The efficacy and safety of Grafix® for the treatment of chronic diabetic foot ulcers: results of a multi-centre, controlled, randomised, blinded, clinical trial

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Abstract
In a randomised, controlled study, we compared the efficacy of Grafix®, a human viable wound matrix (hVWM) (N = 50), to standard wound care (n = 47) to heal diabetic foot ulcers (DFUs). The primary endpoint was the proportion of patients with complete wound closure by 12 weeks. Secondary endpoints included the time to wound closure, adverse events and wound closure in the crossover phase. The proportion of patients who achieved complete wound closure was significantly higher in patients who received Grafix (62%) compared with controls (21%, P = 0.0001). The median time to healing was 42 days in Grafix patients compared with 69.5 days in controls (P = 0.019). There were fewer Grafix patients with adverse events (44% versus 66%, P = 0.031) and fewer Grafix patients with wound-related infections (18% versus 36.2%, P = 0.044). Among the study subjects that healed, ulcers remained closed in 82.1% of patients (23 of 28 patients) in the Grafix group versus 70% (7 of 10 patients) in the control group (P = 0.419). Treatment with Grafix significantly improved DFU healing compared with standard wound therapy. Importantly, Grafix also reduced DFU-related complications. The results of this well-controlled study showed that Grafix is a safe and more effective therapy for treating DFUs than standard wound therapy.

Introduction
There is a worldwide epidemic of diabetes. According to data from the World Health Organization, the global prevalence of diabetes among adults was 6.4% in 2010, affecting 285 million people worldwide. The prevalence of diabetes is expected to

Key Messages
• diabetic foot ulcers are a worldwide epidemic, leading to significant morbidity and rising health care costs
• the aim of the study was to evaluate the efficacy and safety of a cryopreserved placental membrane, Grafix
(N = 50), compared with standard wound care (N = 47) to heal chronic diabetic foot ulcers.

- this multi-centre, randomised clinical study showed a significantly higher wound healing rate, faster healing and fewer wound-related infections among patients who received cryopreserved placental membrane compared with standard wound care, indicating that Grafix is a safe and more effective therapy for treating DFUs than standard wound therapy.

Methods

The study was a prospective, multi-centre, randomised, single-blinded clinical trial to evaluate the safety and efficacy of Grafix for the treatment of chronic DFUs. The primary hypothesis was that Grafix was superior to standard wound care for the primary endpoint of complete wound closure. Patients were enrolled from May 2012 through April of 2013. Key inclusion criteria included confirmed type I or type II diabetes, patients age between 18 and 80 years, index wound present between 4 and 52 weeks, wound located below the malleoli on plantar or dorsal surface of the foot and ulcer between 1 and 15 cm². Main exclusion criteria included haemoglobin A1c above 12%, evidence of active infection including osteomyelitis or cellulitis, inadequate circulation to the affected foot defined by an ankle brachial index <0.70 or >1.30, or toe brachial index ≤0.50 or Doppler study with inadequate arterial pulsation, exposed muscle, tendon, bone or joint capsule and reduction of wound area by ≥30% during the screening period.

Following a 1-week screening period, patients were randomised to the Grafix or control treatment arm in a 1:1 ratio. Patients randomised to Grafix treatment received an application of Grafix once a week (±3 days) for up to 84 days (blinded treatment phase). Patients in the control group received standard wound therapy once a week (±3 days) for up to 84 days. All wounds were appropriately cleaned and surgically debrided to remove all non-viable soft tissue from the wound by scalpel, tissue nippers and/or curettes at each weekly visit. For patients randomised to the Grafix group, the hVWM was placed to come in full contact with the wound and edges. Wounds in both groups received standard wound care that included surgical debridement, off-loading and non-adherent dressings. All patients received a non-adherent dressing: Adaptic® (Systagenix, Gatwick, UK) and either saline moistened gauze or Allevyn® (Smith & Nephew, London, UK) for moderately draining wounds. An outer dressing was then applied. Patients were provided walking boots for wounds on the sole of the foot or a post-op shoe if the wound was on the dorsum of the foot or at the ankle. Custom off-loading boots could be prescribed at the discretion of the site investigator. In addition, the off-loading device used could be changed as needed to accommodate changes in wound size or position.

Patients were evaluated weekly at the clinical site. Patients who achieved complete wound closure then continued to be evaluated during the follow-up phase, twice during the first month and then monthly for two additional visits. Control patients whose wounds were not closed by the end of the blinded treatment phase were able to receive Grafix in the open-label treatment phase, in which Grafix was applied weekly for up to 84 days. Outcome and safety assessments occurred at each visit during the blinded treatment phase, follow-up visits, as well as during the open-label treatment phase.

The primary endpoint of the study was evaluation of complete wound closure of the index wound. Complete wound closure was defined as 100% re-epithelialisation with no wound drainage as determined by the site investigator. Confirmation of wound closure was confirmed at an initial follow-up visit 2 weeks later. Wound closure was independently confirmed via a central wound core laboratory with two blinded wound care...
Table 1 Patient demographics and baseline characteristics

|                      | Grafix (n=50) | Control (n=47) | P-value | 95% confidence interval |
|----------------------|--------------|----------------|---------|-------------------------|
| Mean age, in years (SD) | 55.5 (11.5)  | 55.1 (12.0)    | 0.849   | –5.2 – 4.3               |
| Age ≥65 years (N, %)   | 11 (22%)     | 13 (27.7%)     | 0.521   | 0.292 – 1.861            |
| Male (N, %)           | 33 (66%)     | 35 (74.5%)     | 0.365   | 0.276 – 1.603            |
| Mean years DM (SD)    | 15.4 (11-1)  | 14.0 (11-0)    | 0.549   | –5.80 – 3.10             |
| Mean BMI (SD)         | 33.5 (7.7)   | 32.2 (7.9)     | 0.419   | 4.40 – 1.90              |
| BMI ≥30 (N, %)        | 36 (72%)     | 25 (53.2%)     | 0.057   | 0.975 – 5.253            |
| Race (N, %)           |              |                |         |                         |
| White or Caucasian    | 35 (70%)     | 32 (68-1%)     | 0.581   | –1.847 – 2.073           |
| Black or African American | 13 (26%)   | 12 (25.5%)     | 0.521   | –1.866 – 2.054           |
| American Indian or Alaska Native | 1 (2%) | 1 (2.1%)  | 0.482   | –1.932 – 1.988           |
| Other                 | 1 (2%)       | 2 (4.3%)       | 0.263   | –1.947 – 1.973           |
| Mean wound size at baseline (cm², SD) | 3.41 (3.23) | 3.93 (3.22)   | 0.433   | –0.80 – 1.80              |
| Wound duration (days, SD) | 115.0 (72.6) | 122.9 (83.9)  | 0.621   | –23.7 – 39.5             |
| Glycated haemoglobin >9% (N, %) | 14 (28%) | 13 (27.7%) | 0.970   | 0.418 – 2.473            |
| Mean albumin (g/dl) (SD) | 4.0 (0.4)  | 4.0 (0.3)      | 0.418   | 0.20 – 0.10              |
| Albumin >3.5 g/dl (N, %) | 44 (88%)  | 42 (89.4%)     | 0.263   | –1.947 – 1.973           |
| Ankle bachiial index (ABI) |              |                |         |                         |
| ABI 0.07–0.90 (N, %)  | 10 (21-7%)   | 10 (22.2%)     | 0.44    | –1.89 – 2.03             |
| ABI >0.90            | 36 (78-3%)   | 35 (77.8%)     | 0.39    | –1.89 – 2.00             |

DM, diabetes mellitus.

Efficacy and safety of Grafix®

Sample size and statistical analysis

The study sample size was based on an assumed closure rate of 30% in the control arm and 50% in the Grafix group with a 30% drop-out rate. Under these assumptions, 94 patients, who completed the treatment, in each treatment arm were required to meet the two-sided type 1 error rate of 0.05 with 80% power. A pre-specified interim analysis was planned at 50% enrollment. The interim analysis used a one-sided superiority design based on an Emerson–Fleming symmetric group sequential design using an O’Brien-Fleming boundary shape [Emerson and Fleming (13)]. The analysis was performed by an unblinded statistician and reported to the blinded review committee. Following the interim analysis, the blinded review committee recommended to terminate study enrollment because of overwhelming superiority of the Grafix arm versus the control arm.

The statistical analyses were performed using SAS version 9.2 on an intent-to-treat basis. Baseline demographic and clinical variables were summarised for each treatment arm of the study. Descriptive summaries of the distribution of continuous variables included the mean, standard deviation, median and subject counts; categorical variables were summarised in terms of frequencies and percentages. Treatment group summaries were constructed across all study sites. Statistical comparisons between treatment groups were performed using \( \chi^2 \) testing for categorical variables and analysis of variance (ANOVA) techniques for continuous measures. A Cox proportional hazard regression analysis was performed on time-to-event (wound closure) data.

Results

Patient demographics and baseline characteristics are presented in Table 1. During screening, 139 patients were evaluated. There were 42 patients who failed screening, of which 6 were disqualified after the 1 week run-in period because there was a 30% or greater wound area reduction. Ninety-seven patients were subsequently randomised: 50 received Grafix and 47 received standard wound therapy. Among the 97 patients evaluated, there were 85 plantar foot wounds and 12 dorsal foot wounds. There were eight dorsal wounds in the Grafix treatment group and four dorsal wounds in the control arm of the study. There were no significant differences in baseline characteristics among the two treatment groups. The planned interim analysis showed overwhelming efficacy among patients who received Grafix for the primary and secondary endpoints when compared with the control group (Table 2). Following the interim analysis, the blinded review committee recommended to terminate study enrollment because of overwhelming superiority.

Efficacy evaluation

Blinded treatment phase

The proportion of patients who achieved complete wound closure was significantly higher in patients who received
The probability of closure for the Grafix group was 67% during the 12-week evaluation period for Grafix (Figure 1). A statistically greater probability of complete wound healing was achieved in both groups. The Kaplan–Meier analysis illustrated that 31 of 50 patients (62% vs 10 of 47, P = 0.001) in the Grafix group compared with controls (10 of 47, 21.3%; P = 0.0001). The odds ratio for complete healing for a Grafix patient compared with control was 6.037 (95% CI 2.449–14.882). The Grafix group had significantly faster median time to complete wound closure compared with controls (42.0 days, P = 0.019) indicating superior odds of closure with Grafix relative to standard wound therapy.

Safety evaluation

Overall, fewer Grafix patients experienced at least one adverse event compared with control patients (44.0% vs 66.0%, P = 0.031). Among the patients randomised to Grafix, there were significantly fewer patients with wound-related infections (Grafix, 9 of 50, 18.0%; versus control, 17 of 47, 36.2%; P = 0.044) and fewer hospitalisations related to infections in the Grafix group than control (6% versus 15%, P = 0.15).

Discussion

The results of this study demonstrate that weekly application of Grafix led to improved healing rates of DFUs and

Table 2 Wound healing and safety clinical outcomes

| Wound healing | Grafix (n = 50) | Control (n = 47) | P-value |
|---------------|----------------|-----------------|---------|
| Healed wounds, (N, %) | 31 (62%) | 10 (21%) | <0.001 |
| Median time to wound closure (days) | 42.0 | 69.5 | 0.019 |
| 50% wound area reduction at day 28 (N, %) | 31 (62%) | 19 (40.4%) | 0.035 |
| Median study visits (single blind phase) | 6 | 12 | <0.001 |

Adverse events

| Events | Grafix (N, %) | Control (N, %) | P-value |
|--------|--------------|----------------|---------|
| Subjects experiencing at least one adverse event* (N, %) | 22 (44%) | 31 (66%) | 0.031 |
| Subjects with an infection (N, %) | 13 (26%) | 21 (44.7%) | 0.055 |
| Subjects with a skin or subcutaneous tissue disorder (N, %) | 7 (14%) | 8 (17%) | NS |
| Subjects with injury, poisoning and procedural complications (N, %) | 5 (10%) | 7 (14.9%) | NS |
| Subjects with general disorders (N, %) | 4 (8%) | 3 (6.4%) | NS |
| Subjects with musculoskeletal and connective tissue disorders (N, %) | 4 (8%) | 2 (4.3%) | NS |
| Subjects with a wound-related infection (N, %) | 9 (18%) | 17 (36.2%) | 0.044 |
| Subjects with a serious adverse event due to wound infection (N, %) | 4 (8%) | 10 (21.3%) | 0.084 |
| Subjects having an amputation due to an adverse event (N, %) | 0 (0%) | 1 (2.1%) | NS |

NS, non-significant.

*Included overall number of subjects experiencing at least one adverse event and those with at least 5% adverse events.

Wound recurrence of DFUs closed during the initial 12-week study period was assessed. Follow-up every 4 weeks for an additional 12 weeks showed that ulcers remained closed in 82.1% of patients (23 of 28 patients) in the Grafix group versus 70.0% (7 of 10 patients) in the control group (P = 0.42).

Open-label phase

Patients in the control arm who failed to heal during the initial 12-week treatment period could cross over to receive up to 12 weeks of Grafix therapy (n = 26). After receiving treatment with Grafix, the probability of closure was 67.8% with a mean time to closure for these patients of 42 days.

Regression analysis

Cox proportional hazard regression analysis was performed with treatment group, duration of ulcer, baseline ulcer area, glucose control (glycated haemoglobin), ulcer location and BMI as covariates. Following adjustment for these variables, Grafix was found to have a significant effect on time to closure (P < 0.0001). The hazard ratio was 4.77 (95% CI 2.279, 9.971), indicating superior odds of closure with Grafix relative to standard wound therapy.

Figure 1 Kaplan–Meier analysis of probability of 100% closure for Grafix versus control.

Grafix (31 of 50, 62.0%) compared with controls (10 of 47, 21.3%, P = 0.0001). The odds ratio for complete healing for a Grafix patient compared with control was 6.037 (95% CI 2.449–14.882). The Grafix group had significantly faster median time to complete wound closure compared with controls (42.0 days, P = 0.019), among the wounds that closed in both groups. The Kaplan–Meier analysis illustrated a statistically greater probability of complete wound healing during the 12-week evaluation period for Grafix (Figure 1). The probability of closure for the Grafix group was 67.1% compared with 27.1% for the standard care group (Log-Rank, P < 0.0001). Grafix patients also required fewer study visits (i.e. applications) to achieve closure compared with patients in the control arm (6 versus 12, P < 0.001). Comparison of patients with at least a 50% reduction in wound size by day 28 showed that 31 of 50 patients (62.0%) in the Grafix group achieved this reduction versus 19 of 47 (40.4%) in the control group (P = 0.035). There were 8 (16%) patients who withdrew from the study prior to completion in the Grafix group versus 11 (23.4%) patients who withdrew from the control group.
reduced diabetic foot complications compared with standard wound therapy. In this study, all primary and secondary endpoints showed clinical benefit of Grafix, in the only multi-centre DFU trial to date to meet statistically significant pre-specified interim analyses. This is also the first report of a multi-centre randomised controlled trial (RCT) to investigate the use of human amniotic membrane for the treatment of DFUs. In addition, to the authors’ knowledge, this is the first large, multi-centre RCT for advanced skin substitutes in which the primary endpoint, 100% re-epithelialisation, was confirmed by third-party blinded wound care experts, further removing potential bias and increasing reliability of the endpoint results.

Despite the introduction of numerous advanced wound care products including bioengineered skin substitutes for the treatment of DFUs over the last 15 years, only a few have demonstrated significant clinical efficacy compared with control in multi-centre RCTs (9,10,14), which are considered the best level 1 evidence in trial conduct. Of these studies to date, this study has shown the best relative effect (191% compared with standard of care), among these multi-centre RCTs. In addition, the primary outcome for this trial, which showed a healing rate of 62% at 12 weeks, compares favourably to the RCTs of a bilayered, cell-based product (56%) and human fibroblast-derived dermal substitute (30%) (10). The healing rate in the control arm of this study was 21-3%, which is similar to the average rate of DFU healing in phase 3 randomised clinical trials with a 12-week study duration (24%) (8). Grafix is also available in multiple sizes to allow the product to decrease as the wound size decreases, thereby reducing waste and cost. The healing rates of 62–68% in this trial are consistent with previous published reports of healing of 85% and 68% by 12 weeks among DFU and venous leg ulcer patients, respectively, in an open-label, retrospective study using Grafix (15).

The standard of care treatments selected for this study included surgical wound debridement, high-quality off-loading devices and non-adherent dressings that were provided uniformly to all patients in both treatment groups making the differences in the treatment effect from these interventions alone unlikely. Off-loading is one of the most important elements in DFU treatment (16,17). Several studies have reported significant differences in pressure reduction (18) and wound healing based on the off-loading strategy (17,19–21). In this study, better quality and more effective off-loading devices were provided for all patients than what is commonly provided to patients with DFUs (22). For wounds on the sole of the foot, removable cast walkers were provided; for wounds on the dorsum of the foot and ankle, a post-op shoe was used. Several studies have shown that a higher proportion of wounds heal with removable cast walkers (16) compared with healing sandals or shoes (17,23); however, in community practice only about 15% of patients with DFUs receive this quality of pressure reduction therapy (22). The majority of patients receive less effective off-loading with healing sandals, post-op shoes or therapeutic shoes and insoles as was done in previous randomised trials (21,24). Debridement is another important element in wound therapy. In this study, surgical debridement was done for every patient at every study visit. Other studies have reported that a minority of DFUs received surgical debridement in phase 3 DFU trials (25,26), and that the frequency of wound debridement was associated with differences in wound healing (25) (Table 3). Despite the high-quality standard of care that patients in both groups received, there were overwhelming differences in outcomes between the Grafix-treated group and the control group. In addition, the probability of closure was 67.8% among patients who crossed over from the control group to Grafix after failing to heal. These outcomes, therefore, can only be explained by the effect that Grafix had on these chronic wounds as the significant variable factor between groups. Grafix would be an ideal product to combine with more rigorous off-loading strategies such as total contact casts because it is applied once a week and does not require additional dressing changes (17,19–21). Furthermore, off-loading in a cast would guarantee compliance with off-loading.

There was a significant reduction in wound-related adverse events in patients treated with Grafix. The lower incidence of wound-related infections was likely related to the faster rate of healing and the higher proportion of wounds that healed. The longer the duration of the wound, the greater is the risk of developing a soft tissue or bone infection (27,28). Use of Grafix reduced the median time to wound healing by 4 weeks compared with standard DFU care. Reducing the risk of infection is a key factor in reducing amputations and the cost of DFUs. DFU patients who develop infection are about 56 times more likely to require hospitalisation and 155 times more likely to require amputation (28). Once a patient has an amputation of the foot or leg, the risk of repeated ulcers, infections and amputations increases dramatically (29), as do the associated health care costs.

Placental membranes, first reported as a treatment for wounds in 1910, have a combination of growth factors, collagen-rich extracellular matrix and viable cells including MSCs, neonatal fibroblasts and epithelial cells (12,30–32). However, preservation of cellular viability has previously

### Table 3 Comparison of standard of care among multi-centre, controlled wound care trials

|                         | Grafix* | Dermagraft* | Apligraf* |
|-------------------------|---------|-------------|----------|
| Mean wound size (cm²)   | 3.7     | 2.3         | 3.0      |
| Healed % treatment versus control | 62% versus 21%* | 30% versus 18%* | 56% versus 38%* |
| Time to closure          | 42 versus 70 days* | Not stated | 65 versus 90 days* |
| All adverse events       | 44% versus 66%* | 67% versus 73% | Not stated |
| Wound-related infection  | 18% versus 36-2%* | 19% versus 32%* | 22% versus 32% |
| Off-loading              | Walking boot or Post-op shoe | Therapeutic shoes and custom insoles or healing sandals ad lib | Custom sandal ad lib |
| Debridement              | Every visit | Not stated | Not stated |
|                          |          |             |          |
| *P < 0.05.               |          |             |          |

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limited widespread use (12,15). Grafix, which maintains the native structure of the human tissue, contains living components, which in vitro studies and wound repair models have demonstrated advantages in angiogenic, anti-inflammatory and anti-oxidant effects over dehydrated amniotic membrane (dHAM) products, further support the use of this cellular wound matrix. Arnold et al. reported a 7.5-fold increase in the angiogenic growth factor vascular endothelial growth factor (VEGF) compared to dHAM, important for blood vessel formation. Additional studies performed by Arnold et al. identified reductions in pro-inflammatory cytokines and enhanced anti-oxidant activity as measured by a reduction in oxidant-induced apoptosis of human dermal fibroblasts with cellular wound matrices versus dHAMS (33–35).

Despite advances in advanced wound therapy, chronic DFUs continue to increase in frequency causing significant associated morbidity and rising health care costs. The results of this randomised clinical study demonstrated that weekly application of Grafix increases the proportion of DFUs that heal, accelerates the time to heal, decreases the number of treatments and reduces infections and infection-related hospitalisations when compared with high-quality standard wound care. This is the first clinical product with viable stem cells studied in a randomised clinical trial to successfully show statisitically greater closure rates of chronic DFUs. Based on these findings, Grafix is an important treatment option for health care providers and their patients for the safe and effective treatment of chronic DFUs.

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**APPENDIX**

**The Grafix Diabetic Foot Ulcer Study Group**

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