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Microneedling and epidermal growth factor (EGF) as strategies for the acne scars treatment

Microagulhamento e fator de crescimento epidérmico (EGF) como estratégias para o tratamento de cicatrizes de acne

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ABSTRACT

Background: The microneedling technique and the transdermal drug delivery are indicated to treat acne scars.

Objective: Evaluation of the microneedling technique associated with the drug delivery of the epidermal growth factor (EGF).

Methods: Randomized double-blind clinical trial of 30 patients divided into two groups: (1) - two microneedling sessions with a 30-day interval, (2) - two microneedling sessions with the same interval but associated with EGF drug delivery. The patients were evaluated clinically (global acne scarring grading system - Goodman and Baron) global acne scarring grading system via multispectral image and through self-perception questionnaires. The statistical analysis (Student T-test, SNK test, analysis of variance) was performed with the SisVar software (UFLA, 1996).

Results: The groups were homogeneous regarding age, gender, and phototype. Clinical assessments showed a reduction in severity scores for both groups. The multispectral analysis revealed a decrease in porphyrins ($p = 0.0296$) and an improvement in skin texture in group two subjects.

Conclusion: Microneedling therapy was effective and safe for the acne scars treatment, and EGF demonstrated to be a promising strategy as well.

Keywords: Acne vulgaris; Cicatrix; Skin

RESUMO

Introdução: a técnica de microagulhamento e aplicação de drug delivery transdérmico é indicada para o tratamento das cicatrizes de acne.

Objetivos: avaliar a técnica de microagulhamento associada a aplicação de fator de crescimento epidérmico (EGF) em drug delivery.

Métodos: ensaio clínico duplo-cego randomizado, com seleção de 30 pacientes, divididos em dois grupos: (1) duas sessões de microagulhamento com intervalo de 30 dias e (2) duas sessões de microagulhamento com mesmo intervalo e associação de drug delivery de EGF. Os pacientes foram submetidos à avaliação clínica (escala global de cicatriz de acne - Goodman e Baron, 2006), a avaliação por imagem multiespectral e por questionários de autopercepção. A avaliação estatística (Teste T Student, Teste SNK, análise de variância) foi realizada com o software estatístico SisVar (UFLA, 1996).

Resultados: os grupos foram homogêneos quanto à idade, sexo e fototipo. Na avaliação clínica, houve redução dos escores de gravidade para ambos os grupos. A análise multiespectral revelou redução das porfirinas ($p=0,0296$) e melhora da textura da pele, ambas para o grupo 2.

Conclusão: a terapia com microagulhamento foi eficaz e segura para o tratamento de cicatrizes de acne, e o EGF demonstrou ser um ativo promissor.

Palavras-chave: Acne vulgar; Cicatriz; Pele

Original article

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INTRODUCTION

Acne is an inflammatory disorder, and its primary complication is the development of scars.¹ These can occur at any stage of the disease, but early intervention in the treatment of acne is believed to be the most effective way to prevent them.² They cause a negative aesthetic impact and generate psychosocial damage, reducing the quality of life of affected patients.³ The treatment of acne scars still represents a challenge and instigates the search for other therapies and/or safer and more effective procedures.

The microneedling technique, also known as percutaneous collagen induction,⁴ is considered a minimally invasive procedure, where a device pierces the skin, physically breaking up compact bands of collagen in the superficial layer of the dermis, leading to the formation of microchannels, thus allowing the transdermal administration of substances (drug delivery).⁵ This technique can induce an inflammatory response and stimulate neovascularization and formation of type III collagen, later replaced by type I collagen.⁶ Also, it promotes the release of transforming growth factors (TGF) alpha and beta, connective tissue growth factors (CTGF), platelet-derived growth factors (PDGF), and fibroblast-derived growth factors (bFGF), as well as epidermal growth factor (EGF).⁷

EGF decreases sebum production by suppressing lipogenesis. It also demonstrates an anti-inflammatory effect, modulating the expression of cytokines in keratinocytes, inducing changes in the differentiation and maturation of suprabasal keratinocytes, and promoting downregulation of pro-fibrotic factors such as TGF beta.^{1,8} These effects suggest that its use is effective and that it is a promising therapeutic option for acne scars.

It is the first clinical trial that assessed the influence of EGF in drug delivery associated with microneedling technique, simultaneously comparing these strategies through multispectral analysis of skin conditions and acne scar assessment tools: global acne scarring grading system (Goodman, Baron, 2006), quality of life assessment questionnaires – Dermatology Quality of Life Index (DLQI), and adapted Cardiff Acne Disability Index (CADI).

METHODS

It was a randomized, double-blind, experimental clinical study with a quantitative research model, conducted in a group of patients with acne scars who underwent two microneedling sessions associated or not with EGF drug delivery, performed by a dermatologist. The Ethics Committee in Research with Human Beings of the Federal University of Juiz de Fora approved this study under protocol number 2,702,622. Informed consent was obtained from all participants included in the study.

We performed randomization using Excel software, allocating patients randomly and equally (n=15) into group one (intervention = two microneedling sessions) or group two (intervention = two microneedling sessions associated with drug

delivery of EGF), with subsequent loss to follow-up of two patients, one from each group.

The study design was entirely randomized, with responses repeated over time. The criteria analyzed to obtain the results were: analysis of the parameters obtained in the technological device (VISIA®) that performs multispectral analysis of skin conditions; responses obtained through the adapted CADI and DLQI questionnaires; and the values obtained in the Goodman and Baron's (2006) global acne scarring grading system. The groups analyzed over time were formed by a combination of treatments (one and two) and sex (woman and man), constituting four groups.

We included patients with a clinical diagnosis of acne scars, of both sexes, aged between 18–45 years, Fitzpatrick's skin phototype IV, and who did not use any type of systemic and/or topical dermatological or aesthetic treatment in the last six months in the face. We excluded individuals who were pregnant/lactating, who presented photosensitivity, immunosuppression, active infection (such as herpes simplex, impetigo, among others), severe types of acne (acne conglobata and fulminant), predisposition to keloid formation, presence of cutaneous malignancies, self-declared allergy to EGF and/or anesthetic, and those who did not complete all stages of the study.

In group one, two microneedling sessions were performed as a single strategy and with an interval of 30 days between sessions, and in group two, in addition to the two microneedling sessions of the same interval, application of 1 ml of epidermal growth factor (EPIfactor® in 30g of vehicle – 4000 ng/g) after the procedure. Only the use of sunscreen was indicated for home care.

The disposable manual device used was a dermaroller (DrRoller MTS Roller, MiRoll, Korea), containing a mobile cylinder with eight rows of 2 mm stainless steel needles, totaling 192 needles.

The facial skin surface was sanitized with 70% ethanol, and then an anesthetic block was performed using 2% lidocaine and 4% lidocaine cream (topical) to minimize discomfort.

Each facial region was pierced eight times in different directions (vertical, up and down, horizontal, right and left, and both diagonal directions) to achieve the endpoint of uniform petechiae and purpura across the entire face. We did not conduct skin preparation with retinoid derivatives or depigmenting agents to avoid confounding bias in the study.

At time zero (T0), we perform the clinical assessment and classification of acne scars using the Goodman and Baron scale (2006), multispectral skin analysis, and quality of life assessment using the adapted DLQI and CADI questionnaires. At time one (T1), corresponding to three to seven days after T0, we performed the first session of microneedling alone or associated with EGF. We conducted the second microneedling session, multispectral imaging assessment, clinical assessment, and quality of life assessment at time two (T2 – 30 days after T1). At time three

(T3 – 60 days after T1) and time four (T4 – 90 days after T1), we performed multispectral analysis and clinical evaluation. The last quality of life assessment was performed applying the described questionnaires at time five (T5 – 180 days after T1). The same investigator conducted all these instruments for each volunteer.

For the statistical analysis, the transformation of the mean values and the tests were conducted: T-Test (to compare the groups between the proposed treatments) and SNK Test (to compare the treatments over the study period).

The simple stratified sampling method with stratification defined for four groups, aiming to meet the analysis domains established for the study (treatment strategies: one [microneedling] and two [microneedling associated with EGF drug delivery], in addition to gender [man and woman]), was used as a strategy to reduce the coefficient of variation and, thus, allow intragroup and intergroup comparisons over time. Therefore, each individual became his or her control over time.

RESULTS

Demographic characteristics

There were no statistically significant differences regarding the demographic characteristics (age, sex, and skin type) of the patients between the two treatment groups (Table 1). Regarding the clinical pattern of acne scar classification, all patients had a combination of atrophic scar subtypes in the studied period: icepick, boxcar, and rolling.

Clinical assessment

The classification of acne scars severity by Goodman and Baron (2006) global acne scarring grading system scores four grades that classify them into grade one (macular scar); grade two (mild), grade three (moderate), and grade four (severe). Before performing any treatment strategy (time zero or baseline), two (7.1%) patients were classified as grade two (mild) and

26 (92.8%) as grade three (moderate), considering the studied population (n=28). At the end of treatment (90 days – T4), 10 (35.71%) patients were classified as mild and 18 (64.28%) as moderate, considering the population studied. Table 2 summarizes the results regarding the stratification by groups, and no statistically significant differences were found between the initial and final scores for both strategies employed ($p=0.25$ for group one and $p=0.12$ for group two).

Multispectral analysis

There was a reduction in porphyrin means, which may reflect a reduction in bacterial colonization by *Cutibacterium acnes* (*C. acnes*) ($p=0.0296$). Porphyrins means in group one in both sexes did not change significantly over time, but for treatment group two, they tended to decrease at time 60 (men) and time 30 (women). At time 60, the porphyrin means in both groups was significantly different for men, with a higher mean for group one than for group two, indicating that the use of EGF in group two was important to control the proliferation of *C. acnes* and thus reduce the total amount of porphyrins on the face (Table 3 and Figures 1A and 1B).

For the variables pores, wrinkles, red area, and spots on the entire face, the comparison of means between the times and treatments performed did not show a statistically significant difference ($p>0.05$). However, regarding texture in the frontal region, the overall analysis without stratification by sex and skin phototype showed a trend towards improved skin quality ($p=0.059$) for the treatment effect. Regarding the texture in the lateral area, when performing stratification by sex and skin phototype, a slight increase in the mean values of this parameter was observed, which clinically translates into a skin improvement (Chart 1) without, however, being statistically significant ($p=0.18$). In figure 2, it is possible to notice the improvement in the skin texture of patients belonging to the proposed treatments.

TABLE 1: Demographic characteristics of the population studied as a function of the treatment strategies used for acne scars.

	Treatments		Total (%)
	Microneedling (%)	Microneedling + EGF (%)	
Age (mean) $\pm$SD	28.3 $\pm$ 5.2	27.4 $\pm$ 4.8	27.9 $\pm$ 4.9
Gender			
Women	57.14	61.2	60.7
Men	42.8	35.7	39.2
Fitzpatrick Skin Phototype			
II	14.2	7.14	10.7
III	64.2	57.1	60.7
IV	21.4	35.7	28.5
Total of participants	14	14	28

Caption: EGF = epidermal growth factor; SD = standard deviation

TABLE 2: Grading of severity of acne scars by Goodman and Baron (2006) between treatment groups.

Clinical Evaluation			
	Microneedling		Microneedling + EGF
Baseline (T0)	Grade 2: 14.2% (2/14) Grade 3: 85.7% (12/14)	Baseline (T0)	Grade 2: 0 Grade 3: 100% (14/14)
T4 (90 days)	Grade 2: 42.8% (6/14) Grade 3: 57.1% (8/14)	T4 (90 days)	Grade 2: 28.5% (4/14) Grade 3: 71.4% (10/14)

Caption: EGF = epidermal growth factor

TABLE 3: Mean score for the presence of porphyrins obtained by multispectral analysis of facial skin (laterals: right and left) over time for participants submitted to the proposed treatments and stratified by sex.

Time (days) / Porphyrin (mean score)	0	30	60	90
Treatment 1 Men	2537.6 Aa	2133.8 Aa	2706.6 B*a	2567 Aa
Treatment 1 Women	1637.75 Aa	1265.5 Aa	1575 AB*a*	1975.5 Aab*
Treatment 2 Men	1656 Ab	1590 Ab	718.6 Aa*	1291 Ab
Treatment 2 Women	1777.6 Ab	1072.8 Aa	1455.8 AB*ab*	1436.2 Aab

Caption: Capital letters (A, B) compare treatments (1 and 2) within the same time, and lowercase letters (a, b) compare the times within the same treatment (1 or 2). Analysis with root transformation. Tests performed to compare the means: T and SNK test. Markings with the symbol (*) refer to different means ($p < 0.05$)

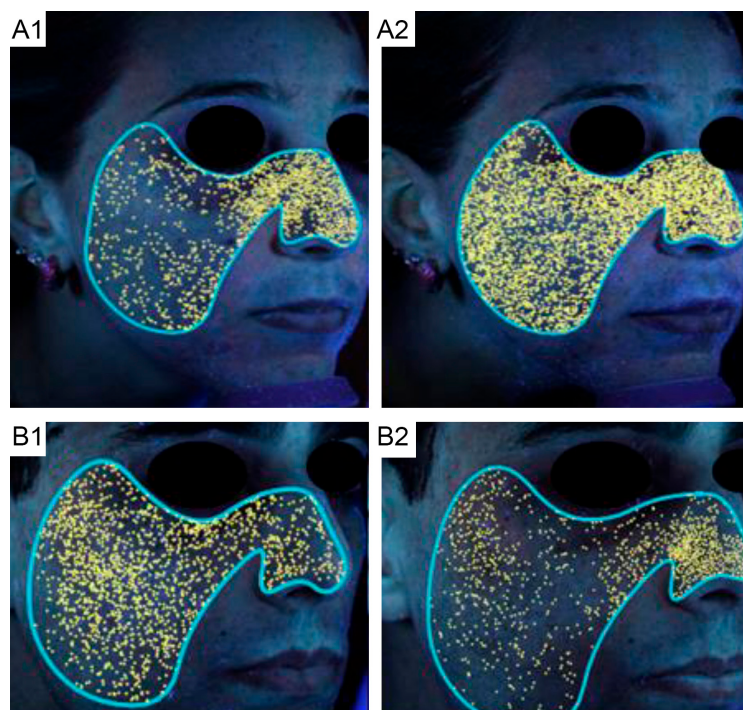


FIGURE 1: Image showing the reduction in the presence of porphyrins in the treatment group: microneedling with drug delivery of EGF.

A = microneedling.

B = microneedling + EGF

1 = start of treatment – Time 0/baseline

2 = end of treatment - Time 4/90 days

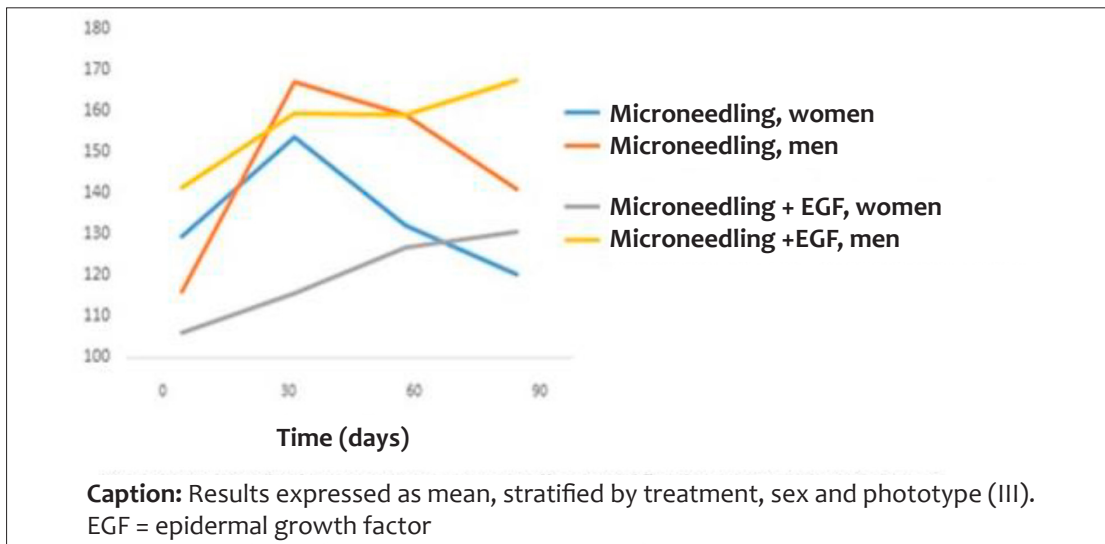


CHART 1: Texture score assessed on patients' faces over time

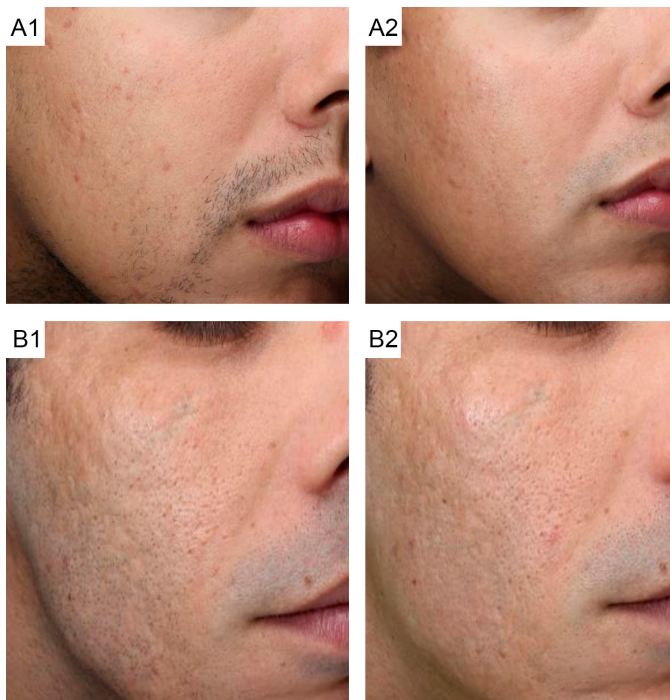


FIGURE 2: Comparative image of skin conditions of patients in treatment groups 1 and 2 at baseline and end of follow-up.

A = microneedling.

B = microneedling + EGF

1 = start of treatment – Time 0/baseline

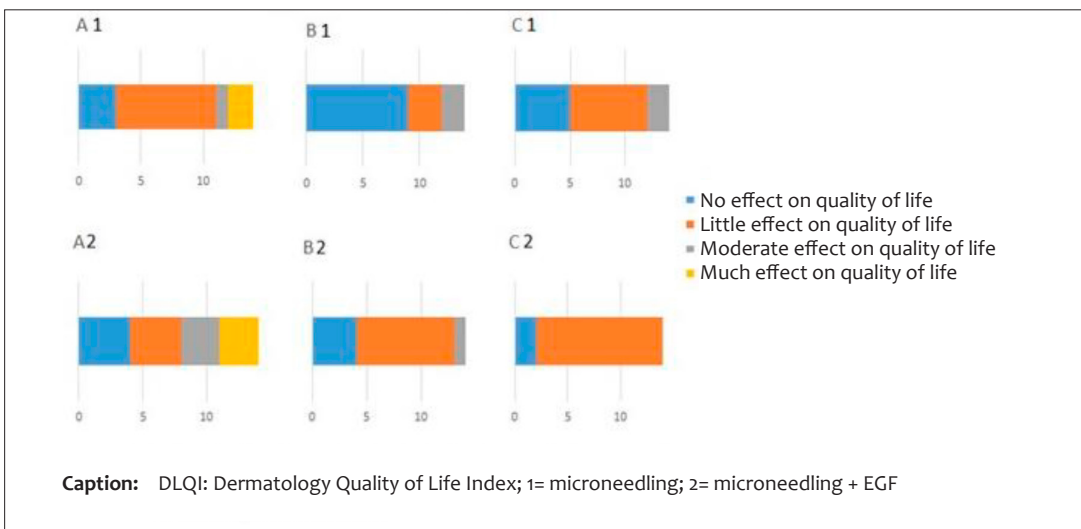
2 = end of treatment - Time 4/90 days

Self-perception assessment

DLQI is a questionnaire that assesses skin diseases in general and grades them according to the following score: 1 (0–1 point) the skin disease does not interfere with the patient's qua-

lity of life; 2 (2–5 points), the skin disease interferes little with the quality of life; 3 (6–10 points), the interference is moderate; 4 (11–20 points), there is a lot of interference in the quality of life; and 5 (21–30 points), there is huge interference in the quality of life of those affected. The results showed no significant differences in the total variation of scores (final score – initial score) for both treatments used ($p=0.25$ for treatment one and $p=0.12$ for treatment two). Before the intervention (time zero or baseline), of the 28 patients who participated of the study, 7 (25%) were classified as score 1; 12 (42.85%) as score 2; 4 (14, 28%) patients as score 3; 5 (17.85%) as score 4; and no patient was classified as score 5 ($n=28$). Based on the stratification of the groups by treatment, in treatment one (microneedling), 3 (21.42%) patients were classified as grade 1; 8 (57.14%) as grade 2; 1 (7.14%) as grade 3; and 2 (14.28%) as grade 4 ($n=14$). For treatment two (microneedling associated with drug delivery of EGF), 4 patients (25.57%) were classified as grade 1; 4 (25.57%) as grade 2; 3 (21.42%) as grade 3; and 3 (21, 42%) as grade 4 ($n=14$). At the end of the treatment, 7 (25%) patients scored 1; 19 (67.85%) scored 2; 3 (10.71%) scored 3; and no patients scored 4 and 5, considering the population studied ($n=28$). Based on the stratification of the groups by treatments, in treatment one (microneedling), 5 (35.71%) patients were classified as grade 1; 7 (50%) as grade 2; and 2 (14.28%) as grade 3 ($n=14$). For treatment two (microneedling associated with drug delivery of EGF), 2 (14.28%) patients were classified as grade 1 and 12 (85.71%) as grade 2 ($n=14$) (Chart 2).

The score obtained in the CADI questionnaire grades acne as mild (1) when the sum of the points obtained in the questionnaire varies from 0–5; moderate (2), when the sum varies from 6–10; and severe (3), when this sum varies from 11–15 points. Before conducting the intervention (time 0 or baseline), 15 (53.57%) patients were ranked as mild; 12 (42.85%) as moderate; and 1 (3.57%) as severe. In the last period of evaluation (time 5/180 days), 26 (92.85%) patients were graded as mild,



and 2 (7.14%) as moderate ($n=28$). In treatment group one (time 0 or baseline), 8 (57.14%) patients were classified as mild, 5 (35.71%) as moderate, and 1 patient (7.14%) as severe. At the end of the study (time 5/180 days), 13 (92.85%) patients were graded as mild and only 1 (7.14%) as moderate ($p=0.062$). For treatment group two, 7 (50%) patients were ranked as mild and 7 as moderate, at time 0–baseline. At the end of treatment (time 5/180 days), 13 (92.85%) patients were classified as mild and only 1 (7.14%) as moderate ($p = 0.12$) (Chart 3).

DISCUSSION

Acne scar management has been a challenging task and a focus of interest for dermatologists. Skin microneedling is a modality for remodeling acne scars, with minimal damage to the epidermis, few adverse events, and shorter recovery time after the procedure, compared to other methods with the same purpose.^{10,5,11} Regarding patient selection, the literature presents several clinical trials with microneedling treatment for acne scars in young patients of both sexes. As verified by Harris *et al.* (2015), the average number of microneedling sessions needed to achieve satisfactory results was three sessions with mean intervals of four weeks (two to eight weeks), chiefly using 1.5 mm needles (1 mm to 3 mm) in length.¹² In agreement with most of the studies analyzed, we selected 30 patients with acne scars, with a mean age of 27.9 ± 4.9 years, skin phototypes IV, of both sexes. After the loss to follow-up ($n=2$), 17 women and 11 men completed the study ($n=28$). Two microneedling sessions were performed using a 2 mm needle at an interval of four weeks between them.

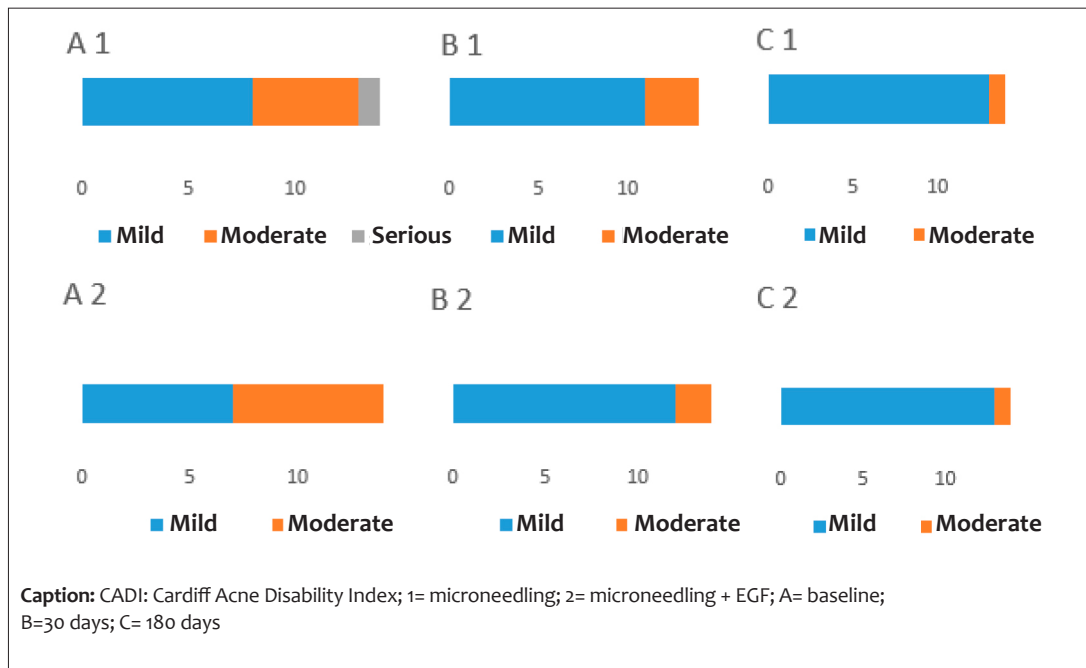
Kalil, Frainer *et al.* (2015) selected 10 patients aged 20–40 years, of both sexes, with atrophic acne scars, who underwent three sessions of microneedling using a 2mm needle. The study used anatomopathological analysis and digital photographs.¹³ There was application drug delivery of growth factors (EGF, IGF, TGF- β 3) using masks. The authors did not find improvement in icepick scars, but they did notice an overall improvement in skin texture and a slight improvement in acne scars. In our study,

we also observed an overall improvement in skin texture, especially in those patients who had EGF drug delivery after two microneedling sessions. Additionally, icepick scars slightly improved with shallowing of their depths.

The literature describes that EGF decreases sebum production and has an anti-inflammatory effect, reducing follicular hyperkeratosis. In addition to acting on active acne, it also stimulates the production of dermal matrix constituents, stimulating the production of organized collagen, downregulating TGF- β 1, which has a pro-fibrotic action. With its application in drug delivery, it is expected not only improvement of acne scars but also an enhancement in those patients who have associated active acne (Kim, Yeo, Li *et al.*, 2014; Draelos, 2016; Lian and Li, 2016).^{8,14,15}

Based on these data, applying EGF in drug delivery after microneedling is justified due to its direct effect on the pathogenesis of acne and acne scars and its anti-inflammatory action, which can be beneficial for a more efficient repair of the lesion caused by the procedure.

Al Qarqaz *et al.* (2018) evaluated 48 patients with skin phototypes III–VI treated with microneedling (Dermastamp[®] electronic device) and noticed a statistically significant improvement when comparing the treatment scores (before and after) obtained by the Goodman and Baron scales and the post-acne hyperpigmentation index (PAHPI).¹⁶ Our study also used different methods to assess the effectiveness of the procedure: Goodman and Baron's global acne scarring grading system, digital photographs, and adapted generic (DLQI) and specific (CADI) self-perception questionnaires. The generic questionnaires assess the quality of life outside the clinical context (Halioua, Beumont, and Lunel, 2000).¹⁷ The specific ones, in turn, are used for a particular disease and, considering that they are manifestations of a determined clinical condition, they are more sensitive when compared to generics. Also, our study demonstrated a reduction in the global acne scarring grading system when comparing the scores before and after treatment, but without statistically significant



differences between the grades obtained at the beginning and the end of the treatment.

A literature review performed in consultation with PubMed from 1946–2015 and Embase from 1947–2015, by Harris *et al.* (2015)¹², for microneedling research to treat acne scars, assessed the effectiveness of the procedure alone, combined with other therapies, histological changes, and adverse events. When assessing the technique in isolation, all studies showed improvement with treatment with a reduction in scar severity, based on the Goodman and Baron scale, one of which decreased from 11.7 points to 6.5. Our study showed a reduction in the means with this same assessment instrument in both treatment groups and for both sexes. However, this difference between the beginning (baseline, time zero) in which 7.1% of the patients were classified as grade 2 (mild), and 92.8% as grade 3 (moderate), and the end of the follow-up (90 days, time four) where 35.7% of

patients were classified as grade 2 (mild) and 64.2% as grade 3 (moderate), was not statistically significant. However, it is worth noting that the 28.6% reduction in cases classified as grade 3 in both groups may translate into clinical improvement.

CONCLUSION

Microneedling therapy was effective and safe for treating acne scars, with minimal adverse events and short recovery time. The clinical response, expressed through the variation of the global acne scarring grading system, showed that all patients improved to varying degrees. EGF, used in drug delivery, proved to be a promising active pharmaceutical ingredient as an adjuvant in the acne scars treatment and most patients presented an improvement in quality of life, expressed by the reduction of the values obtained in the adapted DLQI and CADI questionnaires when compared to the initial values. ●

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Statistical analysis; approval of the final version of the manuscript; study design and planning; preparation and writing of the manuscript; data collection, analysis, and interpretation; active participation in research orientation; intellectual participation in propaedeutic and/or therapeutic conduct of studied cases; critical literature review; critical revision of the manuscript.

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Author's contribution: Statistical analysis; approval of the final version of the manuscript; study design and planning; preparation and writing of the manuscript; data collection, analysis, and interpretation; active participation in research orientation; intellectual participation in propaedeutic and/or therapeutic conduct of studied cases; critical literature review; critical revision of the manuscript.

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Approval of the final version of the manuscript; preparation and writing of the manuscript; active participation in research orientation; intellectual participation in propaedeutic and/or therapeutic conduct of studied cases; critical literature review; critical revision of the manuscript.

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