Rosacea: a clinical review

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Abstract

Rosacea is a field within dermatology with new insight within immunological research and new treatment-algorithm. Patient education on rosacea and appropriate treatments is an important aspect in helping patients succeed with therapy. Treatment should be tailored to each individual patient, taking into account: symptoms, trigger factors, patients’ wishes, most bothersome symptoms, psychological aspect, individual needs. A combination of clinical therapies to treat different symptoms concomitantly may offer the best possible outcomes for the patient. In this review article we describe these aspects.

Introduction

Rosacea is a commonly encountered chronic inflammatory skin disease in adults with a predilection for highly visible areas of the skin such as the face. It is characterized by flushing, redness, pimples, pustules and dilated blood vessels. The eyes are often involved, and thickening of the skin with enlargement (phymas), especially of the nose, can occur in some people. Combination of symptoms and signs focused around the central face can be divided in primary and secondary features.

Although rosacea can occur in anyone, it most commonly affects middle-aged women with fair skin, blue eyes and blonde hair and less frequent in skin phototypes V and VI. Studies have shown up to 10% in the Swedish population and 2-3% in France and Germany. Familial background is described in 15-40% of the patients. Three human leukocyte antigen – HLA alleles (MHC class II) are significantly associated with rosacea (HLA-DRB1, HLA-DQB1 and HLA-DQA1).

Because of the potential complexity of rosacea, it has been classified into the following 4 subtypes according to its signs and symptoms that often occur together: erythematotelangiectatic, papulopustular, phymatous and ocular.

Erythematotelangiectatic rosacea

Erythematotelangiectatic rosacea (ETR) is characterized by flushing and persistent redness of the central face, and often occurs before or at the same time as the bumps and pimples of subtype 2 (papulopustular) rosacea. Visible blood vessels may also be present. People with these signs of rosacea tend to have very sensitive skin, and may feel as if their skin stings or burns at times (Figure 1A).

Papulopustular rosacea

The symptoms of rosacea type 2 (papulopustular rosacea, PPR) may occur along with the facial redness and flushing of rosacea subtype 1 (Figure 1B).

Symptoms of rosacea subtype 2 includes: papules and/or pustules that come and go, combined with transient or persistent facial redness, primarily on the central face; burning and stinging; small visible blood vessels (telangiectasia); raised, scaly red patches known as plaques. While the papules and pustules of subtype 2 rosacea may resemble acne, there are generally no blackheads or whiteheads (comedones) present in rosacea alone. Acne more frequently affects also the back, shoulders, and chest.

This type of rosacea occurs most commonly in middle age and affects more commonly women than men.

Rhinophyma (phymatous rosacea)

Phymatous rosacea can affect nose (rhinophyma), chin (gnathophyma), forehead (metoophyma), ears (otophyma) and eyelids (blepharophyma). Rhinophyma is the most frequent location shows marked skin thickenings and irregular surface nodularities especially of the nose. Telangiectasia can also be presented. Fibrosis is present and increased volume of sebaceous glands is observed.

Histopathological signs are characteristic dilated infundibulum and dense inflammatory infiltrate.

There are 4 histological types of rhinophyma that include glandular, fibrous, fibroangiomatos and actinic.

Rhinophyma was once thought to be caused by heavy alcohol use, but rhinophyma occurs equally in people who do not use alcohol and in those who drink heavily. The problem is much more common in men than in women (Figure 1C).

Ocular rosacea

Ocular rosacea range from minor irritation, foreign body sensation, dryness, and blurry vision to severe ocular surface disruption and inflammatory keratitis. Patients frequently describe a gritty feeling, and they commonly experience blepharitis and conjunctivitis. Other ocular findings include lid margin and conjunctival telangiectasias, eyelid thickening, eyelid crusts and scales, chalazion and hordeolum, punctate epithelial erosions, corneal infiltrates, corneal ulcers, corneal scars, and vascularization. Sight-threatening disease is rare with rosacea.

Ocular rosacea is most frequently diagnosed when patients also suffer from cutaneous disease. However, ocular signs and symptoms may occur prior to cutaneous man-

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A vascular disease

Edema of the upper dermis leading to increased vascular permeability could be observed. Redness is frequent and confirmed most bothersome regardless of subtypes.8

There are many factors contributing to facial redness. Transient flushing may have variable intensity and frequency. Inflammatory lesions, lesionsal or perilesional redness, telangiectasia, persistent macular background erythema are independent of lesions.

Inflammation from papules and pustules or dry inflamed skin may obscure the level of non-transient erythema.

Different treatments depend on type of erythema

Neurovascular component

The following is observed: increased skin sensitivity to noxious heat stimuli in rosacea-affected skin, more prominent in those with PPR vs ETR; lower heat pain threshold in affected vs non-affected areas (based on heating the skin with a probe, from 32°C to 50°C); enhanced perception of noxious heat stimulus; subjective burning perception increased (based on VAS) in patients with rosacea vs control subjects; elevated skin blood flow in PPR-affected vs non-affected skin (based on LDI). This component is not significant for ETR-affected skin.9

The facial hypersensitivity is based on vascular changes due to: stasis, increased blood flow, inflammation, lower pain threshold (as described above), higher skin temperature and hypersensitivity (non-allergic).10

Neurogenic rosacea is a distinct clinical subtype requiring a modified approach to treatment with drugs like gabapentin, pregabalin and duloxetine.11

Genetic profile

Transcriptome analysis has shown a possible distinct gene profile for each subtype. More than 50% have a combination of the different subtypes. This is of importance since this patient group needs combination therapy for their rosacea.

Psychological impact

Rosacea can have a significant psychosocial impact on patients and cause anxiety, embarrassment and low self-esteem:7 77% of patients are affected emotionally, 63% are affected at work, 67% are affected socially, 53% are affected in their relationships and dating behavior.

Pathophysiology of rosacea

The following implications associated with rosacea were investigated: i) vascular disease; ii) neurovascular component; iii) inflammation; innate immunity; iv) demodex folliculorum.12

A vascular disease

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Inflammation: innate and adaptive immunity

Both the innate and adaptive immune system, are involved in the development of rosacea at a very early stage. Initially T-cells and macrophages infiltrate the skin, releasing factors leading to prolonged vasodilation, seen as erythema. They are also responsible for the recruitment of neutrophils and other cells, which later are clinically seen as pustules.

The critical cells involved in the inflammatory response in rosacea are the Th1 and Th17 cells, the mast cells, the macrophages, the antibody-presenting B cells, and the neutrophils.

Increased serine protease activity and cathelicidin promotes skin inflammation in rosacea.12

Mastcells are key mediators of cathelicidin-initiated skin inflammation in rosacea,13 but anti-histaminerg treatment is rarely effective in its treatment.

This knowledge of inflammation is, in our opinion, important to correct stepwise treatment-algorithm with initially treatment with anti-inflammatory drugs.

Demodex

Demodex are ubiquitous in the normal adult population, but not in neonates. It is considered as commensals with no clinical signs in most individuals. They are microscopic mites, commonplace all over the world with parasitic existence in hair follicles, sebaceous glands and eyelid glands. In immunocompromised people they may be found elsewhere. Demodex can be transmitted directly in areas rich in sebaceous glands, but also by indirect contaminated subjects such as towels, bed clothing etc. The life cycle is 14-18 days.

Two distinct species of Demodex mites are found in humans: Demodex folliculorum and D. brevis.14 The mites vary in size from 0.1 mm to 0.4 mm long.

A higher prevalence of D. folliculorum and mean mite density was found in rosacea patients especially with a higher density of mites in involved areas compared with controls, acne and LED. Demodex induces neutrophilic and granulomatous inflammation. Demodex in the eye can cause conjunctival inflammation, superficial corneal vascularization and scarring.15 Mite density increased sig-
nificantly with the length of treatment with topical steroids (P<0.001).16
Neither doxycycline nor topical metronidazole 7.5-10 mg/g have a significant anti-parasitic effect. Demodex infestation does not decrease in parallel with improvement under tetracycline treatment.

Mite-related bacterial antigens such as B. oleronius sonicate might stimulate inflammatory cells in rosacea. Bacillus oleronius is a Gram-negative bacteria isolated in Demodex.27 The bacteria are sensitive to tetracyclines. There is a very complex microbiota in Demodex mites from rosacea patients. Pityriasis folliculorum (with spiky changes, discrete, fine whitish, partly yellowish small scales and papules) can be seen in the sebaceous hair follicles as a result of high density of D. folliculorum. If inflammation of the follicle occurs, a Rosacea-like condition is seen. Mechanical blockage of hair follicle and sebaceous ducts increases number of mites and initial hyperkeratinization of the infundibulum and stagnation may promote bacterial overgrowth.

**Trigger avoidance**

Certain foods and beverages – spicy or caffeinated, and other triggers as sunlight, stress, strenuous exercise, some types of cosmetics, waterproof cosmetics, heavy foundations that require, makeup remover should be avoided.14
An UK-based study on 60,000 patients with rosacea vs 60,000 controls showed a lower risk of rosacea in smokers: OR: 0.65 (95%CI: 0.62-0.67) = protective.19

**Photo protection in rosacea**

UV light is dangerous for rosacea since it stimulates innate inflammation; that’s the reason why photo protection should be integrated into rosacea management (at least SPF 30). Unfortunately rosacea patients are susceptible to irritation caused by sunscreen ingredients. Appropriate protective ingredients (dimethicone, cyclomethicone) in the vehicle can minimize irritation. Physical, inorganic sun blocks (titanium, zinc oxide) are usually well tolerated. Newer products utilizing micro fine particle are under evaluation.20

**Make-up-in rosacea**

Selecting appropriate cosmetics is essential. Waterproof and opaque make-up is preferred. A make-up containing ingredient that provides sun protection and decrease inflammation is recommended. Make-up with lower allergic potential should be used. Mineral make-up is well tolerated. Formulations containing silica and talc are usually used. The aim is to give a matte finish to the complexion.21

**Treatment**

**Patient education on rosacea and appropriate treatments is an important aspect in helping patients succeed with therapy.**22 Treatment should be tailored to each individual patient, taking into account: symptoms, trigger factors, patients’ wishes; most bothersome symptoms; psychological aspect; individual needs.

A combination of clinical therapies to treat different symptoms concomitantly may offer the best possible outcomes for the patient.

Erythema was universally reported as the most bothersome feature of rosacea in a study by Tan et al. (82% of 135 subjects).

**Erythema**

It is important to differentiate the underlying erythema, as its management will differ. Lesional/perilesional erythema can resolve with successful anti-inflammatory treatment. Telangiectatic rosacea is amenable to lasers/IPL management.23,24 To treat persistent macular background erythema, adrenergic receptor agonists can be used.

**Treatment considerations (chronologically based)**

**Papulopustular rosacea**

Our suggestion for initial treatment would be to use a highly well tolerated topical or an oral anti-inflammatory agent first. Topical ivermectin, which appears to have less cutaneous irritation than the metronidazole products, may be a reasonable first choice. In patients who have more inflammatory lesions, we would probably opt for the addition of a cyclin such as a subantimicrobial-dose doxycycline.

In our experience it’s important to treat the inflammatory activity first and typically for about 4 weeks before treating the telangiectasia or erythema. Isotretinoin 0.3 mg/kg is effective for the widespread inflammatory lesion.

**Dye-laser therapy**

Dye-laser therapy is very beneficial for the treatment of telangiectasia. However, if you apply it too early in very inflamed skin, it can lead to stinging and burning. If we first get the inflammation under control with a systemic drug in combination with a topical, and then perform the laser therapy later, these side effects occurs much more rarely. After the telangiectasia was well treated with laser only the background erythema is left.

**Brimonidine**

In the end, brimonidine could be used to minimize the background erythema, but only after the inflammatory lesions and telangiectasias are gone.

**A stepwise approach**

A stepwise approach therefore includes: i) getting the inflammatory lesions under control; ii) using laser modality to get rid of the blood vessels (under control), iii) and finally use brimonidine to (just) minimize the background erythema (Figure 2).

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GRADE (Grading of Recommendations,
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Review

Table 1. Clinical description of rosacea, according to the GRADE classification.

<table>
<thead>
<tr>
<th>GRADE</th>
<th>Score</th>
<th>Clinical description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clear</td>
<td>0</td>
<td>No inflammatory lesions present, no erythema</td>
</tr>
<tr>
<td>Almost clear</td>
<td>1</td>
<td>Very few small papules/pustules, very mild erythema present</td>
</tr>
<tr>
<td>Mild</td>
<td>2</td>
<td>Few small papules/pustules, mild erythema</td>
</tr>
<tr>
<td>Moderate</td>
<td>3</td>
<td>Several small or large papules/pustules, moderate erythema</td>
</tr>
<tr>
<td>Severe</td>
<td>4</td>
<td>Numerous small and/or large papules/pustules, severe erythema</td>
</tr>
</tbody>
</table>

Assessment, Development and Evaluation) scale helps in making judgments about quality of evidence and strength of recommendations (Table 1).

Primarily anti-inflammatory drugs

High-quality evidence for all outcomes, supports that ivermectin is effective and safe for papulopustular rosacea in two trials in 1371 participants. High-quality evidence supports that ivermectin was more effective than metronidazole in papulopustular rosacea one trial in 962 participants. Modified-release doxycycline 40 mg might be as effective as doxycycline 100 mg (quality of evidence: very low), with a quarter of the side effects. Azithromycin might be as effective as doxycycline 100 mg (quality of evidence: very low). Isotretinoin 0.3 mg/kg versus doxycycline 100 mg (after 2 weeks, tapered to 50 mg/day). High-quality evidence for all outcomes supporting that isotretinoin was more effective than doxycycline in papulopustular rosacea one trial in 262 participants.

Treatment of telangiectasia

Nine randomized controlled trials with dye and laser and/or light-based therapies appeared to be effective, but limited data are provided and small sample sizes are considered. Effects on telangiectasia and to a lesser extent on erythema are of low quality of evidence.

Primarily against erythema, brimonidine is a highly selective 2-adrenergic receptor agonist with vasoconstrictive activity. It starts working within 30 minutes and works up to 12 hours. There is high-quality evidence for all outcomes, supporting that brimonidine is twice as effective as vehicle for erythema. In rosacea it is the first and only proven efficacious treatment for persistent erythema.

Lastly, we still need more RCTs for recommendation on treatment of ocular rosacea.

Conclusions

Targeting symptoms and signs with novel treatments (stepwise) and combinations of therapy will significantly improve rosacea. Clear instructions and expectations are required to optimize therapy. Advice on cosmetics, camouflage and photo protection, as well as avoiding triggers, should be part of any consultation.

References

1. Lanoue J, Goldenberg G. Therapies to improve the cosmetic symptoms of rosacea. Cutis 2015;96:19-26