Versatile use of dermal substitutes: A retrospective survey of 127 consecutive cases

Giovanni Nicoletti1,2, Marco Mario Tresoldi1,2, Alberto Malovini3, Marco Visaggio1, Angela Faga4,5, Silvia Scevola2

1Department of Clinical Surgical Diagnostic and Pediatric Sciences, Plastic and Reconstructive Surgery Unit, University of Pavia, 2Advanced Technologies for Regenerative Medicine and Inductive Surgery Research Center, University of Pavia, 3Laboratory of Informatics and Systems Engineering for Clinical Research, Maugeri Clinical Scientific Institutes, 4Plastic and Reconstructive Surgery Unit, Maugeri Clinical Scientific Institutes, 5Department of Molecular Medicine, University of Pavia, Pavia, Italy

Address for correspondence: Dr. Giovanni Nicoletti, Department of Clinical Surgical Diagnostic and Pediatric Sciences, Plastic and Reconstructive Surgery Unit, University of Pavia, Viale Brambilla, 74, 27100 Pavia, Italy. E‑mail: giovanni.nicoletti@unipv.it

ABSTRACT

Background: Dermal substitutes are currently largely used for the treatment of huge skin loss in patients in critical general health conditions, for the treatment of severe burns and to promote the healing process in chronic wounds. Aims: The authors performed a retrospective assessment of their experience with bioengineered skin to possibly identify the most appropriate clinical indication and management for each substitute. Materials and Methods: The study involved 109 patients with 127 skin defects repaired with dermal substitutes over a 9 years period, from 2007 to 2016. Hyalomatrix® was used in 63 defects, whereas Integra® and Nevelia® were used in 56 and 8 defects, respectively. Results: The statistical analysis failed to reveal a correlation between the choice of a specific dermal substitute and any possible clinical variable except in the soft‑tissue defects of the scalp where Hyalomatrix® was electively used. Conclusions: In the authors’ experience, the scalp defects followed a radical excision of skin tumours that included the periosteum. Here, the preliminary cover with a hyaluronan three‑dimensional scaffold constantly allowed for the regeneration of a derma‑like layer with a rich vascular network fit for supporting a split‑thickness skin graft. Nevertheless, the authors still prefer Integra® when the goal is a better cosmetic outcome and Hyalomatrix® when a faster wound healing is required, especially in the management of deep wounds where the priority is a fast obliteration with a newly formed tissue with a rich blood supply. However, these clinical indications still are anecdotally based.

KEY WORDS

Clinical application; collagen; dermal substitutes; hyaluronan; skin reconstruction

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How to cite this article: Nicoletti G, Tresoldi MM, Malovini A, Visaggio M, Faga A, Scevola S. Versatile use of dermal substitutes: A retrospective survey of 127 consecutive cases. Indian J Plast Surg 2018;51:46‑53.
INTRODUCTION

Dermal substitutes are currently largely used for the treatment of huge skin loss in patients in critical general health conditions,[1,2] for the treatment of severe burns[3-5] and to promote the healing process in chronic wounds.[6]

In our clinical practice, the most frequently used dermal substitutes are Integra® (Integra LifeSciences, Plainsboro, NJ, USA), Hyalomatrix® (Anika Therapeutics, Bedford, MA, USA) and more recently, Nevelia® (SYMATESE Biomateriaux, ZI Les Troques, Chaponost, France), as well.

The choice among the different substitutes was originally made according to their reputation for popularity, reliability, availability and cost convenience. Subsequently, as our learning curve progressed to a stage of established clinical confidence, the use of bioengineered skin was extended to several innovative indications and currently, it has replaced the traditional indications for flap surgery for both functional and aesthetic indications in an increasing number of cases, too.[7-9]

In a patients’ cohort of ours, we already performed an observational study to assess the clinical and histological long-term outcomes of Hyalomatrix®[8] used for revision of retracting scars; similarly, we also carried out a long-term objective in vivo instrumental assessment of the skin properties after reconstruction with either Integra® or Hyalomatrix®.[9]

In the present study, we performed a retrospective assessment of the whole of our experience with bioengineered skin and a comprehensive overview of our evidence-based clinical indications. The aim of the survey was an attempt at possibly identify the most appropriate clinical indication and management for each substitute.

MATERIALS AND METHODS

The study was carried at the Plastic Surgery Unit, University of Pavia, Salvatore Maugeri Clinical Scientific Institutes and involved 109 patients, 59 females and 68 males, with an average age of 70 years (range 11–93).

The skin defect was considered the experimental unit of the study irrespective of the number of defects per patient. An overall of 127 skin defects were repaired with bioengineered skin over a period 9 years, from 2007 to 2016.

Analysed data included patients’ sex and age, type of dermal substitute, anatomical sites, wound aetiology, skin loss area and depth, the time interval between the application of the substitute and its coverage with a skin graft, the time interval between the skin graft application and the complete clinical healing of the defect.

The anatomical sites were grouped into five main areas: lower limb including the foot, upper limb including hand and axilla, face and neck, scalp and trunk including the gluteal region.

The area of each soft tissue loss was calculated using the ImageJ software, and its depth was routinely measured intraoperatively with a ruler.

Statistical methods

Categorical variables’ distribution was described by counts and frequencies. Since quantitative variables’ distribution deviated from the normality assumptions (based on the visual inspection of quantile-quantile plots), they were described by median [25th–75th percentiles, interquartile range (IQR)]. The presence of statistically significant differences in terms of categorical variables’ distribution between substitutes’ types was assessed using the Fisher’s exact test; while the Kruskal–Wallis test and Wilcoxon Rank Sum test were employed to test for the presence of statistically significant differences in terms of quantitative variables between substitutes’ types as appropriate. The significance threshold was set to $P < 0.05$. Statistical analyses were performed using the R software version 3.3.0 (www.r-project.org, Free Software Foundation, Boston, MA, USA).

RESULTS

Hyalomatrix® was used in 63 defects, while Integra® and Nevelia® were used in 56 and 8 defects, respectively.

The distribution of the bioengineered skin repair per anatomical site was:
- Lower limb including the foot: 27
- Upper limb including hand and axilla: 20
- Face and neck: 44
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Results showed that the median age of individuals whose skin defects were treated by Nevelia® was significantly higher than that of those whose defects were treated by Integra® (Median = 75.5, IQR = 73.5–82.5 vs. Median = 65.5, IQR = 38.5–77.25, \( P = 0.031 \)).

The skin defects of the scalp were significantly more frequently treated by Hyalomatrix® compared to Integra® (31.75% vs. 12.5%, \( P = 0.016 \)).

No other statistically significant differences were observed.

DISCUSSION

A broad age range was observed in our sample. We tended to use the bioengineered skin for the correction of scar contractures in patients in the stage of development to both save more complex reconstructive options for later stages and to limit the overall burden of surgical scars [Figure 2]. In the cosmetically demanding young patients, we also used the bioengineered skin for the reconstruction of defects of the face as an alternative to traditional skin flaps to minimise the amount of visible scars [Figure 3]. Nevertheless, the vast majority of our sample included elderly patients suffering from advanced skin cancers that are far more frequent in this age group. In our experience, bioengineered skin progressively became the gold standard for reconstruction of skin defects in the aged patients as the comorbidities frequently reported in this patient cohort generally contraindicate complex surgical procedures. Regenerative surgery,

- Scalp: 27
- Trunk including the gluteal region: 9.

The reported aetiologies of the skin defects were as follows: difficult to heal wounds (25), excision of skin tumours (87) and surgical revision of contractures and scars (15).

The characteristics of the analysed samples are summarised in Table 1.

The dermal substitutes demonstrated a trend for progressively increasing frequency of use along the period of study [Figure 1].

The characteristics of the analysed samples by substitutes’ type are depicted in Table 2.

Table 1: Characteristics of the analysed samples

| Variable                  | Level     | Distribution |
|---------------------------|-----------|--------------|
| Sex                       | M         | 68 (53.54%)  |
|                           | F         | 59 (46.46%)  |
| Age                       | Years     | 70 (50-82)   |
| Dermal substitute         | Integra®  | 56 (44.09%)  |
|                           | Hyalomatrix® | 63 (49.61%) |
|                           | Nevelia®  | 8 (6.3%)     |
| Site                      | Lower limb| 27 (21.26%)  |
|                           | Upper limb| 20 (15.75%)  |
|                           | Face and neck | 44 (34.65%) |
|                           | Scalp     | 27 (21.26%)  |
|                           | Trunk     | 9 (7.09%)    |
| Aetiology                 | Difficult-to-heal wounds | 25 (19.69%) |
|                           | Skin tumors | 87 (68.5%)   |
|                           | Retracting scars | 15 (11.81%) |
| Skin graft time after substitute application | Days | 28 (26.5-40) |
| Skin graft engraftment time | Days | 14 (14-21)   |
| Wound depth               | Cm        | 0.5 (0.5-1)  |
| Wound area                | Cm²       | 19.25 (8-41.62) |

Figure 1: Trend for the clinical use of the different dermal substitutes along the period of study

Figure 2: (a) Retracting post-burn scars in the posterior aspect of the left lower limb. (b) Soft tissue loss following scar release in the lower left gluteal area and in the left popliteal fossa. (c) The defects are temporarily repaired with Hyalomatrix® dermal substitute. (d) Stable repair with split thickness skin grafts
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Table 2: Samples' characteristics by dermal substitute type

| Variable                  | Level          | Integra® (56) | Hyalomatrix® (63) | Nevelia® (8) | Global          | Integra® vs. Hyalomatrix® | Integra® vs. Nevelia® | Hyalomatrix® vs. Nevelia® |
|---------------------------|----------------|---------------|-------------------|--------------|-----------------|--------------------------|-----------------------|--------------------------|
| Sex                       | M              | 27 (48.21%)   | 36 (57.14%)       | 5 (62.5%)    | 0.284           | 0.362                    | 0.708                 | 1                        |
|                           | F              | 29 (51.79%)   | 27 (42.86%)       | 3 (37.5%)    |                 |                          |                       |                          |
| Age                       | Years          | 65.5          | 73 (57.5-82.5)    | 75.5         | 0.037*          | 0.066                    | 0.031*                | 0.187                    |
|                           | (38.5-77.25)   |               |                   | (73.5-82.5)  |                 |                          |                       |                          |
| Site                      | Lower limb     | 11 (19.64%)   | 14 (22.22%)       | 2 (25%)      | 0.324           | 0.140                    | 1                     | 1                        |
|                           | Upper limb     | 12 (21.43%)   | 7 (11.11%)        | 1 (12.5%)    |                 |                          |                       |                          |
|                           | Face and neck  | 22 (39.29%)   | 19 (30.16%)       | 3 (37.5%)    |                 |                          |                       |                          |
|                           | Scalp          | 7 (12.5%)     | 20 (31.75%)       | 0 (0%)       | 0.013*          | 0.016*                   | 0.582                 | 0.095                    |
|                           | Trunk          | 4 (7.14%)     | 3 (4.76%)         | 2 (25%)      | 0.116           | 0.705                    | 0.159                 | 0.094                    |
|                           |                |               |                   |             |                 |                          |                       |                          |
| Aetiology                 | Difficult-to-heal wounds | 9 (16.07%) | 15 (23.81%) | 1 (12.5%) | 0.587 |
|                           | Skin tumors    | 41 (73.21%)   | 39 (61.9%)        | 7 (87.5%)   | 0.239           | 0.241                    | 0.667                 | 0.246                    |
|                           | Retracting scars| 6 (10.71%) | 9 (14.29%) | 0 (0%)   | 0.615           | 0.593                    | 1                     | 0.584                    |
| Skin graft time after substitute application | Days          | 29.5 (26-42)  | 28 (26-35)       | 32.5 (31.75-36.25) | 0.315 | 0.259 | 0.776 |
|                           |                |               |                   |             |                 |                          |                       |                          |
| Skin graft engraftment time| Days          | 14 (13-20.25) | 14 (14-21) | 15.5 (13.25-20) | 0.583 | 0.295 | 0.837 |
| Wound depth               | Cm             | 0.5 (0.5-1)   | 0.8 (0.5-1)       | 0.5 (0.5-0.62) | 0.299 | 0.150 | 0.923 |
|                           | Cm²            | 17 (5.38-33.75) | 20.5 (11.12-44.5) | 18.3 (8.75-27) | 0.239 | 0.097 | 0.707 |

Variable: Analyzed variable; Level: Variables’ level; Distribution: Count (%) or median (25th - 75th percentiles) by substitute’s type; Comparisons: P values from the comparison among the substitute’s types or from the pairwise comparison between specific substitute’s types; *: P value<0.05

Therefore, provides low invasive and biologically effective procedures in the so-called fragile patient. Furthermore, not only clinical but also possible economic advantages in choosing dermal substitutes may exist, as surgical costs and surgical time are lower than in traditional reconstructive procedures.[10]

The use of dermal substitutes was sporadic and prudent in the early phase of our experience but as our confidence with both the procedures and the indications improved it was progressively included within the Unit’s routine. Our experience with Nevelia® is limited because it has been in use in our Unit since 1 year only [Figure 1].

A split thickness skin graft transfer was performed after an average time of 28 days after the application of the dermal substitute, and there was no statistically significant difference among the different substitutes. Such an outcome demonstrates similar biological integration times for all of the substitutes under study.

The engraftment time for a skin graft did not show significant differences amongst the different dermal substitutes, thus confirming in all of them the same fitness for supplying an autologous graft.

A very broad wound area range (range: 1.7–1400 cm²) was also observed in our sample and no correlation was demonstrated between the wound size and the choice of any specific dermal substitute. Such a figure confirms the extreme versatility of all of the dermal substitutes under study and their fitness for any wound size.

The statistical analysis demonstrated a correlation between the elective choice of Hyalomatrix® in the soft-tissue defects of the scalp following a radical excision...
of skin tumours that included either the periosteum only or the periosteum plus the outer table of the calvarium.

Although the engraftment of a skin graft can successfully take place on the exposed trabecular bone, its long-term outcome is generally poor due to the high risk of ulceration following minimal trauma. Therefore, the preliminary hyaluronan induced regeneration of a derma-like layer before skin graft transfer on the exposed diploe allowed for a thicker and more reliable bioengineered skin cover [Figure 4].

On the other hand, a skin graft on the bare bone is unlikely if not impossible. Here, the preliminary cover with a hyaluronan three-dimensional scaffold constantly allowed for the regeneration of a derma-like layer with a rich vascular network fit for supporting a split-thickness skin graft. In such a circumstance the revascularisation likely took place both from the periphery and from the fine capillaries lying in the inner and outer tables of the adult cranium thus allowing the integration of the substitute in broad areas of exposed bone, too[13] [Figure 5].

Hyalomatrix® is a dermal substitute made of a non-woven pad of hyaluronic acid benzyl ester layered on top by a semi-permeable membrane of silicone. Hyaluronan provides hydration and maintenance of the extracellular space in the skin. It establishes complex interactions with matrix components and cells and its role ranges from a purely structural function to the regulation of cellular development. Hyaluronan also plays a relevant biological role in the process of wound healing[12] by creating a favourable environment for cell migration. Furthermore, it provides protection against free radical and proteolytic damage to both cells and extracellular matrix molecules, due to its free-radical scavenging, protein-exclusion properties and antioxidant effect.[13-15] In a previous study of ours[9] Hyalomatrix® derived bioengineered skin electively demonstrated to closely approach the hydration and transepidermal water loss of normal skin. Such a figure was supposed to be related to both a hyaluronan induced stimulation and regulation of the sweat gland remnants and a better epidermis–matrix interaction. It also revealed a lively hyaluronan induced neoangiogenesis. Therefore, Hyalomatrix® would be the most appropriate dermal substitute when the water regulation-related skin features and the neoangiogenetic boost are relevant issues.

Due to the limited size of our sample, the statistical analysis failed to reveal any further correlation between the choice of a specific dermal substitute and the remaining clinical variables. Nevertheless, we still prefer Hyalomatrix® when a faster wound healing is required, especially in the management of deep wounds where the priority is a fast obliteration with a new formed tissue provided with a rich blood supply [Figure 6] and Integra® when the goal is a better cosmetic and functional outcome, especially in visible areas of the face as there is no donor site scarring and morbidity [Figure 7]. However, these indications in our clinical practice still are anecdotally based.

Integra® is a matrix of fibres of crosslinked bovine tendon collagen and a glycosaminoglycan (GAG;

**Figure 3:** (a) Congenital melanocytic naevus of the right temple. (b) Temporary repair with Integra® dermal substitute following radical excision. (c) Stable repair following early engraftment of a split thickness skin graft. (d) Long-term outcome of stable split thickness skin graft

**Figure 4:** (a) Soft tissue loss following radical excision of locally infiltrating basal cell carcinoma of the vertex of the scalp. The defect includes the periosteum and a portion of the outer cortex of the skull bone. (b) The defect is temporarily repaired with Hyalomatrix® dermal substitute. (c) Regeneration of a derma-like layer with a rich vascular network fit for supporting a split-thickness skin graft. (d) Stable repair with a split thickness skin graft
Figure 5: (a) Soft tissue loss following radical excision of locally infiltrating squamous cell carcinoma of the vertex of the scalp. The defect includes the periosteum in the lower left area of the excision. (b) Regeneration of a derma-like layer with a rich vascular network fit for supporting a split-thickness skin graft. (c) Long-term outcome after repair with a split-thickness skin graft.

Figure 6: (a) Post-traumatic degloving injury of the Achilles region in the right foot. (b) Soft tissue loss following the wound debridement. (c) The defect is temporarily repaired with Hyalomatrix® dermal substitute. (d) Stable repair with a split-thickness skin graft.

**chondroitin 6-sulphate**) derived from shark cartilage covered on top by a silicone semi-permeable membrane. The bovine collagen and GAG serve as a template for the infiltration of fibroblasts, macrophages, lymphocytes and for the regeneration of a new vascular network. The host fibroblasts provide deposition of newly formed collagen and elastic fibres, which progressively replace the native bovine collagen and GAG three-dimensional scaffold.[16] The silicone layer mimics the natural epidermis providing adequate moisture control. Histological animal and human studies demonstrated good tissue compatibility and integrity, controlled biodegradation and no adverse immunological reactions.[17] In a previous study of ours[9] Integra® derived bioengineered skin electively demonstrated physical, mechanical and optical properties that best approximate to normal skin. Such a figure is likely to be related to its dermal structural organisation, which more closely resembles that of normal skin.

Therefore, Integra® should be the most appropriate solution when the best possible skin color and texture match is required.

Nevelia® was recently introduced in our clinical practice to broaden the spectrum of therapeutical options within the field of bioengineered skin repair.

Nevelia® is a novel three-dimensional porous matrix of stabilised bovine origin type I collagen covered on top by a semi-permeable membrane of silicone. It is made of a specific native collagen with a large fibrous proportion to keep cell adhesion signals and mechanical structure to support regeneration. *In vitro* tests reveal an optimised fibroblast colonisation due to the recognition of collagen fibres and a fast initiation of the revascularisation process.[18]

Although our preliminary results with the Nevelia® induced bioengineered skin would appear to be good, it is too short a clinical experience to make a long-term assessment on a large sample of cases and to identify its most appropriate clinical indications.

The most relevant complication in our sample was the melting graft syndrome that occurred in three cases of Hyalomatrix® skin reconstruction, two in the scalp and one in the gluteal area. Such a figure was related to the poor patients' compliance in the scalp reconstruction and to the objectively difficult management of the gluteal area. One scalp skin reconstruction had to undergo a second skin graft transfer while the other two cases eventually healed with spontaneous re-epithelisation from the wound margins after a course of advanced dressing. Indeed, in potentially contaminated devascularised wounds, and in large areas of bare bone autologous tissue transfer still is an appropriate indication as the bioengineered skin is particularly prone to infection due to the lack of skin adnexa.

In six cases, the bioengineered skin reconstruction could not be completed by a skin graft transfer because severe comorbidities contraindicated any further
Figure 7: (a) Basal cell carcinoma of the dorsum of the nose. (b) Temporary repair with Integra® dermal substitute following tumour radical excision. (c) Stable repair with a split thickness skin graft

surgical procedure. These cases eventually healed with re-epithelisation from the wound margins after a course of advanced dressing in a time ranging from 10 to 18 weeks.

CONCLUSIONS

In our experience, the dermal substitutes were essential and highly versatile reconstructive options providing effective solutions in a wide number of clinical problems with a significant reduction of the surgical complexity.

In our opinion, a hyaluronan-based dermal substitute is the most appropriate choice for coverage of exposed bone without periosteum and for deep soft tissue loss when a fast obliteration of dead space is the primary goal.

Similarly, a dermal substitute with a structured three-dimensional connective fibre organisation is the most appropriate indication when skin color and texture match is a relevant issue in specific clinical cases. Our both experimental[9] and clinical results would suggest a relevant role for the dermal substitute in the modern reconstructive ladder.[19]

Nevertheless, we still need the dermal substitute of the future designed to integrate the demonstrated properties and advantages of both hyaluronan-based and collagen and GAG based scaffold to get cell, and extracellular matrix regulation joined to stimulation of the regenerative process with the establishment of a natural dermal fibres organisation.

Financial support and sponsorship
Nil.

Conflicts of interest
There are no conflicts of interest.

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