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Nasogastric tube versus postpyloric tube feeding for critical illness:

A systematic review and meta-analysis

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Running title: Nasogastric tube versus postpyloric tube feeding

Liru Li MS¹, Jie Huang MR²

¹Department of emergency medicine, Shanghai Fengxian District Central Hospital, Shanghai, China ²Department of Neurology, Shanghai Municipal Hospital of Traditional Chinese Medicine, Shanghai, China

Authors' email addresses and contributions:

LRL carried out the studies, participated in collecting data, and drafted the manuscript. JH performed the statistical analysis and participated in its design. All authors read and approved the final manuscript.

Corresponding Author: Liru Li, Department of emergency medicine, Shanghai Fengxian District Central Hospital, No.6600 Nanqiao Xincheng Nanfeng Highway, Fengxian District, Shanghai, 201499 China. Tel: 13764336577. Email: liliru_724@126.com; Jie Huang, Department of Neurology, Shanghai Municipal Hospital of Traditional Chinese Medicine, No.274 Jingan District Zhijiang West Road, Shanghai, 200010, China. Tel:. Email: Hjie1970@163.com

ABSTRACT

Background and Objectives: Gastric tube feeding and postpyloric tube feeding are two common forms of enteral nutrition in critically ill patients. This study aimed to compare the efficacy and safety of gastric tube feeding with that of postpyloric tube feeding in critically ill patients. Methods and Study Design: PubMed, Embase, and Cochrane Library were systematically searched for eligible trials from their inception until March 2023. Relative risks (RRs) or weighted mean differences (WMDs) with 95% confidence intervals (CIs) were used to estimate categorical and continuous outcomes using the random-effects model. **Results:** Sixteen trials involving 1,329 critically ill patients were selected for the final metaanalysis. Overall, we noted that gastric tube feeding showed no significant difference from post-pyloric tube feeding in mortality (p = 0.891), whereas the risk of pneumonia was significantly increased in patients who received gastric tube feeding (RR: 1.45; p = 0.021). Furthermore, we noted that gastric tube feeding was associated with a shorter time required to start feeding (WMD: -11.05; p = 0.007). Conclusions: This research revealed that initiating feeding through the gastric tube required less time compared to postpyloric tube feeding. However, it was also associated with a heightened risk of pneumonia among critically ill patients.

Key Words: enteral nutrition, nutritional support, pneumonia, critical illness, systematic review

INTRODUCTION

Patients with critical illnesses, including severe acute illnesses such as sepsis, severe trauma, or major surgery, are admitted to the intensive care unit (ICU). The characteristics of critical illnesses result in malnutrition and are complicated by other diseases or dysfunctions. Moreover, the generalized inflammatory response could caused by this situation owing to the release of endogenous stress hormones and cytokines.¹ Unmet nutritional needs are significantly associated with energy-protein malnutrition and the breakdown of muscle mass.^{1,2} Although clinical practice guidelines have addressed the importance of nutritional support for critically ill patients, only 40–60% of patients meet the recommended nutritional goals.^{3,4} Studies have already found that malnutrition is associated with an increased risk of nosocomial infection and mortality in critically ill patients and that patients should receive enteral feeding as long as gastrointestinal function permits.⁵⁻⁷

Enteral nutrition (EN) is considered the preferred means of nutritional support owing to its enhancement of gut immune function, lower cost, and lower risk of septic complications.^{8,9} The EN could be provided via various methods, and the two common forms are gastric tube feeding and small intestinal feeding.^{10,11} The use of gastric tube feeding showed that slow gastric emptying could increase the residual gastric volume; in addition, the risk of bacterial colonization and aspiration pneumonia increased in critically ill patients. One study found that the use of a postpyloric tube could overcome the shortcomings of gastric tube feeding and was associated with high absorptive capacity.¹² The nutritional status of ICU patients is significantly associated with the clinical prognosis. However, whether the use of postpyloric tube feeding remained unclear.

Several systematic reviews and meta-analyses have compared the efficacy and safety of gastric tubes with those of postpyloric tube feeding in critically ill patients.¹³⁻¹⁵ Zhang et al.¹³ identified 17 randomized controlled trials (RCTs) and found that postpyloric tube feeding was associated with higher proportions of estimated energy requirements and reduced residual gastric volume, whereas no significant differences were found between groups for the risk of mortality, new-onset pneumonia, and aspiration. Based on a Cochrane review,¹⁴ RCTs were identified. The review indicated that postpyloric tube feeding was linked to a reduced risk of pneumonia and an enhanced delivery of nutrition.¹⁴ In another study by Liu et al.,¹⁵ involving 41 investigations, post-pyloric tube feeding demonstrated an association with diminished risks of pulmonary aspiration, gastric reflux, pneumonia, or gastrointestinal complications, along with more optimal gastrointestinal nutrition. Nevertheless, it's important to highlight several limitations in prior research, including errors in study inclusion, data extraction, failure to include the latest relevant studies meeting the inclusion criteria, and lack of exploratory analysis results. Therefore, the current study was performed to update the efficacy and safety of gastric tube versus post-pyloric tube feeding in critically ill patients.

MATERÍALS AND METHODS

Search strategy and selection criteria

The Preferred Reporting Items for Systematic Reviews and Meta-Analysis (PRISMA) guidelines were used to guide the study and report this systematic review and meta-analysis.¹⁶ Our study was retrospective registered in INPLASY platform and the registered number was INPLASY202380104. This meta-analysis included RCTs designed to compare the effectiveness and safety of gastric tube feeding with postpyloric tube feeding in critically ill

patients. The publication language was restricted to English, while the publication status was not restricted. We systematically searched the PubMed, Embase, and Cochrane Library databases to identify eligible RCTs from their inception until March 2023 using the core search terms, "enteral nutrition" and "critically ill". Details of the search strategy in each database are provided in the Supplementary Table 1. The websites of ClinicalTrials.gov (US NIH) were searched to identify unpublished trials that had already been completed but had not yet been published. We also manually searched the reference lists of relevant reviews and original articles to identify new eligible trials.

Two reviewers independently performed the literature search and study selection, and inconsistent results were resolved by mutual discussion until a consensus was reached. Studies that met the following inclusion criteria were included: (1) Patients: all patients with critical illness and admitted to the ICU; (2) Intervention and control: gastric tube feeding and postpyloric tube feeding; (3) Outcomes: the primary endpoints were mortality and pneumonia, while the secondary endpoints included abdominal distension, diarrhea, vomiting, bacteremia, constipation, gastrointestinal bleeding, high gastric residual volume, pulmonary aspiration, percentage of total nutrition delivered to the participant, time required to achieve the full nutritional target, time required to start feeding, length of ICU stay, length of hospital stay, and length of mechanical ventilation; and (4) Study design: the study had to have RCT design.

Data collection and quality assessment

The abstracted data were independently analyzed by two reviewers, and the collected information included the first author's name, publication year, country, sample size, mean age, proportion of male participants, disease status, Acute Physiology and Chronic Health Evaluation, intervention, control, enteral feeding protocol, and investigated outcomes. The two reviewers independently assessed the methodological quality of the included trials using the risk of bias described by the Cochrane Collaboration, which was based on random sequence generation, allocation concealment, blinding of participants and personnel, blinding of outcome assessment, incomplete outcome data, selective reporting, and other bias. 17 Any disagreement between the reviewers regarding data collection and quality assessment was resolved by referring to the full text of the article.

Statistical analysis

The investigated outcomes were divided into categorical and continuous outcomes. The categorical outcomes were assessed using events/sample size per group, while the mean,

standard deviation, and sample size per group were applied to assess continuous outcomes. The pooled relative risk (RR) or weighted mean difference (WMD) with 95% confidence intervals (CI) was calculated using a random-effects model, which considered the underlying variations across the included trials.^{18,19} Heterogeneity among the included trials was assessed using I² and Q statistics, and significant heterogeneity was defined as I² \geq 50.0% or p < 0.10.^{20,21} The robustness of the pooled conclusions for mortality and pneumonia was assessed using sensitivity analysis through the sequential removal of a single trial.²² Subgroup analyses for mortality and pneumonia were performed according to country, age, the proportion of male participants, and postpyloric tube, and differences between subgroups were assessed using funnel plots and Egger and Begg tests.^{24,25} The reported *p* value for the pooled effect estimates was two-sided, and the inspection level was 0.05. The analyses in this study were performed using STATA software (version 14.0; Stata Corporation, College Station, TX, USA).

RESULTS

Literature search and study selection

The initial electronic search yielded 1,524 studies, of which 553 articles were removed because of duplicate titles. A total of 927 studies were removed because they reported irrelevant articles, and the remaining 44 studies were retrieved for full-text evaluation. Reviewing the reference lists of relevant studies yielded 13 studies, and detailed evaluations were performed for 57 studies; of these, 41 studies were removed owing to a lack of appropriate controls (n = 20), insufficient data (n = 16), and reviews (n = 5) (Supplementary Table 2). The remaining 16 RCTs were selected for meta-analysis,²⁶⁻⁴¹ and the study selection process is presented in Figure 1.

Study characteristics

The baseline characteristics of the identified trials and the patients involved are summarized in Table 1. A total of 1,329 critically ill patients from 16 RCTs were identified, and the sample sizes ranged from 25 to 180. The mean age of the included patients ranged from 34.2 to 82.0 years, and the proportion of male participants ranged from 48.7% to 77.5%. Among the trials included, three employed a duodenal tube, eight utilized a jejunal tube, and the remaining five opted for a smaller intestinal tube for postpyloric tube feeding. The methodological quality of the included trials is presented in the Supplementary Table 3. Overall, the included trials reported a low risk of bias for random sequence generation, allocation concealment, incomplete outcome data, selective reporting, and other biases, whereas there was a high risk of bias for the blinding of participants, personnel, and outcome assessment.

Primary endpoints

Thirteen trials reported the effects of gastric tube versus postpyloric tube feeding on the risk of mortality. There was no significant difference between gastric tube and postpyloric tube feeding for the risk of mortality (RR: 0.99; 95% CI: 0.83–1.17; p = 0.891; Figure 2), and no evidence of heterogeneity across included trials was observed ($I^2 = 0.0\%$; p = 0.872). Sensitivity analysis indicated that the pooled conclusion was stable after the sequential removal of individual trials (Supplementary Figure 1). The results of subgroup analyses were consistent with those of the overall analysis of all subsets (Table 2). No evidence of publication bias for mortality was observed (p value for Egger: 0.087; p value for Begg: 1.000; Supplementary Figure 2).

Twelve trials reported the effect of gastric tube feeding versus postpyloric tube feeding on the risk of pneumonia. The summary of the results indicated that gastric tube feeding was associated with an increased risk of pneumonia as compared with postpyloric tube feeding (RR: 1.45; 95% CI: 1.06–1.99; p = 0.021; Figure 3), and unimportant heterogeneity was observed among included trials (I² = 22.5%; p = 0.223). Considering the lower limit of 95% CI was approximately 1.00, the pooled conclusion was not stable after the sequential removal of a single trial (Supplementary figure 3). Subgroup analyses found that gastric tube versus postpyloric tube feeding showed an elevated risk of pneumonia if patients' age was ≥ 55.0 years, the proportion of male participants was $\geq 70.0\%$, and used duodenal tube feeding as the control group. No other significant difference was observed between gastric tube versus postpyloric tube feeding for the risk of pneumonia in subgroup analyses (Table 2). There was no significant publication bias for pneumonia (p value for Egger: 0.059; p value for Begg: 0.150; Supplementary Figure 4).

Secondary endpoints

The breakdown of the number of trials reporting the effects of gastric tube versus postpyloric tube feeding on the risk of abdominal distension, diarrhea, and vomiting was 5, 11, and 7 trials, respectively (Figure 4). Overall, gastric tube feeding has no significant effects on the

risk of abdominal distension (RR: 1.36; 95% CI: 0.99–1.88; p = 0.061), diarrhea (RR: 0.93; 95% CI: 0.74–1.18; p = 0.571), and vomiting (RR: 1.34; 95% CI: 0.85–2.12; p = 0.210). There was no evidence of heterogeneity for abdominal distension (I² = 0.0%; p = 0.478) or diarrhea (I² = 0.0%; p = 0.912), whereas significant heterogeneity was observed for vomiting (I² = 47.3%; p = 0.077).

The breakdown of the number of trials reporting the effects of gastric tube versus postpyloric tube feeding on the risk of bacteremia, constipation, gastrointestinal bleeding, high gastric residual volume, and pulmonary aspiration was three, two, four, two, and four trials, respectively (Figure 5). There were no significant differences between gastric tube and postpyloric tube feeding for the risk of bacteremia (RR: 0.94; 95% CI: 0.45–1.97; p = 0.871), constipation (RR: 1.39; 95% CI: 0.70–2.78; p = 0.349), gastrointestinal bleeding (RR: 0.67; 95% CI: 0.33–1.36; p = 0.269), high gastric residual volume (RR: 3.77; 95% CI: 0.07–215.21; p = 0.520), and pulmonary aspiration (RR: 0.91; 95% CI: 0.44–1.88; p = 0.792). No significant heterogeneity was observed for bacteremia (I² = 0.0%; p = 0.636), constipation (I² = 0.0%; p = 0.498), while substantial heterogeneity was observed for high gastric residual volume (I² = 92.1%; p < 0.001).

The breakdown of the number of trials reporting the effects of gastric tube versus postpyloric tube feeding on the percentage of total nutrition delivered to the participant, time required to achieve the full nutritional target, and time required to start feeding was six, four, and five trials, respectively (Figure 6). There were no significant differences between gastric tube and postpyloric tube feeding for a percentage of total nutrition delivered to the participant (WMD: -7.62; 95% CI: -15.49-0.26; p = 0.058), and time required to achieve the full nutritional target (WMD: -1.28; 95% CI: -6.51-3.95; p = 0.631), while gastric tube feeding was associated with shorter time required to start feeding as compared with postpyloric tube feeding (WMD: -11.05; 95% CI: -19.05 to -3.05; p = 0.007). We noted substantial heterogeneity across the included trials in the percentage of total nutrition delivere the full nutritional target (I² = 85.1%; p < 0.001), and the time required to start feeding (I² = 90.0%; p < 0.001).

The breakdown of the number of trials reporting the effects of gastric tube versus postpyloric tube feeding on the length of ICU stay, length of hospital stay, and length of mechanical ventilation was 10, 7, and 8 trials, respectively (Figure 7). No significant differences between gastric tube and postpyloric tube feeding for length of ICU stay (WMD:

0.66; 95% CI: -0.96–2.28; p = 0.423), length of hospital stay (WMD: 1.63; 95% CI: -0.65–3.91; p = 0.162), and length of mechanical ventilation (WMD: 1.01; 95% CI: -0.89–2.91; p = 0.296) were observed.

DISCUSSION

The use of an enteral route for EN can reduce gastric motility, which is responsible for limited caloric intake and is associated with an increased risk of aspiration pneumonia. Postpyloric tube feeding, which delivers feed to the duodenum or jejunum, could overcome these shortcomings. This comprehensive, quantitative study was performed to compare the efficacy and safety of gastric tubes with postpyloric tube feeding for critically ill patients, and a total of 1,329 critically ill patients across a broad range of patients' characteristics, especially disease status, from 16 RCTs were included. This study found that gastric tube feeding significantly increased the risk of pneumonia and the time required to start feeding compared with post-pyloric tube feeding, whereas there were no significant differences between the groups in terms of mortality, abdominal distension, diarrhea, vomiting, bacteremia, constipation, gastrointestinal bleeding, high gastric residual volume, pulmonary aspiration, percentage of total nutrition delivered to the participant, time required to achieve the full nutritional target, length of ICU stay, length of hospital stay, and length of mechanical ventilation.

Our study reported similar effects of gastric tube and postpyloric tube feeding on the risk of mortality, which is consistent with prior meta-analyses.¹³⁻¹⁵ Moreover, the results of the sensitivity and subgroup analyses indicated no significant difference between gastric and postpyloric tube feeding on the risk of mortality, and all included trials reported similar conclusions. The potential reason for this could be that these trials were designed with nutritional and safety outcomes as the primary outcomes, and the sample size was not sufficient to detect potential differences in the risk of mortality between groups.

Our study found that gastric tube feeding was associated with an increased risk of pneumonia compared to postpyloric tube feeding, which was consistent with previous metaanalyses.^{14,15} Studies have illustrated that inhibited gastrointestinal motility, reduced gastric emptying, a pressure drop at the gastroesophageal junction, and abnormal esophageal motility are significantly associated with the progression of pneumonia.^{42,43} The end of the tube in postpyloric tube feeding was placed post-pylorus, which was associated with reduced gastric residual volume and inhibition of gastrointestinal peristalsis. Moreover, postpyloric tube feeding can prevent nutrients from flowing back into the stomach, thereby reducing the risk of aspiration. Furthermore, subgroup analyses found an increased risk of pneumonia in patients receiving gastric tube feeding, when patients' age was ≥ 55.0 years, the proportion of male participants was $\geq 70.0\%$, and used duodenal tube feeding as control. These results suggest that differences are mainly observed in patients at high risk for pneumonia.

Our study did not find significant differences between gastric tube and postpyloric tube feeding for the risk of gastrointestinal complications, which is inconsistent with a previous study.¹⁵ The potential reasons for these differences could be explained by the following: (1) the incidence of gastrointestinal complications could be affected by the dosage, type, and dropping rate of the nutrient solution; (2) the disease status varied between gastric tube and postpyloric tube feeding and could be affected by the progression of gastrointestinal complications; and (3) the incidence of most specific gastrointestinal complications was lower, and a smaller number of trials reported these outcomes; thus, the power was not sufficient to detect potential differences between the groups. In addition, although the use of postpyloric tube feeding could provide more nutrition and reduce the risk of complications, there were no significant differences between the groups in terms of the lengths of ICU stay, hospital stay, and mechanical ventilation.

This study has several limitations. First, most of the included trials reported a high risk of bias for blinding of participants, personnel, and outcome assessment. Second, substantial heterogeneity was observed for several outcomes, particularly the continuous outcomes. Third, the severity of critically illness and disease status differed across the included trials, which could have affected the prognosis of critically ill patients. Fourth, meta-analyses based on published articles have inherent limitations, including inevitable publication bias and restricted detailed analyses.

In conclusions, our study found that gastric tube feeding required a shorter time to start feeding than postpyloric tube feeding but related to a significantly increased risk of pneumonia in critically ill patients. Therefore, post-pyloric tube feeding should be applied for critically ill patients to prevent the risk of pneumonia in clinical practice.

CONFLICT OF INTEREST AND FUNDING DISCLOSURE

The authors declare no competing interests.

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Country	Sample size	Age (years)	Male (%)	Disease status	APACHE	Intervention	Control
USA	38 (19/19)	47.7 (44.8/50.5)	60.5	Critical patients	II: 16.9	Gastric	Jejunal
Canada	80 (43/37)	34.2 (34.7/33.6)	77.5	Ventilated blunt trauma	II: 18.0	Gastric	Duodenal
USA	44 (23/21)	54.4 (49.0/54.0)	68.2	Critical patients	II: 21.0	Gastric	Small intestinal
USA	25 (11/14)	56.7 (60.6/53.6)	56.0	Neurological disease	III: 47.7	Gastric	Duodenal
Spain	101 (51/50)	58.0 (59.0/57.0)	70.3	Critical patients	II: 18.0	Gastric	Jejunal
Australia	73 (39/34)	54.5 (53.5/55.7)	68.5	Critical patients	II: 18.2	Gastric	Jejunal
USA	60 (30/30)	58.9 (58.1/59.6)	50.0	Critical patients	NA	Gastric	Small intestinal
Scotland	49 (27/22)	60.8 (64.0/58.0)	53.1	Severe acute pancreatitis	NA	Gastric	Jejunal
Australia	104 (54/50)	52.1 (54.0/50.0)	50.0	Critical patients	II: 27.1	Gastric	Small intestinal
China	121 (62/59)	68.9 (62/59)	70.2	Critical patients	II: 20.4	Gastric	Duodenal
Australia	180 (89/91)	52.5 (54.0/51.0)	73.9	Critical patients	II: 20.0	Gastric	Jejunal
India	78 (39/39)	39.4 (39.1/39.7)	67.9	Severe acute pancreatitis	II: 8.3	Gastric	Jejunal
Brazil	115 (61/54)	61.4 (60.0/63.0)	48.7	Critical patients	II: 22.0	Gastric	Jejunal
China	70 (35/35)	52.4 (52.0/52.7)	68.6	Critical patients	NA	Gastric	Jejunal
UK	50 (25/25)	52.0 (51.0/53.0)	76.0	Critical patients	II: 19.0	Gastric	Small intestinal
China	141 (71/70)	82.0 (82.0/82.0)	62.4	Critical patients	II: 27.9	Gastric	Small intestinal
	USA Canada USA Spain Australia USA Scotland Australia China Australia India Brazil China UK	USA 38 (19/19) Canada 80 (43/37) USA 44 (23/21) USA 25 (11/14) Spain 101 (51/50) Australia 73 (39/34) USA 60 (30/30) Scotland 49 (27/22) Australia 104 (54/50) China 121 (62/59) Australia 180 (89/91) India 78 (39/39) Brazil 115 (61/54) China 70 (35/35) UK 50 (25/25)	USA 38 (19/19) 47.7 (44.8/50.5) Canada 80 (43/37) 34.2 (34.7/33.6) USA 44 (23/21) 54.4 (49.0/54.0) USA 25 (11/14) 56.7 (60.6/53.6) Spain 101 (51/50) 58.0 (59.0/57.0) Australia 73 (39/34) 54.5 (53.5/55.7) USA 60 (30/30) 58.9 (58.1/59.6) Scotland 49 (27/22) 60.8 (64.0/58.0) Australia 104 (54/50) 52.1 (54.0/50.0) China 121 (62/59) 68.9 (62/59) Australia 180 (89/91) 52.5 (54.0/51.0) India 78 (39/39) 39.4 (39.1/39.7) Brazil 115 (61/54) 61.4 (60.0/63.0) China 70 (35/35) 52.4 (52.0/52.7) UK 50 (25/25) 52.0 (51.0/53.0)	USA38 (19/19)47.7 (44.8/50.5)60.5Canada80 (43/37) 34.2 (34.7/33.6)77.5USA44 (23/21) 54.4 (49.0/54.0)68.2USA25 (11/14) 56.7 (60.6/53.6) 56.0 Spain101 (51/50) 58.0 (59.0/57.0)70.3Australia73 (39/34) 54.5 (53.5/55.7)68.5USA60 (30/30) 58.9 (58.1/59.6)50.0Scotland49 (27/22)60.8 (64.0/58.0)53.1Australia104 (54/50)52.1 (54.0/50.0)50.0China121 (62/59)68.9 (62/59)70.2Australia180 (89/91)52.5 (54.0/51.0)73.9India78 (39/39)39.4 (39.1/39.7)67.9Brazil115 (61/54)61.4 (60.0/63.0)48.7China70 (35/35)52.4 (52.0/52.7)68.6UK50 (25/25)52.0 (51.0/53.0)76.0	USA38 (19/19)47.7 (44.8/50.5)60.5Critical patientsCanada80 (43/37)34.2 (34.7/33.6)77.5Ventilated blunt traumaUSA44 (23/21)54.4 (49.0/54.0)68.2Critical patientsUSA25 (11/14)56.7 (60.6/53.6)56.0Neurological diseaseSpain101 (51/50)58.0 (59.0/57.0)70.3Critical patientsAustralia73 (39/34)54.5 (53.5/55.7)68.5Critical patientsUSA60 (30/30)58.9 (58.1/59.6)50.0Critical patientsScotland49 (27/22)60.8 (64.0/58.0)53.1Severe acute pancreatitisAustralia104 (54/50)52.1 (54.0/50.0)50.0Critical patientsAustralia104 (54/50)52.1 (54.0/51.0)73.9Critical patientsAustralia180 (89/91)52.5 (54.0/51.0)73.9Critical patientsIndia78 (39/39)39.4 (39.1/39.7)67.9Severe acute pancreatitisBrazil115 (61/54)61.4 (60.0/63.0)48.7Critical patientsUK50 (25/25)52.0 (51.0/53.0)76.0Critical patients	USA38 (19/19)47.7 (44.8/50.5)60.5Critical patientsII: 16.9Canada80 (43/37)34.2 (34.7/33.6)77.5Ventilated blunt traumaII: 18.0USA44 (23/21)54.4 (49.0/54.0)68.2Critical patientsII: 21.0USA25 (11/14)56.7 (60.6/53.6)56.0Neurological diseaseIII: 47.7Spain101 (51/50)58.0 (59.0/57.0)70.3Critical patientsII: 18.0Australia73 (39/34)54.5 (53.5/55.7)68.5Critical patientsII: 18.2USA60 (30/30)58.9 (58.1/59.6)50.0Critical patientsNAScotland49 (27/22)60.8 (64.0/58.0)53.1Severe acute pancreatitisNAAustralia104 (54/50)52.1 (54.0/50.0)50.0Critical patientsII: 20.4Australia180 (89/91)52.5 (54.0/51.0)73.9Critical patientsII: 20.4Australia180 (89/91)52.5 (54.0/51.0)73.9Critical patientsII: 20.0India78 (39/39)39.4 (39.1/39.7)67.9Severe acute pancreatitisII: 8.3Brazil115 (61/54)61.4 (60.0/63.0)48.7Critical patientsII: 22.0China70 (35/35)52.4 (52.0/52.7)68.6Critical patientsII: 22.0China70 (35/35)52.4 (52.0/52.7)68.6Critical patientsII: 19.0	USA38 (19/19)47.7 (44.8/50.5)60.5Critical patientsII: 16.9GastricCanada80 (43/37)34.2 (34.7/33.6)77.5Ventilated blunt traumaII: 18.0GastricUSA44 (23/21)54.4 (49.0/54.0)68.2Critical patientsII: 21.0GastricUSA25 (11/14)56.7 (60.6/53.6)56.0Neurological diseaseIII: 47.7GastricSpain101 (51/50)58.0 (59.0/57.0)70.3Critical patientsII: 18.0GastricAustralia73 (39/34)54.5 (53.5/55.7)68.5Critical patientsII: 18.2GastricUSA60 (30/30)58.9 (58.1/59.6)50.0Critical patientsII: 18.2GastricScotland49 (27/22)60.8 (64.0/58.0)53.1Severe acute pancreatitisNAGastricAustralia104 (54/50)52.1 (54.0/50.0)50.0Critical patientsII: 27.1GastricAustralia104 (54/50)52.1 (54.0/51.0)73.9Critical patientsII: 20.4GastricAustralia180 (89/91)52.5 (54.0/51.0)73.9Critical patientsII: 20.0GastricIndia78 (39/39)39.4 (39.1/39.7)67.9Severe acute pancreatitisII: 8.3GastricIndia70 (35/35)52.4 (52.0/52.7)68.6Critical patientsII: 22.0GastricUK50 (25/25)52.0 (51.0/53.0)76.0Critical patientsII: 19.0Gastric

Table 1. The baseline characteristics of included studies and involved patients

Study	Enteral feeding protocol	Follow-up duration
Montecalvo 1992 ²⁶	Began at 25 mL/h/d for the first 24 h and then were increased by 24 mL/h/d until the protein/caloric intake goals were reached	42.0 days
Kortbeek 199927	Starting at 25 mL/h and increasing the rate by 25 mL/h every 4 h until the volume required to meet caloric support was achieved	28.0 days
Kearns 2000 ²⁸	Infusion was stopped for residuals >150 mL, and once the residual was < 150 mL, feeding resumed	42.0 days
Day 2001 ²⁹	The Harris Benedict equation with activity and stress factors were used to calculate the total energy and protein requirement	10.0 days
Montejo 2002 ³⁰	Feedings were started in the first 36 h after admission and delivered continuously to achieve half of the estimated caloric needs in 24 h	16.0 days
Davies 2002 ³¹	At a rate of 20 mL/hr and increased by 20 mL/h every 4 hrs until the target nutrition rate was reached	12.0 days
Neumann 2002 ³²	Starting at 30 mL/h, then advanced to a patient-specific goal rate by 10 mL/h every 6 h	14.0 days
Eatock 2005 ³³	Rate of 30 mL/h increasing to 100 mL/h over 24-48 h. The caloric target was 2,000 kcal per day	16.0 days
White 200934	Enteral feeds were commenced at 40 mL/h. The nasogastric tube was aspirated every 4 h. If the gastric residual was less than 200 mL	5.0 days
	after 4 h, the rate was increased to the recommended target rate	
Hsu 2009 ³⁵	starting at 20 mL/h. The rate was increased by 20 mL/h every 4 h until the patient's goal rate was achieved	34.0 days
Davies 2012 ³⁶	The initial commencement rate and advancement rate toward the hourly target were determined by each hospital's standard practice, but	22.0 days
	the aim was to meet estimated energy requirements as soon as possible by following a locally developed evidence-based algorithm	·
Singh 2012 ³⁷	Nutrient goal (25 kcal/kg per day) in 3 to 4 days	18.0 days
Friedman 201538	The individual energy needs and the formulation of enteral nutrition were determined by clinical staff (doctors and nutritionists)	28.0 days
Wan 2015 ³⁹	Rate of 30 mL/h increasing to 100 mL/h over 24-72 h, the caloric target was set at 25 kcal/kg of ideal bodyweight/day for women and	14.0 days
	30 kcal/kg of ideal bodyweight/day for men	·
Taylor 2016 ⁴⁰	Increased from 40 mL feed/h or current rate to full rate whenever tolerated	5.0 days
Zhu 201841	Energy goals were set at 25 kcal per kg of ideal body weight per day, and the protein target was 1.2-2.0 g per kg of ideal body weight	7.0 days
	per day	

APACHE II: Acute Physiology and Chronic Health Evaluation II; NA: not available

Outcomes, factors and subgroup	No of trials	RR and 95%CI	p value	I^{2} (%)	Q statistic	<i>p</i> value between subgroups
Mortality						
Country						0.608
Eastern	3	1.02 (0.79-1.31)	0.903	7.8	0.338	
Western	10	0.95 (0.74-1.21)	0.668	0.0	0.887	
Age (years)						0.320
≥ 55.0	5	1.03 (0.85-1.25)	0.761	0.0	0.633	
< 55.0	8	0.85 (0.58-1.23)	0.376	0.0	0.864	
Male (%)						1.000
≥ 70.0	5	0.99 (0.75-1.30)	0.916	0.0	0.954	
< 70.0	8	0.99 (0.80-1.23)	0.928	0.0	0.511	
Postpyloric tube						0.809
Duodenal	2	0.90 (0.60-1.35)	0.604	0.0	0.723	
Jejunal	7	0.96 (0.73-1.25)	0.765	0.0	0.890	
Small intestinal	4	0.97 (0.64-1.47)	0.875	23.6	0.270	
Pneumonia						5
Country					$-(Z_{\Lambda})$	0.391
Eastern	4	2.07 (0.84-5.06)	0.113	68.8	0.022	
Western	8	1.28 (0.94-1.75)	0.113	0.0	0.800	
Age (years)				a z		0.195
≥ 55.0	4	1.83 (1.11-3.02)	0.017	25.1	0.261	
< 55.0	8	1.23 (0.82-1.85)	0.311	17.6	0.291	
Male (%)					1	1.000
≥ 70.0	5	1.39 (1.02-1.89)	0.036	0.0	0.474	
< 70.0	7	1.70 (0.79-3.64)	0.173	45.3	0.089	
Postpyloric tube				\sim		0.190
Duodenal	3	1.94 (1.16-3.27)	0.012	0.0	0.407	
Jejunal	6	1.19 (0.76-1.86)	0.450	27.3	0.230	
Small intestinal	3	1.69 (0.87-3.28)	0.125	8.4	0.336	

Table 2. Subgroup analyses for mortality and pneumonia

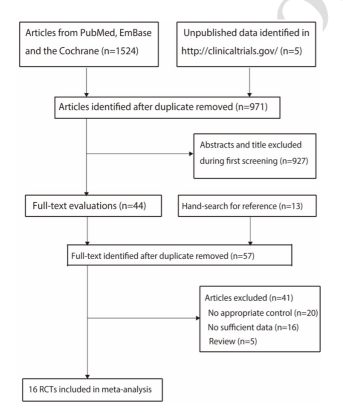
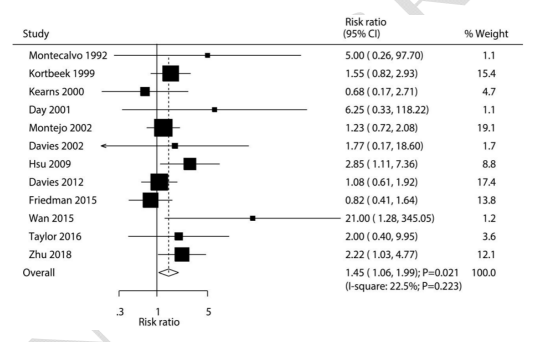
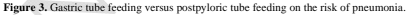


Figure 1. Details of the literature search and trial selection processes.

Study		Risk ratio (95% Cl)	% Weight
Montecalvo 1992		1.00 (0.35, 2.90)	2.6
Kortbeek 1999		1.15 (0.27, 4.80)	1.4
Kearns 2000		- 1.10 (0.39, 3.07)	2.7
Montejo 2002		1.14 (0.71, 1.82)	12.9
Davies 2002		1.09 (0.32, 3.73)	1.9
Eatock 2005		0.58 (0.21, 1.58)	2.9
White 2009 🖌 🛶	_	0.42 (0.16, 1.13)	3.0
Hsu 2009		0.88 (0.57, 1.34)	16.1
Davies 2012		0.94 (0.46, 1.95)	5.5
Singh 2012 —		0.57 (0.18, 1.80)	2.2
Friedman 2015		0.97 (0.60, 1.58)	12.5
Taylor 2016		— 1.00 (0.28, 3.56)	1.8
Zhu 2018		1.15 (0.86, 1.53)	34.5
Overall		0.99 (0.83, 1.17); P=0.891 (I-square: 0.0%; P=0.872)	100.0
	.3 1 Risk ratio	5	

Figure 2. Gastric tube feeding versus postpyloric tube feeding on the risk of mortality.





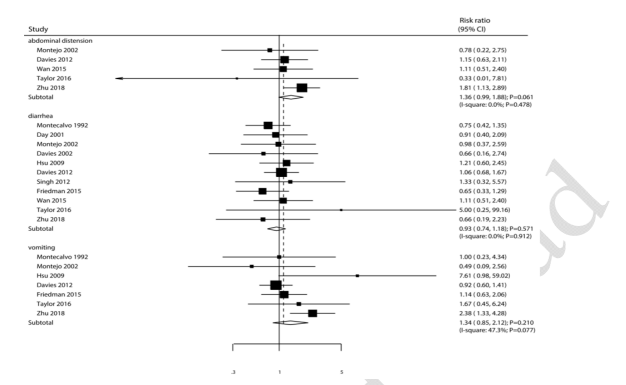


Figure 4. Gastric tube feeding versus postpyloric tube feeding on the risk of abdominal distension, diarrhea, and vomiting.

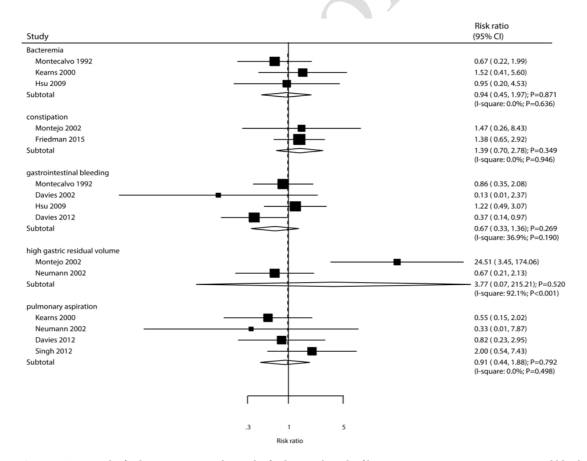


Figure 5. Gastric tube feeding versus postpyloric tube feeding on the risk of bacteremia, constipation, gastrointestinal bleeding, high gastric residual volume, and pulmonary aspiration

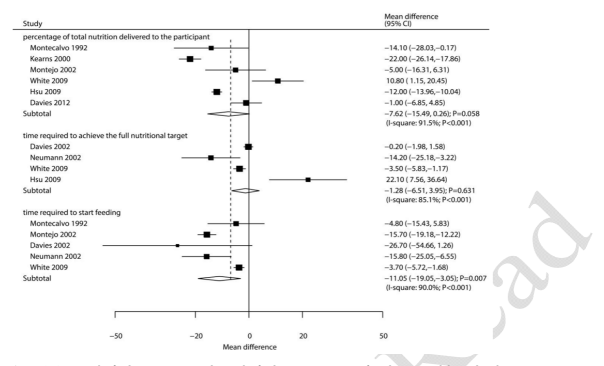


Figure 6. Gastric tube feeding versus postpyloric tube feeding on percentage of total nutrition delivered to the participant, time required to achieve the full nutritional target, and time required to start feeding

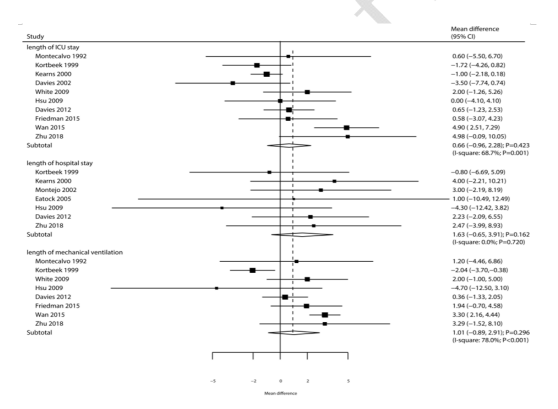


Figure 7. Gastric tube feeding versus postpyloric tube feeding on length of ICU stay, length of hospital stays, and length of mechanical ventilation. ICU, intensive care unit

Supplementary Tables and Figures

Supplementary Table 1. Search strategy in PubMed, EmBase, and the Cochrane Library

Search strategy in PubMed #1: (enteral nutrition OR duodenostomy OR gastrostomy OR jejunostomy OR intubation, gastrointestinal) [MeSH] #2: (duodenostom* OR gastrostom* OR PEJ OR PEG OR jejunostom* OR jtube* OR g-tube* OR ng-tube* OR njtube*):[ab,ti,kw] OR ((nutrition* OR feed* OR fed OR tube* OR intub*) #3: #1 OR #2 #4: (nasogastr* OR duoden* OR gastr* OR nasoduoden* OR jejun* OR nasojejun* OR post-pylor* OR bowel* OR trans-pylor* OR intestine* OR gavage OR orogastric OR stomach OR nasoenter*):[ab, ti, kw]). #5: #3 AND #4 #6: (intensive care OR critical care OR critical illness OR pneumonia OR burn OR respiratory failure OR craniocerebral trauma OR burns OR pancreatitis) #7: (intensive care OR ICU OR critical* ill* OR critical patients OR critical* care OR pneumonia OR burn OR pancreatitis OR trauma OR injur*):[ab, ti, kw]. #8: #6 AND #7 #9: #5 AND #8 Search strategy in EmBase 1. enteric feeding/ or artificial feeding/ or nose feeding/ or nasogastric tube/ or stomach tube/ or stomach intubation/ or gastrostomy/ or percutaneous endoscopic gastrostomy/ or duodenum intubation/ or duodenostomy/ or jejunostomy/ or (gtube* or ng-tube* or gastrostom* or PEG or duodenostom* or jejunostom* or PEJ or j-tube* or nj-tube*).ti,ab. or ((nutrition* or fed or feed* or tube* or intub*) adj5 (gastr* or nasogastr* or stomach or duoden* or nasoduoden* or jejun* or nasojejun* or bowel* or intestine* or post?pylor* or trans? pylor* or nasoenter* or orogastric or gavage)).ab,ti. 2. exp pancreatitis/ or injury/ or burn/ or "head and neck injury"/ or multiple trauma/ or critical illness/ or intensive care/ or intensive care unit/ or pneumonia/ or aspiration pneumonia/ or (pneumonia* or critical* ill* or critical* care or intensive care or ICU or burn* or trauma* or head injur* or pancreatitis).ab,ti. 3.1 and 2. 4. (infant* or child* or adolescent*).af. 5. (adult* or aged).af. 6. 3 not (4 not (5 and 4)) 7. (placebo.sh. or controlled study.ab. or random*.ti,ab. or trial*.ti,ab.) not (animals not (humans and animals)).sh. 8.6 and 7 Search strategy in Cochrane library #1 MeSH descriptor Enteral Nutrition explode all trees #2 MeSH descriptor Gastrostomy explode all trees #3 MeSH descriptor Duodenostomy explode all trees #4 MeSH descriptor Jejunostomy explode all trees #5 MeSH descriptor Intubation, Gastrointestinal explode all trees #6 (gastrostom* or duodenostom* or jejunostom* or PEG or g-tube* or ng-tube* or j-tube* or nj-tube* or PEJ):ab,ti #7 ((nutrition* or fed or feed* or tube* or intub*) near (gastr* or nasogastr* or stomach or duoden* or nasoduoden* or jejun* or nasojejun* or bowel* or intestine* or post?pylor* or trans?pylor* or nasoenter* or orogastric or gavage)):ab,ti #8 (#1 OR #2 OR #3 OR #4 OR #5 OR #6 OR #7) #9 MeSH descriptor Pneumonia explode all trees #10 MeSH descriptor Pneumonia, Aspiration explode all trees #11 MeSH descriptor Intensive Care Units explode all trees #12 MeSH descriptor Burn Units explode all trees #13 MeSH descriptor Respiratory Care Units explode all trees #14 MeSH descriptor Critical Care explode all trees #15 MeSH descriptor Intensive Care explode all trees #16 MeSH descriptor Critical Illness explode all trees #17 MeSH descriptor Craniocerebral Trauma explode all trees #18 MeSH descriptor Burns explode all trees #19 MeSH descriptor Wounds and Injuries explode all trees #20 MeSH descriptor Pancreatitis explode all trees #21 (pneumonia* or critical* ill* or critical* care or intensive care or ICU or burn* or trauma* or head injur* or pancreatitis):ti,ab #22 (#9 OR #10 OR #11 OR #12 OR #13 OR #14 OR #15 OR #16 OR #17 OR #18 OR #19 OR #20 OR #21) #23 (#8 AND #22) #24 (infant* or child* or adolescent*) #25 (adult* or aged) #26 (#24 AND NOT (#25 AND #24)) #27 (#23 AND NOT #26)

Supplementary Table 2. The full bibliography list of the 41 excluded articles

Lack of appropriate controls

1. Landais M, Nay MA, Auchabie J, et al. Continued enteral nutrition until extubation compared with fasting before
extubation in patients in the intensive care unit: an open-label, cluster-randomised, parallel-group, non-inferiority trial.
Lancet Respir Med. 2023;11(4):319-328.

2. Kagan I, Cohen J, Bendavid I, et al. Effect of Combined Protein-Enriched Enteral Nutrition and Early Cycle Ergometry in Mechanically Ventilated Critically Ill Patients-A Pilot Study. Nutrients. 2022;14(8):1589.

4. Chinda P, Poomthong P, Toadithep P, et al. The implementation of a nutrition protocol in a surgical intensive care unit; a randomized controlled trial at a tertiary care hospital. PLoS One. 2020;15(4):e0231777.

5. Mahran G, Mahgoup A, Kamel EZ, et al. Effect of 2 Enteral Feeding Schedules on Intra-abdominal Pressure in Patients Receiving Mechanical Ventilation: A Randomized Controlled Trial. Crit Care Nurse. 2019;39(6):29-35.

6. Brown AM, Fisher E, Forbes ML. Bolus vs Continuous Nasogastric Feeds in Mechanically Ventilated Pediatric Patients: A Pilot Study. JPEN J Parenter Enteral Nutr. 2019;43(6):750-758.

7. Mistraletti G, Umbrello M, Salini S, et al. Enteral versus intravenous approach for the sedation of critically ill patients: a randomized and controlled trial. Crit Care. 2019;23(1):3.

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13. Chen YC, Chou SS, Lin LH, et al. The effect of intermittent nasogastric feeding on preventing aspiration pneumonia in ventilated critically ill patients. J Nurs Res. 2006;14(3):167-80.

14. Roy PM, Person B, Souday V, et al. Percutaneous radiologic gastrostomy versus nasogastric tube in critically ill patients. Clin Nutr. 2005;24(2):321-5.

15. Preiser JC, Peres-Bota D, Eisendrath P, et al. Gut mucosal and plasma concentrations of glutamine: a comparison between two enriched enteral feeding solutions in critically ill patients. Nutr J. 2003;2:13.

16. Schwab D, Mühldorfer S, Nusko G, et al. Endoscopic placement of nasojejunal tubes: a randomized, controlled, prospective trial comparing suitability and technical success for two different tubes. Gastrointest Endosc. 2002;56(6):858-63.

17. Boivin MA, Levy H. Gastric feeding with erythromycin is equivalent to transpyloric feeding in the critically ill. Crit Care Med. 2001;29(10):1916-9.

18. Yavagal DR, Karnad DR, Oak JL. Metoclopramide for preventing pneumonia in critically ill patients receiving enteral tube feeding: a randomized controlled trial. Crit Care Med. 2000;28(5):1408-11.

19. Yavagal DR, Karnad DR, Oak JL. Metoclopramide for preventing pneumonia in critically ill patients receiving enteral tube feeding: a randomized controlled trial. Crit Care Med. 2000;28(5):1408-11.

20. Gharpure V, Meert KL, Sarnaik AP, et al. Indicators of postpyloric feeding tube placement in children. Crit Care Med. 2000;28(8):2962-6

Insufficient data

1. Seifi N, Rezvani R, Sedaghat A, et al. The effects of synbiotic supplementation on enteral feeding tolerance, protein homeostasis, and muscle wasting of critically ill adult patients: a randomized controlled trial. Trials. 2022;23(1):846.

2. Deng LX, Lan-Cao, Zhang LN, et al. The effects of abdominal-based early progressive mobilisation on gastric motility in endotracheally intubated intensive care patients: A randomised controlled trial. Intensive Crit Care Nurs. 2022;71:103232.

3. Lew CCH, Lee ZY, Day AG, et al. Correlation between gastric residual volumes and markers of gastric emptying: A post hoc analysis of a randomized clinical trial. JPEN J Parenter Enteral Nutr. 2022;46(4):850-857.

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critically ill patients: a non-inferiority randomized controlled trial. Chin Med J (Engl). 2021;134(14):1695-1700. 5. Lv B, Hu L, Chen L, et al. Blind bedside postpyloric placement of spiral tube as rescue therapy in critically ill patients:

a prospective, tricentric, observational study. Crit Care. 2017;21(1):248.

6. Peake SL, Davies AR, Deane AM, et al. Use of a concentrated enteral nutrition solution to increase calorie delivery to critically ill patients: a randomized, double-blind, clinical trial. Am J Clin Nutr. 2014;100(2):616-25.

^{3.} Kagan I, Cohen J, Bendavid I, et al. Effect of Combined Protein-Enriched Enteral Nutrition and Early Cycle Ergometry

in Mechanically Ventilated Critically III Patients-A Pilot Study. Nutrients. 2022;14(8):1589.

Supplementary Table 2. The full bibliography list of the 41 excluded articles (cont.)

Insufficient data

7. Holzinger U, Brunner R, Miehsler W, et al. Jejunal tube placement in critically ill patients: A prospective, randomized trial comparing the endoscopic technique with the electromagnetically visualized method. Crit Care Med. 2011;39(1):73-7.

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Pinilla JC, Samphire J, Arnold C, et al. Comparison of gastrointestinal tolerance to two enteral feeding protocols in critically ill patients: a prospective, randomized controlled trial. JPEN J Parenter Enteral Nutr. 2001;25(2):81-6.
Booker KJ, Niedringhaus L, Eden B, et al. Comparison of 2 methods of managing gastric residual volumes from feeding tubes. Am J Crit Care. 2000;9(5):318-24.

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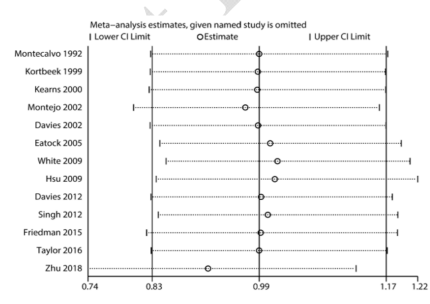
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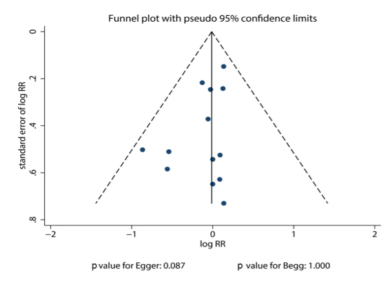
Study	Random sequence generation	Allocation concealment	Blinding of participants and personnel	Blinding of outcome assessment
Montecalvo 1992 ²⁶	Low	Unclear	High	Low
Kortbeek 199927	Low	Low	High	High
Kearns 2000 ²⁸	Low	Low	Low	Low
Day 2001 ²⁹	Low	Low	High	Low
Montejo 2002 ³⁰	Low	Unclear	High	High
Davies 2002 ³¹	Unclear	Low	High	High
Neumann 200232	Low	Low	High	Low
Eatock 200533	Low	Low	High	Unclear
White 2009 ³⁴	Low	Low	High	High
Hsu 200935	Low	Low	Low	Low
Davies 2012 ³⁶	Low	Low	High	High
Singh 201237	Low	Low	High	Low
Friedman 201538	Unclear	Low	Unclear	Unclear
Wan 2015 ³⁹	Low	Low	High	Unclear
Taylor 2016 ⁴⁰	Low	Low	High	High
Zhu 201841	Low	Low	High	Low

Supplementary Table 3. Quality assessment of included trials

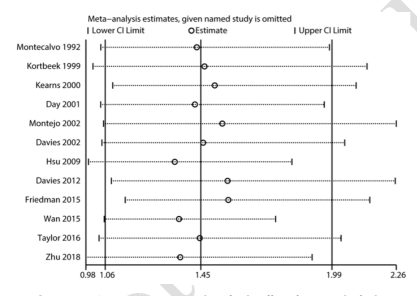
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Study	Incomplete outcome data	Selective reporting	Other bias
Montecalvo 1992 ²⁶	Low	Low	Unclear
Kortbeek 1999 ²⁷	Low	Low	Low
Kearns 2000 ²⁸	Low	Low	Low
Day 2001 ²⁹	Low	Low	Low
Montejo 2002 ³⁰	Low	Low	Low
Davies 2002 ³¹	Low	Low	Low
Neumann 2002 ³²	Low	Low	Low
Eatock 200533	High	Low	Low
White 2009 ³⁴	Low	Low	Unclear
Hsu 2009 ³⁵	Low	Low	Low
Davies 2012 ³⁶	Low	Low	Low
Singh 2012 ³⁷	Low	Low	Low
Friedman 201538	Low	Low	Unclear
Wan 2015 ³⁹	Low	Low	Low
Taylor 2016 ⁴⁰	Low	Low	Low
Zhu 201841	Low	Low	Unclear

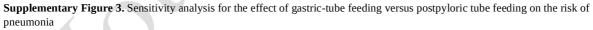


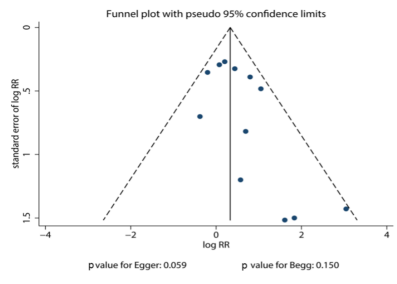
Supplementary Figure 1. Sensitivity analysis for the effect of gastric-tube feeding versus postpyloric tube feeding on the risk of mortality



Supplementary Figure 2. Funnel plot for the effect of gastric-tube feeding versus postpyloric tube feeding on the risk of mortality







Supplementary Figure 4. Funnel plot for the effect of gastric-tube feeding versus postpyloric tube feeding on the risk of pneumonia

23