A Myringotomy and Ventilating Tube Applicator: New Look at a Five-Century-Old Procedure

ABSTRACT

Objectives: To fabricate a single instrument that can be used to perform myringotomy and insert a pressure equalizing tube at almost the same time.

Methods: Design: Surgical Instrumentation Setting: Tertiary Private Hospital Subject: A chicken egg membrane was used as a tympanic membrane model

Results: The fabricated instrument was able to perforate the egg membrane and apply the modified polyethylene pressure equalizing tube in less than one minute without complications.

Conclusion: The prototype applicator can facilitate myringotomy and pressure equalizing (PE) tube insertion at only a fraction of the time it usually takes to do the standard myringotomy and subsequent ventilating tube insertion.

Keywords: Myringotomy, pressure equalizing tube, ventilating tube, acute otitis media, otitis media with effusion, polyethylene tube

Myringotomy with pressure equalizing (PE) or ventilating tube (VT) insertion is one of the most common ambulatory ear surgeries performed in the United States with 19.7 thousand (92.6%) performed in the out-patient setting in 2012 alone.1 It is a relatively rapid procedure and may be performed under topical anaesthesia. The standard procedure is to perform myringotomy initially, then insert the pressure equalizing or ventilating tube subsequently. The myringotomy may cause some pain or discomfort to the patient and affect his/her cooperation when the tube insertion is done next. Moreover, commercial tubes are costly and cannot be readily obtained in many settings. We describe our fabricated applicator and surgical technique to facilitate myringotomy with PE tube insertion and reduce operative time using an egg model.
METHODS

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<th>Tools and equipment:</th>
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<td>a. Alligator forceps</td>
<td>a. 254 mm Hacksaw</td>
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<td>b. Medical grade Polyethylene tube, PE 205</td>
<td>b. Carbide stone</td>
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<td>c. Metal stopper from leftover titanium mesh</td>
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<td>d. Chicken Eggs</td>
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<td>e. Cyanoacrylate (super glue)</td>
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<td>g. Straight forward endoscope 0°, 4mm</td>
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<td>h. Magnifying lens</td>
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FABRICATION

MYRINGOTOMY TUBE APPLICATOR

1. The alligator forceps was grasped with the plastic grip. The distal tip of the alligator forceps was cut to remove 1cm using a hacksaw.

2. The cut tip of the alligator forceps was sharpened using a carbide stone.

3. The corners of the shaft were blunted using a round file.
4. A metal stopper was attached on top of the shaft near the sharpened edge using super glue.

**Figure 5.** Attaching metal stopper on top of shaft using super glue

**PRESSURE EQUALIZING TUBE**

1. A 6mm segment of polyethylene with a bevelled edge was cut using a blade or sharp scissors.

**Figure 6.** Cutting the polyethylene tube

2. The non-bevelled edge was melted over an open flame until the tube was about 4mm in height and the melted edge rolled up and measured around 4mm in diameter, forming a flange.

**Figure 7.** Flanging the tube

**RESULTS**

1. The plunger was retracted by opening the forceps.

**Figure 8 A.** Forceps closed (above) with close up of tip (below); B. Forceps opened (above) with close up of tip (below) showing plunger tip retracted from applicator tip.

2. The polyethylene tube was loaded on the applicator.

**Figure 9.** Loading fabricated ventilating tube on applicator
3. The loaded forcep applicator was inserted through an ear speculum abutting the exposed membrane of a partly-shelled chicken egg under endoscopic guidance.

4. The exposed membrane of a partly-shelled chicken egg was pierced making sure the flange touched the membrane lightly and then the forceps were closed, pushing the tube and leaving it in the membrane.

5. The forceps applicator was retracted, and fluid was suctioned through the ventilating tube. The procedure was repeated 10 times with similar results and without complications.

DISCUSSION

Otitis media with effusion is a common disorder of the middle ear caused by poor clearance of middle ear fluid usually following an episode of acute otitis media. In our current clinical practice guidelines (CPG), a 3-month waiting period of conservative management is preferred. However, in cases where there is persistence of disease and significant hearing loss or for children with recurrent episodes of AOM usually defined as three or more episodes of AOM in 6 months or four or more episodes in 12 months, myringotomy with pressure equalizing (PE) or ventilating tube (VT) insertion is the usual management. It is also performed in adults and may be done under topical anesthesia.
Myringotomy is a very old procedure. The first recorded myringotomy was in 1649 when Jean Riolan the Younger accidentally perforated the eardrum of a patient complaining of ear pain. Astley Cooper presented two papers to the Royal Society in 1801; based on his observation that myringotomy could improve hearing. However, due to injudicious use of the procedure, the practice of myringotomy was almost abandoned. The use of myringotomy was again reintroduced during the 19th century and is presently the most common surgical procedure in children that requires general anesthesia. Innovations have been introduced and different types of pressure equalizing (PE) tubes are currently available in the market. (Figure 13)

The area of the middle ear cavity on the average is 2 cm³ and the widest diameter does not exceed 10 mm. The dimensions of the tympanic membrane along its two major perpendicular axes are 9 to 10 mm (0.35 to 0.39 in) and 8 to 9 mm. With this, it is possible to create an instrument that can serve the purpose of perforating and applying a tube to equalize the pressure in the middle ear and drain fluid from the cavity.

The length of the sharp portion of the fabricated applicator to the metal flange is 6 mm and its widest diameter is 1.5 mm. The dimensions of the fabricated PE tube are 4 mm in length, 2 mm outer diameter, 1.5 mm inner diameter and the flange is 4 mm in diameter. (Figure 14) The size of the applicator and tube are adequate to perforate and fit a ventilation tube in the tympanic membrane. The 1 mm forward and backward motion of the upper part of the forceps is sufficient to push the PE tube and release it from the rest of the mechanism.

The standard myringotomy with pressure-equalizing tube insertion is generally performed as follows: Firstly, the antero-inferior portion of the TM is incised using a myringotomy knife perpendicular to the fibers of the membrane. Secondly, the tube is inserted through the perforation using an alligator forceps. With our innovation, both steps were combined to reduce the time it takes to perform the procedure by half. This could therefore minimize discomfort of the patient and allow ease of application.

The entire procedure was endoscopically guided to allow superior visualization of the ear canal and the tympanic membrane. The upper part of the applicator was simply retracted and the PE tube loaded. The applicator loaded with the PE tube was then inserted through the canal to
perforate the TM making sure that the flange touched the tympanic membrane. The flange was then pushed forward by closing the forceps and releasing the tube. As the whole contraption was initially retracted, the PE tube was automatically left in place once released. We repeated the procedure 10 times with similar results and no complications.

The procedure was performed on an egg membrane to simulate the tympanic membrane as it also contains fluid and somewhat comparable to the consistency of the tympanic membrane. However, the exact pressure needed to perforate the egg membrane was not measured and may not be similar to that of the tympanic membrane. The pressure needed to perforate an ear drum at the minimum is 5 psi; however, 50% will perforate at 15 psi. To our knowledge, this is the first report that investigated using egg membrane as a model for prototype myringotomy applicators. Our innovation has yet to be tested on a human subject. However, the test procedure showed that our prototype applicator was able to perforate the membrane and leave the PE tube in place.

With the tube in place, middle ear fluid can be suctioned and middle ear pressure maintained equal with atmospheric pressure. Suctioning was performed in our model to demonstrate patency of the fabricated ventilating tube. Depending on the type and size of tube used, spontaneous extrusion may occur from 12-18 months, a property we have yet to test using our fabricated tube.

There are several limitations to this study. The ideal size and shape of commercially available tubes which are uniformly beveled and tapered can only be approximated by our method. The most anterior end of the tube may not always conform to the shape of the instrument and may cause difficulty perforating the membrane. The tip of the instrument was also not surgically sharpened, and minute metal fragments may remain. More advanced equipment and tools would allow us to fabricate a better applicator and tube. Following this, the prototype applicator and tube can be tested in human cadaver tympanic membranes and on live subjects. Sterilization by autoclave (forceps applicator) and/or activated Glutaraldehyde solution (applicator and tube) will have to be tested for such a trial.

In conclusion, our prototype forceps applicator and ventilating tube may simplify myringotomy and pressure equalizing (PE) tube insertion and decrease the surgical time to only a fraction of the time needed for the standard procedure, which may take 10-15 minutes. It is inexpensive to fabricate and may potentially reduce the total cost of the procedure as well and we recommend considering its development and use especially in low- and middle-income country settings.

REFERENCES