Percutaneous Endoscopic Gastrostomy Does Not Prolong Survival in Patients With Dementia

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Background: Artificial feeding by a percutaneous endoscopic gastrostomy (PEG) tube in patients with dementia has increased since the introduction of the endoscopic method of tube placement. Few studies have documented survival benefit from this intervention. This report reviews our experience with PEG tube placement for feeding patients with dementia.

Methods: All consultations for PEG tube placement were evaluated by a certified nutrition support nurse (L.M.M.) in consultation with a member of the gastroenterology physician staff (T.O.L.) for 24 months. Evaluation included the attainment of a brief medical history, a physical examination, and a review of comorbid conditions, laboratory variables for nutrition status, and bleeding risk. Interviews with patients or surrogates were conducted, including an explanation of the risks and benefits of PEG tube placement. A Kaplan-Meier survival curve was used to compare the median survival between patients with dementia who received a PEG tube and patients with dementia in whom PEG tube placement was refused.

Results: We received 41 consultations for PEG tube placement in patients with dementia. Percutaneous endoscopic gastrostomy was performed in 23 patients; 18 patients met the medical criteria for PEG tube placement, but surrogates refused placement. The median survival for the 23 patients who underwent PEG was 59 days; the median survival for the 18 patients who did not undergo PEG was 60 days.

Conclusion: There seems to be no survival benefit in patients with dementia who receive artificial feeding by a PEG tube.

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We received 41 requests for PEG in patients with dementia. All patients met our criteria for PEG: advanced dementia had been documented in the medical record, the patient had documentation of dysphagia, life expectancy was considered to be at least 30 days, there was no contraindication to conscious sedation, and there was no other disease contributing to dysphagia. Percutaneous endoscopic gastrostomy tubes were placed in only 23 of these patients with dementia. The other 18 similar patients with dementia did not undergo PEG because the procedure violated advance directives or the appropriate surrogate refused on the patient’s behalf.

The median survival for the 23 patients with dementia who underwent PEG was 59 days (range, 2-219 days). The median survival for the 18 patients with dementia who did not undergo PEG was 60 days (range, 2-229 days). There was no statistically significant difference in survival between the groups ($P = .37$, $df = 1$) using the Kaplan-Meier survival curve (Figure). There was one major complication in the group that underwent PEG—an intra-abdominal abscess, resulting in sepsis and death, making our complication rate 4.3%.

Our findings support the hypothesis that there is no survival benefit to the placement of a PEG tube for artificial feeding in patients with advanced dementia. All patients would have been candidates for PEG tube placement, but the procedure was refused in 18 (44%). Thus, in a similar cohort of patients, PEG or no PEG did not enhance survival.

Our complication rate of 4.3% in those who underwent PEG does not differ from those in other literature reports.$^8$-$^{10}$

More reports$^{11-13}$ question the utility of PEG tube placement in patients with dementia; the 1-month mortality is as high as 54%. Little benefit from PEG has been established for any variable studied—aspiration pneumonia, nutrition status, pressure sores, functional status, patient comfort, or survival.$^{14-16}$

Nevertheless, despite a bleak prognosis for survival in patients with advanced dementia undergoing PEG, the alternative—no feeding—would seem worse. Surrogate decision makers are often presented with a bleak choice—agree to PEG or “let your loved one starve to death.” Ideally, to determine whether PEG-based artificial feeding enhances survival, a prospective, controlled, randomized study would have to be performed. However, such a study would be questioned ethically, and it is highly unlikely that a sufficient number of volunteers would agree to be randomized to PEG or no PEG tube insertion. We are then left with trying to ascertain benefit using indirect means. Three studies$^{5-7}$ evaluating information from Health Care Financing Administration–required data in nursing homes have come to disparate conclusions—artificial nutrition by tube enhances, worsens, or does not alter survival in nursing home residents with chewing and swallowing disorders. However, in all of these studies, it is not clear that the nursing home residents studied had dementia or even underwent PEG for tube feeding. Furthermore, it is quite plausible that the groups were not comparable, with sicker patients receiving feeding tubes. If sicker patients receive feeding tubes, then it is not possible to determine whether feeding tube or PEG tube placement enhance survival. Meier et al$^{13}$ determined that the median survival in a cohort of patients examined for palliative care and undergoing PEG tube placement during a short-term hospitalization did not differ from patients in the same cohort who left the hospital without undergoing PEG (median survival, 195 vs 189 days; $P = .90$). However, it is not clear from the data why patients did or did not undergo PEG tube placement.

The evidence against PEG tube placement for artificial feeding in patients with dementia is substantial. Hospice literature suggests that avoiding artificial nutrition and allowing the patient to consume food and fluids ad lib may enhance comfort. We need to separate the need for the nurturing aspect of food from the provision of artificial nutrition.$^{17}$ The provision of artificial nutrition may lead to an increase in urine and fecal incontinence and increased pulmonary secretions. Incontinence has been associated with an increased risk of pressure ulcers. The
patient with dementia may need physical or chemical restraint to avoid self-extubation. The act of patient restraint also has been identified as a risk factor for pressure ulcers. Furthermore, physical restraint may be seen as a violation of the patient’s right to dignity.

The limitations of our study include the small sample size, the lack of a prospective randomized approach, an all-male sample, and the inability to generalize the results to all patients with dementia. In addition, we do not have clinical information for the 2 groups of patients documenting similarity. However, the patients were similar in that all had advanced dementia, we were requested to place PEG tubes to facilitate hospital discharge, and we would have placed the PEG tube in all patients for whom we were consulted if we had surrogate permission. The difference between the groups was in the refusal of PEG tube placement by the patients (based on advance directives) or their surrogates (based on a careful and thorough explanation of potential risks and benefits).

In summary, in patients with advanced dementia and dysphagia, placement of a PEG tube neither enhances survival nor prevents death by starvation. This does not mean that patients with dementia should not have the right to receive artificial feeding via PEG tube placement. However, this intervention should be undertaken only after full discussion and understanding of the risk and lack of the following benefits: comfort, treatment and prevention of pressure sores, prevention of aspiration, and enhancement of survival.

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REFERENCES

that one limitation of prospective, randomized data is a selection bias in which outcomes are preordained. Thus, the results may very well be valid but are not necessarily generalizable for the population studied.

A subsequent analysis of a subgroup of patients in this trial known to have ischemic heart disease demonstrated an inversion of the survival curves, so that the liberally transfused patient had better survival rates (albeit not statistically significant, since the trial was not powered to detect a difference in this subgroup). With the current ease and safety with which anemia can be treated, we would suggest that waiting for data only from prospective, randomized studies before establishing or revising clinical care guidelines may not be in the best interests of all patients, particularly those at risk.

As we hope readers are aware, the Archives publisher made a serious error in changing Goodnough from first to second author in our commentary published in the June 23 issue; the Archives has published a correction, and we also note this correction in this exchange of letters.

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**Correction**

In the Original Investigation by Murphy and Lipman titled “Percutaneous Endoscopic Gastrostomy Does Not Prolong Survival in Patients With Dementia,” published in the June 9 issue of the Archives (2003; 163:1351-1353), an error occurred in the Figure on page 1352. In that figure, the lines in the figure key were reversed. The patients who underwent percutaneous endoscopic gastrostomy (PEG) (n=23) should have been indicated by the dashed line, and the patients who did not undergo PEG (n=18) should have been indicated by the solid line. The journal regrets the error.