PN-HPT® (Polynucleotides Highly Purified Technology) in facial middle third rejuvenation. Exploring the potential

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Abstract

Background: Wasting of soft tissues leads to flattening and deflation of the aging midface skin. Polynucleotides Highly Purified Technology (PN-HPT®) demonstrated dermal hydration and elasticity as well as fibroblasts vitality and activity.

Aims: To probe the value of PN-HPT® in middle third rejuvenation in an open-design, exploratory prospective cohort study in 40 real-life ambulatorily treated women.

Methods: Three treatment sessions—at baseline (2-ml prefilled syringe containing 10 mg/ml PN-HPT®, 10 mg/ml hyaluronic acid, 200 mM mannitol) as intradermic gel, and after 3 weeks (2-ml prefilled syringe containing 20 mg/ml PN-HPT® intradermic gel) and 6 weeks (same treatment as baseline). The protocol allowed supplemental treatment with specifically formulated PN-HPT® (7.5 mg/ml) when needed in periorcular and eyelid areas. Assessments: sequential photographs of the facial middle third at baseline, third treatment session and 6–8 weeks after the third treatment session; scoring of overall skin quality and texture, skin quality determinants (wrinkles and skin roughness, skin elasticity, skin brightness), scar appearance, and subjective satisfaction with impromptu 10-cm Visual Analogue Scales.

Results: Significant improvement of overall skin quality and texture (from 7.0 ± 1.06 at baseline session to 7.8 ± 0.99 at follow-up), associated with highly significant improvements of wrinkles and skin roughness, elasticity, and brightness (-17.1%, +39.6%, and +51.1%, respectively). The severity scores of post-acne scars decreased from 7.6 ± 1.32 to 4.2 ± 2.13. Individual satisfaction score at the end of treatment: 0.8 ± 0.28.

Conclusions: PN-HPT® candidate as a valuable option for facial middle third rejuvenation. Further trials will hopefully confirm these early results.

KEYWORDS
acne scars, middle-third rejuvenation, Polynucleotides Highly Purified Technology, PN-HPT™, skin booster, collagen deposition
INTRODUCTION

All strategies aimed at rejuvenating the aging midface skin must compare the youthful and elderly midface skin characteristics. The facial middle third aging look appears segmented by dark concavities and shadows due to the changes in how soft tissues distribute and the persistent tethering tensions by the orbital ligament, malar septum, and McGregor’s patch. The youthful middle third of the face appears convex, with skin appearing luminous from the lower eyelid down to the nasolabial fold. In the aging midface, soft tissues no longer mask the inferior orbital rim due to wasting over time, and the lower eyelid does not gently transition into the cheek. The same is true of the concavities in the malar area and the dark shadows due to soft tissue loss over the zygomatic arch. Non-cutaneous age-related modifications also contribute, as is the case for the nasolabial fold deepening associated with the canine fossa recession due to lost bone tissue. Reversing the effects of the soft tissue loss in the inferior orbital rim will reunify the lower eyelid and cheek segments, reduce the tethering effect caused by facial ligaments, and minimize the segmenting shadows and concavities of those areas (Figure 1A,B). Concomitantly, counteracting the soft tissue waste in the malar septal depression will recreate a convex cheek with an intense highlight.

The agents used as usual lifters in the aging facial middle third—hyaluronic acid, calcium hydroxyapatite with lidocaine, polymethylmethacrylate microspheres, poly-L-lactic acid—are volume enhancers of transient efficacy. Sometimes they have limitations, such as the risk of nodules after poly-L-lactic acid infiltration in the medial and lateral cheeks, the malar and submalar areas, the preauricular region, and earlobes. Natural-origin, highly purified polynucleotides from trout gonads, known as Polynucleotides Highly Purified Technology (PN-HPT®), are another option that has emerged over the last years to counteract the aesthetic shortcomings of the soft tissue waste associated with aging. Thanks to advanced purification and sterilization techniques, PN-HPT® are free of protein contaminants and allergic potential.

Highly hydrophilic PN-HPT® have hydrating effects and rapidly act as volume enhancers. In addition, PN-HPT® also prime dermal fibroblasts to produce new matrix glycosaminoglycans and new collagen and elastin fibrils (Figure 2). Based on this evidence, it is rational to expect PN-HPT® to induce favorable and persistent benefits in the middle third skin quality and texture.

The report presents and discusses the aesthetic outcomes in a prospective cohort of women with face middle third aging skin, treated with several PN-HPT® and PN-HPT® + hyaluronic acid gel co-formulations for general rejuvenation and specific treatment of eyelids and other delicate areas. The authors chose to administer PN-HPT® and hyaluronic acid co-formulations at the first treatment session to exploit the transient but immediate replenishing effect and rapid extracellular matrix hydrating action of hyaluronic acid—as shown by previous studies, a prerequisite for the full deployment of the favorable PN-HPT® effects on fibroblasts.
Aims and procedures of face middle third rejuvenation per district or imperfection

1.1.1 | Tear trough deformities

The general aim was to eliminate infraorbital hollows and other local areas of tissue depression. A complementary goal was softening prominent malar bags with infiltrations onto the orbital rim and in the proximity of the zygomatic-cutaneous ligament. Due to anatomical difficulties and the risk of damage to the lacrimal apparatus, investigators usually performed intradermal injections (PN-HPT® + hyaluronic acid and single-agent PN-HPT®) by the micro-wheal technique followed by digital pressure, performed with care and patience. The study protocol also allowed injections according to the linear threading technique with needles or the fanning technique with cannulas. Application of a low-concentration lidocaine cream was helpful when occasional pain developed in the lower lid area or other middle third districts. Most relevant in this sensitive area, a general recommendation was to inform and reassure the subject about some expected temporary bruising and swelling, sometimes appearing after some days of delay.

1.1.2 | Inferior orbital rim

The general aim in this area was to eliminate shadows and hollows; it was also to restore the youthful cheek convexity by re-establishing the smooth transition between the lower eyelid and the cheek. Infiltrating PN-HPT® + hyaluronic acid and single-agent PN-HPT® in the malar septal depression was the usual procedure, complemented by injections all around the arch to restore the soft contours of the zygomatic arch. Some occasional bruising and swelling may develop in the lower lid area after infiltrations of large doses.

1.1.3 | Nasolabial fold and elongated earlobes

Linear injections along the nasolabial fold were the choice PN-HPT® administration technique, with possibly more injections laterally in women with ptotic cheeks. Care was maximum in the nasal alar crease area crossed by the angular artery. Some individuals also underwent ear lobe rejuvenation, mainly if earring drooping was a problem, with minute injections underneath the pierced earlobe hole.

1.1.4 | Malar prominence and submalar area

Associating both PN-HPT® and hyaluronic acid could reduce the risk of exaggerated midface skin lifts and jowl effacement. In some subjects, the large skin area allowed subcutaneous injections in the pre-periosteal plane by the fanning technique with cannulas. Avoiding the
injected mic gel designed explicitly for eye contour in 2-ml prefilled syringes).

To restore the youthful cheek fullness, the point receiving the higher doses of PN-HPT® plus hyaluronic acid or single-agent PN-HPT® was that of the maximum malar cheek protrusion, located at the intersection of the two virtual lines drawn from the tragus to the nasal ala and from the lateral ocular canthus to the oral commissure. Some subjects also needed rejuvenating injections in the submalar area after losing the soft contour between the lower lid and the cheek. Injections near the nose required care to avoid the risk to pierce the angular artery.

2 | METHODS

2.1 | Rationale, study purpose, and study design

A real-life exploratory investigation was the concept that led to the study design, the first one with PN-HPT® explicitly aimed at facial middle third rejuvenation. The authors accepted that the exploratory study concept could justify the open, prospective cohort design, with the provision that further controlled randomized studies will be crucial to confirm any tentatively favorable outcome arising from this investigation.

The study enrolled 40 real-life women seeking aesthetic treatment for facial rejuvenation who were ambulatorily treated over three months, from April until June 2020. All enrolled women agreed with written informed consent to the anonymous elaboration and research use of their data and photographic evidence. All women also agreed to the anonymous publication of the study outcomes, while the authors carried out all procedures in complete agreement with the Helsinki Declaration principles.

2.2 | PN-HPT® formulations and rationale of use at different sessions; injection technique

The investigators underwent a preliminary half-day practical training meeting to uniform their intradermal injection procedures and avoid introducing a bias related to variable injection techniques. All intradermal formulations used in the study were Class III CE 0373 medical devices (Mastelli Srl).

At the first treatment session, all subjects received PN-HPT® co-formulated with hyaluronic acid intradermally to maximize the immediate volume-enhancing effect and prime the dermal fibroblasts to full PN-HPT® activation over the following treatment sessions.9,10 Using specifically designed, low-concentration, single-agent PN-HPT® formulations in delicate areas such as eyelids and the periorcular district minimized the incidence of wheals occasionally occurring with more concentrated PN-HPT® formulations with or without hyaluronic acid.9,11 The unique low-dose PN-HPT® formulations used in such delicate areas were Plinest® fast (7.5 mg/ml of PN-HPT® in 2-ml prefilled syringes) and Plinest® Eye (7.5 mg/ml of PN-HPT® intradermic gel designed explicitly for eye contour in 2-ml prefilled syringes).

Intradermal infiltrations usually required thin 30-G or 32-G needles and the depot technique with micro-wheals spaced 0.5–1 cm. If esteemed of benefit, the study protocol also allowed using the retrograde linear infiltration technique, mainly when confronted with superficial and medium dermis wrinkles and Langer lines and broader areas such as the nasolabial folds. The cross-link (reticular) technique (net of linear intersecting infiltrations) was also valuable over large skin areas such as the cheeks.

The use of flexible cannulas (diameter 27-G, length 37 mm) fanning from the same entry point was helpful in some subjects to minimize the needle-associated discomfort and bruises, mainly in the periorcular and periallabial areas, the repeated repositioning of needles to treat extensive skin areas, the slow recovery, and the less uniform technique.12

2.3 | Doses according to the facial middle third area undergoing treatment

2.3.1 | General middle third and perioral rejuvenation

The sequential treatment schedule began with Newest® (10 mg/ml PN-HPT® + 10 mg/ml hyaluronic acid in 2-ml prefilled syringes), one vial at the baseline treatment session, followed by high-dose PN-HPT®-only Plinest® (20 mg/ml PN-HPT® in 2-ml prefilled syringes), one vial after three weeks, and a second Newest® injection after three more weeks (i.e., 6 weeks after the baseline session). The treatment schedule agreed with the recent suggestions by the Polynucleotides-HPT™ Priming Board—PN-HPT® or PN-HPT® + hyaluronic acid every 2–3 weeks for 3–4 infiltrations followed by Plinest® or Newest® every one or two months for long-term maintenance.13

At-home self-treatment for two months with nucleotides-based creams and other formulations, including oral food supplements such as Plinest® Care In (Mastelli Srl), an anti-oxidant nutritional supplement with nucleotides, vitamin C, vitamin E, Coenzym Q, folic acid, zinc (two tablets per day), complemented the ambulatory intradermal injection program.

2.3.2 | Eyelid and periorcular revitalization

The same mainstay treatment applied with single-agent, low-dose PN-HPT® formulations specifically designed for thin and sensitive areas (Plinest® fast and Plinest® Eye; composition previously described) and used on-demand according to local skin aging and rejuvenation needs. The authors used such specific formulations in 18 study women in the first session, 21 in the second, and 23 in the third. According to the Consensus Report by The Polynucleotides-HPT™ Priming Board, the usual treatment schedule of single-agent PN-HPT® formulations for sensitive areas is one session weekly or every two weeks for 3–4 infiltrations, repeating the cycle every 1–3 months for long-term maintenance.13
2.4 Assessments

Investigators used specifically designed 10-cm Visual Analogue Scales to assess the aesthetic impact of PN-HPT® treatment on “Overall skin quality and texture,” “Wrinkles and skin roughness,” “Skin elasticity,” “Skin brightness,” and “Post-acne scars (aesthetic severity).” Although subjective and lacking formal validation and demonstrated internal consistency and reproducibility, impromptu VAS scales were deemed adequate for an exploratory investigation. Evaluations included a novel subjective satisfaction questionnaire specifically designed for the aesthetic practice (Table 1)¹⁰ and an impromptu four-score pain, discomfort, irritation, and intolerance assessment (0 = None, 1 = Mild, 2 = Moderate, 3 = Severe). One aesthetic medicine specialist per center collaborating with the authors as an independent evaluator not directly active in treatment procedures performed assessments at baseline and the third treatment session after six weeks and 6–8 weeks after the end of the treatment cycle. The authors agreed on definitive scores only after jointly re-examining the high-definition images taken at each visit at three re-assessment meetings during the study.

2.5 Statistics

The overall analysis adopted a conservative, paired-difference approach (within-group Wilcoxon sign-rank test to assess the differences in population mean ranks. The non-continuous nature of the parameters (semi-quantitative scores), the lack of assumptions about the normal distribution of baseline and final data and the difference between two samples’ means, and the relatively small number of subjects in the prospective cohort justified the conservative non-parametric approach. The test statistics W, sums of the signed-ranks distributed around an expected value of zero and estimated with a standard Excel spreadsheet for Mac (version 16.49, update 2021), were compared to tabulated critical values.¹⁴

Probing for differences in PN-HPT® efficacy according to baseline skin aging (covariate) was the second step of the analysis, based on post hoc stratification (skin aging Goglau score ≤3 vs skin aging score ≥4) and analysis of covariance.

3 RESULTS

Table 2 illustrates the cohort baseline demographics and the baseline severity of the real-life women’s aging facial middle third. Fifteen women (37.5%) complained of aesthetically severe post-acne scars at baseline (mean severity VAS score attributed by the investigators, 7.6 ± 1.32). All enrolled women completed the study.

3.1 The aesthetic impact of treatment

Figure 3A (cohort women younger than 70) and 3B (cohort women older than 70) show some examples of the aesthetic outcomes after

| Table 1 | New rapid individual satisfaction questionnaire used in the exploratory study¹⁰ |
| --- | --- |
| **Satisfaction category** | **Subject’s statement** | **Score** |
| Not satisfied | I feel I look worse than before treatment | 0 |
| | My relatives and I do not appreciate any difference before and after treatment | 1 |
| | I perceive only minimal differences before and after treatment—My relatives do not see any difference | 2 |
| | I see moderate differences before and after treatment—My relatives perceive only minimal differences | 4 |
| Moderately satisfied | My relatives and I see moderate aesthetic differences before and after treatment | 6 |
| Satisfied | I see significant aesthetic differences before and after treatment—My relatives see moderate differences | 8 |
| Very satisfied | The aesthetic outcome goes beyond my expectation—All my relatives see significant aesthetic improvements | 10 |

| Table 2 | Baseline demographics of enrolled real-life women and baseline aesthetic severity of the women's facial middle third |
| --- | --- |
| **Demographic parameter** | **Baseline score** |
| Age, years | 52.7 ± 11.60 |
| Weight, kg | 60.2 ± 4.34 |
| Skin phototype (Fitzpatrick scale, score range 1–5) | 3.0 ± 0.42 |
| Skin photoping (Goglau scale, score range 1–4) | 2.9 ± 0.69 |
| Post-acne scars (aesthetic severity, n = 15; 10-cm VAS scale) | 7.6 ± 1.32 |
| Overall skin quality and texture (10-cm VAS scale) | 7.0 ± 1.06 |
| Wrinkles and skin roughness (10-cm VAS scale) | 8.2 ± 1.42 |
| Skin elasticity (10-cm VAS scale) | 4.8 ± 1.21 |
| Skin brightness (10-cm VAS scale) | 4.5 ± 1.11 |
the sequential treatment with PN-HPT® + hyaluronic acid and PN-HPT® alone. The examples show the baseline middle third picture with the situation at follow-up after the treatment cycle. The examples illustrated in Figure 4 compare the aesthetic picture at baseline, the final session, and follow-up.

At the end of the PN-HPT® treatment cycle, the individual satisfaction score was 8.0 ± 0.87 out of a theoretical maximum of 10. Based on independent evaluations of photographs and overall assessment of skin texture and skin roughness, elasticity, and brightness, Figure 5 illustrates how the overall skin quality and texture comprehensively improved between baseline and follow-up (+10.1%; \( p < 0.05 \) vs. baseline). The VAS scores for “Wrinkles and skin roughness,” “Skin elasticity,” and “Skin brightness” as determinants of overall skin quality also showed highly significant improvements at follow-up compared with baseline (-17.1%, +39.6%, and +51.1%, respectively; Figure 6A,B). The same was true for the severity of post-acne scars in the 15 women with aesthetically severe baseline lesions (scars score at follow-up: 4.2 ± 2.13, -44.7%; \( p < 0.01 \) vs. baseline).

Regarding baseline skin aging, the PN-HPT® efficacy appeared somewhat higher in women with more severe skin aging with baseline Goglau scores ≥4. Covariance analysis after post hoc stratification showed significant improvements for all skin quality determinants in women with more advanced skin aging, but improvements only for skin elasticity and skin brightness in women with Goglau skin aging score ≤3 (Table 3).

3.2 | Safety

The mean tolerability score at the end of the treatment cycle was 0.8 ± 0.28. When occasionally observed (in 11 women out of 40), all side effects were no more severe than mild and transient discomfort, irritation or pain at the injection site. No woman reported systemic adverse events.

4 | DISCUSSION

The facial middle third is often the starting point for countering the soft tissue loss in the aging face. Traditional fillers rapidly provide gratifying outcomes as volume enhancers of facial folds and throughs and fine rhytides, although well-known side effects may be a burden. In the lower eyelid and tear trough area, fillers based
on hyaluronic acid may have the risk of post-injection edema, the disturbing bluish hue through the skin known as the Tyndall effect and of clumping in the tear trough area when smiling. On the other side, visible fibroplasia and nodules are always a risk with poly-L-lactic acid fillers. Aesthetic benefits with mere volumetric fillers are also more or less rapidly ephemeral.

PN-HPT®, formulated as Class III CE 0373 medical devices, might help with those problems thanks to their activating potential on dermal and connective fibroblasts. The working hypothesis that inspired the exploratory study was that PN-HPT® priming of dermal connective and subcutaneous fibroblasts would lead to long-term re-modeling and natural restoration of depleted facial middle third soft tissues.

PN-HPT® must not be confused with another class of polydeoxyribonucleotide derivatives, synthetically described with the acronym PDRN, which are biochemically and clinically much different from the PN-HPT® so far discussed. PDRN are available only on prescription (they are not available as non-prescription medical devices). Their main approved indication is the treatment of torpid wounds and diabetic foot ulcers; PDRN development is on course also in other disease conditions such as inflammatory bowel diseases. Even more importantly, PDRN stimulate Vascular-Endothelial Growth Factor (VGEF) production in pathological conditions of low tissue perfusion and have a distinctly specific molecular mechanism of action via the A2 purinergic receptor. Moreover, PDRN administration is intramuscular.

Conversely, PN-HPT® are only available as no-prescription medical devices for topical and intradermal administration. Besides filling intradermal spaces, providing hydration, and plumping up tissues, most likely PN-HPT® act (indirectly) by passively replenishing the dermal pool of nitrogen bases, nucleosides, and nucleotides, as discussed in a recent consensus report in aesthetic medicine and several PN-HPT® clinical studies.

Previous studies have shown that PN-HPT® might promote a more robust extracellular matrix and collagen and elastin production, which may help restore the youthful midface convexity persistently over the long term. Concomitantly, hydration of infiltrated tissues would rapidly improve the midface aesthetics thanks to the hydrating and volume-enhancing effects on coarse wrinkles,
shadows caused by the depletion of soft tissues, and fine rhytides and facial folds. In addition, as polyanion polymers, PN-HPT® are effective water binders that re-orient their spatial structure and orientation after binding water molecules in dermal tissues. The resulting three-dimensional viscoelastic PN-HPT® gel showed to hydrate the extracellular matrix for prolonged periods, although no direct comparison with hyaluronans exists. The co-formulation of PN-HPT® and hyaluronic acid injected at the first treatment session is likely to hydrate even more effectively.

Although conceived only as exploratory, this first study with PN-HPT® in facial middle third rejuvenation supports these hypotheses. The overall skin quality and texture improved significantly from baseline until the third treatment session and after 6–8 further weeks. Skin quality and texture determinants—skin roughness, skin elasticity, skin brightness—improved even more significantly. The women’s photographs confirm the tendency of the aging midface platform, appearing segmented by local concavities and shadows, to shift to a more youthful convex platform, with the re-creation of

**TABLE 3** PN-HPT® efficacy on determinants of overall skin quality stratified for baseline facial middle third skin aging

| Skin parameter                  | Goglau skin aging score ≤3 baseline vs end-of-treatment | Goglau aging score ≥4 baseline vs. end-of-treatment |
|--------------------------------|--------------------------------------------------------|--------------------------------------------------|
| Wrinkles and skin roughness    | 7.9 ± 1.15 vs. 6.5 ± 1.15                               | 9.7 ± 0.49* vs. 7.5 ± 0.53                       |
| Skin elasticity                | 5.4 ± 1.07* vs. 7.0 ± 0.73                               | 3.7 ± 0.49* vs. 6.0 ± 0.0                        |
| Skin brightness                | 4.9 ± 1.27** vs. 7.3 ± 0.86                              | 4.0 ± 0.0** vs. 6.0 ± 0.0                        |

*p < 0.05 vs. baseline.

**p < 0.01 vs. baseline
the dominant cheek highlight typical of the younger women’s face complexion. The improvement was most substantial for post-acne scars, with the relevant VAS scores almost halving at the end of the treatment cycle.

Regarding the overall skin quality determinants, PN-HPT® always appeared effective, although with minor differences related to baseline middle third skin aging severity. In women with baseline Goglau skin aging scores ≥4, improvements were invariably significant or highly significant for every overall skin quality and texture determinant—wrinkles and skin roughness, skin elasticity, and skin brightness. Comparatively, only wrinkles and skin roughness improved significantly in women with baseline Goglau skin aging scores no higher than three. The lack of clinically significant side effects, with only mild, transitory discomfort at the injection site and occasional bruising in less than one-fourth of women, confirms the absence of clinically significant side effects, with only mild, transitory discomfort at the injection site and occasional bruising in less than one-fourth of women, confirms the previous observations about the high PN-HPT® safety in real-life aesthetic practice and clinical studies.13

The study has several weak points. The lack of controls is the most important one. The hypothesis-generating exploratory nature could provisionally justify this liability, but confirming the tentative outcomes arising from this probing investigation will need studies with a sounder controlled and randomized design. A second methodological weakness was using arbitrary VAS scales with no formally validated reproducibility and possibly lacking internal consistency for all skin quality assessments. Once again, using impromptu VAS scales may be temporarily acceptable in exploratory studies like this one—but only temporarily. The preliminary training sessions organized to uniform how investigators performed intradermal injections might be another weak point. These joint sessions probably helped reduce the bias related to the variable techniques of each investigator, but possibly some bias persisted—unknown and impossible to control.

The real problem in assessing skin quality is balancing the regulatory requirements for medical devices with the women’s subjective perception of aesthetic benefits. In other words, the real question is confirming that the objective clinical assessment and subjective perception are mutually consistent. Human studies investigating skin quality and texture have already used VAS scales and scores, but only in combination with validated tools and assessment devices to assess skin hydration and skin surface profilometry.25 Future non-exploratory investigations on PN-HPT® in facial middle third rejuvenation will have to use such validated instruments.26–29 Future controlled trials will also help define the “priming” concept and the PN-HPT® and HA relative contributions to outcomes. Finally, an interesting point that deserves further investigation is whether the PN-HPT® properties on fibroblast vitality and dermal trophism could partially substitute autologous lipofat transfer to correct the midface and lower orbit tissue loss, the most visible age-related facial changes.

5 | CONCLUSIONS

Could the properties and outcomes of PN-HPT® on dermal trophism and hydration, fibroblast vitality, and overall skin quality and texture go beyond the mere volumetric effect of traditional fillers in facial middle third rejuvenation? This very first exploratory investigation with PN-HPT®, as a single active principle or in combination with hyaluronic acid, explored the issue. The study suggests that PN-HPT®-based medical devices may be valuable tools to correct the periorbital hollowing, accentuated tear trough deformity, the facial middle third flattening, the lengthening of the lower eyelid, and the too highlighted bony projections—all aesthetically disturbing features of the aging facial middle third. However, only future well-designed controlled studies will confirm such provisional study outcomes.

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Mastelli S.r.l., Sanremo, Italy, is the producer of PN-HPT® (Highly Purified Technology) as well as Newest®, Plinest®, Plinest® fast and Plinest® Eye, the gel formulations of injectable polynucleotides with or without hyaluronic acid used in this exploratory study. The authors wish to acknowledge the contribution of Mastelli S.r.l. for financial support with the publication costs.

CONFLICT OF INTEREST

The authors declare they have no conflict of interest inappropriately influencing the representation or interpretation of reported research results. Mastelli Srl, Sanremo, Italy, holder of the PN-HPT® patents and producer of the investigated medical devices, will limit its support to contributing to the manuscript publication costs.

AUTHOR CONTRIBUTION

All authors sought and got informed consents from the women subjects seeking aesthetic treatment of their aging facial middle third and enrolled in the study. All women received informed consents about the benefits and risks they could reasonably expect from aesthetic procedures with PN-HPT® formulations with or without hyaluronic acid. All authors performed all aesthetic procedures under the first author’s supervision. All authors are accountable for the clinical and editorial work’s accuracy and integrity, leading to the manuscript’s submission to the Journal of Cosmetic Dermatology, including all comments on outcomes.

ETHICAL APPROVAL

The authors performed the study in complete agreement with the Declaration of Helsinki. No unusual risk was reasonably foreseeable in participating subjects other than those incurred with the ambulatory aesthetic medicine procedures routinely performed over the aging facial middle third. Furthermore, the study protocol excluded any unusual practice different from infiltrations of PN-HPT®, a long-established aesthetic medicine procedure supported by a wealth of scientific literature. Regulations allowed waiving the requirement for formal approval by an Ethics Committee.

PHOTOGRAPH CONSENT STATEMENT

All photographs belong to the authors, who permitted their submission with the patients’ agreement after making them unrecognizable with black bars.
DATA AVAILABILITY STATEMENT

The datasets generated and analyzed in the study are not publicly available. However, the author named the second one currently archives all datasets with full details of participating subjects according to current regulations. After conversion in an anonymous form, all datasets are available from the named author on reasonable request.

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