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Skin graft donor

Short-term outcomes following amniotic membrane and conventional dressing in skin graft donor site; a randomized clinical trial

Houshang Soleimani, Jafar Kazemzadeh* (http://orcid.org/0000-0001-7766-9544)

Department of General Surgery, School of Medicine, Urmia University of Medical Sciences, Urmia, Iran

*Corresponding author: Jafar Kazemzadeh, Email: jafarkazemzade48@gmail.com, kazemzadeh.j@umsu.ac.ir

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Key point

In a randomized clinical trial study on 33 patients in the burn ward, we found the use of amniotic membrane for dressing second-degree burn wounds had better results and benefits such as less pain, faster epithelialization, shorter length of stay at hospital and better patient acceptance in comparison with routine dressings.

Abstract

Introduction: Burn injuries have been associated with a bad prognosis throughout history. Nowadays, the treatment of burn wounds is one of the problems of the country's health care system. There are several treatments for localized burns, such as antimicrobial dressings (using topical antibiotic drugs) and biological dressings (using tissue from living organisms). Biologic dressing with an amniotic membrane is one of the treatments for burned tissues in these patients.

Objectives: This study aimed to investigate the short-term outcomes of dressing with amniotic membrane and routine dressing in graft donor site.

Patients and Methods: This clinical trial study was conducted in the burn ward of Imam Khomeini hospital, Urmia, Iran in 2019. The patients were treated with two amniotic membrane dressings (the patient's left leg) and a local antimicrobial dressing (the patient's right leg). The amniotic membrane was purchased from Sina Cell Company and used to dress the second-degree wound. Patients' right foot was covered with gauze soaked in Nitrofurazone dressing since 2 micrograms of ointment was used for each percent of graft site. The recovery and epithelialization time was assessed one month after the study.
Results: The rate of infection in the two groups did not differ significantly. Furthermore, the rate of wound healing (epithelialization) and post-dressing pain was much better in the group of dressings with amniotic membrane than the group of routine dressings, which was significant (P < 0.01) and showed that amniotic membrane dressings are much more effective in healing wounds and reducing pain compared to the routine dressings.

Conclusion: This study showed that the use of amniotic membrane for dressing second-degree burn wounds, compared to the routine dressings, has better results and benefits such as less pain, faster epithelialization, shorter length of stay at hospital and better patient acceptance.

Trial Registration: Registration of trial protocol has been approved in Iranian registry of clinical trials (RCT20181216041996N1, https://en.irct.ir/trial/35985, ethical code; IR.UMSU.REC.1397.245).

Keywords: Amniotic membrane, burns, wound healing, antimicrobial dressing

Introduction
Burn is a complication that destroys the skin and subcutaneous tissues in different ways with different intensity and extent. Every year, many people suffer from burns requiring hospitalization, and this causes a lot of medical expenses and the resulting complications can affect patients for several years (1). According to annual estimates, there are 700,000 emergency room visits in the United States due to burn injuries (2, 3).

Burn treatment involves several steps; in the acute stage, the main goal is to rehabilitate patients and prevent side effects (kidney failure, acid and base disorders and infectious complications) and in the next stage, to eliminate their physical, mental and rehabilitation complications (4). There are various methods for topical treatment of second and third degree burns, including antimicrobial dressings (administration of topical antibiotics) and biological dressings (use of living tissue) (5-7).

Usually after two weeks of hospitalization and necessary local treatments in patients with grade II burns, re-epithelialization occurs and wound secretions are reduced or completely eliminated (8). Current treatments for burn wounds can be divided into three stages; evaluation, treatment, and rehabilitation (9). Wound dressing is very important and depends on the characteristics of the wound. Grade I wounds with minimal tissue damage do not require dressing. Grade II burns can be treated with daily dressings, topical antibiotics, gauze, and elastic bands. Grade III or deep grade II burns require excision and grafting, and the choice of initial dressing is based on inhibiting bacterial growth and providing a closed dressing until surgery (10, 9).

The amniotic membrane was used in 1910 to cover wounds (11). This method was used in 1974 to treat grade III burns and had good results (12). In other studies, the use of amniotic membranes in the treatment of burns reduced pain and prevented severe water and electrolyte disturbances (13). This method also prepares the wound bed for transplantation (14-18). Studies show that the use of biological dressing with amniotic membrane increases tissue epithelialization (16) and also prevents local infection (19).

Objectives
Due to the use of antimicrobial method in Imam Khomeini hospital in Urmia and the high rate of complications due to burns and high mortality and mortality, the present study aimed to compare the short-term consequences of amniotic membrane dressing and routine dressing at
Patients and Methods

Design and settings
This study was a clinical trial which carried out on 33 patients in the burn ward of Imam Khomeini hospital, Urmia, Iran in 2019. The patients underwent amniotic membrane dressing (patient's left foot) and local antimicrobial dressing (patient's right foot) (Figure 1). Patients with underlying diseases (diabetes, kidney failure, hepatitis, immunodeficiency and cardiovascular diseases) and concomitant trauma, as well as patients who died during treatment or did not consent to biological dressing, and those who lost follow-up, were excluded from the study.

Intervention
The amniotic membrane was purchased from "Sina Cell" and used as a second-degree wound dressing at the donor site. The amniotic membrane was prepared sterile from a pair of pregnant women who delivered by cesarean section and had no history of infectious diseases and placed in buckets containing gentamicin and normal saline (80 mg/l). Placental blood samples were sent to the laboratory to check HBsAg, HIV-Ab and HCV Ab tests. After separation from the chorion and purification, the amniotic membrane was placed in normal saline solution containing gentamicin (80 mg/l) and stored at 4°C. Vaseline gas and then wet gas were placed on the amniotic membrane and bandaging was conducted. Evaluations were performed on the first, seventh and fourteenth days after dressing the area and also at discharge. Possible complications and unpredictable changes were accordingly included in these questionnaires. Patients’ right foot was gauze impregnated with nitrofurazone and 2 μg of ointment was applied per graft. Local infection was based on clinical criteria and in case of symptoms of infection such as wound discharge, symptoms of cellulite and fever, it was prepared from the culture site.

Data analysis
For descriptive statistics, quantitative variables, central indices and dispersion (mean and standard deviation) were calculated, and for qualitative variables, frequency and percentage were calculated. T-test, chi-square, and Fisher’s exact test were applied. A P-value less than of 0.05 was considered significant. All statistical analysis were conducted by SPSS version 18.
Figure 1. CONSORT (consolidated standard of reporting trial) chart for study

Assessed for eligibility (n=41)
- Excluded (n=8)
  - Not meeting inclusion criteria (n=5)
  - Declined to participate (n=3)
- Randomized (n=33) (66 feet)
- Lost to follow-up (n=0)

Allocation

Enrollment
- Control group receiving routine dressing (n=33 left feet)
- Intervention group receiving amniotic membrane dressing (n=33 right feet)

Follow-up
- Lost to follow-up (n=0)

Analysis
- Analysis (n=33)
Results

A total of 33 patients, 23 (69.8%) males and 10 (30.3%) females with a mean age of 26.82±3.37 years were included in the study (33 right legs with amniotic membrane as intervention group and 33 left legs as control group). On the seventh day in the intervention group, one case (3%) of wound infection (positive culture) was observed. In the control group, wound infection was reported in three patients (9.1%). According to the results of Fisher’s exact test, no significant difference between the symptoms of wound infection on the seventh day among groups was detected (P = 0.30). On the fourteenth day, no case of infection was reported in routine dressing and amniotic membrane dressing.

Pain severity in patients reported with Visual Analogue Scale (VAS). In the amniotic membrane group, the mean VAS on the first day was 3.78±1.45 and in the routine dressing group was 4.03±1.53. This difference between the two groups regarding pain on the first day according to T-test was not significant (P=0.48; Table 1). The mean VAS was 1.78±0.97 on the seventh day and in the routine dressing group was 2.70±1.41, therefore according to the T-test, it was significant (P=0.04; Table 2). The severity of pain in the amniotic membrane group was 1.50±0.54 on the fourteenth day and in the routine dressing group was 2.72±1.10 (P=0.02; Table 3).

In the right leg, when the amniotic fluid was employed for dressing, the need for analgesia was much less due to less pain. The mean recovery time in the amniotic membrane dressing group was 2.94±1.18 days and in the routine dressing group was 4.14±2.04 day. This difference
between the two was significant (P=0.004; Table 4). On the seventh day, three patients in the study group (9.1%) and seven patients (21.2%) in the control group had pruritus. This difference was not statistically significant according to Fisher’s exact test (P = 0.15). On the fourteenth day, a significant difference in the frequency of pruritus was observed between the two groups, thereby in the intervention group, one patient (3%) and in the control group, six patients (18.2%) had itching. This difference was statistically significant according to chi-square test (P = 0.04; Table 5). The frequency of burn wound healing or epithelialization on the seventh day after dressing was 81.8% in the amniotic group and 48.5% in the routine dressing group. According to the statistical test with Fisher’s exact test, this difference was statistically significant (P=0.004). The frequency of burn wound healing or epithelialization on the fourteenth day after dressing in the amniotic group was 100% and in the routine dressing group was 78.8% (P = 0.005; Table 6).

| Table 1. Evaluation of pain score with VAS on the first day in the two groups |
|------------------------|---------|-----------------|-------|
| VAS score             | Mean    | Standard deviation | P-value |
| Amniotic group        | 3.78    | 1.45             | 0.48   |
| Routine group         | 4.03    | 1.53             |        |

| Table 2. Evaluation of pain score with VAS on the seventh day in the two groups |
|------------------------|---------|-----------------|-------|
| VAS score             | Mean    | Standard deviation | P-value |
| Amniotic group        | 1.78    | 0.97             | 0.04   |
| Routine group         | 2.70    | 1.41             |        |

| Table 3. Evaluation of pain score with VAS on the fourteenth day in the two groups |
|------------------------|---------|-----------------|-------|
| VAS score             | Mean    | Standard deviation | P-value |
| Amniotic group        | 1.50    | 0.54             | 0.02   |
| Routine group         | 2.72    | 1.10             |        |
Table 4. Comparison of mean recovery time in the two groups

<table>
<thead>
<tr>
<th>Recovery time (day)</th>
<th>Mean</th>
<th>Standard deviation</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amniotic group</td>
<td>2.94</td>
<td>1.18</td>
<td>0.004</td>
</tr>
<tr>
<td>Routine group</td>
<td>4.14</td>
<td>2.04</td>
<td></td>
</tr>
</tbody>
</table>

Table 5. Frequency of pruritus in amniotic membrane and routine dressing at the graft site

<table>
<thead>
<tr>
<th>Recovery time (day)</th>
<th>Amniotic group</th>
<th>Routine group</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Seventh day</td>
<td>3 (9.1%)</td>
<td>7 (21.2%)</td>
<td>0.15</td>
</tr>
<tr>
<td>Fourteenth day</td>
<td>1 (3%)</td>
<td>6 (18.2%)</td>
<td>0.04</td>
</tr>
</tbody>
</table>

Table 6: The rate of burn wound healing and epithelialization in the studied patients on the seventh and fourteenth days

<table>
<thead>
<tr>
<th>Healing and epithelialization</th>
<th>Frequency</th>
<th>Percent</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Seventh day</td>
<td>Amniotic group</td>
<td>27</td>
<td>81.8</td>
</tr>
<tr>
<td></td>
<td>Routine group</td>
<td>16</td>
<td>48.5</td>
</tr>
<tr>
<td>Fourteenth day</td>
<td>Amniotic group</td>
<td>33</td>
<td>100</td>
</tr>
<tr>
<td></td>
<td>Routine group</td>
<td>26</td>
<td>78.8</td>
</tr>
</tbody>
</table>

Discussion

Burn injury is a complication that is one of the most important incidents in human life related to health due to severe complications and very high mortality. Burn injuries are still one of the leading causes of death and disability in the world (20). Considering the use of antimicrobial method in Imam Khomeini hospital in Urmia and also the high rate of complications due to burns and high mortality and mortality, this clinical trial study aimed to compare the short-term consequences of amniotic membrane dressing and routine dressing in graft donor site in 33 patients with mean age of 26.82±3.37 years, of whom 69.7% were male and 30.3% were female.

In our study, there was no significant difference regarding infection between the two methods of using the amniotic membrane and routine dressing. In the study by Branski et al (21), the use of amniotic membrane was not associated with an increased risk of local infection. Several experimental animal studies have confirmed the antimicrobial properties of amniotic membrane in reducing wound infection (21-25). In another study by Bujang et al (24), silver sulfadiazine dressing was used as well but the rate of wound infection was lower in the amniotic dressing group again. Their results were also obtained in our study and no effect of infection was observed in our patients, which could confirm the safety of the amniotic membrane in dressing wounds caused by burns.

In 2015, Ullah et al (25) used the amniotic membrane to dress 370 burn patients. Amniotic membrane creates a dry environment which reduces infection. These results were consistent with the findings of our study. In our study, the mean pain score on the seventh and fourteenth days was significantly different between the two groups, hence patients complained of more pain in the routine dressing. As a result, less pain was reported in the amniotic membrane dressing group, which could be due to the effect of different cytokines in the amniotic membrane, such as transforming growth factor beta (TGF-β), which leads to less analgesic use. In a study by Mostaque et al (26), the pain status showed a significant difference between the two groups, in which amniotic membrane treatment was more accepted by patients or parents. Our study showed that the mean recovery time in the amniotic membrane dressing group was 2.94±1.18 days and in contrast in the routine dressing group was 4.14±2.04 days; the difference
between the two was significant. In a study by Pakel et al (23), they concluded that the amniotic membrane causes rapid epithelialization of the burn without the risk of metalloprotein accumulation, which was consistent with the results of our study. Accordingly, Subrahmanyam et al (17) showed that using the amniotic membrane helps premature epithelialization and accelerates wound healing, which was consistent with the results of our study.

**Conclusion**

Finally, it can be concluded that the employment of amniotic membrane for dressing grade II burn wounds, compared to routine dressing, has good results and benefits including less pain, faster epithelialization, duration, better recovery and reception. Another advantage of amniotic membrane dressing is that in case of burns below 10% that patients do not have the indication for hospitalization, dressing with amniotic membrane which can be conducted on an outpatient admission in the emergency room without the need for anesthesia and operating room.

**Limitations of the study**

One of the limitations of this study was the need for almost long-term follow-up of the patients and justification and cooperation of as many patients as possible.

**Authors’ contribution**

JK and HS designed the study. HS performed the experiments. HS collected data from patients and helped in performance of experiments. JK and HS prepared the primary draft after analysis. Both authors read and signed the final paper.

**Conflicts of interest**

There are no any conflicts of interest.

**Ethical approval**

The research followed the tents of the Declaration of Helsinki. Accordingly, written informed consent taken from all participants before any intervention. This study is a randomized controlled trial which registered in the Iranian Registry of Clinical Trials (IRCT20181216041996N1, https://en.irct.ir/trial/35985). The study was reviewed and approved by the committee of Urmia University of Medical Sciences (IR.UMSU.REC.1397.245) too. This study was extracted from general surgery residency thesis of Houshang Soleimani at this university (Thesis#2582). Besides, ethical issues (including plagiarism, double publication) have been completely considered by the authors.

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