The Importance of Assessing Burning and Stinging when Managing Rosacea: A Review

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Rosacea, a chronic condition usually recognized by its visible presentation, can be accompanied by invisible symptoms, such as burning and stinging. The aim of this review is to gather the most recent evidence on burning and stinging, in order to further emphasize the need to address these symptoms. Inflammatory pathways can explain both the signs and symptoms of rosacea, but available treatments are still evaluated primarily on their ability to treat visible signs. Recent evidence also highlights the adverse impact of symptoms, particularly burning and stinging, on quality of life. Despite an increasing understanding of symptoms and their impact, the management of burning and stinging as part of rosacea treatment has not been widely investigated. Clinicians often underestimate the impact of these symptoms and do not routinely include them as part of management. Available therapies for rosacea have the potential to treat beyond signs, and improve burning and stinging symptoms in parallel. Further investigation is needed to better understand these benefits and to optimize the management of rosacea.

Key words: rosacea; stinging; burning; management; symptom; invisible.

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Rosacea is a chronic inflammatory skin disease predominantly affecting the central face, which is most often diagnosed through assessment of visible signs alone (1, 2). These include erythema, inflammatory lesions, phymatous changes and/or ocular manifestations (1, 3). The disease is also characterized by an array of invisible symptoms, of which burning and stinging sensations are common (4). Other invisible symptoms of rosacea, such as itching, are beyond the scope of this review. Burning in rosacea is defined as “an uncomfortable or painful feeling of heat, typically in the centro-facial region” (1). Stinging in rosacea is defined as “an uncomfortable or painful sharp, pricking sensation, typically in the centro-facial region” (1).

SIGNIFICANCE

Rosacea, which is usually recognized by its visible presentation, can be accompanied by symptoms such as burning and stinging, which are not new features of this chronic disease. By gathering the most recent evidence on burning and stinging in rosacea, it is hoped that clinicians will begin to consciously consider the impact of these symptoms on patients’ quality of life and start to include them as part of their rosacea assessment and disease management approach. Future clinical studies will provide further evidence on the ability of available therapies for rosacea to improve burning and stinging symptoms in parallel with visible signs.

A global survey performed across 6 countries (Canada, France, Germany, Italy, Poland and the USA), which included patients with rosacea (n = 300) and patients with psoriasis with facial involvement (n = 318) revealed that more than 60% of patients surveyed regardless of disease were mostly bothered by the physical discomfort of their disease. The symptoms assessed in the survey included burning, stinging, itching and pain (2). Another global survey (N = 710) which evaluated the impact of rosacea signs and symptoms on overall disease burden found that symptoms were significantly associated with a higher disease burden. These high-burden patients also spent significantly more time, on average, on daily skincare because of their rosacea (p < 0.01) (5). Collectively, this evidence demonstrates the level of impact that symptoms such as burning and stinging can have on patients with rosacea.

Furthermore, most clinical practice guidelines on rosacea recommend that the primary treatment objective is clearing the visible signs of the disease; the invisible symptoms are not always addressed (6, 7). The “Beyond the Visible 2018” report investigating the burden faced by patients with rosacea illustrated the disconnect between physicians and patients – physicians often overestimate the visible signs of rosacea, while underestimating and overlooking the symptom experience for patients (6).

Given the multiple signs and symptoms experienced by patients with rosacea, there is a need to address symptoms that are bothersome to patients but not always addressed. There is also a need to alleviate these symptoms.
alongside the visible signs of the disease. All treatments currently available for rosacea are indicated to treat the visible signs of the disease; however, comprehensive management of the disease, including both signs and symptoms, may serve to further improve therapeutic outcomes and the quality of patients’ lives (4, 6, 7).

For this review, published literature on burning and/or stinging in patients with any type of rosacea over the past 5 years were searched and reviewed with the following criteria on PubMed: (i) rosacea AND burning in any field (79 results); (ii) rosacea AND stinging in any field (43 results); (iii) rosacea AND burning OR stinging in any field (88 results); (iv) rosacea AND sensitive in any field (363 results); and (v) rosacea AND sensitive AND burning OR stinging in any field (14 results).

This review aims to address the available evidence on burning and stinging in patients with rosacea and explores the need to address these symptoms in the treatment and management of rosacea, in order to improve patient outcomes.

BURNING AND STINGING ARE NOT NEW FEATURES

In 2002, the National Rosacea Society (NRS) in the USA categorized rosacea into 4 major subtypes: erythematotelangiectatic (ETR), papulopustular (PPR), phymatous (PHY) and ocular (OR) (8). This subtype-based classification formed the foundation of subsequent evidence generation and publications in rosacea. Although this classification is focused on the visible presentation of the disease, primarily for greater diagnostic accuracy, it does not emphasize the clinical relevance of associated symptoms, such as burning and stinging, experienced by the patient (8). Evidence shows burning and stinging commonly reported by patients with rosacea may be experienced across all subtypes of rosacea.

• ETR is characterized by persistent centro-facial erythema, flushing and telangiectasia (8). In a 2017 study, 13.9% of patients with ETR (n = 409) reported burning or stinging as bothersome symptoms of their disease (p < 0.01 vs PPR; 24.1%, n = 191) (9).
• The PPR subtype is characterized by persistent centrofacial erythema with transient papules or pustules in a central facial distribution (8). In a study exploring the benefit of combination treatment for subjects with severe PPR (N = 273), at least 85% of subjects experienced burning and stinging at baseline. In this study, rosacea severity was measured by the Investigator’s Global Assessment (IGA) score, which is a 5-point scale in which IGA 0 is “clear” of rosacea lesions or erythema and IGA 4 is indicative of severe rosacea (4).
• The PHY subtype rosacea includes thickening skin, irregular surface nodularities, and enlargement, which occurs most commonly as rhinophyma, especially in the nasal area (8). There are limited data available on burning and stinging in PHY; however, according to the NRS, this subtype is frequently observed in combination with ETR and PPR, hence burning and stinging symptoms may also be present, although further investigation is needed (8).
• OR is characterized by a foreign body sensation in the eye, burning or stinging, dryness, itching, ocular photosensitivity, blurred vision, telangiectasia of the sclera or other parts of the eye, or periorbital oedema (8). A global survey (N = 300) found that OR rosacea patients (20% of rosacea patients surveyed) reported burning (53%) and stinging (38%) as 2 common skin symptoms (2).

Burning or stinging was also common in neurogenic rosacea, an additional rosacea subtype proposed in a 2011 study. Patients in this group experienced prominent neurological symptoms, of which burning or stinging was the most common symptom, experienced by 100% of patients (N = 14) alongside erythema (10). Although a small sample size, these findings warrant further investigation.

With advances in available evidence and a need to take on a more patient-centric approach to rosacea treatment, there was a need to progress beyond the subtype classification. A new approach would help to better encompass all signs and symptoms experienced by rosacea patients. In 2013, the American Acne and Rosacea Society (AARS) proposed a phenotype approach to better diagnose and treat patients with rosacea for all signs and symptoms experienced (11). Subsequently, the global ROSacea COnsensus (ROSCO) panel in 2017 supported the transition to a phenotype approach to provide a basis for local guideline development and to help to improve outcomes in all patients with rosacea by individualizing management (12). In 2019, the panel incorporated burning and stinging as secondary diagnostic criteria of rosacea, to be considered as part of any severity assessment (1).

Although the presence of burning and stinging across rosacea subtypes is clear, clinical evidence on these symptoms with proven treatment and management guidance are not readily available. Given this lack of guidance, the need to include these symptoms as a part of diagnosis and management of rosacea may often be overlooked in clinical practice (2, 8, 11).

Role of neurovascular dysregulation

In 1999, researchers were already discussing the pathophysiology of stinging sensations in patients with rosacea and a potential link with sensitive skin (13). Neurovascular dysregulation and an altered immune response are both thought to be key pathophysiological elements involved in the burning and stinging response (14). Patients with rosacea are believed to have an increased density of
Burning and stinging in rosacea: a review

Fig. 1. (a) Previous vs (b) current understanding of rosacea pathophysiology and the believed role of neurovascular transient receptor potential ion channels of vanilloid types (TRPVs) in disease pathogenesis. (a) Adapted from: Steinhoff et al. (16), (b) Buddenkotte and Steinhoff (17). ATP: adenosine triphosphate; CGRP: calcitonin gene-related peptide; ET: endothelin; ETAR: endothelin A receptor; FGF: fibroblast growth factor; IL: interleukin; KLK: kallikrein; LL-37: cathelicidin peptide; MMP: matrix metalloproteinase; PACAP: pituitary adenylate cyclase-activating peptide; PAR: protease activated receptor; pH: potential of hydrogen; SP: substance P; TGF-β: transforming growth factor-beta; T_H: T-helper cell; TLR: toll-like receptor; TNF-α: tumour necrosis factor-alpha; TRPA: transient receptor potential ion channels of ankyrin type; TSLP: thymic stromal lymphopoietin; UVB: ultraviolet B-rays; VEGF: vascular endothelial growth factor. Permissions have been obtained from authors and journals for the use of these images.
transient receptor potential ion channels of vanilloid type (TRPV) on sensory neurones, vascular cells and immune cells. TRPV activation can lead to transient experiences of flushing and burning pain as a result of the release of vasoactive neuropeptides (15, 16). In patients with rosacea, TRPV levels can be elevated and eventually become hyperactive, impacting local immune function, vascular regulation, nociception, and epidermal barrier integrity (15). TRPV hyperactivity can also result in sustained flushing (a feeling of warmth), neurogenic inflammation (with oedema) and inflammatory cell infiltration, which can be extensive (Fig. 1) (16, 17). Dysregulation of TRPVs may play a critical role in the mediation of pathophysiological cascades in rosacea and associated symptoms, such as inflammation, flushing, hypersensitive skin, burning and stinging. These findings suggest that the TRPV ion channels may be a possible target for the treatment of both signs and symptoms of rosacea (15, 16).

**Impact of Demodex mite proliferation**

In 1993, studies using the standardized skin surface biopsy (SSSB) approach observed that *Demodex* mite density was between 3.5 times and 18 times higher in subjects with PPR than in healthy controls (18, 19). A comparison between one of these studies and a 2005 study using SSSB demonstrated *Demodex* mite density was as much as 51 times higher in subjects with PPR than in healthy controls (Fig. 2) (18, 20, 21). The presence of higher *Demodex* mite densities in rosacea compared with control skin has also been confirmed by other authors and using different techniques, including a 2012 PCR study that found density was 5.7 times and 2.9 times higher in subjects with rosacea using ETR/facial PPR or facial PPR than in healthy controls, respectively (22). The causal relationship between *Demodex* mites and rosacea is based not only on their higher density in patients with rosacea compared with healthy controls; other factors include histological damage and perifollicular infiltrate caused by the mite, possible activation of the toll-like receptor 2 (TLR-2) pathway leading to inflammatory skin changes, clinical improvement with pure acaricidal treatments and other arguments explained below (21, 23).

There is evidence that proliferation of *Demodex* mites in facial skin can contribute to the emergence of clinical manifestations of rosacea (21, 24, 25). In a prospective cohort study, the authors showed a highly significant reduction in *Demodex* mites within the first week of topical ivermectin treatment, but also a significant down-regulation of the pro-inflammatory genes interleukin-8 (IL-8), cathelicidin peptide LL-37, beta-defensin-3 (HBD3), TLR-4 and tumour necrosis factor-alpha (TNF-α), concomitantly with significant clinical improvement of both signs and symptoms of rosacea (24).

Proliferation of *Demodex* mites causes cutaneous barrier damage by eroding the epithelium. This, in turn, can lead to skin hypersensitivity, which can be reversible when the *Demodex* mite density is reduced through treatment (21). The underlying pathogenesis is not fully understood; however, there is speculation that chitin, which constitutes part of the *Demodex* mite’s outer shell, plays a role in inducing inflammatory reactions in rosacea (26). TLRs act as sensors of microbial pathogens and trigger downstream immune responses, in order to eliminate the pathogen (15). Chitin induces the expression of TLR-4, suggesting that chitin is sensed by TLRs on keratinocytes and thus could play a role in how *Demodex* mite infiltration contributes to cutaneous inflammation observed in rosacea (26).

There is evidence to support that individuals with ETR have higher mean densities of *Demodex* mites than healthy controls, but lower densities than patients with PPR (27). The association between *Demodex* mites and cutaneous inflammation is controversial, as the skin of individuals without rosacea may also be colonized by *Demodex* mites as a commensal organism (28). As a result, it is difficult to confirm that high *Demodex* mite density is a primary cause of burning and stinging in patients with rosacea; however, the stimulation of the immune response by a higher *Demodex* mite density may play a contributory role (26–30).

Proliferation of *Demodex* mites is reported to play an important role in the pathophysiology of OR. An inflammatory immune response and an increased density of *Demodex* mites causes substantial upregulation in pro-inflammatory cytokines involved in ocular inflammation and vasoregulation, along with enlarged, dilated vessels in the upper dermis. This promotes leukocyte infiltration and post-capillary oedema, which can manifest as blepharitis (31).

**Epidermal barrier damage**

Evidence also shows that patients with PPR experience impaired epidermal barrier function. Despite recent developments in the role of *Demodex* mite colonization...
and neurovascular dysregulation in rosacea, there is still a limited understanding of the correlation between the clinical manifestations of rosacea and dysfunction of the epidermal barrier (32).

Epidermal barrier dysfunction can reduce skin resistance to irritants and allergens, which can exacerbate signs and symptoms of rosacea (33–36). Although the mechanisms are not fully understood, evidence suggests that the interactions between rosacea-prone skin and physical, chemical and microbial factors play important roles in the pathophysiology of rosacea (15, 37). PPR is often misdiagnosed as acne vulgaris. In a study comparing the 2 conditions, significantly more subjects with PPR (n = 463) presented with erythema, burning, dryness and itching vs acne vulgaris subjects (n = 412) (p < 0.05) (33). Although data are limited, the consequences of epidermal barrier dysfunction appear to contribute to the pathophysiology, signs, and symptoms of rosacea.

Transepidermal water loss (TEWL) has been shown to be significantly higher in PPR sites vs healthy subjects (p < 0.001) (33). A 2020 study found that the epidermis of patients with PPR (n = 25) had a lower water content and higher TEWL in lesions than in control areas (38). Mild cleansers and moisturizers have been found to relieve burning and stinging symptoms and improve epidermal barrier function by increasing skin water content and decreasing TEWL (34, 39, 40). Improvement in epidermal barrier function can reduce signs and symptoms of rosacea by mitigating inflammation and potentially decreasing adverse interactions with cutaneous irritants (30, 32).

**BURNING AND STINGING ARE AN IMPORTANT PART OF DIAGNOSIS**

As previously discussed, a phenotype approach to diagnosis of rosacea has been strongly proposed, along with a patient-centric approach to disease management by shifting the focus to presenting signs and symptoms (Table I) (8, 12, 41, 42). There has been a growing acceptance of this approach in diagnosing and managing rosacea, but many clinicians and authors still depend on subtyping when communicating about rosacea (1, 30, 41, 43, 44).

Although diagnostic features are mainly visible signs, the 2019 ROSCO panel highlighted that non-visible cutaneous features, which are the symptoms of rosacea, need to be considered as part of any severity assessment, of which burning and stinging are addressed (Table II). It is also important to note that the 2019 ROSCO consensus recommendations included burning and stinging

### Table I. Summary of recommendations for rosacea diagnosis according to the 2002 National Rosacea Society (NRS) subtype approach and 2017 global ROSacea Consensus panel (ROSCO)/NRS phenotype approaches. (Adapted from: Tan et al. (41))

| Year | Subtype System | Criteria |
|------|----------------|----------|
| 2002 NRS (8) | Major features: Transient centro-facial erythema, Inflammatory papules/pustules | In the absence of a diagnostic feature, a combination of at least 2 of the major features is sufficient to diagnose rosacea. |
| 2017 ROSCO/2017 NRS (12, 42) | Diagnostic features: Persistent centro-facial erythema with periodic intensification by potential trigger factors | The presence of 1 of the 2 diagnostic features is sufficient to diagnose rosacea. |

### Table II. Diagnostic features of rosacea as defined by the 2019 global ROSacea Consensus panel. (Adapted from: Schaller et al. (1))

| Diagnostic (≥ 1 feature is diagnostic) (12) | Major (≥ 2 features are diagnostic) (12) | Secondary |
|------------------------------------------|------------------------------------------|------------|
| Persistent erythema | Inflammatory papules and pustules | Burning sensation |
| Fixed centro-facial redness in a characteristic pattern that may periodically intensify in response to triggers. This may be difficult to assess in darker skin phototypes (V and VI) | Flushing/transient centro-facial erythema | Uncomfortable feeling of heat |
| Phymatous changes | Temporary increase in centro-facial redness. Can include warm, heat, burning and/or pain sensations | Duration, frequency, intensity, extent (areas involved), associations with flushing, triggers, impact on daily life |
| Facial skin thickening due to fibrosis and/or sebaceous glandular hyperplasia. Most commonly affecting the nose where it can cause a bulbous appearance | Telangiectasia | Stinging sensation |
| Ocular manifestations (12) | Visible vessels in the centro-facial region extending beyond the alar area | Uncomfortable or painful sharp, pricking sensation |
| | Oedema | Duration, frequency, intensity, extent (areas involved), triggers, characteristic of the sensation, impact on daily life |
| | Telangiectasia | Facial swelling. Can be soft or firm (non-pitting) and may be self-limited in duration or persistence |
| | Dryness | Duration, degree of swelling (depth, pitting and distortion), extent (areas involved), daily fluctuation, impact on daily life |
| | Feeding of roughened skin. May be tight, scaly and/or itchy | Duration, frequency, intensity, extent (areas involved), pruritus, roughness, scale, tightness, peeling, how often moisturizers needed to be applied, impact on daily life |
as non-visible symptoms that can commonly co-exist with other features of rosacea (1).

The 2019 ROSCO update includes additional considerations to be made by assessing physicians when reviewing the signs and symptoms of rosacea to better treat patients with rosacea (Table II) (1). Adopting such an approach could improve patient outcomes by addressing an individual patient’s current clinical presentation and additional concerns (i.e. effects on quality of life) (12).

Despite the acknowledgement of burning and stinging through the phenotype approach in diagnosis and management, a practical tool to specifically score burning and stinging related to rosacea currently does not exist and highlights an opportunity for practical improvements. In addition, many clinicians have not fully incorporated the impact of non-visible symptoms to adequately recognize and address them when treating patients with rosacea (Fig. 3) (6).

**BURNING AND STINGING ARE OFTEN NOT RECOGNIZED AND ADDRESSED**

Clinicians often do not recognize or incorporate the impact of symptoms of rosacea. They will, however, strongly emphasize the visible signs of the disease, as compared with patients’ perceptions about their rosacea (Fig. 3) (6).

A recent survey (2020) highlighted that when doctors (N=361) were asked what they typically investigated in their patients with rosacea and psoriasis seen in consultation for the first time, only a minority cited symptoms. Doctors were approximately 30% more likely to investigate non-visible symptoms in their new patients with psoriasis vs those with rosacea (absolute reduction: 13%, 40% psoriasis vs 27% rosacea; p<0.01). When physicians were also asked what they typically investigated in their new and follow-up patients with rosacea as an open-ended question, there were very few mentions of burning and stinging (Table III) (2).

These data demonstrate the need to consistently include symptoms such as burning and stinging during assessment of patients presenting with rosacea. Furthermore, they highlight the need to address these symptoms when designing the management plan.

**MANAGEMENT NEEDS TO ENCOMPASS ASSESSMENT OF SYMPTOMS OF ROSACEA**

Most available guidelines and consensus recommendations in rosacea focus on treating visible signs of the disease (Table IV) (1,45‒47). The aim now is to improve our approach to rosacea management even further by consistently assessing rosacea symptoms, especially burning and stinging. It is important to recognize that reducing the impact of burning and stinging may also improve patient QoL (4, 48).

Given the adverse physical and psychosocial factors experienced by patients with rosacea, there is a need to address this burden beyond just alleviating the visible signs of the disease by aiming for “clear” (IGA 0) (6). Although this treatment approach focuses on visible improvement, the benefits of reaching “clear” (IGA 0) can extend beyond the visible benefits including improvement in quality of life (1, 49).

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**Table III. Invisible symptoms reported by patients with rosacea with moderate disease impact vs invisible symptoms investigated by dermatologists/physicians. (Adapted from: Steinhoff et al. (2))**

| Rosacea patients (n=300) | Doctor-investigated symptoms (n=361) |
|--------------------------|--------------------------------------|
| **Patient experienced ever** | **Average impact on QoL out of 10 (n)**<sup>a</sup> | **New patients** | **Follow-up patients** |
| Burning                  | 44% | 6.7 (129) | 16% | 13% |
| Stinging                 | 29% | 6.6 (88)  | 2%  | 1%  |
| Itching                  | 55% | 6.5 (162) | 10% | 8%  |
| Pain/soreness<sup>b</sup> | 21% | 6.7 (60)  | 5%  | 3%  |

<sup>a</sup>Quality of life (QoL) impact reported by patients was measured on a scale from 0 to 10, where 0 meant the symptom had no impact on their QoL, while a score of 10 meant the symptom extremely impacted their QoL. <sup>b</sup>Soreness at affected areas, around patches.

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Fig. 3. Over/underestimation of signs vs symptoms of rosacea. Calculated using patient perception on symptom impact vs doctor (patient’s rate – physician’s rate). Adapted from: Tan et al. (6).
In a pooled analysis of 1,366 subjects with rosacea from 4 randomized controlled trials, data at the end of treatment (no particular rosacea treatment) showed that more “clear” (IGA 0) than “almost clear” (IGA 1) subjects had a clinically meaningful difference in their change in DLQI score of at least 4 points is considered a moderate disease burden (4). 42% reported that itching and/or burning was a reason for non-compliance with their prescribed treatment. As such, clinicians should consider the impact of each treatment option on the occurrence of these events when deciding the treatment approach (1). Burning and/or stinging experienced by patients needs to be explored in future studies in order to better understand the therapeutic benefits of treatment on reducing the symptoms of rosacea.

### AVAILABLE TREATMENTS CAN INDUCE BURNING AND/OR STINGING

Many prescription treatments available for rosacea can cause burning and/or stinging, further increasing the burden on patients with rosacea (50, 51–62). In a recent global survey exploring patients with rosacea with at least a moderate disease burden (n=300), 42% reported that itching and/or burning was a reason for non-compliance with their prescribed treatment. As such, clinicians should consider the impact of each treatment option on the occurrence of these events when deciding the treatment approach (2). Burning and/or stinging are reported as adverse reactions in the summary of product characteristics (SmPCs)/prescribing information (PI) for common rosacea treatments (Table V) (51–53, 55–59, 62).

### Azelaic acid

Reports of burning and/or stinging exist in the published literature for patients treated with azelaic acid 15% gel (54, 63, 64). This has been well defined as a neurosensory property that can affect some patients. In the vehicle-controlled, phase III azelaic acid 15% gel studies in

### Table IV. 2019 ROSCO phenotype-based treatment approach (Adapted from: Schaller et al. (1) Schaller et al. (45))

| Erythema | Inflammatory papules/pustules | Telangiectasia | Phyma |
|----------|-----------------------------|----------------|-------|
| Transient | Persistent | Mild | Moderate | Severe | Mild, moderate & severe | Clinically inflamed |
| Alpha-adrenergics | Alpha-adrenergics: | Ivermectin | Ivermectin | Ivermectin | Clinically non-inflamed |
| Alpha-adrenergics: | brimonidine | Azelaic acid | Azelaic acid | Metronidazole |
| Alpha-adrenergics: | oxymetazoline (48) | Metronidazole | | |
| Oral | Beta-blockers | Doxycycline | Doxycycline | Doxycycline | Isotretinoin |
| Betaxolol | | | | | |
| Metoprolol | | | | | |
| Other | IPl | IPl | Electrodessication | Lasers |
| | | | | |

General skincare for all patients (sun protection factor 30+, moisturizers, gentle cleansers, trigger avoidance).

There is limited evidence to support the use of topical alpha-adrenergic modulating agents or oral beta blockers for treatment of flushing/transient erythema. However, clinical experience suggests that they could be considered in certain situations. Persistent centro-facial erythema associated with periodic intensification by potential trigger factors. Doxycycline 40 mg modified-release superior to placebo; doxycycline 40 mg modified-release inferior to doxycycline 100 mg. No inference possible from indirect comparison. Use of intense pulsed light (IPL) and vascular lasers in darker skin phototypes may require consideration by a healthcare provider with experience in this situation, e.g. pulsed-dye laser (PDL) and 532-nm potassium-titanyl-phosphate laser.

### Table V. Rosacea treatments and burning and/or stinging-related adverse effect

| Erythematous rosacea | Brimonidine tartrate 3 mg/g gel (51) | Application site erythema, pruritus, flushing and skin burning sensation are common adverse reactions, reported in 1.2–3.3% of patients in clinical studies |
|----------------------|-------------------------------------|----------------------------------------------------------------------------------------------------------------------------------|
|                      | Oxymetazoline hydrochloride 1% cream (48) | No mention of stinging or burning in the PI |
|                      | Papulopustular rosacea | Burning/stinging – mild: 13.3% of patients, moderate: 2.8% of patients, none reporting severe burning and stinging |
|                      | Minocycline 1.5% foam (indicated for inflammatory lesions) (52) | Application site burning – very common (≥1/10) |
|                      | Azelaic acid 15% gel (53, 54) | Application site burning – very common (≥1/10) |
|                      | Azelaic acid 20% cream (licensed for the topical treatment of acne) (55) | Application site burning – very common (≥1/10) |
|                      | Azelaic acid 15% foam (56) | Application site burning & stinging, paraesthesia and tenderness in 6.2% of patients |
|                      | Ivermectin 10 mg/g cream (57, 58) | Skin burning sensation – common (≥1/100 to <1/10) |
|                      | Metronidazole 0.75% gel (59) | Irritation will subside after 1 week |
|                      | Doxycycline 40 mg hard capsules (60) | Burning, pain of skin/stinging – common (≥1/100 to <1/10) |
|                      | Doxycycline 100 mg dispersible tablets (61) (licensed for the treatment of acne) | No mention of stinging or burning in SmPCs |
|                      | Isotretinoin 10 mg soft capsules (licensed for the treatment of acne) (62) | Skin irritation (pruritus, rash, dermatitis, dry skin, skin fragility, skin exfoliation) which may have associated symptoms – very common (≥1/10) |

PI: prescribing information; SmPCs: summary of product characteristics.
patients with moderate PPR (N = 333), the most common treatment-related, cutaneous adverse events (incidence \(\geq 10\%\)) included burning and stinging. Burning, stinging or itching were experienced by 38\% (statistical significance not available) of patients in the azelaic acid group, an incidence higher than in the vehicle-controlled group. In 70\% of this subset of patients, these adverse events were transient in nature and mild to moderate in intensity. Furthermore, 0.6\% of patients reported persistent and severe burning and stinging. Skin care was not controlled in these pivotal studies with azelaic acid 15\% gel (54). The use of moisturizers is recommended to reduce the severity and duration of burning and stinging in patients treated with azelaic acid 15\% gel (63).

Patients with rosacea being treated with azelaic acid foam (N = 54, 74.1\% of whom were PPR patients), a newer formulation of this topical treatment, completed an online survey of 3 questionnaires including the Rosacea Treatment Preference Questionnaire (RTPQ), Treatment Satisfaction with Medicines Questionnaire (SATMED-Q), and Dermatology Life Quality Index (DLQI). Patient-reported side-effects still included burning (7.4\%) or burning (3.7\%), with mean importance scores of 2.5 and 7.0, respectively (on a 10-point scale, with 0 meaning no importance and 10 meaning most important) (65).

**Metronidazole or azelaic acid**

In patients with rosacea who had at least one prescription for topical metronidazole gel/cream or azelaic acid gel (60.7\% of whom were patients with PPR) who completed a one-time online survey, the most common treatment-related concerns included burning sensations (8.8\% of total responders). A total of 9.2\%, 6.8\% and 11.1\% of patients receiving metronidazole gel, metronidazole cream and azelaic acid gel were concerned about burning sensations, respectively. Stinging sensations were less of a concern (3.2\% of total responders), with 6.2\% of patients in the metronidazole gel group expressing concern and no patients in the other 2 groups. Using the RTPQ across the study cohort to assess treatment-related concerns, burning received a mean score of 7.4/10, comparable with soreness (7.6/10), itching (7.5/10) and dryness (7.3/10) (66).

**Ivermectin**

In the 2 pivotal, vehicle-controlled, 12-week, phase III ivermectin 10 mg/g cream studies in moderate-to-severe PPR, treatment-emergent burning related to the study drug was reported less frequently in the ivermectin vs vehicle group (67, 68). The most common treatment-related adverse event in study 1 was skin burning, although the frequency of occurrence was lower than the vehicle group (1.8\% vs 2.6\%) (67). In study 2, skin burning was reported in 0.2\% of the ivermectin group vs 1.7\% of the vehicle group (68).

In a 40-week, investigator-blinded extension of the phase III studies, treatment-emergent burning related to the study drug was lower in the ivermectin 10 mg/g cream vs vehicle/azelaic acid 15\% gel group. Over the 40-week extension in study 1, burning was reported in 0.2\% of the ivermectin group vs 0.5\% of the azelaic acid gel group, and in study 2 was reported in 0.5\% of the ivermectin group vs 1.4\% of the azelaic acid gel group (69). It is important to note that this data cannot be directly compared as the data refer to 52 weeks of continuous ivermectin 10 mg/g cream therapy vs 40 weeks of continuous azelaic acid 15\% gel therapy after 12 weeks of pre-treatment on vehicle. However, the overall reduction in burning with ivermectin over the long-term might provide further treatment benefits (67–69).

**CHANGE IN BURNING AND/OR STINGING OVER TIME**

In clinical studies of PPR treatment, published data on burning and/or stinging in subjects at both baseline and as outcomes at follow-up visits to address potential improvement of these symptoms over time are more limited.

**Metronidazole**

In a phase IV, open-label, multicentre, community-based study of subjects with mild to moderately severe PPR treated with topical metronidazole gel, stinging was evaluated at baseline and week 12 using question 1 in the DLQI. Subjects reported a 25\% improvement in itching, pain, soreness or stinging overall from baseline to week 12 (70).

**Ivermectin**

In the phase III, investigator-blinded study comparing the efficacy and safety of ivermectin cream 10 mg/g once-daily with metronidazole 7.5 mg/g cream twice-daily (N = 962) (ATTRACT study), the incidence of worsening from baseline to week 16 for burning/stinging was higher in the metronidazole group vs the ivermectin group (15.5\% vs 11.1\%) (71). When including a 36-week extension, the incidence of worsening from baseline was still lower for ivermectin vs metronidazole (10.6\% vs 13.6\%) (72).

Previously unpublished data from the ivermectin 10 mg/g cream phase III studies and their 40-week extension demonstrated that the proportion of subjects reporting no burning/stinging increased from baseline to week 12 in both groups (Fig. 4). Study 1 demonstrated a greater improvement in the ivermectin vs vehicle group. Study 2 suggesting a more comparable increase (68).

In the ivermectin group, a further increase from week 12 to week 52 in the proportion of subjects reporting no burning/stinging was observed in both studies; an increase in the vehicle/azelaic acid group was observed.
in only one study (Fig. 4). Treatments cannot be directly compared with one another when assessing reduction in burning/stinging, as subjects in the azelaic acid 15% gel group were initially treated with vehicle cream for 12 weeks (68).

At week 12, more subjects treated with vehicle vs ivermectin had burning/stinging scores worse than baseline; ~9% vs ~4%, respectively, in study 1 and ~6% vs ~4%, respectively, in study 2. In both studies, the percentage of subjects with burning/stinging scores worse than baseline at each study visit across the 40-week extension was higher with vehicle/azelaic acid vs ivermectin, which persisted to week 52 (68).

The ANSWER phase IIIb/IV study evaluated ivermectin 10 mg/g cream as monotherapy vs combination therapy with doxycycline 40 mg modified-release capsules (sub-antibiotic dose doxycycline) in severe rosacea subjects (IGA 4) (N = 273) and systematically included burning/stinging as a secondary endpoint. Ivermectin significantly decreased burning/stinging sensations alone, or in combination with sub-antibiotic dose doxycycline, in most subjects. At baseline, more than 85% of subjects reported burning/stinging. Most subjects experienced a reduction in burning/stinging from baseline, with approximately 74% of subjects in both groups being symptom-free at week 12 (p < 0.001 for both treatments; post-hoc analysis). At baseline, subjects experienced approximately 5 episodes of flushing per week and the proportion of subjects without flushing increased from baseline to week 12 by 47.2% with combination therapy and by 41.9% with monotherapy (p < 0.001 for both treatments; post-hoc analysis) (4). The 2019 ROSCO consensus recommendations include burning as a symptom that can co-exist with flushing/transient erythema (1).

The ANSWER post-hoc analysis established a significant positive correlation between burning/stinging sensations and the total DLQI score, as well as individual parameters of the DLQI (Fig. 5). Significant correlations between the reduction in burning/stinging and individual DLQI parameters were found for itchy/painful skin, problems with partner/friends, feeling embarrassed, and social activities. Significant correlations between “clear” (IGA 0) or “almost clear” (IGA 1) and individual DLQI parameters were found for feeling embarrassed (−0.253; p = 0.000), interference with shopping/home and garden (−0.161; p = 0.012), and social activities (−0.143; p = 0.026) (49).

These findings are indirectly confirmed by the results of the ANSWER study, which included subjects with papules and pustules (inflammatory lesions). As expected, there was an improvement in inflammatory lesions, but interestingly also of erythema, flushing, burning and stinging in both treatment arms although ivermectin or ivermectin in combination with doxycycline are primarily used for the treatment of inflammatory lesions (4).

These data demonstrate that continued treatment can alleviate burning and stinging over time, highlighting the need for further exploration of the impact of different PPR treatments on these symptoms in a clinical trial setting. This should be highlighted to the patient when addressing rosacea treatment to emphasize the importance of adherence in managing both the signs and symptoms of rosacea.

CONCLUSION

Although increasing evidence is available to highlight the negative effect of burning and stinging on patients’ QoL, clinicians often do not consciously consider their impact and are not consistently including them as part of their rosacea assessment and disease management approach (6, 49). The 2019 ROSCO panel highlighted
that non-visible rosacea features, such as burning and stinging, need to be considered as part of any severity assessment (1); however, a practical tool to score burning and stinging related to rosacea does not exist.

Recent scientific evidence from clinical studies highlights that some treatment options have the potential to improve burning and stinging symptoms. Most of these data have been studied with topical ivermectin, as discussed above (4, 68, 70). Furthermore, evidence also supports the importance of managing epidermal barrier dysfunction associated with rosacea with appropriate skincare, which, in turn, can help to relieve burning and stinging symptoms (34–36, 39, 40).

Further investigation is needed to fill data gaps; improve our understanding of how to reduce the impact of burning, stinging, and other symptoms of rosacea; and to better incorporate consistent assessment of these symptoms so they are included as part of rosacea management. It is believed, based on information available to date, that this approach will improve patient outcomes in rosacea.

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