Evaluating the role of small particle hyaluronic acid fillers using micro-droplet technique in the face, neck and hands: a retrospective chart review

Andreas Nikolis1–4
Kaitlyn M Enright1,3–5

1Victoria Park Medispa, Montreal, QC, Canada; 2Department of Plastic Surgery, Université de Montréal, Montréal, QC, Canada; 3VP Research Unit, Montreal, QC, Canada; 4Erevna Innovations Inc., Montreal, QC, Canada; 5Department of Experimental Surgery, McGill University, Montreal, QC, Canada

Background: Loss of the viscoelastic properties of the skin is a primary sign of aging and contributes to the appearance of wrinkles. Hyaluronic acid (HA) fillers are one of the most commonly used treatments for age-related soft-tissue reduction and volume loss. Evidence is also emerging that HA fillers rejuvenate the skin.

Methods: A retrospective chart review was completed on 20 subjects treated with small particle HA (SP-HA), to investigate its effects on skin properties. Subjects having received three treatments in the face, neck, and/or hands were considered in the analyses. Skin hydration, trans-epidermal water loss (TEWL), and pH were assessed at baseline (injection #1), Week 4 (injection #2), Week 8 (injection #3), and Week 12 (follow-up).

Results: Treatment with SP-HA significantly improved hydration levels in the face, neck, and hands. Significant results were seen in the face following the first three treatments, with subjects moving up to the next hydration level (ie, hydration went from dry to moisturized) and by the second treatment in the neck and hands. TEWL scores on the face and neck remained within healthy values throughout all visits. At baseline, TEWL scores on the hands were within critical condition and after three injections they recuperated to healthy values, while pH values remained within the normal range throughout treatment.

Conclusion: A treatment regimen consisting of three SP-HA injections was safe and well tolerated. SP-HA use demonstrated a hydrating effect while positively impacting the skin’s ability to retain moisture.

Keywords: restylane, skinboosters, hyaluronic-acid, HA, skin, hydration

Background

The use of injectable hyaluronic acid (HA) fillers for facial esthetics is among the most frequent non-surgical cosmetic procedure performed in the US,1,2 with more than a million HA procedures performed annually.3 To provide more tailored treatment options to physicians and subjects, a small-particle HA (SP-HA) was developed to address skin revitalization and was originally launched in Europe in 2004 (Restylane® Vital; Galderma, Uppsala, Sweden). Its low degree of cross-linking gives SP-HA viscoelastic properties that make these products especially pliable and moldable, which is important for achieving an optimal esthetic outcome with injections into the dermal layer of the skin.4

A SP-HA has recently been approved in Canada as a skin booster for rejuvenation of the face, neck, and hands (Restylane® Skinboosters Vital 20 mg/mL and Vital Light 12 mg/mL; Galderma). Injections of SP-HA have been shown to be
SP-HA is typically injected using an intradermal micro-
droplet technique that deposits small amounts of prod-
uct (eg, 10µL) throughout the treatment area. However,
SP-HA can be injected using multiple different techniques
(eg, threading, fanning, puncture, etc), depending on the
injector’s preference after clinical assessment. While the
ability of HA to improve the appearance of fine lines is
well established, clinical observation also suggests that
SP-HA may rejuvenate the skin by increasing hydration
to the treatment area. Skin requires sufficient moisture for
biomechanical functions which can impact normal skin
physiology. As such, adequate hydration is paramount to
establishing skin health. Furthermore, dehydration can
cause dry, flaky, and tight skin that increases the appearance
of age. Increasing moisture and hydration can therefore
rejuvenate the skin by improving the appearance and health
of skin. The mechanism of action is presumably HA’s known
hydrophilic properties. However, studies published to date
have reported varied results on the efficacy of SP-HA in
improving skin hydration.5,6,9 The difficulty in accurately
measuring hydration may account for these variations in
findings. For example, environmental influences (eg, room
temperature, humidity) and patient behavior (eg, smoking,
alcohol consumption) could account for the mixed findings
found in the relevant literature. The authors of this article
thus set out to investigate the following research question
under standardized conditions: Do micro-aliquot injections
of SP-HA increase cutaneous hydration in the face, neck,
and hands in a reproducible and safe manner?

This article presents the first reported safety and efficacy
evaluation of SP-HA in a Canadian population, as well as
the impact of SP-HA on various skin parameters (ie, hydra-
tion, pH, and trans-epidermal water loss [TEWL]) following
treatment of the face, neck, and hands.

Study design

The following study received unconditional approval from
the centralized Institutional Review Board “Veritas IRB”. The
A waiver of informed consent was granted by the IRB, in
accordance with normative guidelines and Health Canada
regulations. The procedures in this study conform to the
guiding principles of the Declaration of Helsinki.

All active patient charts within one center were reviewed
for inclusion criteria in a retrospective chart review to identify
subjects treated with SP-HA in a skin health improvement
program. As part of the program, it was standard for the
center to collect data on skin parameters such as hydration,
pH, and TEWL at each visit. Patient selection was based on
a convenience sampling method. Identifiable medical charts
were coded and a blinded study coordinator used the non-
identifiable coded charts to evaluate eligibility. All research
procedures were performed on non-identifiable patient data.
Females between the ages of 30 and 75 years who underwent
SP-HA treatments to improve skin smoothness, appear-
ance, and elasticity in the lower cheek/jawline, neck, and/or
dorsal hands were eligible. Exclusion criteria included
known hypersensitivity to HA fillers, lidocaine, amide local
anesthetics, or streptococcal proteins; a history of bleeding
disorders, porphyria, or keloids; and treatment with therapies
thought to be confounding (ie, other HA fillers in the same
areas in the last 6 months). Subjects having received three
treatments, between 4–5 weeks apart, were considered in the
final analyses. A total of four visits were assessed for each
patient: Visit 1 – Baseline (injection #1), Visit 2 – Week 4
(injection #2), Visit 3 – Week 8 (injection #3), and Visit 4 –
Week 12 (follow-up).

Charts meeting the following criteria were excluded:
males, as gender-related differences in the skin’s matrix has
been shown to have an effect on the viscoelastic properties
considered in this review;10 subjects under 30 or over 75
years old, due to age-related differences in skin properties;11 a
Fitzpatrick skin type V;12 smokers, as smoking can dehydrate
the skin;13 refusal to have signed the photo release consent
form; and concomitant medications or diseases that affect
skin hydration or cause an increased level of swelling or
bruising.14 Only subjects with complete Corneometer, pH,
and TEWL data sets were reviewed. Figure 1 displays a flow
chart outlining the protocol methods.

Condition of measurements: The room which housed
the devices measuring the various skin parameters remained
constant as numerous evaluations are carried out requiring
multiple daily recordings of temperature (fixed between 22°C
and 24°C) and humidity (between 20% and 30%). As with all
subjects who undergo skin measurements at the center, sub-
jects were adapted to room conditions 20–30 minutes prior to
testing. This was usually the time required to fill in the patient
diaries and undergo photography sessions. Measurements
were performed within the same 4-hour window to exclude
diurnal variations. Skin measurements were taken at each
visit, just prior to receiving their next scheduled injections.

Injection technique: Areas of injection were precon-
ditioned with ice for 5 minutes prior to treatment and for
10–15 minutes following injections. The areas injected were
standardized and included bilateral measurements for: 1)
the face (cheek region) with measurements 3 cm lateral to the modiolus; 2) the neck with measurements 4 cm lateral to the midline, within each prominent horizontal wrinkle at a point equidistant from the midline; and 3) the hands with injections 1 cm proximal to the metacarpal phalangeal joints and 1 cm distal to the extensor retinaculum. Face, neck, and hands were injected at 1 cm intervals. Hand injections required the use of a skin tenting technique. Injections were performed on either side of the tendons, with special care not to inject the tendons, tendon sheath, and vascular structures. The maximal volumes included 0.5 mL per each cheek, 0.3 mL for each side of the neck, and 0.5 mL for each hand.

Outcome measures: The primary outcome of interest was the measure of skin hydration as evaluated by the Corneometer * CM 825 using the Multi-Probe Adapter (MPA) System by Courage + Khazaka (Cologne, Germany). The Corneometer measures the skin moisture content in arbitrary units between 0 and 130, with <30 indicating very dry; 30–40 dry; and >40 sufficiently moisturized. Individual values assessed per area were represented by an average of three Corneometer determinations. Other measures evaluated on an exploratory basis included pH and TEWL. TEWL was evaluated using the Tewameter® TM 300, Courage + Khazaka (Cologne, Germany). The Tewameter measures the rate of water lost through the skin in g/h/m² and is an estimate of the skin’s ability to retain moisture. Higher TEWL values indicate greater water loss and are consistent with increased damage of the barrier function of the stratum corneum. A TEWL score below 25 is indicative of healthy skin, as per manufacturer guidelines. Individual TEWL values were represented by an average of 20 determinations. TEWL scores were considered for safety analyses, as treatment with SP-HA was not expected to cause an increase in water
loss through the stratum corneum. pH was assessed using the pHmeter® PH 905, Courage + Khazaka (Cologne, Germany) and was considered for safety analyses, as treatment with SP-HA was not expected to cause a pH imbalance (eg, a normal skin pH range is between 4.0 and 7.0). Data on adverse events (AEs) were also collected, as well as 2D and 3D photos.

Statistical analyses
Descriptive statistics such as the mean, SDs, and mean differences from baseline were collected on all skin variables assessed (ie, hydration, pH, TEWL), at each visit. A two-tailed paired-sample t-test was used to compare means between all visits. Significance for all analyses was set at <0.05.

Results
Of all patient charts reviewed (N=1,938), 20 met all eligibility requirements and were considered for evaluation. The included charts were comprised of women with a mean age of 58.92 years (range: 46–70, SD: 9.27). Nineteen subjects received SP-HA treatments in the face (Figures 2–4), 15 received treatments in the neck, and 12 received treatments in the hands (Figures 5 and 6). The majority of the 20 subjects

Figure 2 Patient at baseline (left) and 4 weeks after receiving one SP-HA injection in the face (right).
Notes: Patient displays a reduction in her deep wrinkles and fine lines, and the Corneometer assessment revealed an amelioration in her skin’s biophysical parameters such as hydration.
Abbreviation: SP-HA, small particle hyaluronic acid.

Figure 3 Patient at baseline (left) and Week 8 after receiving two SP-HA injections in the face (right).
Notes: Patient displays a reduction in the lines of her perioral area, and the Courage + Khazaka assessments revealed an amelioration in her skin’s biophysical parameters.
Abbreviation: SP-HA, small particle hyaluronic acid.
Figure 4: Patient at baseline (left) and Week 12 after receiving three SP-HA injections in the face (right).

Notes: Patient displays a reduction in her moderate to deep lines, and the Courage + Khazaka assessments revealed an amelioration in her skin’s biophysical parameters.

Abbreviation: SP-HA, small particle hyaluronic acid.

Figure 5: Patient at baseline (left) and Week 12 after receiving three SP-HA injections in the hands (right).

Notes: Patient displays a reduction in the signs of aging in her hands (eg, loss of fat, visible veins, muscles, and tendons), and the Courage + Khazaka assessments revealed an amelioration in her skin’s biophysical parameters.

Abbreviation: SP-HA, small particle hyaluronic acid.

Figure 6: Patient at baseline (left) and Week 12 after receiving three SP-HA injections in the hands (right).

Notes: Patient displays a reduction in the signs of aging in her hands (eg, loss of fat, visible veins, muscles, and tendons), and the Courage + Khazaka assessments revealed an amelioration in her skin’s biophysical parameters.

Abbreviation: SP-HA, small particle hyaluronic acid.
received treatments in multiple areas with the following combinations being present in the sample: n=8, face, neck, and hands; n=6, face and neck; n=3, face and hands; n=2, face only; n=1, neck and hands. All subjects received bilateral treatments.

**Corneometer (hydration)**

Table 1 displays the mean, SDs, and mean differences from baseline for the hydration values, as determined by the Corneometer, for the face, neck, and hands (bilaterally). Face: At Visit 1, mean values were considered to be “dry” (ie, between 30 and 40), as per manufacturer guidelines. At Visits 2, 3, and 4, mean values were considered to be “sufficiently hydrated” (ie, >40), as per manufacturer guidelines.

Neck: For all visits, mean values were considered to be “sufficiently hydrated” (ie, >40), as per manufacturer guidelines. Hydration values at Visit 3 were significantly higher than during Visit 1 on both sides of the neck (P=0.045, right side; P=0.001, left side). All other comparisons were not statistically significant (P>0.05).

Hands: At Visit 1, mean values were considered to be “very dry” (ie, <30). At Visits 2, 3, and 4, mean values were considered to be “dry” (ie, between 30 and 40), as per manufacturer guidelines. Visit 3 was statistically significant to Visit 1 on both the right (P=0.006) and left (P=0.046) sides. Visit 4 was also statistically significant to Visit 1 on both the right (P=0.013) and left sides (P=0.025). All other comparisons were not statistically significant (P>0.05).

**Tewameter (trans-epidermal water loss)**

Table 2 displays the mean, SDs, and mean differences from baseline of the TEWL values, as determined by the Tewameter, for the face, neck, and hands (bilaterally).

**Face:** At all visits, mean pH values fell within normal ranges (ie, 4.0–7.0). There were no statistically significant differences between visits (P>0.05).

**Neck:** At all visits, mean pH values fell within normal ranges (ie, 4.0–7.0). Visit 2 was approaching statistical insignificance from Visit 1 on the left side (P=0.049). Visit 4 was statistically different from Visit 1 on both right (P=0.027) and left sides.

**Hands:** At Visit 1 and Visit 2, mean TEWL values were above the 25 g/h/m² critical threshold. At Visit 3 and Visit 4, mean TEWL values were below the 25 g/h/m² critical threshold. TEWL values at Visit 3 and Visit 4 significantly decreased from Visit 1 (P=0.000) and Visit 2 (P=0.000), on both sides. Visit 2 was not statistically significant to Visit 1, on either side (P>0.05).

### Table 1 Mean, SDs, and mean differences from baseline of the hydration values (as determined by the Corneometer), for the face, neck, and hands (bilaterally)

<table>
<thead>
<tr>
<th></th>
<th>V1 Right side</th>
<th>V2 Right side</th>
<th>V3 Right side</th>
<th>V4 Right side</th>
<th>V1 Left side</th>
<th>V2 Left side</th>
<th>V3 Left side</th>
<th>V4 Left side</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Face</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean ±SD</td>
<td>33.28±14.82</td>
<td>42.30±13.41</td>
<td>46.77±8.31</td>
<td>49.45±14.95</td>
<td>39.10±12.78</td>
<td>46.14±13.23</td>
<td>45.11±8.14</td>
<td>48.31±15.15</td>
</tr>
<tr>
<td>Mean diff from V1</td>
<td>–</td>
<td>9.02</td>
<td>13.47</td>
<td>16.31</td>
<td>–</td>
<td>7.04</td>
<td>5.21</td>
<td>8.80</td>
</tr>
<tr>
<td><strong>Neck</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean ±SD</td>
<td>48.04±7.00</td>
<td>52.85±8.05</td>
<td>57.58±10.39</td>
<td>57.02±15.99</td>
<td>48.44±7.55</td>
<td>56.45±9.14</td>
<td>61.87±10.87</td>
<td>51.12±8.82</td>
</tr>
<tr>
<td>Mean diff from V1</td>
<td>–</td>
<td>4.36</td>
<td>9.48</td>
<td>7.76</td>
<td>–</td>
<td>7.62</td>
<td>13.28</td>
<td>0.70</td>
</tr>
<tr>
<td><strong>Hands</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean ±SD</td>
<td>24.38±9.90</td>
<td>32.12±8.02</td>
<td>36.22±8.78</td>
<td>35.97±8.44</td>
<td>25.84±8.04</td>
<td>31.98±7.35</td>
<td>34.68±7.30</td>
<td>39.45±11.10</td>
</tr>
<tr>
<td>Mean diff from V1</td>
<td>–</td>
<td>6.14</td>
<td>9.54</td>
<td>11.02</td>
<td>–</td>
<td>4.74</td>
<td>6.89</td>
<td>11.94</td>
</tr>
</tbody>
</table>

**Notes:** Significance <0.05; bsignificance <0.005. **Abbreviation:** V, visit.
No severe AEs were reported. Reported AEs were all mild and included the typical and associated with the procedure and not to the products themselves. Such as temporary erythema, swelling, and bruising associated with an injection procedure, expected incidents associated with an injection procedure, differences between visits (P > 0.05).

P side (P = 0.030). All other comparisons were not statistically significant (P > 0.05).

Significance < 0.005.

**Notes:** Significance < 0.05; significance < 0.005.
**Abbreviation:** V, visit.

and left (P = 0.019) sides, Visit 3 on both right (P = 0.019) and left (P = 0.012) sides and also from Visit 2 on the right side (P = 0.030). All other comparisons were not statistically significant (P > 0.05).

Hands: At all visits, mean pH values fell within normal ranges (ie, 4.0–7.0). There were no statistically significant differences between visits (P > 0.05).

Reported AEs were all mild and included the typical and expected incidents associated with an injection procedure, such as temporary erythema, swelling, and bruising associated with the procedure and not to the products themselves. No severe AEs were reported.

**Discussion**

Despite the known dehydrating effects of increasing age, we allowed for the inclusion of wide range of age groups in this study (ie, 30–75 years old). Our significant findings within such a large range of age groups attest to the suitability of SP-HA in young, middle-aged, and older women.

Based on results of the Corneometer, treatment with SP-HA was able to move subjects up to the next hydration level (ie, face went from being dry to moisturized and hands went from being very dry to dry). For the face, significant results were seen after only one of the three treatment sessions. For the neck and hands, two treatments were needed in order to significantly increase hydration levels. While over the three treatments, there was a general trend in the face, neck, and hands to continuously improve in hydration levels, our findings suggest that two SP-HA treatments may be sufficient to observe clinical effects. As such, a discussion may ensue with the patient with respect to the need for a third treatment. Our finding of a general trend to improve in hydration levels following multiple injections of SP-HA is in alignment with the findings of other researchers, who have investigated the use of HA fillers manufactured using different technologies.5,8,18

Throughout the study, the neck had the highest moisture content, followed by the face and hands. It was observed that subjects who were sufficiently hydrated at baseline (ie, >40 corneometer reading), had hydration scores that remained relatively stable. However, subjects who were dehydrated at baseline (ie, <20 corneometer reading) had the greatest change in Corneometer values between visits. This difference in baseline values accounts for the large SD between subjects. Future studies should investigate treatment with SP-HA in a strictly dehydrated population, by using a Corneometer score at baseline of <20 as an inclusion criterion, to amplify the observed effects described in the present study. Furthermore, it is expected that in clinical practice, subjects with dehydrated skin are those most likely to experience a significant rejuvenating effect of SP-HA. The ability of SP-HA to increase skin hydration in the face, neck, and hands supports its function as a skin health adjunct in addition to being a HA filler. Our findings contribute to the body of literature increasingly investigating skin rejuvenation using HA, beyond simply treating deep wrinkles and folds.18

**Table 2** Mean, SDs, and mean differences from baseline of the trans-epidermal water loss values (as determined by the Tewameter), for the face, neck, and hands (bilateral)

<table>
<thead>
<tr>
<th></th>
<th>V1 Right side</th>
<th>V2 Right side</th>
<th>V3 Right side</th>
<th>V4 Right side</th>
<th>V1 Left side</th>
<th>V2 Left side</th>
<th>V3 Left side</th>
<th>V4 Left side</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Face</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean ± SD</td>
<td>12.32±3.09</td>
<td>18.53±12.08</td>
<td>13.37±3.46</td>
<td>12.32±2.49</td>
<td>14.99±11.16</td>
<td>17.28±11.73</td>
<td>12.75±2.73</td>
<td>12.52±2.87</td>
</tr>
<tr>
<td>Mean dif from V1</td>
<td>–</td>
<td>6.18</td>
<td>1.32</td>
<td>–0.05</td>
<td>–</td>
<td>2.26</td>
<td>–2.96</td>
<td>–3.47</td>
</tr>
<tr>
<td><strong>Neck</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean ± SD</td>
<td>8.54±2.11</td>
<td>8.42±2.68</td>
<td>7.91±2.13</td>
<td>9.53±2.16</td>
<td>7.85±2.38</td>
<td>8.31±2.97</td>
<td>7.24±2.16</td>
<td>8.63±1.98</td>
</tr>
<tr>
<td>Mean dif from V1</td>
<td>–</td>
<td>0.10</td>
<td>–0.29</td>
<td>1.26</td>
<td>–</td>
<td>0.43</td>
<td>–0.54</td>
<td>0.72</td>
</tr>
<tr>
<td><strong>Hands</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean ± SD</td>
<td>24.38±9.90</td>
<td>30.65±10.97</td>
<td>12.12±5.70</td>
<td>10.11±3.23</td>
<td>25.84±8.04</td>
<td>30.34±10.97</td>
<td>8.92±4.12</td>
<td>9.22±3.2</td>
</tr>
</tbody>
</table>
Results from the TEWL analyses revealed that SP-HA was safe and well tolerated and did not damage the stratum corneum’s ability to retain moisture or effectively act as a barrier. These data were in accordance with other investigations into the safety of multiple HA injections and their effects on the stratum corneum’s barrier function.19 This was exemplified by the fact that the treatment regimen did not cause a significant increase in the TEWL scores throughout the study. Additionally, results from TEWL scores on the hands indicate that SP-HA may actually increase the skin’s ability to retain moisture. This was evidenced by the fact that after two and three injections of SP-HA, the TEWL scores on the hands significantly decreased to below critical levels. The finding that the stratum corneum had greater water retention following treatment with SP-HA leads to the possibility that HA may reverse possible damage to the skin’s water-barrier function vs retaining hydration from the injection of micro- aliquots of product in the deep dermis. Further studies are required to identify the mechanism through which TEWL scores are positively affected by SP-HA.

The slightly variable skin pH values found in this study fell within the normal acidic range reported in the literature (ie, between pH 4.0 and 7.0), providing evidence that treatment with SP-HA did not disturb the balance of the skin microflora or cause a pH imbalance. This consistent physiologic marker demonstrates the skin remained within its homeostatic condition throughout treatment. While we found statistically significant differences between the pH values of some visits, the differences were minimal and would likely not result in clinically significant effects.

A limitation of this study arises from the study design. Given that this was a retrospective chart review, there were inherent disadvantages that could have been avoided with a prospective study. These included possible selection bias, subjects not being adequately matched on certain characteristics, weaknesses in exposure and outcome assessment due to improper recordkeeping, and missing incomplete or poorly documented information. Another limitation was that only one site was used and subjects were usually seen by the same physician at this setting; therefore, possible variations may occur at different sites when physicians with other diagnostic and treatment trends are assessed.

Overall, the findings of this retrospective chart review support that a treatment regimen consisting of three SP-HA injections, spaced 1 month apart, is safe and well tolerated by subjects. SP-HA demonstrated a major hydrating effect (as exemplified by an increase in Corneometer scores) and possible evidence of increasing the skin’s ability to retain moisture after only two treatments (as exemplified by a decrease in TEWL scores). Future directions include prospective studies validating these results and further investigating the optimal treatment regimen for improving and/or maintaining skin health with SP-HA.

Statement of institutional review board approval

In compliance with normative documents governing research with humans, the following study including its protocol, supplemental documents, such as patient consent to photography, and the Principle Investigator and clinical site received unconditional approval from the independent review board “Veritas IRB”. A waiver of informed consent was granted by the IRB based on normative guidelines and Health Canada regulations of the Tri-Council Policy Statement articles 3.1–3.5, as this study posed no risk to the welfare of the participants.

Acknowledgments

Presented at: IMCAS AMERICAS, March 2017, Cancun, Mexico. This study was funded by an unrestricted educational grant from Galderma Canada.

Author contributions

AN: principal investigator, article preparation; KME: study coordinator, data abstraction, analyses, article preparation. All authors contributed toward data analysis, drafting and critically revising the paper and agree to be accountable for all aspects of the work.

Disclosure

The authors wish to disclose the following facts which may be considered as potential conflicts of interest: Dr. Andreas Nikolis is a consultant speaker and research collaborator for Allergan, Galderma, and Merz Pharma. Kaitlyn M Enright reports no conflicts of interest in this work.

References


