Hyaluronan Hydrogels for Injection in Superficial Dermal Layers: An In Vitro Characterization to Compare Performance and Unravel the Scientific Basis of Their Indication

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Abstract: Background: Skinboosters represent the latest category of hyaluronan (HA) hydrogels released for aesthetic purposes. Different from originally developed gels, they are intended for more superficial injections, claiming a skin rejuvenation effect through hydration and possibly prompting biochemical effects in place of the conventional volumetric action. Here, three commercial skinboosters were characterized to unravel the scientific basis for such indication and to compare their performances. Methods: Gels were evaluated for water-soluble/insoluble-HA composition, rheology, hydration, cohesivity, stability and effect, in vitro, on human dermal fibroblasts towards the production of extracellular matrix components. Results: Marked differences in the insoluble-hydrogel amount and in the hydrodynamic parameters for water-soluble-HA chains were evidenced among the gels. Hydration, rigidity and cohesivity also varied over a wide range. Sensitivity to hyaluronidases and Reactive Oxygen Species was demonstrated allowing a stability ranking. Slight differences were found in gels’ ability to prompt elastin expression and in ColIV/ColII ratio. Conclusions. A wide panel of biophysical and biochemical parameters for skinboosters was provided, supporting clinicians in the conscious tuning of their use. Data revealed great variability in gels’ behavior notwithstanding the same clinical indication and unexpected similarities to the volumetric formulations. Data may be useful to improve customization of gel design toward specific uses.

Keywords: glycosaminoglycans; hyaluronan; dermal filler; hydration; rheological behavior; hyaluronidase; degradation by reactive oxygen species; human dermal fibroblasts

1. Introduction

Facial injection of hyaluronan (HA)-based gels for aesthetic purposes is well-established and HA crosslinked with 1,4 butandiolediglycidylether (BDDE), suspended in physiologic solution, is generally employed [1–25]. The use of the BDDE-HA hydrogel typically relies on its elastic behavior, high hydrophilicity and improved resistance to hyaluronidase action, compared to unmodified HA. The elastic behavior is responsible for gel capacity to maintain volume while deforming under the stress of facial movements thus assuring a filling, natural-looking effect; network hydrophilicity improves tissue hydration at the injection site while the higher (but not complete) resistance to enzymatic degradation, compared to natural-occurring HA, assures longer in vivo permanence while preserving bio-absorption.

Since the launch of the first products, the suggested clinical use for this type of hydrogels has changed. Volume restoration, based on the gel physical “filling” effect, has been...
considered the main indication for more than two decades \cite{2,10–14,19,22}. Considering that gel-projection capacity is related to its stiffness, \(G'\) (Storage Modulus) has long been the only parameter considered in selecting the most suitable gel for the specific clinical need. A more recent approach has been extended to the whole fluido-dynamic gel behavior as well as to its sensitivity to enzymatic and radical degradation and to other properties (e.g., hydration capacity, cohesivity) \cite{1,5–8,14–25}. All of the latter features are known to contribute to the in vivo effect after injection and/or over time, and they are therefore generally studied to predict relative in vivo performance and to provide clinicians with valuable information to select treatments and optimize outcome \cite{1,5–8,14–25}. Additionally, due to these studies, HA-BDDE gel design has been improving. As a consequence, hydrogels are now tuned towards more specific and differentiated uses to meet diverse clinical needs finally aiming at a full face restoration \cite{7,8,21,26,27}. “Skinboosters” are the latest category of HA gels that entered the market. Unlike typical “volumetric” fillers, skinboosters are intended for the injection into more superficial (intradermal vs. deep dermis-periosteum) skin layers and are expected to improve skin appearance and texture rather than performing by a projection effect. Specifically, improvement in skin hydration and elasticity, and possibly a boost in extracellular matrix molecules biosynthesis is claimed after the application of such gels \cite{28–33}. A stretching effect, referring to wrinkle distension from gel hydration, that is anyway filling the volume loss to a certain extent, is also predicted \cite{30}.

Former studies and literature report extensively analyzed volumetric gels and compared available formulations, suggesting opportunities toward design optimization, and supporting their appropriate selection and use \cite{1,5–8,14–25}. On the contrary, no similar studies have been carried out so far to scientifically assist this recent “skinboosting” approach. The aim of the study was to assess a panel of biophysical and biochemical features for this specific type of hydrogel. For this purpose, three gels among the currently marketed crosslinked HA-based skinboosters (i.e., Restylane Vital (RV), Juvederm Volite (JV), Visco-derm Hydrobooster (HB)) were evaluated for their composition in water-soluble/insoluble HA, rheological behavior, hydration capacity, cohesivity, sensitivity to degradation and ability to induce, in vitro, human dermal fibroblasts towards enhanced production of extracellular matrix components. Beyond increasing our knowledge of these gels, results allow the comparison of gel performance. Further, they may shed light on the possible specific features for the HA-BDDE gels to be classified as a skinbooster.

2. Results

2.1. Water-Soluble Fraction of Fillers: Quantitative Determination and Hydrodynamic Analysis

Total HA concentration in each gel is provided by the manufacturer (Table 1). The water-soluble fraction of the biopolymer was quantified here, allowing us to derive the specific composition in water-insoluble/water-soluble HA (Table 1). Water-soluble HA was found in each formulation and in a comparable amount (about 4 mg/mL; \(p > 0.05\)). As a result of the diverse biopolymer total concentration, the gels significantly differ for the water-insoluble HA content. Specifically, the insoluble HA fraction in HB was far higher than the one found in the other gels, up to 2.6 fold higher, compared to JV. As for RV, the insoluble HA content almost doubled compared to JV.

| HA (mg/mL) | RV | JV | HB |
|-----------|----|----|----|
| total     | 20 | 12 | 25 |
| Water-soluble | 4.7 ± 0.6 | 4.0 ± 0.6 | 4.2 ± 0.9 |
| Water-insoluble | 15.4 ± 0.6 | 8.0 ± 0.6 | 20.8 ± 0.9 |
| Soluble HA (wt%) | 23 ± 3 | 33 ± 5 | 17 ± 4 |

HA in each gel and hydrodynamic parameters for the water-soluble HA fractions as derived from the Size Exclusion Chromatography-Triple Detector-Array (SEC-TDA) analyses. The total HA concentration reported is the value indicated in the product’s package insert.
The SEC-TDA analyses revealed the presence of a 525 ± 23 kDa HA in H_B (Table 2). Shorter HA chains, of about 260 kDa M_w, were found in R_V and HA of ~160 kDa was found in J_V. The intrinsic viscosity and the hydrodynamic radius values varied consistently. The M_w/M_n values suggested for R_V and, especially J_V broader distributions compared to H_B one.

### Table 2. Weight average molecular weight (M_w), numeric average molecular weight (M_n), polydispersity index (M_w/M_n), intrinsic viscosity ([η]) and hydrodynamic radius (R_h).

| Sample | M_w (kDa) | M_n (kDa) | M_w/M_n | [η] (dL/g) | R_h (nm) |
|--------|-----------|-----------|---------|-----------|---------|
| R_V    | 266 ± 29  | 109 ± 19  | 2.5 ± 0.6 | 5.7 ± 0.1 | 26.8 ± 1.1 |
| J_V    | 161 ± 21  | 49 ± 5    | 3.3 ± 0.2 | 3.4 ± 0.2 | 17.9 ± 1.0 |
| H_B    | 525 ± 23  | 305 ± 55  | 1.7 ± 0.2 | 7.5 ± 0.4 | 37.2 ± 1.7 |

By comparing the Mark–Houwink–Sakurada (MHS) curves (log intrinsic viscosity vs. log molecular weight) derived for the soluble fractions to the ones obtained for linear HA samples (Figure S1), a lower intrinsic viscosity over all the molecular weight distribution, was found for the analyzed samples, thus indicating a more compact conformation, consistent with modified chains. Thus, we can assert that soluble HA fractions contain BDDE-HA molecules.

#### 2.2. ^1^H-NMR Analyses

The ^1^H-NMR spectra obtained for the gels are shown in the Supplementary Material (Figure S2). The BDDE/HA (disaccharide unit) molar ratio was quantified by integrating the signal at δ 1.6 ppm, due to the aliphatic (CH₂)₂ moiety of the BDDE molecule, with respect to the HA N-acetyl signal at δ 1.9 ppm (ref). The BDDE/HA ratio (mol %) was 1.1, 7.0 and 9.5 for R_V, J_V and H_B respectively.

#### 2.3. Hydration Capacity

Water absorption was measured to compare the gel hydration capacity. Data reported in Figure 1 demonstrated that gels were able to hydrate and, therefore, to expand, when incubated in a physiological medium. This indicated that the commercialized formulations are not at the equilibrium swelling. Specifically, as reported in Figure 1, when allowed to equilibrate in Dulbecco’s Phosphate Buffered Saline (PBS), R_V almost doubled its volume and H_B more than tripled it. J_V showed the lowest hydration extent (p < 0.05 vs. R_V and p < 0.001 vs. H_B).

![Figure 1.](image_url) Gels’ hydration ability. Volumetric expansion (gel volume at equilibrium/gel volume in the syringe) occurring when the formulations are equilibrated in PBS; data indicate the final volume reached by 1 mL of the formulations when allowed to hydrate in PBS.
2.4. Rheological Parameters

The rheological characterization confirmed, as expected, an elastic behavior for all the gels, with tan delta values in the range 0.2–0.6. Further, results indicated that the gels were not comparable for rigidity and, surprisingly, G' values varied over a wide range (Figure 2a). RV's stiffness was far higher, while HB behaved as the less rigid gel. Specifically, G' values at 0.7 Hz frequency were about 40 Pa for HB, about 120 Pa for JV and as high as 430 Pa for RV. The complex viscosity data confirmed the typical profile of this type of hydrogels with values constantly decreasing with frequency. RV behaved as the most viscous gel while HB proved the least viscous one (Figure 2b).

![Figure 2. Rheological characterization. G' values (a) and complex viscosity (b) as a function of the frequency. Measurements were performed at 37 °C and 0.1% strain.](image-url)
2.5. Degradation Studies

The degradation studies revealed that all the gels were sensitive to both Reactive Oxygen Species (ROS) and Bovine Testicular Hyaluronidase (BTH) action (Figures 3 and 4). Gel sensitivity to ROS was demonstrated by the rheological behavior in the presence of a ROS generating system (Figure 3), compared to a control. Specifically, G' values were recorded during filler incubation with the H$_2$O$_2$/Cu$^{2+}$ system and after diluting the gel, to the same extent, with water (control) (Figure 3a–c). Dilution with water reflected in a 1.1–1.4 fold decrease in G' values, compared to the data in Figure 2 and, gel rigidity remained constant over time (Figure 3a–c). In the presence of ROS, a marked drop in G' was observed indicating depolymerization (Figure 3a–c). The gels showed diverse degradation profiles. Specifically, RV's stiffness rapidly decreased from about 400 Pa to values lower than 1 Pa, close to the minimum measurable values, thus causing a scattered signal (Figure 3c). A more gradual reduction in G' was observed for JV and for HB with the latter degrading faster (Figure 3a,b). In particular, as indicated in Figure 3d, at 2 min of incubation with the ROS generating system, G' loss (% vs. ctr) was already higher than 99% for RV. At the same incubation time, JV still preserved 98 ± 1% of its stiffness while, for HB, around 20% residual G' was recorded, compared to control. At 5 min of incubation, JV still retained 47 ± 6% G'; about 93% G' loss could be measured only at the longer time tested (8 min).

When the gels were incubated in the presence of BTH, a certain extent of HA solubilization was registered for each sample, indicating sensitivity to enzymatic hydrolysis (Figure 4a). In particular, around 1 mg/mL HA solubilized in 3 h incubation with BTH (2 U/mL) while the amount rose to 1.2–1.4 mg/mL when higher enzyme concentration was used (5 U/mL BTH) for the same time interval. Under the most drastic hydrolysis
conditions tested, around 1.7–2.1 mg/mL of water-soluble fraction increase was recorded. No significant differences in the solubilization extent \( (p > 0.1) \) could be detected among the gels under the applied conditions, only very slightly lower solubilization of \( \text{H}_B \) was recorded under the most drastic conditions. However, based on these solubilization rates and on the initial gels’ composition (Table 1), \( \text{H}_B \) showed the highest amount of residual water-insoluble HA during incubation with BTH, regardless of the enzyme concentration and incubation time (Figure 4b).

The hydrodynamic parameters for the soluble fractions obtained after incubation with BTH are reported in Figure 4c. After 3 h incubation with 2 U/mL BTH, longer soluble chains were found for \( \text{H}_B \) while under the most drastic hydrolysis conditions, slightly lower \( M_w \) values were found for \( \text{R}_v \). With the increase in BTH concentration and incubation time, an increase of the HA amount in solution and of polydispersity and a decrease of the \( M_n \) values were recorded.

| sample | HA (mg/mL) solubilized in the presence of BTH |
|--------|-----------------------------------------------|
|        | 2 U/mL (3h) | 5 U/mL (3h) | 5 U/mL (6h) |
| \( \text{J}_v \) | 1.0±0.2 | 1.5±0.2 | 2.1±0.3 |
| \( \text{R}_v \) | 0.8±0.1 | 1.3±0.1 | 2.1±0.1 |
| \( \text{H}_B \) | 0.9±0.1 | 1.2±0.1 | 1.7±0.1 |

Figure 4. Enzymatic degradation. (a) Amount of HA (mg/mL) solubilized due to the BTH action after 3 h incubation with BTH 2 U/mL and after 3 and 6 h incubation with BTH 5 U/mL. (b) Residual water-insoluble HA (mg/mL) under the same hydrolysis conditions as in (a). (c) SEC-TDA data for the HA water-soluble fraction after 3 h incubation with 2 U/mL and 6 h incubation with BTH 5 U/mL.

2.6. Cohesivity

The cohesivity test revealed great differences in gels’ behavior. \( \text{H}_B \) completely retained its structure in the whole interval time of observation, therefore it was assigned with the highest cohesivity score (“fully cohesive”) in the Gavard–Sundaram Cohesivity Scale (Figure 5) proving the most cohesive gel. On the contrary, \( \text{J}_v \) progressively completely loses definition behaving as a “fully dispersed” gel (cohesivity score less than 1). An intermediate cohesive behavior was found for \( \text{R}_v \) that behaved as a “mostly cohesive” (cohesivity score 4) at the early time of observation then showing an increasing extent of fragmentation (cohesivity score 3, “partially cohesive” gel, at 70 and 90 s of observation).
2.7. Biological Studies

As shown in Figure 6, the gel treatments on fibroblasts showed the absence of any toxicity. Images of cells after 24 h incubation with the gels and of untreated cells are shown in Figure 6a. It was evident that fibroblasts in presence of filler, present a very similar morphology to the untreated cells and also the cell density in each well was comparable. Quantitative results on metabolic activity, reported in Figure 6b, confirmed the total absence of cytotoxicity for all the gels.

To elucidate the effect on matrix synthesis due to HA treatments on human dermal fibroblasts, Type I collagen (COLI), type IV collagen (COLIV) and elastin (ELS) gene expression was quantified by qRT-PCR (Figure 7). Overall, the three biomarkers investigated were upregulated, specifically for HB and RV with respect to untreated cells. COLI was about 2.5-fold and 1.5-fold expression vs CTR in HB and RV treated cells. On the contrary, it was down-regulated by JV. COLIV was upregulated by about 3-fold, 4.5-fold and 5-fold expression over the control following HB, JV and RV treatments, respectively (Figure 7). Finally, ELS increased its expression by about 6-fold, 5.8-fold and 3.8-fold expression in fibroblasts treated with HB, JV and RV respectively. Regarding integrins, mRNA expression showed that all samples increased integrins expression (INTα1 and β1). Specifically, RV upregulated INTα1 by about 4-fold and INTβ1 by 3-fold expression. Instead, INTα1 increased by about 30-fold and 7-fold vs. CTR in presence of JV and HB, respectively. While, INTβ1 mRNA resulted in about a 50-fold increase vs. CTR for both JV and HB, samples.
Figure 6. (a) Morphological observation at MO of Human Dermal Fibroblasts (HDF) in the control and in presence of filler, after 24 h treatment. Scale bar 50 µm; (b) Cell vitality with respect to the untreated cells in presence of JV, RV and HB respectively.

Figure 7. Gene expression analyses: the results are expressed as fold change of HA treated cells respect to untreated cells (CTR) for COLI, COLIV and ELS (a) in human dermal fibroblasts. In addition, INTα1 and β1 (b) were accomplished. Data showed as the averages ±SD.
At 48 h of incubation, HDF were lysed to accomplish Western blotting analyses on specific biomarkers. Results are reported in Figure 8. The graph shows a slight increase in elastin expression in response to RV and HB treatments (respectively 1.09 and 1.31-fold vs. CTR), the latter proved significant vs. control and JV treatment. RV, being less effective than HB proved superior with respect to elastin biosynthesis in JV treatment. This latter showed Col I production to a similar extent with respect to the control, while HB slightly upregulated its expression (1.20-fold vs. CTR). Finally, the cells treated with RV and HB presented a Col IV higher expression in comparison to control and also JV treated cells. Specifically, RV increased Col IV protein level by about 1.38-fold vs. CTR and HB improved it by about 1.52-fold instead, JV seemed to be not effective in the modulation of this biomarker.

![Western blotting](image)

**Figure 8.** Western blotting for expression of collagen I, collagen IV and Elastin normalized to actin in the densitometry reported as average and SD. * stands for significant (p < 0.05) variations respect to CTR, # stands for p < 0.05 for elastin expression in HB treatment vs. RV and JV ones.

### 3. Discussion

Since the first application of hyaluronic acid (HA) injectable implants, the purpose for these treatments has been widened and patients and clinicians now share more consciousness. The final aim of the clinician to satisfy patient expectations without radically changing their face but improving their aspect is obtained by searching for the right product, and the right injection technique within a proposed line of formulations. It has to be considered that the biological diversity of patients and everyday life habits are the major responsible for treatment effectiveness: age, kind of skin, weight loss (or fat compartment), smoking attitude, UV irradiation, have to be considered. However, it has been established that a more precise biophysical description of the HA-based formulations may help in the right product choice according to patient needs. In this respect, scientific evidence has been improving not only to address rheological behavior or chemical aspects (e.g., HA modification degree, molecular weight, etc.) but also biochemical and biological features of the product itself. To this aim, three hyaluronan-based hydrogels were characterized in this research study to bridge the biophysical and biochemical properties to their potential performance as skinboosters and to compare them.

With regard to the gels’ composition, according to the labels, the three skinboosters greatly differ in total HA concentration [31–33]. The amount of total HA even doubles from one gel to another. As found for other similar formulations intended for skin rejuvenation, a certain amount of water-soluble HA was present (Tables 1 and 2) [7,8]. In particular, data indicated comparable concentration in soluble HA (about 4 mg/mL) among the skinboosters but diverse hydrodynamic parameters. Specifically, RV and JV soluble HA chains were more similar in length (160–270 kDa $M_w$) while, the soluble HA fraction in HB presented far higher molecular weight (525 kDa $M_w$), therefore potential diverse biochemical effect could be expected for this gel [34]. For all the samples, the analyses of the MHS curves suggested water-soluble HA chains conformation other than linear thus indicating a certain extent of chemical modification. This is in agreement with previous studies on similar products [15]. Rationally, the water-soluble HA fraction may derive
from the crosslinking process leaving HA chains that, even if crosslinked/modified, are still soluble in an aqueous medium due to low(er) molecular weight.

Gel composition studies also highlighted, among the skinboosters, a far higher amount of water-insoluble HA (hydrogel fraction) for HB thus suggesting potential different biophysical behavior for this gel. The hydrogel fraction in HB was far higher even compared to volumetric gels while, in general, similar or lower water-soluble HA concentrations were found in the skinboosters [7]. Collected data indicated an unexpected similarity (in total HA concentration, composition and soluble HA chains’ hydrodynamic parameters) between the “volumetric” and the “skinbooster” Restylane formulations [7]. Even if less concentrated in total HA, insoluble/soluble-HA composition Juvederm skinbooster resulted also similar to the corresponding volumetric gel with comparable hydrodynamic parameters, as well [7].

Data on gel-water uptake support the claimed in vivo tissue hydration effect. The gels were able to absorb water even more than tripling their weight/volume (HB). However, it is worth underlining that JV increased its volume only by about 30% that was unexpected for the pursued application as a skinbooster. This result is peculiar if compared to the water up-take values generally reported for other dermal fillers with volumetric indication (up to 280% volume increase due to hydration) [7]. HB showed water up-take values close to the highest ones reported for volumetric preparations, while the other two skinbooster preparations showed lower hydration ability, even with respect to the volumetric gels. It has to be considered that being injected at a superficial layer, a very high water uptake is undesirable as it may result in edema, which will reduce patient compliance [21,35]. However, the products that are proposed for more superficial injections need to be easily spreadable, this will ensure distribution of the gel in a larger area, and the water recall in the tissue will be more physiological. Finally, it is worth underlining that collected data do not directly translate into the fillers relative in vivo expansion at the injection site: compression forces exerted by the surrounding tissue, counteracting gel expansion, should also be considered for a more accurate prediction. The rheological data were also surprising since evidencing a huge difference in rigidity notwithstanding the specific proposed application as skinboosters: more than one order magnitude G’ variation was measured. Further, while JV and HB rigidity was, as expected, far lower than the one reported for the volumetric dermal fillers, the G’ value registered for RV was higher than the majority of the volumetric gels and close to the RLift stiffness (RLift is the volumetric gel of the Restylane fillers family) [7]. A low rigidity is associated with high deformability under applied stress, thus reflecting in a reduced or even absent “palpability” of the gels, which is crucial to obtain a natural-looking effect when gels are delivered in superficial skin layers [8,9,14,36,37].

Investigation of gel degradation profile is key since it is related to the in vivo longevity of the gels. All the samples showed sensitivity to BTH and ROS action thus ensuring in vivo resorbability. BTH was selected for this study because of the reported similarities to human hyaluronidases (HAse) [38,39] and because it is commercially available in pure form. BTH action is expected to have two effects: solubilization of the water-insoluble HA hydrogel and hydrolysis of the water-soluble HA chains (reduction of the average molecular weight). Despite the comparable rate of water-insoluble fraction solubilization, when equal volumes of the gels are injected, a longer in situ persistence can be predicted for HB, while JV is expected as the shorter-lasting gel. This is related to the initial differences in total HA amount and insoluble hydrogel fraction that will resist enzymatic degradative conditions, based on the comparable degradation rate. Hydrodynamic analyses revealed that the molecular weight distribution of the soluble HA chains is widened during enzymatic hydrolysis. This is rationally related to the depolymerization of the water-soluble HA chains while progressive solubilization of the hydrogel may be responsible for the passage of longer polymeric chains to the solution. This would also explain the observed variation in the average molecular weight during incubation with the enzyme. The increase in c/Mn ratio (c is the water-soluble polymer concentration (mg/mL) and Mn is the number average
molecular weight) proves, as expected, the increase in the polymer mol number in solution accompanying BTH action [40].

Data on the rheological behavior of the gels under in vitro oxidative stress conditions indicated markedly diverse sensitivity to ROS action. Specifically, based on collected data, the shorter permanence may be predicted for RV while a more gradual loss of the clinical effect may be expected for H8 and, finally, JV showed the highest resistance.

In vitro degradation studies either with ROS or with BTH serve as support to characterize the gels in comparable situations, however, the two phenomena are contemporary acting in vivo, besides the mechanical stress of face muscle movements, therefore the lifetime of the injected gels may be better derived through in vivo study. However, considering the more superficial delivery for skinboosters, and the intradermal penetration of UV radiation, sensitivity to ROS action, poorly investigated so far, has a key role for the gel stability when injected [41].

Skinboosters showed far lower resistance to ROS action than volumetric gels. Compared to the less sensitive volumetric gel, the most resistant skinbooster retained 50% rigidity in about the same time interval but in the presence of half the amount of ROS [7].

Cohesivity was recently defined as gel capacity to not dissociate [20]. Dermal fillers are lately investigated for their cohesive behavior referring to the pilot study by Sundaram and collaborators establishing a scale for gel rating [20]. Availability of cohesivity data on commercialized gels is helpful to clinicians in the selection of the most appropriate product to achieve the specific clinical objective. Fillers with higher cohesivity are considered a better choice for more superficial treatment. They allow spreadability and, therefore, a gel homogeneous distribution within the tissue, without fragmentation, thus avoiding palpability and even the eventually occurring formation of nodules [6,20,21]. H8 was the most cohesive. A better tissue integration pattern can be predicted for this gel based on its high cohesivity/low viscosity profile [6,20,21]. Results obtained in the framework of this study indicated that, despite the similar indication of use, the skinboosters here-characterized showed wide variability in terms of cohesivity (from 1 to 5) similar to our previous findings on volumetric preparations [7]. Even if the parameters affecting the gel’s cohesivity still need to be fully clarified, the strategy used for HA crosslinking may reasonably have an effect on gel’s behavior in this specific analysis.

Soft gels that are aiming at tissue revitalization should improve skin texture besides their effect in filling wrinkles. Their expected lower G’ and viscosity gives less performance than other gels to fill void volume, but this helps in reducing palpability and the Tyndall effect [19,21]. However, it can be argued that a biological effect is desirable beyond the physical effect. For this reason, we studied the gels in contact with human dermal fibroblasts. Data on the metabolic activity of cells, cultured on Tissue Cultured Polystyrene, in the presence of the hydrogels were demonstrated to be consistent with the CE approval of these class III medical devices, with an absence of any toxicity. Biological data showed a slight improvement in key biomarkers for HB and RV and a lower if none effect for JV. Specifically, elastin expression for HB was superior with respect to the other treatments, and ColIV expression with respect to Coll was improved in RV and H8 treatments. Comparison with data obtained for volumetric gels did not reveal a stronger biochemical effect for skinboosters on Coll and elastin production, however, the ratio collagen IV/Col I, besides elastin biosynthesis may support, as the final outcome, an improvement of flexible cutaneous basement membrane structure and, therefore, of skin (dermal tissue) quality and texture.

Biological data obtained on dermal fibroblasts at transcriptional and protein levels proved that lightly crosslinked HA-based gels, aiming at superficial treatments, were able to prompt collagen synthesis. Alteration in the extracellular matrix and especially collagen expression are generally associated with skin aging [42]. In this respect, improved biosynthesis of ECM proteins has a beneficial effect on skin/dermal rejuvenation procedures, besides wrinkles filling, leading to tissue hydration and consistency improvement [43]. It has been reported that integrins play a key function in cell adhesion as signaling receptors,
acting as a bridge between ECM and cytoskeleton proteins in order to allow cell movement [44,45]. Activation of the integrin complex (α1β1) by an extracellular ligand (e.g., collagen, laminin and elastin) leads to specific intracellular signaling, involving phosphorylation and dephosphorylation reactions for matrix remodeling [46,47]. Our results allowed us to evaluate a beneficial effect of the analyzed samples on proteins matrix hypothesizing an involvement of the integrins receptor.

Finally, it is here demonstrated that biosynthetic pathways related to extracellular macromolecules are indeed affected by modified HA, even if this aspect was less investigated for these kinds of formulations generally used for their biomechanical effects.

Overall, it is worth underlying that the increasing number of in vitro studies characterizing commercial gels is surely useful to increase clinicians’ awareness in the selection and use of these gels and they are even more valuable in case of newly-developed gels for which clinical data are still lacking. However, attention has to be paid to translate these in vitro data to in vivo performance since additional aspects such as the mechanical properties of the tissue surrounding the implanted gel that can affect both hydration and projection capacity of the gels as well as individual inflammatory reactions and other in vivo mechanisms affecting degradation, and biological effect dependence on products in vivo degradation should be also considered.

4. Materials and Methods

4.1. Materials

Restylane Vital (RV) is a Q-Med AB product (Q-Med AB, Uppsala, Sweden). Juvederm Volite (JV) is distributed by Allergan S.P.A. (Pringy, France). Viscoderm Hydrobooster (HB) is distributed by IBSA Farmaceutici Italia srl (Lodi, Italy). They are HA-based dermal fillers intended for use as skinboosters. They all consist of BDDE-crosslinked HA, suspended in physiological medium. The total HA concentration, as reported in the package inserts, is 20 mg/mL for RV, 12 mg/mL for JV and 25 mg/mL for HB. Bovine testicular hyaluronidase, BTH (EC 3.2.1.35), salt-free lyophilized powder with a specific activity of 890 U/mg was purchased from Sigma-Aldrich S.R.L. (Milan, Italy) (cat. N.H3884, lot. SLBF8562V). Dulbecco’s Phosphate Buffered Saline (PBS) without calcium and magnesium was purchased from Lonza Sales Ltd., Switzerland (cat. N.BE17-516F, lot. N.3MB191). Hydrogen peroxide, 30% w/w in water was purchased from Sigma Aldrich S.R.L. (Milan, Italy), cat. N.H1009. Copper (II) sulfate (≥99%) Fluka, cat. N.61230, was purchased from Sigma Aldrich S.R.L. (Milan, Italy).

4.2. Biophysical and Chemical Characterization

4.2.1. Soluble Fraction Quantification and Hydrodynamic Characterization

Filler soluble fraction was quantified as reported elsewhere [7,8]. Briefly, each filler was diluted to 4 mg/mL in PBS (1.0 mL final volume). The resulting suspension was kept under stirring (1000 rpm) for 24 h at 37 °C. The sample was then centrifuged at 13,000 × g for 5 min, and the supernatant was removed and filtered on 0.22 µm. Filtered samples were then quantitatively analysed for the HA content by carbazole. Based on the total HA concentration indicated in the package inserts, the amount of water-insoluble HA (mg/mL) in each gel was calculated. Hydrodynamic characterization of gel’s soluble fractions was also performed by using the SizeExclusion Chromatography–Triple Detector Array (SEC-TDA) equipment by Viscotek (Viscotek, Malvern, UK). A detailed description of the SEC-TDA system, of its potential to analyse biopolymers such as HA, and of the analysis conditions are reported elsewhere [40]. Sample’s molecular weight (Mw, Mn, Mw/Mn), molecular size (hydro-
dynamic radius—$R_h$) and intrinsic viscosity ($[\eta]$) distributions were derived. The Mark–Houwink–Sakurada (MHS) curve (log$[\eta]$ vs. log$M_w$) was directly derived for each sample.

4.2.2. $^1$H-NMR Analyses

Gels were investigated for the BDDE/HA content by $^1$H-NMR, as previously described, with slight modifications [25]. Briefly, gels were diluted to 4 mg/mL in HCl 0.01 M and incubated for 72 h at 70 °C under stirring (400 rpm). The samples were neutralized by adding Na$_2$HPO$_4$ and lyophilized. The dried samples were dissolved in D$_2$O and $^1$H-NMR analyses were performed using a Bruker DRX-400 ($^1$H NMR: 400 MHz) instrument at 298 K. Data were processed using the data analysis packages integrated with Bruker TopSpin® 4.0.5 software.

4.2.3. Hydration Capacity

Gels were diluted to 4 mg/mL in PBS and incubated overnight in thermostaker at 37 °C under stirring for 16 h. After centrifugation ($13,000 \times g$, 5 min) and supernatant removal, the pellet was weighed obtaining the hydrated sample mass (g), corresponding to the hydrated sample volume (mL) (density equal to 1 g/mL). The hydration extent of each gel was calculated as:

$$\text{gel's hydration capacity (mL/mL)} = \frac{\text{hydrated sample volume (mL)}}{\text{initial sample volume (mL)}}$$ (1)

Such values represent the volume expansion for each formulation when allowed to reach the equilibrium swelling in PBS.

4.2.4. Rheological Characterization

Rheological measurements were performed as reported elsewhere [7,8] with slight modifications. A Physica MCR301 oscillatory rheometer (Anton Paar, Ostfildern, Germany) equipped with a parallel plate geometry, 25 mm plate diameter ($R_v$, $J_v$) and 50 mm ($H_B$), and Peltier temperature control was used. Measurements were performed at 37 °C. Oscillation frequency sweep tests were carried out over a frequency range from 0.159 to 10 Hz (a range of frequencies considered physiologically relevant for the specific application), at a constant strain selected within the linear viscoelastic range (0.1%). $G'$ and $G''$ were measured and reported as a function of frequency. Complex viscosity values were also registered in the frequency range exploited. Representative curves are reported.

4.2.5. Stability to Degradation

Gels were evaluated for their sensitivity to degradation due to ROS- and BTH-action. Stability of the gels to ROS action was studied using the H$_2$O$_2$/Cu$^{2+}$ system for generating radicals. Experiments were carried out as previously reported, with minor modification [7]. Briefly, aqueous solutions of H$_2$O$_2$ and CuSO$_4$ were added to each gel to have H$_2$O$_2$ 187 mM and CuSO$_4$ 1.87 mM while diluting the gel 1:1.3 ($w/w$). The suspensions were mixed and rapidly placed on the lower plate of the rheometer. A PP25 geometry, (25 mm plate diameter) was used. A time oscillatory test was carried out at 37 °C. Specifically, the storage modulus value of the samples was measured as a function of the time while maintaining constant the frequency (1.59 Hz) and the strain (0.1%). The delay between the addition of the H$_2$O$_2$/Cu$^{2+}$ system to the samples and the acquisition of the first $G'$ value was 1–3 min. For each gel, the same experiment was performed adding water in place of the H$_2$O$_2$/Cu$^{2+}$ system (control). Degradation was monitored by measuring the $G'$ decrease (% in respect to the control) as a function of the incubation time (up to 8′) with the ROS generating system. Experiments were carried out in duplicate.

Sensitivity of the gels to enzymatic degradation was evaluated as previously reported with slight modification [7,8,25]. Specifically, the amount (mg/mL) of HA solubilized due to BTH action was determined as a measure of degradation. Gels were diluted in PBS to 4 mg/mL final concentration and incubated in the presence of BTH 2 U/mL (3 h) and
5 U/mL (3 h and 6 h) at 37 °C under stirring. Testing was interrupted by boiling the sample for 10 min to inactivate the enzyme. Samples were then centrifuged at 13,000 $\times$ g for 5 min. The supernatant was removed, filtered on 0.22 µm, opportunistically diluted in water and then quantitatively analyzed for the HA content by carbazole assay. The amount of soluble HA already existing in the gel (determined as indicated in Section 4.2.1) was subtracted to obtain the amount of HA solubilized due to BTH action. Further, the water-soluble fraction derived from the less and the most drastic enzymatic hydrolysis conditions tested underwent hydrodynamic characterization using the SEC-TDA system (see also Section 4.2.1).

4.2.6. Cohesivity

Product cohesivity was evaluated following the validated protocol reported by Sundaram and collaborators [20]. Specifically, 1 mL of gel was stained with 10 µL of toluidine blue (0.1%) and filled in a 1 mL syringe. The sample was extruded in a 1 L beaker with 700 mL of water mQ, while stirring (160 rpm) with the aid of the magnetic stirrer (2.5 cm). Photos at diverse time intervals and videos were taken. Cohesivity was evaluated independently by 4 raters that assigned, for each sample, at each time, a value of cohesivity (from 1 to 5) referring to the Gavard–Sundaram Cohesivity Scale [20]. Results were reported as the mean score ± SD.

4.3. Biological Evaluation

4.3.1. Cell Cultures

A human dermal fibroblast cell line immortalized with hTERT (HDF cells, BJ-5ta, ATCC CRL-4001) was cultured in a 4:1 mixture of Dulbecco’s Modified Eagle Medium (DMEM) and Medium199 supplemented with 0.01 mg/mL hygromycin B and 10% (v/v) FBS. All materials for HDF culture were purchased from Gibco and Invitrogen. The cells were grown on tissue culture plates using an incubator with a humidified atmosphere (95% air/5% CO$_2$ v/v) at 37 °C.

4.3.2. Cell Viability (MTT Test)

MTT test was accomplished according to La Gatta et al., 2018. Briefly, cells were seeded at a density equal to $2 \times 10^4$ cells/cm$^2$ in 12 wells. Twenty-four hours after seeding, the treatments were added to the medium at 0.16% w/w concentration. After 24 h from gels addition, the cells were observed at inverted optical microscope (MO). Then, cell viability was assessed by measuring the reduction of the tetrazolium dye 3-(4,5-dimethylthiazol-2-yl)-5-(3-carboxymethoxyphenyl)-2-(4-sulfophenyl)-2H–tetrazolium (MTT). Medium was removed and cells were treated three times with PBS to remove any residual suspended hydrogels before adding the MTT solution. Cell viability in presence of fillers was reported with respect to untreated cells (viability %).

4.3.3. Type I Collagen (COLIA1), Type IV Collagen (COLIVA1), Elastin (ELS) and Integrins ($\alpha_1$ and $\beta_1$) mRNA Quantification Using qRT-PCR Analyses

Total RNA was extracted using TRIzol RNA Isolation Reagents (Thermofischer scientific, Waltham, MA USA), and reverse transcribed by Reverse Transcription System Kit (Promega, Milan, Italy). Quantitative real-time polymerase chain reactions (qRT-PCR) were performed in duplicate for all genes of interest using IQ™ SYBR® Green Supermix (Bio-Rad Laboratories, Milan, Italy) and internal control (glyceraldehyde-3-phosphate dehydrogenase, (GAPDH) housekeeping gene). Results are expressed as fold change ($2^{-\Delta\Delta CT}$ method) in treated cells vs. untreated cells (the control), and normalized to transcript levels of housekeeping gene [48]. qRT-PCR was performed using custom primers reported in Table 3.
Table 3. Primers sequence used for qRT-PCR analyses.

| Gene Name (Symbol)                             | PCR Primer Sequence 5'-3'                  | Amplicon Length (bp) |
|------------------------------------------------|--------------------------------------------|----------------------|
| Glyceraldehyde3-phosphate dehydrogenase (GAPDH) | TTCCACGGCAACGTCGAAGGTCATCA                | 115                  |
| Type I collagen (COLIA1)                       | CCAGAAGAACTGTTACTCATCA                    | 103                  |
| Type IV collagen (COLIVA1)                     | GGATCGGCTACTCTTTTGTGATCA                  | 104                  |
| Elastin (ELS)                                  | AGGTGTATACCCAGGCGTGACAA                   | 105                  |
| α-1 integrin (INTα1)                           | TCAGGATCCGGGAAAGTCAT                     | 108                  |
| β-1 integrin (INTβ1)                           | ACTGTGATCGGCGTATATTGACAC                  | 110                  |

4.3.4. Western Blotting for Collagen Type 1, Elastin, and Actin

Western blotting analyses were performed after 48 h of treatment. Cells were lysed in Radio-Immunoprecipitation Assay (RIPA buffer 1×; Cell Signaling Technology) and intracellular protein concentration was determined through the Bradford method. Specifically, 30 µg of proteins for each sample were resolved on a 10% SDS–PAGE gel and transferred onto nitrocellulose membrane (GE, Amersham, UK). Then, the membrane was blocked by 5% non-fat milk in Tris-buffered saline and 0.05% Tween-20 (TTBS) for 30 min and primary antibodies against Elastin (Santa Cruz, Dallas, TX, USA), Col I (Elabscience, Houston, TX, USA) and Col IV (Abcam, Cambridge, UK) were diluted 1:500 and incubated overnight at 4°C. Secondary antibodies horseradish peroxidase-conjugated donkey anti-mouse and goat anti-rabbit were diluted 1:5000 and incubated for 2 h at room temperature. Anti-β-Actin antibody used at 1:1000 dilution was used as the loading control. The signal was detected using the ECL system (Chemicon-Millipore, Milano, Italy) and the semi-quantitative analyses of protein expression were carried out with the ImageJ program.

4.4. Statistical Analysis

Unless otherwise indicated, each experiment was performed at least in triplicate and results are reported as the mean value ± standard deviation. Data were statistically evaluated by performing One-way ANOVA tests followed by post hoc correction for multiple comparison. The level of significance was fixed at 0.05.

5. Conclusions

Biophysical and biochemical parameters were derived for three crosslinked HA-based gels proposed for intradermal injection as skinboosters. Collected data indicated H_B as the gel with the highest concentration in insoluble HA and the longer soluble HA chains and with the highest water uptake, suggesting deeper hydration properties. H_B showed the lowest rigidity/viscosity and the highest cohesivity. J_V was found to be the most resistant to degradation by ROS while longer permanence in the presence of BTH was recorded for H_B. HA-based samples determined a significant improvement of matrix biomarkers expression at the transcriptional level and strong upregulation of integrins was found for J_V and H_B treated fibroblasts. However, Western blotting analyses showed only a slight boosting effect of gels on Human Dermal Fibroblasts in terms of Col IV/Col I and elastin production. Comparison with conventional volumetric formulations revealed differences far lower than expected in relation to composition, hydration capacity and biochemical effects. However, except for R_V, rigidity was decisively lower indicating higher deformability as the most important requirement for more superficial implantation.
These results are thought to be useful to practitioners for improving the use of these gels and valuable to adjust gel design towards even more specific performance.

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