thead>
<tr>
<th>Author, year</th>
<th>n/dose/treatment time</th>
<th>Outcomes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hernandez, 2000\textsuperscript{159}</td>
<td>Group 1 (n = 60): 10–20 mg 3 × /week</td>
<td>Clinical: improvement of wrinkles, skin thickness and color, pores, elasticity, and pigmented lesions</td>
</tr>
<tr>
<td></td>
<td>Group 2 (n = 60): placebo Cosmetic procedures 2 groups, 2 months</td>
<td></td>
</tr>
<tr>
<td>Kalil, 2008\textsuperscript{160}</td>
<td>Single group (n = 50): 20 mg 3 × /week, 3 months</td>
<td>Clinical: improvement in the general appearance of the skin, wrinkles, color, and texture Histological: improvement of collagen and elastic fibers of solar elastosis</td>
</tr>
<tr>
<td>Rabello-Fonseca, 2008\textsuperscript{161}</td>
<td>Group 1 (n = 15): 10 mg 3 × /week</td>
<td>Clinical: improvement in the general appearance of the skin, wrinkles, color, and texture Histological: improvement of collagen and elastic fibers of solar elastosis</td>
</tr>
<tr>
<td></td>
<td>Group 2 (n = 15): 20 mg 3 × /week</td>
<td>No difference between doses</td>
</tr>
<tr>
<td></td>
<td>3 months</td>
<td></td>
</tr>
<tr>
<td>Bagatin, 2010\textsuperscript{162}</td>
<td>Group 1 (n = 16): 20 mg 3 × /week + photoprotector Group 2 (n = 16): photoprotector 3 months</td>
<td>Clinical: clinical improvement, profilometry, corneometry, and viscoelastic measures Histological and immunohistochemical: slight improvement, no significant reduction in p53 protein expression</td>
</tr>
<tr>
<td>Bagatin, 2014\textsuperscript{163}</td>
<td>Group 1 (n = 12): 20 mg 3 × /week + photoprotector</td>
<td>Clinical: improvement in patient opinion, blinded photographic assessment, quality of life Histological and immunohistochemical: decreased corneal layer thickening of the dermis, decreased expression of p53 protein, increased type 1 collagen</td>
</tr>
<tr>
<td></td>
<td>Group 2 (n = 12): 0.05% tretinoin cream, on alternate nights + photoprotector</td>
<td></td>
</tr>
<tr>
<td></td>
<td>6 months</td>
<td></td>
</tr>
<tr>
<td>Bravo, 2015\textsuperscript{164}</td>
<td>Single group (n = 20): 20 mg 3 × /week 3 months</td>
<td>Clinical: improved skin quality in the opinion of the patient and researcher Histological: 60% increase in the thickness of collagen fibers in 65% of patients; improved elastic tissue fragmentation</td>
</tr>
</tbody>
</table>
Table 5  Studies involving oral isotretinoin, non-melanoma skin cancer, actinic keratoses, and field cancerization

<table>
<thead>
<tr>
<th>Author, year</th>
<th>Study design (n): indication</th>
<th>Dose/duration of treatment</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hayday, 1980</td>
<td>Case report (1): multiple keratoacanthomas</td>
<td>From 2–6 mg/kg/day, 16 weeks</td>
<td>No new lesions were observed</td>
</tr>
<tr>
<td>Peck, 1982</td>
<td>Case series (3): multiple BCC</td>
<td>Mean of 1.5 mg/kg/day, 2.5–4 years</td>
<td>Regression of 9/65 lesions; no new lesions in 2–4 years</td>
</tr>
<tr>
<td>Levine, 1984</td>
<td>Case report (1): SCC and multiple keratoacanthomas</td>
<td>2 mg/kg/day, 16 weeks</td>
<td>Regression of various lesions and reduction of new tumors</td>
</tr>
<tr>
<td>Peck, 1987</td>
<td>Case reports (2): multiple BCC</td>
<td>2 mg/kg/day, 7 and 8 years</td>
<td>Regression of 15% of the lesions, decrease in the number of new lesions in the patient exposed to arsenic</td>
</tr>
<tr>
<td>Lippman, 1987</td>
<td>Case series (4): keratoacanthoma and SCC</td>
<td></td>
<td>Patient 1 – total regression of the keratoacanthoma; 2 – partial regression; 3 – partial regression of subcutaneous mass; 4–70% reduction in SCC</td>
</tr>
<tr>
<td>Kraemer, 1988</td>
<td>Case series (5): xeroderma pigmentosum</td>
<td>2 mg/kg/day, 2 years</td>
<td>121 tumors before treatment</td>
</tr>
<tr>
<td>Peck, 1988</td>
<td>Case series (12): BCC</td>
<td>Mean of 3.1 mg/kg/day, 8 months</td>
<td>Reduction to 21 tumors in 2 years of treatment Suspension, follow-up for 1 year: 25 tumors Doses of 0.25–1.5 mg/kg/day were ineffective</td>
</tr>
<tr>
<td>Mosheil, 1989</td>
<td>Case series (5): xeroderma pigmentosum</td>
<td>2 mg/kg/day, 2 years</td>
<td>63% reduction in 25 tumors</td>
</tr>
<tr>
<td>Lippman, 1992</td>
<td>Series of cases (32/28): SCC</td>
<td>1 mg/kg/day + interferon alfa, 2 months</td>
<td>Suspension: 8 times more tumors Partial response: 68%; total response: 25%</td>
</tr>
<tr>
<td>Tangrea, 1992</td>
<td>Randomized, placebo-controlled clinical study (951): multiple SCC</td>
<td>10 mg/day, 3 years</td>
<td>No difference with placebo</td>
</tr>
<tr>
<td>Majewski, 1994</td>
<td>Case series (4): multiple AK</td>
<td>10.4–0.5 mg/kg/day + calcitriol, 12 months</td>
<td>Patient 1: complete response; 2 and 3: 50–80% regression</td>
</tr>
<tr>
<td>Levine, 1997</td>
<td>Randomized, placebo-controlled clinical study (525): 4 or more BCC or SCC</td>
<td>Retinol (25,000) × isotretinoin (5–10 mg/day) × placebo</td>
<td>No difference between drugs and placebo</td>
</tr>
<tr>
<td>Feldman, 2007</td>
<td>Case report (1): multiple keratoacanthomas</td>
<td>40 mg/day, followed by acitretin and topical retinoid</td>
<td>Regression of some lesions</td>
</tr>
<tr>
<td>Trojanova, 2018</td>
<td>Case report (1): epidermodysplasia verruciformis</td>
<td>0.33–1 mg/kg/day, 18 years</td>
<td>Reduction in the number of SCC</td>
</tr>
<tr>
<td>Ianhez, 2019</td>
<td>Randomized clinical study (60): AK</td>
<td>10 mg/day × 0.05% tretinoin cream</td>
<td>Reduction in the number of AKs</td>
</tr>
</tbody>
</table>

SCC, squamous cell carcinoma; BCC, basal cell carcinoma; AK, actinic keratosis.

...and third months, the facial papules improved and regressed after 15 days. However, signs of disease activity, erythema, and perifollicular scaling remained. A later study reported reduction of facial papules after two to four months with isotretinoin, 10mg every other day, in ten patients. Recently, two cases treated with isotretinoin, 10mg/day, presented improvement in the papules after 30–45 days of treatment. In the last three studies, the progression of FFA was not evaluated. To date, data in the literature do not allow an absolute conclusion about the efficacy of this drug in FFA. Further studies are needed.

Dissecting cellulitis (DC). DC is a neutrophilic primary scarring alopecia with follicular pustules, nodules, intercommunicating abscesses, and irreversible follicular destruction. It can constitute the tetrad of follicular occlusion when associated with pilonidal cyst, hidradenitis, and acne conglobrata.

The first report of therapeutic success with isotretinoin, at a dose of 0.5 mg/kg/day for three months, showed relapse and the need for two more cycles of 1 mg/kg/day until remission. Three patients with DC received 1 mg/kg/day, then 0.75 mg/kg/day, for maintenance, for nine to 11
months, without recurrence after ten months (one patient) and after two years and six months (two patients). The authors suggested high doses and prolonged treatment to reduce relapses.\textsuperscript{106} A retrospective study, including seven patients treated with a dose of 0.75 mg/kg/day for nine to 12 months, found no recurrence at 16–42 months.\textsuperscript{207} A retrospective study of 51 patients treated with 0.5–0.8 mg/kg/day observed complete remission in 92% of the patients after three months and frequent relapses.\textsuperscript{208} In another report of 28 patients treated with a mean dose of 30 mg/day, seven had reduced inflammatory activity; relapse and need for retreatment were not specified.\textsuperscript{108}

Doses of 10 mg/day to 1 mg/kg/day, duration, maintenance doses, and varying associations have been reported in the literature. Despite the small number of reports, frequent relapses, and few studies with long follow-up, a SR concluded that, even without evidence, oral isotretinoin is considered the treatment of choice for DC.\textsuperscript{204} Quinquaud folliculitis decalvans (QFD). Quinquaud folliculitis decalvans is a rare, chronic, and recurrent neutrophilic scarring alopecia that affects young adults of both sexes. It is characterized by fibrotic plaques of alopecia with tufts of hair on the periphery, erythema, follicular pustules, flaking, and crusts. Its etiology is unclear, and no therapy is capable of inducing prolonged remission. Isotretinoin can act by inhibiting the migration of neutrophils and modulating innate immunity against Gram-positive bacteria, through negative regulation of TLR-2. Isotretinoin is widely cited, with differences regarding efficacy, safety, time to remission, and relapses.\textsuperscript{209}

In a retrospective study involving 82 patients, 16 (20%) used isotretinoin; eight (50%) improved, but the duration of the response was only three months.\textsuperscript{210} A multicenter, prospective study included 60 patients with QFD and different treatments; 15 (25%) were treated with isotretinoin for three months, with no difference in efficacy compared with the combination of rifampicin and clindamycin in the five-year follow-up. The authors developed a therapeutic protocol for QFD, suggesting isotretinoin only for severe cases, when a response is not maintained with other treatments.\textsuperscript{211} Another study assessed 39 patients treated with isotretinoin 0.1–1.02 mg/kg/day for a mean of 2.5 months; 82% presented a partial or complete response. Doses above 0.4 mg/kg/day and lasting more than three months have been associated with the best response.\textsuperscript{212}

Recent SRs have shown controversial results. One concluded that isotretinoin is the treatment with the largest number of publications, despite the limited response\textsuperscript{213}; the other concluded that the ideal option is the combination of clindamycin and rifampicin, with level of evidence 3.\textsuperscript{214}

Other diseases with keratinization and inflammation disorders

The modulation of the inflammatory response and keratinocyte hyperproliferation and differentiation justifies the indication of isotretinoin for keratinization disorders with inflammation and difficult treatment. There are case reports and citations in reviews, with no conclusions about dose and duration. Genodermatoses need continuous treatment and there are no data on long-term risks.

Results of studies justifying recommendation

Pityriasis rubra pilaris

A chronic, papular-desquamative disease, of unknown familial or acquired etiology. Its treatment is difficult, and includes UVB associated with coal tar, topical corticosteroids, calcipotriene, keratolytics, oral retinoids, methotrexate, azathioprine, and cyclosporine. The use of isotretinoin has been reported since the 1980s, with good results.\textsuperscript{215,216} A recent SR included 182 studies and 475 patients. Among those treated with retinoids, isotretinoin led to a good response in 61%; etretinate in 47%; and acitretin in 24%. The authors suggested that the first-line treatment is isotretinoin, followed by methotrexate and immunobiologicals. Cutaneous xerosis is aggravated by the drug and requires the use of emollients.\textsuperscript{217}

Cutaneous lupus erythematosus (LE)

Isotretinoin 0.2–1 mg/kg/day was indicated as an option for refractory cases of subacute LE, chronic LE, and hyperkeratotic forms, with efficacy similar to hydroxychloroquine. However, adverse events and faster relapse are more frequent; in practice, it is little used. Contraindicated in the association of LE and Sjogren’s syndrome.\textsuperscript{218}

Generalized granuloma annular

A non-infectious granulomatous disease, with papules and plaques, of unknown cause. There is no effective treatment. There are reports in the literature of the use of isotretinoin 0.5 mg/kg/day for two to six months, in the case of generalized forms, with good response but with recurrence; maintenance at a low daily dose is suggested.\textsuperscript{219}

Human papilloma virus (HPV)/condyloma acuminatum

Oral isotretinoin 0.5–1 mg/kg/day was effective for condyloma of the cervix and mucocutaneous warts, especially when flat and recalcitrant.\textsuperscript{220,221} There may be no response, even at a dose of 1 mg/kg/day; however, it contributes to the reduction of the volume or multiplicity of lesions, favoring supporting treatments.

Darier’s disease

A genetic dermatosis, with extensive areas of hyperkeratotic papules and plaques. Case reports and a review article reported improvement with isotretinoin at daily doses of 0.2–0.7 mg/kg. As a chronic disease, there is a need for continuous use with surveillance of hepatotoxicity, hypertriglyceridemia, and teratogenicity.\textsuperscript{222–224}

Others

Varied and off-label indications, with no possibility of conclusions on efficacy and safety as they are single reports,
Table 6  Summary of doses and treatment time for approved and unapproved indications for oral isotretinoin, according to clinical studies, case series, case reports, and consensuses

<table>
<thead>
<tr>
<th>Indications</th>
<th>Oral isotretinoin dose</th>
<th>Treatment time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acnes grades III and IV or unresponsive to previous treatments^1,2,4,5,6,36,40-43,46,51-54,56,58,59,64,65,67,71-74,79,80,83</td>
<td>0.5–1 mg/kg/day</td>
<td>Up to a dose of 120–150 mg/kg/day or until complete regression of the lesions</td>
</tr>
<tr>
<td>Acne – “low daily dose”^10,38,48,55,61,62,69,71,80-82</td>
<td>0.1–0.5 mg/kg/day</td>
<td>Up to 18 months or up to 1 to 2 months after lesion resolution</td>
</tr>
<tr>
<td>Acne flares^100,101</td>
<td>Low dose (up to 8 weeks) associated with prednisone</td>
<td>8 weeks low dose oral isotretinoin</td>
</tr>
<tr>
<td>Severe and extensive acne associated with macrocomedones^36,42,100,101</td>
<td>0.1–0.2 mg/kg, 8 weeks, may or may not be increased gradually with fractionated prednisone used in the first 2 to 4 weeks</td>
<td>Prednisone: 0.5–1 mg/kg/day for 2 weeks or until resolution of the flare Low dose: 8 weeks</td>
</tr>
<tr>
<td>Rosacea 112–119</td>
<td>0.25 to 0.3 mg/kg</td>
<td>Prednisone: 2–4 weeks</td>
</tr>
<tr>
<td>Rosacea fulminans^120</td>
<td>0.2–0.5 mg/kg, increased up to 0.5–1 mg/kg, associated with prednisone 40–60 mg/day</td>
<td>Microdose: up to 33 months</td>
</tr>
<tr>
<td>Phymatous rosacea^121</td>
<td>0.1–0.5 mg/kg</td>
<td>3–4 months</td>
</tr>
<tr>
<td>Facial papules of frontal fibrosing alopecia^107–109</td>
<td>0.5–1 mg/kg/day</td>
<td>6 months for active phase</td>
</tr>
<tr>
<td>Dissecting cellulitis^212–221</td>
<td>0.5–1 mg/kg/day</td>
<td>4 months</td>
</tr>
<tr>
<td>Pityriasis rubra pilaris^122–124</td>
<td>0.5–2 mg/kg/day</td>
<td>Mean of 33 weeks</td>
</tr>
<tr>
<td>Cutaneous lupus erythematosus^225</td>
<td>0.2–1 mg/kg/day</td>
<td>4 months</td>
</tr>
<tr>
<td>Generalized granuloma annulare^226</td>
<td>0.5 mg/kg/day</td>
<td>2–6 months</td>
</tr>
<tr>
<td>Condyloma acuminatum and recalcitrant warts^227,228</td>
<td>0.5 mg/kg/day (cervical condyloma)^227</td>
<td>2–4 months, maintenance at lower doses</td>
</tr>
<tr>
<td>Darier’s disease^229,230</td>
<td>0.5–1 mg/kg/day^228</td>
<td>Mean of 33 weeks</td>
</tr>
<tr>
<td></td>
<td>0.2 mg/kg at the beginning, increase to 0.5–1 mg/kg, according to the hue of the lesion</td>
<td>12 weeks</td>
</tr>
</tbody>
</table>

include: agnogenic keratoderma, oral mucosa ulcer perioral dermatitis, Galli-Galli disease, acne-like rash secondary to vemurafenib, dermatophytosis, lichen planopilaris and lichen planus pigmentosus, Cushing’s disease, sebaceous hyperplasia, Fordyce granules, multiple steatocystoma, reticulated confluent papillomatosis (Gougerot-Carteaud), and erosive postular dermatosis of the scalp.125–129

Table 6 presents a summary of approved and off-label indications for oral isotretinoin in dermatology, regarding doses and treatment times, reported in clinical studies, guidelines for conduct, and consensuses.

Perspectives

For the cure of acne, the present authors consider it relevant to expand the prescription of isotretinoin for adolescents and adults, as well as the prescription of anti-androgens (contraceptives and spironolactone) for adult women, thus reducing the prescription of oral antibiotics, considering the growing alert about bacterial resistance. The still very high use of these drugs is worrisome, lasting from over six months up to one year or more (mean: 331 days), according to a 2016 study that highlights the lack of knowledge of or disregard of the recommendations on the rational use of...
It is known that there is no minimum age to prescribe isotretinoin, since acitretin is indicated for children of any age to treat severe keratinization disorders. However, it is necessary to guide the patient and family that use in pre-adolescents may imply the need for new treatment cycles; a new cycle can begin after three months.\textsuperscript{241} Until now, and considering that the patent of isotretinoin has expired, there appears to be no interest from the pharmaceutical industry in conducting multicenter, randomized and controlled studies aimed at future approvals for other dermatoses. Only studies with appropriately-sized samples and high quality methodology will allow approval by regulatory agencies and the possibility of establishing levels of evidence in accordance with international standards.\textsuperscript{242-245} Perhaps the development of a new oral retinoid that can meet other indications, in addition to acne and psoriasis, could expand the use of these drugs in dermatology.

### Conclusions

This consensus aims to guide dermatologists on the use of oral isotretinoin for the benefit of patients. There is only level I evidence (SR and meta-analysis) with respect to efficacy and safety, ensured by adverse event monitoring, in the treatment of acne vulgaris. For rosacea, its use in low daily doses is mentioned in one SR, without mentioning the level of evidence. For the other indications, the literature is scarce, generally based on case reports, some even anecdotal, and rare randomized clinical studies with small samples (seborrheic dermatitis, photaging), with no possibility of determining the level of evidence. However, some dermatological conditions that are difficult to control and for which oral isotretinoin was attempted due to its multiple mechanisms of action are worth mentioning. There was a 100% consensus among the authors of this manuscript that off-label indications are expanding and should be included. In turn, in the opinion of the authors, indications for purely esthetic purposes or oil control are not recommended, particularly for women of childbearing age. Finally, common sense is needed to prescribe a teratogenic drug, particularly for off-label prescriptions, in which the responsibility lies entirely with the physician.

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### Authors’ contributions

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### Conflicts of interest

None declared.

### References


