Online-Only Supplement:

Early parenteral nutrition in critically ill patients with short term relative contraindications to early enteral nutrition:

A randomized controlled trial.


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Note: Additional information may be found at the study web site:
www.EvidenceBased.net/EarlyPN.
**Early PN Trial Conduct and Management:**


*PN protocol sub-committee:* Gordon S. Doig (Chair), Fiona Simpson, Michael O’Leary.  

*Infectious complications sub-committee:* Gordon S. Doig (Chair), Tom Solano, Fiona Simpson.  

*Data Quality and Management:* Jennifer L. Hannam (Northern Clinical School Intensive Care Research Unit, University of Sydney, Australia).  

*Statistical analysis:* Gordon S. Doig.  

*Independent Data Safety and Monitoring Committee:* John Moran (Chair, Dept of Intensive Care, The Queen Elizabeth Hospital, Adelaide, Australia), Petra Graham (Dept of Statistics, Macquarie University, Sydney, Australia) and Andrew Bersten (Dept of Critical Care Medicine, Flinders University, Adelaide, Australia).
Table 1: Complete Eligibility Criteria:

Inclusion Criteria (All YES answers for enrolment)

1. Is the patient expected to remain in ICU today and tomorrow?
2. Is the patient 18 years of age or older?
3. Has the patient been admitted to the study ICU less than 24 hours?
4. Does the patient have a central venous access line through which parenteral nutrition could be delivered?
5. Is this patient not expected to receive enteral, parenteral or oral nutrition today or tomorrow?

Exclusion Criteria (All NO answers for enrolment, YES to any for exclusion)

1. Known pregnancy or currently breastfeeding.
2. Has the patient previously been enrolled and randomised into this study?
3. Is the patient to receive palliative care only and is not expected to survive ICU or hospital discharge?
4. Was the patient admitted to this, or another, ICU during this current hospitalisation?
5. Was the patient admitted to the study ICU directly from another ICU?
6. Is the patient moribund and not expected to survive 24 hours?
7. Is the patient brain dead or suspected to be brain dead?
8. Are there long term contraindications to enteral or oral nutrition such that the patient would normally be supported with parenteral nutrition (Ex. Home TPN patient)?
9. Does the patient require treatment of thermal injury to greater than 20% of total body surface area?
10. Is the primary reason for admission to the ICU for the treatment of a condition that requires timely nutritional support (Ex. Anorexia nervosa.)?
11. Body weight < 35 Kg
12. Height < 140 cm (Demi armspan < 59 cm)
13. Is there a contraindication to treatment with Kabiven G19%?

NB - see next page for contraindications to Kabiven G19% based on TGA licensing indications.
Contraindications to Kabiven G19% based on TGA Licensing Indications.

c1. Known hypersensitivity to egg or soya protein or to any of the ingredients of the study PN (for full ingredients see Product Information, MIMS TGA Document Appendix 1).

c2. Severe hyperlipidaemia (Documented serum total cholesterol >7mmol/L and/or triglycerides >3 mmol/L).

c3. Severe liver insufficiency (Biopsy proven cirrhosis, or documented portal hypertension with a known past history of either upper GI bleeding attributed to portal hypertension or of hepatic failure leading to encephalopathy / coma.)

c4. Severe blood coagulation disorders (Documented INR > 3.0 not due to coumarin therapy, platelet count <15,000).

c5. Inborn errors of amino acid metabolism (Ex. PKU etc)

c6. Severe renal insufficiency without access to haemofiltration or dialysis.

c7. Acute shock as defined by arterial systolic blood pressure ≤ 90mmHg or mean arterial pressure ≤ 70mmHg despite adequate fluid resuscitation (i.e. following rapid infusion of ≥ 500mL crystalloid or 200mL colloid solution and /or PAOP ≥ 12mmHg, CVP ≥ 8mmHg) or increasing need for noradrenaline / adrenaline / dopamine to maintain blood pressure where the infusion rate has increased by more that 50% over the previous hour to greater than 0.6mg/hour (10 mcg/min) norad / adrenaline or 30mg dopamine.

 Patients are not excluded if their initial shock responds to fluid therapy or if the catecholamine infusion rate has not increased by more than 50% over the previous one hour period or if the current infusion rate is less than 0.6mg norad / adrenaline per hour .

c8. Hyperglycaemia (blood sugar > 10 mmol/L) that currently requires the administration of more than 6 units of insulin/hour at the time of enrolment.

c9. Pathologically elevated serum levels of any of the electrolytes included in Kabiven G19% at the time of enrolment. Documented Sodium >155 mmol/L, potassium > 6.2 mmol/L, magnesium > 2.0 mmol/L, ionised calcium > 1.5 mmol/L, phosphate >2.0 mmol/L, chloride > 120 mmol/L.

 The patient may become eligible if these pathologically elevated electrolyte levels can be corrected within 24 hours of admission to the study ICU.

c10. General contraindications of infusion therapy: acute pulmonary oedema, hyperhydration, decompensated cardiac insufficiency and hypotonic dehydration

 The patient may become eligible if these general contraindications to fluid therapy can be corrected within 24 hours of admission to the study ICU.

c11. Haemophagocytic syndrome

c12. Severe trauma with acute shock (see Exclusion Criteria c7 for definition of acute shock).

c13. Diabetes mellitus with ketoacidosis or non-ketotic hyperosmolar state.

c14. Acute myocardial infarction with acute shock (see Exclusion Criteria c7 for definition of acute shock) or pulmonary oedema.

c15. (Metabolic acidosis or severe sepsis) with acute shock (see Exclusion Criteria c7 for definition of acute shock. Use Bone Criteria for definition of Severe Sepsis).

c16. Coma (GCS ≤8) in association with hyperosmolarity of the blood (≥320mOs/kg) from any cause.
eTable 2: PN Protocols

Study PN Protocol A: ALL Early PN PATIENTS EXCEPT MALNOURISHED

Feeding Day 1 (first 24 hours of PN)
- Commence Kabiven G19% at 60ml/hr (or goal rate, whichever is lower).
- Consider trace element, mineral and vitamin needs as clinically appropriate.

Feeding Day 2 (second 24 hours of PN)
- Increase Kabiven G19% to 80ml/hr (or goal rate, whichever is lower).
- Consider trace element, mineral and vitamin needs as clinically appropriate.

Feeding Day 3 (next 24 hours)
- Increase Kabiven G19% to goal rate, as appropriate.
- Consider trace element, mineral and vitamin needs, as clinically appropriate.
- Recommend trialing enteral/oral nutrition, if clinically appropriate.
- Once the patient tolerates ≥ 475 kcal/day EN, complete remainder of 24 hour Kabiven infusion and do not hang another bag.
- If patient tolerates any oral caloric intake from food, complete remainder of 24 hour Kabiven infusion and do not hang another bag.

Feeding Day 4 (next 24 hours) plus all additional days after Day 4
- May switch to parenteral nutrition solution tailored to patient’s specific clinical needs. Goals not to exceed 25–35 kcal/kg and 1.0–1.5 g protein/kg.
- Consider long term needs regarding trace element, mineral and vitamins as clinically appropriate.
- Recommend trialing enteral/oral nutrition, if clinically appropriate.
- Once the patient tolerates ≥ 475 kcal/day EN, complete remainder of 24 hour Kabiven infusion and do not hang another bag.
- If patient tolerates any oral caloric intake from food, complete remainder of 24 hour Kabiven infusion and do not hang another bag.

INSULIN / GLUCOSE PROTOCOL: Early PN Patients

If glucose levels exceed 10 mmol/L an insulin infusion should be commenced and titrated to achieve peak serum glucose levels of < 10 mmol/L. Frequent monitoring of the patient’s blood glucose should be initiated as per your ICU’s usual practice for patients receiving an insulin infusion.

If insulin infusion is required at ≥ 6 units/hr to maintain glucose target:
- Reduce Kabiven G19% to 40ml/hr for 24 hours.
- At the end of 24 hours, if insulin needs are reduced below 6 units/hr, increase Kabiven G19% to 80mls (or original goal rate, whichever is lower) for 24 hours.
- At the end of this second 24 hour period, if insulin needs remain below 6 units/hr, increase Kabiven G19% to goal rate.
- If insulin requirements exceed 6 units/hr at any time during the above process, reduce PN to previously tolerated rate, or 40 mls/hr (whichever is higher), for 24 hours. Begin increasing rate every 24 hours as above, if tolerated.
eTable 2 continued: PN Protocols

Study PN Protocol B: MALNOURISHED Early PN PATIENTS (Ex. BMI ≤ 17):

Feeding Day 1 (first 24 h of PN)
- Commence Kabiven G19% at 40ml/hr (or goal rate, whichever lower).
- Strongly recommend administering 100mg thiamine, commencing at least 30 minutes prior to initiation of Kabiven G19% infusion, as clinically indicated as per product licensing indications.
- Recommend daily administration of other vitamins, minerals and trace elements, as clinically appropriate.

Feeding Day 2 (second 24 hours of PN)
- Increase Kabiven G19% to 60ml/hr (or goal rate, whichever is lower).
- Recommend daily administration of vitamins, minerals and trace elements, as clinically appropriate.

Feeding Day 3 (next 24 hours)
- Increase Kabiven G19% to goal rate, as appropriate.
- Recommend daily administration of vitamins, minerals and trace elements, as clinically appropriate.
- Recommend trialing enteral/oral nutrition, if clinically appropriate.
- Once the patient tolerates ≥ 475 kcal/day EN, complete remainder of 24 hour Kabiven infusion and do not hang another bag.
- If patient tolerates any oral caloric intake from food, complete remainder of 24 hour Kabiven infusion and do not hang another bag.

Feeding Day 4 (next 24 hours) plus all additional days after Day 4
- May switch to parenteral nutrition solution tailored to patient’s specific clinical needs. Goals not to exceed 25–35 kcal/kg and 1.0–1.5 g protein/kg.
- Strongly recommend addressing long term needs regarding trace elements, minerals and vitamins as clinically appropriate.
- Recommend trialing enteral/oral nutrition, if clinically appropriate.
- Once the patient tolerates ≥ 475 kcal/day EN, complete remainder of 24 hour Kabiven infusion and do not hang another bag.
- If patient tolerates any oral caloric intake from food, complete remainder of 24 hour Kabiven infusion and do not hang another bag.

INSULIN / GLUCOSE PROTOCOL: Early PN Patients

If glucose levels exceed 10 mmol/L an insulin infusion should be commenced and titrated to achieve peak serum glucose levels of < 10 mmol/L. Frequent monitoring of the patient’s blood glucose should be initiated as per your ICU’s usual practice for patients receiving an insulin infusion.

If insulin infusion is required at ≥ 6 units/hr to maintain glucose target:
- Reduce Kabiven G19% to 40ml/hr for 24 hours.
- At the end of 24 hours, if insulin needs are reduced below 6 units/hr, increase Kabiven G19% to 80mls (or original goal rate, whichever is lower) for 24 hours.
- At the end of this second 24 hour period, if insulin needs remain below 6 units/hr, increase Kabiven G19% to goal rate.
- If insulin requirements exceed 6 units/hr at any time during the above process, reduce PN to previously tolerated rate, or 40 mls/hr (whichever is higher), for 24 hours. Begin increasing rate every 24 hours as above, if tolerated.
**eTable 3: Harris-Benedict equations and adjustment factors used by study web site**

For males:

Target metabolic needs (kcals/day) = (66.5 + (13.75 × Wt) + (5.003 × Ht) - (6.775 × Age)) × adjustment factor

For females:

Target metabolic needs (kcals/day) = (655.1 + (9.563 × Wt) + (1.85 × Ht) - (4.676 × Age)) × adjustment factor

Where:
- **Wt** = weight in Kg
- **Ht** = height in cm
- **Age** = Age in years

**Adjustment Factors** (most severe was selected).

**Other, not listed below**
- Any other problem, not listed below.

**Infection, mild**
- Ex. mild skin, line or surgical wound infection. Local redness, heat and swelling but no systemic signs.

**Operation, minor**
- Any surgical procedure that does not require general anaesthesia or respiratory support.

**Operation, major**
- Any surgical procedure that does require general anaesthesia or respiratory support.

**Infection, peritonitis (non-septic)**
- Peritonitis based on visual inspection or culture. Patient does not have systemic signs of sepsis.

**Cancer**
- Patient is known to have an active tumour. May or may not be undergoing active or palliative treatment.

**Trauma, single fracture (skeletal)**
- Patient has trauma resulting in a single skeletal fracture of any bone except long bones.

**Infection, moderate**
- Infections that would normally require ICU admission for treatment. Ex. Community acquired pneumonia, Ventilator Associated Pneumonia.

**Trauma, single long-bone fracture**
- Trauma with a fracture to a long bone (femur, humerus, tibia, fibula, radius and ulna).

**Trauma, multiple fractures**
- Trauma with multiple fractures to any bones, including at least one long bone.

**Trauma, blunt with or without fractures**
- Blunt trauma, such as a motor vehicle crash and fall from height. Includes Penetrating trauma.

**Infection, severe**
- Any infection, or suspected infection, that expresses itself systemically as sepsis.

**Burns, less than or equal to 20% TBSA**
- Chemical or thermal burns to less than 20% of total body surface area.

**Malnourished (high risk of refeeding syndrome)**
- Body mass index of less than 17 or history and physical exam consistent with malnourishment or high risk of malnourishment. Based on clinical grounds decided by attending clinician.

**Note:** Harris-Benedict calculated targets were capped at 35 kcal/kg/day and obese patients (BMI ≥ 30 kg/m²) used ideal body weight (BMI = 21 kg/m²) in all Harris-Benedict calculations.
<table>
<thead>
<tr>
<th></th>
<th>Standard Care</th>
<th>Early PN</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>682 patients</td>
<td>681 patients</td>
</tr>
<tr>
<td>Patients receiving PN at any time, n (%)</td>
<td>254 (37.2%)</td>
<td>679 (99.7%)</td>
</tr>
<tr>
<td>Patients receiving EN at any time, n (%)</td>
<td>298 (43.7%)</td>
<td>274 (40.23%)</td>
</tr>
<tr>
<td>Patients starting PN first, n (%)</td>
<td>186 (27.3%)</td>
<td>679 (99.7%)</td>
</tr>
<tr>
<td>Time from enrolment to starting PN first, mean (95% CI)</td>
<td>1.99 days (1.45 to 2.70)</td>
<td>44 minutes (36 to 55)</td>
</tr>
<tr>
<td>Patients commencing EN after starting PN, n (%)</td>
<td>80/186 (43.0%)</td>
<td>405/679 (59.6%)</td>
</tr>
<tr>
<td>Time to commencing EN after starting PN, mean (95% CI)</td>
<td>5.08 days (4.66 to 5.52)</td>
<td>3.83 days (3.52 to 4.17)</td>
</tr>
<tr>
<td>Patients starting EN first, n (%)</td>
<td>199 (29.2%)</td>
<td>0 (0%)</td>
</tr>
<tr>
<td>Time from enrolment to starting EN first, mean (95% CI)</td>
<td>1.98 days (1.43 to 2.78)</td>
<td></td>
</tr>
<tr>
<td>Patients receiving PN after starting EN, n (%)</td>
<td>48/199 (24.1%)</td>
<td>0 (0%)</td>
</tr>
<tr>
<td>Time to receiving PN after starting EN, mean (95% CI)</td>
<td>4.59 days (4.09 to 5.17)</td>
<td></td>
</tr>
<tr>
<td>Patients starting EN+PN together, n (%)</td>
<td>19 (2.8%)</td>
<td>0 (0%)</td>
</tr>
<tr>
<td>Time from enrolment to starting EN+PN together, mean (95% CI)</td>
<td>5.58 days (3.90 to 7.96)</td>
<td></td>
</tr>
<tr>
<td>Patients who never received EN or PN, n (%)</td>
<td>278 (40.8%)</td>
<td>2 (0.3%)</td>
</tr>
<tr>
<td>Time from enrolment to ICU discharge or death in patients never receiving EN or PN, mean (95% CI)</td>
<td>3.72 days (2.65 to 5.20)</td>
<td>4.50 days (3.21 to 6.30)</td>
</tr>
</tbody>
</table>

**PN**: parenteral nutrition, **EN**: enteral nutrition, **CI**: confidence interval
### eTable 5: Clinically Significant Organ Failure and Concomitant Interventions, crude event rates, not adjusted for time at risk (ICU stay)

<table>
<thead>
<tr>
<th>Organ system failures, Not adjusted for time at risk of failure (ICU stay)</th>
<th>Standard Care 682 patients mean (95% CI)</th>
<th>Early PN 681 patients mean (95% CI)</th>
<th>mean difference (95% CI)</th>
<th>p-value*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Renal failure, creatinine &gt; 2.0 mg/dL (170 μmol/L)</td>
<td>1.63 (1.41 to 1.88)</td>
<td>1.74 (1.51 to 2.01)</td>
<td>0.11 (-0.32 to 0.69)</td>
<td>0.65</td>
</tr>
<tr>
<td>Pulmonary failure, PaO₂:FiO₂ ratio &lt; 301</td>
<td>7.94 (7.55 to 8.33)</td>
<td>7.31 (6.97 to 7.69)</td>
<td>-0.62 (-1.30 to 0.14)</td>
<td>0.11</td>
</tr>
<tr>
<td>Hepatic failure, total bilirubin &gt; 2.0 mg/dL (32.5 μmol/L)</td>
<td>1.29 (1.09 to 1.52)</td>
<td>1.20 (1.01 to 1.42)</td>
<td>-0.09 (-0.43 to 0.40)</td>
<td>0.69</td>
</tr>
<tr>
<td>Coagulation failure, platelets &lt; 81 x10⁹/L</td>
<td>1.91 (1.77 to 2.04)</td>
<td>1.48 (1.38 to 1.58)</td>
<td>-0.43 (-0.62 to -0.21)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Cardiovascular systolic blood pressure &lt; 90mmHg, not fluid responsive</td>
<td>1.08 (0.97 to 1.21)</td>
<td>0.85 (0.76 to 0.95)</td>
<td>-0.23 (-0.40 to 0.02)</td>
<td>0.04</td>
</tr>
<tr>
<td>MODs Number of days with two or more organ system failures on the same day</td>
<td>3.78 (3.53 to 4.06)</td>
<td>3.57 (3.34 to 4.83)</td>
<td>-0.21 (-0.67 to 0.32)</td>
<td>0.42</td>
</tr>
<tr>
<td>Number of patients developing MODS during ICU stay, n (%)</td>
<td>542 (79.5%)</td>
<td>526 (77.2%)</td>
<td>-2.2% (-7.5% to 3.1%)</td>
<td>0.32**</td>
</tr>
</tbody>
</table>

### Concomitant therapies and tertiary outcomes, adjusted for time at risk (ICU stay)

<table>
<thead>
<tr>
<th></th>
<th>days per 10 patient•ICU days</th>
</tr>
</thead>
<tbody>
<tr>
<td>Renal replacement therapy</td>
<td>1.58 (0.94 to 1.43)</td>
</tr>
<tr>
<td>Invasive mechanical ventilation</td>
<td>7.41 (6.99 to 7.85)</td>
</tr>
<tr>
<td>Pressure ulcer treatment Treatment for Stage 1 or greater</td>
<td>1.28 (1.04 to 1.58)</td>
</tr>
<tr>
<td>Low serum albumin &lt; 2.5 g/dL (25 g/L)</td>
<td>5.44 (5.11 to 5.78)</td>
</tr>
<tr>
<td>Systemic antibiotic use</td>
<td>7.35 (7.00 to 7.71)</td>
</tr>
<tr>
<td>Witnessed aspiration events per patient</td>
<td>0.015 (0.012 to 0.017)</td>
</tr>
<tr>
<td>Witnessed aspiration with new pulmonary infiltrates events per patient</td>
<td>0.004 (0.002 to 0.010)</td>
</tr>
</tbody>
</table>

*p-values from Negative Binomial model.  ** p-value from exact Pearson chi-square. PN: Parenteral Nutrition, CI: Confidence Interval, MODs: Multiple organ dysfunction syndrome, ICU: Intensive Care Unit.
eFigure 1. Energy intake from enteral nutrition and/or parenteral nutrition during ICU stay

PN: parenteral nutrition, EN: enteral nutrition, ICU: intensive care unit, Kcals: kilocalories. Day 1 equals day of study enrolment.
eFigure 2: Subjective Global Assessment of muscle wasting during ICU stay.

Legend: P-values from fully factorial repeated measures ANOVA: p < 0.0001 change over time, p =0.014 difference between groups (0.16 grade per week). Grey shaded area represents test based 95% confidence interval from fully factorial repeated measures ANOVA analysis between groups. 

ICU: Intensive Care Unit. PN: parenteral nutrition.
eFigure 3: Subjective Global Assessment of fat loss during ICU stay.

**Legend:** P-values from fully factorial repeated measures ANOVA: $p < 0.0001$ change over time, $p = 0.045$ difference between groups (0.13 grade per week). Grey shaded area represents test based 95% confidence interval from fully factorial repeated measures ANOVA analysis between groups. **ICU:** Intensive Care Unit. **PN:** parenteral nutrition.