The Fistula Adapter - an Evolving Device in Treatment of Enteroatmospheric Fistulæ

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Abstract — Negative pressure wound therapy is established in the management of the open abdomen but displays major problems in case of an accompanying intestinal fistula. Here we present a novel device for standardized treatment of enteroatmospheric fistulæ - the fistula adapter.

Keywords — enteroatmospheric fistula, intestinal fistula, negative pressure wound therapy, open abdomen, vacuum therapy.

I. INTRODUCTION

The introduction of negative pressure wound therapy (NPWT) has improved the treatment of an open abdomen in many ways. However, enteroatmospheric fistulæ (EAF) remain a serious problem. NPWT, applied via foam to the wound surface, often leads to enlargement of the fistula or excrescence of the bowel mucosa. Additionally, viscous stool might obstruct the foam. This results in a separation of the foam from the wound surface. The suction starts to leak and this may lead to contamination or infection of the wound. Several drain-out systems using NPWT have been published. Unfortunately none of them could completely solve this problem. In most cases individual, time-consuming modifications of the wound dressing are necessary to avoid fistulas and obstruction of NPWT devices. In a first publication in 2011 we introduced the fistula adapter (FA) for treatment of EAF. Since then three new modifications have been developed to overcome some of the experienced problems. In this work we introduce these strategies.

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Fig. 1. Four available versions of the PPM-fistula adapter™-set (Fa. Phametra, Pharma & Medica-Trading GmbH, Herne/Ruhrstadt, Germany)

Fig. 2. Two fistula adapters inserted in a foam, the lower brim is cut to match the outline of the foam

II. TECHNICAL DESCRIPTION

The FA consists of a soft thermoplastic elastomer to prevent pressure on the underlying wound surface. The original version had an inner diameter of 1.5 cm and a height of 3 cm (adjusted to commercially available polyurethane foams). A lower brim serves as interface to the wound surface and the upper brim as connection to an ostomy bag. Because of limitations with the original FA there are now four versions available (fig. 1).

Indication for use of the FA is a stage 4 open abdomen (frozen abdomen) with an enteroatmospheric fistula. In selected cases use in stage 3 open abdomen might also be indicated.7 Treatment goal is the healing of the wound area surrounding the EAF. This enables conventional ostomy therapy and discharge of the patient for further outpatient treatment. 6-12 month later a surgical resection of the fistula and final closure of the abdomen might be attempted.
After measurement of the fistula and of the wound size, a polyurethane foam is cut to match the wound surface. The opening in the foam for insertion of the FA can be made by using scissors or scalpel. The foam is easily manageable, so remaining parts can be stripped away without much effort. Then the adapter is inserted into the preformed hole in the foam. The hole should be a little smaller than the FA. If the adapter will be deformed after insertion, it can be easily reshaped by a little pressure. The fistula might be positioned next to the wound edge. The lower brim tends to overlap in this case. We usually cut it with the scissors to match the outline of the foam. Also more than one FA can be used in a foam (fig. 2). Next a silicone gauze will be aligned to the wound surface. In this way direct contact of the foam with the wound surface is prevented (fig. 3). Now the foam is placed. The FA has to enclose the underlying fistula. Afterwards the prepared foils are affixed to keep foam and FA in place. It is best practice to remove the coverfilm from the foil and put the middle of the foil onto the fistula adapter. Then the foil is carefully fixed laterally. Next a hole for the suction pad is cut into the foil (fig. 4).

In case of a small wound area the suction pad cannot be placed directly beside the ostomy base plate. We therefore build a so called air-bridge. In this area the foil is already covering the skin, so there is no direct contact between foam and skin. This second small foam also has to be covered by a foil (fig. 5-7).
In a next step the suction pad is fixed. Afterwards the appliance can be put above the adapter after incision of the foil covering the FA. Then NPWT is activated. By this the FA is reliably fixed to the wound surface (fig. 8). According to the wound conditions a negative pressure of 75 to 125 mmHg should be used. To check the wound dressing we first look at the foam. It has to be contracted at all areas. To functionally control leak tightness we open the two-pieces appliance. If the foam remains contracted, we have proven leak tightness of the system (fig. 9). If the ostomy bag is filled with fluid or stool it can be emptied at any time or continuously via drain (fig. 10).

III. CONCLUSION

Here a novel device for managing EAF in a frozen abdomen using NPWT is presented. In most cases stable separation of the fistula from the negative pressure acting on the surrounding wound is achieved. The system can be easily applied and supports early mobilization of the patient as well as split thickness skin grafting. Most patients can be discharged for outpatient treatment with a conventional ostomy bag and, if desired, surgical revision 6–12 month later. With help of the newly developed versions of the FA some of the remaining problems could be solved. These are:

- large fistulae up to 4 cm in diameter
- multiple fistulae lying next to another
- high output-volume fistulae
- colonic fistulae with formed stools
- deep positioned fistulae

For the problem of fistulae directly located at the wound margin an additional version of the FA is being developed.

REFERENCES:


