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Solving Periprosthetic Leakage With a Novel Prosthetic Device

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INTRODUCTION

Total laryngectomy (TL) is still an indispensable surgical procedure for advanced and recurrent larynx and hypopharynx cancer. Tracheoesophageal prosthetic voice rehabilitation is the method of choice for restoring oral communication in most Western countries, with success rates of around 90%.1 However, with an increasing rate of salvage TLs performed after failed chemo(radiotherapy), more attention is needed to maintain durable results.

Voice prostheses (VPs) have a median device lifetime of around 2 to 3 months, and the main indication for replacement is transprosthetic leakage, which is solvable by replacing the VP.2–4 Recurrent periprosthetic leakage is, however, a problem requiring more attention.3,4 It can be caused by normal gradual subsiding of postsurgical edema, making the VP too long and thereby permitting periprosthetic leakage, which is solvable by downsizing the VP.1 Later in time, comorbidities such as gastric reflux, local infection, radiation effects, or recurrence of tumor can have a profound effect on the surrounding tissue. This can lead to atrophy and/or widening of the tracheoesophageal puncture (TEP) tract, also resulting in periprosthetic leakage.6,7 Therefore, comorbidities such as reflux first should be treated adequately when possible in order to prevent periprosthetic leakage on the long term.

Often, the easiest short-term solution is placing a thin silicone washer on the tracheal side of the VP if the VP is still functioning properly. This is simple, effective, and cheap solution because there is no need to replace the current VP. It is, however, obvious that in case of periprosthetic leakage the fluids originate from the esophageal side. A washer on that side is more effective than a washer on the tracheal side; otherwise, the fluids are still able to penetrate the TEP tract up to the tracheal side. However, this means replacement of the VP and thus higher costs. Patients known with recurrent periprosthetic leakage could benefit from instantly placing a VP with an enlarged esophageal flange. Thus, a new VP with an extraesophageal flange (Provox Vega XtraSeal [PVX], Atos Medical, Hörby, Sweden) was developed, which we tested at our institute.

MATERIALS AND METHODS

We performed a prospective evaluation on the efficacy, satisfaction, and ease of placement of the PVX among a consecutive cohort of patients seen in the outpatient clinic with periprosthetic leakage. After placement of the PVX, patients and physicians were asked to fill in a study-specific questionnaire regarding the satisfaction of (placement of) the PVX with regard to the handling of the insertion device and the procedure. Patients were excluded from follow-up (FU) if they had received two successive VPs other than a PVX.

Prosthesis

The PVX is an adjustment of the regular Provox Vega with an additional enlarged esophageal flange glued to the VP at the flange-shaft crossing (see Figure 1). The flange is angled, thin, and flexible, which should enhance its adherence to the surface around the TEP-tract to prevent leakage around the VP. The prosthesis is inserted with the regular insertion device, with special attention to the proper unfolding of the enlarged esophageal flange by inserting the entire VP into the esophagus (overshooting) and pulling the tracheal flange back in position.

Statistics

Descriptive statistics were used to report patient characteristics and Kaplan-Meier analysis to assess device lifetime. All analyses were done in SPSS Statistics 20.0 (IBM Corp., Armonk, NY).

This study does not fall under the scope of the Medical Research Involving Human Subjects Act, which was confirmed by the institutional review board (MREC16.1202).

RESULTS

We included 13 patients (85% male). The mean age at TL was 59 years, and median FU after TL was 117 months (see Table I).
The reason for placement of the first PVX in each patient was periprosthetic leakage (n = 11), a too wide TEP tract (n = 1), or a lost VP (n = 1). These latter two replacements were performed in two patients who were known with recurrent leakage around the VP and were therefore included in this study. In these 13 patients, 26 PVXs were placed. Five patients received multiple PVXs during FU, with a maximum of seven PVXs in one patient (see Table II).

After replacement, the seal was checked by the patient drinking water. The seal was sufficient in 25 of 26 placements. In the remaining replacement, calcium hydroxyapatite (Radiesse; Merz Pharmaceuticals, Germany) was injected in the oval-shaped TEP-tract, which solved the persistent periprosthetic leakage.

Results from the study-specific questionnaire indicated that loading of the PVX in the insertion device went well in all cases except one, for which more force than usual was needed during the overshooting phase. All but one patient reported no difference in ease and discomfort during placement; this latter patient favored placement of the new VP.

**Device Lifetime**

The median device lifetime of the PVX was 68 days (95% CI 56–80). Median device lifetime of the former VP before placement of the first PVX was 38 days (95% CI 1–76). One patient died 3 days after placement of the PVX from a metastasized esophageal cancer. Former VPs led to aspiration problems; with the PVX, the patient was aspiration-free. A second patient with an irresectable tracheal recurrence also died with the third PVX in situ. The patient was free from periprosthetic leakage since insertion of the first PVX; the first two PVXs lasted 79 and 62 days, respectively.

**Reason for Removal**

The main reason for removal of the PVX was transprosthetic leakage in 50% (13 of 26), followed by leakage not otherwise specified in 15% (4 of 26), which probably all were cases of transprosthetic leakage. In one patient (4%) the PVX had to be removed because of periprosthetic leakage. See Table II for the other reasons. Two patients still had PVX in situ at last date of FU (June 2018), with device lifetime of 504 and 835 days, respectively. No adverse events occurred during the study period.

During FU, in seven patients the PVX was replaced with a Provox Vega, in three cases combined with a washer at the tracheal side. One patient went back to the usual ActiValve Light. The median in situ time of the subsequent non-PVX VP was 62 days. The reasons for removal of these non-PVX VPs were periprosthetic leakage (n = 3), transprosthetic leakage (n = 3), and surgical revision (n = 1). Of the five remaining patients, two still had a PVX in situ at last date of FU; two died; and the TEP-tract was closed in one patient.

**DISCUSSION**

In this prospective evaluation of the PVX, we were able to test the device lifetime, efficacy, and ease of placement. The median device lifetime was 68 days, comparable to the median device lifetimes of the Provox2 (63 days) or Provox Vega (66 days), which we recently found in a consecutive cohort of patients for over 13 years and is in line with other literature. Only one PVX needed replacement due to periprosthetic leakage, although in one patient the reason for removal was unknown and in four patients the leakage problem was not otherwise specified.

A recent meta-analysis reported an average rate of 7.2% of patients suffering from an enlarged TEP tract and/or periprosthetic leakage. The most commonly used

![Fig. 1. Schematic drawing and actual photo of the Provox Vega XtraSeal (Atos Medical, Hörby, Sweden) showing (A) the location of the normal esophageal flange, (B) the extra thin, angled esophageal flange, (C) and the tracheal flange. [Color figure can be viewed in the online issue, which is available at www.laryngoscope.com.]](image_url)
treatments were temporary removal of the VP and injections at the TEP tract. Temporary removal and placement of a nasogastric feeding tube and cuffed canula is, however, quite cumbersome for the patient, and it might take several days before sufficient shrinkage is observed. Placement of a silicone washer is usually an elegant and conservative solution to manage periprosthetic leakage, especially when the VP is still functioning properly. If the insertion of a washer fails, other strategies such as injection of a filler-like hydroxyapatite, fat, or collagen; the application of a purse string suture; or temporary removal of the VP to allow for shrinkage could be tried to prevent unwanted surgical closure of the TE fistula.

Earlier studies have reported success rates of 77% to 88% in managing periprosthetic leakage with an enlarged flange on the tracheal side. Kress et al. described 76 patients with periprosthetic leakage who were managed with custom fit VPs with an enlarged flange on the esophageal side and were highly successful (97%). Choussy et al. evaluated 28 Blom-Singer large esophageal and tracheal flange VPs (InHealth technologies, Carpinteria, CA, USA) in 18 patients and reported success in all patients, with a median device lifetime of 70 days (range 24–219). It indeed seems logical that an extra flange on the esophageal side is more successful than a flange on the tracheal side because it provides a better seal to the mucosa. However, a tracheal flange can be placed on an existing VP, whereas an esophageal flange usually necessitates replacement of the VP and thus higher costs.

Due to local reimbursement differences and costs of VPs in various countries, it is difficult to give exact numbers, but on average the costs of a Provox Vega combined with a silicone washer are quite comparable with that of a PVX, ranging from 90% to 110% of the costs of a PVX (communication by manufacturer, Atos Medical, Hörby, Sweden). If there is need for replacement of the VP, a washer on the esophageal side/PVX is most effective; however, if there is no need for replacement, a washer on the tracheal side is most cost-efficient.

**CONCLUSION**

With this prospective study, we have demonstrated that the new PVX adds a valuable new tool to solving...
periprosthetic leakage, diminishing the burden of this uncomfortable adverse event both for the patient and the clinician. We were able to solve almost all cases of periprosthetic leakage and were able to reach an adequate median device lifetime of 68 days, comparable to current device lifetime of modern voice prostheses.

BIBLIOGRAPHY


