Home parenteral nutrition (HPN) is needed for patients with acute or chronic intestinal failure in whom nutritional and / or water and electrolyte balance cannot be corrected by oral or enteral feeding and in whom PN is feasible at home (Messing B et al in Intestinal Failure 2001). While HPN was first described in 1970 (via an arterio-venous fistula), it has been used as a treatment via a central catheter in the UK since the 1980s and the number of patients have been increasing so that the total number exceeded 1000 in 2012 (British Artificial Nutrition Survey (BANS) reports). Over the last 5 years there has been a greater than 200% increase in new HPN patients.

The reasons for starting PN include a short, leaking (perforated), obstructed or dysfunctional gut or extensive small intestinal mucosal disease. Occasionally it is due to no enteral access or after unsuccessful enteral nutrition. Consideration for stopping HPN may be when oral / enteral intake is resumed, a problem has resolved (e.g. obstruction or entero-cutaneous fistula), or the gut has adapted (e.g. after colon continuity has been restored) or it is no longer in the patients’ best interest”.

The aims of HPN include preventing or treating undernutrition and/or dehydration, helping a patient become fit for surgery, reducing diarrhoea/vomiting and reducing abdominal pain caused by oral/enteral feeding.

A patient having HPN or his/her family or carers have much to learn before/while a patient is at home on PN. They need to learn about the principles of asepsis, basic gut and vascular anatomy and the complications of PN, and about their underlying illness. They have to learn the skills of hand washing, putting on gloves, connecting and disconnecting a feed, changing a dressing, flushing a line and managing a pump. To do this they need to be physically fit (to lift pump and bags), with good manual dexterity and vision. They need to be psychologically motivated and manage the PN in an appropriate home environment. They need to learn how to adjust to lifestyle events such as bathing, showering, swimming, partaking of sports and travel. In addition to the PN they may also have to cope with wound/stoma management, chronic pain and much medication (e.g. analgesics, proton pump inhibitors, anti-diarrhoeal and anti-emetic drugs).

This document was requested by Michael Stroud when Chairman of the British Association of Parenteral and Enteral Nutrition (BAPEN) to help Nutrition Support Teams (NST) setting up Intestinal Failure (IF)/HPN centres and it addresses the key practicalities. While not directed at commissioners, it may be referenced by them.
It addresses the facilities and quality of care expected at an HPN/IF centre for infants, children, young people and adults. It has been written for guidance in the United Kingdom. It can be read in conjunction with the Healthcare Improvement Scotland Document on Complex Nutritional Care Standards (December 2015). This BIFA position statement does cover 5 of the 6 standards quoted in the document; however it does not address the enteral nutrition one. It was drafted by the BIFA Committee and widely distributed via BAPEN. This final version has been modified after comments from BAPEN council and members, and from the British Society of Paediatric Gastroenterology, Hepatology and Nutrition (BSPGHAN) and IF Working Group. It will be reviewed again in 2 years’ time.
BIFA Position Statement
Home Parenteral Nutrition

Unit
1. Patients receiving home parenteral nutrition (HPN) should be cared for by an Intestinal Failure (IF) or HPN designated unit with a multidisciplinary nutrition support team (NST).
2. HPN should only be practiced in centres where there is a governance framework (with Trust Board support) including finance in place for caring for these patients.
3. The unit should ideally have at least 20 adult patients (or 10 paediatric patients) receiving HPN of which more than 5 adults have been receiving it for more than 5 years. New units may develop under the guidance of an established unit meeting the criteria in this document.
4. All patients starting HPN or fluids must be registered via the Bluteq high cost drugs system and their Bluteq number should be added to all prescriptions and invoices (in NHS England only).
5. HPN units should work together to standardise practice.

Team
6. The adult team should consist of a consultant (with dedicated sessions), one (ideally 2 or more) nutrition nurse specialist, a specialist dietitian (with dedicated sessions) and a specialist pharmacist (with dedicated sessions). Children’s teams should have a paediatric gastroenterologist, paediatric trained nurses, a dietitian and a pharmacist.
7. There should be arrangements for cross cover with skilled staff of appropriate expertise when any member of the team is away.
8. Within the team there should be dedicated staff (nutrition nurse, dietitian, pharmacist or IF/HPN co-ordinator/administrator) to liaise directly with the home care companies.
9. There should be a member of the team or another specified expert available for 24 hours emergency specialist advice (with access to an intensive therapy or a high dependency unit when required).
10. There should be on site expertise in stoma care/tissue viability that includes managing a dehisced abdominal wound and when appropriate giving distal enteral feeding.

Practice
11. There must be a central venous catheter insertion service that can insert tunnelled central lines, peripherally inserted central catheters (PICCs) and implantable ports. It should be possible to achieve PN access within 24 hours for a short term central venous catheter and within 3 working days for a long term central venous catheter (except an implantable port).
12. There must be a clear pathway to admit a patient if an emergency (e.g. superior vena cava occlusion, catheter fracture or sepsis).
13. There must be written protocols with pathways, agreed by teams involved with central venous catheter care, for the management of complications (see 11 above). Information leaflets that outline procedures, complications and actions to take including contact telephone numbers/emails should be given to patients/carers.
14. Units must have agreed pathways and the expertise to repair or unblock a central venous catheter.
15. Staff (includes medical) must have access to a training program (includes both knowledge and skills) in complex nutritional care as relevant to IF/HPN patients.
16. Once a decision for HPN is made the patient/parents or carers should be assessed for their ability to perform PN procedures. This assessment will have to be repeated when situations change.
17. The team should have the specialist skills, when appropriate, to train a patient, carer, friend or family member to do all the procedures or facilitate training by a home care company.
18. When a patient is medically stable and a home care HPN company has an agreed contact, the patient should be able to go home within 5 working days of the formulation request (final script) submission to the home care company (as specified in the NHS National Framework for the Supply of HPN in England).
19. Patients with cancer and/or needing palliative care have their individual needs/circumstances considered quickly and are fast tracked home or to a hospice within 14 days of admission providing they are medically and psychologically stable.
20. Wherever possible scripts for adults should aim to have the least amounts of additions to licensed standard bags. This may reduce both the compounding time and costs.
21. HPN is only supplied by an accredited NHS HPN national framework supplier (in England). However, the Trust and the prescriber are still overall responsible for the patient’s care and they need to monitor the performance of the home care companies.
22. Patients should be reviewed in a dedicated outpatient department (or sometimes by telephone or video link) at least every 1-26 weeks. The frequency is determined by the patients’ underlying medical problems, their nutritional/fluid status and complexity of the regimen/medical treatments/psychosocial issues. The clinic appointment should be with the MDT (at least nutrition doctor, specialist nurse and dietitian).

**Relationships**

**Internal**

23. Good links and established pathways with an onsite interventional radiology service for patients with difficult venous access or for re-establishing patency if a central vein becomes occluded.
24. Good relationship with microbiology service and regular/when appropriate meetings to discuss patients (esp central venous catheter sepsis, discitis etc).
25. Have good relationships and links with an aseptic unit.
26. Have access to a psychological medicine department and for children/young adults to have access to children and adolescent mental health services (CAMHS) both of which should have experience in managing patients with intestinal failure.
27. Have easy (ideally on site) links to urology, gynaecology, vascular surgery, plastic surgery, upper and lower GI surgery, nephrology (haemodialysis unit), a hepato-biliary team, ophthalmology, dental care, occupational therapy, physiotherapy, social care and speech and language therapy (SALT). In addition for children and young people paediatric surgery, young people play/youth workers and paediatric physiotherapy.

**External**

28. Good links and established pathways with a centre doing intestinal failure surgery (e.g. enterocutaneous fistula repairs and re-establishing bowel continuity) and that can pre-assess patients considered for intestinal transplantation and make timely referrals.
29. Good relationship with commercial HPN service providers (as per NHS HPN national framework in England) and have regular meetings.

30. Services for young people should provide an adolescent and transition service with pathways and a dedicated transition process held with the adult provider including joint meetings between the patients, the parents or carers and NSTs from both the paediatric and adult centres.

31. A discharge planning service which engages with community services (e.g. community nurses and dietitians, health visitors, stoma care/tissue viability, pain team etc).

32. If a patient is not suitable for homecare or an approved homecare provider cannot be found there should be an escalation process to relevant NHS personnel.

Outcomes

33. The team must conduct regular audits of their outcomes. Units should be aspiring to an adult/young people inpatient catheter related sepsis rate of less than 3/1000 catheter days and an outpatient one of less than 1/1000 catheter days. Infants on parenteral nutrition have higher rates of catheter related sepsis.

34. Clear evidence of participation in relevant audits and clinical governance. These may include waiting times, length of stay, readmission rates, central vein thrombosis, central venous catheter occlusion, abnormal liver function tests and numbers weaned from HPN.

35. The views and quality of life of HPN patients, carers and families and the views of staff should be regularly audited.

36. Effective networking with nearby hospitals to where a patient may be admitted (e.g. with sepsis or CVC occlusion) and dissemination of shared care guidelines (e.g. catheter related sepsis or CVC occlusion) to the nearby hospitals.

37. All requested data about HPN patients must be reported to eBANS.
Comments not inco-operated

20. I have certainly seen patients where the need for extra fluid has led to a doubling up of "standard" bags so the patient gets extra calories as well as extra fluid and electrolytes, so is over fed! So I wonder whether a small comment about patients' needs for fluid and electrolytes not always matching their energy and nitrogen requirements would be helpful, just to try to preclude this.

35. How often audited and by whom. HCC or Trust?

34. Where should these be shared?

Use of term HPS for home parenteral support to include nutrition and fluid.