# 6.2 Enteral Nutrition (Other): Probiotics

## May 2015

2015 Recommendation: Based on 4 level 1 studies and 24 level 2 studies, the use of probiotics should be considered in critically ill patients.

**2015 Discussion:** The committee noted that with the addition of the data from 5 new studies (Cui 2013, Tan 2013, Wang 2013, Lopez de Toro 2014 and Sanaie 2014), the reduction of infections was stronger and the trend towards a reduction in VAP still remained, especially in patients with high mortality risk. These estimates were still found to be sensitive to the quality of the primary trials. This reduction in infections disappeared when only high quality studies were considered. The signal for a trend towards a reduction in ICU LOS was accompanied by significant statistical heterogeneity. The committee agreed that the earlier harm seen with the use of probiotics in the PROPATRIA study was not evident in the remaining 27 studies. Based on this, the committee agreed that the weak recommendation for the use of probiotics was still appropriate although no recommendation could be made for the dose or a particular type of probiotic, with the exception of Saccharomyces boulardii which should not be used as it is considered unsafe in ICU patients (1).

(1) Lherm T, Monet C, Nougiere B, Soulier M, Larbi D, Le Gall C, Caen D, Malbrunot C. Seven cases of fungemia with Saccharomyces boulardii in critically ill patients. Intensive Care Med. 2002 Jun;28(6):797-801.

2013 Recommendation: Based on 3 level 1 and 20 level 2 studies, the use of probiotics should be considered in critically ill patients.

**2013 Discussion:** The committee noted the trend towards a reduction in VAP with the use of probiotics and the modest treatment effect of reducing overall infections, especially in patients with high mortality risk. However, these estimates of effect are sensitive to the quality of the primary trials. This reduction in infections disappeared when only high quality studies were considered. The committee agreed that the interpretation of the earlier PROPATRIA trial, which showed increased harm with the use of probiotics, was confounded by the concomitant use of fiber and jejunal feeding. With the exception of Saccharomyces boulardii, a recent mega-synthesis showed that probiotics are not associated with increased risk (1). Based on this, the committee agreed to make a weak recommendation for their use, however, no recommendation for dose or a particular type of probiotic could be made with the exception of Saccharomyces boulardii which should not be used as it is considered unsafe in ICU patients (2) .

(1) Agency for Health Care Research and Quality, US Department of Health and Human Services. Safety of Probiotics Used to Reduce Risk and Prevent or Treat Disease April 2011

(2) Lherm T, Monet C, Nougiere B, Soulier M, Larbi D, Le Gall C, Caen D, Malbrunot C. Seven cases of fungemia with Saccharomyces boulardii in critically ill patients. Intensive Care Med. 2002 Jun;28(6):797-801.

| Semi Quantitative | Scoring |
|-------------------|---------|
|-------------------|---------|

| Values                            | Definition  | 2013 Score  | 2015 Score  |
|-----------------------------------|---|---|---|
| Effect size                       | Magnitude of the absolute risk reduction attributable to the intervention listeda higher score indicates a larger effect size   | Infections<br>1 (overall)<br>0 (for high quality) | Infections<br>1 (overall)<br>0 (for high quality)<br>2 (high mortality<br>RCTS) |
| Confidence<br>interval            | 95% confidence interval around the point estimate of the absolute risk reduction, or the pooled estimate (if more than one trial)a higher score indicates a smaller confidence interval   | 2   | 3   |
| Validity                          | Refers to internal validity of the study (or studies) as measured by the presence of concealed randomization,<br>blinded outcome adjudication, an intention to treat analysis, and an explicit definition of outcomesa higher<br>score indicates presence of more of these features in the trials appraised | 2   | 2   |
| Homogeneity or<br>Reproducibility | Similar direction of findings among trialsa higher score indicates greater similarity of direction of findings among trials   | 1   | 1   |
| Adequacy of<br>control group      | Extent to which the control group represented standard of care (large dissimilarities = 1, minor dissimilarities=2, usual care=3)   | 1   | 1   |
| Biological<br>plausibility        | Consistent with understanding of mechanistic and previous clinical work (large inconsistencies =1, minimal inconsistencies =2, very consistent =3)  | 3   | 3   |
| Generalizability                  | Likelihood of trial findings being replicated in other settings (low likelihood i.e. single centre =1, moderate likelihood i.e. multicentre with limited patient population or practice setting =2, high likelihood i.e. multicentre, heterogenous patients, diverse practice settings =3.                  | 2   | 2   |
| Low cost                          | Estimated cost of implementing the intervention listeda higher score indicates a lower cost to implement the intervention in an average ICU   | 2   | 2   |
| Feasible                          | Ease of implementing the intervention listeda higher score indicates greater ease of implementing the intervention in an average ICU  | 3   | 3   |
| Safety                            | Estimated probability of avoiding any significant harm that may be associated with the intervention listeda higher score indicates a lower probability of harm  | 2   | 2   |

## 6.2 Enteral Nutrition (Other): Probiotics

### Question: Does the addition of probiotics to enteral feeding result in better outcomes in critically ill patients?

Summary of evidence: There were 4 level 1 and 24 level 2 studies that were reviewed. Of the 28 included trials, 17 enrolled heterogeneous critically ill (medical and surgical) ICU patients (Spinder 2008, Barraud 2010, Frohmader 2010, Morrow 2010, Ferrie 2011, Tempe 1983, Heimburger 1994, Bleichner 1997, Kecskes 2003, Jain 2004, Klarin 2005, McNaught 2005. Forestier 2008, Klarin 2008, Knight 2008, Lopez de Toro 2014, Sanaie 2014), 6 enrolled patients with acute pancreatitis (Besselink 2008, DerSimonian 1986, Li 2007, Olah 2007, Cui 2013, Wang 2013), 1 enrolled trauma patients (Kotzampassi 2006), 1 enrolled head injury patients (Tan 2011, Tan 2013) and 2 enrolled burn patients (Schlotterer 1987, Lu 2004). Three trials studied the effects of the addition of *saccharomyces boulardii* to enteral nutrition, four studied the effects of Lactobacillus plantarum, three studied the effects of Lactobacillus rhamnosus, three studied the effects of VSL #3, one studied the effects of Trevis ™ (combination of probiotics+ prebiotics), four studied the effects of Synbiotic 2000 (combination of probiotics and prebiotics), one studied Ecologic 641 (probiotics) plus prebiotics (Besselink 2008), and nine studies used probiotics of varying strains . In one study, synbiotics were compared to a prebiotic (vs. placebo/conventional therapy), hence the data from this trial was not included in the meta-analysis (Olah 2007). Bleichner only reported on diarrhea while the other studies reported on clinical outcomes. In most of the studies patients received either enteral or parenteral nutrition, but no further details were provided.

**Mortality:** Probiotics had no effect on hospital mortality when the data from 16 trials were pooled (RR 0.98, 95% CI 0.8, 1.18, p=0.80, heterogeneity  $I^2=0\%$ ; figure 1) and no effect on ICU mortality pooling results from 7 trials (RR0.84, 95% CI 0.64, 1.1, p=0.21, heterogeneity  $I^2=0\%$ ; figure 2).

**Overall infections and VAP:** Infectious complications were reported in 12 trials. Pooled results show that probiotics were associated with a significant reduction in infectious complications (RR 0.82, 95% CI 0.69, 0.97, p=0.02, heterogeneity  $l^2=41\%$ ; figure 3). When the data from the 6 trials reporting VAP were pooled, probiotics were associated with a trend towards a decrease in the incidence of VAP (RR 0.74, 95% CI 0.55, 1.01, p=0.06, heterogeneity  $l^2=45\%$ ; figure 4).

**Subgroup analyses:** Several subgroup analyses were done to elucidate the effects of probiotics on infections (see figure 5). The details are as follows:

**Dose of probiotics:** Subgroup analyses showed similar rates of infectious complications in trials using high dose probiotics ( $\geq$ 5 x 10<sup>9</sup> CFU/day) (0.87, 95% CI 0.72, 1.06, p = 0.18) as those using a lower dose (<5 x 10<sup>9</sup> CFU/day) (RR 0.40, 95% CI 0.11, 1.50, p=0.18; p-value for the difference between groups: p=0.25).

*Lactobacillus plantarum:* Subgroup analyses showed that *L. plantarum*, either alone or in combination with other probiotics, was associated with a significant reduction in overall infections (RR 0.70, 95% CI 0.50, 0.97, p=0.03). However, this was not significantly different from the aggregated results of trials of that did not include *L. plantarum* (RR 0.88, 95% CI 0.72, 1.09, p=0.25; p-value for the difference between groups: p=0.23).

*Lactobacillus rhamnosus* GG: Subgroup analyses showed that effect of trials using LGG was not different from trials that did not include LGG (RR 0.86, 95% CI 0.67, 1.10 compared to RR 0.76, 95% CI 0.58, 1.01; p-value for the difference between groups: p=0.53). Higher mortality: The median mortality rate (hospital mortality or ICU mortality if hospital not reported) in the control groups of all studies was 15%. Subgroup analyses showed that probiotics were associated with a significant reduction in overall infections among patients with higher risk of death (>15% mortality in the control group) (RR 0.75, 95 % CI 0.57, 0.98, p=0.03). There was no significant effect in overall infections observed for trials of patients with a lower mortality ( $\leq$ 15% mortality) in the control group (RR 0.88, 95% CI 0.66, 1.18, p=0.40) and the test of subgroup differences was not significant (p-value for the difference between groups: p=0.41).

**Methodological score**: The median method score was 10. We compared trials with a methods score of less than 10 with those with a score of 10 or more. Trials with a higher score showed no effect on infection (RR 0.93, 95% CI 0.76, 1.15, p=0.51), whereas trials with a lower methods score showed a significant reduction in infectious complications (RR 0.70, 95% CI 0.58, 0.85, p=0.0003, p-value for the difference between groups: p=0.05).

**Length of Stay:** Probiotics had no impact on hospital LOS when data from 12 trials were pooled (WMD -1.23, 95% CI -4.21, 1.74, p=0.42, heterogeneity I<sup>2</sup>=66%; figure not shown). There was a trend towards a decrease in ICU LOS when results of 14 trials were pooled (WMD -3.26, 95% CI -7.82, 1.31, p=0.16, heterogeneity I<sup>2</sup>=93%; figure 6).

**Other:** The impact on diarrhea, reported variably as days of diarrhea, diarrhea rates and/or duration of diarrhea was reported in 12 trials. Pooling results from 8 trials that reported patients who developed diarrhea, probiotics had no effect (RR 0.95, 95% CI 0.80, 1.13, p=0.54; heterogeneity  $I^2=5\%$ ; figure 7). Data were too sparse to aggregate other reported individual infections (see table 1).

### Conclusions:

1) The addition of probiotics to enteral nutrition has no effect on hospital or ICU mortality.

2) The addition of probiotics to enteral nutrition is associated with a significant reduction in overall infectious complications and a trend towards a reduction in the incidence of VAP. This was seen only in the subgroup of lower quality studies.

3) The addition of probiotics to enteral nutrition had no effect on hospital length of stay or diarrhea, but is associated with a trend in reduction of ICU LOS.

Level 1 study: if all of the following are fulfilled: concealed randomization, blinded outcome adjudication and an intention to treat analysis. Level 2 study: If any one of the above characteristics are unfulfilled

|   | Study            | Population   | Methods Score   |  | Type of Probiotic/Interventio   | n   |
|---|------------------|--|---|--|---|---|
|   | Sludy            | Fopulation   | Methous Score   | Delivery Vehicle   | Intervention/Dose/Duration  | Control   |
| 1 | Tempe 1983       | ICU patients<br>N=40                                   | C.Random: yes<br>ITT: yes<br>Blinding: double<br>Score: 10<br>Viability (intervention): NR      | EN tube  | EN (unknown) + Ultra-Levure ( <i>Saccharomyces boulardii</i> ), 10 <sup>10/</sup> 1L solution for 11-21 days                                | EN (unknown) + Placebo<br>(sterile solution)              |
| 2 | Schlotterer 1987 | Burn patients<br>N=18                                  | C.Random: no<br>ITT: no<br>Blinding: double<br>Score: 8<br>Viability (intervention): NR         | NG tube  | EN (Polydiet or Nutrigil) + <i>Saccharomyces boulardi</i><br>500 mg QID for 8-28 days   | EN (Polydiet or Nutrigil) +<br>Placebo                    |
| 3 | Heimburger 1994  | Mixed ICU patients<br>83% received antibiotics<br>N=62 | C.Random: no<br>ITT: no<br>Blinding: double<br>Score: 9<br>Viability (intervention): NR         | EN tube  | EN (standard) + 1g of Lactinex ( <i>Lactobacillus acidophilus &amp; Lactobaccilus bulgaricus</i> ) 2 X 10 <sup>6</sup><br>TID for 5-10 days | EN (standard) + placebo<br>(0.5g dextrose + 0.5g lactose) |
| 4 | Bleichner 1997   | Mixed ICU patients<br>N=128                            | C.Random: not sure<br>ITT: yes<br>Blinding: double<br>Score: 13<br>Viability (intervention): NR | EN tube  | EN (unknown) + Saccharomyces boulardii<br>500 mg QID for 21 days or until EN stopped  | EN (unknown) + Placebo<br>(powder)                        |
| 5 | Kecskes 2003     | ICU patients on antibiotics<br>N=45                    | C.Random: no<br>ITT: no<br>Blinding: double<br>Score: 8<br>Viability (intervention): yes        | NJ tube  | NJ tube EN (Nutrison fibre) + fermented oatmeal formula kith <i>Lactobacillus plantarum</i> 299 10 ° BID and fibre for 7 days 2             |   |
| 6 | Jain 2004        | ICU patients<br>N=90                                   | C.Random: no<br>ITT: yes<br>Blinding: double<br>Score: 10<br>Viability (intervention): NR       | Oral or NG tube EN or PN + Trevis <sup>™</sup> 1 capsule TID + 7.5g<br>Raftilose (oligofructose)<br>BID until hospital discharge |   | EN or PN + Placebo<br>(powdered sucrose capsules)         |

 Table 1. Randomized studies evaluating Probiotics in critically ill patients

| 7  | Lu 2004          | Burn patients<br>N=40  | C.Random: no<br>ITT: yes<br>Blinding: double<br>Score: 9<br>Viability (intervention): NR | NR  | EN + synbiotics (4 types of probiotics & 4 types<br>of unspecified prebiotics) for 21 days   | EN + 4 types of prebiotics                                    |
|----|------------------|--|--|---|--|---|
| 8  | Klarin 2005      | Critically ill patients on<br>antibiotics<br>N=17  | C.Random: no<br>ITT: no<br>Blinding: no<br>Score: 6<br>Viability (intervention): NR      | Mixed in fermented<br>oatmeal, given via NG<br>tube | EN + <i>Lactobacillus plantarum</i> 299v, 10 <sup>9</sup> /day<br>50ml every 6 hours x 3 days then 25 ml every 6<br>hours until ICU discharge  | EN (Impact or Nutrodrip<br>Fibre). Some patients<br>needed PN |
| 9  | McNaught 2005    | ICU patients on antibiotics<br>N=130   | C.Random: no<br>ITT: yes<br>Blinding: no<br>Score: 7<br>Viability (intervention): NR     | Oral, NJ tube                                       | EN or PN + Proviva, (oatmeal & fruit drink) 5 x<br>10 <sup>7</sup> CFU/ml of L. plantarum 299v X 500 mls until<br>hospital discharge or beyond   | EN or PN alone  |
| 10 | Kotzampassi 2006 | Multiple trauma patients<br>from 5 ICUs<br>N=77<br>Blinding: double<br>Score: 8<br>Viability (intervention): NR<br>VAP determination: clinical                       |  | Endoscopic<br>gastrostomy or NG<br>tube             | EN or PN + Synbiotic 2000 Forte 10 <sup>11</sup> , 1<br>sachet/day for 15 days until ICU discharge   | EN or PN + Placebo<br>(Maltodextrin), mixed in tap<br>water   |
| 11 | Alberda 2007     | erda 2007 ICU patients<br>N=28 C.Random: no<br>ITT: yes;<br>Blinding: double<br>Score: 10<br>Viability (intervention): No for VSL # 3;<br>Yes for bacteria sonicates |  | NG tube   | Jevity Plus (EN) (10 g<br>fructooligosaccharides/1000 mL and 12 g of<br>soluble and insoluble fiber blend) +<br>VSL # 3, 1 package BID,<br>9 x 10 <sup>11</sup> /day for 7 days until ICU discharge or<br>EN discontinuation | Jevity Plus + Placebo   |
| 12 | Li 2007          | Severe acute pancreatitis<br>patients<br>N=25  | C.Random: no<br>ITT: yes<br>Blinding: no<br>Score: 7<br>Viability (intervention): NR     | Given enterally                                     | Jinshuangqi ( <i>bifidobacteria, lactobacillus and streptococcus</i> ) 2.0 g TID on basis of traditional treatment Duration: NR  | Traditional treatment   |

| 13 | Olah 2007      | Severe acute pancreatitis<br>patients<br>N=83                           | C.Random: no<br>ITT: no<br>Blinding: no<br>Score: 9<br>Viability (intervention): NR   | NJ tube   | EN (Nutricion Fibre) + Synbiotic 2000, 4 X 1010<br>CFU for 7 days   | EN (Nutricion Fibre) + 10g<br>plant fibres ((2.5 g each of<br>Betaglucan, Inulin, Pectin &<br>Resistant starch) (Prebiotics)<br>BID for at least 2 days |  |
|----|----------------|---|---|---|---|---|--|
| 14 | Forestier 2008 | Mixed ICU patients, 50%<br>on antibiotics<br>N=208                      | C.Random: not sure<br>ITT: no<br>Blinding: double<br>Score: 8<br>Viability (intervention): NR<br>VAP determination: objective | NG tube or Oral (after<br>tube removal)                         | <i>Lactobacillus casei rhamnosum</i> , 10 <sup>9 CFU</sup> BID until<br>ICU discharge   | Placebo (growth medium never exposed to bacteria).  |  |
| 15 | Besselink 2008 | Acute pancreatitis patients<br>from 15 ICUs<br>N=298                    | C.Random: not sure<br>ITT: yes<br>Blinding: double<br>Score:11<br>Viability (intervention): NR<br>VAP determination: clinical | NJ tube or Oral   | EN (Nutrison Multifibre)<br>+ Ecologic 641<br>10 <sup>10</sup> CFU BID for 28 days  | EN (Nutrison Multifibre) +<br>Placebo (cornstarch +<br>maltodextrins)   |  |
| 16 | Klarin 2008    | ICU patients from 5 ICUs,<br>on antibiotics for c. Difficile<br>N=68    | C.Random: yes<br>ITT: no<br>Blinding: double<br>Score: 10<br>Viability (intervention): NR                                     | Mixed in fermented<br>oatmeal added to<br>enteral feeds NG tube | 299 Lactobacillus plantarum,<br>8 x 10 <sup>8</sup> CFU/ml given as 6 x 100 ml doses every<br>12h & after 50 ml given BID until ICU discharge | Same oatmeal gruel mixed with lactic acid   |  |
| 17 | Knight 2009    | General ICU patients<br>N=300   | C.Random: yes<br>ITT: no<br>Blinding: double<br>Score: 10<br>Viability (intervention): NR<br>VAP determination: clinical      | NJ or OG (orogastric)<br>tube                                   | EN (Nutrition Energy) +<br>Synbiotic 2000 FORTE<br>4 x10 <sup>11</sup> species/sachet<br>BID for 28 days or ICU discharge                     | EN (Nutrison Energy) +<br>Placebo   |  |
| 18 | Barraud 2010   | Mechanically ventilated<br>ICU patients, 80% on<br>antibiotics<br>N=167 | C.Random: yes<br>ITT: yes;<br>Blinding: double<br>Score: 12<br>Viability (intervention): NR<br>VAP determination: objective   | NG tube   | EN (Fresubin) + Ergyphilus<br>2 x 10 <sup>10</sup> per capsule + potato starch 5 caps/day<br>for 28 days                                      | EN (fresubin) + Placebo<br>capsules (excipient of potato<br>starch)   |  |

| 19 | Morrow 2010    | ICU patients<br>N=146                             | C.Random: no;<br>ITT: yes;<br>Blinding: double; Score:10<br>Viability (intervention): yes<br>VAP determination: objective | Oropharynx and NG<br>tube | EN (routine care) + <i>Lactobacillus rhamnosus</i><br>GG, 2X10 <sup>9</sup> BID as lubricant and mixed with<br>water until extubation   | EN (routine care) + inert plant<br>starch inulin (prebiotic) BID as<br>as lubricant and mixed with<br>water |
|----|----------------|---|---|---------------------------|---|---|
| 20 | Frohmader 2010 | General ICU patients<br>on antibiotics<br>N=45    | C.Random: yes<br>ITT: yes<br>Blinding: double<br>Score: 11<br>Viability (intervention): yes                               | NG or NJ tube             | EN (Standard) + VSL #3<br>mixed in nutritional supplement (Sustagen), BID<br>until hospital discharge   | EN (Standard) + placebo<br>mixed in nutritional<br>supplement (Sustagen), BID                               |
| 21 | Ferrie 2011    | Critically ill patients with<br>diarrhea,<br>N=36 | C.Random: no<br>ITT: yes<br>Blinding: double<br>Score: 10<br>Viability (intervention): yes                                | NG tube                   | EN (Standard) + Culturelle ( <i>Lactobacillus rhamnosus GG</i> ), 10 <sup>10</sup> species/capsule + 280 mg inulin powder for 7 days  | EN (Standard) + Raftiline,<br>gelatin capsule with 280 mg<br>inulin powder (prebiotic)                      |
| 22 | Sharma 2011    | Acute pancreatitis patients<br>N=50               | C.Random: yes<br>ITT: yes<br>Blinding: double<br>Score:11<br>Viability (intervention): yes                                | Oral, NJ or NG            | EN (standard) or oral<br>4 sachets each 2.5 X 10 <sup>9</sup><br>Lactobacillus acidophilus, Bifidobacterium<br>longus, Bifidobacterium bifidum &<br>Bifidobacterium infantalis + 25 gms fructose for<br>7 days  | EN (Standard) + placebo   |
| 23 | Tan 2011       | Closed head injury patients<br>N=52               | C.Random: yes<br>ITT: yes<br>Blinding: single<br>Score:10<br>Viability (intervention): yes<br>VAP determination: clinical | NG tube                   | EN (standard)<br>total of 10 <sup>9</sup> bacteria i.e.<br>7 sachets each 0.5 x 10 <sup>8</sup> <i>Bifidobacterium</i><br><i>longum</i> , 0.5 X 10 <sup>7</sup> 1 <i>Lactobacillus bulgaricus</i> and<br>0.5 X 10 <sup>7</sup> <i>Streptococcus thermophilus</i> for 21<br>days | EN (standard)   |
| 24 | Cui 2013       | Severe acute pancreatitis<br>N=70                 | C.Random: no<br>ITT: yes<br>Blinding: no<br>Score:9<br>Viability (intervention): yes                                      | EN                        | EN + bifidobacterium, 4 capsules (each 210 mg, 2.604 x 10 <sup>9</sup> ) every 12 hours, given through nasal gastric tube. Total dose per day 20.832 x 10 <sup>9</sup> .  | EN  |

| 25     | Tan 2013  | Severe craniocerebral<br>trauma   | C.Random: no<br>ITT: other<br>Blinding: no<br>Score:11<br>Viability (intervention): yes  | NG tube   | EN + 1×10 <sup>9</sup> bacteria of viable probiotics (Golden<br>Bifid, 3.5 g for 3 times per day) per day for 21<br>days.   | EN (standard)           |  |
|--------|---|---|--|---|---|-------------------------|--|
| 26     | Wang 2013   | Severe acute pancreatitis<br>with intestinal ileus or<br>abdominal distention.<br>N=183 | C.Random: no<br>ITT: yes<br>Blinding: no<br>Score: 6<br>Viability (intervention): NR   | SBFT  | EN (standard) + capsules 0.5g TID containing<br>Bacillus subtilis and Enterococcus faecium (5.0<br>x 10 <sup>7</sup> Bacillus subtilis and 4.5 x 10 <sup>8</sup><br>Enterococcus faecium per 250 g capsule).<br>Unclear timeframe.    | EN (standard)           |  |
| 27     | Lopez de Toro 2014  | Medical and surgical ICU<br>pts with multi-organ failure<br>N=89                        | C.Random: yes<br>ITT: yes<br>Blinding: no<br>Score:11<br>Viability (intervention): NR  | EN  | EN + symbiotic drink with streptococcus<br>thermophilus, lactobacillus bulgaricus,<br>lactobacillus casei, lactobacillus acidophilus,<br>bifidobacterium, Escherichia coli, coliformes x 7<br>days (max 4.8 x10 <sup>9</sup> UFC/ml). | EN and PN               |  |
| 28     | Sanaie 2014   | Critically ill pts, SIRS,<br>expected LOS <u>&gt;</u> 7 days<br>N=40                    | C.Random: yes<br>ITT: yes<br>Blinding: double<br>Score:9<br>Viability (intervention): yes  | NG tube EN (standard) + 2 sachets VSL#3 BID x 7 days. |   | EN (standard) + placebo |  |
| EN: er | dom: concealed randomization<br>iteral nutrition<br>sojejunal |   | NG: nasogastric     CFU: Colony forming units       OG: orogastric     NR: not reported       FOS: fructooligosaccharides     NR: not reported |   |   |                         |  |

Trevis™: 1 capsule = Lactobacillus acidophilus La5, Bifidobacterium lactis Bb12, Streptococcus thermophilus, Lactobacillus bulgaricus, 4 x 10<sup>9</sup>/total

Synbiotic 2000 Forte: 10<sup>11</sup> CFU of each: Pediococcus pentoseceus 5-33:3, Leuconostoc mesenteroides 32-77:1, L. paracasei ssp paracasei 19, L. plantarum 2362 & 2.5 g each of: inulin, oat bran, pectin and resistant starch

Ergyphilus: 1010 Lactobaccilus rhamnosus GG, Lactobacillus casei, Lactobacillus acidophilus, Bifidobacterium bifidus,

VSL #  $3: > 10^{10}$  Bifidobacterium longum, Bifidobacterium breve,  $> 10^{10g}$  Bifidobacterium infantis,  $> 10^{11g}$  Lactobacillus acidophulus, plantarum, casei, bulgaris & Streptococcus thermophilus

Jinshuangqi: Bifidobacterium longum > 10<sup>7</sup> CFU, Lactobacillus bulgaricus > 10<sup>6</sup> CFU & Streptococcus Thermophilus > 10<sup>6</sup> CFU

Ecologic 641: Lactobacillus acidophilus, Lactobacillus salivarius, Lactococcus lactis, Bifidobacterium bifidum & Bifidobacterium lactis

Synbiotic 2000: 10<sup>10</sup> CFU of each: Pediococcus pentoseceus 5-33:3, Leuconostoc mesenteroides 32-77:1, L. paracasei ssp paracasei 19, L. plantarum 2362 & 2.5 g each of: betaglucan, inulin, pectin and resistant starch Golden Bifid: Bifidobacterium bifidum, Lactobacillus bulgaricus and Streptococcus thermophilus triple human probiotics supplemented oligosaccharides FOS (bifidus factor)

|   | Study            | Mort                                    | - V                                     | Infec                          | •                              | Length  | of Stay                                       | Diar   | rhea   |
|---|------------------|---|---|--------------------------------|--------------------------------|---|---|--|--|
|   | Sludy            | Intervention                            | Control                                 | Intervention                   | Control                        | Intervention                                  | Control                                       | Intervention   | Control  |
| 1 | Tempe 1983       | 3/20 (15)                               | 3/20 (15)                               | NR                             | NR                             | NR  | NR  | <b>Diarrhea days</b><br>34/389 (9)                                     | <b>Diarrhea days</b><br>63/373 (17)                        |
| 2 | Schlotterer 1987 | NR                                      | NR                                      | NR                             | NR                             | NR  | NR  | Diarrhea days<br>3/150 (2)   | Diarrhea days<br>19/143 (13)                               |
| 3 | Heimburger 1994  | NR                                      | NR                                      | NR                             | NR                             | NR  | NR  | <b>Diarrhea</b><br>5/16 (31)   | <b>Diarrhea</b><br>2/18 (11)                               |
| 4 | Bleichner 1997   | NR                                      | NR                                      | NR                             | NR                             | NR  | NR  | Diarrhea<br>18/64 (28) <sup>i</sup><br>Days w/ diarrhea<br>91/648 (14) | Diarrhea<br>24/64 (38)<br>Days w/ diarrhea<br>134/683 (20) |
| 5 | Kecskes 2003     | Hospital<br>1/22 (5)                    | Hospital<br>2/23 (9)                    | Septic Compl<br>1/22 (5)       | Septic Compl<br>7/23 (30)      | <b>Hospital</b><br>13.7 ± 8.7                 | Hospital<br>21.4 ± 17.9                       | NR   | NR   |
| 6 | Jain 2004        | Hospital<br>22/45 (49)                  | Hospital<br>20/45 (45)                  | Septic Compl<br>33/45 (73)     | Septic Compl<br>26/45 (58)     | Hospital<br>24.0 ± 31.5<br>ICU<br>11.9 ± 13.1 | Hospital<br>18.7 ± 13.5<br>ICU<br>9.0 ± 8.9   | NR   | NR   |
| 7 | Lu 2004          | Hospital<br>2/20 (10)                   | Hospital<br>1/20 (5)                    | Infectious Compl<br>8/20 (40)  | Infectious Compl<br>11/20 (55) | NR  | NR  | NR   | NR   |
| 8 | Klarin 2005      | Hospital<br>2/8 (25)<br>ICU<br>1/8 (12) | Hospital<br>2/7 (29)<br>ICU<br>2/7 (29) | NR                             | NR                             | Hospital<br>48.3 ± 30.4<br>ICU<br>14.2 ± 10.6 | Hospital<br>34.3 ± 15.4<br>ICU<br>16.3 ± 15.7 | NR   | NR   |
| 9 | McNaught 2005    | 18/52 (35)                              | 18/51 (35)                              | Septic morbidity<br>21/52 (40) | Septic morbidity<br>22/51 (43) | ICU<br>5 (2-9)                                | ICU<br>4 (2-7)                                | NR   | NR   |

 Table 1. Randomized studies evaluating Probiotics in critically ill patients (continued)

| 10 | Kotzampassi 2006 | ICU<br>5/35 (14)     | ICU<br>9/30 (30)      | Infections<br>22/35 (63)<br>VAP<br>19/35 (54)<br>Septic Compl<br>17/35 (49)<br>Central venous<br>line infections<br>13/35 (37)<br>Wound Infections<br>6/35 (17)<br>UTI<br>6/35 (17) | Infections<br>27/30 (90)<br>VAP<br>24/30 (80)<br>Septic Compl<br>23/30 (77)<br>Central venous<br>line infections<br>20/30 (66)<br>Wound Infections<br>8/30 (26)<br>UTI<br>13/30 (43) | ICU<br>27.7 ± 15.2     | ICU<br>41.3 ± 20.5        | <b>Diarrhea</b><br>5/35 (14) | <b>Diarrhea</b><br>10/30 (30) |
|----|------------------|----------------------|-----------------------|---|--|------------------------|---------------------------|------------------------------|-------------------------------|
| 11 | Alberda 2007     | ICU<br>1/10 (10)     | ICU<br>1/9 (11)       | NR  | NR   | NR                     | NR                        | <b>Diarrhea</b><br>1/10 (14) | Diarrhea<br>2/9 (23)          |
| 12 | Li 2007          | NR                   | NR                    | Infections<br>8/14 (58)   | Infections<br>10/11 (91)   | Hospital<br>42 ± 5.0   | Hospital<br>49 ± 6.8      | NR                           | NR                            |
| 13 | Olah 2007        | Hospital<br>2/33 (6) | Hospital<br>6/29 (21) | Infections<br>9/33 (27)<br>Septic Compl<br>7/33 (12)<br>Pancreatic<br>Abscess<br>2/33 (6)<br>Infected<br>Pancreatic<br>Necrosis<br>2/33 (6)<br>UTI<br>3/33 (9)                      | Infections<br>15/29 (52)<br>Septic Compl<br>17/29 (28)<br>Pancreatic<br>Abscess<br>2/29 (7)<br>Infected<br>Pancreatic<br>Necrosis<br>6/29 (21)<br>UTI<br>3/33 (9)                    | Hospital<br>14.9 ± 3.3 | Hospital<br>19.7 ± 4.5    | NR                           | NR                            |
| 14 | Forestier 2008   | NR                   | NR                    | <b>VAP</b><br>19/102 (19)   | <b>VAP</b><br>21/106 (20)  | ICU<br>22.5 ± 20.6     | <b>ICU</b><br>19.7 ± 16.7 | NR                           | NR                            |

| 15 | Besselink 2008 | 24/152 (16)   | 9/144 (6)   | Infections<br>46/152 (30)<br>VAP<br>24/152 (16)<br>Bacteremia<br>33/152 (22)<br>Infected necrosis<br>21/152 (14)<br>Urosepsis<br>1/52 (2)      | Infections<br>41/144 (28)<br>VAP<br>16/144 (11)<br>Bacteremia<br>22/144 (15)<br>Infected necrosis<br>14/144 (10)<br>Urosepsis<br>2/144 (1)       | Hospital<br>28.9 ± 41.5<br>ICU<br>6.6 ± 17    | Hospital<br>23.5 ± 25.9<br>ICU<br>3.0 ± 9.3   | <b>Diarrhea</b><br>25/152 (16)   | <b>Diarrhea</b><br>28/144 (19)   |
|----|----------------|---|---|--|--|---|---|--|--|
| 16 | Klarin 2008    | Hospital<br>3/22 (5)<br>ICU<br>2/22 (9)                             | Hospital<br>2/22 (0)<br>ICU<br>2/22 (9)                             | c. difficile+ fecal<br>samples<br>0/71   | c. difficile+ fecal<br>samples<br>4/80   | Hospital<br>25.8 ± 19.4<br>ICU<br>8.0 ± 5.4   | Hospital<br>50.3 ± 75.2<br>ICU<br>11.6 ± 14   | NR   | NR   |
| 17 | Knight 2009    | Hospital<br>35/130 (27)<br>ICU<br>28/130 (22)                       | Hospital<br>42/129 (33)<br>ICU<br>34/129 (26)                       | <b>VAP</b><br>12/130 (9)   | <b>VAP</b><br>17/129 (13)  | ICU<br>6 (3-11)                               | ICU<br>7 (3-14)                               | <b>Diarrhea</b><br>7/130 (5)   | <b>Diarrhea</b><br>9/129 (7)   |
| 18 | Barraud 2010   | ICU<br>21/87 (24)<br>28 days<br>22/87 (25)<br>90 days<br>27/87 (31) | ICU<br>21/80 (26)<br>28 days<br>19/80 (24)<br>90 days<br>24/80 (30) | All infections<br>30/87 (34)<br>Infection > 96 hr<br>26/87 (30)<br>VAP<br>23/87 (26)<br>Catheter related<br>BSI<br>3/87 (4)<br>UTI<br>4/87 (5) | All infections<br>30/80 (38)<br>Infection > 96 hr<br>29/80 (36)<br>VAP<br>15/80 (19)<br>Catheter related<br>BSI<br>11/80 (14)<br>UTI<br>4/89 (5) | Hospital<br>26.6 ± 22.3<br>ICU<br>18.7 ± 12.4 | Hospital<br>28.9 ± 26.4<br>ICU<br>20.2 ± 20.8 | <b>Diarrhea</b><br>48/87 (55)  | <b>Diarrhea</b><br>42/80 (53)  |
| 19 | Morrow 2010    | 12/68 (18)  | 15/70 (21)  | <b>VAP</b><br>13/73 (18)   | VAP<br>28/73 (38)  | Hospital<br>21.4 ± 14.9<br>ICU<br>14.8 ± 11.8 | Hospital<br>21.7 ± 17.4<br>ICU<br>14.6 ± 11.6 | Non C. Difficile<br>Diarrhea<br>42/68 (62)<br>C. difficile<br>diarrhea<br>4/68 (6) | Non C. Difficile<br>Diarrhea<br>44/70 (63)<br>C. difficile<br>diarrhea<br>13/70 (19) |

| 20 | Frohmader 2010        | 5/20 (25)                                      | 3/25 (12)                                      | NR   | NR  | ICU<br>7.3 ± 5.7                                  | <b>ICU</b><br>8.1 ± 4                             | Diarrhea<br>episodes/pt/day<br>0.53 ± 0.54  | Diarrhea<br>episodes/pt/day<br>1.05 ± 1.08                                |
|----|-----------------------|--|--|--|---|---|---|---|---|
| 21 | Ferrie 2011           | Hospital<br>2/18 (11)<br>6 months<br>7/18 (39) | Hospital<br>2/18 (11)<br>6 months<br>5/18 (28) | Infections<br>14/18 (78)   | Infections<br>16/18 (89)                              | Hospital<br>54.50 ± 31.26<br>ICU<br>32.04 ± 24.46 | Hospital<br>59.04 ± 33.92<br>ICU<br>29.75 ± 18.81 | Duration of<br>Diarrhea<br>$3.83 \pm 2.39$<br>Loose stools/day<br>$1.58 \pm 0.88$ | Duration of<br>Diarrhea<br>2.56 ± 1.85<br>Loose stools/day<br>1.10 ± 0.79 |
| 22 | Sharma 2011           | Hospital<br>2/24 (8)                           | Hospital<br>2/26 (8)                           | NR   | NR  | Hospital<br>13.23 ± 18.19<br>ICU<br>4.94 ± 9.54   | Hospital<br>9.69 ±9.69<br>ICU<br>4.0 ± 5.86       | NR  | NR  |
| 23 | Tan 2011              | <b>28 day</b><br>3/26 (12)                     | <b>28 day</b><br>5/26 (19)                     | Infections<br>9/26 (35)<br>VAP<br>7/26 (27)                      | Infections<br>15/26 (58)<br>VAP<br>13/26 (50)         | ICU<br>6.8 ± 3.8                                  | ICU<br>10.7 ± 7.3                                 | NR  | NR  |
| 24 | Cui 2013              | Hospital<br>1/23 (4)                           | <b>Hospital</b><br>1/25 (4)                    | N/A  | N/A   | Hospital<br>10.4 ± 3.9 (23)                       | Hospital<br>13.4 ± 5.2 (25)                       | NR  | NR  |
| 25 | Tan 2013              | <b>28 day</b><br>23/26 (12)                    | <b>28 day</b><br>5/26 (19)                     | NR   | NR  | ICU<br>6.8 ± 3.8 (26)                             | ICU<br>10.7 ± 7.3 (26)                            | NR  | NR  |
| 26 | Wang 2013             | Unspecified<br>1/62 (8.1)                      | Unspecified<br>3/61 (9.8)                      | Pancreatic sepsis<br>8/62 (13)<br>MODS<br>7/62 (11.3)20<br>14.16 | Pancreatic sepsis<br>13/61 (21)<br>MODS<br>15/61 (25) | NR  | NR  | NR  | NR  |
| 27 | Lopez de Toro<br>2014 | Hospital<br>19/46 (41)<br>ICU<br>15/46 (33)    | Hospital<br>18/43 (42)<br>ICU<br>14/43 (33)    | Hospital acquired<br>infections<br>9/46 (20)                     | Hospital acquired<br>infections<br>13/43 (30)         | Hospital<br>18.5 (10-36)<br>ICU<br>9 (3-19)       | Hospital<br>24.5 (10-38)<br>ICU<br>8 (2.5-16.5)   | NR  | NR  |

#### Canadian Clinical Practice Guidelines

| 28     | Sanaie 2014                         | <b>28 day</b><br>2/20 (10) | <b>28 day</b><br>5/20 (25) | Bacteremia<br>2/20(10) | Bacteremia<br>5/20(25) | ICU<br>13.85 ± 6.96 | ICU<br>14.16 ± 5.97    | NR | NR |
|--------|-------------------------------------|----------------------------|----------------------------|------------------------|------------------------|---------------------|------------------------|----|----|
| NR: N  | NR: Not Reported UTI: Urinary Tract |                            |                            |                        |                        | BSI:                | Blood Stream Infection |    |    |
| VAP: Y | Ventilator Associated Pneun         | ICU: Int                   | ensive Care Unit           |                        |                        |                     |                        |    |    |

Figure 1. Hospital Mortality

,

Figure 2. ICU Mortality

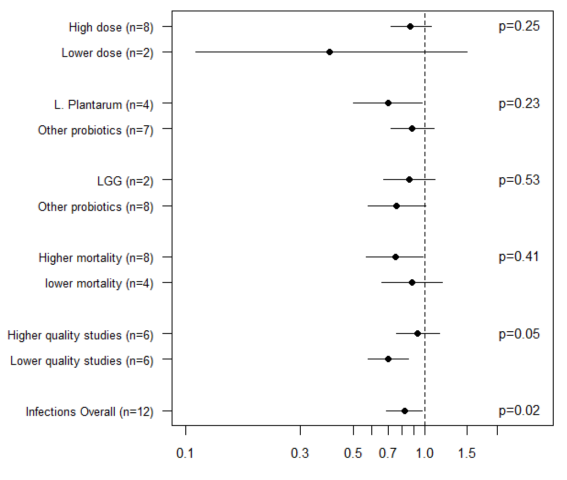
•

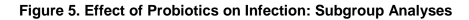
Figure 3. Infections

.

# Figure 4. VAP

|                                   | Probiotics             |         | Control     |          | Risk Ratio  |                     |                                    | Risk Ratio          |
|-----------------------------------|------------------------|---------|-------------|----------|-------------|---------------------|------------------------------------|---------------------|
| Study or Subgroup                 | Events                 | Total   | Events      | Total    | Weight      | M-H, Random, 95% Cl | Year                               | M-H, Random, 95% Cl |
| Kotzampassi 2006                  | 19                     | 35      | 24          | 30       | 25.4%       | 0.68 [0.48, 0.97]   | 2006                               | -=-                 |
| Forestier 2008                    | 19                     | 102     | 21          | 106      | 16.9%       | 0.94 [0.54, 1.64]   | 2008                               | -4-                 |
| Knight 2009                       | 12                     | 130     | 17          | 129      | 12.9%       | 0.70 [0.35, 1.41]   | 2009                               |                     |
| Morrow 2010                       | 13                     | 73      | 28          | 73       | 16.5%       | 0.46 [0.26, 0.82]   | 2010                               |                     |
| Barraud 2010                      | 23                     | 87      | 15          | 80       | 16.4%       | 1.41 [0.79, 2.51]   | 2010                               |                     |
| Tan 2011                          | 7                      | 26      | 13          | 26       | 11.9%       | 0.54 [0.26, 1.13]   | 2011                               |                     |
| Total (95% CI)                    |                        | 453     |             | 444      | 100.0%      | 0.74 [0.55, 1.01]   |                                    | •                   |
| Total events                      | 93                     |         | 118         |          |             |                     |                                    |                     |
| Heterogeneity: Tau <sup>2</sup> = | 0.06; Chi <sup>2</sup> | = 9.10  | , df = 5 (F | ° = 0.11 | ); I² = 45% | 6                   |                                    | 0.01 0.1 1 10 100   |
| Test for overall effect: 2        | Z = 1.90 (             | P = 0.0 | 6)          |          |             |                     | Favours Probiotics Favours control |                     |





Favours Experimental RR Favours control

Legend: Numbers in brackets indicate the number of studies. RR: Risk ratio

p values for the subgroups indicate the differences in the subgroup effect of probiotics on infections.

LGG= Lactobacillus rhamnosus GG

Figure 6. ICU LOS

•

## Figure 7. Diarrhea

| 5                                 | Experim                  | ental   | Contr     | ol      |            | Risk Ratio          |  | Risk Ratio          |
|-----------------------------------|--------------------------|---------|-----------|---------|------------|---------------------|--|---------------------|
| Study or Subgroup                 | Events                   | Total   | Events    | Total   | Weight     | M-H, Random, 95% Cl | Year   | M-H, Random, 95% Cl |
| Heimburger 1994                   | 5                        | 16      | 2         | 18      | 1.3%       | 2.81 [0.63, 12.54]  | 1994   |                     |
| Bleichner 1997                    | 18                       | 64      | 24        | 64      | 11.3%      | 0.75 [0.45, 1.24]   | 1997   | +                   |
| Kotzampassi 2006                  | 5                        | 35      | 10        | 30      | 3.2%       | 0.43 [0.16, 1.12]   | 2006   | <b>-</b>            |
| Alberda 2007                      | 1                        | 10      | 2         | 9       | 0.6%       | 0.45 [0.05, 4.16]   | 2007   |                     |
| Besselink 2008                    | 25                       | 152     | 28        | 144     | 11.9%      | 0.85 [0.52, 1.38]   | 2008   |                     |
| Knight 2009                       | 7                        | 130     | 9         | 129     | 3.2%       | 0.77 [0.30, 2.01]   | 2008   |                     |
| Barraud 2010                      | 48                       | 87      | 42        | 80      | 32.3%      | 1.05 [0.79, 1.39]   | 2010   | +                   |
| Morrow 2010                       | 42                       | 68      | 44        | 73      | 36.1%      | 1.02 [0.79, 1.33]   | 2010   | +                   |
| Total (95% CI)                    |                          | 562     |           | 547     | 100.0%     | 0.95 [0.80, 1.13]   |  | •                   |
| Total events                      | 151                      |         | 161       |         |            |                     |  |                     |
| Heterogeneity: Tau <sup>2</sup> = | = 0.00; Chi <sup>z</sup> | = 7.37, | df = 7 (P | = 0.39) | ); I² = 5% |                     |  |                     |
| Test for overall effect:          | P = 0.54                 | )       |           |         |            | F                   | 0.01 0.1 1 10 100<br>avours experimental Favours control |                     |